Cover photos: Upper left.—Mechanical vacuum extractor (Sampletek, Mavco Industries, Lincoln, Nebraska) used for chemical extractions and analyses, e.g., cation exchange capacity, NH₄OAc, pH 7. Upper right.—Shaw pipette used for particle-size analysis of the <2-mm fraction. Lower left.—Polarizing petrographic microscope used for optical analysis of a sand or silt fraction. Lower right.—Saran-coated natural and reconstituted clods on tension table prior to 33-kPa desorption.


Trade names are used in this manual solely for the purpose of providing specific information. Mention of a trade name does not constitute a guarantee of the product by USDA nor does it imply an endorsement by USDA.
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PREFACE

Laboratory data are critical to the understanding of the properties and genesis of a single pedon, as well as to the understanding of fundamental soil relationships based on many observations of a large number of soils. The development of an analytical database and the soil relationships based on those data are the cumulative effort of several generations of soil scientists.

The purpose of Soil Survey Investigations Report (SSIR) No. 42, Kellogg Soil Survey Laboratory Methods Manual, is to document methodology and to serve as a reference for the laboratory analyst. This document is expected to continue to change over time as new knowledge and technologies result in the development of new methods and the modification or retirement of old methods. This manual documents current methods and archives obsolete methods. It provides a historical perspective, documenting the contributions of many soil scientists who have gone before us. Many of these scientists are noted in the section on contributors.

Dr. Rebecca Burt, author of the Soil Survey Information Manual (SSIR No. 45, 2011), served as technical editor for all versions of SSIR No. 42 (1989, 1992, 1996, 2004, and 2014). Her contribution was significant in scope. She wrote some of the methods; was responsible for the review process, encompassing additions, corrections, and consistency of other methods; and provided leadership in assembling this document.
FOREWORD

The standard methods described in Soil Survey Investigations Report (SSIR) No. 42, “Kellogg Soil Survey Laboratory Methods Manual,” are those used by the Kellogg Soil Survey Laboratory (KSSL), National Soil Survey Center (NSSC). Included in SSIR No. 42 are descriptions of both current and obsolete methods, all of which are documented by method codes and linked with analytical results that are stored in the National Cooperative Soil Survey (NCSS) Soil Characterization Database. This linkage between laboratory method codes and the respective analytical results is reported on the KSSL data sheets.

The methods currently in use at the laboratory are described in enough detail that they can be performed at many laboratories without reference to other sources. An introduction to each group of related methods describes common characteristics. However, some repetition is included in each method description to make them complete in themselves and to minimize the need to reference to other parts of the manual.

Some analytical results in the NCSS Soil Characterization Database were obtained using procedures that are no longer used at the KSSL. Descriptions for these procedures are located in a section following the current methods. Information is not available to describe these procedures in the same detail as used to describe the current methods.

Since the publication of the SSIR No. 42 in 1996, the number and kinds of methods performed at the laboratory have increased significantly, resulting in a re-structuring of the method codes. The past and current method codes are hierarchical and alphanumerical. The code structure for the obsolete methods, however, allowed only a maximum of four characters, e.g., 6A2b. The new structure allows more characters, carrying more information about each method, e.g., particle-size and sample-weight basis for reporting data. This version of SSIR No. 42 includes not only the new method codes but also the older codes. This linkage between the two code systems is important to maintain because the older codes are reported in many older data sheets and scientific publications.

The KSSL data have historically been provided in printed reports, e.g., Primary and Supplementary Characterization Data Sheets, and later on electronic media, such as tapes and disks. More recently, other reports have been developed, e.g., Soil Taxonomy Characterization Data Sheets, and data are available from the NRCS Soils website http://soils.usda.gov/.

As in the past, this version of SSIR No. 42 assigns method codes and described methods only for those data reported on the Primary Characterization Data Sheets. With the exception of some KSSL primary analytical data that are included for user convenience, the Supplementary and Taxonomy Characterization Data Sheets show derived data, using analytical data as a basis for calculation, and do not carry method codes. Data on the Supplementary and Taxonomy Characterization Data Sheets are not described in this manual. For
more detailed information about the calculation and application of these derived values, refer to the other important source documents that are provided by the United States Department of Agriculture, Natural Resources Conservation Service (USDA–NRCS), and referenced in this manual.

The methods described herein identify the specific type of analytical or calculated data. Most of the methods are analytical in nature, i.e., they provide quantitative or semi-quantitative measurements, and include physical, chemical, mineralogical, and biological analyses. Sample collection and preparation in the field and the laboratory are also described. Historically, SSIR No. 42 has described some derived values, e.g., coefficient of linear extensibility (COLE) and water retention difference (WRD), and reported these values along with the analytical data on the Primary Characterization Data Sheets. This version of SSIR No. 42 follows this tradition. Since the publication of SSIR No. 42 in 1996, several more derived values have been added to the Primary Characterization Data Sheets. Although these data have been assigned method codes, detailed descriptions are not included in this version of SSIR No. 42. For more detailed information about the calculation and application of these derived values, refer to other important source documents that are provided by USDA–NRCS and referenced in this manual.

The purpose of this manual is to document methodology and to serve as a reference for the laboratory analyst. The standard methods described in this version (5.0) replace all earlier versions of SSIR No. 42 (1989, 1992, and 1996, respectively) and SSIR No. 1, “Procedures for Collecting Soil Samples and Methods of Analysis for Soil Survey” (1972, 1982, and 1984). All KSSL methods are performed with methodologies appropriate for their specific purpose. The KSSL standard operating procedures are standard methods, peer-recognized methods, KSSL-developed methods, and/or methods specified in “Keys to Soil Taxonomy” (2014). This manual also serves as the primary document from which a companion manual, Soil Survey Laboratory Information Manual (SSIR No. 45, 2011), was developed. SSIR No. 45 describes in greater detail the application of KSSL data.

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Research Soil Scientist
Soil Survey Research and Laboratory
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Lincoln, Nebraska
CONTRIBUTORS

Laboratory data are critical to the understanding of the properties and genesis of individual pedons as well as to the understanding of fundamental soil relationships based on many observations of a large number of soils. The development of laboratory methods, the analytical database, and the understanding of soil relationships based on those data are the cumulative effort of several generations of scientists. These efforts may be categorized as methods development, database design and development, and investigations of data relationships. Methods development at the Kellogg Soil Survey Laboratory results from a broad knowledge of soils, encompassing topical areas of pedology, geomorphology, micromorphology, physics, chemistry, mineralogy, biology, and field and laboratory sample collection and preparation. Listed below are many of the contributing scientists, some of whom have since retired from USDA–NRCS or are deceased. Some of the contributions are cited in this manual as scientific publications. The following list is not comprehensive for the developmental work that preceded the formation of the Kellogg Soil Survey Laboratory (KSSL) in Lincoln, Nebraska. In addition to the scientists, the current KSSL technicians are also listed. These analysts play a key role in performing the analyses and in developing new methods, modifying old methods, and ensuring data quality. Scientists (past and current) and physical science technicians (current) are listed alphabetically.

Scientists

Physical Science Technicians
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CURRENT METHODS OF ANALYSIS

SAMPLE COLLECTION AND PREPARATION (1)

Field Sample Collection and Preparation (1A)
Site Selection (1A1)
  Geomorphology (1A1a)
  Pedon (1A1b)
  Water (1A1c)
  Biological (1A1d)

1. Application

  General: The United States National Cooperative Soil Survey (NCSS) Program has prepared soil maps for much of the country. Both field and laboratory data are used to design map units and provide supporting information for scientific documentation and predictions of soil behavior. A soil map delineates areas occupied by different kinds of soil, each of which has a unique set of interrelated properties characteristic of the material from which it is formed, its environment, and its history (Soil Survey Division Staff, 1993). The soils mapped by the NCSS are identified by names that serve as references to a national system of soil taxonomy (Soil Survey Staff, 2014a). Coordination of mapping, sampling-site selection, and sample collection in this program contributes to the quality assurance process for laboratory characterization (Burt, 1996). Requisites to successful laboratory analysis of soils occur long before the sample is analyzed (Soil Conservation Service, 1984; Soil Survey Staff, 1996). In the field, these requisites include site selection, descriptions of site and soil pedon, and careful sample collection. A complete description of the sampling site not only provides a context for the various soil properties determined but it is also a useful tool in the evaluation and interpretation of the soil analytical results (Patterson, 1993). Documentation of the landscape, landform, and pedon at the sampling site serves as a link in a continuum of analytical data, sampled horizon, pedon, landscape, and overall soil survey area.

  The objectives of a project or study form the basis for designing the sampling strategy. A carefully designed sampling plan is required to provide reliable samples for the purpose of the sampling. The plan needs to address site selection, depth of sampling, type and number of samples, details of collection, and sampling and subsampling procedures to be followed. The Kellogg Soil Survey Laboratory (KSSL) primarily serves the NCSS, which is conducted jointly by USDA Natural Resources Conservation Service (NRCS), USDI Bureau of Land Management (BLM), USDA Forest Service, and representatives of U.S.
universities and agricultural experiment stations. In this context, the primary objective of KSSL sampling programs has been to select sites and pedons that are representative of a soil series or landscape segment and to collect samples that are representative of horizons within the pedon.

There are various kinds of sampling plans, e.g., intuitive and statistical, and many types of samples, e.g., representative, systematic, random, and composite. In the field, the KSSL has more routinely used intuitive sampling plans to obtain representative samples. An intuitive sampling plan is one that is based on the judgment of the sampler, wherein general knowledge of similar materials, past experience, and present information about the universe of concern, ranging from knowledge to guesses, are used (Taylor, 1988). A representative sample is one that is considered to be typical of the universe of concern and whose composition can be used to characterize the universe with respect to the parameter measured (Taylor, 1988).

In the laboratory, the primary objectives of sample collection and preparation are to homogenize and obtain a representative soil sample to be used in chemical, physical, and mineralogical analyses. The analyst and the reviewer of data assume that the sample is representative of the soil horizon being characterized. Concerted effort is made to keep analytical variability small. Precise laboratory work means that the principal variability in characterization data resides in sample variability, i.e., sampling is the precision-limiting variable. As a result, site-selection and sample collection and preparation are critical to successful soil analysis.

**Geomorphic Considerations:** Soils form a vital, complex continuum across the Earth’s landscape. The prime goal of the Soil Survey is to segregate the soil continuum into individual areas that have similar properties, and therefore, similar use and management. Soils cannot be fully understood or studied using a single observation scale. Instead, soil scientists use multiple scales to study and segregate soils and to transfer knowledge to soil users. To accomplish the task of Soil Survey at reasonable cost and time, soil scientists extend knowledge from point observations and descriptions to larger land areas.

Soil map-unit delineations are the individual landscape areas defined during and depicted in a soil survey. Soil observation, description, and classification occur at the pedon scale (1 to ≈7 m) and represent a small portion of any map unit (10’s to 1000’s hectares). Further, pedons selected, described, and sampled for laboratory analysis represent only a small subset of the observation points. Pedon descriptions and classifications along with measured lab data, however, accurately apply to a named soil map unit or landscape areas (soil component) within the map unit. Soil scientists can reliably project (“scale up”) pedon information to soil map units based on experience and the strong linkages among soils, landforms, sediment bodies, and geomorphic processes.

Thus, soil geomorphology serves several key functions in Soil Survey. These functions can be summarized as:
1. Provides a scientific basis for quantitatively understanding soil landscape relationships, stratigraphy, parent materials, and site history.

2. Provides a geologic and geographic context or framework that explains regional soil patterns.

3. Provides a conceptual basis for understanding and reliably predicting soil occurrence at the landscape scale.

4. Communicates effectively and succinctly soil location within a landscape.

During a Soil Survey, soil scientists achieve these functions both tacitly and by deliberate effort. Geomorphic functions are best explained by citing examples. The first function listed above involves planned, detailed soil landscape studies (e.g., Ruhe et al., 1967; Daniels et al., 1970; Gamble et al., 1970; Parsons et al., 1970; Gile et al., 1981; Lee et al., 2001, 2003a, 2003b), which are an important component of the Soil Survey. Such studies quantify and explain the links between soil patterns and stratigraphy, parent materials, landforms, surface age, landscape position, and hydrology. Studies of this nature provide the most rigorous, quantitative, and complete information about soil patterns and landscapes. The required time and effort are significant, but they are justified by the quantitative information and scientific understanding acquired. Soil survey updates by MLRA can and should involve similar studies.

The three other geomorphic functions listed are tacit and to a degree inherent in a soil survey. A number of earth science sources (Fenneman, 1931, 1938, 1946; Hunt, 1967; Wahrhaftig, 1965) identify and name geomorphic regions, which are grouped by geologic and landform similarity. The value of relating soil patterns to these regions is self-evident. Such terms as Basin and Range, Piedmont, Columbia Plateau, and Atlantic Coastal Plain provide both a geologic and geographic context for communicating regional soil and landform knowledge.

Soils occurrence can be accurately predicted and mapped using observable landscape features (e.g., landforms, vegetation, slope inflections, parent material, bedrock outcrops, stratigraphy, drainage, and photo tonal patterns). During a soil survey, soil scientists develop a tacit knowledge of soil occurrence generally based on landscape relationships. Soil occurrence is consistently linked to a number of geomorphic attributes. Among these are landform type, landscape position, parent material distribution, slope shape, slope gradient, and drainage pattern. This tacit soil landscape knowledge model is partially encapsulated in block diagrams and map unit and pedon descriptions. In turn, a clear, concise geomorphic description effectively conveys soil location within a landscape to other soil scientists and soil users. The Geomorphic Description Systems (GDS) is not discussed here. Refer to Wysocki et al., 2000; Schoeneberger et al., 2012; and Schoeneberger and Wysocki, 2012, for discussion of a comprehensive and consistent system for describing geomorphic and landscape attributes for soil

Geomorphology is an integral part of all soil survey processes and stages. Preliminary or initial knowledge about soil patterns is commonly based on landscape or geomorphic relationships. Observations during a soil survey refine existing landscape models or sometimes compel and create new models. Map unit design includes recognition and naming of landforms and also includes observations of landscape position, parent materials, and landscape and soil hydrology. Soil scientists capture this observational and expert knowledge through soil map unit and pedon descriptions, which should convey relationships among soil properties, soil horizons, landscapes, geomorphology, and parent materials.

Any study plan, site selection, or pedon sampling must also consider and address the geomorphology. Study or sampling objectives can vary. Every sampled pedon should include a complete description of both soil and geomorphology. In a characterization project, the sample pedons should be representative of the landscape unit (e.g., stream terrace, backslope) they occur upon. Note that the landscape unit that is sampled can be multi-scale. The unit could be a landform (e.g., stream terrace, dune, or drumlin), a geomorphic component (e.g., nose slope), a hillslope position (e.g., footslope), or all of these.

Sampled pedons represent both a taxonomic unit and a landscape unit. Both the landscape unit and taxonomic unit should be considered in site selection. Note that a single landscape unit (e.g., backslope) may contain more than one taxonomic units. A landscape unit is a easier to recognize in the field and to map than a soil taxonomic unit. For a characterization project, select the dominant taxonomic unit within a given landscape unit. The existence of other soils or taxa can and should be included in the soil description and the map unit description.

Soil patterns on landscapes follow catenary relationships. It is important to characterize not only the individual pedon properties but also the soil relationships both above and below on the landscape. This goal requires that soils be sampled as a catenary sequence (i.e., multiple samples across the same hillslope). This sampling scheme appears intensive, but it serves multiple purposes. A sample pedon or set of pedons provides vital characterization data and also can quantify the catenary pattern and processes. As such, it is an efficient use of sampling time and effort as well as laboratory resources. Moreover, it provides an understanding of the entire soil landscape.

Lastly, and perhaps most importantly, soil geomorphic relationships deserve—and sometimes demand—specific study during a soil survey. Crucial problems can be addressed by appropriately designed study of geomorphology, stratigraphy, or parent material. For example, a silty or sandy mantle over adjacent soils and/or landforms may be of eolian origin. A well-designed geomorphic study can test this hypothesis. In another geomorphic setting, soil distribution and hydrology may be controlled by stratigraphic relationships rather than by elevation or landscape patterns. A drill core or backhoe pit sequence can address this hypothesis. These
studies need not be elaborate, but they require forethought and planning. Such studies are applicable and necessary to the Major Land Resource Area (MLRA) approach to soil survey.

Pedon, Water, and Biological Sampling: The pedon is presented in Soil Taxonomy (Soil Survey Staff, 2014a) as a unit of sampling within a soil, i.e., the smallest body of one kind of soil large enough to represent the nature and arrangement of horizons and variability in the other properties that are preserved in samples (Soil Survey Division Staff, 1993). In the NCSS Program, laboratory pedon data combined with field data (e.g., transects and pedon descriptions) are used to define map unit components, establish ranges of component properties, establish or modify property ranges for soil series, and answer taxonomic and interpretive questions (Wilson et al., 1994).

Water samples are analyzed by the KSSL on a limited basis in support of specific research projects. These projects are typically in conjunction with soil investigations and have involved monitoring seasonal nutrient flux to evaluate movement of N and P via subsurface and overland flow from agricultural lands into waterways and wetlands.

Biological samples are also collected for analysis at the KSSL, either in conjunction with pedon sampling or for specific research projects. Measurable biological indices have been considered as a component to assess soil quality (Gregorich et al., 1997; Pankhurst et al., 1997). A large number of soil biological properties have been evaluated for their potential use as indicators of soil quality/health (Doran and Parkin, 1994; Pankhurst et al., 1995). The USDA–NRCS has utilized soil biology and carbon data in macro-nutrient cycling, soil quality determinations, resource assessments, global climate change predictions, long-term soil fertility assessments, impact analysis for erosion effects, conservation management practices, and carbon sequestration (Franks et al., 2001). Soil Quality was identified as an emphasis area of the USDA–NRCS in 1993. All soil quality publications and technical notes are available online at http://soils.usda.gov/.

2. Summary of Method

A site is selected that meets the objectives of the laboratory sampling. The site and the soil pedon are described and georeferenced, using such instruments as wide area augmentation system, global positioning system (WAAS GPS). A complete soil and geomorphic description are completed. The soil descriptions include observations of specific soil properties, such as texture, color, slope, and depth. Descriptions may also include inferences of soil quality (soil erodibility and productivity) as well as soil-forming factors (climate, topography, vegetation, and geologic material). The sampled pedons should be representative of the landscape unit and can be multi-scale (fig. 1A-1).

A soil pit is often excavated with a back-hoe (fig. 1A-2). Depth and breadth of pit depend on the soil material and the objectives of sampling. Soil horizons or
Figure 1A-1.—Landscape of selected site for sampling.

Figure 1A-2.—Excavated pit for pedon sampling.
zones of uniform morphological characteristics are identified for sampling (fig. 1A-3). Photographs are typically taken of the landform or landform segment and the soil profile. Photographs of the soil profile include photo tapes showing vertical scale. The photographs are taken after the layers have been identified (fig. 1A-4) but before the extraction of the vertical section by the sampling process (fig. 1A-5).

The variable nature or special problems of the soil itself, e.g., Vertisols, Histosols, or permafrost-affected soils, may require the use of specific excavation and sampling techniques. For example, the shear failure that forms slickensides in Vertisols also disrupts the soil to the point that conventional soil horizons do not adequately describe the morphology.

Representative samples are collected and mixed for chemical, physical, and mineralogical analyses. A representative sample is collected using the boundaries of the horizon to define the vertical limits and the observed short-range variability to define the lateral limits. The tag on the sample bag is labeled to identify the site, pedon, and soil horizon for the sample.

In the field, the 20- to 75-mm fraction is generally sieved, weighed, and discarded. In the laboratory, the <20-mm fraction is sieved and weighed. The KSSL estimates weight percentages of the >2-mm fractions from volume estimates of the >20-mm fractions and weight determinations of the <20-mm fractions.
Figure 1A-4.—Photographs are typically taken of a soil profile after the layers have been identified but before the vertical section by the sampling process. Note scale in metric units.

Undisturbed clods are collected for bulk density and micromorphological analysis. Clods are obtained in the same part of the pit as the mixed, representative sample. Bulk density clods are used for water retention data; to convert from a weight basis to a volume basis; to determine the coefficient of linear extensibility (COLE); to estimate saturated hydraulic conductivity; and to identify compacted horizons. Microscope slides prepared from other clods are used for micromorphology to identify fabric types, skeleton grains, weathering intensity, illuviation of argillans, and to investigate genesis of soil or pedological features.

Water samples may also be collected for laboratory analyses at the same time as pedon sampling. Choice of water sampling site depends not only on the purpose of the investigation but also on local conditions, depth, and the frequency of sampling (Velthorst, 1996). Specific recommendations are not applicable, as the details of collection can vary with local conditions. Nevertheless, the primary objective of water sampling is the same as with soil and biological sampling, i.e., to obtain a representative sample for laboratory analyses. Water samples require
Biological samples may also be collected for analysis at the laboratory, either in conjunction with pedon sampling or for specific research projects. As with pedon sampling, sampling for root biomass includes selecting a representative site, sampling by horizon, and designating and sampling a sub-horizon if root mass and morphology change. The same bulk sample collected for soil mineralogical, physical, and chemical analyses during pedon sampling can also be used for some soil biological analyses. Alternatively, a separate bio-bulk sample can be collected in the field. Surface litter and O horizons are sampled separately, as with pedon sampling. If certain biological analyses, e.g., microbial biomass, are requested, these samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory to avoid changes in the microbial communities.

3. Interferences

In the process of sampling, a number of obstacles may arise from external sources, e.g., weather, accessibility, steep terrain, wet terrain, insects, and large rock fragments. Sometimes pits have to be hand-excavated. Common sense
and the guidelines for obtaining representative samples are applied to the extent possible.

Preservation of sample integrity, i.e., avoiding changes or contamination, during sampling and transport is important. Sampling for trace element analysis requires the use of clean, non-metallic equipment. Extreme care and precision are required for samples with low natural elemental concentrations.

Do not allow soils to dry; some soils irreversibly harden upon drying, affecting some laboratory analyses, such as particle size (Kubota, 1972; Espinoza et al., 1975; and Nanzyo et al., 1993). High temperatures can also alter microbial populations and activity (Wollum, 1994).

Avoid contamination of water samples; do not touch the inner part of the sample container, screw cap, or sample water. Gloves (powderless) may be used. Water samples are affected by microbial activity, resulting in a change in the concentration of some elements (e.g., nitrate, phosphate, and ammonium); the reduction of sulfate to sulfide and chlorine to chloride; and the loss of iron through precipitation or oxidation (Velthorst, 1996). The addition of microbial inhibitors may be necessary.

In general, for most biological samples, plastic bags will suffice because they are generally permeable to CO$_2$ and O$_2$, preventing sample drying, i.e., aerobic samples will remain aerobic during transport to the laboratory (Wollum, 1994). The KSSL recommends double-bagging zip-lock plastic bags to prevent loss of water from biological samples.

The kind of water-sample container (adsorption, desorption) as well as the bottle volume can affect the analytical results. For example, polyethylene bottles increase the chlorine content with time or adsorb organic material; errors increase with the permeability of bottle wall; glass bottles release sodium and silicon with time; and small sample volume has more contact with larger bottles compared to small bottles (Velthorst, 1996). Water-sample containers should be acid washed and capped in the laboratory prior to collection in the field. The drying of these containers should also be considered with regards to interferences or contaminants. Ceramic cups for collection of soil and water may require an acid pretreatment prior to installation in the field, as these cups have a small cation exchange capacity, sorbing dissolved organic carbon and releasing aluminum and silica (Velthorst, 1996). Refer to the respective manufacturer’s manual, e.g., Soil Moisture Corporation, for the appropriate treatment of these cups.

Avoid long periods between collection and laboratory analysis of water and some types of biological (e.g., microbial biomass) and soil samples (e.g., sulfidic materials). To prevent significant changes (e.g., degradation, volatilization, alteration in microbial community), these samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory. Do not allow water samples to freeze, which can influence pH and the separation of dissolved organic matter from the water phase.
4. Safety

Several hazards can be encountered in the field during sample collection. Examples include sharp-edged excavation tools, snake bites, and falls. To meet U.S. Department of Labor Occupational Safety and Health Administration (OSHA) standards, which are available online at http://www.osha.gov/, sampling pits deeper than 125 cm (5 feet) need to be shored or have one side that is opened and sloped upward to prevent entrapment. Take precautions when operating or in the proximity of machinery, e.g., backhoe, drill rig, or hydraulic probe, and when lifting sample bags. Acetone is highly flammable. Avoid open flames and sparks. Acetone should be used downwind from a site to keep fumes from collecting in the bottom of the pit. Use extra care when storing and transporting acetone. Refer to the Material Safety Data Sheets (MSDS) for information on the chemical make-up, use, storage, emergency procedures, and potential health effects of the hazardous materials associated with this method.

5. Equipment

5.1 Plastic bags, for mixed soil samples
5.2 Zip-lock plastic freezer bags, for biological samples
5.3 Tags, for bagged samples
5.4 Plastic bags, for bulk density and thin section clods
5.5 Aluminum case, for shipping clod boxes
5.6 Shipping bags (canvas, leather, or burlap) for mixed samples
5.7 Clod boxes, cardboard with dividers
5.8 Core boxes, to transport cores from drill rig or hydraulic probe
5.9 Stapler, with staples
5.10 Hair nets
5.11 Rope
5.12 Clothespins
5.13 Felt markers, permanent
5.14 Sampling pans
5.15 Sampling knives
5.16 Chisel
5.17 Rock hammer
5.18 Nails
5.19 Measuring tape
5.20 Photo tape
5.21 Sieves (3-inch and 20-mm)
5.22 Plastic sheets
5.23 Canvas tarp
5.24 Camera
5.25 Frame, 50 cm x 50 cm
5.26 Garden clippers
5.27 Pruning shears
5.28 Bucket
5.29 Scale, 100-lb capacity, for rock fragments
5.30 Electronic balance, ±0.01 g sensitivity, for weighing roots and plant residue
5.31 Cooler, with ice or gel packs, for biological samples
5.32 Containers, with screw caps, acid-washed, for water samples
5.33 Gloves, plastic, powderless
5.34 Bulk density equipment, if natural clods are not appropriate technique, e.g., bulk density frame or ring excavations, compliant cavity, and cores
5.35 First-aid kit
5.36 Dust mask
5.37 Hardhat
5.38 Hand lens

6. Reagents
6.1 Acetone
6.2 Water, in spray bottle
6.3 Dow Saran F-310 resin, available from Dow Chemical Company
6.4 1 N HCl
6.5 Material Safety Data Sheets (MSDS)

7. Procedures

**Project Categories**

The number and types of samples collected from a site are governed in part by the objectives of the information needed. At the KSSL, the sampling and analysis requests are used as a basis for categorizing projects, with reference and characterization projects being the most common.

**Reference Projects:** These projects are designed to answer specific questions on mapping or soil classification, provide data for transect of a mapping unit, or collect calibration standards. Samples are typically collected from specific horizons in three to five locations, which either relate to the sampling question or are representative of the map unit. Typically, a limited number of analyses, specific to the questions asked, are performed on these samples.

If a transect is used to test map unit composition, an appropriate sample from each transect point may be collected for analyses that are critical to distinguishing
between map unit components. Also, samples may be collected as standards for the survey project for texture, for organic carbon, or for calibration of field office analyses, such as base saturation.

**Characterization Projects:** These projects are designed to obtain comprehensive soil characterization data for a representative pedon of a map unit or a pedon that is included in a research study. Samples collected from each horizon include bulk samples of approximately 3 kg, as well as clods of natural fabric for bulk density and micromorphology. A standard suite of laboratory analyses are performed on each horizon. In addition, specific analyses, such as mineralogy or andic properties, may be requested to provide more complete information on the specific pedon sampled.

**Geomorphology and Stratigraphy Projects:** These research projects are designed to study relationships between soils, landforms, and/or the stratigraphy of parent materials. For example, a specific project may be designed to study the relationships between a catena of soils, their morphological properties, e.g., redoximorphic features, and the hydrology of the area. Another study may be designed to determine the lateral extent of stratigraphic breaks. Site or pedon selection is governed by the objectives of the study but often is intended to represent typical segments of the landform. Sampling and analytical requests may be similar to the scheme used in a characterization or reference project. Often, core samples may be collected to several meters in depth through the use of a hydraulic probe.

### Pedon Sampling Equipment

**Excavated Pits:** A pit may be excavated by hand or with a backhoe. Hand-digging may be necessary due to specific characteristics of the site location, type of soil material, or availability of a backhoe. Pedons are generally excavated either through the solum and into the parent material or to a maximum depth of 2 meters. If a backhoe is used, the pit should be dug in the form of an arc with a minimum working face deeper than about 150 cm (5 ft). Slope the pit upward toward the backhoe for an escape route. The pit can also be modified from the back side to form a “T” with the back of the trench opened and widened as an escape route. If this is not practical and the pit is deeper than 125 cm (5 ft), shoring is required to meet OSHA standards.

The sampling procedure is the same for hand-dug and backhoe pits. Mark horizons or zones to be sampled. Take a representative sample from boundary to boundary of a horizon. The lateral extent should include the observed short-range variability. Unless the soil exhibits little short range variability, the best procedure is to place 4 to 5 kg soil on the plastic sheet or canvas tarp, mix thoroughly by rolling action, and place a representative subsample, minimum of 3 kg (3 qt), in a plastic sample bag. Label a tag with soil name, soil survey number, horizon (zone), and depth (as a minimum). Double fold the top of the plastic bag (forward and reverse) and staple the top of the tag under the folds. The depth of sampling
may be extended by bucket auger or hydraulic probe as appropriate to meet the
objectives of the project. If the soil has rock fragments in one or more horizons,
the soil and coarse fragments need to be sieved and weighed as described below.

Collect three bulk density clods from each horizon. Two clods are used in the
primary analysis. The third clod is reserved for a rerun, if needed. Clods should
be roughly fist sized and should fit into the cell (8 x 6 x 6 cm) of a clod box fairly
snugly. Take the clods from the same vicinity of the pit as the mixed sample.
Carve out a working section in the pit wall to remove an undisturbed block. Break
the block into fist sized pieces and pare into an ovoid (egg-shaped) clod. Place
the clod in a hair net. Place a staple on top of clod to note orientation. If the
clod is dry, mist the clod with water just until the surface glistens. The water
inhibits saran penetration of the clod. Dip the clod once, briefly, in saran mix to
coat the clod, and hang it from a rope with a clothes pin to dry. Clods can be
dipped and then hung or can be hung and then dipped by raising the container
up to immerse the clod, briefly. To prevent acetone evaporation, keep the saran
container covered except when dipping clods. Coat the clod only once in the field.
Additional coats are applied in the laboratory. When the clod is dry (bottom is not
sticky to the touch), place the clod in a plastic bag and then into a cell of a clod
box. Label the appropriate cell on the inside of the lid of the box to identify the
soil survey number and horizon (zone) for the clod. Clod boxes are designed to
identify sequences of three clods per horizon.

Collect two clods from each horizon for preparation of thin sections and
micromorphological examination. Place a staple in the top of each clod to denote
orientation. Clods should be roughly fist sized but otherwise unmodified. If the
soil fabric is fragile, the clod can be placed in a hairnet and dipped briefly in saran
as described above. Place the clod in a plastic bag and then into a cell of a clod
box. Make special note of any features to be studied by thin section. Label the
appropriate cell on the inside of the lid of the box to identify the soil survey number
and horizon (zone) for the clod.

If the material is too sandy and/or too dry to hold together in a clod, bulk
density samples can be collected with an aluminum can or other small can of
known volume. Sampling is easier if the bottom of the can has a small hole that
allows air to escape as the can is inserted. Smooth a planar area in the pit face;
or, if sampling from the top down, smooth a planar horizontal area. In either case,
choose an area that appears representative of the horizon. With the palm of your
hand, gently push the can into the smoothed area until the bottom of the can is
flush with the wall. If resistance stops you before the bottom of the can is flush
with the pit wall, lay a board across the bottom of the can and tap lightly with a
hammer or geology pick until the can is flush. Dig out the sampling can plus extra
sample and, with a knife blade, smooth off the sample flush with the top of the
can. Empty the contents of the can into a plastic bag, tie the top of the bag in a
single knot, and put the bag into a cell in a clod box. Label the appropriate cell
on the inside of the lid of the box to identify the soil survey number and horizon
(zone) for the sample. Collect two samples per horizon. Indicate the volume of the sampling can in the sampling notes. It is assumed that there is no volume change with water content in sandy soils. Therefore, one density is representative for all water contents of coarse-textured soils.

Do not leave empty cells in a clod box. Fill empty cells with a wadded paper. This prevents clods from shifting in transit. Tape down the top of a filled clod box with nylon filament tape (one short piece on each end and two short pieces in front). Label the top of the box to identify type of sample (bulk density or thin section) and appropriate soil survey numbers and horizons (zones) for the samples. Place six clod-boxes in an aluminum case for shipment. Single clod-boxes also ship well.

**Hand Probe:** Remove surface if it is not suitable for coring. Remove core sections, and lay them in order on plastic sheet. Measure core length against depth in hole to determine if the core has been compressed. Mark horizon breaks on the plastic. Mix the horizon or zone to be sampled. Place sample in a plastic bag and label with soil survey number, horizon (zone), and depth for the core. Samples need to be a minimum of 500 g (1 pt) and are generally suitable for only a limited number of analyses.

**Hydraulic Probe:** Remove surface if it is not suitable for coring. Remove core sections, and then lay in order on plastic sheet. With a sharp knife, trim the exterior to remove any oil and contaminating soil material. Split one core open to mark horizons, describe, and then sample. Measure core length against depth in hole to determine if the core has been compressed. Mark horizon breaks on the plastic. Mix the horizon or zone to be sampled. Place sample in a plastic bag and label with soil survey number, horizon (zone), and depth for the core. Obtain a minimum of 500 g (1 pt) for a reference sample or 3 kg (3 qt) for a characterization sample.

If the core has not been compressed and the core diameter is 3 inches or more, samples for bulk density can be taken from a second core. Mark an 8-cm long segment on an undisturbed section and slice a cylindrical segment. Measurements of core diameter and length can be used to calculate volume and density at the field-state water content. Core segments can be placed in a hair net, dipped once briefly in saran mix to coat the clod, hung from a rope with a clothes pin to dry, placed in a plastic bag, and then put into a cell of a clod box.

**Rotary Drill (Hollow Stem):** Remove drill core sections and lay in order on plastic sheet. Measure core length against depth in hole to determine if the core has been compressed. Mark horizon breaks on the plastic. Mix the horizon or zone to be sampled. Place sample in a plastic bag and label with soil survey number, horizon (zone), and depth for the core. Obtain a minimum of 500 g (1 pt) for a reference sample or 3 kg (3 qt) for a characterization sample.

If the core has not been compressed and the core diameter is 3 inches or more, samples for bulk density can be taken from the core. Mark an 8-cm long segment on an undisturbed section and slice a cylindrical segment. Note the core
diameter and length in the soil description. Place the core segment in a plastic bag, and place bag into a bulk density (clod) box for shipment. Measurements of core diameter and length can be used to calculate volume and density at the field-state water content. Core segments can be placed in a hair net, dipped once briefly in saran mix to coat the clod, hung from a rope with a clothes pin to dry, placed in a plastic bag, and then put into a cell of a clod box. Label the appropriate cell number on the inside of the box lid to identify the site, pedon, and horizon.

A core segment can be taken for thin section. Place a staple in the top of the core, place the core in a plastic bag and put the bag into a cell in a clod box. Label the appropriate cell number on the inside of the box lid to identify the site, pedon, and horizon.

**Bucket Auger:** Remove surface if it is not suitable for auguring. Remove auger loads and lay in order on plastic sheet. When horizon breaks are detected, measure depth in hole and mark it on the plastic. Mix the horizon or zone to be sampled. Place sample in a plastic bag and label with soil survey number, horizon (zone), and depth for the sample. Obtain a minimum of 500 g (1 pt) for a reference sample or 3 kg (3 qt) for a characterization sample. Sampling depth in a pit can be extended by the use of an auger in the pit bottom.

**Pedon Sampling Types**

**Soils with Rock Fragments:** If coarse fragments up to 75 mm (3 in) in diameter are to be weighed in the field, weigh excavated sample in a bucket of known weight (tare). Sieve the sample through both a 75-mm and 20-mm sieve (¾ in) onto a canvas tarp that can be suspended from a scale. Estimate the coarse fragment volume percent of both the 75- to 250-mm (10 in) fraction and >250-mm fraction, and record these values in the description or sampling notes. Weigh the 20- to 75-mm and the <20-mm fractions in pounds or kilograms, and record these weights. Weights are calculated to an oven-dry base in the laboratory. Place a minimum of 4 kg (1 gal) in a plastic bag, double fold the bag, and staple. The water content is determined on the sample in the laboratory. If the 20- to 75-mm fraction is not weighed in the field, estimate the volume percent and record in the sampling notes or description. Refer to method 3A2 of this manual on the analysis of particles >2 mm.

**Organic Soils:** If the soils are drained or the natural water table is below the surface, obtain samples of upper layers from a pit. If the hydraulic conductivity is slow enough, dig and remove samples below the water table as far as practical with due haste and place on a plastic sheet in an orderly fashion for describing and processing. If undisturbed blocks can be removed for bulk density, carve out cubes of known dimension (e.g., 5 cm on a side), place the block in a plastic bag and tie the top in a knot. Place in a second plastic bag if soil is saturated, tie the top in a knot. Put the double bagged sample in a clod box and label the
appropriate cell on the inside of the lid to identify the soil survey number and horizon (zone) for the sample. Note the sample dimensions in the sampling notes.

Collect samples from below the water table with a Macaulay peat sampler. If the samples appear undisturbed, mark 10-cm segments, slice with a knife, and place a single segment in a plastic bag. Tie the top in a knot, place in a second plastic bag, and tie the top of that bag in a knot. Put the double-bagged sample in a clod box and label the appropriate cell on the inside of the lid to identify the soil survey number and horizon (zone) for the sample. Note the sampler diameter and length of core in sampling notes. The sample shape is a half-cylinder. As an alternative, carve a block to fit snugly in a tared water can. Place lid on can, put can in a plastic bag, tie the top, and put the bag in a clod box. Identify the can number, depth, and tare weight in sampling notes. Take replicate samples for the mixed sample, as necessary.

Larger samples can be taken below the water table by removing the surface mat with a spade and sampling lower layers with a post-hole digger. Place samples of each layer on plastic for examination. Transfer samples to small plastic bags and knead to remove air. Put two small bags of sample into one large plastic bag, fold top, staple, and tag.

**Sulfidic Soil Materials:** These materials, as defined in the “Keys to Soil Taxonomy” (Soil Survey Staff, 2014a), commonly occur in intra-tidal zones adjacent to oceans and are saturated most or all of the time. Use containers with an airtight cover. Mason jars and plastic containers with a positive sealing mechanism work well. Glass containers must be adequately packed for shipment to prevent breakage. Fill the container nearly full of sample and add the ambient mixture of soil and water so that all air is eliminated when the lid is secured. Keep containers in the dark and cool. Sulfidic soil samples require expedited transport in a cooler and are refrigerated (4 °C) immediately upon arrival at the laboratory. Once the container is in the lab, if it appears air remained in the container, nitrogen gas can be bubbled through the sample for a few minutes to displace air, then the lid can be replaced. The intent is to keep the material at the field pH prior to running the (incubation) oxidized pH test and other analyses whose results may change upon oxidation.

**Permafrost Affected Soils:** Soils that have permafrost present two special sampling problems. The permafrost is very resistant to excavation, and the cryoturbation disrupts horizon morphology. In many cases, the surface layers are organic materials. The following sampling approach is suggested.

Test the depth to the frost table with a small (1 to 2 mm) diameter steel rod. Excavate a small pit (about 0.7 by 1.3 m) to leave about 10 cm of unfrozen material over the permafrost. If a cyclic pattern (up to a few meters) is evident in the surface topography, extend the pit through at least one cycle to the depth of sampling. The organic layers can be carved out with a sharp knife or shovel in many cases and removed. Save the large chunks, if possible.
The objective is to record the morphology of the unfrozen soil before the permafrost is disturbed. Examine the surface and designate horizons. If the soil is disrupted to the extent that lateral horizons do not represent the morphology, impose a grid over the pit face and sketch the morphology on graph paper. Describe the soil down to the frost table. When the description of the unfrozen material is complete, remove all unfrozen material to examine the conformation of the frost table. Note on graph paper if necessary and photograph.

Frozen earth can be removed in successive steps with a gasoline-powered jackhammer. Place pieces from each step on a separate plastic sheet. Examine pieces and describe the morphology as they are removed. Note thickness of segregated ice lenses and make a visual estimate of relative volume of segregated ice. Place representative pieces into a water-tight container so that the sample can be weighed, dried, and weighed again to calculate the amount of water and volume of ice. Excavate to a depth of 30 to 50 cm below the frost table, if practical. Clean off the pit face and be ready to photograph immediately. Sample each horizon or zone for mixed sample, bulk density, and thin section as is practical.

Vertisols: The shear failure that forms slickensides in Vertisols also disrupts the soil to the point that conventional horizons do not adequately describe the morphology. A gilgai surface topography is reflected in the subsurface by bowl-shaped lows and highs. One convention is to sample pedons out of the low and the high areas, which represent extremes in the cyclic morphology.

In order to examine morphology and associated soil properties in more spatial detail, the following procedure is suggested. Dig a trench long enough to cover two or three cycles of morphological expression. From the bottom of the pit, remove soil from the nonwork face so it slopes up and away. Use nails and string to outline boundaries of morphological cells. Assign a number and a horizon designation to each cell.

Construct a level line about 1 meter below the highest point on the surface. Hammer a spike into the wall at one end of the pit. Tie a loop in string, place the loop over the spike, and run the string to the far end of the pit. Place a line level on the string, tie another loop in the string, place a second spike through the loop, pull the string taut, raise or lower the spike until the string is level, and hammer the spike into the pit face.

Place a marker at each meter along the string from one end to the other. Transfer the morphology outlined by the string to graph paper by measuring the x-coordinate along the string and the y-coordinate above or below the string, both in centimeters. Use a level or a plumb bob to make the y measurement vertical.

Sample each cell for characterization analysis as described above. The sampling scheme can include traditional pedon sequences by sampling vertical sequences of cells at low, high, and intermediate positions along the cycle.

Subaqueous Soils: Sampling of subaqueous soils is conducted during both winter and summer months. These soils are typically sampled to an average
depth of 100 to 150 cm, usually in water \(<2.5\) m deep. Soils are sampled using the standard bucket auger, Macaulay peat sampler, or vibracorer. Soils are described and classified using USDA–NRCS soil survey methods (Soil Survey Staff, 2014a; Schoeneberger et al. 2012). Common measurements in a subaqueous soil survey include but are not limited to bathymetry, water-quality measurements (e.g., pH, dissolved oxygen, salinity water temperature), and soil-quality measurements (e.g., reaction to \(H_2O_2\), reaction by oxidized pH, fluidity, electrical conductivity, bulk density satiated). For more information on water column measurements, soil profile descriptions, and soil profile measurements, refer to Schoeneberger et al. (2012). For additional information on subaqueous soils, refer to Demas and Rabenhorst (1999), Bradley and Stolt (2003), and Erich et al. (2010).

**Pedon Sampling Schemes**

**Horizon, Incremental, and Fixed-depth Sampling:** Horizon sampling has been the most common sampling scheme in soil survey. Other sampling schemes include incremental and fixed-depth sampling. Incremental sampling may be used when project objectives (e.g., soil genesis or archeological) require within-horizon detail (Schoeneberger et al., 2012). Property variation or trends within horizons require samples at specified increments (e.g., 10-cm depths). Incremental samples should be taken within horizons, without crossing horizon boundaries. Incremental sampling provides more detail than horizon sampling but adds time and expense. It is generally limited to special projects (Schoeneberger et al., 2012). Fixed-depth sampling may be used when specified objectives (e.g., surface compaction studies) address properties by fixed depths (e.g., 0 to 5 cm or 5 to 10 cm) instead of by horizons (Schoeneberger et al., 2012). This approach is appropriate for certain purposes but precludes data comparison by horizon. Data collected by depth are comparable within a study and to other studies employing the same depths. Fixed-depth samples may cross horizons that contain contrasting materials (e.g., sandy over clayey strata). Resulting data represent neither horizon and are difficult to interpret. Caution is advised when this approach is used (Schoeneberger et al., 2012).

**Paired Pedons:** In the early 1950s, field and laboratory soil scientists of the Soil Conservation Service began sampling “paired pedons” selected from the middle of the range of a single phase of a series (Mausbach et al., 1980). Paired pedons were morphologically matched as closely as possible through field observations given practical restrictions of time, size of area, access to site, and inherent variability of the parent material, with variability within these pairs representing variability within a narrow conceptual range (Mausbach et al., 1980). Evaluation of vertical distribution of properties of important horizons has been performed in soil survey by sampling one complete pedon plus satellite samples of these horizons. The efficient assessment of a single horizon requires that the horizon be sampled in several pedons. Sampling of paired pedons was
considered a good first-approach technique to study soils in an area. Important early literature on soil variability includes Robinson and Lloyd (1915), Davis (1936), and Harradine (1949). After series concepts narrowed, variability studies of properties and composition of mapping units included Powell and Springer (1965), and Wilding et al. (1965), McCormack and Wilding (1969), Beckett and Webster (1971), Nielsen et al. (1973), Crosson and Protz (1974), Amos and Whiteside (1975), and Bascomb and Jarvis (1976). Studies of the variability of properties within a series include Nelson and McCracken (1962), Andrew and Stearns (1963), Wilding et al. (1964), Ike and Clutter (1968), and Lee et al. (1975). Refer to the Soil Survey Investigations Report No. 51, “Soil Survey Field and Laboratory Methods Manual” (Soil Survey Staff, 2014a), for more detailed discussion of sampling strategies.

Biological Sampling

Biological samples can also be collected for laboratory analysis, either in conjunction with pedon sampling or for specific research projects. At the time of sampling for above-ground biomass, the plants should be identified either in the field or later using a plant identification key so as to determine which plants are associated with the soil microbial communities. Typically, a 50- x 50-cm area is sampled. All vegetation is clipped to the soil surface and separated by genus or species and by live and dead fractions. Each plant fraction is weighed, dried, and reweighed to determine above-ground biomass. As with pedon sampling, sampling for root biomass includes selecting a representative site, sampling by horizon, and designating and sampling a subhorizon if root mass and morphology change. The sampling area is approximately 1 m². These samples are weighed, dried, and reweighed to determine root biomass. Typically, the roots are separated by hand sieving at the laboratory. The same bulk sample collected for soil mineralogical, physical, and chemical analyses during pedon sampling can also be used for some soil biological analyses, e.g., particulate organic matter (POM) and total N, C, and S. Alternatively, a separate bio-bulk sample can be collected in the field. As with pedon sampling, surface litter and O horizons are sampled separately for bulk density determinations by cutting out a 50- x 50-cm area in a square to a measured depth. Include replicate samples in the sampling plan, the primary purpose of the replicate samples is to identify and/or quantify the variability in all or part of the sampling and analysis system. Properly label samples to show important information, e.g., soil, depth, and horizon. If certain biological analyses, e.g., microbial biomass, are requested, these samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory to avoid changes in the microbial communities.

USDA–NRCS field procedures and sampling protocols for samples that do not require analysis at the KSSL are not covered in this manual. Refer to http://soils.usda.gov/ or contact State land-grant institutions and soil survey offices for more detailed discussion of these topics.
Water Sampling

Water samples can be collected for laboratory analyses, either in conjunction with pedon sampling or for specific research projects. The amount and composition of water samples vary strongly with small changes in location. Choice of water-sampling site depends not only on the purpose of the investigation but also on local conditions, depth, and frequency of sampling (Velthorst, 1996). Specific recommendations are not applicable, as the details of collection can vary with local conditions. Nevertheless, the primary objective of water sampling is the same as that of soil sampling, i.e., to obtain a representative sample for use in laboratory analyses. USDA–NRCS projects requiring collection of water samples have typically been conducted in conjunction with special soil investigations. For more detailed discussion of sampling protocols and investigations of water quality, refer to the U.S. Geological Survey field manual, available online at http://pubs.water.usgs.gov/. Detailed information about the elements of a water-quality monitoring and assessment program are available at the U.S. Environmental Protection Agency’s website http://www.epa.gov/.

Preserve samples in the field-state until analysis at the laboratory. Prevent the introduction of change or contamination. Before collecting the water samples in the field, rinse the containers several times with the sample water. Completely fill the container and screw cap with the sample water. Avoid touching the sample water, the inner part of the container, or the screw cap. Gloves (powderless) may be used. Include blank samples in the sampling plan. The primary purpose of blank samples is to identify potential sources of sample contamination and assess the magnitude of contamination with respect to concentration of target analytes. There are many possible types of blanks (e.g., source-solution, equipment, trip, ambient, and field blanks). Include replicate samples in the sampling plan, the primary purpose of the replicate samples is to identify and/or quantify the variability in all or part of the sampling and analysis system. Common types of replicate samples include concurrent, sequential, and split. Refer to Wilde et al. (1993) for more detailed descriptions of the purpose and processing procedures for blanks and replicate samples. Properly label sample containers to show important information, e.g., location, depth, and time. Water samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory.

Some water analyses, e.g., electrical conductivity, total C, and inorganic C, need to be performed promptly because optimal preservation is not possible (Velthorst, 1996). Upon completion of these analyses, sample filtration (0.45-µm membrane) is used to separate dissolved from suspended material. The sample is then split into two subsamples, with one acidified to pH 2 for cation analyses (e.g., Al, Fe, Mn) and the other for anion analyses. These other water analyses also need to be performed as promptly as possible.
8. References


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**Laboratory Sample Collection and Preparation (1B)**

**Soil (1B1)**

**Samples (1B1a)**

The purpose of any soil sample is to obtain information about a particular soil and its characteristics. Sampling provides a means to estimate the parameters of these soil characteristics with an acceptable accuracy at the lowest possible cost (Petersen and Calvin, 1986). Subsampling also may be used, as it permits the estimation of some characteristics of the larger sampling unit without the
necessity of measurement of the entire unit. Subsampling reduces the cost of the investigation, but usually decreases the precision with which the soil characteristics are estimated. Efficient use of subsampling depends on a balance between cost and precision (Petersen and Calvin, 1986).

Soil variability and sample size are interferences to sample collection and preparation. The objective of laboratory preparation is to homogenize the soil samples used in chemical, physical, and mineralogical analyses. At each stage of sampling, an additional component of variability, the variability within the larger units, is added to the sampling error (Petersen and Calvin, 1986). Soil material needs to be adequate in amount and thoroughly mixed in order to obtain a representative sample.

KSSL receives bulk soil samples from across the U.S. and internationally for a wide variety of chemical, physical, and mineralogical analyses. KSSL also typically receives natural fabrics, clods, and cores. Undisturbed clods are used to investigate micromorphology and determine some physical properties, e.g., bulk density.

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**Laboratory Sample Collection and Preparation (1B)**

**Soil (1B1)**

**Bulk Samples (1B1a1)**

Laboratory identification numbers and preparation codes are assigned to bulk soil samples. These identification numbers are unique client- and laboratory-assigned numbers that carry important information about the soil sample (e.g., pedon, soil horizon, location, and year sampled). Laboratory preparation codes depend on the properties of the sample and on the requested analyses. These codes carry generalized information about the characteristics of the analyzed fraction, i.e., the water content (e.g., air-dry, field-moist) and the original and final particle-size fraction (e.g., sieved <2-mm fraction processed to 75 µm) and, by inference, the type of analyses performed. Identification numbers and preparation codes are reported on the KSSL Primary Characterization Data Sheets. Refer to the Soil Survey Investigations Report No. 45, “Soil Survey Laboratory Information Manual” (Soil Survey Staff, 2011), for a detailed explanation of sample identification numbers. Since the publication of SSIR No. 42, version 3 (1996), there has been a significant revision of these preparation codes. This version (5) of SSIR No. 42 (2014) does not describe in detail the revised preparation codes. Detailed information on the current preparation codes as they appear on the Primary Characterization Data Sheets may be obtained from the KSSL upon request.

All soils from quarantined areas are strictly controlled under quarantine regulations 7 CFR 330 of the United States Department of Agriculture, Animal and Plant Health Inspection Service (APHIS). For preparation methods for soil bulk samples, proceed to 1B1b.
Laboratory Sample Collection and Preparation (1B)
Soil (1B1)
   Natural Fabrics, Clods, and Cores (1B1a2-4)

   Laboratory identification numbers and preparation codes are assigned to natural fabrics (NF), clods, and cores. These identification numbers typically relate to a corresponding bulk sample. Refer to 1B1a1 for information on these identification numbers, preparation codes, and soil quarantine regulations.

Laboratory Sample Collection and Preparation (1B)
Soil (1B1)
   Soil Sample Preparation (1B1b)

1. Application

   In some previous editions of SSIR No. 42, laboratory preparation procedures were described as stand-alone methods based on various procedures summarized by specific preparation codes that are reported on the KSSL Primary Characterization Data Sheets. In this version of SSIR No. 42 (2014), these procedures are described more as a procedural process. A process approach is appropriate in that any one sample received from the field may result in a number of laboratory subsamples being collected and prepared based on analytical requests and type of materials. This approach provides the hierarchy whereby laboratory procedures are described in this version of SSIR No. 42. The intent herein is not to detail all possibilities but to describe some of the master preparation procedures that are typically requested for laboratory analyses.

   Laboratory analyses of soil samples are generally performed on the air-dry, fine-earth (<2-mm) fraction (method 1B1b2b1). Air-dry is generally the optimum water content to handle and to process soil. In addition, the weight of air-dry soil remains relatively constant and the biological activity is low during storage. For routine soil analyses, most U.S. and Canadian laboratories homogenize and process samples to pass a 2-mm sieve (Bates, 1993). For some standard air-dry analyses, the <2-mm fraction is further processed so as to be in accordance with a standard method, e.g., Atterberg limits (1B1b2c1); to meet the sample preparation requirements of the analytical instrument, e.g., total C, N, and S (1B1b2d1); or to achieve greater homogeneity of sample material, e.g., total elemental analysis (1B1b2e1) and carbonates and/or gypsum (1B1b2d2). Additionally, some standard air-dry analyses require non-sieved material, e.g., whole-soil samples for aggregate stability (1B1b2a1).

   A field-moist, <2-mm sample is prepared for some analyses, e.g., water retention (1B1b1b2), particle-size analysis, and plasticity index for Andisols and Spodosols (1B1b1c1), if the physical properties of a soil are irreversibly altered by
air-drying, and/or if moist chemical analyses (1B1b1b1) are appropriate. Some biological analyses require field-moist samples (1B1b1a2), as air-drying may cause significant changes in the microbial community. The decomposition state of organic materials is used in “Keys to Soil Taxonomy” (Soil Survey Staff, 2014) to define sapric, hemic, and fibric organic materials, and thus the evaluation of these materials (Histosol analysis) requires a field-moist, whole-soil sample (1B1b1a1). Knowing the amount of rock fragments is necessary for several applications, e.g., available water capacity and linear extensibility. Generally, the >2-mm fractions are sieved, weighed, and discarded (1B1b2f1a) and are excluded from most chemical, physical, and mineralogical analyses. Some exceptions include, but are not limited to, samples containing coarse fragments with carbonate- or gypsum-indurated material or material from Cr and R soil horizons. In these cases, the coarse fragments may be crushed to <2 mm and the analytical results reported on that fraction, e.g., 2 to 20 mm (1B1b2f1a2a), or the coarse fragments and fine-earth material are homogenized and crushed to <2 mm and laboratory analyses are made on the whole-soil (1B1b2f1b1a). Additionally, depending on the type of soil material, samples can be tested for the proportion and particle-size of air-dry rock fragments that resist abrupt immersion in tapwater (1B1b2f1a3).

2. Summary of Method

Any one soil sample received from the field may result in a number of laboratory subsamples being collected and prepared based on the properties of the sample and on the requested analyses. For most standard chemical, physical, and mineralogical analyses, the field sample is air-dried, crushed, and sieved to <2 mm. Field-moist, fine-earth fraction samples are processed by forcing the material through a 2-mm screen by hand or with a large rubber stopper and then are placed in a refrigerator for future analysis. Depending on the nature of the soil material and requested analyses, air-dry and/or field-moist samples may also be prepared as whole-soil samples or processed further to fractions finer than <2 mm, e.g., <0.425 mm for Atterberg limits and ≈180 µm for chemical analysis of organic materials. Air-dry, <75 µm subsamples for major and trace elements are processed metal-free.

Generally, weight measurements are made and recorded on the 20- to 75-mm, 5- to 20-mm, and 2- to 5-mm fractions. These fractions are then discarded. In some cases, these weight measurements are not recorded. Additionally, some or all of these >2-mm fractions may be processed to a finer fraction and saved for chemical, physical, and mineralogical analysis. For example, after the respective weights of the 5- to 20-mm and 2- to 5-mm fractions are recorded, these fractions may be recombined, crushed to <2 mm in a laboratory jaw crusher, and saved for laboratory analysis. In other cases, the fine-earth fraction and the >2-mm fractions are homogenized, passed through a laboratory jaw crusher to reduce all material to pass a 2-mm sieve, and saved for laboratory analysis.
3. Interferences

Soil variability and sample size are interferences to sample collection and preparation. At each stage of sampling, an additional component of variability, the variability among smaller elements within the larger units, is added to the sampling error (Petersen and Calvin, 1986). Soil material needs to be in adequate amount and thoroughly mixed to obtain a representative sample.

Soil is mixed by moving it from the corners to the middle of the processing area and then redistributing the material. This process is repeated four times. Enough soil material needs to be sieved and weighed if a statistically accurate rock fragment content is to be obtained. In order to accurately measure rock fragments with a maximum particle diameter of 20 mm, the minimum specimen size (“dry” weight) that needs to be sieved and weighed is 1.0 kg. Refer to American Society for Testing and Materials (ASTM) Standard Practice D 2488 (ASTM, 2012). A homogenized soil sample is more readily obtained from air-dry material than from field-moist material. Whenever possible, “moist” samples or materials should have weights two to four times larger than those of “dry” specimens (ASTM, 2012).

4. Safety

Dust from the sample processing is a nuisance. Use a mask, face-shield, goggles, and/or respirators when operating dust generating equipment. Keep clothing and hands away from the crusher and pulverizer when these machines are in use. Use downdraft tables for sample processing procedures. The HCl used to check carbonates can destroy clothing and irritate skin. Immediately use water to rinse acid from clothing or skin; seek professional medical help if needed.

5. Equipment

5.1 Electronic Balance, ±1-g sensitivity and 15-kg capacity
5.2 Cardboard trays for sample storage
5.3 Trays, plastic, tared
5.4 Sieves, square-hole, stainless steel
   5.4.1 80 mesh, 180 µm
   5.4.2 10 mesh, 2 mm
   5.4.3 4 mesh, 4.75 mm
   5.4.4 19 mm, ¾ in
   5.4.5 76 mm, 3 in
5.5 200-mesh, 75 µm, nylon cloth sieve
5.6 40-mesh, 0.425 mm
5.7 Pulverizer
5.8 Wooden rolling pin
5.9 Rubber roller
5.10 Laboratory jaw crusher, Model BB200, Retsch Inc., Newton, PA
5.11 Metal plate, 76 x 76 x 0.5 cm
5.12 Containers, paper, 12-oz, with lids
5.13 Containers, plastic, 1-pint, 1, 4, and/or 8 oz with tops
5.14 Scintillation glass vials, 20-mL
5.15 Metal weighing cans, 2-oz
5.16 Brown Kraft paper
5.17 Air compressor, Cast-iron Series, SpeedAire, Campbell Hausfeld Mfg. Co., Harrison, OH
5.18 Planetary ball mill, Fritsch, Model P-5, Lab Synergy, VWR, Radnor, PA
5.19 Rotor mill, Fritsch, Model P-14, VWR, Radnor, PA
5.20 Stein Mill, Steinlite Corp., Atchison KS
5.21 Syalon balls, 12- to 15-mm, and bowls, 80-mL
5.22 Metal weighing cans, 2 oz
5.23 Cross beater mill, Fritsch, Model P-16, VWR, Radnor, PA
5.24 Downdraft tables, 36 in x 72 in, Dynamo Downdraft Cartridge Table, Air Cleaning Specialists, Inc., Fenton, MO

6. Reagents

6.1 Reverse osmosis (RO) water
6.2 1 N HCl
6.3 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate \( (\text{NaPO}_3)_6 \) and 7.94 g of sodium carbonate \( (\text{Na}_2\text{CO}_3) \) in 1 L of RO water.

7. Procedures

**Soil Bulk Sample Preparation (1B1b)**

7.1 Weigh soil sample in sample bag to nearest g when logged-in and record weight.
7.2 Weigh sample in sample bag to nearest g before air-drying and record weight.
7.3 Remove soil sample from sample bag and distribute on a plastic tray. Thoroughly mix soil material.
7.4 If field-moist subsamples are requested, proceed to method 1B1b1.
7.5 If only air-dry subsamples are requested, proceed to method 1B1b2.
Field-Moist Preparation (1B1b1)

7.6 If field-moist, whole-soil subsamples are requested, proceed to method 1B1b1a.

7.7 If only field-moist, <2-mm subsamples are requested, proceed to method 1B1b1b.

Whole-Soil (1B1b1a)

7.8 Methods for field-moist, whole-soil subsamples include but are not limited to 1B1b1a1 and 1B1b1a2 as follows:

Histosol Analysis (1B1b1a1)

7.9 For Histosol analysis, select material for representative subsamples from at least five different areas on the plastic tray. Prepare a subsample of the field-moist, whole soil prepared in an 8-oz container. Store in the refrigerator for future analysis.

Biological Analysis (1B1b1a2)

7.10 For biological analyses, select material for representative subsamples from at least five different areas on the plastic tray. Prepare a subsample of field-moist, whole soil prepared in a plastic container. This subsample is termed a biology bulk sample, which may be obtained from the bulk soil sample upon arrival at the laboratory. Store the biology bulk sample in the refrigerator for future analysis. For additional information on biology collection and preparation methods, proceed to 1B3.

<2-mm Fraction (1B1b1b)

7.11 Methods for field-moist, <2-mm subsamples include but are not limited to 1B1b1b1, 1B1b1b2, 1B1b1b3, and 1B1b1c1 as follows:

Chemical and Selected Physical Analysis (1B1b1b1)

7.12 For moist chemical analysis, select material for representative subsamples from at least five different areas on the plastic tray. Process a subsample of field-moist material by forcing the material through a 2-mm screen by hand or with a large rubber stopper and then place in a 1-pint plastic container. Store in the refrigerator for future analysis.

1500-kPa Water Content (1B1b1b2)

7.13 For moist 1500-kPa water content of mineral and organic material, select material for representative subsamples from at least five different areas on
the plastic tray. Process a subsample of field-moist material by forcing the material through a 2-mm screen by hand or with a large rubber stopper and then place in a 4-oz plastic container. Store in the refrigerator for future analysis.

**Field-Moist/Oven-Dry Ratio (1B1b1b3)**

7.14 For field-moist/oven-dry (FMOD) ratio (which is required if any moist analyses are determined), select material for representative subsamples from at least five different areas on the plastic tray. Process a subsample of field-moist material by forcing the material through a 2-mm screen by hand or with a large rubber stopper and then place in a 2-oz metal weighing can.

**<2-mm Fraction Sieved to <0.425 mm (1B1b1c)**

**Atterberg Limits (1B1b1c1)**

7.15 For moist Atterberg limits, select material for representative subsamples from at least five different areas on the plastic tray. Process a subsample of field-moist material by forcing the material through a 2-mm screen by hand or with a large rubber stopper and then sieve to 40-mesh (0.425 mm). Place subsample in a 12-oz plastic container and store in the refrigerator for future analysis.

**Air-Dry Preparation (1B1b2)**

7.16 Before air-drying, weigh sample on a tared tray (tray weight) to nearest g and record weight.

7.17 Typically, air-dry the sample in an oven at 30 to 35 °C for 3 to 7 days. In some instances, depending on field-moisture content at the time of sample arrival at the laboratory as well as the nature of the material, this drying period may be shortened.

7.18 Weigh sample to nearest g after air-drying and record weight.

7.19 If air-dry, whole-soil subsamples are requested, proceed to method 1B1b2a.

7.20 If only air-dry, <2-mm subsamples are requested, proceed to method 1B1b2b.

**Whole-Soil (1B1b2a)**

7.21 Methods for air-dry, whole-soil subsamples include but are not limited to 1B1b2a1 and 1B1b2a2 as follows:

**Aggregate Stability Analysis (1B1b2a1)**

7.22 For aggregate stability analysis, select material for representative subsamples from at least five different areas on the plastic tray. Prepare an air-dry, whole-soil sample in a 12-oz paper container.
Repository Samples (1B1b2a2)

7.23 For comprehensive sampling projects, prepare a 12-oz paper container of fine-earth material. Also prepare a 12-oz container for natural fabric (NF) sample. Store soil samples in repository. If a NF sample was not selected in the field, select one from air-dried bulk sample. Generally, do not select NF samples for reference projects.

<2-mm Fraction (1B1b2b)

7.24 Weigh sample to nearest g and record weight. This weight includes the >2-mm fractions.

7.25 Roll soil material on a flat, metal plate that is covered with brown Kraft paper using a wooden rolling pin and/or rubber roller to crush clods to pass a 2-mm sieve. For samples that have easily crushed coarse fragments, substitute a rubber roller for a wooden rolling pin. Roll and sieve until only the coarse fragments that do not slake in sodium hexametaphosphate solution remain on sieve. Crush clayey soils that contain no coarse fragments in a laboratory jaw crushe.

7.26 Process air-dry soil by sieving to <2-mm. Thoroughly mix material by moving the soil from the corners to the middle of the processing area and then redistributing the material. Repeat process four times. For preparation of the >2-mm fractions, proceed to method 1B1b2f.

7.27 Methods for air-dry, <2-mm fractions include but are not limited to 1B1b2b1, 1B2b2b2, 1B2b2b3, 1B2b2b4, 1B2b2b6, 1B2b2c1, 1B2b2d1, 1b2b2d2, 1B2b2d3, 1B2b2d4, 1B2b2d5, and 1B2b2e1 as follows:

Standard Chemical, Physical, and Mineralogical Analysis (1B1b2b1)

7.28 For standard chemical, physical, and mineralogical analysis, select material for representative subsamples from at least five different areas on the plastic tray. Prepare one subsample of the air-dry, sieved <2-mm fraction in a 12-oz paper container.

Salt Analysis (1B1b2b2)

7.29 For a saturation paste when salt analyses are requested, select material for representative subsamples from at least five different areas on the plastic tray. Prepare one subsample of the air-dry, sieved <2-mm fraction in a 1-pint plastic container.

1500-kPa Water Content (1B1b2b3)

7.30 For air-dry, 1500-kPa water content analysis of mineral and organic materials, select material for representative subsamples from at least five
different areas on the plastic tray. Prepare a subsample of air-dry, sieved <2-mm fraction in a 1-oz plastic cup.

**Air-Dry/Oven-Dry Ratio (1B1b2b4)**

7.31 For air-dry/oven-dry (AD/OD) ratio analysis (required if any air-dry analysis are determined), select material for representative subsamples from at least five different areas on the plastic tray. Prepare a subsample of the air-dry, sieved <2-mm fraction in a 2-oz metal weighing can.

**Visible and Near-Infrared Diffuse Reflectance Spectroscopy (VNIR–DRS) (1B1b2b6)**

7.32 For visible and near-infrared diffuse reflectance spectroscopy (VNIR–DRS) analysis, select material for representative subsamples from at least five different areas on the plastic tray. Prepare a subsample of air-dry, sieved <2-mm fraction in a 1-oz plastic cup.

**<2-mm Fraction Sieved to <0.425 mm (1B1b2c)**

7.33 For Atterberg limits, select material for representative subsamples from at least five different areas on the plastic tray. Prepare one subsample of the air-dry, sieved <2-mm material sieved to 40-mesh (0.425 mm) in a 12-oz paper container.

**<2-mm Fraction Processed to ≈180 µm (1B1b2d)**

7.34 For total C, N, and S analyses, select material for representative subsamples from at least five different areas on the plastic tray. Prepare one subsample of the air-dry, sieved <2-mm fraction processed in a cross beater mill to ≈80 mesh (180 µm) in a 20-mL scintillation glass vial.

**Calcium Carbonate and Gypsum (1B1b2d2)**

7.35 For the determination of the amount of carbonates and/or gypsum, use the prepared subsample (method 1B1b2d1).

**Chemical Analysis of Organic Materials (1B1b2d3)**

7.36 For chemical analysis of organic materials, select material for representative subsamples from at least five different areas on the plastic tray. Prepare a subsample of the air-dry, sieved <2-mm fraction processed in a cross beater to a fine grind (≈80 mesh, 180 µm) in a 12-oz paper container.
Presence of Carbonates (1B1b2d4)

7.37 To check for the presence of carbonates, use the prepared subsample (method 1B1b2d1). Reference samples (“knowns”) are available for comparisons. Place 1 g of the air-dry fine-earth fraction in porcelain spot plate, add reverse osmosis water, and stir to remove entrapped air. Add 1 N HCl to soil (method 3A2a1, reagent 6.2), observe amount of effervescence, and record as follows:

*None.*—No visual effervescence.

*Very Slight.*—Bubbles rise at a few points in the sample and consistently appear at the same point in either a steady stream of tiny bubbles or in a slower stream of larger bubbles. Do not mistake trapped air bubbles for a positive test. Generally, these air bubbles appear immediately after the addition of 1 N HCl.

*Slight.*—More small bubbles, and possibly a few larger bubbles, appear throughout the sample than with a *very slight* reaction.

*Strong.*—More large bubbles are evident than with a *slight* reaction. Often the reaction is violent at first and then quickly decreases to a reaction that produces many small bubbles.

*Violent.*—The sample effervesces violently. Many large bubbles appear to burst from the spot plate.

Mid-Infrared Diffuse Reflectance Spectroscopy (MIR–DRS) (1B1b2d5)

7.38 To determine mid- and near-infrared diffuse spectroscopy (MIR–DRS) analysis, use the prepared subsample (method 1B1b2d1).

<2-mm Fraction Processed to 75 µm (1B1b2e)

Total Major and/or Trace Elements (1B1b2e1)

7.39 For total major and/or trace element analyses, select material for representative subsamples from at least five different areas on the plastic tray. Prepare one subsample of the air-dry, metal free, sieved <2-mm fraction processed in a planetary ball mill for 2 min and sieved to <75 µm (200 mesh) in a 20-mL scintillation glass vial.

>2-mm Fractions (1B1b2f)

7.40 The following methods are used for samples with >2 mm fractions. These fractions include mineral coarse fragments as well wood fragments that are >20 mm in cross section and cannot be crushed and shredded with the fingers. If the >2-mm fractions are to be weighed, recorded, and discarded with no further laboratory analysis, proceed to method 1B1b2f1a. When the data for >2-mm fractions are not recorded, proceed
to method 1B1b2f1b. If the >2-mm fractions contain carbonate- or gypsum-indurated material, and laboratory analysis is requested, proceed to method 1B1b2f1a1a. If the >2-mm fractions are Cr or R material (Soil Survey Staff, 2014), and laboratory analysis is requested, proceed to method 1B1b2f1b1a. If testing is requested for the proportion and particle-size of the air-dry, >2-mm fraction that resist abrupt immersion in tapwater, proceed to method 1B1b2f1a3.

**Particle-Size Analysis (1B1b2f1)**

**Particle-Size Analysis, Recorded (1B1b2f1a)**

7.41 Process the air-dry soil by sieving to <2 mm as described in method 1B1b2b. In this method (1B1b2f1a), weight measurements are made on the 20- to 75-mm, 5- to 20-mm, and 2- to 5-mm fractions. Weigh soil material with diameters of 2 to 5 mm. If separating the <2-mm fraction from fragments is difficult, soak (100 g of 2- to 5-mm fraction) in sodium hexametaphosphate solution for 12 h. Air-dry and weigh the material that does not slake, record weight, and discard. Weigh, record weight, and discard particles with diameters of 20 to 75 mm and 5 to 20 mm. The <2-mm fraction is saved for chemical, physical, and mineralogical analysis (1B1b2b)

**Particle-Size Analysis, Not Recorded (1B1b2f1b)**

7.42 This method (1B1b2f1b) is the same as described in method 1B1b2f1a, except the weight of the >2-mm fractions are not recorded and all analytical results are reported on a <2-mm basis.

**2- to 20-mm Fraction Processed to <2-mm (1B1b2f1a1) Chemical, Physical, and Mineralogical Analysis (1B1b2f1a1a)**

7.43 This method (1B1b2f1a1a) is commonly used for samples that contain carbonate- or gypsum-indurated material. Process the air-dry soil by sieving to <2 mm as described in method 1B1b2b. Weigh soil material with diameters of 20- to 75-mm, 5 to 20-mm, and 2 to 5-mm and record weights as described in method 1B1b2f1a. The 5- to 20-mm and 2- to 5-mm fractions are then recombined after their respective weights are recorded. The recombined, 2- to 20-mm, material is crushed to <2 mm in a laboratory jaw crusher. This material is saved for laboratory analysis, and analytical results are reported on the 2- to 20-mm basis. The <2-mm material is also saved for chemical, physical, and mineralogical analysis as described in method 1B1b2b. If carbonate or gypsum accumulations are soft and easily pass a 2-mm sieve, the standard method (1B1b2b) is usually requested.
2- to 20-mm Fraction Processed to <2-mm and Recombined with <2-mm Fraction (1B1b2f1a2)
Chemical, Physical, Mineralogical Analysis (1B1b2f1a2a)

7.44 This method (1B1b2f1a2a) is rarely requested. Process the air-dry soil by sieving to <2 mm as described in method 1B1b2b. Weigh soil material with diameters of 20 to 75 mm, 5 to 20 mm, and 2 to 5 mm and record weights as described in method 1B1b2f1a. The 5- to 20-mm and 2- to 5-mm fractions are then recombined after their respective weights are recorded. The recombined, 2- to 20-mm, material is crushed to <2 mm in a laboratory jaw crusher. This material is then recombined with the <2-mm fraction. The analytical results are reported on the <20-mm basis or are calculated to the <2-mm basis.

Particle-Size Analysis, Not Recorded (1B1b2f1b)
Whole-Soil Processed to <2-mm (1B1b2f1b1)
Chemical, Physical, and Mineralogical Analysis (1B1b2f1b1a)

7.45 This method (1B1b2f1b1a) is mainly used to prepare samples from Cr or R soil horizons (Soil Survey Staff, 2014). Homogenize particles with diameters >2 mm and the fine-earth material (<2-mm) and crush to <2-mm in a laboratory jaw crusher. This material is saved for laboratory analysis, and analytical results are reported on the whole-soil basis.

8. Calculations
Calculations for coarse fragments are reported in 3A2.

9. Report
Reported data include but are not limited to the following:

9.1 Weight (g) of field-moist soil sample
9.2 Weight (g) of air-dry soil sample
9.3 Weights (g) of processed air-dry soil
9.4 Weight (g) of 20- to 75-mm fraction
9.5 Weight (g) of 5- to 20-mm fraction
9.6 Weight (g) of 2- to 5-mm fraction
9.7 Weight (g) of subsample of 2- to 5-mm fraction before slaking
9.8 Weight (g) of subsample of 2- to 5-mm fraction after slaking
9.9 Effervescence with HCl (none, very slight, slight, strong, violent)

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.
11. References


Laboratory Sample Collection and Preparation (1B)

Soil (1B1)

Soil Bulk Sample Preparation (1B1b)

Air-Dry Preparation (1B1b2)

>2-mm Fractions (1B1b2f)

Particle-Size Analysis (1B1b2f1)

Particle-Size Analysis, Recorded (1B1b2f1a)

Proportion and Particle-Size of Air-Dry Rock Fragments that Resist Abrupt Immersion in Tap Water (1B1b2f1a3)

1. Application

It is widely accepted that the mechanical preparation of soil for laboratory analysis is difficult to standardize for samples that contain >2-mm particles with rupture resistance intermediate between earthy bodies and fragments of highly resistant rock (Grossman, 2004). A procedure is needed that is both reproducible and that subjects the >2-mm fraction to an intermediate stress less than standard mechanical preparation. Abrupt immersion in tap water of the initially air-dry >2-mm fraction has been chosen as the manner in which stress is applied. For some situations, comparison of the results to the use of a dispersing agent may be valuable, but another procedure would need to be established.
This method (1B1b2f1a3) provides a comparison to mechanical preparation methods and should aid in the development of a more precise mechanical preparation that is dependent on sample properties (Grossman, 2004). Certain analyses can be run on the >2-mm fraction after separation by the water-immersion method. The resulting values can be subtracted from analyses on the whole ground sample to obtain estimates for the <2-mm fraction as removed by the water immersion treatment.

2. Summary of Method

A representative subsample of the air-dry field sample is weighed and passed through a No.10 sieve. The >2-mm fraction is placed abruptly in tapwater, left overnight, gently agitated, passed through a No. 10 sieve, and the wet >2-mm material is passed through a nest of No. 10, No. 4, 9.5-mm (nominally 10 mm), and 20-mm sieves.

3. Interferences

There are no known interferences.

4. Safety

Dust from sample processing is a nuisance. A mask should be worn in order to avoid breathing dust.

5. Equipment

5.1 Buckets, plastic, 10-L with 1-L marks
5.2 Bucket, plastic, in which 20-cm diameter sieve fits snugly
5.3 Bucket, 19-L (5-gal), straight-sided, with 30-cm diameter
5.4 Drying trays, fiberglass, 35 x 48 cm
5.5 Cake pans, aluminum foil, 20 x 20 x 4 cm
5.6 Sieves: 30-cm diameter No. 10, 20-mm diameter No. 4, 9.5- and 20-mm plus bottom pan
5.7 Top loading balance, 1-g sensitivity and >10,000-g capacity, with pan large enough to mount the drying trays listed above in equipment item 5.4
5.8 Pan, plastic, 60 x 40 x 15 cm
5.9 Plastic sheet, 8-mm, large enough to line plastic container
5.10 Plastic beads, 9 x 6 mm, segregated by different colors
5.11 Teaspoon, tablespoon

6. Reagents

6.1 CaCl₂•2H₂O
6.2 Tapwater of acceptable dispersibility (taken as Zone A in Flanagan and Holmgren, 1977).

7. Procedure

7.1 From the field sample, remove and weigh a representative subsample that is roughly one-fourth of the sample and inclusive of the >20-mm fraction.

7.2 Pass subsample through a No. 10 sieve. Weigh the >2-mm particles and discard the <2-mm fraction. Immerse the >2-mm fraction abruptly in tapwater in a 10-L bucket in which there is approximately 1-L of water for roughly every 500 g of >2-mm particles. Leave the sample overnight.

7.3 Insert your hand to the bottom of the bucket and rotate the mass at 1 rotation per second 20 times. In the sink, place the 30-cm diameter No. 10 sieve over the top of a 19-L bucket. Quantitatively transfer the >2-mm particles into the No.10 sieve and wash with a water stream to remove the material that is not resistant to the immersion treatment. Use a minimum amount of water.

7.4 After the 19-L bucket is full, add about 30 g of CaCl$_2$. Allow to stand until settled out. Pour off as much of the water as possible without transferring solids to the sink. Dry the sediment in the bucket by evaporation, followed by disposal as a solid.

7.5 This step follows 7.3 directly if no strength measurements are taken. Quantitatively transfer from the 30-cm diameter sieve to a tray. Air-dry the sample on the tray at 30 to 35 °C. Pass through a nest of 20-mm, No. 4 and No. 10 sieves.

7.6 Weigh each separate to the nearest g and report. Subtract the >20-mm weight from the initial >2-mm weight. Obtain the 2- to 5-mm, 5- to 20-mm, and 2- to 20-mm particles as a percent of the <20-mm fraction.

8. Calculations

8.1 $A = \frac{B}{(C-D)} \times 100$

8.2 $E = \frac{F}{(C-D)} \times 100$

8.3 $G = A + E$

where:
$A =$ Weight percentage 2- to 5-mm fraction on <20-mm basis (%)
$B =$ Air-dry weight of 2- to 5-mm fraction after water immersion (g)
$C =$ Air-dry total weight of whole soil subsample (g)
$D =$ Air-dry weight of >20-mm fraction after water immersion (g)
$E =$ Weight percentage of 5- to 20-mm fraction on <20-mm basis (%)
$F =$ Air-dry weight of 5- to 20-mm fraction after water immersion (g)
$G =$ Weight percentage 2- to 20-mm fraction on <20-mm basis (%)
9. Report
   Report the 2- to 5-mm, 5- to 20-mm, and 2- to 20-mm fractions as weight percentages of the <20-mm fraction.

10. Precision and Accuracy
   Precision and accuracy data are available from the KSSL upon request.

11. References

Laboratory Sample Collection and Preparation (1B)
Water (1B2)
Samples (1B2a)
   As for soil samples, laboratory identification numbers and preparation codes are assigned to water samples. Refer to 1B1a1 for information on these identification numbers and preparation codes.

Laboratory Sample Collection and Preparation (1B)
Water (1B2)
Water Sample Preparation (1B2b)
   Avoid long periods between collection and laboratory analysis of water. To prevent significant changes (e.g., degradation, volatilization), water samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory. Do not allow water samples to freeze, which can influence pH and the separation of dissolved organic matter from the water phase.
   Some water analyses (e.g., electrical conductivity, total C, inorganic C) need to be performed promptly, as optimal preservation is not possible (Velthorst, 1996). Upon completion of these analyses, sample filtration (0.45-µm membrane) is used to separate dissolved from suspended material. The sample is then split into two subsamples. One subsample is acidified to pH 2 for cation analyses (e.g., Al, Fe, Mn) and the other for anion analyses. These other water analyses also need to be performed as promptly as possible.
Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)

Samples (1B3a)
  Biology Bulk Sample (1B3a1)
  Microbial Biomass Sample (1B3a2)

As for soil and water samples, laboratory identification numbers and preparation codes are assigned to biology samples. Refer to 1B1a1 for information on these identification numbers and preparation codes. Some biology samples arrive at the laboratory as part of the soil bulk sample. If this is the case, biological subsamples are collected and prepared as described in 1B1b1a2. In other cases, biology bulk samples may be split in the field (1B3a1) and are separate sampling units from the soil bulk sample (1B1a1). Additionally, some biological samples, e.g., microbial biomass (1B3a2), are separate units from the soil bulk or other biology samples. These samples require expedited transport under ice or gel packs and are refrigerated (4 °C) immediately upon arrival at the laboratory.

Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)

Biological Material Preparation (1B3b)

Avoid long periods between collection and laboratory analysis of biological samples to prevent significant changes (e.g., microbial community). Store biology samples in the refrigerator (4 °C) for future analysis. Biological preparation includes but is not limited to the following.

Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)

Biological Material Preparation (1B3b)
  Field-Moist Preparation (1B3b1)
    <2-mm Fraction (1B3b1a)
      Microbial Biomass (1B3b1a1)

Refer to the section on soil biological and plant analyses (6) for additional information on the further processing and preparation of these biological samples for laboratory analysis.
Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)
  Biological Material Preparation (1B3b)
    Air-Dry Preparation (1B3b2)
      <2-mm Fraction Sieved to <53 µm, with ≥ 53 µm (Particulate Organic Matter) and <53 µm Processed to ≈180 µm (1B3b2a)
        Total Carbon, Nitrogen, and Sulfur (1B3b2a1)
    Refer to the section on soil biological and plant analyses (6) for additional information on the further processing and preparation of these biological samples for laboratory analysis.

Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)
  Biological Material Preparation (1B3b)
    Air-Dry Preparation (1B3b2)
      <2-mm Fraction (1B3b2b)
        Other Biological Analyses (1B3b2b1)
    Refer to the section on soil biological and plant analyses (6) for additional information on the further processing and preparation of biological samples for laboratory analysis.

Laboratory Sample Collection and Preparation (1B)
Biological Materials (1B3)
  Biological Material Preparation (1B3b)
    Dry (50˚C) Preparation (1B3b3)
      Roots (1B3b3a)
        Root Biomass (1B3b3a1)
          Roots Processed to ≈180 µm (1B3b3a1a)
            Total Carbon, Nitrogen, and Sulfur (1B3b3a1a1)
      Plants (1B3b3b)
        Plant Biomass (1B3b3b1)
          Plants Processed to ≈180 µm (1B3b3b1a)
            Total Carbon, Nitrogen, and Sulfur (1B3b3b1a1)
    Refer to the section on soil biological and plant analyses (6) for additional information on the further processing and preparation of these biological samples for laboratory analysis.
CONVENTIONS (2)

Methods and Codes (2A)

The KSSL ensures continuity in its analytical measurement process with the use of standard operating procedures (SOPs). A standard method is defined herein as a method or procedure developed by an organization, based on consensus opinion or other criteria and often evaluated for its reliability by a collaborative testing procedure (Taylor, 1988). A SOP is written in a standard format and adopted for repetitive use when performing a specific measurement or sampling operation, i.e., a SOP may be a standard or one developed by a user (Taylor, 1988).

The use of SOPs provides consistency and reproducibility in soil preparations and analyses and helps to ensure that these preparations and analyses provide results of known quality. SSIR No. 42, “Soil Survey Laboratory Methods Manual,” version 5.0, replaces as a methods reference all earlier versions of SSIR No. 42 (1989, 1992, 1996, and 2004) and SSIR No. 1, “Procedures for Collecting Soil Samples and Methods of Analysis for Soil Survey” (1972, 1982, and 1984). All KSSL methods are performed with methodologies appropriate for the specific purpose. The KSSL SOPs are standard methods, peer-recognized methods, KSSL-developed methods, and/or methods specified in “Keys to Soil Taxonomy” (Soil Survey Staff, 2014). This manual also serves as the primary document from which a companion manual, SSIR No. 45, “Soil Survey Laboratory Information Manual” (Soil Survey Staff, 2011), was developed. SSIR No. 45 describes in greater detail the application of KSSL data.

Included in SSIR No. 42 are descriptions of current as well as obsolete methods, both of which are documented by method codes and linked with analytical results that are stored in the KSSL database. This linkage between laboratory method codes and the respective analytical results is reported on the KSSL data sheets. Reporting the method by which the analytical result is determined helps to ensure user understanding of KSSL data. In addition, this linkage provides a means of technical criticism and traceability if data are questioned in the future.

The methods in current use at the KSSL are described in SSIR No. 42 in enough detail that they can be performed in many laboratories without reference to other sources. Descriptions of the obsolete methods are located at the back.
of this manual. Information is not available to describe some of these obsolete procedures in the same detail as used to describe the current methods in the laboratory.

Since the 1996 publication of SSIR No. 42 (Soil Survey Staff, 1996), there has been a significant increase in the number and kinds of methods performed at the KSSL, resulting in a re-structuring of the method codes. As in past versions of SSIR No. 42, the current method codes are hierarchical and alphanumerical. The older method code structure, however, used only a maximum of four characters, e.g., 6A1b. The new structure has more characters, carrying more information about the method, e.g., particle-size and sample weight bases for reporting data. This version of SSIR No. 42 carries not only the new method codes but also the older ones as well. This linkage between the two method code systems is important to maintain, as many older KSSL data sheets and scientific publications report these older codes.

The KSSL data have been provided in reports, e.g., Primary and Supplementary Characterization Data Sheets, and in electronic forms, including tapes, disks, and CD-ROMs. More recently, other reports have been developed, e.g., Soil Taxonomy Characterization Data Sheets, and data are available via the NRCS Soils website http://soils.usda.gov/. Historically, SSIR No. 42 described and assigned method codes to only those data reported on the Primary Characterization Data Sheets. This tradition is followed in this version of the SSIR No. 42. With the exception of some KSSL primary analytical data included for user convenience, the data on the Supplementary and Taxonomy Characterization Data Sheets are derived data. They use analytical data as a basis for calculation and do not carry method codes. Data on the Supplementary and Taxonomy Characterization Data Sheets are not described in this manual. Other calculated data appear on the Primary Characterization Data Sheets as “Pedon Calculations”, e.g., Weighted Average Clay. These calculated data are neither assigned method codes nor described in this manual. For more detailed information about the calculation and application of these derived values, refer to SSIR No. 45, “Soil Survey Laboratory Information Manual” (Soil Survey Staff, 2011) and “Keys to Soil Taxonomy” (Soil Survey Staff, 2014). Additional information may also be obtained from the KSSL upon request.

Data Types (2B)

The methods described herein identify specific types of analytical or calculated data. Most of these methods are analytical in nature, i.e., quantitative or semiquantitative measurements, and include physical, chemical, mineralogical, and biological analyses. Sample collection and preparation in the field and the laboratory are also described. Historically, SSIR No. 42 described some derived values, e.g., coefficient of linear extensibility (COLE) and water retention
difference (WRD). These values were reported along with the analytical data on the Primary Characterization Data Sheets. This version of SSIR No. 42 follows this tradition. In recent publications of SSIR No. 42, a few more derived values have been added to the Primary Characterization Data Sheets. Although these data have been assigned method codes, detailed descriptions are not included in this version of SSIR No. 42. For more detailed information about the calculation and application of these derived values, refer to SSIR No. 45 (Soil Survey Staff, 2011) and “Keys to Soil Taxonomy” (Soil Survey Staff, 2014).

**Particle-Size Fraction Base for Reporting Data (2C)**

**Particles <2 mm (2C1)**

**Particles >2 mm (2C2)**

The reporting conventions for particle-size fractions for the <2-mm and >2-mm fractions are herein designated as 2C1 and 2C2, respectively. Unless otherwise specified, all data are reported on the basis of the <2-mm material. Other size fractions reported on the Primary Characterization Data Sheets include, but are not limited to, the <0.4, <20, <75 mm, and whole-soil bases. The maximum coarse-fragment size for the >2-mm base varies. The base usually includes those fragments as large as 75 mm (3 in) present in the soil. The maximum size for fragments >75 mm, commonly termed “whole soil,” includes boulders with maximum horizontal dimensions less than those of the pedon. The maximum particle-size set is recorded in the parentheses in the column heading of the data sheet. The base with which to calculate the reported >2-mm percentages includes all material in the sample smaller than the particle size recorded in the column heading.

**Sample Weight Base for Reporting Data (2D)**

**Air-Dry/Oven-Dry (2D1)**

**Field-Moist/Oven-Dry (2D2)**

**Correction for Crystal Water (2D3)**

Unless otherwise specified, all data are reported on an oven-dry weight or volume basis for the designated particle-size fraction. The calculation of the air-dry/oven-dry (AD/OD) ratio is used to adjust AD results to an OD weight basis. If required for a procedure, the AD/OD ratio is also used to calculate the sample weight that is equivalent to the required OD soil weight. The AD/OD ratio, unless otherwise specified, is determined on a <2-mm sieved sample by method 3D1. Currently, if a <2-mm sample is “fine grind” (e.g., <180 µm for total C or <75 µm for trace elements) or if a subset of a <2-mm sample is further prepared (≥0.53- and <0.53-µm fractions for hyper-particulate organic matter or the 2- to 0.5-mm fraction for aggregate stability), the AD/OD of the <2-mm sieved sample as
prepared in method 3D1 is used in KSSL method calculations (if required). For the methods of analysis and calculations of the AD/OD and FM/OD ratios, refer to methods 3D1 and 3D2, respectively. For correction of crystal water, refer to method 2D3.

AD and OD weights are defined herein as constant sample weights obtained after drying at 30 ±5 °C (≈3 to 7 days) and at 110 ±5 °C (≈12 to 16 h), respectively. As a rule of thumb, air-dry soils contain about 1 to 2 percent water and are drier than soils at 1500-kPa water content. FM weight is defined herein as the sample weight obtained without drying prior to laboratory analysis. In general, these weights are reflective of the water content at the time of sample collection.

**Significant Figures and Rounding (2E)**

Unless otherwise specified, the KSSL uses the procedure of significant figures to report analytical data. Historically, significant figures are said to be all digits that are certain plus one, which contains some uncertainty. If a value is reported as 19.4 units, the 0.4 is not certain, i.e., repeated analyses of the same sample would vary more than one-tenth but generally less than a whole unit.

**Data Sheet Symbols (2F)**

The analytical result of “zero” is not reported by the KSSL. The following symbols are used or have been used for trace or zero quantities and for samples not tested.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>tr, Tr, TR</td>
<td>Trace, either is not measurable by quantitative procedure used or is less than reported amount.</td>
</tr>
<tr>
<td>tr(s)</td>
<td>Trace, detected only by qualitative procedure more sensitive than quantitative procedure used.</td>
</tr>
<tr>
<td>-</td>
<td>Analysis run but none detected.</td>
</tr>
<tr>
<td>--</td>
<td>Analysis run but none detected.</td>
</tr>
<tr>
<td>-(s)</td>
<td>None detected by sensitive qualitative test.</td>
</tr>
<tr>
<td>blank</td>
<td>Analysis not run.</td>
</tr>
<tr>
<td>nd</td>
<td>Not determined, analysis not run.</td>
</tr>
<tr>
<td>&lt;</td>
<td>Either none is present or amount is less than reported amount, e.g., &lt;0.1 is in fact &lt;0.05 since 0.05 to 0.1 is reported as 0.1.</td>
</tr>
</tbody>
</table>
SOIL PHYSICAL AND FABRIC-RELATED ANALYSES (3)

Particle-Size Distribution Analysis (3A)
Particles <2 mm (3A1)
  Pipette Analysis (3A1a)

1. Application

  General: One of the most requested KSSL characterization analysis is particle-size distribution analysis (PSDA). The behavior of most soil physical properties and many chemical properties are sharply influenced by presence and relative abundance of the particle-size distribution classes. Precise meaning is given to the term soil texture only through the concept of particle-size distribution (Skopp, 1992).

  Particle-size distribution analysis is a measurement of the size distribution of individual particles in a soil sample. These data may be presented on a
cumulative PSDA curve. These distribution curves are used in many kinds of investigations and evaluations, e.g., geologic, hydrologic, geomorphic, engineering, and soil science (Gee and Bauder, 1986). In soil science, particle-size is used as a tool to explain soil genesis, quantify soil classification, and define soil texture.

In the USDA classification system (Soil Survey Division Staff, 1953, 1993), soil texture refers to the relative proportions of clay, silt, and sand on a <2-mm basis. The system also recognizes proportions of five subclasses of sand. Other classification systems include the particle-size classes for differentiation of families in soil taxonomy (Soil Survey Staff, 2014) and the systems of the International Union of Soil Science (IUSS); the Canadian Soil Survey Committee (CSSC); and the American Society for Testing and Materials (ASTM). When data are reported and interpreted, it is important to recognize that these other classification systems are frequently cited in the literature, especially engineering systems, e.g., American Association of State Highway and Transportation Officials (AASHTO) and the ASTM Unified Soil Classification System (USCS) (Gee and Bauder, 1986).

**Standard KSSL PSDA (3A1a1):** The procedure described herein covers the destruction or dispersion of <2-mm diameter soil aggregates into discrete units by chemical, mechanical, or ultrasonic means, followed by the separation or fractionation of these particles according to size limits through sieving and sedimentation (Gee and Bauder, 1986). Upon isolation of these particle-sizes or size increments, the amount of each size-fraction is then gravimetrically measured as a percent of the total sample weight on an oven-dry basis. The Kilmer and Alexander (1949) pipette method was chosen by the USDA Soil Conservation Service because it is reproducible in a wide range of soils.

The KSSL routinely determines the soil separates of total sand (0.05 to 2.0 mm), silt (0.002 to 0.05 mm), and clay (<2 µm), with five subclasses of sand (very coarse, coarse, medium, fine, and very fine) and two subclasses of silt (coarse and fine). Coarse silt is a soil separate with particle diameter of 0.02 to 0.05 mm. In the International Classification system, 0.02 mm (20 µm) is the break between sand and silt. The particle-size separation at 20 µm also has significance in optical microscopy. This class limit represents the optical limits of the polarizing light microscope. Particle-size classes are a compromise between engineering and pedological classes (Soil Survey Staff, 2014). In engineering classifications, the limit between sand and silt is a diameter of 0.074 mm. The break between sand and silt is 0.05 and 0.02 mm in the USDA and International classification systems, respectively. In engineering classes, the very fine sand (VFS) separate is split. In particle-size classes of soil taxonomy (Soil Survey Staff, 2014), the VFS is allowed to “float,” i.e., VFS is treated as sand if the texture is fine sand, loamy fine sand, or a coarser class and is treated as silt if the texture is very fine sand, loamy very fine sand, sandy loam, silt loam, or a finer class (Soil Survey Staff, 2014). Refer to the Soil Survey Staff (2014) for additional discussion on particle-size classes.
**Fine-Clay and Carbonate Clay:** In addition to routinely determining soil separates of sand, silt, and clay, the KSSL determines the fine-clay and/or carbonate clay fractions, depending on analytical requests and properties of the sample. The fine-clay fraction consists of mineral soil particles that have an effective diameter of <0.2 µm. The percentage fine clay is determined for soils that are suspected of having illuviated clay. Fine-clay data can be used to determine the presence of argillic horizons or as a tool to help explain soil genesis. The carbonate-clay fraction is considered important in PSDA because clay-size carbonate particles have properties that are different from non-carbonate clay. The cation-exchange capacity of carbonate clay is very low compared to non-carbonate clay. The water holding capacity of carbonate clay is about two-thirds that of non-carbonate clay. Because carbonate clay is a diluent, it is often subtracted from the total clay in order to make inferences about soil genesis and clay activities.

**Pretreatment and Dispersion Techniques:** The phenomena of flocculation and dispersion (deflocculation) are very important in determining the physical behavior of the colloidal fraction of soils and thus, indirectly, have a major bearing on the physical properties that soils exhibit (Sumner, 1992). In the standard KSSL method 3A1a1, soils are pretreated to remove organic matter and soluble salts. Samples are chemically treated with hydrogen peroxide and sodium hexametaphosphate to improve dispersion. The primary objectives of dispersion are the removal of cementing agents, the rehydration of clays, and the physical separation of individual soil particles (Skopp, 1992). The hydrogen peroxide oxidizes the organic matter. The sodium hexametaphosphate complexes any calcium in solution and replaces it with sodium on the exchange complex. Upon completion of the chemical treatments, mechanical agitation through shaking is used to enhance separation of particles and to facilitate fractionation.

Complete dispersion by method 3A1a1 may be prevented in the presence of cementing agents, such as calcium carbonate, Fe, and Si. In these instances, special pretreatment and dispersion methods (3A1a2-5) may be performed upon request by the project coordinator as follows:

**Carbonate Removal (3A1a2):** Soils high in carbonate content do not readily disperse. Pretreatment of these soils with acid removes the carbonates (Grossman and Millet, 1961; Jackson, 1969; Gee and Bauder, 1986; Gee and Or, 2002). The determination of particle-size distribution after the removal of carbonates is used primarily for studies of soil genesis and parent material.

**Iron Removal (3A1a3):** Iron oxides and other oxides coat and bind particles of sand, silt, and clay to form aggregates. Soils with iron cementation do not readily disperse. The iron oxides are removed using bicarbonate-buffered, sodium dithionite-citrate solution (Mehra and Jackson, 1960; Gee and Bauder, 1986; Gee and Or, 2002).

**Silica Removal (3A1a4):** Soils that are cemented by Si do not completely disperse with hydrogen peroxide pretreatment and sodium hexametaphosphate.
A pretreatment with a weak base dissolves the Si bridges and coats and increases the soil dispersion. This determination is used for soil parent material and genesis studies.

**Ultrasonic Dispersion (3A1a5):** Soils that are not completely dispersed by standard PSDA can be dispersed using ultrasonic dispersion (Gee and Bauder, 1986; Gee and Or, 2002). Pretreatments coupled with ultrasonic dispersion yield maximum clay concentrations (Mikhail and Briner, 1978). This is a developmental procedure because no standard method has been adopted using ultrasonic dispersion.

**Water Dispersible PSDA (3A1a6):** This method provides a means of evaluating the susceptibility of a soil to water erosion. The degree to which a soil disperses without the oxidation of organic matter, the removal of soluble salts, or the addition of a chemical dispersant may be compared with results from chemical dispersion (Bouyoucos, 1929).

**Gypseous PSDA (3A1a7):** This PSDA method is primarily intended for soils that have ≥40% gypsum but is available upon request for soils with lesser amounts of gypsum. Soil taxonomy was recently amended to more comprehensively recognize properties of gypseous soils (Pearson et al., 2014). Among the revisions was the addition of substitute particle size classes (family level of classification) for soils with ≥40% gypsum in the <20 mm size fraction (gypseous soils) (Soil Survey Staff, 2014). The added substitute particle-size classes were “gypseous-skeletal” for gypseous soils that have ≥35% coarse fragments and two classes for gypseous soils that have <35% coarse fragments: "coarse-gypseous" (≥50% particles 0.1 to 2.0 mm diameter) and “fine-gypseous” (<50% particles 0.1 to 2.0 mm diameter) (Soil Survey Staff, 2014). These taxa were established to enable better interpretations of soils with high amounts of gypsum, especially interpretations of those properties related to water retention and movement. These definitions apply to “particles” with any mineralogy, including gypsum (Pearson et al., 2014).

**Field-Moist PSDA (3A1a1b):** The standard KSSL procedure for particles with <2-mm diameter is the air-dry method (3A1a1a). Although a homogenized sample is more easily obtained from air-dry material than from moist material, some soils irreversibly harden when dried. In such cases, moist PSDA (3A1a1b) may be used. The phenomenon of aggregation through oven- or air-drying is an important example of irreversibility of colloidal behavior in the soil-water system (Kubota, 1972; Espinoza et al. 1975). Drying such soils decreases the measured clay content. This decrease can be attributed to the cementation upon drying (Maeda et al., 1977). The magnitude of the effect varies with the particular soil (Maeda et al., 1977).

**KSSL PSDA Process:** In SSIR No. 42 (1996), stand-alone methods were described for the non-routine pretreatment and dispersion techniques as well as for the analysis of particles not routinely reported, e.g., fine and/or carbonate-clay fractions. In subsequent versions of SSIR No. 42, these procedures are described.
more as a procedural process. This approach is appropriate in that certain procedural steps may be modified, omitted, or enhanced by the investigator, depending on the properties of the sample and on the requested analyses. The process by which specific procedural steps are selected for sample analysis is based upon knowledge or intuition of certain soil properties or related to specific questions, e.g., special studies of soil genesis and parent material.

2. Summary of Method

Standard KSSL PSDA: The standard KSSL procedure for analysis of particles with diameters <2-mm is the air-dry method (method 3A1a1a), whereby a 10-g sample is pretreated to remove organic matter and soluble salts. The sample is dried in an oven to obtain the initial weight, dispersed with a sodium hexametaphosphate solution, and mechanically shaken. The sand fraction is removed from the suspension by wet sieving and then fractionated by dry sieving. The clay and fine silt fractions are determined using the suspension remaining from the wet sieving process. This suspension is diluted to 1 L in a sedimentation cylinder and stirred, and then 25-mL aliquots are removed with a pipette at calculated, predetermined intervals based on Stokes' law (Kilmer and Alexander, 1949). The aliquots are dried at 110 °C and weighed. Coarse silt is the difference between 100% and the sum of the sand, clay, and fine silt percentages.

Fine-Clay Determination: The soil suspension from method 3A1a1a is used to determine the fine-clay fraction. This suspension is stirred, poured into a centrifuge bottle, and centrifuged at 1500 rpm. A 25-mL aliquot is withdrawn with a pipette. The aliquot is dried in an oven and weighed, and the percentage of fine clay is calculated based on the total sample weight. The time of centrifugation is determined from the following equation modified from Stokes’ law (Jackson, 1969).

\[ t_m = \frac{63.0 \times 10^8 \eta \log(rs^{-1})}{(N_m^2 \Delta \rho)} \]

where:
- \( t_m \) = Time in minutes
- \( \eta \) = Viscosity in poises
- \( r \) = Radius in cm from center of rotation to sampling depth (3 cm + s)
- \( s \) = Radius in cm from center of rotation to surface of suspension
- \( N_m \) = rpm (1500)
- \( D\mu \) = Particle diameter in microns (0.2 \( \mu \)m)
- \( \Delta \rho \) = Difference in specific gravity between solvated particles and suspension liquid

63.0 \( \times 10^8 \) = Combination of conversion factors for convenient units of time in minutes, \( t_{min} \), \( N_m \) as rpm, and particle diameter in microns, \( D\mu \).

Carbonate-Clay Determination: The residue from method 3A1a1a is used to determine the carbonate-clay fraction. This residue is treated with acid in a closed
system. The pressure of the evolved gas is measured. The pressure is related linearly to the CO₂ content in the carbonates. A manometer is used to measure the pressure.

**Pretreatment and Dispersion Techniques:** In the standard PSDA, an air-dry soil sample is pretreated to remove organic matter and soluble salts (3A1a1a). Additional non-routine chemical pretreatments are available for the removal of cementing agents that commonly prevent complete dispersion. These pretreatments may be requested by the project coordinator as follows:

**Carbonate Removal (3A1a2a):** Carbonates are destroyed with a 1 N NaOAc solution buffered to pH 5. The NaOAc solution is added to the sample until carbonate bubbles no longer evolve. The NaOAc solution is then washed from the sample. After destruction of carbonates, the remainder of method 3A1a1 is followed.

**Iron Removal (3A1a3a):** Soil samples are pretreated with H₂O₂ to remove organic matter. Iron oxides are removed with bicarbonate-buffered, sodium dithionite-citrate solution and heated until the sample changes to a grayish color. The suspension is flocculated with saturated NaCl solution and filtered to remove soluble salts. After removal of iron oxides, the remainder of method 3A1a1 is followed.

**Silica Removal (3A1a4a):** Soils are pretreated with H₂O₂ to remove organic matter. Soils with Si cementation or coatings are pretreated with a weak NaOH solution overnight. After removal of siliceous cementing agents, the remainder of method 3A1a1 is followed.

**Ultrasonic Dispersion (3A1a5a):** A soil sample is pretreated to remove organic matter and soluble salts. The sample is dried in an oven and weighed to obtain the initial weight. Sodium hexametaphosphate solution is added to the sample, which is then made to 100-mL volume with RO water. The sample is subjected to ultrasonic vibration for 5 min. After dispersion with the ultrasonic probe, method 3A1a1 is followed.

**Water Dispersible PSDA (3A1a6a):** Water dispersible particle-size distribution analysis may also be determined from a soil suspension without the removal of organic matter or soluble salts or without the use of a chemical dispersant. Upon omitting these procedural steps, the remainder of method 3A1a1 is followed.

**Gypseous PSDA (3A1a7):** Clay percentages are determined using the standard KSSL PSDA in which gypsum is removed. These clay percentages are then recalculated on a <2-mm “gypsum-intact” basis. Sand fractions are determined on a separate 10-g sample using sonication in 50 mL 70/30 ethanol-aqueous solution. Sands are wet-sieved through a 300-mesh screen (0.047-mm opening), rinsed with 800 mL 70/30 ethanol-aqueous solution, and collected. Sands are then dried at 35 ºC, dry sieved, and weighed by fraction. Silt percentages are calculated as the difference between 100% and the combined recalculated clay plus sand percentages.
**KSSL Field-moist PSDA**: For soils that irreversibly harden when dried, moist particle-size analysis (method 3A1a1b) may be requested by project coordinator. This procedure requires two 10-g samples of <2-mm, moist soil to be pretreated for removal of organic matter and soluble salts. One sample is dried in the oven to obtain the oven-dry weight, and the other sample is further processed as outlined in method 3A1a1a. Similarly to an air-dry sample, a field-moist soil may also be subjected to special pretreatment and dispersion techniques (3A1a1-6b). Both the air-dry and moist PSDA data are determined as percent of the <2-mm fraction on an oven-dry basis.

3. Interferences

**Standard KSSL PSDA**: The sedimentation equation that is used to measure the settling rates of particles of different sizes is as follows:

$$v = 2r^2g(\rho_s - \rho_l)/(9\eta)$$

where:

- $v$ = Velocity of fall
- $r$ = Particle radius
- $g$ = Acceleration due to gravity
- $\rho_s$ = Particle density
- $\rho_l$ = Liquid density
- $\eta$ = Fluid viscosity

This formula results from an application of Stokes' law and is referred to as Stokes' law. Assumptions used in applying Stokes' law to soil sedimentation measurements are as follows:

1. Terminal velocity is attained as soon as settling begins.
2. Settling and resistance are entirely due to the viscosity of the fluid.
3. Particles are smooth and spherical.
4. There is no interaction between individual particles in the solution (Gee and Bauder, 1986; Gee and Or, 2002).

Since soil particles are not smooth and spherical, the radius of the particle is considered an equivalent rather than an actual radius. In this method, particle density is assumed to be 2.65 g cc$^{-1}$.

Gypsum interferes with PSDA by causing flocculation of particles. Gypsum is removed by stirring and washing the soil with reverse osmosis water. This procedure is effective if the soil contains <25% gypsum.

Partial flocculation may occur in some soils if excess $H_2O_2$ is not removed from the soil after its use in organic matter oxidation.

Treatment of micaceous soils with $H_2O_2$ causes exfoliation of the mica plates and a matting of particles when dried in the oven. Since exfoliation occurs in
these soils, a true measurement of fractions is uncertain (Drosdoff and Miles, 1938).

**Fine-Clay Determination:** In the fine-clay determination, the distance from the center of rotation to the surface of the suspension must be constant for each centrifuge bottle. The particle density ($\rho_p$) of the fine clay is assumed to be 2.5 g cc$^{-1}$ (Jackson, 1969). The suspension temperature must be known to enter the correct liquid viscosity in the equation. Position the bottle under pipette without sudden movement of the centrifuge rotor, which causes disturbance of solution. The withdrawal rate with pipette should be constant.

**Carbonate-Clay Determination:** The carbonate-clay analysis is semiquantitative. It is assumed that all of the carbonates are converted to CO$_2$. This method measures all forms of carbonates. In addition to calcium carbonate, the carbonates of Mg, Na, and K also react with the acid. Analytical interferences may be caused by temperature changes within the reaction vessel. The analyst should not touch the glass of the vessel when reading the pressure. When sealing the vessel, the analyst should not hold onto the vessel any longer than necessary to tighten the cap. The internal pressure must be equalized with the atmosphere. Approximately 3 to 5 s are required to equalize the internal pressure of the bottle when piercing the septa with a needle. The analyst should replace septa and O rings at regular intervals because they develop leaks after extensive use.

**Pretreatment and Dispersion Techniques:** The PSDA results are dependent on the pretreatments used to disperse the soil. The presence of cementing agents, such as carbonates, Fe, and Si, often prevent complete dispersion. In these cases, special pretreatment and dispersion procedures may be performed upon request on either an air-dry or field-moist sample. However, these special techniques in themselves may interfere with PSDA as follows:

**Carbonate Removal (3A1a2):** The removal of carbonates with 1 N NaOAc (pH 5) results in sample acidification. This pretreatment can destroy the primary mineral structure of clay (Gee and Bauder, 1986).

**Iron Removal (3A1a3):** If the temperature of the water bath exceeds 80 °C during Fe removal, elemental S can precipitate (Mehra and Jackson, 1960). This pretreatment can destroy primary mineral grains in the clay fraction (El-Swaify, 1980).

**Silica Removal (3A1a4):** The effects of Si removal with 0.1 N NaOH on the clay fraction and particle-size distribution are unknown.

**Ultrasonic Dispersion (3A1a5):** Ultrasonic dispersion has been reported to destroy primary soil particles. Watson (1971) summarized studies that reported the destruction of biotite and breakdown of microaggregates by ultrasonic dispersion. However, Saly (1967) reported that ultrasonic vibration did not cause the destruction of the clay crystalline lattice or the breakdown of primary grains. The samples ranged from sandy to clayey soils. The cementing agents represented humus, carbonates, and hydroxides of Fe and Al. No standard procedures have been adopted using ultrasonic dispersion.
Field-Moist PSDA: Soils that irreversibly harden when dried are difficult to disperse. The PSDA for these soils can be determined on moist samples (method 3A1a1b) upon the request of the project coordinator.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when handling acid and \( \text{H}_2\text{O}_2 \). Mix acids in ventilated fume hoods. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Heat samples in ventilated fume hoods for removal of organic matter or cementing agents. Handle heated samples with leather gloves. Perform the transfer of acid to gelatin capsules near a sink in case of leakage or spills.

Users should be familiar with centrifuge operation. Opposite centrifuge bottles need to be balanced. Centrifuge should not be opened until centrifuge rotor has completely stopped.

5. Equipment

5.1 Fleakers, 300-mL, tared to 1 mg
5.2 Ceramic filter candles, .3 \( \mu \)m absolute retention (source currently unavailable)
5.3 Rack to hold ceramic filter candle and sample container
5.4 Mechanical shaker, horizontal, 120 oscillations min\(^{-1}\), 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.5 Cylinders, 1-L, white line fused onto glass at 1-L mark
5.6 Oven, 110 °C
5.7 Hot plate, 100 °C
5.8 Vacuum, 0.8 bars (80 kPa)
5.9 Thermometer, 0 to 150 °C
5.10 Desiccator
5.11 Motor driven stirrer, (Kilmer and Mullins, 1954)
5.12 Hand stirrer, perforated disk fastened to a rod
5.13 Adjustable pipette rack (Shaw, 1932; fig.3A1–3A3)
5.14 Lowy pipettes, 25-mL, with overflow bulb
5.15 Polyurethane foam, pipe insulation that fits snugly around cylinder
5.16 Sieve shaker with 12.7-mm (½ in) vertical and lateral movement at 500 oscillations min\(^{-1}\), accommodates a nest of 76-mm (3 in) sieves
5.17 Weighing bottles, 90-mL, with screw caps, tared to 1 mg
5.18 Weighing bottles, 90-mL, tared to 1 mg
5.19 Weighing bottles, 90-mL, tared to 0.1 mg
5.20 Drying dishes, aluminum
Figure 3A-1.—Shaw Pipette (Shaw, 1932).
5.21 Timer or clock with second hand
5.22 Electronic balance, ±0.10-mg sensitivity
5.23 Electronic balance, ±1.0-mg sensitivity
5.24 Watch glasses, 50- and 65-mm diameters
5.25 Evaporating dish, porcelain, 160-mm diameter, 31-mm height, with lip
5.26 Set of 76-mm (3 in) sieves, square weave phosphor bronze wire cloth, except 300 mesh, which is twilled weave. U.S. series and Tyler Screen Scale equivalent designations are as follows:

<table>
<thead>
<tr>
<th>Sand Size</th>
<th>Opening (mm)</th>
<th>U.S. No.</th>
<th>Tyler Mesh Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very coarse sand (VCS)</td>
<td>1.0</td>
<td>18</td>
<td>16</td>
</tr>
<tr>
<td>Coarse sand (CS)</td>
<td>0.5</td>
<td>35</td>
<td>32</td>
</tr>
<tr>
<td>Medium sand (MS)</td>
<td>0.25</td>
<td>60</td>
<td>60</td>
</tr>
<tr>
<td>Fine sand (FS)</td>
<td>0.105</td>
<td>140</td>
<td>150</td>
</tr>
<tr>
<td>Very fine sand (VFS)</td>
<td>0.047</td>
<td>300</td>
<td>300</td>
</tr>
</tbody>
</table>
5.27 Centrifuge, International No. 11, with No. 949 rotor head, International Equip. Co., Boston, MA
5.28 Centrifuge bottle, 500 mL
5.29 Torsion balance
5.30 Manometer, hand-held gauge and differential pressure, PCL-200A/C Series, Omega Engineering, Stamford, CT
5.31 Gelatin capsules, 5 mL
5.32 Threaded weighing bottles, 90 mL
5.33 Machined PVC caps for threaded 90-mL weighing bottles, 3.2-cm (1¼ in) diameter with 1.1-cm (7/16 in) diameter hole drilled in center, O-ring seal
5.34 O-rings, 3.2 x 38.1 mm (⅛ x 1½ in)
5.35 Septa, rubber, 7.9-mm (5/16 in) diameter. Place in machined cap.
5.36 Hypodermic needle, 25.4 mm (1 in), 23 gauge
5.37 Ultrasonic probe, 19-mm (¾ in) horn, 20 kHz, 600 watts

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Hydrogen peroxide \((H_2O_2)\), 30 to 35%
6.3 Sodium hexametaphosphate \((NaPO_3)_6\), reagent grade
6.4 Sodium carbonate \((Na_2CO_3)\), reagent grade
6.5 Sodium hexametaphosphate solution. Dissolve 35.7 g of \((NaPO_3)_6\) and 7.94 g of \(Na_2CO_3\) in 1 L of RO water. See Section 7.12 for standardization of sodium hexametaphosphate solution.
6.6 Ethyl alcohol
6.7 Calcium sulfate (anhydrous) or equivalent desiccant
6.8 Hydrochloric acid \((HCl)\), 6 \(N\), technical grade. Dilute 1 L of concentrated HCl with 1 L of RO water
6.9 Sodium carbonate \((Na_2CO_3)\) reagent. Dissolve 10.6 g \(Na_2CO_3\) in RO water and make to 1 L (10 mg CaCO_3)
6.10 1 \(N\) sodium acetate \((NaOAc)\) solution, buffered to pH 5. Dissolve 680 g of \(NaOAc\) in 4 L of RO water. Add ≈250 mL of acetic acid. Make to 5-L volume with RO water.
6.11 Sodium citrate solution, 0.3 \(M\) \(Na_3C_6H_5O_7\cdot2H_2O (88.4 \text{ g L}^{-1})\)
6.12 Sodium bicarbonate buffer solution, 1 \(M\) \(NaHCO_3 (84 \text{ g L}^{-1})\)
6.13 Sodium dithionite \((Na_2S_2O_4)\), hydrosulphite
6.14 Saturated NaCl solution (solubility at 20 °C; 360 g L\(^{-1}\))
6.15 Sodium hydroxide solution (NaOH), 0.1 \(N\). Dissolve 4 g NaOH pellets in 1 L of RO water.
6.16 Ethanol-aqueous solution, 70/30
7. Procedures

Pipette Analysis (3A1a)

7.1 If the standard air-dry PSDA procedure for the <2-mm fraction is requested, proceed to method 3A1a1a.

7.2 If PSDA is requested on a field-moist sample, proceed to method 3A1a1b.

7.3 If special pretreatment and dispersion techniques (removal of carbonate, Fe, or Si; ultrasonic dispersion; or water dispersible) are requested on air-dry soil samples, proceed to methods 3A1a2-6a, respectively.

7.4 If special pretreatment and dispersion techniques (removal of carbonate, Fe, or Si; ultrasonic dispersion; or water dispersible) are requested on field-moist soil samples, proceed to methods 3A1a2-6b, respectively.

Standard Pretreatments and Dispersion (3A1a1)

Air-Dry (3A1a1a)

7.5 Weigh 10 g of <2-mm, air-dry soil to nearest mg on an electronic balance and place into a numbered, tared, 300-mL fleaker. Wash and tare these fleakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.6 Add ≈50 mL of RO water and 5 to 7.5 mL of H$_2$O$_2$ to the soil sample at ambient temperature. Cover the soil sample with a 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place sample on hot plate. If the froth from the reaction exceeds the capacity of the fleaker, transfer the sample to a larger beaker.

7.7 Place the sample on a hot plate and heat to 90 °C. Add four 5- to 7.5-mL increments of H$_2$O$_2$ at 30-min intervals. If oxidation is incomplete, add additional H$_2$O$_2$ until organic matter oxidation is complete. Heat the sample for an additional 45 min to decompose excess H$_2$O$_2$. If the reaction is violent, do one or any combination of the following: (a) add small increments of ethyl alcohol to the sample, (b) remove the sample from the hot plate to slow the reaction, (c) transfer sample to a 1000-mL beaker, or (d) reduce the amount of H$_2$O$_2$ to sample. Record any unusual sample reactions.

7.8 Place the sample on the filter rack. Add 150 mL of RO water. Insert a filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Aspirate until liquid is removed and only slightly dampened sample remains. Wash the sample four additional times with ≈150 mL of RO water. Stir the sample with filter candle to ensure all soil particles will be rinsed. During aspiration, it may be necessary to occasionally apply back-pressure to filter candle and remove build-up of soil that inhibits aspiration. If the sample contains gypsum and flocculates, then the
following additional washings may be used. If the sample contains 1 to 5% gypsum, stir the sample with a magnetic stirrer for 5 min and wash five times with ≈250 mL of RO water each time. If the sample contains >5% gypsum, place the sample in a 1000-mL beaker and stir the sample with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.9 Place sample in oven. Dry the sample overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.10 Record the total weight (TW) of the sample to the nearest mg.

7.11 Add the exact volume of sodium hexametaphosphate solution (≈10 mL), equivalent to 0.4408 g of sodium hexametaphosphate, to each sample. Subtract the weight of the sodium hexametaphosphate (DW) that is contained in the extracted aliquot from the silt and clay weights to calculate silt and clay percentages. To determine the exact volume of sodium hexametaphosphate to add to each sample, refer to Section 7.12. Let stand until sample is completely moistened by sodium hexametaphosphate. Add ≈175 mL of RO water.

7.12 A sodium hexametaphosphate standardization is performed with each new batch of solution. Use only designated weighing bottles for standardization. Wash and tare these bottles after each standardization. Add duplicate aliquots (8.5, 9.0, 9.3, 9.6, 10.0, 10.3, 10.6, 11.0 mL) of sodium hexametaphosphate solution to numbered, tared, 90-mL weighing bottles. Oven-dry the aliquots overnight and record dry residue weight of sodium hexametaphosphate. Determine the exact volume of sodium hexametaphosphate solution to add to each sample by regressing the volume of sodium hexametaphosphate solution against the dry residue weight of sodium hexametaphosphate and then by predicting the volume needed to dispense 0.4408 g of sodium hexametaphosphate into each sample.

7.13 Place the sample in a horizontal shaker set at 120 oscillations min⁻¹ and shake for 15 h (overnight).

7.14 Remove the sample from the shaker and pour through a 300-mesh (0.047-mm) sieve mounted on a ring stand. The sample may need to be finger rubbed during transfer to speed washing. Place a funnel below the sieve and a 1-L cylinder below the funnel. Collect the silt and clay in the 1-L cylinder. Avoid using jets of water when washing the sample. Wash and rub all particles from the fleaker into the sieve. Continue to wash until the suspension volume in the cylinder is ≈800 mL. Sand and some of the coarse silt remain on the sieve. Rinse all <20-µm particles into the cylinder. Fill the cylinder to 1 L and cover with a 65-mm watch glass. Place pipe insulation around cylinders (sample and blank) to prevent rapid changes
in temperatures. Prepare a RO water blank to measure temperature fluctuations. Allow the cylinder to stand overnight to equilibrate the suspension with the room temperature. Wash the sand into an evaporation dish and dry the sand at 110 °C overnight.

7.15 Transfer the dried sand to a nest of sieves that has a top-to-bottom order of 1.0, 0.5, 0.25, 0.1, and 0.047 mm. Shake the sand for 3 min on a shaker that has 1.3-cm vertical and lateral movements and oscillates at 500 strokes min⁻¹. Record the weight of each separate sand fraction (SWᵢ) to the nearest mg. If optical analysis is requested, place the very fine sand and fine sand fractions in gelatin capsules and the remaining sand fractions in a labeled vial. Store capsules in the labeled vial. Wash sand dishes after every use.

7.16 Determine the percentage of fine silt and clay gravimetrically by using a Lowy, 25-mL pipette to remove an aliquot from the suspension in the 1-L cylinder. Periodically, gravimetrically calibrate the delivery volume of the pipette by weighing the amount of RO water dispensed from the pipette. Record the delivery volume (DV) and use the value to calculate the results. Regulate the vacuum such that the pipette fills in ≈12 s. Record temperature (T₁) of blank. Mount the pipette on an adjustable pipette rack (Shaw, 1932). Stir the silt and clay suspension with mechanical stirrer for at least 5 min. Place the cylinder on a stable, vibrationless table and stir with a hand stirrer in an up-and-down motion for 30 s. Timing is started upon completion of the stirring. Record the time that stirring is stopped. For the <20-µm fraction, slowly lower the closed pipette to a 10-cm depth in the suspension, turn on the vacuum, and withdraw an aliquot at the calculated time (table 1). Dispense the aliquot into a tared and numbered, 90-mL weighing bottle. Rinse the pipette twice with RO water and dispense into the tared weighing bottle with the aliquot. For the <2-µm fraction, pipette after 4.5, 5, 5.5, or 6.5 h. Record temperature (T₂) of blank. Use the average of T₁ and T₂ to adjust the pipette depth in the suspension as indicated in table 2. Repeat the procedure described for the <20-µm fraction. If determination of carbonate is required, use weighing bottle with screw threads. Dry the aliquots at 110 °C overnight and cool in a desiccator that contains calcium sulfate or an equivalent desiccant. Record the weight of the residue (RW) to the nearest 0.1 mg.

7.17 Use the 90-mL, round-bottomed weighing bottles for the <20-µm aliquots. Wash and tare after every fourth use. Use the 90-mL, square-bottomed weighing bottles for the <2-µm aliquots. Wash and tare after every use.

7.18 If optical mineralogy, fine-clay, and/or carbonate-clay determinations are not requested, the procedural aspects of the standard air-dry PSDA for the <2-mm fraction are complete. If optical mineralology is requested, save the sediment and proceed to the section on optical mineralogy. If fine-clay and/
or carbonate-clay determinations are requested, proceed to the sections on these analyses.

**Optical Mineralogy**

**7.19** If optical mineralogy is requested, decant the suspension and transfer the sediment to a 400-mL beaker. Fill the beaker to a height of 5.5 cm. Stir the sediment and allow to settle for 5 min. Discard the supernatant. Refill the beaker to a height of 5.5 cm. Stir again, allow to settle for 3 min, and then decant. Repeat the filling, stirring, 2-minute settling, and decanting until top half of suspension is clear. Transfer the sediment, which is dominantly 20 to 50 µm, to a labeled drying dish. Wash with ethanol, air-dry, and save in the drying dish for optical mineralogy.

**Fine Clay Determination (<0.2 µm)**

**7.20** Stir the silt and clay suspension with mechanical stirrer for 5 min. Remove sample from mechanical stirrer and place on table. Stir with the hand stirrer in an up-and-down motion for 30 s and allow the suspension to settle for 15 min.

**7.21** Pour the suspension into a centrifuge bottle and fill to the line marked on the bottle. The line marked on each bottle is 13 cm, which is the distance from the center of rotation to the surface of the suspension. Stopper and shake well to mix the suspension.

**7.22** Balance opposite centrifuge loads, which consist of centrifuge bottle, trunnion carrier, and bucket. Place loads on a torsion balance and add water to the lighter bucket until both loads weigh the same.

**7.23** Read the temperature of the suspension.

**7.24** Centrifuge at 1500 rpm.

**7.24.1** For methods 3A1a1-5a and 3A1a1-5b, vary the centrifuge time according to the temperature as follows (based on Stokes’ Law, \( s = 15 \text{ cm}, r = 18 \text{ cm}, N_m = 1500 \text{ rpm}, \text{ and } \rho_p = 2.5 \text{ g cm}^{-1}):

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>Viscosity ((\eta))</th>
<th>Delta-Density ((\Delta\rho))</th>
<th>Time (Min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18</td>
<td>0.01055</td>
<td>1.501</td>
<td>39.0</td>
</tr>
<tr>
<td>19</td>
<td>0.01029</td>
<td>1.501</td>
<td>38.0</td>
</tr>
<tr>
<td>20</td>
<td>0.01004</td>
<td>1.502</td>
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<td>(Δρ)</td>
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### 7.24.2
For methods, 3A1a6a and 3A1a6b, vary the centrifuge time according to the temperature as follows (based on Stokes’ Law, \( s = 15 \) cm, \( r = 18 \) cm, \( N_m = 1500 \) rpm, and \( \rho_p = 2.5 \) g cc\(^{-1}\)):

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### 7.25
After centrifuging, lower the pipette to a 3-cm depth in the suspension. Withdraw a 25-mL aliquot at a rate of ≈12 s. Avoid turbulence. Transfer the aliquot to a weighing bottle.
7.26 Place weighing bottle with aliquot in oven. Dry overnight at 110 °C. Remove sample from oven, place in desiccator with calcium sulfate or equivalent desiccant, and cool to ambient temperature.

7.27 Weigh residue weight (RW) to nearest 0.1 mg.

7.28 Use the 90-mL, round-bottomed weighing bottles for the <0.2 µm aliquots. Wash and tare after every fourth use.

**Carbonate Clay (<2 µm): Manometer Calibration**

7.29 Calibrate the manometer quarterly or whenever equipment changes. Calibrate by placing replicated aliquots of 0.0 to 20.0 mL of the Na$_2$CO$_3$ reagent into numbered, tared, 90-mL weighing bottles. Dry the standard samples in the oven overnight at 110 °C. Remove samples from oven, place in desiccator, and cool to ambient temperature. Record the weight of the standard samples to nearest 0.1 mg.

7.30 With a thin film of glycerin, lubricate the lip of the 90-mL weighing bottle that contains the Na$_2$CO$_3$. Dispense 3 mL of 6 N HCl into a gelatin capsule and place the top on the capsule. If HCl leaks from the capsule, discard the capsule. Place the capsule into the glass bottle and immediately cap the bottle. Release pressure in the bottle by piercing the septa with a hypodermic needle that is not connected to the manometer. Allow 3 to 5 s for internal pressure in bottle to equalize.

7.31 After the gelatin capsule has dissolved (several minutes), slowly tip and rotate the bottle to saturate the standard sample adhering to the sides of the bottle. Handle only the cap to avoid changing the temperature of the container. Allow sample to stand for at least 30 min.

7.32 Adjust the manometer to zero before taking measurements. Insert the hypodermic needle in the septa stopper, which is connected to the transducer. Measure the pressure inside the weighing bottle. Record the manometer readings (mm Hg) to the nearest whole number.

7.33 Calculate the linear regression equation, i.e., the dependent variable is the weight of Na$_2$CO$_3$ (regressed or predicted values) and the independent variable is the corresponding manometer readings.

**Carbonate Clay: Analysis**

7.34 Determine the presence of carbonates in <2-mm soil by placing soil on a spot plate and adding two or three drops of 1 N HCl. The rate of CO$_2$ evolution indicates the relative amount of carbonates (1B1b2b5).

7.35 If the soil contains more than a “slight” amount of carbonates, determine the amount of carbonate clay in the <2-µm dry residue. Use the 90-mL,
square-bottomed weighing bottles for the <2-µm aliquots and carbonate determination. Wash and tare after every use.

7.36 With a thin film of glycerin, lubricate the lip of the 90-mL weighing bottle that contains the <2-µm residue. In each analysis batch, include an empty weighing bottle as a blank. Dispense 3 mL of 6 N HCl into a gelatin capsule and place the top on the capsule. If HCl leaks from the capsule, discard the capsule. Place the capsule into the glass bottle and immediately cap the bottle. Release any pressure in the bottle by piercing the septa with a hypodermic needle that is not connected to the manometer. Approximately 3 to 5 s are required to equalize the internal pressure of the bottle.

7.37 After the gelatin capsule has dissolved (several minutes), slowly tip and rotate the bottle to saturate the clay adhering to the sides of the bottle. Handle only the cap to avoid changing the temperature of the container. Allow sample to stand for at least 30 min.

7.38 Adjust the manometer to zero before taking measurements. Insert the hypodermic needle in septa stopper which is connected to the transducer. Measure the pressure inside the weighing bottle and record the manometer readings (MR) to the nearest whole number (mm Hg). Begin readings with the blank (BR).

7.39 Compare the sample readings with those of a standard curve prepared by measuring CO₂ evolved from a series of Na₂CO₃ aliquots with a range 0 to 200 mg.

**Carbonate Removal (3A1a2)**

**Air-Dry (3A1a2a)**

7.40 Weigh sufficient sample to yield 10 g of <2-mm, air-dry, carbonate-free soil sample, e.g. if the sample contains 50% carbonates, weigh 20 g of soil. Place the <2-mm, air-dry sample into a 300-mL, tared fleaker.

7.41 Add ≈200 mL of the 1 N NaOAc solution to the sample, mix with a stirring rod, and cover with a watch glass. Allow the sample to stand overnight.

7.42 Place the sample on the hot plate and heat to ≈90 °C until bubbles are no longer evident. Do not boil. Heating accelerates reaction. Decant the solution and add more 1 N NaOAc solution. If a reaction occurs, repeat the heating procedure. Continue to decant, add NaOAc solution, and heat until all the carbonates are removed. The speed of dissolution can be increased by lowering the pH of the 1 N NaOAc solution (Rabenhorst and Wilding, 1984).

7.43 When no more carbonate bubbles are observed, insert the ceramic filter candle into the solution. Apply vacuum and candle the sample to dryness. Rinse once with 200 mL of RO water. Proceed to method 3A1a1a for remaining PSDA procedural steps.
Iron Removal (3A1a3)  
Air-Dry (3A1a3a)

7.44 A maximum of ≈0.5 g of Fe\textsubscript{2}O\textsubscript{3} can be dissolved in 40 mL of the citrate solution. Adjust the weight of the <2-mm, air−dry soil sample so that no single fleaker contains more than 0.5 g of Fe\textsubscript{2}O\textsubscript{3}. Split the sample into different fleakers if necessary. Total sample weight after dissolution should be ≈10 g. Place the sample into a tared, labeled, 300–mL fleaker.

7.45 Add ≈50 mL of RO water and 5 mL of H\textsubscript{2}O\textsubscript{2} to the soil sample at ambient temperature. Cover the soil sample with a 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place sample on hot plate. If the froth from the reaction exceeds the capacity of the fleaker, transfer the sample to a larger beaker.

7.46 Place the sample on a hot plate and heat to 90 °C. Add 5-mL increments of H\textsubscript{2}O\textsubscript{2} at 45-min intervals until oxidation has completed or until 30 mL of H\textsubscript{2}O\textsubscript{2} have been added. Heat the sample for an additional 45 min to decompose excess H\textsubscript{2}O\textsubscript{2}. If the reaction is violent, add small increments of ethyl alcohol to the sample or remove the sample from the hot plate to slow the reaction.

7.47 Add 40 mL of the citrate solution and 5 mL of the sodium bicarbonate. Heat to 80 °C in a water bath, but do not exceed 80 °C. Add 1 g of sodium dithionite powder with a calibrated scoop. Stir constantly with a glass rod for 1 min and then occasionally for 15 min. Add 10 mL of saturated NaCl solution and mix.

7.48 Centrifuge or candle the sample to remove the dissolved Fe\textsubscript{2}O\textsubscript{3}. Combine the split samples into fewer fleakers.

7.49 If the sample contains less than 0.5 g of Fe\textsubscript{2}O\textsubscript{3}, repeat the dissolution treatment. For samples that contain more than 0.5 g of Fe\textsubscript{2}O\textsubscript{3}, repeat the dissolution treatment two more times. Upon completion of Fe removal, proceed to method 3A1a1a for remaining PSDA procedural steps.

Silica Removal (3A1a4)  
Air-Dry (3A1a4a)

7.50 Weigh 10 g of <2-mm, air-dry soil to nearest mg on an electronic balance and place in a 300-mL, tared fleaker.

7.51 Add ≈50 mL of RO water and 5 mL of H\textsubscript{2}O\textsubscript{2} to soil sample at ambient temperature. Cover the soil sample with 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place sample on hot plate. If the froth from the reaction exceeds the capacity of the fleaker, transfer the sample to a larger beaker.
7.52 Place the sample on a hot plate and heat to 90 °C. Add 5-mL increments of \( \text{H}_2\text{O}_2 \) at 45-min intervals until oxidation has completed or until 30 mL of \( \text{H}_2\text{O}_2 \) have been added. Heat the sample for an additional 45 min to decompose excess \( \text{H}_2\text{O}_2 \). If the reaction is violent, add small increments of ethyl alcohol to the sample or remove the sample from the hot plate to slow the reaction.

7.53 Soak the sample overnight in 100 mL of 0.1 \( N \) NaOH. Upon removal of siliceous cementing agents, proceed to method 3A1a1a for remaining PSDA procedural steps.

**Ultrasonic Dispersion (3A1a5)**

**Air-Dry (3A1a5a)**

7.54 Weigh 10 g of <2-mm, air-dry soil to nearest mg on an electronic balance and place into a numbered, tared, 300-mL fleaker. Wash and tare these fleakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.55 Add ≈50 mL of RO water and 7.5 mL of \( \text{H}_2\text{O}_2 \) to the soil sample at ambient temperature. Cover the soil sample with a 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place sample on hot plate. If the froth from the reaction exceeds the capacity of the fleaker, transfer the sample to a larger beaker.

7.56 Place the sample on a hot plate and heat to 90 °C. Add four 5- to 7.5-mL increments of \( \text{H}_2\text{O}_2 \) at 30-min intervals. If oxidation is incomplete, add additional \( \text{H}_2\text{O}_2 \) until organic matter oxidation is complete. Heat the sample for an additional 45 min to decompose excess \( \text{H}_2\text{O}_2 \). If the reaction is violent, do one or any combination of the following: (a) add small increments of ethyl alcohol to the sample, (b) remove the sample from the hot plate to slow the reaction, (c) transfer sample to a 1000-mL beaker, or (d) reduce the amount of \( \text{H}_2\text{O}_2 \) to sample. Record any unusual sample reactions.

7.57 Place the sample on the filter rack. Add 150 mL of RO water. Insert a filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Aspirate until liquid is removed and only slightly dampened sample remains. Wash the sample four additional times with ≈150 mL of RO water. Stir the sample with filter candle to ensure all soil particles will be rinsed. During aspiration, it may be necessary to occasionally apply back-pressure to filter candle and remove build-up of soil that inhibits aspiration. If the sample contains gypsum and flocculates, then the following additional washings may be used. If the sample contains 1 to 5% gypsum, stir the sample with a magnetic stirrer for 5 min and wash five times with ≈250 mL of RO water each time. If the sample contains >5% gypsum, place the sample in a 1000-mL beaker and stir the sample with a...
magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.58 Place sample in oven. Dry the sample overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.59 Record the total weight (TW) of the sample to the nearest mg.

7.60 Add ≈100 mL of RO water and the exact volume of sodium hexametaphosphate solution (≈10 mL) equivalent to 0.4408 g of sodium hexametaphosphate to sample. Subtract the weight of the sodium hexametaphosphate (DW) that is contained in the extracted aliquot from the silt and clay weights to calculate silt and clay percentages. To determine the exact volume of sodium hexametaphosphate solution to add to each sample, refer to Section 7.61.

7.61 A sodium hexametaphosphate standardization is performed with each new batch of solution. Use only designated weighing bottles for standardization. Wash and tare these bottles after each standardization. Add duplicate aliquots (8.5, 9.0, 9.3, 9.6, 10.0, 10.3, 10.6, 11.0 mL) of sodium hexametaphosphate solution to numbered, tared, 90-mL weighing bottles. Oven-dry the aliquots overnight and record dry residue weight of sodium hexametaphosphate. Determine the exact volume of sodium hexametaphosphate solution to add to each sample by regressing the volume of sodium hexametaphosphate solution against the dry residue weight of sodium hexametaphosphate and then by predicting the volume needed to dispense 0.4408 g of sodium hexametaphosphate into each sample.

7.62 Disperse the suspension with ultrasonic vibrations. Ensure the power supply is properly tuned. Consult the instruction manual. Immerse the probe in the suspension to a 3-cm depth. Set timer to 5 min. Press start button. Adjust output control as required. Between samples, clean the probe by placing it in water or alcohol and energizing it for a few seconds.

7.63 After ultrasonic dispersion, add ≈75 mL RO water and pour the suspension through a 300-mesh (0.047-mm) sieve mounted on a ring stand. The sample may need to be finger rubbed during transfer to speed washing. Place a funnel below the sieve and a 1-L cylinder below the funnel. Collect the silt and clay in the 1-L cylinder. Avoid using jets of water in washing the sample. Wash and rub all particles from the fleaker into the sieve. Continue to wash until the suspension volume in the cylinder is ≈800 mL. Sand and some of the coarse silt remain on the sieve. Rinse all <20-µm particles into the cylinder. Fill the cylinder to 1 L and cover with a 65-mm watch glass. Place pipe insulation around cylinders (sample and blank) to prevent rapid changes in temperatures. Prepare a RO water blank to measure temperature fluctuations. Allow the cylinder to stand overnight.
to equilibrate the suspension with the room temperature. Wash the sand into an evaporation dish and dry the sand at 110 °C overnight. Proceed to method 3A1a1a for remaining PSDA procedural steps.

**Water Dispersion (3A1a6)
Air-Dry (3A1a6a)**

7.64 Weigh 10 g of <2-mm, air-dry soil to nearest mg on an electronic balance and place into a numbered, tared, 300-mL fleaker. Wash and tare these fleakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.65 Dry the sample in an oven at 110 °C overnight. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.66 Record the total weight (TW) of the sample to the nearest mg.

7.67 Add ≈175 mL of RO water to sample. Place the sample in a horizontal shaker set at 120 oscillations min\(^{-1}\) and shake for 15 h (overnight).

7.68 Remove the sample from the shaker and pour through a 300-mesh (0.047-mm) sieve mounted on a ring stand. The sample may need to be finger rubbed during transfer to speed washing. Place a funnel below the sieve and a 1-L cylinder below the funnel. Collect the silt and clay in the 1-L cylinder. Avoid using jets of water in washing the sample. Wash and rub all particles from the fleaker into the sieve. Continue to wash until the suspension volume in the cylinder is ≈800 mL. Sand and some of the coarse silt remain on the sieve. Rinse all <20-µm particles into the cylinder. Fill the cylinder to 1 L and cover with a 65-mm watch glass. Place pipe insulation around cylinders (sample and blank) to prevent rapid changes in temperatures. Prepare a RO water blank to measure temperature fluctuations. Allow the cylinder to stand overnight to equilibrate the suspension with the room temperature. Wash the sand into an evaporation dish and dry the sand at 110 °C overnight.

7.69 Transfer the dried sand to a nest of sieves that has a top-to-bottom order of 1.0, 0.5, 0.25, 0.1, and 0.047 mm. Shake the sand for 3 min on a shaker that has 1.3-cm vertical and lateral movements and oscillates at 500 strokes min\(^{-1}\). Record the weight of each separate sand fraction (SW\(_i\)) to the nearest mg. If optical analysis is requested, place the very fine sand and fine sand fractions in gelatin capsules and the remaining sand fractions in a labeled vial. Store capsules in the labeled vial. Wash sand dishes after every use.

7.70 Determine the percentage of fine silt and clay gravimetrically by using a Lowy, 25-mL pipette to remove an aliquot from the suspension in the 1-L cylinder. Periodically, gravimetrically calibrate the delivery volume of the pipette by weighing the amount of RO water dispensed from the pipette.
Record the delivery volume (DV) and use the value to calculate the results. Regulate the vacuum such that the pipette fills in ≈12 s. Record temperature ($T_1$) of blank. Mount the pipette on an adjustable pipette rack (Shaw, 1932). Stir the silt and clay suspension with mechanical stirrer for at least 5 min. Place the cylinder on a stable, vibrationless table and stir with a hand stirrer in an up-and-down motion for 30 s. Timing is started upon completion of the stirring. Record the time that stirring is stopped. For the <20-µm fraction, slowly lower the closed pipette to a 10-cm depth in the suspension, open the pipette, turn on the vacuum, and withdraw an aliquot at the calculated time (table 3). Dispense the aliquot into a tared and numbered, 90-mL weighing bottle. Rinse the pipette twice with RO water and dispense into the tared, weighing bottle with aliquot. For the <2-µm fraction, pipette after 4.5, 5, 5.5, or 6.5 h. Record temperature ($T_2$) of blank. Use the average of $T_1$ and $T_2$ and adjust the pipette depth in the suspension as indicated in table 4. Repeat the procedure described for the <20-µm fraction. Dry the aliquots at 110 °C overnight and cool in a desiccator that contains calcium sulfate or an equivalent desiccant. Record the residue weight (RW) to the nearest 0.1 mg. Proceed to method 3A1a1a for remaining PSDA procedural steps.

Gypseous Soils (3A1a7)

Air-Dry (3A1a7a)

7.71 Weigh two 10-g samples of air-dry soil. Weigh to nearest mg on an electronic balance. Place one 10-g sample into numbered, tared, 300-mL fleaker. Place the other 10-g sample into numbered, tared, 50-mm diameter beaker. Wash and tare these fleakers and beakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.72 For the 10-g sample placed in the 300-mL fleaker, proceed to method 3A1a1a for all PSDA procedural steps.

7.73 For the 10-g sample placed in the 50-mm diameter beaker, add 50 mL 70/30 ethanol-aqueous solution.

7.73.1 Allow solution to stand for 5 min.

7.73.2 Insert ultrasonic probe ≈20 mm below the surface of solution and apply energy for 5 min.

7.73.3 Pass solution through the 300-mesh screen (0.047-mm opening), rinsing with ≈800 mL 70/30 ethanol-aqueous solution. Collect sands.

7.73.4 Dry sands at 35°C.

7.73.5 Proceed to method 3A1a1a for remaining procedural steps to sieve and weigh sand fractions.
**Field-Moist (3A1a1b)**

7.74 Weigh enough <2 mm, moist soil to achieve two ≈10-g samples of air-dry soil. Weigh to nearest mg on an electronic balance and place into numbered, tared, 300-mL flasks. Wash and tare these flasks once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.75 Add ≈50 mL of RO water and 7.5 mL of H$_2$O$_2$ to both soil subsamples at ambient temperature. Cover the soil samples with a 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place sample on hot plate. If the froth from the reaction exceeds the capacity of the flask, transfer the sample to a larger beaker.

7.76 Place the samples on a hot plate and heat to 90 °C. Add four 5- to 7.5-mL increments of H$_2$O$_2$ at 30-min intervals. If oxidation is incomplete, add additional H$_2$O$_2$ until organic matter oxidation is complete. Heat the sample for an additional 45 min to decompose excess H$_2$O$_2$. If the reaction is violent, do one or any combination of the following: (a) add small increments of ethyl alcohol to the sample, (b) remove the sample from the hot plate to slow the reaction, (c) transfer sample to a 1000-mL beaker, or (d) reduce the amount of H$_2$O$_2$ to sample. Record any unusual sample reactions.

7.77 Place the sample on the filter rack. Add 150 mL of RO water. Insert a filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Aspirate until liquid is removed and only slightly dampened sample remains. Wash the sample four additional times with ≈150 mL of RO water. Stir the sample with filter candle to ensure all soil particles will be rinsed. During aspiration, it may be necessary to occasionally apply back-pressure to filter candle and remove build-up of soil that inhibits aspiration. If the sample contains gypsum and flocculates, then the following additional washings may be used. If the sample contains 1 to 5% gypsum, stir the sample with a magnetic stirrer for 5 min and wash five times with ≈250 mL of RO water each time. If the sample contains >5% gypsum, place the sample in a 1000-mL beaker and stir the sample with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.78 Place one of the samples in the oven and dry overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.79 Record the total weight (TW) of the H$_2$O$_2$-treated, oven-dry sample to the nearest mg. The H$_2$O$_2$-treated, oven-dry sample is not used in the remaining PSDA procedural steps. Proceed to method 3A1a1a and use the H$_2$O$_2$-treated sample that was not dried in oven for remaining PSDA procedural steps.
Carbonate Removal (3A1a2)
Field-Moist (3A1a2b)

7.80 Weigh two 10-g samples of <2-mm, moist soil to nearest mg on an electronic balance and place into 300-mL, tared flekers. Weigh sufficient samples to yield 10 g of <2-mm, moist, carbonate-free soil sample, e.g., if the sample contains 50% carbonates, weigh 20 g of soil.

7.81 Add ≈200 mL of the 1 N NaOAc solution to both samples, mix with a stirring rod, and cover with a watch glass. Allow the samples to stand overnight.

7.82 Place the samples on the hot plate and heat to ≈90 °C until bubbles are no longer evident. Do not boil. Heating accelerates reaction. Decant the solution and add more 1 N NaOAc solution. If a reaction occurs, repeat the heating procedure. Continue to decant, add NaOAc solution, and heat until all the carbonates are removed. The speed of dissolution can be increased by lowering the pH of the 1 N NaOAc solution (Rabenhorst and Wilding, 1984).

7.83 When no more carbonate bubbles are observed, insert the ceramic filter candle into the solution. Apply vacuum and candle the samples to dryness. Rinse once with 200 mL of RO water.

7.84 Add ≈50 mL of RO water and 5 mL of H₂O₂ to both soil samples at ambient temperature. Cover the soil samples with 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place samples on hot plate. If the froth from the reaction exceeds the capacity of the flekers, transfer the samples to larger beakers.

7.85 Place the samples on the hot plate and heat to ≈90 °C. Add 5-mL increments of H₂O₂ at 45-min intervals until oxidation has completed or until 30 mL of H₂O₂ have been added. Heat the sampled for an additional 45 min to decompose excess H₂O₂. If the reaction is violent, add small increments of ethyl alcohol to the samples or remove the samples from the hot plate to slow the reaction.

7.86 Place the samples on the filter rack. Add 150 mL of RO water. Insert filter candle, connect to the vacuum trap assembly with tubing, and turn on vacuum. Wash the samples four additional times with ≈150 mL of RO water. If the samples contain gypsum and flocculates, then the following additional washings may be used. If the samples contain 1 to 5% gypsum, stir the samples with a magnetic stirrer for 5 min and wash 5 times with ≈250 mL of RO water each time. If the samples contain >5% gypsum, stir the samples with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.87 Place one of the samples in the oven and dry overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.
7.88 Record the total weight (TW) of the H₂O₂-treated, oven-dry sample to the nearest mg. The H₂O₂-treated, oven-dry sample is not used in the remaining PSDA procedural steps. Proceed to method 3A1a1a and use the H₂O₂-treated sample that was not dried in oven for remaining PSDA procedural steps.

Iron Removal (3A1a3)
Field-Moist (3A1a3b)

7.89 Weigh two 10-g samples of <2-mm, moist soil to nearest mg on an electronic balance and place into 300-mL, tared flakers. As a maximum of ≈0.5 g of Fe₂O₃ can be dissolved in 40 mL of the citrate solution, adjust the weight of the <2-mm, moist soil sample so that no single flaker contains more than 0.5 g of Fe₂O₃. Split the sample into different flakers if necessary. Total sample weight after dissolution should be ≈10 g.

7.90 Add ≈50 mL of RO water and 5 mL of H₂O₂ to both soil samples at ambient temperature. Cover the soil samples with 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place samples on hot plate. If the froth from the reaction exceeds the capacity of the flaker, transfer the samples to larger beakers.

7.91 Place the samples on a hot plate and heat to 90 °C. Add 5-mL increments of H₂O₂ at 45-min intervals until oxidation has completed or until 30 mL of H₂O₂ have been added. Heat the samples for an additional 45 min to decompose excess H₂O₂. If the reaction is violent, add small increments of ethyl alcohol to the samples or remove the samples from the hot plate to slow the reaction.

7.92 Add 40 mL of the citrate solution and 5 mL of the sodium bicarbonate. Heat to 80 °C in a water bath, but do not exceed 80 °C. Add 1 g of sodium dithionite powder with a calibrated scoop. Stir constantly with a glass rod for 1 min and then occasionally for 15 min. Add 10 mL of saturated NaCl solution and mix.

7.93 Centrifuge or candle the samples to remove the dissolved Fe₂O₃. Combine the split samples into fewer flakers.

7.94 If the samples contain less than 0.5 g of Fe₂O₃, repeat the dissolution treatment. For samples with more than 0.5 g of Fe₂O₃, repeat the dissolution treatment two more times.

7.95 Place the samples on the filter rack. Add 150 mL of RO water. Insert a filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Wash the samples four additional times with ≈150 mL of RO water. If the samples contain gypsum and flocculates, then the following additional washings may be used. If the sample contains 1 to 5% gypsum, stir the sample with a magnetic stirrer for 5 min and wash five times with ≈250 mL of RO water each time. If the sample contains >5% gypsum, stir
the sample with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.96 Place one sample in the oven and dry overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.97 Record the total weight (TW) of the sample to the nearest mg. The H$_2$O$_2$ oven-dry sample is not used in the remaining PSA procedural steps. Proceed to method 3A1a1a and use the H$_2$O$_2$-treated sample that was not dried in oven for remaining PSDA procedural steps.

**Silica Removal (3A1a4)**

**Field-Moist (3A1a4b)**

7.98 Weigh two 10-g samples of <2-mm, moist soil to nearest mg on an electronic balance and place in 300-mL, tared flakers.

7.99 Add ≈50 mL of RO water and 5 mL of H$_2$O$_2$ to both soil samples at ambient temperature. Cover the soil samples with 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place samples on hot plate. If the froth from the reaction exceeds the capacity of the flaker, transfer the sample to larger beakers.

7.100 Place the samples on a hot plate and heat to 90 °C. Add 5-mL increments of H$_2$O$_2$ at 45-min intervals until oxidation has completed or until 30 mL of H$_2$O$_2$ have been added. Heat the samples for an additional 45 min to decompose excess H$_2$O$_2$. If the reaction is violent, add small increments of ethyl alcohol to the samples or remove the samples from the hot plate to slow the reaction.

7.101 Soak the samples overnight in 100 mL of 0.1 N NaOH.

7.102 Place the samples on the filter rack. Add 150 mL of RO water. Insert filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Wash the samples four additional times with ≈150 mL of RO water. If the sample contains gypsum and flocculates, then the following additional washings may be used. If the samples contain 1 to 5% gypsum, stir the samples with a magnetic stirrer for 5 min and wash 5 times with ≈250 mL of RO water each time. If the samples contain >5% gypsum, stir the sample with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.103 Place one sample in the oven and dry overnight at 110 °C. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.104 Record the total weight (TW) of the sample to the nearest mg. The H$_2$O$_2$-treated, oven-dry sample is only used for calculation of results and is not used in the remaining PSDA procedural steps. Proceed to method 3A1a1a
and use the H$_2$O$_2$-treated sample that was not dried in oven for all of the remaining PSDA procedural steps.

**Ultrasonic Dispersion (3A1a5)**

**Field-Moist (3A1a5b)**

7.105 Weigh enough <2 mm, moist soil to achieve two ≈10-g samples of air-dry soil. Weigh to nearest mg on an electronic balance and place into numbered, tared, 300-mL fleakers. Wash and tare these fleakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.106 Add ≈50 mL of RO water and 7.5 mL of H$_2$O$_2$ to the soil samples at ambient temperature. Cover the soil samples with 50-mm watch glass. Allow initial oxidation of organic matter to complete and then place samples on hot plate. If the froth from the reaction exceeds the capacity of the fleaker, transfer the samples to a larger beaker.

7.107 Place the samples on a hot plate and heat to 90 °C. Add four 5- to 7.5-mL increments of H$_2$O$_2$ at 30-min intervals. If oxidation is incomplete, add additional H$_2$O$_2$ until organic matter oxidation is complete. Heat the sample for an additional 45 min to decompose excess H$_2$O$_2$. If the reaction is violent, do one or any combination of the following: (a) add small increments of ethyl alcohol to the sample, (b) remove the sample from the hot plate to slow the reaction, (c) transfer sample to a 1000-mL beaker, or (d) reduce the amount of H$_2$O$_2$ to sample. Record any unusual sample reactions.

7.108 Place the sample on the filter rack. Add 150 mL of RO water. Insert a filter candle, connect to the vacuum trap assembly with tubing, and turn on the vacuum. Aspirate until liquid is removed and only slightly dampened sample remains. Wash the sample four additional times with ≈150 mL of RO water. Stir the sample with filter candle to ensure all soil particles will be rinsed. During aspiration, it may be necessary to occasionally apply back-pressure to filter candle and remove build-up of soil that inhibits aspiration. If the sample contains gypsum and flocculates, then the following additional washings may be used. If the sample contains 1 to 5% gypsum, stir the sample with a magnetic stirrer for 5 min and wash five times with ≈250 mL of RO water each time. If the sample contains >5% gypsum, place the sample in a 1000-mL beaker and stir the sample with a magnetic stirrer for 5 min then wash five times with ≈750 mL of RO water each time to remove soluble gypsum.

7.109 Place one sample in oven and dry overnight at 110 °C. Remove sample from oven, place in a desiccator, and cool to ambient temperature.

7.110 Record the total weight (TW) of the H$_2$O$_2$-treated, oven-dry sample to the nearest mg. The H$_2$O$_2$-treated, oven-dry sample is only used for the calculation of results and is not used in the remaining PSDA procedural
steps. Use the \( \text{H}_2\text{O}_2 \)-treated sample that was not dried in oven for the following PSDA procedural steps.

7.111 Add \( \approx100 \text{ mL} \) of RO water and the exact volume of sodium hexametaphosphate solution (\( \approx10 \text{ mL} \)) equivalent to 0.4408 g of sodium hexametaphosphate to sample. Subtract the weight of the sodium hexametaphosphate (DW) that is contained in the extracted aliquot from the silt and clay weights to calculate silt and clay percentages. To determine the exact volume of sodium hexametaphosphate to add to each sample refer to Section 7.112.

7.112 A sodium hexametaphosphate standardization is performed with each new batch of solution. Use only designated weighing bottles for standardization. Wash and tare these bottles after each standardization. Add duplicate aliquots (8.5, 9.0, 9.3, 9.6, 10.0, 10.3, 10.6, 11.0 mL) of sodium hexametaphosphate solution to numbered tared, 90-mL weighing bottles. Oven-dry the aliquots overnight and record dry residue weight of sodium hexametaphosphate. Determine the exact volume of sodium hexametaphosphate solution to add to each sample by regressing the volume of sodium hexametaphosphate solution against the dry residue weight of sodium hexametaphosphate and then by predicting the volume needed to dispense 0.4408 g of sodium hexametaphosphate into each sample.

7.113 Disperse the suspension with ultrasonic vibrations. Ensure the power supply is properly tuned. Consult the instruction manual. Immerse the probe in the suspension to a 3-cm depth. Set timer to 5 min. Press start button. Adjust output control as required. Between samples, clean the probe by placing it in water or alcohol and energizing it for a few seconds.

7.114 After ultrasonic dispersion, add \( \approx75 \text{ mL} \) RO water and pour the suspension through a 300-mesh (0.047-mm) sieve mounted on a ring stand. The sample may need to be finger rubbed during transfer to speed washing. Place a funnel below the sieve and a 1-L cylinder below the funnel. Collect the silt and clay in the 1-L cylinder. Avoid using jets of water in washing the sample. Wash and rub all particles from the fleaker into the sieve. Continue to wash until the suspension volume in the cylinder is \( \approx800 \text{ mL} \). Sand and some of the coarse silt remain on the sieve. Rinse all <20-\( \mu \text{m} \) particles into the cylinder. Fill the cylinder to 1 L and cover with a 65-mm watch glass. Place pipe insulation around cylinders (sample and blank) to prevent rapid changes in temperatures. Prepare a RO water blank to measure temperature fluctuations. Allow the cylinder to stand overnight to equilibrate the suspension with the room temperature. Wash the sand into an evaporation dish and dry the sand at 110 °C overnight. Proceed to method 3A1a1a for remaining PSDA procedural steps.
Water Dispersion (3A1a6)
Field-Moist (3A1a6b)

7.115 Weigh enough <2 mm, moist soil to achieve two ≈10-g samples of air-dry soil. Weigh to nearest mg on an electronic balance and place into numbered, tared, 300-mL fleakers. Wash and tare these fleakers once every 2 months. A quality control sample is included in each batch (≤24 samples).

7.116 Dry one sample in an oven at 110 °C overnight. Remove the sample from the oven, place in a desiccator, and cool to ambient temperature.

7.117 Record the total weight (TW) of the sample to the nearest mg. The oven-dry sample is not used in the remaining PSA procedural steps.

7.118 Add ≈175 mL of RO water to sample that was not dried in the oven. Place sample in a horizontal shaker set at 120 oscillations min\(^{-1}\) and shake for 15 h (overnight).

7.119 Remove the sample from the shaker and pour through a 300-mesh (0.047-mm) sieve mounted on a ring stand. The sample may need to be finger rubbed during transfer to speed washing. Place a funnel below the sieve and a 1-L cylinder below the funnel. Collect the silt and clay in the 1-L cylinder. Avoid using jets of water in washing the sample. Wash and rub all particles from the fleaker into the sieve. Continue to wash until the suspension volume in the cylinder is ≈800 mL. Sand and some of the coarse silt remain on the sieve. Rinse all <20-µm particles into the cylinder. Fill the cylinder to 1 L and cover with a 65-mm watch glass. Place pipe insulation around cylinders (sample and blank) to prevent rapid changes in temperatures. Prepare a RO water blank to measure temperature fluctuations. Allow the cylinder to stand overnight to equilibrate the suspension with the room temperature. Wash the sand into an evaporation dish and dry the sand at 110 °C overnight.

7.120 Transfer the dried sand to a nest of sieves that has a top-to-bottom order of 1.0, 0.5, 0.25, 0.1, and 0.047 mm. Shake the sand for 3 min on a shaker that has 1.3-cm vertical and lateral movements and oscillates at 500 strokes min\(^{-1}\). Record the weight of each separate sand fraction (SW\(_i\)) to the nearest mg. If optical analysis is requested, place the very fine sand and fine sand fractions in gelatin capsules and the remaining sand fractions in a labeled vial. Store capsules in the labeled vial. Wash sand dishes after every use.

7.121 Determine the percentage of fine silt and clay gravimetrically by using a Lowy, 25-mL pipette to remove an aliquot from the suspension in the 1-L cylinder. Periodically, gravimetrically calibrate the delivery volume of the pipette by weighing the amount of RO water dispensed from the pipette. Record the delivery volume (DV) and use the value to calculate the
results. Regulate the vacuum such that the pipette fills in ≈12 s. Record temperature \( T_1 \) of blank. Mount the pipette on an adjustable pipette rack (Shaw, 1932). Stir the silt and clay suspension with mechanical stirrer for at least 5 min. Place the cylinder on a stable, vibrationless table and stir with a hand stirrer in an up-and-down motion for 30 s. Timing is started upon completion of the stirring. Record the time that stirring is stopped. For the <20-µm fraction, slowly lower the closed pipette to a 10-cm depth in the suspension, turn on the vacuum, open the pipette, and withdraw an aliquot at the calculated time (table 1). Place the aliquot into a tared and numbered, 90-mL weighing bottle. Rinse the pipette with RO water into the tared weighing bottle. For the <2-µm fraction, pipette after 4.5, 5, 5.5, or 6.5 h. Record the temperature \( T_2 \) of blank. Use the average of \( T_1 \) and \( T_2 \) and adjust the pipette depth in the suspension as indicated in table 2. Repeat the procedure described for the <20-µm fraction. Dry the aliquots at 110 °C overnight and cool in a desiccator that contains calcium sulfate or an equivalent desiccant. Record the residue weight (RW) to the nearest 0.1 mg. Proceed to method 3A1a1a for remaining PSDA procedural steps. Also proceed to method 3A1a1a for determination of fine-clay and/or carbonate clay.

8. Calculations

Use calculations in sections 8.1–8.6 for all PSDA methods (3A1a1-5a and 3A1a1-5b), except water dispersible PSDA and gypsiferous/gypseous soils PSDA. Use calculations in sections 8.7–8.10 for water dispersible PSDA (3A1a6a and 3A1a6b). Use calculations in sections 8.11–8.14 for gypseous soils PSDA (3A1a7a).

Calculations 8.1–8.6 for Methods 3A1a1-5a and 3A1a1-5b

8.1 Clay % = \( \frac{100 \times [(RW_2 - DW) \times (CF / TW)]}{RW_2} \)

where:
- \( RW_2 \) = Residue weight (g), <2-µm fraction
- \( DW \) = Dispersing agent weight (g) = \( \frac{0.4408}{CF} \)
- \( CF = 1000 \text{ mL/DV} \)
- \( DV = \text{Dispensed pipette volume} \)
- \( TW = \text{Total weight (g), H}_2\text{O}_2\text{-treated, oven-dry sample} \)

8.2 Fine Silt % = \( \frac{100 \times [(RW_{20} - DW) \times (CF / TW)]}{RW_2} - \text{Clay %} \)

where:
- \( RW_{20} \) = Residue weight (g) of <20-µm fraction

8.3 Sand % = \( \sum (SW_i / TW) \times 100 \)

where:
- \( SW_i \) = Weight of sand fractions (1.0, 0.5, 0.25, 0.1, and 0.047 mm)
8.4 Coarse silt % = 100 − (Clay % + Fine Silt % + Sand %)

8.5 Fine Clay (%) = 100 x [(RW − DW) x (CF / TW)]

where:
- RW = Residue weight (g) of <0.2-μm fraction
- DW = Dispersing agent weight (g) = (0.4364 / CF)
- CF = 1000 mL/DV
- DV = Dispensed pipette volume
- TW = Total weight of H₂O₂-treated, oven-dry sample

8.6 Calculate carbonate clay percentage as follows:

8.6.1 Correct the manometer reading as follows:
- CR = (MR − BR)

where:
- CR = Corrected reading
- MR = Manometer reading
- BR = Blank reading. Three blanks are run with each batch (≤24 samples). The average of three blanks is used as BR.

8.6.2 Calculate two regression equations, i.e., one for corrected manometer readings <100 and another for corrected readings ≥100. Use the Na₂CO₃ weights as the dependent variable (regressed or predicted values) and the corresponding manometer readings as the independent variable.

8.6.3 Use the corrected (CR) linear regression equations to estimate the g of CaCO₃ in the sample.

8.6.4 Carbonate Clay Equivalent (<2 μm) (%) = [g CaCO₃ x 100 x CF] / TW

where:
- CF = 1000 mL/dispensed pipette volume (mL)
- TW = Total weight of H₂O₂-treated oven-dry sample

8.6.5 Noncarbonate Clay (<2 μm) (%) = Total Clay (%) − Carbonate Clay Equivalent (%)

Calculations 8.7–8.10 for Methods 3A1a6a and 3A1a6b

8.7 Clay % = 100 x [(RW₂ x CF) / TW]

where:
- RW₂ = Residue weight (g), <2-μm fraction
- CF = 1000 mL/DV
DV = Dispensed pipette volume
TW = Total weight (g), oven-dry sample

8.8 Fine Silt % = [100 x RW_{20} x CF] / TW - Clay %
   where:
   RW_{20} = Residue weight (g) of <20-µm fraction

8.9 Sand % = \sum (SW_i / TW) x 100
   where:
   SW_i = Weight of sand fractions (1.0, 0.5, 0.25, 0.1, and 0.047 mm)

8.10 Coarse silt % = 100 - (Clay % + Fine Silt % + Sand %)

Calculations 8.11–8.14 for Method 3A1a7a

8.11 Clay_{GR} % = 100 x [(RW_2 - DW) x (CF / TW)]
   where:
   Clay_{GR} = Clay (%) with gypsum removed
   RW_2 = Residue weight (g), <2-µm fraction
   DW = Dispersing agent weight (g) = (0.4408 / CF)
   CF = 1000 mL / DV
   DV = Dispensed pipette volume
   TW = Total weight (g), H_2O_2-treated, oven-dry sample

8.12 Clay_{GI} % = Clay_{GRA1} x Clay_{GRA2}
   where:
   Clay_{GI} = Clay (%) with gypsum intact
   Clay_{GRA1} = Clay with gypsum removed, adjusted to g clay/100 g insoluble soil basis
   Clay_{GRA2} = Clay with gypsum removed, adjusted to g insoluble soil/10 g <2-mm intact soil basis

8.13 Sand % = \sum (SW_i / TW) x 100
   where:
   SW_i = Weight of sand fractions (1.0, 0.5, 0.25, 0.1, and 0.047 mm)

8.14 Total Silt % = 100 - (Total Clay_{GI} % + Total Sand %)

9. Report
   Report each particle-size fraction to the nearest 0.1 percent.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.
11. References


Table 1.—Sampling times at 10-cm depth, 0.4408 g L\(^{-1}\) NaHMP solution, and 2.65 g cc\(^{-1}\) particle density.\(^1\)

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1 Use this table with methods 3A1a1-5a and 3A1a1-5b.
Table 2.—Sampling depths (cm) for 2-µm clay, 0.4408 g L⁻¹ NaHMP solution, and 2.65 g cc⁻¹ particle density.¹

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Table 2.—Sampling depths (cm) for 2-µm clay, 0.4408 g L⁻¹ NaHMP solution, and 2.65 g cc⁻¹ particle density.—Continued

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¹ Use this table with methods 3A1a1-5a and 3A1a1-5b.
Table 3.—Sampling times at 10-cm depth and 2.65 g cc\(^{-1}\) particle density.\(^1\)

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\(^1\) Use this table with methods 3A1a6a and 3A1a6b.
Table 4.—Sampling depths (cm) for 2-µm clay and 2.65 g cc⁻¹ particle density.¹

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Table 4.—Sampling depths (cm) for 2-µm clay and 2.65 g cc\(^{-1}\) particle density.—Continued

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\(^{1}\) Use this table with methods 3A1a6a and 3A1a6b.

Particle-Size Distribution Analysis (3A)
Particles >2 mm (3A2)

Rock and pararock fragments are defined as particles >2 mm in diameter and include all particles with horizontal dimensions less than the size of a pedon (Soil Survey Division Staff, 1993). Rock fragments are further defined as strongly cemented or more resistant to rupture. Pararock fragments are less cemented
than the strongly cemented class. Most of the pararock fragments are broken into particles 2 mm or less in diameter during the preparation of samples for particle-size analysis in the laboratory. Rock fragments are generally sieved and excluded from most chemical, physical, and mineralogical analyses. Exceptions are described in method 1B1b2f. It is necessary to know the amount of rock fragments for several applications, e.g., available water capacity and linear extensibility (Grossman and Reinsch, 2002).

The KSSL determines weight percentages of the >2-mm fractions by field and laboratory weight measurements by method 3A2a1. In the field or in the laboratory, the sieving and weighing of the >2-mm fraction are limited to the <75-mm fractions. In the field, fraction weights are usually recorded in pounds, whereas in the laboratory, fraction weights are recorded in grams. The 20- to 75-mm fraction is generally sieved, weighed, and discarded in the field. This is the preferred and usually most accurate method. Less accurately, the 20- to 75-mm fraction is estimated in the field as a volume percentage of the whole soil. If the fraction is sieved and weighed in the laboratory, the results are usually not reliable because of small sample size.

The KSSL estimates weight percentages of the >2-mm fractions from volume estimates of the >20-mm fractions and weight determinations of the <20-mm fractions by method 3A2a2. The volume estimates are visual field estimates. Weight percentages of the >20-mm fractions are calculated from field volume estimates of the 20- to 75-mm, 75- to 250-mm, and >250-mm fractions. The >250-mm fraction includes stones and boulders that have horizontal dimensions that are smaller than the size of the pedon. Weight measurements for the 2- to 20-mm fraction are laboratory measurements. Weight measurements of the 20- to 75-mm fractions in the field are more accurate than visual volume estimates. Weight measurements of this fraction in the laboratory are not reliable. The volume estimates that are determined in the field are converted to dry weight percentages. For any >2-mm fractions estimated by volume in the field, the KSSL calculates weight percentages by method 3A2b. The visual volume estimates of the >20-mm fraction are subjective. The conversion of a volume estimate to a weight estimate assumes a particle density of 2.65 g cc$^{-1}$ and a bulk density for the fine-earth fraction of 1.45 g cc$^{-1}$. Measured values can be substituted in this volume to weight conversion, if required.

Soil variability and sample size are interferences to weight determinations of the >2-mm particles. Enough soil material needs to be sieved and weighed to obtain statistically accurate rock fragment content. In order to accurately measure rock fragments with maximum particle diameters of 20 and 75 mm, the minimum dry specimen sizes that need to be sieved and weighed are 1.0 and 60.0 kg, respectively. Refer to ASTM method D 2488 (ASTM, 2012). Whenever possible, the field samples or “moist” material should have weights two to four times larger (ASTM, 2012). Therefore, sieving and weighing the 20- to 75-mm fraction
should be done in the field. The <20-mm fractions are sieved and weighed in the laboratory.

Procedures for reporting data for a size fraction base are outlined in Section 2C. Unless otherwise specified, the particle-size fractions 2 to 5, 5 to 20, 20 to 75, and 0.1 to 75 mm are reported on a <75-mm oven-dry weight percentage basis. The total >2-mm fraction is reported on a whole soil oven-dry weight percentage base.

Particle-Size Distribution Analysis (3A)
Particles >2 mm (3A2)
   Weight Estimates (3A2a)
      By Field and Laboratory Weighing (3A2a1)

1. Application
   Method 3A2a1 is used to determine weight percentages of the >2 mm fractions by field and laboratory weight measurements. The 20- to 75-mm fraction is generally sieved, weighed, and discarded in the field or is obtained from a field volume percentage estimate. However, the 20- to 75-mm fraction can be sieved and weighed in the laboratory. The <20-mm fractions are sieved and weighed in the laboratory.

2. Summary of Method
   Field weights are determined for the 20- to 75-mm fraction. This is the preferred method. When field determinations are not possible, weight measurements for the 20- to 75-mm fraction can be determined in the laboratory. The <20-mm fractions are sieved and weighed in the laboratory. The percentage of any 2- to 75-mm fraction on a <75-mm oven-dry weight basis is calculated.

3. Interferences
   Soil variability and sample size are interferences to weight determinations of the >2-mm particles. Enough soil material needs to be sieved and weighed to obtain statistically accurate rock fragment content. In order to accurately measure rock fragments with maximum particle diameters of 20 and 75 mm, the minimum specimen sizes (“dry” weights) that need to be sieved and weighed are 1.0 and 60.0 kg, respectively. Refer to ASTM method D 2488 (ASTM, 2012). Samples received in the laboratory generally have a maximum weight of 4 kg. Therefore, sieving and weighing the 20- to 75-mm fraction should be done in the field. The <20-mm fractions are sieved and weighed in the laboratory.

4. Safety
   Several hazards can be encountered in the field during sample collection. Examples include sharp-edged excavation tools, snake bites, and falls.
5. Equipment

5.1 Electronic balance, ±1-g sensitivity and 15-kg capacity
5.2 Trays, plastic, tared
5.3 Sieves, square-hole
   5.3.1 9 mesh, 2 mm
   5.3.2 4 mesh, 4.76 mm
   5.3.3 20 mm, ¾ in
   5.3.4 76 mm, 3 in
5.4 Mechanical shaker with 9-mesh and 4-mesh sieves
5.5 Rubber roller
5.6 Metal plate, 76 x 76 x 0.5 cm
5.7 Scale, 100-lb (45-kg) capacity
5.8 Brown Kraft paper

6. Reagents

6.1 Reverse osmosis (RO) water
6.2 1 N HCl. Dilute 83.3 mL of concentrated HCl in 1 L of RO water.
6.3 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate (NaPO₃)₆ and 7.94 g of sodium carbonate (Na₂CO₃) in 1 L of distilled water.

7. Procedure

Field

7.1 Sieve a representative horizon sample with a 76-mm sieve. Sieve about 60 kg of material to accurately measure rock fragments that have a maximum particle diameter of 75 mm. As a 60-kg sample may not be feasible because of limitations of time and/or soil material, actual sample size may be 30 or 40 kg. Discard the >75-mm material. Weigh and record weight (lbs) of <75-mm fraction. Sieve this material with a 20-mm sieve. Discard the 20- to 75-mm fraction. Weigh and record weight (lbs) of <20-mm fraction. Place a subsample of the <20-mm material in a plastic bag. Label and send to laboratory for analyses.

Laboratory

7.2 Distribute the field sample on a plastic tray, weigh, and record moist weight. Air-dry, weigh, and record weight.
7.3 Process air-dry material on a flat, metal plate that is covered with brown Kraft paper. Thoroughly mix material by moving the soil from the corners
to the middle of the processing area and then by redistributing the material. Repeat process four times. Roll material with wooden rolling pin to crush clods to pass a 2-mm sieve. For samples with easily crushed coarse fragments, substitute rubber roller for wooden rolling pin. Sieve clayey soils that contain many coarse fragments in the mechanical shaker. Roll and sieve until only the coarse fragments that do not slake in sodium hexametaphosphate solution remain on the sieve.

7.4 If more sample is received than is needed for processing, select a subsample for preparation. Weigh subsample and record weight.

7.5 Weigh soil material with diameters of 2 to 5 mm. Soak in sodium hexametaphosphate solution for 12 h. Air-dry, weigh the material that does not slake, and discard. Weigh, record weight, and discard coarse fragments with diameters of 20 to 75 mm and 5 to 20 mm. Most laboratory samples do not contain 20- to 75-mm fragments because this fraction is generally sieved, weighed, and discarded in the field.

8. Calculations

8.1 If field weight measurements are determined for the <75-mm and the 20- to 75-mm fraction, convert these weights in pounds to grams. If laboratory measurements are determined for the <75 mm and the 20- to 75-mm fractions, these weights are already in grams.

8.2 Determine field-moist weight of the subsample as received in the laboratory. Determine air-dry weight of subsample. Air-dry weight is defined as a constant sample weights obtained after drying at 30 ±5 °C (≈3 to 7 days).

8.3 Determine ratio of slaked, air-dried weight (g) to unslaked, air-dried weight (g) for the 2- to 5-mm fraction. Using this ratio, adjust weight of coarse fragments with <5-mm diameters.

8.4 Base coarse-fragment calculation on oven-dry weight. Use the AD/OD (air-dry/oven-dry ratio) (method 3D1) to calculate the oven-dry weight of <2-mm fraction. Use the following equation to determine the percentage of any 2- to 75-mm fraction on a <75-mm oven-dry weight-basis.

\[
\text{Percentage } > 2 \text{ mm fraction } (<75\text{-mm basis}) = \left( \frac{A}{B} \right) \times 100
\]

where:

\( A = \) Weight of 2- to 75-mm fraction (g)
\( B = \) Weight of <75-mm fraction (g)

8.5 Determine oven-dry weight by weighing the sample after oven-drying at 110 °C for 24 h or by calculating as follows:
Oven-dry weight (g) = [Air-dry weight (g)] / AD/OD

where:
AD/OD = Air-dry/oven-dry weight

8.6 Similarly, determine oven-dry weight from the field-moist weight of a sample by calculating as follows:
Oven-dry weight (g) = [Field-moist weight (g)] / [Field-moist weight (g) / Oven-dry weight (g)]

8.7 In calculations of the oven-dry weight percentages of the >2-mm fraction, make corrections for the field-water content of the <75-mm sample at sampling and for the water content of the air-dry bulk laboratory sample. Base the corrections for the field-water content on the difference between the field-moist weight and air-dry weight of the bulk sample.

9. Report

Field

9.1 Weight (lbs) of field-moist, <75-mm fraction
9.2 Weight (lbs) of field-moist, 20- to 75-mm fraction

Laboratory

9.3 Weight (g) of field-moist soil sample
9.4 Weight (g) of air-dry soil sample
9.5 Weight (g) of air-dry processed soil sample
9.6 Weight (g) of 20-to 75-mm fraction
9.7 Weight (g) of 5- to 20-mm fraction
9.8 Weight (g) of 2- to 5-mm fraction
9.9 Weight (g) of subsample 2- to 5-mm fraction before slaking
9.10 Weight (g) of subsample 2- to 5-mm fraction after slaking

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References

Particle-Size Distribution Analysis (3A)

Particulate >2 mm (3A2)
- Weight Estimates (3A2a)
  - From Volume and Weight Estimates (3A2a2)
- Volume Estimates (3A2b)

1. Application

Method 3A2a2 is used to determine weight percentages of the >2 mm fractions from volume estimates and weight determinations. The volume estimates are visual field estimates for any fractions that are >20 mm. The weight estimates are laboratory measurements for the 2-to 20-mm or 2- to 75-mm fractions. The volume estimates for any fractions that are >20 mm are converted to weight percentages. The total >2-mm fraction is reported on an oven-dry weight basis for whole soil. Method 3A2b is the calculations used to derive weight percentages from volume percentages of all the >2-mm material.

2. Summary of Method

Visual field-volume estimates are determined for any fractions that are >20 mm. These volume estimates include, if applicable, the 20- to 75-mm, 75- to 250-mm, and the >250-mm fractions. The >250-mm fraction includes stones and boulders that have horizontal dimensions that are less than those of the pedon. Instead of visual field-volume estimates, field weights for the 20- to 75-mm fraction may be determined. This is the preferred method. If these measurements are unavailable, visual field volume estimates of the 20- to 75-mm fraction are used rather than laboratory weights of this fraction. The <20-mm fractions are sieved and weighed in the laboratory.

3. Interferences

Soil variability and sample size are interferences to weight determinations of the >2-mm particles. Enough soil material needs to be sieved and weighed to obtain statistically accurate rock fragment content. In order to accurately measure rock fragments with maximum particle diameters of 20 and 75 mm, the minimum specimen sizes (“dry” weights) that need to be sieved and weighed are 1.0 and 60.0 kg, respectively. Refer to ASTM Standard Practice D 2488 (ASTM, 2012). Samples received in the laboratory generally have a maximum weight of 4 kg. Therefore, sieving and weighing the 20- to 75-mm fraction should be done in the field.
The visual volume estimates of the >75-mm fractions are subjective. The conversion of a volume estimate to a weight estimate assumes a particle density of 2.65 g cc\(^{-1}\) and a bulk density for the fine-earth fraction of 1.45 g cc\(^{-1}\). If particle density and bulk density measurements are available, they are used in the calculations.

4. Safety

Several hazards can be encountered in the field during sample collection. Examples include sharp-edged excavation tools, snake bites, and falls.

5. Equipment

5.1 Electronic balance, ±1-g sensitivity and 15-kg capacity
5.2 Trays, plastic, tared
5.3 Sieves, square-hole
   5.3.1 9 mesh, 2 mm
   5.3.2 4 mesh, 4.76 mm
   5.3.3 20 mm, ¾ in
   5.3.4 76 mm, 3 in
5.4 Mechanical shaker with 9-mesh and 4-mesh sieves
5.5 Rubber roller
5.6 Metal plate, 76 x 76 x 0.5 cm
5.7 Scale, 45-kg (100-lb) capacity
5.8 Brown Kraft paper

6. Reagents

6.1 Reverse osmosis (RO) water
6.2 1 \(N\) HCl. Dilute 83.3 mL of concentrated HCl in 1 L of distilled water.
6.3 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate (NaPO\(_3\))\(_6\) and 7.94 g of sodium carbonate (Na\(_2\)CO\(_3\)) in 1 L of distilled water.

7. Procedure

**Field**

7.1 Determine volume estimates as percentages of soil mass for the 75- to 250-mm and for the >250-mm fractions. The >250-mm fraction includes stones and boulders with horizontal dimensions less than those of the pedon.
7.2 Determine either weight measurements in pounds or visual field-volume estimates in percentages for the 20- to 75-mm fragments. Weight
measurements for the 20- to 75-mm fraction are the preferred method. However, volume estimates are more accurate than laboratory weights using small samples.

7.3 If field weight measurements are determined for the 20- to 75-mm fraction, sieve an entire horizon sample with a 76-mm sieve. Sieve ≈60 kg of material to accurately measure rock fragments that have a maximum particle diameter of 75 mm. A 60-kg sample may not be possible because of limitations of time and/or soil material. Actual sample size may be 30 or 40 kg. Discard the >75-mm fraction. Weigh and record weight of <75-mm fraction. Sieve this material with a 20-mm sieve. Discard the 20- to 75-mm fraction. Weigh and record weight of <20-mm fraction. Place a subsample of the <20-mm material in an 8-mil plastic bag. Label and send to laboratory for analyses.

Laboratory

7.4 Distribute the field sample on a plastic tray, weigh, and record moist weight. Air-dry, weigh, and record weight.

7.5 Process air-dry material on a flat, metal plate that is covered with brown Kraft paper. Thoroughly mix material by moving the soil from the corners to the middle of the processing area and then by redistributing the material. Repeat process four times. Roll material with wooden rolling pin to crush clods to pass a 2-mm sieve. For samples with easily crushed coarse fragments, substitute rubber roller for wooden rolling pin. Sieve clayey soils that contain many coarse fragments in the mechanical shaker. Roll and sieve until only the coarse fragments that do not slake in sodium hexametaphosphate solution remain on the sieve.

7.6 If more sample is received than is needed for processing, select subsample for preparation. Weigh subsample and record weight.

7.7 Weigh soil material with diameters of 2 to 5 mm. Soak in sodium hexametaphosphate solution for 12 h. Air-dry, weigh the material that does not slake, and discard. Weigh, record weight, and discard coarse fragments with diameters of 20 to 75 mm and 5 to 20 mm. Most laboratory samples do not contain 20- to 75-mm fragments as this fraction is generally weighed, sieved, and discarded in the field.

8. Calculations

From Volume and Weight Estimates (3A2a2)

8.1 Calculate weight percentages from volume percentages using measured bulk density ($D_{b/m}$) and particle density ($D_p$). If measurements are unavailable, assume a $D_{b/m}$ of 1.45 g cc$^{-1}$ and a $D_p$ of 2.65 g cc$^{-1}$.
8.2 Use the following equation to convert all volume estimates to weight percentages for specified fractions.

Percentage >2 mm (wt basis) = \[100 \frac{D_p (x)}{D_p (x) + D_{b\text{m}} (1-x)}\]

where:
\(D_p\) = Particle density (2.65 g cc\(^{-1}\), unless measured)
\(D_{b\text{m}}\) = Bulk density (1.45 g cc\(^{-1}\) for <2-mm fraction, unless measured)
\(x\) = [volume fragments >i mm]/[volume whole soil]

where:
i = size fraction above which volume estimates are made and below which weight percentages are determined, usually 20 or 75 mm in diameter

8.3 Use the preceding equation to calculate any individual fraction >j mm (j = any size fraction) by substituting an appropriate value of \(D_{b\text{m}}\) representing the fabric <j mm.

**Volume Estimates (3A2b)**

8.4 Use the following equation to determine the volume of the <2-mm fraction per unit volume of whole soil.

\[C_m = \frac{[\text{Volume moist <2-mm fabric}]/[\text{Volume moist whole-soil}]}{[D_{p}(1-y)(1-x)]/[D_{p}(1-y) + D_{b\text{m}}(y)]}\]

where:
\(C_m\) = Rock fragment conversion factor
Volume moist whole soil = Volume of fine earth + rock fragments on moist whole-soil basis
\(y\) = [weight material between 2 mm and i mm]/[weight material <i mm]

8.5 Use the following formula to convert laboratory data on a <2-mm weight basis to moist whole-soil volume basis.

\(C_m \times D_{b\text{m}} \times \text{lab datum}\)

8.6 Use the following formula to determine the volume percentage of <2-mm fabric in whole soil.

\(C_m \times 100\)

8.7 Use the following formula to determine the volume percentage of >2-mm fabric in whole soil.

\(100 (1-C_m)\)
8.8 Use the following formula to report weight of <2-mm fabric per unit volume of whole soil for some soils.

\[(Cm \times Db_m)\]

9. Report

Field

9.1 Volume (%) >250-mm fraction (includes stones and boulders with horizontal dimensions smaller than size of a pedon)
9.2 Volume (%) 75- to 250-mm fraction
9.3 Volume (%) 20- to 75-mm fraction (not needed if weighed in field)
9.4 Weight (lbs) <75-mm fraction
9.5 Weight (lbs) 20- to 75-mm fraction

Laboratory

9.6 Weight (g) of field moist soil sample
9.7 Weight (g) of air-dry soil sample
9.8 Weight (g) of air-dry processed soil sample
9.9 Weight (g) of 20-to 75-mm fraction
9.10 Weight (g) of 5- to 20-mm fraction
9.11 Weight (g) of 2- to 5-mm fraction
9.12 Weight (g) of subsample 2- to 5-mm fraction before slaking
9.13 Weight (g) of subsample 2- to 5-mm fraction after slaking

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Bulk Density (3B)

Density is defined as mass per unit volume. Soil bulk density of a sample is the ratio of the mass of solids to the total or bulk volume. This total volume includes the volume of both solids and pore space. Bulk density is distinguished from particle density, which is mass per unit volume of only the solid phase.
Particle density excludes pore spaces between particles. As bulk density (Db) is usually reported for the <2-mm soil fabric, the mass and volume of rock fragments are subtracted from the total mass and volume.

Bulk density may be highly dependent on soil conditions at the time of sampling. Changes in soil volume due to changes in water content alter bulk density. Soil mass remains fixed, but the volume of soil may change as water content changes (Blake and Hartge, 1986). Bulk density, as a soil characteristic, is actually a function rather than a single value. Subscripts are therefore added to the bulk density notation (Db) to designate the water state of the sample when the volume was measured. The KSSL uses bulk density notations of Db<sub>f</sub>, Db<sub>33</sub>, Db<sub>od</sub>, and Db<sub>r</sub> for field-state, 33-kPa equilibration, oven-dry, and rewet, respectively.

The field-state (Db<sub>f</sub>) value is the bulk density of a soil sample including the water content of the soil in the field at the time of sampling. The 33-kPa equilibration (Db<sub>33</sub>) value is the bulk density of a soil sample that has been desorbed to 33 kPa (⅓ bar). The oven-dry (Db<sub>od</sub>) value is the bulk density of a soil sample that has been dried in an oven at 110 °C. The rewet (Db<sub>r</sub>) value is the bulk density of soil sample that has been equilibrated, air-dried, and re-equilibrated. Db<sub>r</sub> is used to determine the irreversible shrinkage of soils and subsidence of organic soils. The determinations of these bulk density values, Db<sub>f</sub>, Db<sub>33</sub>, Db<sub>od</sub>, and Db<sub>r</sub>, are described in methods 3B1a, 3B1b, 3B1c, and 3B1d, respectively.

Bulk density may also be determined for field-moist soil cores of known volume (method 3B6a). The bulk density of a weak or loose soil material for which the clod or core method is unsuitable may be determined by the compliant cavity method (method 3B3a).

In general, there are two broad groupings of bulk density methods. One group is for soil materials coherent enough that a field-sample can be removed, and the other group is for soils that are too fragile to remove a sample and therefore an excavation operation must be performed. In the former group, there are clod methods in which the sample has an undefined volume and is coated and then the volume is determined by submergence. Also under the former there are various methods in which a cylinder of known volume is obtained of soil sufficiently coherent that it remains in the cylinder. The complete cylinder may be inserted (3B6a), or only part of the cylinder is inserted and the empty volume is subtracted from the total volume of the core (e.g., variable height method, Grossman and Reinsch, 2002). In the latter group, three excavation methods have been used to determine Db<sub>r</sub>, as follows: (1) compliant cavity (3B3a), (2) ring excavation (3B4a), and (3) frame excavation (3B5a) (Grossman and Reinsch, 2002). The frame-excavation provides for a larger sample area and is advantageous where there is large, very local variability, as in the O horizons of woodlands (Soil Survey Staff, 2014).

The complication concerning the difference between bulk density of the soil and bulk density of the sample is particularly important for the clod method as presented here because the method permits determination of the volume at
different water contents and hence volumes. If the water content is at or near field capacity, desiccation cracks are closed and the bulk density (Db33 or Dbf if field-water is near field capacity) of the soil and of the sample are considered the same. However, if the sample is at a water content below field-capacity because of drying after sampling or because the sample was taken below field-capacity, then desiccation cracks that occur in place are excluded from the soil and the bulk density of the sample exceeds that of the soil. If the sample is large and inclusive of the desiccation cracks, as in some excavation methods, then again the sample and soil bulk density are the same. The difference between the bulk density of the sample and that of the soil is particularly large for oven-dry clods (D_{oda}) of soils having high extensibility. This difference may be difficult to accurately determine for such soils if they are taken through a rewet cycle. Grossman and Reinsch (2002) discuss the manipulation of clod bulk densities (the sample) at water contents below field capacity to obtain an estimate of the soil bulk density at such water contents. Similarly, estimates of soil bulk density at intermediate field-water contents between field capacity and oven-dryness inclusive of desiccation crack space are discussed by Grossman et al. (1990).

References


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**Bulk Density (3B)**

**Saran-Coated Clods (3B1)**

**Field-State (Db) (3B1a)**

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Method 3B1a determines the bulk density value Db of a soil sample at field-soil water content at time of sampling. Field-state bulk density (Db,) offers the
opportunity to obtain bulk density information relatively cheaply without the expense incurred to obtain water retention data. $D_b$ is particularly useful if the soil layers are at or above field capacity and/or the soils have low extensibility and do not exhibit desiccation cracks even if below field capacity.

2. Summary of Method

Field-occurring fabric (clods) is collected from the face of an excavation. One coat of plastic lacquer is applied in the field. Additional coats of plastic lacquer are applied in the laboratory. In field-water state or after equilibration, the clod is weighed in air to measure its mass and in water to measure its volume. After the clod is dried in an oven at 110 °C, the mass and volume are determined again. A correction is made for the mass and volume of rock fragments and for plastic coatings (Brasher et al., 1966; Blake and Hartge, 1986; Grossman and Reinsch, 2002).

3. Interferences

Errors are caused by nonrepresentative samples. Only field-occurring fabric (clods) should be sampled. The whole-soil bulk density may be overestimated because sampled clods frequently exclude the crack space between clods (Grossman and Reinsch, 2002).

The penetration of plastic lacquer into the voids of sandy or organic soils interferes with the corrections for mass and volume of the plastic coat and with the accuracy of water content determinations. Penetration can be reduced by spraying water on the clod and then immediately dipping the clod in the plastic lacquer.

Loss of soil during the procedure voids the analyses because all calculations are based on the oven-dry soil mass. Holes in the plastic coating, which are detected by air bubbles escaping from submerged clod, introduce errors in volume measurement. An inadequate evaporation of the plastic solvent results in overestimation of the soil mass. A drying time of 1 h is usually sufficient for evaporation of solvent. However, clods with high organic matter content may need longer.

Bulk density is reported for <2-mm soil fabric. Correction for rock fragments >2-mm requires either knowledge or assumption of the rock fragment density. Errors in the estimation or measurement of rock fragment density affect the accuracy of the value for soil bulk density. Rock fragments may contain water, which complicates the application to actual water-holding capacity.

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a
lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Rigid shipping containers. The KSSL uses a corrugated box with compartments.
5.3 Plastic bags, 1-mil, 127 x 89 x 330 mm
5.4 Wire. The KSSL uses a 28-awg coated copper wire.
5.5 Hairnets
5.6 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.7 Hook assembly for weighing below balance
5.8 Plexiglas water tank
5.9 Lift apparatus, powered by compressed air, Hoff’s Machine & Welding, Inc.
5.10 Oven, 110 °C
5.11 Sieve, No. 10 (2-mm openings)
5.12 Rope, 3-m
5.13 Clothespins
5.14 Silt loam soil
5.15 Hot plate
5.16 Spray bottle

6. Reagents

6.1 Acetone (2-propanone; dimethyl ketone)
6.2 Water
6.3 Alcohol
6.4 Dow Saran F-310 resin, available from Dow Chemical Company
6.5 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a
wooden stick. In the laboratory, stir solvent with a non-sparking, high speed stirrer while slowly adding resin. Stir plastic lacquer for 15 min at 25 °C. Store plastic lacquer in covered plastic or steel containers.

**Procedure**

**Field**

7.1 Collect field-occurring clods, ≈100 to 200 cm$^3$ in volume (fist-sized), from the face of the excavation. Three clods per horizon are recommended. It is important that these clods be as representative of the bulk sample volume as possible. Remove a piece of soil larger than a clod from the face of sampling pit. From this piece, prepare a clod by gently cutting or breaking protruded peaks and compacted material from clod. If roots are present, trim roots with shears. No procedure for sampling clods is applicable to all soils. Adjust field-sampling techniques to meet field-conditions at time of sampling.

7.2 Make a clothesline by stretching a rope between two fixed points. Tie clod with fine copper wire or place clod in a hairnet. If clod is dry, moisten surface with a fine mist of water. Quickly dip entire clod into plastic lacquer (fig. 3B1a-1). Suspend clod from clothesline to dry (fig. 3B1a-2). Dry clod for 30 min or until odor of solvent dissipates. If the value of $D_{bf}$ is required, store clods in waterproof plastic bags as soon as coating dries because coating is permeable to water vapor.

7.3 Pack clods in rigid containers to protect them during transport.

**Laboratory**

7.4 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod. Record weight (TAG) of tag and wire. Loop fine copper wire around clod, leaving a tail to which round stock tag is attached. Record weight of clod (CC1).

7.5 Dip clod in 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time and then reweigh clod and record weight (CC2).

7.6 The clod should be waterproof and ready for volume measurement by water displacement. Suspend the clod below the balance, submerge in water, and record weight (WMCW).

7.7 Dry clod in an oven at 110 °C until weight is constant. Weigh oven-dry clod in air (WODC) and in water (WODCW) and record weights.

7.8 If clod contains >5% rock fragments by weight, remove them from clod. Place the clod in a beaker and place on hot plate. Cover hot plate with a
Figure 3B1a-1.—Dipping clods with hairnet in plastic-lacquer.

Figure 3B1a-2.—After dipping, clods are tied to clothesline to dry.
liquid vapor trap. Use a fume hood. Heat clod on hot plate in excess of 200 °C for 3 to 4 h. The plastic coating disintegrates at temperatures above 200 °C. After heating, clod should appear black and charred. Remove clod from hot plate, lightly coat with liquid detergent, and add hot water.

7.9 Wet sieve the cool soil through a 2-mm, square-hole sieve. Dry and record weight (RF) of rock fragments that are retained on the sieve. Determine rock fragment density by weighing them in air to obtain their mass and in water to obtain their volume. If rock fragments are porous and have a density similar to soil sample, do not correct clod mass and volume measurement for rock fragments. Correct for rock fragments if these fragments can withstand breakdown when dry soil is placed abruptly in water or Calgon.

7.10 Correct bulk density for weight and volume of plastic coating. The coating has an air-dry density of ≈1.3 g cm⁻³. The coating loses 10 to 20% of its air-dry weight when dried in oven at 110 °C.

8. Calculations

8.1 \( D_{b_i} = \frac{\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}}{\left[ \left( \frac{\text{CC2} - \text{WMCW}}{\text{WD}} \right) - (\frac{\text{RF}}{\text{PD}}) - (\frac{\text{MPC}}{1.3}) \right]} \)

where:
- \( D_{b_i} \) = Bulk density in g cc⁻¹ of <2-mm fabric at field-sampled water state
- \( \text{WODC} \) = Weight of oven-dry coated clod
- \( \text{RF} \) = Weight of rock fragments
- \( \text{ODPC} = \text{MPC} \times 0.85 \), weight of oven-dry plastic coat
- \( \text{TAG} \) = Weight of tag and wire
- \( \text{CC2} \) = Weight of clod after three laboratory plastic coats
- \( \text{WMCW} \) = Weight of coated clod in water before oven drying
- \( \text{WD} \) = Water density
- \( \text{PD} \) = Density of rock fragments

8.2 \( \text{MPC} = \left[ \frac{\text{CC2} - \text{CC1}}{3} + \text{FCE} \right] \times \text{RV} \)

where:
- \( \text{MPC} \) = Weight of plastic coat before oven-drying
- \( \text{CC1} \) = Weight of clod before three laboratory plastic coats
- \( \text{RV} \) = Percent estimate of remaining clod volume after cutting to obtain flat surface (≈80%)

8.3 \( \text{FCE} = 1.5 \times \left[ \frac{\text{CC2} - \text{CC1}}{3} \right] \)

where:
- \( \text{FCE} \) = Estimate of field-applied plastic coat
8.4 $\text{Db}_{od} = \frac{\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}}{\left(\frac{\text{WODC} - \text{WODCW}}{\text{WD}}\right) - \left(\frac{\text{RF}}{\text{PD}}\right) - \left(\frac{\text{MPC}}{1.3}\right)}$

where:
- $\text{Db}_{od} =$ Bulk density in g cm$^{-3}$ <2-mm fabric at oven dryness
- $\text{WODCW} =$ Weight of oven-dry coated clod in water

8.5 $W_f = \frac{\left(\text{CC2} - \text{MPC}\right) - \left(\text{WODC} - \text{ODPC}\right)}{\left(\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}\right)} \times 100$

where:
- $W_f =$ Percent water weight in sampled clod

9. Report

Bulk density is reported to the nearest 0.01 g cm$^{-3}$.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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**Bulk Density (3B)**

**Saran-Coated Clods (3B1)**

**33-kPA Desorption ($\text{Db}_{33}$) (3B1b)**

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to determine the coefficient of linear extensibility, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Method 3B1b determines the bulk density value ($\text{Db}_{33}$) of a soil sample equilibrated at 33 kPa.

2. Summary of Method

Field-occurring fabric (clods) are collected from the face of an excavation. One coat of plastic lacquer is applied in the field. Additional coats of plastic lacquer
are applied in the laboratory. The clod is desorbed to 33 kPa. After equilibration, the clod is weighed in air to measure its mass and in water to measure its volume. After the clod is dried in an oven at 110 °C, the mass and volume are determined again. A correction is made for the mass and volume of rock fragments and for plastic coatings (Brasher et al., 1966; Blake and Hartge, 1986).

3. Interferences

Errors are caused by nonrepresentative samples. Only field-occurring fabric (clods) should be sampled. The whole-soil bulk density may be overestimated because sampled clods frequently exclude the crack space between clods (Grossman and Reinsch, 2002).

The penetration of plastic lacquer into the voids of sandy and organic soils interferes with the corrections for mass and volume of the plastic coat and with the accuracy of water content determinations. Penetration can be reduced by spraying water on the clod and then immediately dipping the clod in the plastic lacquer. Dipping should be done as quickly as possible to reduce penetration of plastic.

Loss of soil during the procedure voids the analyses because all calculations are based on the oven-dry soil mass. Holes in the plastic coating, which are detected by air bubbles escaping from submerged clod, introduce errors in volume measurement. An inadequate evaporation of plastic solvent results in overestimation of the soil mass. A drying time of 1 h is usually sufficient for evaporation of solvent. However, clods with high organic matter content may need longer.

Clods placed in an unsealed plastic bag can lose moisture during storage prior to analysis. If clods irreversibly dry below 33-kPa-water content, then $D_b^{1/3}$ values for 33 kPa will be erroneous. Completely seal the plastic storage bag to prevent drying.

Bulk density is reported for <2-mm soil fabric. Correction for rock fragments >2-mm requires either knowledge or assumption of the rock fragment density. Errors in the estimation or measurement of rock fragment density affect the accuracy of the value for soil bulk density. Rock fragments may contain water, which complicates the application to actual water-holding capacity.

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).
Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate
5.3 Air pressure, 33-kPa
5.4 Rigid shipping containers. The KSSL uses a corrugated box with compartments.
5.5 Plastic bags, 1-mil, 127 x 89 x 330 mm
5.6 Wire. The KSSL uses a 28-awg coated copper wire.
5.7 Hairnets
5.8 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.9 Hook assembly for weighing below balance
5.10 Plexiglas water tank
5.11 Lift apparatus, powered by compressed air, Hoff’s Machine & Welding, Inc.
5.12 Oven, 110 °C
5.13 Sieve, No. 10 (2-mm openings)
5.14 Rope, 3-m
5.15 Clothespins
5.16 Knife
5.17 Tile cut-off saw with diamond blade
5.18 Hot plate
5.19 Desiccator with ceramic plate
5.20 Vacuum, 80-kPa (0.8 bar)
5.21 Metal probe
5.22 Spray bottle
5.23 Reinforced paper towels or cheesecloth
5.24 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.

6. Reagents

6.1 Acetone (2-propanone; dimethyl ketone)
6.2 Water
6.3 Alcohol
6.4 Dow Saran F-310 resin, available from Dow Chemical Company
6.5 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a wooden stick. In the laboratory, stir solvent with a non-sparking, high-speed stirrer while slowly adding resin. Stir plastic lacquer for 15 min at 25 °C. Store plastic lacquer in covered plastic or steel containers.

7. Procedure

Field

7.1 Collect field-occurring clods, ≈100 to 200 cm$^3$ in volume (fist-sized), from the face of the excavation. Three clods per horizon are recommended. It is important that these clods be as representative of the bulk sample volume as possible. Remove a piece of soil larger than a clod from the face of sampling pit. From this piece, prepare a clod by gently cutting or breaking protruded peaks and compacted material from clod. If roots are present, trim roots with shears. No procedure for sampling clods is applicable to all soils. Adjust field-sampling techniques to meet field-conditions at time of sampling.

7.2 Make a clothesline by stretching a rope between two fixed points. Tie clod with fine copper wire or place clod in a hairnet. If the clod is dry, moisten surface with a fine mist of water. Quickly dip entire clod into plastic lacquer (fig. 3B1a-1). Suspend clod from clothesline to dry (fig. 3B1a-2). Dry clod for 30 min or until odor of solvent dissipates. If the value of Db, is required, store clods in waterproof plastic bags as soon as coating dries because coating is permeable to water vapor.

7.3 Pack clods in rigid containers to protect them during transport.

Laboratory

7.4 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod (fig. 3B1b-1). Record weight (TAG) of tag and wire. Loop fine copper wire around clod, leaving a tail to which round stock tag is attached. Record weight of clod (CC1).

7.5 Dip clod in 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time and then reweigh clod and record weight (CC2).
Figure 3B1b-1.—A round stock tag with sample identification number is prepared. The cut copper wire is looped around the clod.

7.6 With a diamond saw, cut a flat surface on the clod. Place cut clod surface on a tension table, maintained at 5-cm tension (fig. 3B1b-2). Periodically check clod to determine if it has reached equilibrium. Determination can be made by inserting metal probe, touching, or comparing weight. When clod has reached equilibrium, remove clod and record weight (WSC).

7.7 If cut clod does not adsorb water, place clod in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of clod in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until clod has equilibrated at saturation. Remove clod and record weight (WSC).

7.8 Place the clod in a pressure-plate extractor. To provide good contact between clod and ceramic plate, cover ceramic plate with paper towels and saturate with water. Place surface of cut clod on paper towel. Close container and secure lid. Apply gauged air pressure of 33 kPa. When water ceases to discharge from outflow tube, clod is at equilibrium. Extraction usually takes 3 to 4 weeks. Remove clod and record weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, equilibrate clod on tension table and repeat desorption process.
Figure 3B1b-2.—After a flat surface on the clod is cut with a diamond saw, the clod is placed on a tension table, maintained at 5-cm tension.

7.9 Dip clod in the 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod and record weight (CC3). If the clod has adsorbed >3% in plastic by weight or smells excessively of solvent, allow longer drying time and then reweigh clod.

7.10 The clod should be waterproof and ready for volume measurement by water displacement. Suspend clod below the balance, submerge in water, and record weight (WMCW).

7.11 Dry clod in an oven at 110 °C until weight is constant. Weigh oven-dry clod in air (WODC) and in water (WODCW) and record weights.

7.12 If clod contains >5% rock fragments by weight, remove them from clod. Place the clod in a beaker and place on hot plate. Use a fume hood. Heat clod on hot plate in excess of 200 °C for 3 to 4 h. The plastic coating disintegrates at temperatures above 200 °C. After heating, clod should appear black and charred. Remove clod from hot plate and add hot water.

7.13 Wet sieve the cool soil through a 2-mm, square-hole sieve. Dry and record weight (RF) of rock fragments that are retained on the sieve. Determine rock fragment density by weighing them in air to obtain their mass and in water to obtain their volume. If rock fragments are porous and have
a density similar to soil sample, do not correct clod mass and volume measurement for rock fragments. Correct for rock fragments if these fragments can withstand breakdown when dry soil is placed abruptly in water or Calgon.

7.14 Correct bulk density for weight and volume of plastic coating. The coating has an air-dry density of $\approx 1.3 \text{ g cm}^{-3}$. The coating loses 10 to 20% of its air-dry weight when dried in oven at 110 °C.

8. Calculations

8.1 $D_{b_{33}} = \frac{W_{ODC} - RF - ODPC - TAG}{(CC3 - WMCW)/WD} - \frac{(RF/PD) - (MPC1 / 1.3)}{}}$

where:

$D_{b_{33}}$ = Bulk density in g cc$^{-1}$ of <2-mm fabric at 33-kPa tension
$W_{ODC}$ = Weight of oven-dry coated clod
$RF$ = Weight of rock fragments
$TAG$ = Weight of tag and wire
$ODPC$ = MPC1$x0.85$, weight of oven-dry plastic coat
$CC3$ = Weight of equilibrated clod after four additional plastic coats
$WD$ = Water density
$PD$ = Density of rock fragments
$MPC1$ = Weight of plastic coat before oven-drying
$WMCW$ = Weight in water of coated clod equilibrated at 33-kPa tension

8.2 $MPC1 = ((CC2 - CC1) + FCE) \times RV + (CC3 - WMC)$

where:

$MPC1$ = Weight of plastic coat before oven-drying
$CC2$ = Weight of clod after three laboratory plastic coats
$CC1$ = Weight of clod before three laboratory plastic coats
$WMC$ = Weight of coated clod equilibrated at 33-kPa tension
$RV$ = Percent estimate of remaining clod volume after cutting to obtain flat surface ($\approx 80\%$)

8.3 $FCE = 1.5 \times [(CC2 - CC1)/3]$

where:

$FCE$ = Estimate of field-applied plastic coat

8.4 $D_{b_{od}} = \frac{W_{ODC} - RF - ODPC - TAG}{(W_{ODC} - W_{ODCW})/WD} - \frac{(RF/PD) - (MPC1 / 1.3)}{}}$

where:

$D_{b_{od}}$ = Bulk density in g cc$^{-1}$ <2-mm fabric, oven-dry fabric
WODCW = Weight of oven-dry clod coated in water

$$W_{33} = \left( \frac{((CC3 - MPC1) - (WODC - ODPC))/((WODC - RF - ODPC - TAG))}{100} \right) \times 100$$

where:

$W_{33}$ = Percent water weight retained at 33-kPa tension

9. Report
Bulk density is reported to the nearest 0.01 g cm$^{-3}$.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

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**Bulk Density (3B)**

**Saran-Coated Clods (3B1)**

**Oven-Dry ($Db_{od}$) (3B1c)**

1. Application
Bulk density is used to convert data from a weight basis to a volume basis, to determine the coefficient of linear extensibility, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Method 3B1c determines the bulk density value ($Db_{od}$) of an oven-dry soil sample.

2. Summary of Method
Field-occurring fabric (clods) is collected from the face of an excavation. One coat of plastic lacquer is applied in the field. Additional coats of plastic lacquer are applied in the laboratory. The clod is dried in an oven at 110 °C and then weighed in air to measure its mass and in water to measure its volume. A correction is made for the mass and volume of rock fragments and for plastic coatings (Brasher et al., 1966; Blake and Hartge, 1986).
3. Interferences

Errors are caused by nonrepresentative samples. Only field-occurring fabric (clods) should be sampled. The whole-soil bulk density may be overestimated because sampled clods frequently exclude the crack space between clods (Grossman and Reinsch, 2002).

The penetration of plastic lacquer into the voids of sandy or organic soils interferes with the corrections for mass and volume of the plastic coat and with the accuracy of water content determinations. Penetration can be reduced by spraying water on the clod and then immediately dipping the clod in the plastic lacquer. Dipping should be done as quickly as possible to reduce penetration of plastic.

Loss of soil during the method voids the analyses because all calculations are based on the oven-dry soil mass. Holes in the plastic coating, which are detected by air bubbles escaping from submerged clod, introduce errors in volume measurement. An inadequate evaporation of plastic solvent results in overestimation of the soil mass. A drying time of 1 h is usually sufficient for evaporation of solvent. However, clods with high organic matter content may need longer.

Bulk density is reported for <2-mm soil fabric. Correction for rock fragments >2-mm requires either knowledge or assumption of the rock fragment density. Errors in the estimation or measurement of rock fragment density affect the accuracy of the value for soil bulk density. Rock fragments may contain water, which complicates the application to actual water-holding capacity.

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Rigid shipping containers. The KSSL uses a corrugated box with compartments.
5.3 Plastic bags, 1-mil, 127 x 89 x 330 mm
5.4 Wire. The KSSL uses a 28-awg coated copper wire.
5.5 Hairnets
5.6 Stock tags, 25.4-mm (1-in) diameter paper tag with metal rim
5.7 Hook assembly for weighing below balance
5.8 Plexiglas water tank
5.9 Lift apparatus, powered by compressed air, Hoff’s Machine & Welding, Inc.
5.10 Oven, 110 °C
5.11 Sieve, no. 10 (2-mm openings)
5.12 Rope, 3-m
5.13 Clothespins
5.14 Hot plate
5.15 Spray bottle
5.16 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.

6. Reagents
6.1 Acetone (2-propanone; dimethyl ketone)
6.2 Water
6.3 Dow Saran F-310 resin, available from Dow Chemical Company
6.4 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a wooden stick. In the laboratory, stir solvent with a non-sparking, high-speed stirrer while slowly adding resin. Stir plastic lacquer for 15 min at 25 °C. Store plastic lacquer in covered plastic or steel containers.

7. Procedure

Field

7.1 Collect field-occurring clods, ≈100 to 200 cm³ in volume (fist-sized), from the face of the excavation. Three clods per horizon are recommended. It is important that these clods be as representative of the bulk sample volume as possible. Remove a piece of soil larger than a clod from the face of sampling pit. From this piece, prepare a clod by gently cutting or breaking protruded peaks and compacted material from clod. If roots are present, trim roots with shears. No procedure for sampling clods is applicable to all
soils. Adjust field-sampling techniques to meet field conditions at time of sampling.

7.2 Make a clothesline by stretching a rope between two fixed points. Tie clod with fine copper wire or place clod in a hairnet. If the clod is dry, moisten surface with a fine mist of water. Quickly dip entire clod into plastic lacquer (fig. 3B1a-1). Suspend clod from clothesline to dry (fig. 3B1a-2). Dry clod for 30 min or until odor of solvent dissipates. If the value of Db is required, store clods in waterproof plastic bags as soon as coating dries because coating is permeable to water vapor.

7.3 Pack clods in rigid containers to protect them during transport.

Laboratory

7.4 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod. Record weight (TAG) of tag and wire. Loop fine copper wire around clod, leaving a tail to which round stock tag is attached. Record weight of clod (CC1).

7.5 Dip clod in 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time and then reweigh clod and record weight (CC2).

7.6 Dry clod in an oven at 110 °C until weight is constant. Weigh oven-dry clod in air (WODC) and in water (WODCW) and record weights.

7.7 If clod contains >5% rock fragments by weight, remove them from clod. Place the clod in a beaker and place on hot plate. Cover hot plate with a liquid vapor trap. Use a fume hood. Heat clod on hot plate in excess of 200 °C for 3 to 4 h. The plastic coating disintegrates at temperatures above 200 °C. After heating, clod should appear black and charred. Remove clod from hot plate, lightly coat with liquid detergent, and add hot water.

7.8 Wet sieve the cool soil through a 2-mm, square-hole sieve. Dry and record weight (RF) of rock fragments that are retained on the sieve. Determine rock fragment density by weighing them in air to obtain their mass and in water to obtain their volume. If rock fragments are porous and have a density similar to soil sample, do not correct clod mass and volume measurement for rock fragments. Correct for rock fragments if these fragments can withstand breakdown when dry soil is placed abruptly in water or Calgon.

7.9 Correct bulk density for weight and volume of plastic coating. The coating has an air-dry density of ≈1.3 g cm$^{-3}$. The coating loses 10 to 20% of its air-dry weight when dried in oven at 110 °C.
8. Calculations

8.1 \( \text{Db}_{od} = \frac{[\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}]}{[\{\text{WODC} - \text{WODCW}\}/\text{WD}] - (\text{RF}/\text{PD}) - (\text{MPC}/1.3)} \)

where:
- \( \text{Db}_{od} \) = Bulk density in g cm\(^{-3}\) of <2-mm, oven-dry fabric
- \( \text{WODC} \) = Weight of oven-dry coated clod
- \( \text{RF} \) = Weight of rock fragments
- \( \text{TAG} \) = Weight of tag and wire
- \( \text{ODPC} \) = MPC \( \times \) 0.85, weight of oven-dry plastic coat
- \( \text{WD} \) = Water density
- \( \text{PD} \) = Density of rock fragments
- \( \text{WODCW} \) = Weight of oven-dry coated clod in water

8.2 \( \text{MPC} = [(\text{CC2} - \text{CC1}) + \text{FCE}] \times \text{RV} \)

where:
- \( \text{MPC} \) = Weight of plastic coat before oven-drying
- \( \text{CC2} \) = Weight of clod after three laboratory plastic coats
- \( \text{CC1} \) = Weight of clod before three laboratory plastic coats
- \( \text{RV} \) = Percent estimate of remaining clod volume after cutting to obtain flat surface (≈80%)

8.3 \( \text{FCE} = 1.5 \times \frac{(\text{CC2} - \text{CC1})}{3} \)

where:
- \( \text{FCE} \) = Estimate of field-applied plastic coat

9. Report

Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Bulk Density (3B)
Saran-Coated Clods (3B1)
Rewet (Db,) (3B1d)

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to determine the coefficient of linear extensibility, to estimate saturated hydraulic conductivity, and to identify compacted horizons. The rewet bulk density (Db,) is used to determine irreversible shrinkage of soils and subsidence of organic soils. Method 3B1d determines the bulk density value (Db,) of a re-wetted soil sample.

2. Summary of Method

Field-occurring fabric (clods) is collected from the face of an excavation. One coat of plastic lacquer is applied in the field. Additional coats of plastic lacquer are applied at the laboratory. After equilibration, the clod is weighed in air to measure its mass and in water to measure its volume. The clod is air-dried, re-equilibrated, and its mass and volume re-measured. After the clod is dried in an oven at 110 °C, the mass and volume are determined again. A correction is made for the mass and volume of rock fragments and for plastic coatings (Brasher et al., 1966; Blake and Hartge, 1986; Grossman and Reinsch, 2002).

3. Interferences

Errors are caused by nonrepresentative samples. Only field-occurring fabric (clods) should be sampled. The whole-soil bulk density may be overestimated because sampled clods frequently exclude the crack space between clods (Grossman and Reinsch, 2002).

The penetration of plastic lacquer into the voids of sandy or organic soils interferes with the corrections for mass and volume of the plastic coat and with the accuracy of water content determinations. Penetration can be reduced by spraying water on the clod and then immediately dipping the clod in the plastic lacquer. Dipping should be done as quickly as possible to reduce penetration of plastic.

Loss of soil during the procedure voids the analyses because all calculations are based on the oven-dry soil mass. Holes in the plastic coating, which are detected by air bubbles escaping from submerged clod, introduce errors in volume measurement. Inadequate drying results in overestimation of the soil mass. An inadequate evaporation of plastic solvent results in overestimation of the soil mass. A drying time of 1 h is usually sufficient for evaporation of solvent. However, clods with high organic matter content may need longer.

Bulk density is reported for <2-mm soil fabric. Correction for rock fragments >2-mm requires either knowledge or assumption of the rock fragment density.
Errors in the estimation or measurement of rock fragment density affect the accuracy of the value for soil bulk density. Rock fragments may contain water, which complicates the application to actual water-holding capacity.

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate
5.3 Air pressure, 33-kPa
5.4 Rigid shipping containers. The KSSL uses a corrugated box with compartments.
5.5 Plastic bags, 1-mil, 127 x 89 x 330 mm
5.6 Wire. The KSSL uses a 28-awg coated copper wire.
5.7 Hairnets
5.8 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.9 Hook assembly for weighing below balance
5.10 Plexiglas water tank
5.11 Lift apparatus, powered by compressed air, Hoff’s Machine & Welding, Inc.
5.12 Oven, 110 °C
5.13 Sieve, No. 10 sieve (2-mm openings)
5.14 Rope, 3-m
5.15 Clothespins
5.16 Knife
5.17 Tile cut−off saw with diamond blade
5.18 Hot plate
5.19 Desiccator with ceramic plate
5.20 Vacuum, 80-kPa (0.8 bar)
5.21 Metal probe
5.22 Spray bottle
5.23 Reinforced paper towels or cheesecloth
5.25 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.

6. Reagents

6.1 Acetone (2-propanone; dimethyl ketone)
6.2 Water
6.3 Alcohol
6.4 Dow Saran F-310 resin, available from Dow Chemical Company
6.5 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the laboratory, stir solvent with a non-sparking, high-speed stirrer while slowly adding resin. Stir plastic lacquer for 15 min at 25 °C. Store plastic lacquer in covered plastic or steel containers. Acetone may be substituted for MEK (methyl ethyl ketone) as a solvent.

7. Procedure

7.1 Collect field-occurring clods, ≈100 to 200 cm³ in volume (fist-sized), from the face of the excavation. Three clods per horizon are recommended. It is important that these clods be as representative of the bulk sample volume as possible. Remove a piece of soil larger than a clod from the face of sampling pit. From this piece, prepare a clod by gently cutting or breaking protruded peaks and compacted material from clod. If roots are present, trim roots with shears. No procedure for sampling clods is applicable to all soils. Adjust field sampling techniques to meet field conditions at time of sampling.

7.2 Make a clothesline by stretching a rope between two fixed points. Tie clod with fine copper wire or place clod in a hairnet. If the clod is dry, moisten surface with a fine mist of water. Quickly dip entire clod into plastic lacquer (fig. 3B1a-1). Suspend clod from clothesline to dry (fig. 3B1a-2). Dry clod for 30 min or until odor of solvent dissipates. If the value of Db is required, store clods in waterproof plastic bags as soon as coating dries because coating is permeable to water vapor.
7.3 Pack clods in rigid containers to protect them during transport.

**Laboratory**

7.4 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod. Record weight (TAG) of tag and wire. Loop fine copper wire around clod, leaving a tail to which round stock tag is attached. Record weight of clod (CC1).

7.5 Dip clod in 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time and then reweigh clod and record weight (CC2).

7.6 With a diamond saw, cut a flat surface on the clod.

7.7 Place cut clod surface on a tension table, maintained at 5-cm tension. Periodically check clod to determine if it has reached equilibrium. Determination can be made by inserting metal probe, touching, or comparing weight. When clod has reached equilibrium, remove clod and record weight (WSC).

7.8 If cut clod does not adsorb water, place clod in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of clod in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until clod has equilibrated at saturation. Remove clod and record weight (WSC).

7.9 Place the clod in a pressure-plate extractor. To provide good contact between clod and ceramic plate, cover ceramic plate with paper towels and saturate with water. Place surface of cut clod on paper towel. Close container and secure lid. Apply gauged air pressure of 33-kPa. When water ceases to discharge from outflow tube, clod is at equilibrium. Extraction usually takes 3 to 4 weeks. Remove clod and record weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, equilibrate clod on tension table and repeat desorption process.

7.10 Dip clod in the 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod and record weight (CC3). If the clod has adsorbed >3% in plastic by weight or smells excessively of solvent, allow longer drying time and then reweigh clod.

7.11 The clod should be waterproof and ready for volume measurement by water displacement. Suspend clod below the balance, submerge in water, and record weight (WMCW).
7.12 Remove layer of plastic from flat surface of clod. Air-dry clod at room temperature (≈20 to 25 °C) for 4 to 6 days. Dry clod at 40 to 50 °C for 2 to 3 days or until weight is constant.

7.13 Repeat steps 7.7, 7.8, and 7.9. After equilibrium is obtained, remove clod and record weight (WAR).

7.14 Dip clod in the 1:4 plastic lacquer. Wait 7 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and then dip clod in 1:7 plastic lacquer. Wait 12 min and dip clod in 1:7 plastic lacquer. Wait 55 min and then reweigh clod and record weight (CC4). If the clod has adsorbed >3% in plastic by weight or smells excessively of solvent, allow longer drying time and then reweigh clod.

7.15 After coating, record weight of clod suspended in air (CC4) and in water (WARW).

7.16 Dry clod in an oven at 110 °C until weight is constant. Weigh oven-dry clod in air (WODC) and in water (WODCW) and record weights.

7.17 If clod contains >5% rock fragments by weight, remove them from clod. Place the clod in a beaker and place on hot plate. Use a fume hood. Heat clod on hot plate in excess of 200 °C for 3 to 4 h. The plastic coating disintegrates at temperatures above 200 °C. After heating, clod should appear black and charred. Remove clod from hot plate and add hot water.

7.18 Wet sieve the cool soil through a 2-mm, square-hole sieve. Dry and record weight (RF) of rock fragments that are retained on the sieve. Determine rock fragment density by weighing them in air to obtain their mass and in water to obtain their volume. If rock fragments are porous and have a density similar to soil sample, do not correct clod mass and volume measurement for rock fragments. If rock fragments are porous and have a density similar to soil sample, do not correct clod mass and volume measurement for rock fragments. Correct for rock fragments if these fragments can withstand breakdown when dry soil is placed abruptly in water or Calgon.

7.19 Correct bulk density for weight and volume of plastic coating. The coating has an air-dry density of ≈1.3 g cm⁻³. The coating loses 10 to 20% of its air-dry weight when dried in oven at 110 °C.

8. Calculations

8.1 $D_b_{33} = \frac{\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}}{\left[\frac{\text{(CC3} - \text{WMCW})}{\text{WD}} - (\text{RF/PD}) - (\text{MPC1/1.3})\right]}$

    where:
    
    $D_b_{33} =$ Bulk density in g cm⁻³ of <2-mm fabric at 33-kPa tension
    
    WODC = Weight of oven-dry coated clod
RF = Weight of rock fragments
ODPC = MPC1 x 0.85, weight of oven-dry plastic coat
TAG = Weight of tag and wire
CC3 = Weight of equilibrated clod after four additional plastic coats
WD = Water density
PD = Density of rock fragments
WMCW = Weight in water of coated clod equilibrated at 33-kPa tension

8.2 MPC1 = \{[(CC2 - CC1) + FCE] x RV\} + (CC3 - WMC)

where:
MPC1 = Weight of plastic coat before air-drying and rewet
CC2 = Weight of clod after three laboratory plastic coats
CC1 = Weight of clod before three laboratory plastic coats
RV = Percent estimate of remaining clod volume after cutting to obtain flat surface (≈80%)
WMC = Weight of coated clod equilibrated at 33-kPa tension

8.3 FCE = 1.5 \times \frac{[(CC2 - CC1)]}{3}

where:
FCE = Estimate of field-applied plastic coat.

8.4 \( D_{br} = \frac{[WODC - RF - ODPC - TAG]}{[(CC4 - WARW) / WD] - (RF / PD) - (MPC2 / 1.3)} \)

where:
\( D_{br} \) = Bulk density in g cm\(^{-3}\) <2-mm fabric at 33-kPa tension after rewetting
CC4 = Weight of clod after twelve plastic coats
WARW = Weight in water of coated clod equilibrated at 33-kPa tension after rewetting

8.5 MPC2 = \{[(CC2 - CC1) + FCE] x RV2\} + (CC3 - WMC) + (CC4 - WAR)

where:
MPC2 = Weight of plastic coat after rewetting and before oven drying
WAR = Weight of clod after rewet equilibration
RV2 = Percent estimate of remaining clod volume after remaining layer of plastic (≈0.95)

8.6 \( D_{od} = \frac{[WODC - RF - ODPC - TAG]}{[(WODC - WODCW) / WD] - (RF / PD) - (MPC2 / 1.3)} \)

where:
\( D_{od} \) = Bulk density in g cm\(^{-3}\) of <2-mm fabric at oven dryness
WODCW = Weight in water of oven-dry coated clod

8.7 \[ W_{33} = \frac{((CC3 - MPC1) - (WODC - ODPC))}{(WODC - RF - ODPC - TAG)} \times 100 \]

where:

- \( W_{33} \) = Percent water weight retained at 33-kPa tension

8.8 \[ W_r = \frac{((CC4 - MPC2) - (WODC - ODPC))}{(WODC - RF - ODPC - TAG)} \times 100 \]

where:

- \( W_r \) = Percent water weight retained at 33-kPa tension after rewet

9. Report

Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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**Bulk Density (3B)**

**Reconstituted (3B2)**

- 33-kPa Desorption (\( D_{33} \)) (3B2a)
- Oven-Dry (\( D_{od} \)) (3B2b)

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to determine the coefficient of linear extensibility, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Some models and programs require one bulk density to represent a given horizon. The reconstituted bulk density provides a single, reproducible value for horizons that are subject to tillage or other mechanical disturbances followed by an extreme water-state cycle (Reinsch and Grossman, 1995).
2. Summary of Method
   The <2-mm sample is formed into a clod by wetting and desiccation cycles that simulate reconsolidation by water in a field setting. Plastic lacquer is applied in the laboratory to form an impermeable coat on the clod. The clod is desorbed to 33-kPa. After equilibration, the clod is weighed in air to measure the mass and in water to measure the volume. After the clod is oven dried at 110 °C, its mass and volume are determined again (Brasher et al., 1966; Blake and Hartge, 1986; Grossman and Reinsch, 2002).

3. Interferences
   Some samples disintegrate when they are removed from the cells.

4. Safety
   Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).
   Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

5. Equipment
   5.1 Electronic balance, ±0.01-g sensitivity
   5.2 Pressure-plate extractor with porous ceramic plate
   5.3 Air pressure, 33-kPa
   5.4 Clod forming cylinder. The KSSL constructs the cell by attaching a brass ring or schedule 20 or 40 PVC pipe, 5.4-cm diameter and 6 to 7 cm high, to a 100-kPa ceramic plate with waterproof glue and caulk.
   5.5 Anti-sorting device. To reduce natural sorting caused by placing the sample in the cell, the KSSL uses a device created by attaching a perpendicular wire to the center of a 5.2-cm diameter wire screen with 0.5-cm openings.
   5.6 Wire. The KSSL uses 28-awg coated copper wire.
   5.7 Hairnets
   5.8 Stock tags, 1-inch diameter paper tag with metal rim
   5.9 Hook assembly for weighing below balance
   5.10 Plexiglas water tank
   5.11 Lift apparatus, powered by compressed air, Hoff’s Machine & Welding, Inc.
5.11 Oven, 110 °C
5.12 Plastic tub at least 10 cm deep
5.13 Paper discs cut from water-insoluble, permeable paper
5.14 Tile cut-off saw with diamond blade
5.15 Desiccator with ceramic plate
5.16 Vacuum, 80-kPa (0.8 bar)
5.17 Metal probe
5.18 Spray bottle
5.19 Reinforced paper towels or cheesecloth
5.20 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.

6. Reagents

6.1 Acetone (2-propanone; dimethyl ketone)
6.2 Water
6.3 Alcohol
6.4 Dow Saran F-310 resin, available from Dow Chemical Company
6.5 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of the handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and reduce cost. Stir the solvent with a non-sparking, high-speed stirrer while slowly adding the resin. Stir the plastic lacquer for 15 min at 25 °C. In the field, mix with a wooden stick. Store the plastic lacquer in covered plastic or steel containers. Acetone may be substituted for MEK (methyl ethyl ketone) as a solvent.

7. Procedure

Reconstituted Clod Construction

7.1 Drape a hairnet in the cell. Place a paper disc in the bottom of the cell. Place the anti-sorting screen into the cell. Add <2-mm, prepared sample to within a few mm of the top of the cell. Lift the anti-sorting screen from the cell.

7.2 Place the cell on a tension table with the top of the cell 5-cm below the top of the table. After equilibration, place the cell into a tub and add water to a level higher than the surface of the soil in the cell but below the top lip of
the cell. This allows the soil to become inundated from beneath. Allow the sample to equilibrate.

7.3 Remove the cell from the tub and allow to dry at room temperature. After the clod has dried, remove the clod by lifting on the hairnet or inverting the cell and lightly tamping the base of the cell. The reconstituted clod is used to measure bulk density and water retention.

**Bulk Density Measurement**

7.4 Prepare a round stock tag with sample identification number. Cut the copper wire to loop around the clod. Record the weight of the tag and wire (TAG). Loop fine copper wire around the clod leaving a tail to which the round stock tag is attached. Record the weight of the clod (CC1).

7.5 Mist the clod with water to create a film of water on the surface of the clod. Dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 55 min and then reweigh the clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time, then reweigh the clod and record the weight (CC2).

7.6 Cut a flat surface on the clod with a diamond saw. Place the cut clod surface on a tension table maintained at 5–cm tension. Periodically check clod to determine if it has reached equilibrium. Determination can be made by inserting metal probe, touching, or comparing weight. When the clod has reached equilibrium, remove the clod and record the weight (WSC).

7.7 If the clod does not adsorb water, place the clod in a desiccator that has water covering the desiccator plate. Add a few mL of alcohol. Apply suction using in-house vacuum for 24 hours. Remove the clod and record the weight (WSC).

7.8 Place the clod in a pressure-plate extractor. To provide good contact between the clod and ceramic plate, cover the ceramic plate with paper towels and saturate with water. Place the cut surface of the clod on the paper towel. Close the container and secure the lid. Apply gauged air pressure of 33-kPa. When water stops discharging from the outflow tube (usually after 3 or 4 weeks in the extractor), the clod is at equilibrium. Remove the clod and record the weight (WMC). Compare WMC to WSC. If WMC is greater than or equal to WSC, equilibrate the clod on the tension table and repeat the desorption process.

7.9 Dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 55 min and then reweigh the clod and record the weight (CC3). If the
clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time and then reweigh the clod.

7.10 The clod should now be waterproof and ready for the volume measurement by water displacement. Suspend the clod below the balance, submerge in water, and record the weight (WMCW).

7.11 Dry the clod in an oven at 110 °C overnight. Weigh the oven-dry clod in air (WODC) and in water (WODCW) and record the weights.

7.12 It is necessary to correct bulk density for weight and volume of the plastic coating. The coating has an air-dry density of about 1.3 g/cm³. The coating loses 10 to 20 percent of its air-dry weight on oven drying at 110 °C.

8. Calculations

8.1 \[ \text{Db}_{33} = \frac{\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}}{\left[\frac{\text{CC2} - \text{WMCW}}{\text{WD}}\right] - \left(\frac{\text{RF}}{\text{PD}}\right) - \left(\frac{\text{MPC1}}{1.3}\right)} \]

where:
- \( \text{Db}_{33} \) = Bulk density in grams per cubic centimeter of <2-mm fabric at 33-kPa tension
- \( \text{WODC} \) = Weight of oven-dry coated clod
- \( \text{ODPC} \) = MPC x 0.85, weight of oven-dry plastic coat
- \( \text{TAG} \) = Weight of tag and wire
- \( \text{CC3} \) = Weight of equilibrated clod after four additional plastic coats
- \( \text{WMCW} \) = Weight of coated clod equilibrated at 33-kPa tension in water
- \( \text{MPC1} \) = Weight of plastic coat before oven-drying
- \( \text{WD} \) = Water density

8.2 \[ \text{MPC1} = \left[\frac{\text{CC2} - \text{CC1}}{\text{RV}}\right] + (\text{CC3} - \text{WMC}) \]

where:
- \( \text{MPC1} \) = Weight of plastic coat before oven-drying
- \( \text{CC2} \) = Weight of clod after four laboratory plastic coats
- \( \text{CC1} \) = Weight of clod before four laboratory plastic coats
- \( \text{RV} \) = Percent estimate of remaining clod volume after cutting to obtain flat surface (≈95%)
- \( \text{WMC} \) = Weight of coated clod equilibrated at 33-kPa tension

8.3 \[ \text{Db}_{\text{od}} = \frac{\text{WODC} - \text{ODPC} - \text{TAG}}{\left[\frac{\text{WODC} - \text{WODCW}}{\text{WD}}\right] - \left(\frac{\text{MPC1}}{1.3}\right)} \]

where:
- \( \text{Db}_{\text{od}} \) = Bulk density in grams per cubic centimeter of <2-mm fabric at oven-dryness
- \( \text{WODCW} \) = Weight of oven-dry coated clod in water

8.4 \[ \text{W}_{33} = \left[\frac{\text{CC3} - \text{MPC1} - (\text{WODC} - \text{ODPC})}{\text{WODC} - \text{ODPC} - \text{TAG}}\right] \times 100 \]
where:
\[ W_{33} \text{= Weight percentage of water retained at 33-kPa tension} \]

9. Report

Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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**Bulk Density (3B)**

**Compliant Cavity (3B3)**

**Field-State (3B3a)**

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Excavation procedures (e.g., compliant cavity, ring, and frame) have applicability to layers that can be described as cohesionless, high in rock fragments >5 mm, or thin (<5 cm thick) and for which the clod method is unsuitable (Grossman and Reinsch, 2002). The compliant cavity method was designed for fragile, cultivated near-surface layers and O horizons of forestland soils. This method has the important advantage that it is not necessary to flatten the ground surface on steep slopes or to remove irregularities, i.e., the surficial zone is usually not altered (Grossman and Reinsch, 2002).
2. Summary of Method

By this method (3B3a), the cavity volume on the zone surface is lined with thin plastic and water is added to a datum level. Soil is quantitatively excavated in a cylindrical form to the required depth. The difference between the initial volume and that after excavation is the sample volume. The excavated soil is dried in an oven and then weighed. A correction is made for the weight and volume of rock fragments.

3. Interferences

Bulk density by the compliant cavity method can be made on soils with rock fragments but is more complex than other methods (Grossman and Reinsch, 2002).

4. Safety

Follow standard field and laboratory safety precautions.

5. Equipment

5.1 Fabricated Plexiglas rings, 9-mm thick, 130-mm inside diameter, and ≥200-mm outside diameter. Make three 16-mm diameter holes that are 10 mm from the outer edge of ring. Position holes equidistant apart. Use three, 25 x 50 mm, Plexiglas pieces as guides. Attach two pieces on one side to form an “L.” Allow a 15-mm gap to permit removal of soil material. On the other side, position the single piece in line with the longer leg of the “L” so that an adjacent, parallel line forms a diameter.

5.2 Make 50-mm thick foam rings from flexible polyurethane with an “Initial Load Displacement” of 15 to 18 kg. The foam rings have the same inside diameter as the Plexiglas rings.

5.3 Fabricate a 240-mm crossbar from 5 x 18 mm metal stock to which legs (25-mm high and 180 x 180 mm in cross section) are welded. Drill a hole 100 mm from one end of the crossbar and 7 mm from the edge and through which a No. 6 machine bolt is placed.

5.4 Mount hook gauge on crossbar. Make hook gauge from No. 6, round-headed, 100-mm long machine bolts with hexagonal nuts. Obtain the machine bolts from toggle bolt assemblies. Sharpen the machine bolt to a sharp point. Drill a hole in the center of the crossbar. Insert the machine bolt in the hole. Place nuts above and below the crossbar. The two nuts adjust the hook length below the crossbar and provide rigidity. Hold the machine bolt by the tightened nuts and heat the bolt. After softening, sharply bend the bolt upward to form a U-shape.

5.5 Use wing nuts and three, 250- to 400-mm long, 10- to 13-mm diameter, threaded rods to mount and position the compliant cavity. Sharpen the
rods. Place two regular nuts at the end of threaded rod to increase the area of surface struck.

5.6 Syringe, 60 mL
5.7 Plastic film, ½ mil, 380-mm wide or wider; 460-mm wide for larger ring.
5.8 Plastic bags, 110 °C capability, with ties
5.9 Sharpie pen
5.10 Graduate cylinders, plastic, 250 to 2000 mL
5.11 Level, small
5.12 Kitchen knife, small
5.13 Scissors, small, to cut fine roots
5.14 Hack saw blade to cut large roots
5.15 Weights for plastic film
5.16 Clothespins. If conditions are windy, use the clothespins for corners of plastic film.
5.17 Hard rubber or plastic mallet
5.18 Sieve, square-hole, 10 mesh, 2 mm

6. Reagents

6.1 Water

7. Procedure

7.1 Place a ring of plastic foam on ground and cover with rigid ring (130-mm inside diameter). Mount the assembly on the soil surface by securely driving threaded rods into the ground through holes in ring and by tightening ring with wing nuts.

7.2 Line cavity with ½ mil plastic. Fill cavity to tip of hook gauge with a known quantity of water from graduated cylinder.

7.3 Remove plastic film and water. Measure the volume of water to tip of hook gauge. This volume ($V_d$) is the measurement of cavity volume prior to excavation (dead space).

7.4 Excavate soil quantitatively and in a cylindrical form to the required depth. Fill the excavation cavity to tip of hook gauge with water from graduated cylinder. Measure the volume of water. This volume ($V_f$) is the measurement of excavated soil and dead space. The difference between the two water volumes ($V_f - V_d$) is the volume of excavated soil ($V_e$).

7.5 The excavated soil is dried in oven and weighed. If necessary, make a correction for weight and volume of >2-mm material ($V_g$) in sample and compute bulk density. The weight of macroscopic vegetal material ($g \text{ cm}^{-3}$) also may be reported.
8. Calculations

8.1 \[ V_e = V_f - V_d - V_g \]

where:
- \( V_e \): Excavation volume of <2-mm fraction (cc)
- \( V_f \): Water volume measurement of excavated soil and dead space (cc)
- \( V_d \): Water volume measurement of dead space (cc)
- \( V_g \): Gravel volume (>2-mm fraction) (cc). Calculate \( V_g \) by dividing the weight of >2-mm fraction by particle density of the >2-mm fraction. Default value is 2.65 g cc\(^{-1}\).

8.2 \[ W_f = W_o - W_c \]

where:
- \( W_f \): Oven-dry weight of <2-mm soil (g)
- \( W_o \): Oven-dry weight of excavated soil (g)
- \( W_c \): Oven-dry weight of rock fragments (g)

8.3 \[ D_b = \frac{W_f}{V_e} \]

where:
- \( D_b \): Bulk density (g cc\(^{-1}\))
- \( W_f \): Oven-dry weight of <2-mm soil (g)
- \( V_e \): Excavation volume of <2-mm material (cc)

9. Report

Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Bulk Density (3B)
Ring Excavation (3B4)
Field-State (3B4a)

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to estimate saturated hydraulic conductivity, and to identify compacted horizons.
Excavation procedures (e.g., compliant cavity, ring, and frame) have applicability to layers that can be described as cohesionless, high in rock fragments >5 mm, or thin (<5 cm thick) and for which the clod method is unsuitable (Grossman and Reinsch, 2002). The ring excavation method is robust, simple, and rapid. This method is good for O horizons in the woods where local variability is large. The diameter can range down to 15 cm and upwards to 30 cm or more. It is not necessary to excavate from the whole area within the ring. A limit of 2 cm on the minimum thickness of the sample should be considered.

2. Summary of Method

A 20-cm ring is inserted into the ground. A piece of shelf standard is placed across the ring near to a diameter. The distance to the ground surface is measured at eight points equally spaced along the diameter using the depth-measurement tool. The piece of shelf is rotated 90°, and eight more measurements are made. The 16 measurements are then averaged. The soil is excavated to the desired depth, and the distance measurements repeated. The change in distance is calculated on the removal of the soil. This change in distance is then multiplied by the inside cross-sectional area of the ring to obtain the volume of soil. The excavated soil is oven-dried and weighed. If rock fragments are present, the weight and volume of >2-mm material in the sample are corrected and bulk density is computed. Bulk density of soil is reported in g cm⁻³ by method 3B4a.

3. Interferences

Rock fragments may make insertion of ring into the ground impossible.

4. Safety

Follow standard field and laboratory safety precautions.

5. Equipment

5.1 Metallic cylinder, 20-cm diameter, 10- to 20-cm high, and about 1-mm deep
5.2 Shelf standard (slotted rod), 1.5-cm wide, 1-cm high, and 25-cm long
5.3 Piece of retractable ruler, 30-cm long with 0.1-mm divisions
5.4 Piece of wood, 10 x 10 x 30 cm
5.5 Hand digging equipment
5.6 Depth-measurement tool (Grossman and Reinsch, 2002)

6. Reagents

None.
7. Procedure

7.1 Insert a 20-cm diameter ring below the depth of excavation.

7.2 Place a piece of shelf standard across the ring near to or along a diameter. Measure the distance to the ground surface at eight points equally spaced along the diameter using the depth-measurement tool.

7.3 Rotate the piece of shelf standard 90° and make eight more measurements. Average the 16 measurements.

7.4 Excavate the soil to the desired depth. Repeat the distance measurements.

7.5 Calculate the change in distance on removal of the soil. Multiply the change in distance by the inside cross-sectional area of the ring to obtain the volume of the soil (Ve).

7.6 Dry the excavated soil in oven and weigh. If necessary, make a correction for weight and volume of >2-mm material in sample and compute bulk density. The weight of macroscopic vegetal material (g cm$^{-3}$) also may be reported.

8. Calculations

8.1 \[ W_f = W_o - W_e \]

where:

- \( W_f \) = Oven-dry weight of <2-mm soil (g)
- \( W_o \) = Oven-dry weight of excavated soil (g)
- \( W_c \) = Oven-dry weight of rock fragments (g)

8.2 \[ D_b = \frac{W_f}{V_e} \]

where:

- \( D_b \) = Bulk density (g cm$^{-3}$)
- \( W_f \) = Oven-dry weight of <2-mm soil (g)
- \( V_e \) = Excavation volume of <2-mm material (cm$^{-3}$)

9. Report

Bulk density is reported to the nearest 0.01 g cm$^{-3}$.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References

Bulk Density (3B)
Frame Excavation (3B5)
Field-State (3B5a)

1. Application

Bulk density is used to convert data from a weight basis to a volume basis, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Excavation procedures (e.g., compliant cavity, ring, and frame) have applicability to layers that can be described as cohesionless, high in rock fragments >5 mm, or thin (<5 cm thick) and for which the clod method is unsuitable (Grossman and Reinsch, 2002). This method is good for O horizons in the woods where local variability is large and rock fragments are commonly present. The size of 0.1 m² is sufficient to encompass considerable local variability.

2. Summary of Method

The assembled frame is placed on the ground surface. The four threaded rods are pushed through the holes in the corners of the frame deep enough to hold. The frame is then secured onto the soil surface by screwing down wing nuts, and plastic is placed over the frame and secured. The depth-measurement tool is placed on top of a slot to measure the distance to the soil surface. The slots are traversed, and measurements of the distance to the ground surface are made at about 40 regularly spaced intervals. The plate is then removed, and soil is excavated and retained. Measurements of the distance to the ground surface are repeated. The volume of soil is determined by taking the difference in height and multiplying by 1000 cm². The rock fragments up to 20 mm are included in the sample. Excavated soil is oven-dried and weighed. Bulk density of soil is reported in g cm⁻³ by method 3B5a.

3. Interferences

None.

4. Safety

Follow standard field and laboratory safety precautions..

5. Equipment

5.1 Lumber for square wooden frame with 0.1 m² inside area. Frame is made from 8 pieces of wood: 2 pieces, 2 x 4 x 46 cm; 2 pieces, 2 x 4 x 53 cm; and 4 blocks, 4 x 5 x 9 cm.

5.2 Square Plexiglas, 35 cm on edge x 0.6 cm thick, with 5 parallel equally spaced slots, 1.5 cm across x 28 cm long
5.3 Four threaded rods, 50 cm long x 0.6 cm diameter, with wing nuts
5.4 Depth-measurement tool (Grossman and Reinsch, 2002, p. 209)
5.5 Hand digging equipment

6. Reagents
   None.

7. Procedure
   7.1 Assemble the square wooden frame by attaching the 9 cm side of a 4 x 5 x 9 cm block to each end of both 53-cm long pieces. Two-centimeter-wide cuts are made half-way across each of the 46- and 53-cm long pieces to provide half-lap joints. The cuts are 5 cm in for the 46-cm long pieces. Holes 1.0 to 1.5 cm in diameter are drilled in the center of the attached blocks. The four pieces are joined by the vertical half-lap joints to form a square frame.
   7.2 Place the frame on the ground surface. Push the four threaded rods through the holes in the corners of the frame sufficiently deeply to hold. Secure onto the soil surface by screwing down wing nuts.
   7.3 Place the plastic plate over the frame and secure.
   7.4 Place the depth-measurement tool on top of a slot and measure the distance to the soil surface.
   7.5 Traverse the slots, making measurements of the distance to the ground surface at about 40 regularly spaced intervals. Remove the plate.
   7.6 Excavate and retain the soil. The walls of the cavity should be vertical and coincident with the edge of the frame.
   7.7 Repeat the measurements of the distance to the ground surface. Determine the difference in height and multiply by 1000 cm$^2$ to obtain the volume of soil excavated. Usually, rock fragments up to 20 mm are included in the sample.
   7.8 Dry the excavated soil in oven and weigh. If necessary, make a correction for weight and volume of >2-mm material in sample and bulk density computed. The weight of macroscopic vegetal material (g cm$^{-3}$) also may be reported.

8. Calculations
   8.1 \[ W_f = W_o - W_e \]
      where:
      \[ W_f = \text{Oven-dry weight of <2-mm soil (g)} \]
      \[ W_o = \text{Oven-dry weight of excavated soil (g)} \]
      \[ W_e = \text{Oven-dry weight of rock fragments (g)} \]
8.2 \( Db=\frac{Wf}{Ve} \)

\( Db = \text{Bulk density (g cm}^{-3}) \)
\( Wf = \text{Oven-dry weight of <2-mm soil (g)} \)
\( Ve = \text{Excavation volume of <2-mm material (cm}^{-3}) \)

9. Report
Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

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**Bulk Density (3B)**

**Soil Cores (3B6)**

**Field-State (3B6a)**

1. **Application**
Bulk density is used to convert data from a weight basis to a volume basis, to determine the coefficient of linear extensibility, to estimate saturated hydraulic conductivity, and to identify compacted horizons. Method 3B6a determines the bulk density value of a moist soil core of known volume. Field bulk density \( (Db_f) \) offers the opportunity to obtain relatively cheaply bulk density information without the expense incurred to obtain water retention. \( Db_f \) is particularly useful if the soil layers are at or above field capacity and/or the soils have low extensibility and do not exhibit desiccation cracks even if below field capacity.

2. **Summary of Method**
A metal cylinder is pressed or driven into the soil. The cylinder is removed, extracting a sample of known volume. The moist sample weight is recorded. The sample is then dried in an oven and weighed.

3. **Interferences**
During coring process, compaction of the sample is a common problem. Compression can be observed by comparing the soil elevation inside the cylinder with the original soil surface outside the cylinder. If compression is excessive, soil core may not be a valid sample for analysis. Rock fragments in the soil interfere
with core collection. Dry or hard soils often shatter when hammering the cylinder into the soil. Pressing the cylinder into the soil reduces the risk of shattering the sample.

If soil cracks are present, select the sampling area so that crack space is representative of sample, if possible. If this is not possible, make measurements between the cracks and determine the aerial percentage of total cracks or of cracks in specimen.

4. Safety
   No known hazard exists with this procedure.

5. Equipment
   5.1 Containers, air-tight, tared, with lids
   5.2 Electronic balance, ±0.01-g sensitivity
   5.3 Oven 110 °C
   5.4 Sieve, No. 10 (2 mm-openings)
   5.5 Coring equipment. Sources described in Grossman and Reinsch (2002).

6. Reagents
   None.

7. Procedure
   7.1 Record the empty core weights (CW).
   7.2 Prepare a flat surface, either horizontal or vertical, at the required depth in sampling pit.
   7.3 Press or drive core sampler into soil. Use caution to prevent compaction. Remove core from the inner liner, trim protruding soil flush with ends of cylinder, and place in air-tight container for transport to laboratory. If soil is too loose to remain in the liner, use core sampler without the inner liner and deposit only the soil sample in air-tight container. Moisture cans can also be pushed directly into a prepared face. For fibrous organic materials, trim sample to fit snugly into a moisture can.
   7.4 Dry core in an oven at 110 °C until weight is constant. Record oven-dry weight (ODW).
   7.5 Measure and record cylinder volume (CV).
   7.6 If sample contains rock fragments, wet-sieve sample through a 2-mm sieve. Dry and weigh the rock fragments that are retained on sieve. Record weight of rock fragments (RF). Determine density of rock fragments (PD).
8. Calculations

\[
Db = \frac{(ODW - RF - CW)}{[CV - (RF / PD)]}
\]

where:
\(Db\) = Bulk density of <2-mm fabric at sampled, field water state (g cm\(^{-3}\))
\(ODW\) = Oven-dry weight
\(RF\) = Weight of rock fragments
\(CW\) = Empty core weight
\(CV\) = Core volume
\(PD\) = Density of rock fragments

9. Report

Bulk density is reported to the nearest 0.01 g cm\(^{-3}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Water Retention (3C)

Water retention is defined as the soil water content at a given soil water suction. By varying the soil suction and recording the changes in soil water content, a water retention function or curve is determined. This relationship is dependent on particle-size distribution, clay mineralogy, organic matter, and structure or physical arrangement of the particles as well as hysteresis, i.e., whether the water is absorbing into or desorbing from the soil. The data collected in these procedures are from water desorption. Water retention or desorption curves are useful directly and indirectly as indicators of other soil behavior traits, such as drainage, aeration, infiltration, plant-available water, and rooting patterns (Topp et al., 1993).

Two desorption procedures are commonly used to measure water retention: a suction method and a pressure method. The KSSL uses the pressure method (U.S. Salinity Laboratory Staff, 1954) with either a pressure-plate or pressure-membrane extractor. Methods 3C1a-e1 (pressure-plate extraction) are used to determine water retention at 6, 10, 33, 100, or 200 kPa, respectively (0.06, 0.1, ⅓, 1, or 2 bar, respectively) for sieved, <2-mm, air-dry soil samples of non-swelling soils, loamy sand or coarser soil, and for some sandy loams. Methods 3C1a-d2
and 3C1a-d3 (pressure-plate extractions) are used to measure water retention of natural clods or cores that have been equilibrated at 6, 10, 33, or 100 kPa. Methods 3C1a-d2 and 3C1a-d3 are usually used in conjunction with the bulk density method 3B1b.

Method 3C1c4 (pressure-plate extraction) is used to determine the water retention of a clod equilibrated at 33-kPa, air-dried, and re-equilibrated. The resulting data are called rewet water-retention data and are usually used in conjunction with the rewet bulk density data in method 3B1d to estimate changes in physical properties of a soil as it undergoes wetting and drying cycles. Method 3C2a1a (pressure-membrane extraction) is used to determine water retention at 1500 kPa (15 bar) for <2-mm (sieved), air-dry soil samples. Method 3C2a1b is used to measure water retention at 1500 kPa for <2-mm (sieved), field-moist soil samples. Method 3C3 is used to determine field water content at the time of sampling for cores, clods, or bulk samples.

References


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Water Retention (3C)
Pressure-Plate Extraction (3C1)

6, 10, 33, 100, or 200 kPa (3C1a-e)

<2-mm (Sieved), Air-Dry (3C1a-e1a)

1. Application

The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate unsaturated hydraulic conductivity. Methods 3C1a-e1a (pressure-plate extraction) are used to determine water retention at 6, 10, 33, 100, or 200 kPa, respectively, for <2-mm (sieved), air-dry soil samples of non-swelling soils, loamy sand or coarser soil, and for some sandy loams.

2. Summary of Method

The pressure desorption method (U.S. Salinity Laboratory Staff, 1954) is used. A sample of <2-mm (sieved), air-dry soil is placed in a retainer ring sitting on a porous ceramic plate in a pressure-plate extractor. The plate is covered with
water to wet the samples by capillarity. The sample is equilibrated at the specified pressures. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The gravimetric water content is determined.

3. Interferences

A leak in the pressure extractor prevents equilibration of samples. Check for outflow air to verify that the pressure-plate extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity.

After extended use, the porous ceramic plate becomes clogged and water outflow is restricted. Clean the plate by flushing it sequentially with 500 mL of 10% H₂O₂, 1000 mL of 1 N HCl, and 500 mL of reverse osmosis (RO) water. The solutions are pulled through the plate with a vacuum, and the waste is captured in a trap.

Check the rubber membrane on the bottom of the plate for leaks. Inflate the membrane and then submerge it in water. If air bubbles escape from the membrane, remove the plate from service.

Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986). Water retention data for soils with expansive clay is overestimated when sieved samples are used in place of natural soil fabric for tensions of 6, 10, and 33 kPa, respectively (Young and Dixon, 1966).

Aerated 0.005 M CaSO₄ has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).

4. Safety

High pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure extractor lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1 Pressure-plate extractor with porous ceramic plate
5.2 Electronic balance, ±0.01-g sensitivity
5.3 Oven, 110 °C
5.4 Pressure source, regulator, and gauge
5.5 Retainer rings, 10-mm high and 50-mm diameter
5.6 Metal weighing cans with lids
6. Reagents

6.1 Reverse osmosis (RO) water

6.2 Hydrogen peroxide ($H_2O_2$), 10% solution. Dilute 333 mL of 30% $H_2O_2$, technical grade, in 1 L of RO water.

6.3 Hydrochloric acid (HCl), 1 N. Dilute 83.3 mL of concentrated HCl in 1 L of RO water.

6.4 Ethyl alcohol, 95%, technical grade

7. Procedure

7.1 Saturate the ceramic plate by applying RO water through the adapter and apply enough pressure so that the rubber membrane is bulging a few centimeters. Care should be taken to remove all air.

7.2 Place the saturated ceramic plate in a pressure-plate extractor. Place retainer rings on the ceramic plate.

7.3 Fill retaining ring with 10 to 15 g of <2-mm or fine-grind, air-dry soil. Include a quality control (QC) sample in each pressure-plate extractor.

7.4 Add enough water to cover the ceramic plate but not to cover the rings. Continue to add water until all samples have moistened by capillarity. If samples do not moisten, apply ethyl alcohol to the surface of the samples. Close the apparatus and let stand overnight.

7.5 Apply the specified pressure. Monitor the outflow tube for water discharge. Periodically submerge the outflow tube in water to monitor for air bubbles that indicate ceramic plate failure. Samples are equilibrated when water ceases to emit from the outflow tube. The outflow tube can be submerged under water in a buret to measure when water ceases to emit from the outflow tube.

7.6 When samples have equilibrated, quickly transfer the samples to tared water cans ($M_c$), cover with lids, and record the weights ($M_{s+w}$).

7.7 Remove lids, place samples in oven, and dry at 110 °C overnight. Record weights ($M_s$).

8. Calculations

\[ H_2O \text{ } \% = 100 \times \frac{[(M_{s+w} - M_s) / (M_s - M_c)]]}{\}

where:

- $H_2O \text{ } \% =$ Percent gravimetric water content
- $M_{s+w}$ = Weight of solids + $H_2O$ + container
- $M_s$ = Weight of solids + container
- $M_c$ = Weight of container
9. Report
Report water content to the nearest 0.1 percent.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Water Retention (3C)
Pressure-Plate Extraction (3C1)
6, 10, 33, or 100 kPa (3C1a-d)
Natural Clods (3C1a-d2)

1. Application
The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate unsaturated hydraulic conductivity. Methods 3C1a-d2 are used to determine the water retention of natural clods at 6, 10, 33, or 100 kPa, respectively.

2. Summary of Method
The pressure desorption method (U.S. Salinity Laboratory Staff, 1954) is used. Natural clods are placed on a tension table and equilibrated at a 5-cm tension at the base of the sample. The clods are then transferred to a porous ceramic plate, which is placed in a pressure-plate extractor. The sample is equilibrated at the
specified pressures. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The gravimetric water content is determined.

3. Interferences

A leak in the pressure extractor prevents equilibration of samples. Check outflow air to verify that each pressure-plate extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity.

After extended use, the porous ceramic plate becomes clogged and water outflow is restricted. Clean the plate by flushing it sequentially with 500 mL of 10% H₂O₂, 1000 mL of 1N HCl, and 500 mL of RO water. The solutions are pulled through the plate with a vacuum, and the waste is captured in a trap.

Check the rubber membrane on the bottom of the plate for leaks. Inflate the membrane and then submerge it in water. If air bubbles escape from the membrane, remove the plate from service.

Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986).

Aerated 0.005M CaSO₄ has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

High pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure-apparatus lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1  Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate (fig. 3C1a-1)
5.3 Pressure source, regulator, and gauge
5.4 Oven, 110 °C
5.5 Clothespins
5.6 Knife
5.7 Tile cut-off saw with diamond blade
5.8 Desiccator with ceramic plate
5.9 Vacuum, 80-kPa (0.8 bar)
5.10 Needle probe
5.11 Sieve, No. 10 (2-mm openings)
5.12 Hot plate
5.13 Fume hood
5.14 Reinforced paper towels with nylon fibers, GSA
5.15 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.
5.16 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.17 Wire. The KSSL uses a 28-awg coated copper wire.

Figure 3C1a-1.—Clods in pressure-plate extractor following saturation.
6. Reagents

6.1 Reverse osmosis (RO) water

6.2 Hydrogen peroxide ($\text{H}_2\text{O}_2$), 10% solution. Dilute 333 mL of 30% $\text{H}_2\text{O}_2$, technical grade, in 1 L of RO water.

6.3 Hydrochloric acid (HCl), 1 N. Dilute 83.3 mL of concentrated HCl in 1 L of RO water.

6.4 Ethyl alcohol, 95%, technical grade

6.5 Acetone (2-propanone; dimethyl ketone)

6.6 Dow Saran F-310 resin, available from Dow Chemical Company

6.7 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a wooden stick. In the laboratory, stir solvent with a non-sparking, high speed stirrer while slowly adding resin. Stir plastic lacquer for 30 min at 25 °C. Store plastic lacquer in covered plastic or steel containers.

6.9 Microbicide, for tension table, Fisher Scientific

7. Procedure

7.1 This procedure is usually combined with the bulk density method (3B1b).

7.2 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod. Record the weight of the tag and wire (TAG). Loop fine copper wire around the clod, leaving a tail to which the round stock tag is attached. Record the weight of the clod (CC1).

7.3 Dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 55 min and then reweigh the clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time, then reweigh the clod and record the weight (CC2).

7.4 Cut a flat surface on the clod with a tile saw. It is necessary to wet the clod above the initial desorption point. This is accomplished by placing the cut clod surface on a tension table that is maintained at 5-cm tension. Periodically check the clod for equilibrium (clod has wetted up). Determination can be made by inserting a needle probe, touching, or comparing weight. When the clod has reached equilibrium, remove the clod and record the weight (WSC).
7.5 If cut clod does not absorb water, place clod in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of clod in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until clod has equilibrated at saturation. Remove the clod and record the weight (WSC).

7.6 Saturate the ceramic plate by applying RO water through the adapter and apply enough pressure so that the rubber membrane is bulging a few centimeters. Care should be taken to remove all air.

7.7 Place the saturated ceramic plate in a pressure-plate extractor. To provide good contact between the clod and ceramic plate, cover the ceramic plate with paper towels and saturate with water. Place the cut surface of the clod on the paper towel. Prepare a saturated, sieved soil as a quality control (QC) sample. Place several retaining rings in the extractor. Fill the retaining rings with the soil standard. Close the container and secure lid.

7.8 Apply gauged air pressure of 6, 10, 33, or 100 kPa (fig. 3C1a-2). If more than one water retention point is requested, begin with the lowest pressure. Periodically submerge the outflow tube in water to monitor for air bubbles that indicate ceramic plate failure. Samples are equilibrated when water ceases to emit from the outflow tube. The outflow tube can be submerged under water in a buret to measure when water ceases to emit from the

Figure 3C1a-2.—Pressure-plate extraction at 33 kPa for clods.
outflow tube. When water stops discharging from the outflow tube, the clod is at equilibrium. Determine the gravimetric water content of the QC. If the water content of the QC is more than twice the standard deviation, apply pressure for additional time. Recheck the QC. If the water content of the QC is less than twice the standard deviation, rewet the clods and desorb again. If the water content of the QC is within acceptable limits, then the apparatus has functioned properly.

7.9 Remove the clod and record the weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, re-equilibrate the clod on the tension table and repeat the desorption process. If additional water retention points are requested, then repeat the desorption process at the next higher pressure. When the clod is equilibrated at 33 kPa and bulk density is to be measured, continue with method 3B1b.

7.10 Dry the clod in an oven at 110 °C overnight and record oven-dry weight (WODC).

7.11 If the clods contains >5% in rock fragments by weight, remove the rock fragments from the clod. This determination of the percent rock fragments is based on particle-size data of >2-mm fraction. Submerge the remaining soil material in a beaker of water and place on a hot plate. Use a fume hood. Boil ≈1 h. The plastic coating loosens from soil material upon heating. Remove beaker from the hot plate. Allow to cool. Discard plastic coating.

7.12 Wet sieve the cool soil through a 2-mm sieve. Dry and record the weight (RF) of the rock fragments that are retained on the sieve. If the rock fragments have properties similar to those of the soil sample, do not correct the clod mass for the rock fragments.

8. Calculations

8.1 \[ \text{H}_2\text{O} \% = \left(\frac{(\text{WMC} - \text{MPC}) - (\text{WODC} - \text{ODPC}) \times 100}{(\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG})}\right) \]

where:
- \( \text{H}_2\text{O} \% \) = Percent gravimetric water content
- \( \text{WMC} \) = Weight of equilibrated, coated clod
- \( \text{WODC} \) = Weight of oven-dry coated clod
- \( \text{RF} \) = Weight of rock fragments
- \( \text{ODPC} \) = MPC × 0.85, weight of oven-dry plastic coat
- \( \text{TAG} \) = Weight of tag and wire

8.2 \[ \text{MPC} = \left(\left(\text{CC2} - \text{CC1}\right) + \text{FCE}\right) \times \text{RV}\] 

where:
- \( \text{MPC} \) = Weight of plastic coat before oven-drying
CC2 = Weight of clod after three laboratory plastic coats
CC1 = Weight of clod before three laboratory plastic coats
RV = Percent estimate of remaining clod volume after cutting to obtain flat surface (∼80%)

8.3 \[ FCE = 1.5 \times \left( \frac{CC2 - CC1}{3} \right) \]
where:
\[ FCE = \text{Estimate of field-applied plastic coat, if applied} \]

9. Report
Report water content to the nearest 0.1 percent.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

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Water Retention (3C)
Pressure-Plate Extraction (3C1)
   6, 10, 33, or 100 kPa (3C1a-d)
Soil Cores (3C1a-d3)

1. Application
The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate
unsaturated hydraulic conductivity. Methods 3C1a-d3 are used to determine the water retention of soil cores at 6, 10, 33, or 100 kPa, respectively.

2. Summary of Method

The pressure desorption procedure (U.S. Salinity Laboratory Staff, 1954) is used. A metal cylinder is pressed or driven into the soil. Upon removal from the soil, the cylinder extracts a sample of known volume. The sample weight is recorded. The sample is dried in an oven and then weighed. Soil core is placed on a tension table and equilibrated at a 5-cm tension at the base of the sample. The cores are then transferred to a porous ceramic plate, which is placed in a pressure-plate extractor. The sample is equilibrated at the specified pressures. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The gravimetric water content is determined.

3. Interferences

A leak in the pressure extractor prevents equilibration of samples. Check for outflow air to verify that the extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity.

After extended use, the porous ceramic plate becomes clogged and water outflow is restricted. Clean the plate by flushing it sequentially with 500 mL of 10% \( \text{H}_2\text{O}_2 \), 1000 mL of 1 \( \text{N} \text{HCl} \), and 500 mL of RO water. The solutions are pulled through the plate with a vacuum, and the waste is captured in a trap.

Check the rubber membrane on the bottom of the plate for leaks. Inflate the membrane and then submerge it in water. If air bubbles escape from the membrane, remove the plate from service.

Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986).

Compaction of the sample during the sampling process is a common problem. Compression can be observed by comparing the soil elevation inside the cylinder with the original soil surface outside the cylinder. If compression is excessive, soil core may not be a valid sample for analysis. Rock fragments in the soil interfere with core collection. Dry or hard soils often shatter when hammering the cylinder into the soil. Pressing the cylinder into the soil reduces the risk of shattering the sample.

\( \text{Aerated } 0.005 \text{ M CaSO}_4 \) has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).
4. Safety

High pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure-apparatus lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate
5.3 Pressure source, regulator, and gauge
5.4 Oven, 110 °C
5.5 Desiccator with ceramic plate
5.6 Vacuum, 80-kPa (0.8 bar)
5.7 Needle probe
5.8 Sieve, No. 10 (2-mm openings)
5.9 Fume hood
5.10 Coring equipment. Sources described in Blake and Hartge (1986).
5.11 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.
5.12 Reinforced paper towels with nylon fibers, GSA

6. Reagents

6.1 Reverse osmosis (RO) water
6.2 Hydrogen peroxide (H$_2$O$_2$), 10% solution. Dilute 333 mL of 30% H$_2$O$_2$, technical grade, in 1 L of RO water.
6.3 Hydrochloric acid (HCl), 1 N. Dilute 83.3 mL of concentrated HCl in 1 L of RO water.
6.4 Alcohol
6.5 Microbicide, for tension table, Fisher Scientific

7. Procedure

7.1 This procedure can be combined with the bulk density method (3B1b).
7.2 Record the weight (CW) of the sampling cylinders.
7.3 Prepare a flat surface in the sampling pit, either horizontal or vertical, at the required depth. Press or drive the core sampler into the soil. Use caution to prevent compaction. Remove the core from the sample holder, trim the protruding soil flush with the cylinder ends, and place core in an air-tight container for transport to the laboratory.
7.4 To help ensure core remains intact, place cheesecloth on the bottom of the core. It is necessary to wet the core above the initial desorption point. This is accomplished by placing the flat core surface on a tension table maintained at 5-cm tension. Periodically check the core for equilibrium (core has wetted up). Determination can be made by inserting a needle probe, touching, or comparing weight. When the core has reached equilibrium, remove the core and record the weight (WSC).

7.5 If core does not absorb water, place core in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of core in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until core has equilibrated at saturation. Remove the core and record the weight (WSC).

7.6 Saturate the ceramic plate by applying RO water through the adapter and apply enough pressure so that the rubber membrane is bulging a few centimeters. Care should be taken to remove all air.

7.7 Place the saturated ceramic plate in a pressure-plate extractor. To provide good contact between the core and ceramic plate, cover the ceramic plate with paper towels. Place the flat core surface on the paper towel. Prepare a saturated, sieved soil as a quality control (QC) sample. Place several retaining rings in the extractor. Fill the retaining rings with the soil standard. Close the container and secure lid.

7.8 Apply gauged air pressure of 6, 10, 33, or 100 kPa. If more than one water retention point is requested, begin with the lowest pressure. When water stops discharging from the outflow tube, the core is at equilibrium. Determine the gravimetric water content of the standard. If the water content of the QC is more than twice the standard deviation, apply pressure for additional time. Recheck the QC. If the water content of the QC is less than twice the standard deviation, rewet the cores and desorb again. If the water content of the QC is within acceptable limits, then the apparatus has functioned properly.

7.9 Remove core and record the weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, re-equilibrate core on the tension table and repeat the desorption process. If additional water retention points are requested, then repeat the desorption process at the next higher pressure. When the core is equilibrated at 33 kPa and bulk density is to be measured, continue with method 3B1b.

7.10 Dry core in an oven at 110 °C overnight and record oven-dry weight (WODC).

7.11 If sample contains rock fragments, wet sieve the sample through a 2-mm sieve. Dry and weigh the rock fragments that are retained on the sieve. If the rock fragments have properties similar to those of the soil sample, do
not correct the clod mass for the rock fragments. Record the weight of the rock fragments (RF).

8. Calculations

\[ H_2O \% = 100 \times \frac{(WMC - WODC)}{(WODC - CW - RF)} \]

where:
- \( H_2O \% \) = Percent gravimetric water content
- WMC = Weight of solids + H\(_2\)O + container
- CW = Weight of solids + container
- WODC = Weight of container
- RF = Weight of rock fragments

9. Report

Report water content to the nearest 0.1 percent.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Water Retention (3C)
Pressure-Plate Extraction (3C1)
  33 kPa (3C1c)
  Rewet (3C1c4)

1. Application
   The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate unsaturated hydraulic conductivity. The rewet water retention data (3C1c4) are used in conjunction with the rewet bulk density (3B1d) to estimate the change in physical properties of a soil as it undergoes wetting and drying cycles.

2. Summary of Method
   The pressure desorption method (U.S. Salinity Laboratory Staff, 1954) is used. Natural clods are placed on a tension table and equilibrated at a 5-cm tension at the base of the sample. The clods are then transferred to a porous ceramic plate, which is placed in a pressure-plate extractor. The sample is equilibrated at 33 kPa. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The equilibrated clod weight is recorded. The clod is air dried and then placed on a tension table and desorbed again. After the second equilibration, the gravimetric water content is determined.

3. Interferences
   A leak in the pressure extractor prevents equilibration of samples. Check outflow air to verify each pressure-plate extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity.
   After extended use, the porous ceramic plate becomes clogged and water outflow is restricted. Clean the plate by flushing it sequentially with 500 mL of 10% \( \text{H}_2\text{O}_2 \), 1000 mL of 1 \( \text{N} \) HCl, and 500 mL of RO water. The solutions are pulled through the plate with a vacuum, and the waste is captured in a trap.
   Check the rubber membrane on the bottom of the plate for leaks. Inflate the membrane and then submerge it in water. If air bubbles escape from the membrane, remove the plate from service.
   Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986).
   Aerated 0.005 \( M \text{CaSO}_4 \), has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in
samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

High pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure-plate lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate
5.3 Pressure source, regulator, and gauge
5.4 Oven, 110 °C capability
5.5 Clothespins
5.6 Knife
5.7 Tile cut−off saw with diamond blade
5.8 Desiccator with ceramic plate
5.9 Vacuum, 80-kPa (0.8 bar)
5.10 Needle probe
5.11 Sieve, No. 10 (2−mm openings)
5.12 Hot plate
5.13 Fume hood
5.14 Reinforced paper towels with nylon fibers, GSA
5.15 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.
5.16 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.17 Wire. The KSSL uses a 28-awg coated copper wire.
5.18 Retainer rings, 10-mm high and 50-mm diameter
6. Reagents

6.1 Reverse osmosis (RO) water

6.2 Hydrogen peroxide ($\text{H}_2\text{O}_2$), 10% solution. Dilute 333 mL of 30% $\text{H}_2\text{O}_2$, technical grade, in 1 L of RO water.

6.3 Hydrochloric acid (HCl), 0.05 N. Dilute 8 mL of concentrated HCl in 1 L of RO water.

6.4 Ethyl alcohol, 95%, technical grade

6.5 Acetone (2-propanone; dimethyl ketone)

6.6 Dow Saran F-310 resin, available from Dow Chemical Company

6.7 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a wooden stick. In the laboratory, stir solvent with a non-sparking, high speed stirrer while slowly adding resin. Stir plastic lacquer for 15 min at 25 °C. Store plastic lacquer in covered plastic or steel containers.

6.8 Microbicide, for tension table, Fisher Scientific

7. Procedure

7.1 This procedure is usually used in conjunction with the bulk density method 4A1i.

7.2 Prepare a round stock tag with sample identification number. Cut the copper wire. Record the weight of the tag and wire (TAG). Loop fine copper wire around the clod, leaving a tail to which the round stock tag is attached. Record the weight of the clod (CC1).

7.3 Dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 55 min and then reweigh the clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time, then reweigh the clod and record the weight (CC2).

7.4 Cut a flat surface on the clod with a tile saw.

7.5 It is necessary to wet the clod above the initial desorption point. This is accomplished by placing the cut clod surface on a tension table that is maintained at 5-cm tension. Periodically check the clod for equilibrium (clod has wetted up). Determination can be made by inserting a needle probe, touching, or comparing weight. When the clod has reached equilibrium, remove the clod and record the weight (WSC).
7.6 If cut clod does not absorb water, place clod in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of clod in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until clod has equilibrated at saturation. Remove the clod and record the weight (WSC).

7.7 Saturate the ceramic plate by applying RO water through the adapter and apply enough pressure so that the rubber membrane is bulging a few centimeters. Care should be taken to remove all air.

7.8 To provide good contact between the clod and ceramic plate, cover the ceramic plate with paper towels and saturate with water. Place the cut surface of the clod on the paper towel. Prepare a saturated, sieved soil as a quality control (QC) sample. Place several retaining rings in the extractor. Fill the retaining rings with the soil standard. Close the container and secure lid.

7.9 Apply gauged air pressure of 6, 10, 33, or 100 kPa. If more than one water retention point is requested, begin with the lowest pressure. When water stops discharging from the outflow tube, the clod is at equilibrium. Determine the gravimetric water content of the QC. If the water content of the QC is more than twice the standard deviation, apply pressure for additional time. Recheck the QC. If the water content of the QC is less than twice the standard deviation, rewet the clods and desorb again. If the water content of the QC is within acceptable limits, then the apparatus has functioned properly.

7.10 Remove the clod and record the weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, re-equilibrate the clod on the tension table and repeat the desorption process. If additional water retention points are requested, then repeat the desorption process at the next higher pressure. When the clod is equilibrated at 33 kPa and bulk density is to be measured, continue with method 3B1d.

7.11 Air-dry the clod at room temperature (≈20 to 25 °C) for 4 to 6 days. Dry the clods at 40 to 50 °C for 2 to 3 days or until weights are constant.

7.12 Repeat steps 7.5, 7.6, 7.7, and 7.8. Record clod weight after equilibration (WMC2). Determine bulk density as described in method 3B1d.

7.13 Dry the clod in oven at 110 °C overnight and record oven-dry weight (WODC).

7.14 If the clod contains >5% in rock fragments by weight, remove the rock fragments from the clod. This determination of the percent rock fragments is based on particle-size data of >2-mm fraction. Submerge the remaining soil material in a beaker of water and place on a hot plate. Use a fume hood. Boil =1 h. The plastic coating loosens from soil material upon
heating. Remove beaker from the hot plate. Allow to cool. Discard plastic coating.

7.15 Wet sieve the cool soil through a 2-mm sieve. Dry and record the weight (RF) of the rock fragments that are retained on the sieve. If the rock fragments have properties similar to those of the soil sample, do not correct the clod mass for the rock fragments.

8. Calculations

8.1 \[ \text{H}_2\text{O} \% = \frac{[(\text{WMC} - \text{MPC}) - (\text{WODC} - \text{ODPC}) \times 100]}{\text{WODC} - \text{RF} - \text{ODPC} - \text{TAG}} \]

where:
- H\(_2\)O \% = Percent gravimetric water content
- WMC = Weight of equilibrated, coated clod
- WODC = Weight of oven-dry coated clod
- RF = Weight of rock fragments
- ODPC = MPC \times 0.85, weight of oven-dry plastic coat
- TAG = Weight of tag and wire

8.2 \[ \text{MPC} = \frac{[(\text{CC2} - \text{CC1}) + \text{FCE}] \times \text{RV}}{\text{RV}} \]

where:
- MPC = Weight of plastic coat before oven-drying
- CC2 = Weight of clod after three laboratory plastic coats
- CC1 = Weight of clod before three laboratory plastic coats
- RV = Percent estimate of remaining clod volume after cutting to obtain flat surface (≈80%)

8.3 \[ \text{FCE} = 1.5 \times \frac{[(\text{CC2} - \text{CC1})]}{3} \]

where:
- FCE = Estimate of field applied plastic coat, if applied

8.4 \[ \text{H}_2\text{O}_r \% = \frac{[(\text{WMC}2 - \text{MPC}2) - (\text{WODC} - \text{ODPC}2)] \times 100}{\text{WODC} - \text{RF} - \text{ODPC}2 - \text{TAG}} \]

where:
- H\(_2\)O\(_r\) \% = Percent water weight retained at 33-kPa tension after rewetting
- WMC2 = Weight of equilibrated, coated clod after rewetting
- MPC2 = Weight of moist plastic coat after rewetting. Same as MPC unless additional plastic coats were added.
- OPC2 = MPC2 \times 0.85, weight of oven-dry plastic coat
9. Report
   Report water content to the nearest 0.1 percent.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

Water Retention (3C)
Pressure-Plate Extraction (3C1)
   33 kPa (3C1c)
   Reconstituted (3C1c5)

1. Application
   The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate unsaturated hydraulic conductivity. Method 3C1c5 is used to determine the water retention of a reconstituted clod at 33 kPa.

2. Summary of Method
   The pressure desorption method (U.S. Salinity Laboratory Staff, 1954) is used. Natural clods are placed on a tension table and equilibrated at a 5-cm tension at the base of the sample. The clods are then transferred to a porous ceramic plate, which is placed in a pressure-plate extractor. The sample is equilibrated at the specified pressures. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The gravimetric water content is determined.
3. Interferences

A leak in the pressure extractor prevents equilibration of samples. Check outflow air to verify that each pressure-plate extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity.

After extended use, the porous ceramic plate becomes clogged and water outflow is restricted. Clean the plate by flushing it sequentially with 500 mL of 10% H$_2$O$_2$, 1000 mL of 1 N HCl, and 500 mL of RO water. The solutions are pulled through the plate with a vacuum, and the waste is captured in a trap.

Check the rubber membrane on the bottom of the plate for leaks. Inflate the membrane and then submerge it in water. If air bubbles escape from the membrane, remove the plate from service.

Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986).

Aerated 0.005 M CaSO$_4$ has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).

4. Safety

Acetone is highly flammable. Neither open flames nor nearby operation of electrical equipment is permitted when acetone is used. Avoid inhalation or physical contact. Use with adequate ventilation. In closed buildings, use a fume hood. Keep in tightly closed containers. Use safety glasses, proper gloves, and a lab coat. Wash hands immediately after handling acetone. Additional information on the safe handling of acetone is available in the material safety data sheets (MSDS).

Saran F-310 resin decomposes rapidly at temperatures >200 °C, releasing hydrogen chloride gas. Avoid contact with Fe, Zn, Cu, and Al in solution. Avoid all contact with strong bases.

High pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure-apparatus lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Pressure-plate extractor with porous ceramic plate
5.3 Pressure source, regulator, and gauge
5.4 Oven, 110 °C
5.5 Clothespins
5.6 Knife
5.7 Tile cut-off saw with diamond blade
5.8 Desiccator with ceramic plate
5.9 Vacuum, 80-kPa (0.8 bar)
5.10 Needle probe
5.11 Sieve, No. 10 (2-mm openings)
5.12 Hot plate
5.13 Fume hood
5.14 Reinforced paper towels with nylon fibers, GSA
5.15 Tension table. The KSSL constructs a tension table by placing porous firebricks, covered with reinforced paper towels, in a tub of water.
5.16 Stock tags, 25.4-mm (1-in) diameter paper tag, with metal rim
5.17 Wire. The KSSL uses a 28-awg coated copper wire.
5.18 Retainer rings, 10-mm high. Use 50-mm diameter rings for organic soils and 40-mm diameter rings for all other soils.

6. Reagents
6.1 Reverse osmosis (RO) water
6.2 Hydrogen peroxide (H$_2$O$_2$), 10% solution. Dilute 333 mL of 30% H$_2$O$_2$, technical grade, in 1 L of RO water.
6.3 Hydrochloric acid (HCl), 1 N. Dilute 83.3 mL of concentrated HCl in 1 L of RO water.
6.4 Ethyl alcohol, 95%, technical grade
6.5 Acetone (2-propanone; dimethyl ketone)
6.6 Dow Saran F-310 resin, available from Dow Chemical Company
6.7 Plastic lacquer. Prepare plastic lacquer with resin-to-solvent ratios of 1:4 and 1:7 on a weight basis. Fill a 3.8-L (1-gal) metal paint can with 2700 ±200 mL of solvent (fill to the bottom of handle rivet). Add 540 g or 305 g of resin to make 1:4 or 1:7 plastic lacquer, respectively. Use the 1:4 plastic lacquer for the initial field and laboratory coatings. Use the 1:7 plastic lacquer for the last two laboratory coatings. The 1:7 plastic lacquer is used to conserve the resin and to reduce cost. In the field, mix solvent with a wooden stick. In the laboratory, stir solvent with a non-sparking, high speed stirrer while slowly adding resin. Stir plastic lacquer for 30 min at 25 °C. Store plastic lacquer in covered plastic or steel containers. Acetone may be substituted for MEK (methyl ethyl ketone) as a solvent.
6.8 Microbicide, for tension table, Fisher Scientific
7. Procedure

7.1 This procedure is usually combined with the bulk density method (3B2b).

7.2 Prepare a round stock tag with sample identification number. Cut the copper wire and loop around the clod. Record the weight of the tag and wire (TAG). Loop fine copper wire around the clod, leaving a tail to which the round stock tag is attached. Record the weight of the clod (CC1).

7.3 Dip the clod in the 1:4 plastic lacquer. Wait 7 min and then dip the clod in the 1:7 plastic lacquer. Wait 12 min and then dip the clod in the 1:7 plastic lacquer. Wait 55 min and then reweigh the clod. If the clod has adsorbed >3% in plastic by clod weight or smells excessively of solvent, allow longer drying time, then reweigh the clod and record the weight (CC2).

7.4 Cut a flat surface on the clod with a tile saw. It is necessary to wet the clod above the initial desorption point. This is accomplished by placing the cut clod surface on a tension table that is maintained at 5-cm tension. Periodically check the clod for equilibrium (clod has wetted up). Determination can be made by inserting a needle probe, touching, or comparing weight. When the clod has reached equilibrium, remove the clod and record the weight (WSC).

7.5 If cut clod does not absorb water, place clod in a desiccator on a water-covered plate with a 0-cm tension. Submerge only the surface of clod in the water. Add a few mL of alcohol. Use in-house vacuum and apply suction until clod has equilibrated at saturation. Remove the clod and record the weight (WSC).

7.6 Saturate the ceramic plate by applying RO water through the adapter and apply enough pressure so that the rubber membrane is bulging a few centimeters. Care should be taken to remove all air.

7.7 Place the saturated ceramic plate in a pressure-plate extractor. To provide good contact between the clod and ceramic plate, cover the ceramic plate with paper towels and saturate with water. Place the cut surface of the clod on the paper towel. Prepare a saturated, sieved soil as a quality control (QC) sample. Place several retaining rings in the extractor. Fill the retaining rings with the soil standard. Close the container and secure lid.

7.8 Apply gauged air pressure of 33 kPa. If more than one water retention point is requested, begin with the lowest pressure. Periodically submerge the outflow tube in water to monitor for air bubbles that indicate ceramic plate failure. Samples are equilibrated when water ceases to emit from the outflow tube. The outflow tube can be submerged under water in a buret to measure when water ceases to emit from the outflow tube. When water stops discharging from the outflow tube, the clod is at equilibrium. Determine the gravimetric water content of the QC. If the water content of the QC is more than twice the standard deviation, apply pressure for
additional time. Recheck the QC. If the water content of the QC is less than twice the standard deviation, rewet the clods and desorb again. If the water content of the QC is within acceptable limits, then the apparatus has functioned properly.

7.9 Remove the clod and record the weight (WMC). Compare WMC to WSC. If WMC ≥ WSC, re-equilibrate the clod on the tension table and repeat the desorption process. If additional water retention points are requested, then repeat the desorption process at the next higher pressure. When the clod is equilibrated at 33 kPa and bulk density is to be measured, continue with method 3B1b.

7.10 Dry the clod in an oven at 110 °C overnight and record oven-dry weight (WODC).

7.11 If the clods contains >5% in rock fragments by weight, remove the rock fragments from the clod. This determination of the percent rock fragments is based on particle-size data of >2-mm fraction. Submerge the remaining soil material in a beaker of water and place on a hot plate. Use a fume hood. Boil ≈1 h. The plastic coating loosens from soil material upon heating. Remove beaker from the hot plate. Allow to cool. Discard plastic coating.

7.12 Wet sieve the cool soil through a 2−mm sieve. Dry and record the weight (RF) of the rock fragments that are retained on the sieve. If the rock fragments have properties similar to those of the soil sample, do not correct the clod mass for the rock fragments.

8. Calculations

8.1 \( \text{H}_2\text{O} \% = \left[\frac{(\text{WMC} - \text{MPC}) - (\text{WODC} - \text{ODPC}) \times 100}{(\text{WODC} - \text{RF} - \text{ODPC} - \text{TÅG})}\right] \)

where:
- \( \text{H}_2\text{O} \% \): Percent gravimetric water content
- \( \text{WMC} \): Weight of equilibrated, coated clod
- \( \text{WODC} \): Weight of oven-dry coated clod
- \( \text{RF} \): Weight of rock fragments
- \( \text{ODPC} \): \( \text{MPC} \times 0.85 \), weight of oven-dry plastic coat
- \( \text{TÅG} \): Weight of tag and wire

8.2 \( \text{MPC} = \left\{\left[\left(\text{CC2} - \text{CC1}\right) + \text{FCE}\right] \times \text{RV}\right\} \)

where:
- \( \text{MPC} \): Weight of plastic coat before oven-drying
- \( \text{CC2} \): Weight of clod after three laboratory plastic coats
- \( \text{CC1} \): Weight of clod before three laboratory plastic coats
8.3  \[ \text{FCE} = 1.5 \times \left[ \frac{(\text{CC}2 - \text{CC}1)}{3} \right] \]

where:

\[ \text{FCE} = \text{Estimate of field-applied plastic coat, if applied} \]

9. Report

Report water content to the nearest 0.1 percent.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Water Retention (3C)

Pressure-Membrane Extraction (3C2)

1500-kPa (3C2a)

<2-mm (Sieved), Air-Dry or Field-Moist (3C2a1a-b)

1. Application

The data collected are used for the water retention function, water-holding capacity, pore-size distribution, porosity, and saturated conductivity of a soil sample at specific water contents. The data are also used to calculate unsaturated hydraulic conductivity. Water retention at 1500 kPa for <2-mm (sieved), air-dry and field-moist samples are determined by methods 3C2a1a and 3C2a1b, respectively.
2. Summary of Method

The pressure desorption procedure (U.S. Salinity Laboratory Staff, 1954) is used. A sample of <2-mm (sieved), air-dry or field-moist soil is placed in a retainer ring sitting on a cellulose membrane in a pressure-membrane extractor. The membrane is covered with water to wet the samples by capillarity. The sample is equilibrated at 1500 kPa. The pressure is kept constant until equilibrium is obtained (Klute, 1986). The gravimetric water content is determined.

3. Interferences

A leak in the pressure extractor prevents equilibration of samples. Check outflow air to verify that the pressure membrane extractor is functioning properly and does not leak. Monitor the pressure for stability. Equilibration must be done at constant temperature and humidity. Samples that do not wet by capillarity are moistened with ethyl alcohol.

Laboratory-determined, water retention data are usually higher than field-determined, water retention data because the confining soil pressure is not present in the laboratory (Bruce and Luxmoore, 1986).

Aerated 0.005 \( \text{M} \) \( \text{CaSO}_4 \) has also been recommended (Dane and Hopmans, 2002), especially for fine-textured soils that contain significant amounts of swelling clays. Distilled or deionized water can possibly promote dispersion of clays in samples, and freshly drawn tapwater is often supersaturated with air, affecting the water content at a given pressure head (Dane and Hopmans, 2002).

4. Safety

High-pressure plumbing must be maintained in good working order. Ensure that the pressure is zero before removing bolts from the pressure-apparatus lid. Ensure that the bolts are tightened before applying pressure. Do not drop the lid, which is heavy.

5. Equipment

5.1 Pressure membrane extractor (figs. 3C2a-1 and 3C2a-2)
5.2 Cellulose membrane
5.3 Retainer rings, 10-mm height and 40-mm diameter
5.4 Electronic balance, ±0.01-g sensitivity
5.5 Oven, 110 °C
5.6 Pressure source, regulator, and gauge
5.7 Metal weighing cans, tared, with lids
5.8 Vacuum trap assembly
5.9 Vacuum, 80-kPa (0.8 bar)
Figure 3C2a-1.—Sieved (<2-mm) soil placed in pressure-membrane extractor.

Figure 3C2a-2.—Pressure-membrane extraction at 1500-kPa for <2-mm samples.
6. Reagents
6.1 Ethyl alcohol, 95%, technical grade
6.2 Reverse osmosis (RO) water

7. Procedure
7.1 Submerge a cellulose membrane in RO water for 12 h or more before use. Install the wet cellulose membrane in the pressure extractor.
7.2 Add water and retaining rings. Add enough water to keep membrane moist. Water level should be less than height of retaining rings.
7.3 Fill retaining rings with 10 to 15 g of <2-mm or fine-grind, air-dry or field-moist soil sample. Include a quality control (QC) sample with each plate. Continue to add water until all samples have moistened by capillarity. If samples do not moisten, apply ethyl alcohol to the surface of the sample. Allow ethyl alcohol to evaporate. Cover samples with a sheet of plastic to reduce evaporation, close the extractor, and let stand overnight.
7.4 Remove excess water on the plate with a vacuum and trap assembly.
7.5 Assemble the extractor and uniformly tighten the bolts. Torque the bolts on both sides of the hinge to 138.0 kPa (200 psi). Torque the remaining bolts to 103.5 kPa (150 psi).
7.6 Increase air pressure ≈150 kPa every 15 min until 1500 kPa is reached. The next day, apply the pressure differential (±5 psi). This forces the rubber diaphragm against the top of the samples. The samples are equilibrated when water ceases to emit from the outflow tube.
7.7 At equilibrium, open the extractor and quickly transfer the samples to water cans, cover with lids, and record the weights ($M_{s+w}$).
7.8 Remove the lids, place samples in the oven, and dry at 110 °C overnight. Remove samples from the oven, replace the lids, allow cans to cool to ambient temperature, and record the weights ($M_s$).
7.9 Record the weights of the empty cans ($M_c$).

8. Calculations
$$H_2O \% = 100 \times \left[ \frac{(M_{s+w} - M_s)}{(M_s - M_c)} \right]$$

where:
$H_2O \% =$ Percent gravimetric water content
$M_{s+w}$ = Weight of solids + $H_2O$ + container
$M_s$ = Weight of solids + can
$M_c$ = Weight of container
Gypsiferous/gypseous soils are a special case because gypsum (CaSO₄•2H₂O) loses most of its chemically combined water (crystal water) at 105 °C (Nelson et al., 1978). Properties of gypsiferous/gypseous soils, such as 1500-kPa water content, that are reported on an oven-dry weight basis are converted to include the weight of crystal water in gypsum. Refer to method 3D3 for these conversion calculations. The 1500-kPa water content is corrected when the gypsum content of the soil is >1%. Gypsum content of the soil is determined in method 4E2a1a1.

9. Report
Report water content to the nearest 0.1 percent.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Water Retention (3C)
Field-State (3C3)

1. Application
Field-water content can be determined by weighing, drying, and reweighing a soil sample (3C3). The resulting data are used to estimate the water content at the time of sampling.
2. **Summary of Method**
   Soil samples are collected in the field. The samples are stored in plastic or metal containers to prevent drying and then transported to the laboratory. Gravimetric water content is determined (Gardner, 1986).

3. **Interferences**
   Leaks in the plastic or metal storage containers cause the samples to dry, resulting in an underestimation of the field water content.

4. **Safety**
   Use insulated gloves to remove samples from the oven.

5. **Equipment**
   5.1 Electronic balance, ±0.01-g sensitivity
   5.2 Oven, 110 °C
   5.3 Moisture cans, tared

6. **Reagents**
   None.

7. **Procedure**
   7.1 Collect soil samples in the field. Place samples in airtight, metal or plastic containers.
   7.2 Record sample weight ($M_{s+w}$).
   7.3 Dry sample in an oven at 110 °C overnight. Record oven-dry weight ($M_s$).
   7.4 Record weight of container ($M_c$).

8. **Calculations**
   \[H_2O \% = 100 \times \frac{(M_{s+w} - M_s)}{(M_s - M_c)}\]
   where:
   \(H_2O \%\)=Percent gravimetric water content
   \(M_{s+w}\)=Weight of solids + $H_2O$ + container
   \(M_s\)=Weight of solids + container
   \(M_c\)=Weight of container

9. **Report**
   Report water content to the nearest 0.1 percent.
10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Air-Dry/Oven-Dry Ratio (AD/OD) (3D1)
Field-Moist/Oven-Dry Ratio (FM/OD) (3D2)
Correction for Crystal Water (3D3)

1. Application

Soil properties generally are expressed on an oven-dry weight basis. The calculation of the air-dry/oven-dry (AD/OD) ratio (method 3D1) or field-moist/oven-dry (FM/OD) ratio (method 3D2) is used to adjust all results to an oven-dry basis and, if required in a procedure, to calculate the sample weight that is equivalent to the required oven-dry soil weight. The AD/OD ratio, unless otherwise specified, is determined on a <2-mm sieved sample by method 3D1. Currently, if a <2-mm sample is fine grind (e.g., <180 µm for total C or <75 µm for trace elements) or if a subset of a <2-mm sample is further prepared (≥0.53- and <0.53-µm fractions for hyper particulate organic matter or the 2- to 0.5-mm fraction for aggregate stability), the AD/OD of the <2-mm sieved sample as prepared in method 3D1 is used in KSSL method calculations (if required).

Gypsiferous/gypseous soils are a special case because gypsum (CaSO₄•2H₂O) loses most of the its chemically combined water (crystal water) at 105 °C. Properties of gypsiferous/gypseous soils that are reported on an oven-dry weight basis should be converted to include the weight of crystal water in gypsum. In method 3D1, the AD/OD ratio is calculated. This ratio is used to convert soil properties to an oven-dry basis. In method 3D3, the AD/OD ratio is converted to a crystal water basis (Nelson et al., 1978). The inclusion of weight of crystal water in gypsum allows the properties of gypsiferous/gypseous soils to be compared with those properties of nongypsiferous/nongypseous soils. This conversion also avoids the possible calculation error of obtaining >100% gypsum when the data are expressed on an oven-dry basis (Nelson, 1982).

AD and OD weights are defined herein as constant sample weights obtained after drying at 30 ±5 °C (≈3 to 7 days) and at 110 ±5 °C (≈12 to 16 h), respectively. As a general rule, air-dry soils contain about 1 to 2 percent water and are drier than soils at 1500-kPa water content. FM weight is defined herein as the sample
weight obtained without drying prior to laboratory analysis. In general, these weights are reflective of the water content at the time of sample collection.

2. Summary of Method

A sample is weighed, dried to a constant weight in an oven, and reweighed. The moisture content is expressed as a ratio of the air-dry weight to the oven-dry weight (AD/OD) or as a ratio of field-moist weight to the oven-dry weight (FM/OD). Soil properties of gypsiferous/gypseous soils that are reported on an oven-dry weight basis are converted to include the weight of the crystal water. When reporting the water content of gypsiferous/gypseous soils, the crystal water content must be subtracted from the total oven-dry water content. The AD/OD ratio is corrected to a crystal water basis when the gypsum content of the soil is ≥1%. Gypsum content of the soil is determined in method 4E2a1a1.

3. Interferences

Traditionally, the most frequently used definition for a dry soil is the soil mass after it has come to a constant weight at a temperature of 100 to 110 °C (ASTM, 2012). Many laboratory ovens are not capable of maintaining this prescribed temperature range. Temperatures that are >50 °C may promote oxidation or decomposition of some forms of organic matter.

Samples may not reach a constant weight with overnight drying. Do not add moist samples to an oven with drying samples unless the drying samples have been in the oven for at least 12 to 16 h. Soil samples may adsorb significant amounts of moisture from the atmosphere after cooling. Prompt weighing, i.e., <30 min after samples have cooled, helps to eliminate this problem. During the weighing or drying processes, the non-uniform weight of weighing vessels, sample contamination, or sample loss may lead to erroneous results.

The removal of structural water, most commonly in gypsum, can produce a positive error. When reporting the water content of gypsiferous/gypseous soils, the crystal water content must be subtracted from the total oven-dry water content. Gypsum, hydrous oxides, and amorphous material may be affected.

4. Safety

Use heat resistant gloves to remove weighing containers from a hot oven. No other significant hazard is associated with this procedure. Follow standard laboratory procedures.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Oven, thermostatically controlled, 110 ±5 °C
5.3 Thermometer, 0 to 200 °C
5.4 Tin dishes, 4.5-cm diameter x 3-cm height, with covers
6. Reagents
   No reagents are required for this determination.

7. Procedure
   7.1 Tare the moisture dishes. Record each sample number and associated
dish number.
   7.2 Add 10 to 20 g <2 mm, air-dry soil to each moisture dish for AD/OD
determination. For FM/OD determination, add enough <2 mm or fine-grind,
moist soil to achieve ≈10 to 20 g sample of air-dry soil. Weigh the dish plus
the sample and record the weight to the nearest 1 mg. Place the sample
dish in a drying oven set at 110 ±5 °C. Allow the sample to remain in the
oven overnight (12 to 16 h).
   7.3 Remove the sample dish and allow it to cool before reweighing. Record the
oven-dry weight to the nearest 1 mg.
   7.4 Do not allow the sample dish to remain at room temperature for >30 min
before reweighing.
   7.5 Discard the sample.
   7.6 Refer to the calculations for the correction for crystal water of gypsum in
gypsiferous/gypseous soils.

8. Calculations

Calculations 8.1–8.2 for AD/OD ratio (method 3D1)

8.1 \( \text{AD/OD ratio} = \frac{\text{AD}}{\text{OD}} \)
where:
\( \text{AD} = (\text{Air-dry weight}) - (\text{Tin tare weight}) \)
\( \text{OD} = (\text{Oven-dry weight}) - (\text{Tin tare weight}) \)

8.2 \( \text{H}_2\text{O} = \left[ \frac{(\text{AD} - \text{OD}) \times 100}{\text{OD}} \right] \)
where:
\( \text{H}_2\text{O} = \% \text{ Water content} \)
\( \text{AD} = (\text{Air-dry weight}) - (\text{Tin tare weight}) \)
\( \text{OD} = (\text{Oven-dry weight}) - (\text{Tin tare weight}) \)

Calculations 8.3–8.4 for FM/OD ratio (method 3D2)

8.3 \( \text{FM/OD ratio} = \frac{\text{FM}}{\text{OD}} \)
where:
\( \text{FM} = (\text{Field-moist weight}) - (\text{Tin tare weight}) \)
\( \text{OD} = (\text{Oven-dry weight}) - (\text{Tin tare weight}) \)
8.4 \( H_2O = [(FM - OD) \times 100] / OD \)

where:
- \( FM \) = (Field-moist weight) - (Tin tare weight)
- \( OD \) = (Oven-dry weight) - (Tin tare weight)

**Calculations 8.5–8.6 for gypsum \( H_2O \) correction (method 3D3)**

8.5 \( \frac{AD}{OD}_c = \frac{(AD/OD)_{uc}}{[1 + (Gypsum \times 0.001942)]} \)

where:
- \( \frac{AD}{OD}_c \) = Air-dry/oven-dry ratio, corrected basis, gypsiferous/gypseous soils
- \( \frac{AD}{OD}_{uc} \) = Air-dry/oven-dry ratio, uncorrected basis
- Gypsum = % Gypsum uncorrected (method 4E2a1a1)

8.6 \( H_2O_c = \frac{[H_2O_{uc} - (Gypsum \times 0.1942)]}{[1 + (Gypsum \times 0.001942)]} \)

where:
- \( H_2O_c \) = % Water content, corrected basis, gypsiferous/gypseous soils
- \( H_2O_{uc} \) = % Water content, uncorrected basis (calculation 8.2)
- Gypsum = % Gypsum uncorrected (method 4E2a1a1)

**AD/OD Data Use**

The following equation is used to calculate the weight of air-dry soil needed to provide a given weight of oven-dry soil for other analytical procedures.

8.7 \( AD = \frac{(OD_r)}{[1 - (H_2O/100)]} \)

where:
- \( AD \) = Required weight of air-dry soil
- \( OD_r \) = Desired weight of oven-dry soil
- \( H_2O \) = Percent water determined from AD/OD (calculation 8.2)

9. **Report**

Report the AD/OD and/or FM/OD ratio as a dimensionless value to the nearest 0.01 unit.

10. **Precision and Accuracy**

Precision and accuracy data are available from the KSSL upon request.

11. **References**


Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Coefficient of Linear Extensibility (COLE) (3D4)

Coefficient of linear extensibility (COLE) is a derived value that denotes the fractional change in the clod dimension from a moist to a dry state (Franzmeier and Ross, 1968; Grossman et al., 1968; Holmgren, 1968). COLE may be used to make inferences about shrink-swell capacity and clay mineralogy. The COLE concept does not include irreversible shrinkage, such as that occurring in organic soils and some andic soils. Certain soils with relatively high contents of smectite clay have the capacity to swell significantly when moist and to shrink and crack when dry. This shrink-swell potential is important for soil physical qualities (e.g., large, deep cracks in dry seasons) as well as for genetic processes and soil classification (Buol et al., 1980).

COLE can be expressed as percent, i.e., linear extensibility percent (LEP). LEP = COLE x 100. The LEP is not the same as LE. In soil taxonomy (Soil Survey Staff, 2014), linear extensibility (LE) of a soil layer is the product of the thickness, in centimeters, multiplied by the COLE of the layer in question. The LE of a soil is defined as the sum of these products for all soil horizons (Soil Survey Staff, 2014). Refer to Soil Survey Staff (2014) for additional discussion of LE.

Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Coefficient of Linear Extensibility (COLE) (3D4)

Air-Dry or Oven-Dry to 33-kPa Tension (3D4a)

The KSSL calculates COLE for the whole soil (air-dry or oven-dry to 33-kPa suction) by method 3D4a. COLE is reported in units of cm cm$^{-1}$. Calculate COLE when coarse fragments are present as follows:

$$COLE_{ws} = \left\{ \frac{1}{Cm(D_{b_{33<2mm}}/D_{b_{d<2mm}})+(1-Cm)} \right\}^{\frac{1}{3}} - 1$$

where:

$$COLE_{ws} = \text{Coefficient of linear extensibility on a whole-soil base}$$
$Db_{33<2mm} =$ Bulk density at 33-kPa water content on a $<2$-mm base (g cm$^{-3}$)

$Db_{d<2mm} =$ Bulk Density, oven-dry or air-dry, on a $<2$-mm base (g cm$^{-3}$)

$Cm =$ Coarse fragment (moist) conversion factor

If no coarse fragments are present, $Cm = 1$. If coarse fragments are present, calculate $Cm$ as follows:

$$Cm = \frac{Vol_{<2mm}}{Vol_{whole}}$$

where:

$Vol_{<2mm} =$ Volume moist $<2$-mm fabric (cm$^3$)

$Vol_{whole} =$ Volume moist whole soil (cm$^3$)

Or (alternatively)

$$Cm = \frac{(100 - Vol_{>2mm})}{100}$$

where:

$Vol_{>2mm} =$ Volume percentage of the $>2$-mm fraction

If no coarse fragments are present, $Cm = 1$ and the previous equation reduces as follows:

$$COLE_{ws} = \left(\frac{Db_{d<2mm}}{Db_{33<2mm}}\right)^{\frac{1}{3}} - 1$$

where:

$COLE_{ws} =$ Coefficient of linear extensibility on a whole-soil base

$Db_{d<2mm} =$ Bulk Density, oven-dry or air-dry, on a $<2$-mm base (g cm$^{-3}$)

$Db_{33<2mm} =$ Bulk Density at 33-kPa water content on a $<2$-mm base (g cm$^{-3}$)

References


Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Water Retention Difference (WRD) (3D5)

The calculation of the water retention difference (WRD) is considered the initial step in the approximation of the available water capacity (AWC). WRD does not allow for restriction of roots from the soil layer or osmotic pressure. Usually, the volume of rock fragments is considered a diluent containing no water between the suctions that define WRD. WRD is a calculated value that denotes the volume fraction for water in the whole soil that is retained between 1500-kPa suction and an upper limit of usually 33- or 10-kPa suction. The upper limit (lower suction) is selected so that the volume of water retained approximates the volume of water held at field capacity. The 10-, 33- and 1500-kPa gravimetric water contents are then converted to a whole-soil volume basis by multiplying by the bulk density ($D_b_{33}$) and adjusting downward for the volume fraction of any rock fragments present in the soil. The lower suctions, e.g., 10- or 5-kPa, are used for coarse materials. Refer to Soil Survey Staff Division Staff (1993) and Grossman et al. (1994) for additional discussion on coarse materials and the significance of soil water content at lower suctions and for suggestions related to the selection of these lower suctions for the determination of water retention difference (WRD).

Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Water Retention Difference (WRD) (3D5)

Between 33-kPa and 1500-kPa Tension (3D5a)

The KSSL calculates the WRD between 33- and 1500-kPa suctions in the whole soil by method 3D5a. The WRD is reported as centimeters of water per centimeter of depth of soil (cm cm$^{-1}$), but the numbers do not change when other units, e.g., in$^{-1}$ or ft$^{-1}$, are needed. The WRD with $W_{33}$ as the upper limit is reported as cm cm$^{-1}$. This WRD is calculated on a whole-soil base as follows:

$$WRD_{ws} = [\left(W_{33<2mm} - W_{1500<2mm}\right) \times \left(D_b_{33<2mm}\right) \times Cm] / (P_w \times 100)$$

where:

- $WRD_{ws}$ = Volume fraction (cm$^3$ cm$^{-3}$) of water retained in the whole soil between 33-kPa and 1500-kPa suction, reported in cm cm$^{-1}$
- $W_{33<2mm}$ = Weight percentage of water retained at 33-kPa suction on a <2-mm soil basis
- $W_{1500<2mm}$ = Weight percentage of water retained at 1500-kPa suction on a <2-mm soil basis

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If available, moist 1500-kPa (method 3C2a1b) is the first option in the WRD calculation; otherwise, dry 1500-kPa (method 3c2a1a) is used.

\[ D_{b_{33<2mm}} = \text{Bulk density at 33-kPa water content on a } <2\text{-mm base (g cm}^{-3}) \]

\[ P_w = \text{Density of water (1 g cm}^{-3}) \]

\[ C_m = \text{Coarse fragment material conversion factor} \]

If no coarse fragments are present, \( C_m = 1 \). If coarse fragments are present, calculate \( C_m \) as follows:

\[ C_m = \frac{\text{Vol}_{<2\text{mm}}}{\text{Vol}_{\text{whole}}} \]

where:

\[ \text{Vol}_{<2\text{mm}} = \text{Volume moist } <2\text{-mm fabric (cm}^3) \]

\[ \text{Vol}_{\text{whole}} = \text{Volume moist whole soil (cm}^3) \]

Or (alternatively)

\[ C_m = \frac{(100 - \text{Vol}_{>2\text{mm}})}{100} \]

where:

\[ \text{Vol}_{>2\text{mm}} = \text{Volume percentage of the } >2\text{-mm fraction.} \]

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### Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

#### Water Retention Difference (WRD) (3D5)

**Between 10-kPa and 1500-kPa Tension (3D5b)**

The KSSL also calculates the WRD between 10-kPa (\( W_{10} \)) and 1500-kPa suctions (\( W_{1500} \)) by method 3D5b. This WRD value can be calculated by substituting the \( W_{10} \) in place of \( W_{33} \) in the equation for method 3D5a. The \( W_{10} \) may be used as the upper limit of plant-available water for coarse soil materials.

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### Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

#### Water Retention Difference (WRD) (3D5)

**Between 33-kPa Rewet and 1500-kPa (Air-Dry) Tension (3D5c)**

The KSSL also calculates the WRD between 33-kPa rewet (\( W_r \)) and \( W_{1500} \) by method 3D5c. This WRD value can be calculated by substituting the \( W_r \) in place of \( W_{33} \) in the equation for 3D5a. The \( W_r \) is used for organic materials.
References


Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

1500-kPa Water Content/Total Clay (3D6)

Divide the 1500-kPa water retention (method 3C2a) by the total clay percentage (method 3A1a). This ratio is reported as a dimensionless value. In the past, the ratios of 1500-kPa water:clay have been reported as g g$^{-1}$. For more detailed information on the application of this ratio, refer to Soil Survey Staff (2011, 2014).

Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Total Silt Fraction (3D7)

Total silt is a soil separate with 0.002- to 0.05-mm particle diameter. The KSSL determines the fine silt separate by pipette analysis and the coarse silt separate by difference (3A1a). Total silt is reported as a weight percentage on a <2-mm basis (3D7). For more information on these data, refer to Soil Survey Staff (2011).

Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

Total Sand Fraction (3D8)

Total sand is a soil separate with 0.05- to 2.0-mm particle diameter. The KSSL determines the sand fractions by sieve analysis (3A1a). Total sand is the sum of the very fine sand (VFS), fine sand (FS), medium sand (MS), coarse sand (CS), and very coarse sand fractions VCS). The rationale for five subclasses of sand and the expansion of the texture classes of sand, e.g., sandy loam and loamy sand, is that the sand separates are the most visible to the naked eye and the most detectable by “feel” by the field soil scientist. Total sand is reported as a weight percentage on a <2-mm basis (3D8). For more information on the application of these data, refer to Soil Survey Staff (2011).
Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

2- to 5-mm Fraction (3D9)

The KSSL determines coarse fraction with 2- to 5-mm particle diameter by methods outlined in 3A2. The 2- to 5-mm divisions correspond to the size of opening of the No. 10 and No. 4 screen (4.76 mm), respectively, used in engineering. Coarse fractions with 2- to 5-mm particle diameter correspond to the “fine pebbles” rock-fragment division (Soil Survey Division Staff, 1993). Coarse fractions with 2- to 5-mm particle diameter are reported as a weight percentage on a <75-mm basis (3D9). For more information on coarse fraction with >2-mm particle diameters and application of these data, refer to Soil Survey Staff (2011).

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Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

5- to 20-mm Fraction (3D10)

The KSSL determines coarse fraction with 5- to 20-mm particle diameter by procedures outlined in 3A2. The 5- to 20-mm divisions correspond to the size of opening of the No. 4 screen (4.76 mm) and the ¾-in screen (19.05 mm), respectively, used in engineering. Coarse fractions with 5- to 20-mm particle diameter correspond to the “medium pebbles” rock fragment division (Soil Survey Division Staff, 1993). Coarse fractions with 5- to 20-mm particle diameter are reported as a weight percentage on a <75-mm basis (3D10). For more information on these data, refer to Soil Survey Staff (2011).

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Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

20- to 75-mm Fraction (3D11)

The KSSL determines coarse fraction with 20- to 75-mm particle diameter by procedures outlined in 3A2. The 20- to 75-mm divisions correspond to the size of opening of the ¾-in screen (19.05 mm) and the 3-in screen (76.1 mm), respectively, used in engineering. Coarse fractions with 20- to 75-mm particle diameter correspond to the “coarse pebbles” rock fragment division (Soil Survey Division Staff, 1993). Coarse fractions with 20- to 75-mm particle diameter are reported as a weight percentage on a <75-mm basis (3D11). For more information on these data, refer to Soil Survey Staff (2011).
Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

0.1- to 75-mm Fraction (3D12)

The KSSL determines coarse fractions with 0.1- to 75-mm particle diameter by procedures outlined in 3A1a and 3A2. The 75-mm division corresponds to the size of opening in the 3-in screen (76.1 mm) used in engineering. These data are listed for taxonomic placement for particle-size class, i.e., to distinguish loamy and silty family particle-size classes. Refer to Soil Survey Staff (2014, 2011) for additional discussion on particle-size classes. Coarse fractions with 0.1- to 75-mm particle diameter are reported as a weight percentage on a <75-mm basis (3D12).

Ratios and Estimates Related to Particle-Size Analysis, Bulk Density, and Water Retention (3D)

>2-mm Fraction (3D13)

The KSSL determines coarse fractions with >2-mm particle diameter by procedure outlined in 3A2. Coarse fractions with >2-mm particle diameter are reported as a weight percent on a whole-soil basis (3D13). For more information on these data, refer to Soil Survey Division Staff (1993) and Soil Survey Staff (2011, 2014).

References


Micromorphology (3E)

Thin Sections (3E1)

Preparation (3E1a)

1. Application

Micromorphology is used to identify fabric types, skeleton grains, weathering intensity, and illuviation of argillans and to investigate genesis of soil or pedological features.
2. Summary of Method

In this method (3E1a), a soil clod is impregnated with a polymer resin (Innes and Pluth, 1970). A flat surface of the soil sample is glued to a glass slide. The soil sample is cut and ground to a thickness of ≈30 µm. The thin section is examined with a petrographic microscope (Anon. 1987; Cady, et al., 1986).

3. Interferences

Impregnation of the soil sample must be complete or the sample will disintegrate during processing. Air bubbles interfere with petrographic examination. The number of bubbles can be minimized by using proper temperature, pressure, and technique. The final, 30-µm thickness is estimated by examining the slide under polarized light. If the quartz-interference colors are of first order, i.e., white, gray, and pale yellow, the sample is ≈30 µm (Anon. 1987).

4. Safety

Use adequate ventilation when mixing, heating, and applying resins. Use tongs or heat resistant gloves when handling hot slides or resins.

5. Equipment

5.1 Petro-thin, thin sectioning system, Buehler, Lake Bluff, IL
5.2 Metallographic polisher with cast-iron laps
5.3 Diamond saw
5.4 Electric oven
5.5 Hot plate with temperature control or a petrographic slide warmer
5.6 Polarizing microscope
5.7 Vacuum, 0.8-bar (80 kPa)
5.8 Desiccator
5.9 Porcelain crucibles
5.10 Standard petrographic slides
5.11 Cover glass
5.12 Silicon carbide abrasives
5.13 Squares of thick, rough-textured plate glass, 305 mm
5.14 Metal probes or dissecting needles
5.15 Small forceps
5.16 Art brush
5.17 Razor blade
5.18 Small chisel, probe (ice pick), ordinary hacksaw, or jeweler’s hacksaw
6. Reagents

6.1 Scotchcast resin, Industrial Electrical Products Division, 3M Company, 3M Center, St. Paul, MN 55101.
6.2 Epoxide. The KSSL uses EPO-MIX from Buehler.
6.3 Ethylene glycol, automotive coolant/anti-freeze

7. Procedure

Sample Collection

7.1 Collect samples by any procedure that does not disturb the natural structure. Core samplers are commonly used. A satisfactory procedure for some soils is to use a knife or trowel to carve a clod that fits in bulk density box cells, a tin box, matchbox, or a round ice-cream container.

7.2 Place clods in an upright position in the container. Mark the top of clod with a thumb tack, staple, or pin to ensure proper orientation.

7.3 Select clods from bulk samples if orientation is of no interest. Do not select clods that are coated with Saran or other plastic because coatings interfere with the grinding after the clod is sectioned.

7.4 Place clods in small plastic bags to avoid contamination from other samples during transit. Pack irregular clods in box cells or containers with light weight material to avoid breakage. Unless fragility is affected, the prevention of moisture loss is usually unnecessary because most samples are usually dried before impregnation.

7.5 The whole core or clod can be impregnated, but better results are generally obtained with specimens that are ≤5 cm³. Remove specimens from the sample with a small chisel, a probe (ice pick), an ordinary hacksaw, or a jeweler's hacksaw.

Sample Preparation

7.6 Place the soil sample in a disposable heat- and chemical-resistant beaker and place in a glass desiccator. Evacuate the air from the desiccator and dry the sample overnight at 80 °C. The natural structure of the soil samples is better preserved by the technique of freeze-drying. Impregnation of the freeze-dried samples may be an improvement over oven-dried samples. In preparation for the proceeding step, bring the freeze-dried or oven-dried sample to ≈80 °C.

Mixing plastic solution

7.7 Many good resins are on the market. The NSSL uses Scotchcast. Add two parts by weight of part A to three parts of part B. For best results, raise part
A and part B to ≈80 °C before mixing. Weigh parts to an accuracy of 2% before mixing. Mix until a uniform color is obtained.

**Impregnation**

7.8 Open the desiccator that contains the dry samples, add the heated plastic solution, and evacuate all the air from the sample. Release the vacuum and again evacuate the air. Do not mistake boiling solution under evacuated conditions for escaping air bubbles from the sample material.

7.9 Cure the impregnated soil overnight in the oven at 110 °C. After curing, the block is ready for sectioning. Disposable containers can be cut with the cooled samples during sectioning.

**Cutting and Rough Grinding**

7.10 With a diamond saw blade, cut the sample block into 13-mm thick chips that are small enough to fit on a regular petrographic slide.

7.11 With a slurry of successively finer abrasives, grind one surface smooth on the revolving lap until the surface is highly polished. Experience is required to determine the mixture of abrasive and water that gives the best results for each grade of abrasive. If the sample surface tends to pull apart or to react with water, dry and re-impregnate the small chip or polish it by hand on a glass plate. Alternatively, grind the block with an abrasive and lubricate with ethylene glycol.

7.12 Clean the sample free of all abrasive material. An ultrasonic bath is recommended. Dry thoroughly.

**Mounting**

7.13 Burnish petrographic slides to a uniform thickness.

7.14 Firmly attach the chip to the burnished slide with a strong, transparent bonding agent. Use a thermoplastic cement or epoxide. Mix the resin and hardener according to the product instructions. Allow entrapped air to rise to the top. Apply a thin layer of epoxide to the slide and to the chip. Place the chip obliquely on the slide and lower slowly. Move the chip back and forth with moderate pressure to remove entrapped air bubbles and excess epoxide. Clamp with a spring clamp and cure at ≈50 °C for 1 h.

**Final Grinding**

7.15 To ensure proper operation of the Petro-thin, consult the operation and maintenance instructions.

7.16 Clean the glass of excess cement.
7.17 Use the Petro-thin to cut off the excess sample and grind the sample to ≈30 µm with the diamond lap. Examine the section frequently under a polarizing microscope during the final stages of grinding. If quartz is present in the sample, use it to judge thickness. If the sample is ≈0.030 mm thick, the quartz interference colors are of the first order, i.e., white, gray, and pale yellow.

7.18 If a Petro-thin is not available, trim the mounted sample with a diamond saw blade to a thickness of 50 to 100 µm. Begin with the coarse abrasive and lap by hand until the sample is relatively thin. Use successively finer abrasives.

7.19 Care and considerable practice are needed to develop the dexterity required to handle an almost-finished section without over-grinding. Use the finest abrasive to finish grinding on ground-glass plates. Wash the section free of abrasive and dry thoroughly.

Seating Cover Glass

7.20 Heat the finished section and the cover slip to ≈40 °C. Spread a small quantity of epoxide over the surface of the thin section and the cover slip. Wait a few seconds for the air bubbles to escape. Place the cover glass obliquely on one end of the section and lower very slowly. If any air bubbles remain, remove them by pressing lightly on the cover glass with a soft eraser. As the section cools, but before the plastic hardens, remove excess epoxy with a razor blade. After the epoxy hardens, remove the final thin film with a razor blade. A thick film may cause the slide to break when the epoxy is removed.

7.21 Very dense soils and soils with clay fractions that have 30% or more montmorillonite require special handling. Use either a dry-grinding technique or a more penetrating impregnation procedure. Without using water, cut the sample to the appropriate size. Sprinkle a coarse abrasive (American Optical No. 190) on the ground-glass plate and commence to dry-grind one face of the sample by hand. Use a figure “8” motion or a counterclockwise motion for best results. Use successively finer abrasives and continue to grind until the surface is highly polished. Proceed with the standard mounting technique.

7.22 Aroclor 5460, a thermoplastic chlorinated diphenyl resin (Monsanto), seems to give better impregnation of dense soils. Place pieces of air-dry soil material in xylene and evacuate. Submerge the xylene-saturated soil material in molten Aroclor 5460. Hold the sample in the Aroclor at ≈200 °C for 1 to 2 days. Remove impregnated soil material, allow to cool, and prepare thin sections by dry-grinding.
8. Calculations
None.

9. Report
Describe the thin section as outlined in method 3E1b.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Micromorphology (3E)
Thin Sections (3E1)
Interpretation (3E1b)

1. Application
Background: Micromorphology is defined as the study of soils or regolith samples in their natural undisturbed arrangement using microscopic techniques (Cady, et al., 1986; Stoops, 2003). This study involves the use of thin sections of soil fabric, which is soil material that encompasses the spatial arrangement, size, and shape of soil components in a finite, three-dimensional unit (Stoops, 2003). These natural aggregates of soil are impregnated with a resin, sliced, and then ground and polished to a thickness of about 30 \( \mu \)m on a glass slide. This thin section is studied to describe the types and arrangements of features and optical properties of minerals. This technique entails descriptive terminology that has been developed over the past 60 years. The science and terminology of microfabric analysis was initially documented by Kubiena (1938). Since then, important publications documenting terminology have included Brewer (1964), FitzPatrick (1984, 1993), Bullock et al. (1985), Stoops (2003), and Stoops et al.

1 Michael Wilson, research soil scientist, and W. Dennis Nettleton, research soil scientist (retired), wrote the procedure for description and interpretation of soil micromorphology as seen in thin sections.

Examination of thin sections with a polarizing light microscope can be considered an extension of field morphological studies. The level of resolution increases from field examination to optical microscopic examination and finally to submicroscopic techniques (electron microscopy). Because this sequence of techniques increasingly sacrifices field of view (Cady et al., 1986), the best results are obtained when multiple techniques are employed. For example, some features, such as depletions and concentrations of redoximorphic features, are observable with no magnification. Yet, the use of a microscope may be required to develop genetic theories of formation based on mineralogy, nodules morphology, or characteristics of coatings. Thus, the results of micromorphological studies are most useful when they are combined with other field data (landscape and soil morphological descriptions) and laboratory data (Cady, 1965; Stoops et al., 2010).

Micromorphology is used to identify types and sequences of active or past processes occurring in soils via identification of coatings, fabric types, skeleton grains, and weathering intensity. It is an ideal tool to investigate genesis of soil or pedological features.

Initially, the investigator should scan the overall features of a thin section and determine those features that require emphasis. Rarely can a question be answered by examination of a single thin section from one horizon. Thus, initial scanning should include all thin sections from a soil profile or all those related to a particular problem. Different kinds of illumination should be used with each magnification. By using a polarizing light microscope, we can examine how light interacts with grains in order to determine their mineralogy. Strong convergent light with crossed polarizers elucidates structures in dense or weakly birefringent material that may appear opaque or isotropic. Use of reflected light is often advantageous for examination of isotropic material (Wilson et al., 2012) (fig. 3E1-1). Structures in translucent specimens become more clearly visible if plain light is used and the condensers are stopped down. Everything should be viewed in several positions of the stage or during slow rotation with cross-polarized light.

A thin section is a two-dimensional slice through a three-dimensional body. The shapes of mineral grains and structural features are viewed in one plane, and the true shapes must be inferred. A grain that appears needle-shaped may be a needle or the edge of a flat plate. An elliptical pore may be an angular slice through a tube. A circular unit is probably part of a sphere. The best way to accustom oneself to a volume interpretation of shape rather than a planar interpretation of shape is to repeatedly view similar features that appear to be cut at different angles while maintaining a three-dimensional perspective and an awareness of section thickness. A well-prepared section is 20- to 30-µm thick. Grains less than 20-µm (fine silt and clay) in diameter appear stacked and cannot
be viewed individually. Similarly, pores smaller than 20 to 30 µm cannot be seen clearly. A pore diameter of 20-µm equates to a soil moisture tension of 15 kPa (0.15 bar) (Rode, 1969), so visible pores in thin section are mostly drained at water contents below field capacity.

The general analytical approach is the same for grain studies (method 7B1) and micromorphology. Individual particle-size fractions are used for the identification and mineralogical analyses in grain studies, and the primary objective is mineral quantification. Sand and silt grains are identified by standard methods presented in petrography texts (e.g., Kerr, 1977; Dyar et al., 2008). Although grain counts are important, mineral quantification, especially of minor constituents, is less important in micromorphology. Thin sections are primarily used for information about component arrangement. Sample preparation procedures used for grain studies destroy or eliminate the arrangement of components during the separation of sand, silt, and clay. Recognition of aggregates, grain weathering, secondary pseudomorphs, and pedofeatures, such as coatings, infillings, or nodules, is most important in micromorphological analysis.

Birefringence is the transmission of cross-polarized light through a mineral. The color of the transmitted light represents the difference between two refractive indices of the mineral (fig. 3E1-2). The term “isotropic conditions” indicates that the mineral is dark under cross-polarized light. Isotropic conditions occur when there is no observable difference between the two refractive indices, when the mineral has no internal atomic order (lack of crystallinity), or when the mineral is in the isometric crystal system, e.g., halite or garnet.
Clay that is completely dispersed, randomly arranged (lacks orientation), has short-range order (noncrystallinity), is coated by humus or Fe or Mn oxyhydroxides (Stoops, 2003), or is less than 1 µm in size exhibits no birefringence and appears isotropic in cross-polarized light. However, the clay in a soil is seldom all random and isotropic and occurs as aggregates as well as massive interstitial fillings, coatings, and bridges and in groundmass. Clay generally develops in oriented bodies, either during formation or as a result of pressure or translocation. If enough plate-shaped particles are oriented together in a body that is large enough to see under the microscope, birefringence can be observed. Thus, even though clay particles are submicroscopic, they can be described, characterized, and sometimes identified, e.g., the 1:1 and 2:1 lattice clays can be distinguished.

Silicate clay minerals in soils are platy, with the exception of halloysite. The $a$ and $b$ crystallographic axes are within the plane of the clay mineral “plate,” and the $c$ axis is almost perpendicular to this plane. Even though the crystals are members of the monoclinic crystal system, the minerals are pseudo-hexagonal because the unit lengths of the $a$ and $b$ axes are nearly equal, and the $c$ axis is nearly perpendicular to the other axes. The optical properties, crystal structure, and general habit of most clay-sized minerals are analogous to those of the micas, which can be used as a model to analyze and describe clay properties.
The speed of light that travels in the direction of the c axis and vibrates parallel to the a axis is almost the same as that light that vibrates parallel to the b axis. Therefore, the refractive indices are very close, and the interference effects in cross-polarized light are small (birefringence is weak and interference colors are white, yellow, or brown) when observed along the c axis (observed when the grain is lying flat on the microscope stage or in the thin section). Light that vibrates parallel to the c axis travels faster than in other directions. Therefore, the refractive index is lower. If the edge of the crystal or aggregate of crystals is viewed along the a-b plane between crossed polarizers, two straight extinction positions are viewed, and interference colors are stronger (high birefringence) (fig. 3E1-2). If clay-sized grains are concentrated and organized so that most of the plates are parallel, these optical effects can be observed. The degree and quality of optical effects depend on the purity, continuity, and the orientation of the clay body.

Kaolinite has low birefringence (the refractive index is similar along all three axes) and has refractive indices slightly higher than quartz. In a thin section with a thickness of 30-µm, interference colors for kaolinite are gray to pale yellow. In residual soils that are derived from coarse-grained igneous rocks, kaolinite occurs as silt- and sand-sized, book-like and accordion-like aggregates of (fig. 3E1-3). Even though halloysite can form oriented aggregates, it should not show birefringence because of its tubular habit. Halloysite may show very faint, patternless birefringence, which is caused by impurities or by refraction of light at the interfaces between particles.

The 2:1 lattice minerals (fig. 3E1-2) have high birefringence and show bright, intermediate-order interference colors if the edges of aggregates are viewed. This difference in birefringence between 2:1 and 1:1 minerals (such as kaolinite) is

Figure 3E1-3.—A Btvx2 horizon in a pedon of Dothan soil (06N0831) in plane-polarized (left) and cross-polarized (right) light. The horizon has close porphyric c/f related distribution and a micromass that is composed of kaolinite embedded with Fe oxides. Note the yellow vermiform kaolinite in the right center of the image. This grain likely forms via in situ weathering of a sand-sized grain. Frame width is 2.5 mm.
useful to distinguish these two mineral groups. In the clay-size range, distinctions among smectite, mica, vermiculite, and chlorite in thin section are seldom possible. These clay minerals are usually mixed in the soil, seldom occur pure, and are commonly stained and mixed with iron oxide and organic matter.

Residual clay has been in place since its formation by weathering. Although it may have been transported within fragments of weathered material, it remains in place relative to the fabric of these fragments. This clay may be random, have no orientation, and thus be isotropic; however, more often it shows some birefringence. In many residual materials, clay is arranged either in forms that are pseudomorphs of rock minerals or in definite bodies of crystal aggregates, e.g., vermicular or accordion-like kaolin books (fig. 3E1-3). The regular, intact arrangement of these materials is usually diagnostic of residual material.

Location features that distinguish translocated clay from residual clay are its occurrence in separate bodies, usually with distinct boundaries, and its location on present or former pore walls, channel linings, or ped faces. Translocated clay may have a different composition than matrix clay, especially if its origin is another horizon. This clay is more homogeneous and usually finer than matrix clay. Translocated clay displays lamination, which consists of variable bands of differing clay-sized particles. The lamination is indicative of deposition in successive increments (fig. 3E1-4). The bands manifest birefringence and extinction, indicating that these translocated clay bodies are oriented aggregates. If these bodies are straight, they have parallel extinction. If these bodies are curved, a dark band is present wherever the composite $c$ axis and the composite $a$ and $b$ axes are parallel to the vibration planes of the polarizers. When the stage is rotated, these dark bands sweep through the clay aggregate.

Figure 3E1-4.—A clay coating in a 2Btx horizon of a Cowarts soil (pedon 06N0834). Note the lamination visible under plane-polarized light (left). Under cross-polarized light (right), the clay coating exhibits birefringence and developed lines of extinction, both characteristic of a well-oriented illuvial clay deposit. Frame width is 2.5 mm.
Clay rearrangement may result from differentially applied stress that produces shear or stress orientation (fig. 3E1-5). Root pressure, wetting and drying, mass movement, slump, and creep can produce this orientation. Platy particles can become oriented by slippage along a plane, e.g., slickenside faces in a Vertisol. Stress orientation may therefore be inferred if faces on structural units are smooth and do not have identifiable clay coatings. These particles become oriented inside the structural aggregates as well as on faces. In plain polarized light, clay in thin sections appears homogeneous and featureless and stress orientation is not observed. In cross-polarized light, the orientation pattern is speckled or striated, consisting of bright colored streaks or linear zones showing aggregate birefringence. There may be numerous sets of these slippage planes, which appear in different positions as the stage is turned. Stress-oriented clay may also be near rigid bodies, e.g., sand-sized grains, or along root channels (fig. 3E1-6). Stress-oriented clay is often strongly developed on ped faces. Stress can also orient mica flakes and other small platy grains.

Other substances, such as goethite, gibbsite, carbonate minerals, and gypsum, may form pore linings and ped coatings. These substances can be

Figure 3E1-5.—A Btk2 horizon in a pedon of a White House soil (pedon 40A3559) under cross-polarized light. The horizon has a high percentage of clay of expandable aluminosilicate clay-sized minerals. Clays become aligned through shrink-swell processes. This alignment results in preferred orientation of clay particles, making the clay birefringent (visible under cross-polarized light). Shrinking and swelling results in a loss of clay coatings along ped faces in many soils. Frame width is 0.9 mm.
identified by their mineralogical properties. Calcite is distinctive in thin section. It is white under plane-polarized light and has extreme birefringence (very bright colors) under cross-polarized light (fig. 3E1-7). It can be found in several forms or habits, including microcrystalline (micrite forming crystallitic b-fabric), larger needle shaped or prismatic crystals, and carbonate coatings, pendants, nodules,

Figure 3E1-6.—The Bt1 horizon in a Brod soil (pedon 03N1014), a Ustolic Haplocryalf from Montana. Note the oriented clay around coarse grains (granostriated b-fabric) under plane-polarized light (left) and cross-polarized light (right). Frame width is 1.1 mm.

Figure 3E1-7.—Calcium carbonate around quartz and feldspar grains in a chitonic c/f related distribution pattern. The calcite has characteristic high birefringence under cross-polarized light. This is the Bkq2 horizon in a Cax soil (pedon 97P0420) from San Bernardino, CA. Frame width is 0.9 mm.
filaments, and laminar crusts (Durand et al., 2010). Gypsum is generally colorless in thin section and has weak birefringence (first order gray) (fig. 3E1-8). The morphology of gypsum is variable but distinctive, appearing as prismatic, tabular, lenticular, or acicular forms (Poch et al., 2010).

Amorphous coatings of organic matter, with or without admixed Fe and Al, are common, especially in spodic horizons (Wilson and Righi, 2010). This material is dark brown to black, isotropic to faintly birefringent, and commonly flecked with minute opaque grains. Amorphous coatings of organic matter occur as the bridging and coating material in B horizons of sandy Spodosols and as thin coatings or stains on pore and ped faces in other soils.

2. Procedure

Description of Microfabrics

Terminology has developed for the description of soil fabric composition (Brewer and Sleeman, 1963; Brewer, 1964 and 1976; Stoops and Jongerius, 1975; Brewer et al., 1983; Bullock et al., 1985; and Stoops, 2003). As these terms have become more widely adopted in the literature, the KSSL increasingly uses them in Soil Survey Investigations Reports (SSIRs), journal articles, and soil-project correspondence. Micromorphological descriptions commonly contain terminology from different sources to describe properties of the fabric, but it is important to use current, standard terminology for consistency. The terminology emphasized here is taken from Stoops (2003), and this book should be consulted.

Figure 3E1-8.—Lenticular gypsum crystals surrounding a micritic calcite mass under cross-polarized light. This is the Bym1 horizon in a Drygyp soil (pedon 04N0770) from Nevada. Frame width is 2.5 mm.
for details. Stoops et al. (2010) is an excellent reference for application of these
terms for different types of soils and suites of minerals.

The major descriptive separations of materials that compose the soil fabric are
“groundmass” (the base material of coarse and fine materials) and “pedofeatures”
(distinct fabric units that are concentrated and distinct from the groundmass).
These two major components are described as the elements of soil fabric.
The commonly described components of thin section descriptions are (a) soil
microstructure and porosity, (b) groundmass (mineral and organic constituents),
and (c) pedofeatures.

**Soil Microstructure:** This descriptive component concerns the size, shape,
and arrangement of particles and voids in soil material. It is applicable to both
aggregated and non-aggregated soil material. If aggregates are present, types of
peds (spheroidal, blocky, plates, or prisms aggregates) can be described as well
as the degree of ped separation, size, accommodation, roughness, and pattern.

Voids, the space between solid materials, form a continuum in soil but are
treated as individuals and described by morphology. Types of voids are “packing
voids” (formed from the loose packing of unaccommodated faces of grains or
between peds or other compound individuals) (fig. 3E1-9), “vughs” (generally
equidimensional voids with irregular shapes) (fig. 3E1-10), “vesicles” (spherical
voids with smooth perimeters) (fig. 3E1-11), “channels” (elongated voids) (fig.
3E1-6), “chambers” (smooth walled voids intersected by channels), and “planes”
(planer, flat voids) (fig. 3E1-12). In addition to types of voids, the size, abundance,

**Figure 3E1-9.—Simple packing voids, partially coated with
fine materials under plane-polarized light. These
voids arise from loose packing of soil components.
This is the Bt1 horizon in an Endlich soil (pedon
99P0001) from Colorado. Frame width is 1.1 mm.
Figure 3E1-10.—A void (under plane-polarized light) that would be described as a vugh. Vughs are semi-equipartitioned, irregular voids that are not interconnected with other voids. This photomicrograph is of the Bkss horizon of a Typic Epiaquert from Brazoria County, Texas (pedon 98P0582). This horizon is 52% clay and 42% silt. The COLE is 0.011, and the clay fraction is principally composed of smectite. Frame width is 2.5 mm.

Figure 3E1-11.—A Typic Haplargid (Dera soil, pedon 81P0610) from Utah under cross-polarized light. The walls consisting of “smooth, simple curves” indicate that this void is a vesicle. These vesicles were formed in the thin, surface crust (A1 horizon). Frame width is 3.2 mm.
The roughness of walls, accommodation, and arrangement of voids may also be described.

Nomenclature for microstructures has been developed based on ped and void types. The list of microstructure types is fairly extensive; for example, vughy microstructure, granular microstructure, and lenticular microstructure. The most common types are listed in Stoops (2003).

Groundmass: Groundmass is a general term for the coarse and fine materials present in the soils, inclusive of voids. Groundmass components can be considered to form the base materials in the soil and reflect the lithology and weathering of the parent material. Groundmass can be considered the materials left after the identification of the pedofeatures, such as nodules and clay coatings. The term “groundmass” is used similarly to Brewer’s (1976) term “s-matrix,” although some differences exist (Stoops, 2003).

A description of groundmass should include the author’s designated size limit between coarse (c) and fine (f) materials, c/f-related distribution, and characteristics of coarse material and fine material (micromass). The five “coarse-fine (c/f) related distribution patterns,” as described by Stoops (2003), are intended to be broadly defined. There are no restrictions on material type, absolute size, orientation, granulation, or origin. The system may be used to describe the distribution of primary particles, e.g., quartz grains, as well as compound units, e.g., humic micro-aggregates. The size limit between coarse and fine units “floats,” based on the individual soil. The coarser particles may be silt, sand, or gravel, whereas the finer material may be clay, silt, or sand. Figure 3E1-13 shows the average textures, linear extensibilities (LE), and drained pore to filled pore (DP/FP) ratios of some related distribution patterns of a number of U.S. soils.

Figure 3E1-12.—Voids termed “planes” that developed in the platy structure of the A1 horizon in a Frisite soil (pedon 98P0453) from Wyoming. The image on the left is under plane-polarized light, and the image on the right is under cross-polarized light. Frame width is 2.5 mm.
The five coarse-fine (c/f) related distribution patterns are monic, gefuric, chitonic, enaulic, and porphyric. The monic distribution consists of fabric units of only one size group, e.g., pebbles, sand, lithic fragments (coarse monic), or clay (fine monic). The gefuric distribution consists of coarser units linked by bridges of, but not surrounded by, finer material. The chitonic distribution has coarser units surrounded by coatings of finer material (fig. 3E1-7). In the enaulic distribution, larger units support one another, and the interstitial spaces are partially filled with aggregates of finer material (fig. 3E1-14). The enaulic fabric consists of a greater amount of finer material than in either the gefuric or chitonic distributions. The porphyric distribution is the end member of the sequence. This distribution has coarse materials embedded in finer materials, and there is an absence of interstitial pores (figs. 3E1-1 and 3E1-3). The porphyric distribution may be divided into types based on the spacing of the coarser units. Additional information on the coarser material can also be recorded regarding the mineralogy (composition), size, abundance, shape, color, roughness, and sorting.

Figure 3E1-13.—Types of related distribution patterns and their physical properties. Frame width of each idealized kind of fabric is 0.5 mm. The lower size limit of coarse material in the c/f related distribution patterns was set at about 50 µm for most of the slides.
The fine materials (micromass) and the characteristics of the fabric are generally described based on optical characteristics (color, limpidity, and birefringence) because the size and shape of the particles are below the resolution of the microscope. Zones or aggregates of oriented clay display interference colors due to the birefringence of clay particles. Thus, a primary descriptive category of the micromass is orientation and distribution of interference colors from birefringence fabric, or b-fabric. The main types of b-fabric are described as undifferentiated (absence of interference colors); crystallitic (presence of small birefringent mineral grains, e.g., calcite, that result in the color of the b-fabric) (fig. 3E1-7); speckled (small, randomly arranged domains of oriented clay) (fig. 3E1-15); striated (elongated domains of oriented clay that are generally parallel throughout the structural unit); and strial (the entire structural unit exhibits birefringence with uniform parallel extinction throughout the unit. Both speckled and striated b-fabrics are further subdivided. Speckled materials can be either stipple speckled (fig. 3E1-16) or mosaic speckled (fig. 3E1-17). Striated materials can be subdivided into 10 types, e.g., porostriated, parallel striated (fig. 3E1-18), monostriated, granostriated (fig. 3E1-19), or random striated.

**Pedofeatures:** The term *pedofeatures* refers to discrete units of material that are present in the fabric and have apparent differences to the groundmass based on concentration. Examples are clay coatings, nodules, and crystals. The size of a pedofeature has no upper limit. It has a lower limit of about 20 µm, which is the lower limit of resolution with a petrographic microscope. A major distinction in pedofeatures is made between (a) those that formed due to weathering within or intrusion of a feature into the groundmass (*matrix pedofeatures*) and (b) those that formed through processes outside the groundmass and formed in voids or appended to the groundmass (*intrusive pedofeatures*). A matrix pedofeature can

*Figure 3E1-14.—An example of the enaulic c/f related distribution pattern under plane-polarized light (left) and cross-polarized light (right). The image is of an E horizon in a Fuquay soil (pedon 06N0830) from South Carolina. The soil is a Kandiudult. Frame width is 2.5 mm.*
Figure 3E1-15.—A speckled b-fabric under cross-polarized light in an Ap horizon of a Typic Argiudoll. The soil is in the Southridge series (pedon 87P0075) from Allamakee County, Iowa. This horizon has 19% clay and a COLE of 0.016. The clay fraction is dominated by vermiculate and smectite. Frame width is 1.3 mm.

Figure 3E1-16.—A stipple speckled b-fabric under cross-polarized light from the BE horizon in an Adco soil (pedon 87P0771), which is a Vertic Albaqualf, from Macon County, Missouri. The horizon consists of 29% clay and 63% silt. It has a COLE of 0.026 and a clay fraction dominated by smectite with lesser amounts of mica and kaolinite. Frame width is 1.3 mm.
Figure 3E1-17.—A mosaic speckled b-fabric under cross-polarized light of the 2Btg3 horizon in a Leonard soil (Pedon 87P0770), which is a Chromic Vertic Epiaqualf, from Macon County, Missouri. Frame width is 1.1 mm. This horizon has 48% clay, a COLE of 0.132, and a clay fraction dominated by smectite.

Figure 3E1-18.—A parallel striated b-fabric under cross-polarized light of the Bt horizon in a Gloria soil (pedon 40A2845), which is a Durixeralf, from Monterey County, California. Frame width is 1.3 mm. This horizon has 47% clay, a COLE of 0.056, and has a clay fraction dominated by illite with lesser amounts of kaolinite and smectite.
Figure 3E1-19.—A granostriated b-fabric under cross-polarized light of the Bt horizon in a Redding soil (Pedon 40A2847), which is a Durixeralf, from San Diego County, California. Frame width is 1.1 mm. This horizon has 60% clay, a COLE of 0.063, and a clay fraction dominated by kaolinite and smectite.

be (1) an impregnative pedofeature (an infusion of a material, e.g. iron oxides, into the microporosity of the groundmass or preexisting void) (fig. 3E1-20); (2) a depletion pedofeature (a lower concentration or loss of a substance, e.g. iron oxide, in a zone of the ground mass) (fig. 3E1-21); or (3) a fabric pedofeature (an alteration in the groundmass only, e.g., disturbed material filling void via bioturbation of earthworm). Descriptive terms for the types of matrix pedofeatures are hypocoatings, quasicoatings, matrix infillings, intercalations, and matrix nodules. Descriptive terms for intrusive pedofeatures are coatings, infillings, crystals and crystal intergrowths, intercalations, and nodules. The difference between these two sets of descriptive terms (matrix and intrusive pedofeatures) is based on the location of the pedofeature being described. Coatings can be textural coatings (composed of sand, silt, or clay), crystalline, cryptocrystalline, or amorphous. Textural coatings can be described as sorted, unsorted, laminated, or layered. The most common coating described in thin sections is composed of clay (fig. 3E1-4). Several terms for clay coatings have been used in the past, including cutans (Brewer, 1964), clay films, and clay skins. Brewer described cutans according to composition, e.g., argillans, neomangans, and calcitans, but this terminology has been dropped in favor of describing the mineral or textural composition of the coating, e.g., calcite coatings and clay coatings.
Figure 3E1-20.—A ironstone nodule (petroplinthite) from the E horizon in a Dothan soil under plane-polarized transmitted light (left) and reflected light (right). The core of the nodule is regarded as composed of well-crystalline goethite as opposed to plinthite nodules, which are composed of primarily noncrystalline Fe oxides (Eswaran and Raghu Mohan, 1972). Frame width is 16.7 mm.

Figure 3E1-21.—The mottled fabric on the pit face (left) of a Btvx2 horizon in a Fuquay pedon, and the thin section of soil fabric (right) from the same horizon. These photos illustrate the movement of Fe oxides in the soil as it undergoes seasonal saturation and reduction followed by reoxidation.

Usually, the basic descriptive terms for soil fabrics do not imply any specific genesis of the feature. However, modifiers can be added when fabric descriptions are complete enough to understand the means of formation, e.g., “stress-oriented clay coating,” signifying in-place plasma modification resulting from differential forces, such as shearing. This can be compared to the description of an “illuviation clay coating,” formed by movement of material in solution or suspension and later deposited.
Interpretations

Related Distribution Patterns: The average properties of some related distributions are shown in figure 3E1-13. In an experimental study of soil microfabrics by anisotropic stresses of confined swelling and shrinking, Jim (1986) showed that with an increase in the activity and proportion of the clay fraction, the related distribution patterns alter from dominantly monic to enaulic to porphyric.

Some monic fabrics are inherited and include soil fabrics formed in sand dunes, sandy sediments deposited by streams and rivers, beach deposits, and gruss. Fauna can produce monic fabrics that are mostly fecal pellets. Monic fabrics also can form by fracturing and flaking of organic coatings in the upper B horizons of the Spodosols (Flach, 1960) and by freezing and thawing (Brewer and Pawluk, 1975).

Several kinds of finer material (micromass) can bridge the coarser particles (skeleton grains) to form gefuric related distribution patterns. Gefuric patterns are common in weakly developed argillic and spodic horizons and in some duripans. Bridges of material form between skeleton grains. Typically, the cement or clay is material that bond covalently with skeleton grains. These cements commonly include silica (fig. 3E1-22), iron, aluminum, and organic matter (Chadwick and Nettleton, 1990). As the amount of cementing agent or clay increases, the next progression is the chitonic related distribution pattern (Pawluk, 1983). For example, silicate clays can bridge skeleton grains in some argillic horizons (gefuric c/f related distribution pattern) and subsequently develop complete coatings of clay with increasing illuviation (chitonic c/f related distribution pattern).

In spodic horizons, monomorphic Al-Fe organic complexes develop and, with increasing amounts, grade from gefuric to chitonic to close porphyric (Wilson and Righi, 2010; Salem Avad et al., 1982). Even though organic matter has covalent bonds and usually surrounds grains, organic material can form pellets in void spaces between skeleton grains in some spodic horizons. These types of horizons form horizons with enaulic c/f related distribution patterns. These pellets are polymorphic materials, consisting of crumbs or granules of several forms of degraded plant materials (De Coninck and Righi, 1983). In the lower horizons of Spodosols, monomophic materials are dominant. These materials are regarded as soluble organic compounds that are water-transported and precipitated. These deposits coat mineral grains and are present on channel pores (Buurman et al., 2005), forming horizons with close porphyric c/f related distribution patterns.

The enaulic related distribution patterns are more common in soil material in which the cement bonds to itself more strongly than to skeleton grains. For example, ionic-bonded calcite and gypsum tend to bond to themselves more strongly than to skeleton grains in sandy-textured soils (fig. 3E1-8), thereby producing enaulic followed by open porphyric c/f related distribution patterns (Chadwick and Nettleton, 1990).
Figure 3E1-22.—Horizon with duripan exhibiting silica cementation. Fabric has an opal and chalcedony laminar cap. The matrix above and below is composed of durinodes (non-crystalline silica) surrounded by moderately-oriented silicate clays. Clay can provide the initial absorption surface for silica in soil solution. The absorption of silica onto established silica phases leads to formation of nodules. Frame width is 1.1 mm. (Series name not designated; Jefferson Co., OR; pedon 87P0513; 2Bkqm horizon under plane-polarized light).

_Porphyric c/f_ related distribution patterns are common in loessial soils, especially in argillic and petrocalcic horizons, duripans, and ortstein. This pattern can be the end member of a progress sequence, or form from normal packing of grains in materials that have a high proportion of fine material. In precursors of the porphyric related distribution patterns, the silt-to-clay ratio is useful for identifying the kind of sequences by which the porphyric pattern develops (Brewer et al., 1983). In porphyric related patterns, there may or may not be skeleton grains of primary minerals, pedorelicts, organics, lithic fragments of shale, sandstone, or other rocks. The micromass consists of silt and clay, and the interstices tend to be filled with minimal formation of coatings.

**Micromass:** There are at least two origins for oriented clay on coarser textured, sandy soils. One origin is a result of clay illuviation, commonly associated with monic, gefuric, or enaulic c/f related distribution patterns. The other origin is shrink-swell processes and is more commonly found in porphyric c/f related distribution patterns with linear extensibilities >4 % for dryland soils, i.e., soils in aridic, xeric, or ustic soil moisture regimes. The b-fabrics in these latter types of soils are the true _granostriated_ b-fabrics. Shrink-swell forces have been
involved in their formation as shown by the relatively few papules or clay coatings remaining and by areas of *striated* b-fabrics.

*Speckled or striated* b-fabrics have higher clay contents, usually <30% but as much as 70% (Brewer et al., 1983). These b-fabrics are important in many fine-textured B horizons. *Parallel, cross,* and *random striated* b-fabrics are evidence of linear extensibilities >4% in dryland soils that have higher clay contents. Clay content is typically >35% in soils with these b-fabrics, but the threshold amount is dependent on the type of clay mineral and the degree of dryness common to the environment. In these three fabrics, clay coatings are rarely found, but areas of *granostriated* and *porostriated* b-fabrics may be present.

Deformation experiments indicate that the degree of clay orientation increases with an increase in clay percentage, linear extensibility, and applied stress (Clark, 1970; Edil and Krizek, 1976). In an experimental study of soil microfabrics by anisotropic stresses of confined swelling and shrinking, Jim (1986) showed that with an increase in the activity and content of the clay fraction, there is an increase in the long and narrow plasma separations, i.e., a progression from *stippled speckled* to *mosaic speckled* to *parallel striated* b-fabrics. The b-fabric types can form a sequence relative to increasing linear extensibility in soils from a similar climate. With increasing linear extensibility, the micromass sequence has been shown to progress from *stippled speckled* to *mosaic speckled* to *random striated* to *parallel striated* (Nettleton et al., 1969; Holzhey et al., 1974).

*Stippled speckled plasmic* b-fabric is very common in finer-grained porphyric B horizons of a wide range of soil groups (Brewer et al., 1983). Soils with this b-fabric generally have a linear extensibility of <4 percent. In some stipple speckled b-fabrics, clay islands are pseudomorphs of some weatherable mineral. In other stipple speckled fabrics, these islands are fragments of clay coatings or eolian sand-size clay aggregates (Butler, 1974). *Mosaic speckled* b-fabrics commonly contain more islands and therefore have more clay than stipple speckled fabrics. However, in mosaic speckled b-fabrics, linear extensibility also remains low. Shrink-swell forces have not been sufficient or have not operated long enough to have homogenized the islands of clay into the soil matrix.

*Striated* b-fabrics occur in soil horizons that have undergone stress either due to shrink-swell forces or to tillage. Even though root growth has been found to be adequate to increase the percentage of oriented clay near the root-soil interface (Blevins et al., 1970), root growth does not appear adequate to form these highly stressed b-fabrics.

Organic matter or iron stains that result in a flecked distribution pattern can mask the birefringence of crystalline clays. A complete absence of interference colors is what characterizes the *undifferentiated* b-fabric that is common in Spodosols and Andisols. In spodic horizons, the clay is commonly composed of monomorphic organic materials enriched with Al (Wilson and Righi, 2010). In Andisols, the b-fabric is *undifferentiated* due the presence of noncrystalline colloids of allophane or imogolite (Sedov et al., 2010). The water-holding
capacities of these soil horizons are relatively high, and some unweathered volcanic ash may be present.

Crystallitic b-fabrics are common in B horizons of soils that formed in arid climates. These soils have many small birefringent mineral grains that control the interference colors of the whole (Stoops, 2003). Micaceous soils with fine grained sericite are commonly crystallitic. Arid soils that are rich in microcrystalline (micritic) calcite also exhibit this type of fabric (Durand et al., 2010) compared to soil horizons principally composed of microcrystalline gypsum that have nearly undifferentiated b-fabric (Poch et al., 2010). In soil horizons with large areas of interlocking crystals, soil permeability is restricted, unconfined compressive strength is increased, and particle dispersion is limited, depending on the degree of cementation.

Pedofeatures: Most clay coatings (fig. 3E1-4) are formed, at least in part, by illuviation. If clay coatings are present in argillic horizons of dryland soils, the soil linear extensibility is typically <4 percent (Nettleton et al., 1969). In some humid environments, these features may be present even where the linear extensibility is >4 percent due to decreasing amounts of shrink swell. As soils in humid environments do not dry to the same degree as those in the desert, the clay coatings may survive because only part of the linear extensibility is effective.

The content of strongly oriented clay (typically clay coatings, but also including hypocoatings and quasicoatings), in soils that have argillic horizons is usually <5% of the soil volume. In some sandy soils that are low in silt, these three coatings may be up to 30% of the soil material (Brewer et al., 1983). The measured illuvial clay rarely accounts for the difference in clay content between the A and B horizons, indicating that some of the clay may originate from weathering in place and some from a destruction of the three types of clay coatings.

Clay hypocoatings (matrix pedofeatures immediately adjoining a void) and quasicoatings (a pedofeature within the matrix and not adjacent to a void surface) may originate by the weathering of primary minerals, the isolation of clay coatings by the channel, and void migration within the soil matrix (Nettleton et al., 1968; Nettleton et al., 1990) or by the introduction of eolian sands and silts that are composed of clays (Butler, 1974; Brewer and Blackmore, 1976). Internal fabric resemblances, comparison of parent material, and comparison of size and shape of minerals within the hypocoatings and quasicoatings may help to determine if the coatings are pseudomorphs of one of the primary minerals.

Nodules of Fe and Mn oxides are common in soils with fluctuating water table levels, but do not tend to develop in soils with long term saturation (fig. 3E1-20) (Jien et al., 2010). Soluble Fe and Mn migrate via diffusion through the soil matrix, precipitating in small pores. As the small pores grow, they commonly envelop small grains that become entrapped in the interior of the nodules (Wilson et al., 2012; Lindbo et al., 2002). Coatings of Fe oxides and Mn oxides are common along voids of soils that undergo seasonal saturation (fig. 3E1-1). The soluble
Fe moves to the channel or pore, where it precipitates (Ogg et al., 2011). This migration can result in depletion pedofeatures (fig. 3E1-21).

A study of soil voids may be useful in predicting the clay activity and shrink-swell behavior of soils. In an experimental study of soil microfabrics by anisotropic stresses of confined swelling and shrinking, Jim (1986) showed that with an increase in the activity and content of the clay fraction, there is a drastic decrease in void volume, especially the >30 µm voids. Furthermore, the void shapes change from compound packing voids to planar voids and vughs. With an increase in stress from shrink-swell forces, aggregates become flattened at contacts, resulting in more angular aggregates and eventually fused compound units.

Micromorphological studies can measure porosity and predict soil water content at various suctions and hydraulic conductivity. In thin section studies of voids in sands and sandy soils, there is a close correlation between micromorphology and suction methods (Swanson and Peterson, 1942). However, in those soils in which the volume changes with water content, pore size distribution is undefined and no constant void size distribution exists (Brewer, 1976). Furthermore, several invalidated assumptions are commonly made in relating porosity to permeability (Nielsen et al., 1972, p. 11). The assumptions that especially relate to soil fabric are "no pores are sealed off," "pores are distributed at random," and "pores are generally uniform in size." Vepraskas et al. (1991) used micromorphology to demonstrate that fractures and veins in saprolite were generally plugged with clay and oxides and that most water flow through the saprolite was via root channels.

The size, shape, and arrangement of skeleton grains determine the nature of simple packing voids, but the origin of compound packing voids is not so straightforward. The unaccommodated peds of the compound packing voids may be formed by faunal excreta, shrink-swell action, human activities, or unknown causes.

Vughs typically occur in soil materials that have a wide range in size of particles, including silicate clays. Some vughs form by the weathering and removal of carbonate, and others form by faunal activity or the normal packing of plasma and skeleton grains. Blevins et al. (1970) showed that porosity decreased around trees due to root pressure on the surrounding soil. Balbino et al. (2002) found that the loss of porosity on soils cleared of their native savanna vegetation to develop pastures resulted from the loss of earthworms and other fauna rather than the change in land use. Voids characterized as planes (fig. 3E1-12) are produced in relatively uniform fine-textured soils by a relatively regular system of cracking upon drying (Brewer, 1976). Once formed, these joint planes tend to open in the same place during successive drying cycles.

Vesicular pores (fig. 3E1-11) are non-connecting spherical pores that typically occur in soils in arid regions (Nettleton and Peterson, 1983). The process to form these vesicles results from sealing of the soil surface and the entrapment of air during cycles of wetting and drying (Lapham, 1932; McFadden et al., 1998;
Turk and Graham, 2011; Williams et al., 2012). This cycle is also responsible for creation of polygonal cracking and the creation of columnar structure. Over time, the vesicles collapse and form planar voids (Turk and Graham, 2011). Laboratory studies verify this phenomenon (Springer, 1958). If soils that have a high content of silt are allowed to dry before each irrigation, the vesicle size increases with the number of irrigations (Miller, 1971). Studies of infiltration rates and sediment production in rangeland in central and eastern Nevada found that the infiltration rates are lowest and the sediment yields are highest on sites that have vesicular surface horizons (Blackburn and Skau, 1974; Rostagno, 1989). The failure of most vesicles to connect to other voids and the low strength of the crust in which vesicles occur help to explain the low infiltration rates and high sediment yields.

3. References


Wet Aggregate Stability (3F)

Wet-Sieving (3F1)

Air-Dry, 2 to 1 mm, 2- to 0.5-mm Aggregates Retained (3F1a1a)

1. Application

An aggregate is a group of primary particles that cohere to each other more strongly than to other surrounding soil particles (Soil Science Society of America, 1997). Disaggregation of soil mass into aggregates requires the application of a disrupting force. Aggregate stability is a function of the capacity of cohesive forces between particles to withstand the applied disruptive force. The analysis of soil aggregation can be used to evaluate or predict the effects of various agricultural techniques, e.g. tillage and organic-matter additions, and erosion by wind and water (Nimmo and Perkins, 2002). The measurement can serve as a predictor of infiltration and soil erosion potential. This method provides a measure of aggregate stability following a disruption of initially air-dry aggregates by abrupt submergence followed by wet sieving. This procedure was developed for use by the soil survey field offices of the Natural Resources Conservation Service.
2. Summary of Method

This method (3F1a1a) measures the retention of air-dry aggregates (2 to 1 mm) on a 0.5-mm sieve after sample has been submerged in RO water overnight followed by agitation of sample.

3. Interferences

Air bubbles in the sieve can create tension in the water, thereby reducing the percentage of aggregates that are retained on the 0.5-mm sieve. Variation in the moisture content of air-dry soils can affect results. A correction should be made for the sand >0.5 mm resistant to dispersion in sodium hexametaphosphate.

4. Safety

If ovens are used, hot surfaces can be a hazard. Follow standard laboratory safety precautions.

5. Equipment

5.1 Bowls, Rubbermaid or equivalent, 1800 mL
5.2 Electronic balance, ±0.01-g sensitivity and 500-g capacity
5.3 Sieves, square-hole
   5.3.1 0.5 mm, stainless steel, no.35, 125-mm diameter, 50-mm height
   5.3.2 1 mm, brass, 203-mm diameter, 50-mm height
   5.3.3 2 mm, brass, 203-mm diameter, 50-mm height
5.4 Oven, 110 °C
5.5 Camping plate, Coleman, stainless steel, 152-mm diameter, Peak 1, Model 8553-462.
5.6 Aluminum foil dish, 57-mm diameter x 15-mm deep, with lifting tab

6. Reagents

6.1 Reverse osmosis (RO) water
6.2 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate (Na₄P₂O₇) and 7.94 g of sodium carbonate (Na₂CO₃) in 1 L of RO water. Alternatively, use Calgon water softener.

7. Procedure

7.1 Use natural fabric (NF) samples in pint containers. Assemble a 2-mm sieve on top of a 1-mm sieve. Crush the NF sample by hand or with mortar and pestle. Crush sample so as to pass the 2-mm sieve with a minimum reduction in size. Sieve entire NF sample.
7.2 Place the material that is retained on 1-mm sieve in pint container and discard the remaining material.
7.3 Sieve the material again with 1-mm sieve to remove dust and other small particles. Weigh approximately 3.00 ±0.05-g sample of the 2- to 1-mm material in aluminum foil dishes.

7.4 Place 0.5-mm sieve in plastic bowl and fill bowl so that the water level is at a 20-mm height above the base of screen. Remove air bubbles with a syringe.

7.5 Distribute the 3.00-g sample (2 to 1 mm) on the 0.5-mm sieve. Aggregates should not touch. Allow sample to sit overnight.

7.6 Agitate the sample by raising and lowering the sieve in the water bowl 20 times in 40 s. On the upward strokes, drain sieve but do not raise so high that air enters to beneath the sieve.

7.7 Remove sieve from water bowl, place on Coleman plate, and dry in oven for 2 to 2.5 h at 110 °C. During drying process, the plate retains the soil that drops through the sieve.

7.8 Remove the sample from the oven. Weigh sieve, plate, and sample. Record weight (Wt1). If no sand (>0.5 mm) is present, discard sample from sieve and plate by brushing. Weigh sieve and plate. Record weight (W2). Sample is those aggregates retained on 0.5-mm sieve \( W_R = Wt_1 - Wt_2 \).

7.9 If sand (>0.5 mm) is present and no particle-size data is available, discard sample on plate and disperse sample that was retained on the sieve with sodium hexametaphosphate solution. Alternatively, place 3 g of Calgon in plastic bowl and stir until dissolved. Place the 0.5-mm sieve with sample in sodium hexametaphosphate (or Calgon) solution so that the solution line is 35 mm above the base of screen. Gently triturate the dispersing solution with fingers to remove soft <0.5 mm material adhering to the ≥0.5 mm particles. Remove sieve from sodium hexametaphosphate (or Calgon) solution and rinse with RO water until all sodium hexametaphosphate (or Calgon) solution has passed through sieve and only the sand (>0.5 mm) is left on sieve. Place sieve on Coleman plate, place in oven, and dry for 2 to 2.5 h at 110 °C.

7.10 Remove sample from oven. Weigh the sieve, plate, and sample. Record weight (Wt3). Discard sample and brush sieve and plate. Weigh sieve and plate. Record weight (Wt4). Sand weight \( S_W = Wt_3 - Wt_4 \).

7.11 Thoroughly wash sieve and plate with RO water, especially those sieves with sodium hexametaphosphate solution.

8. Calculations

Aggregates (%) = \( \frac{(W_R - S_W)}{(l_w/(AD/OD)) - S_w} \) x 100

where:

\( l_w \) = Initial sample weight (approximately 3 g)
\[ W_R = \text{Total weight of aggregates retained on 0.5-mm sieve} \]
\[ S_W = \text{Weight of 2- to 0.5-mm sand} \]
\[ \text{AD/OD} = \text{Air-dry/oven-dry weight (if not available, use 1.00)} \]

9. Report

Report aggregate stability as a percentage of aggregates (2- to 0.5-mm) retained after wet sieving. Do not report determinations if the 2- to 0.5-mm primary particles are ≥50% of the 2- to 1-mm sample.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Particle Density (3G)
Pycnometer Gas Displacement (3G1)

Oven-Dry, <2 mm (3G1a1)
Oven-Dry, >2 mm (3G1a2)

1. Application

Density is defined as mass per unit volume. Particle density refers to the density of the solid particles collectively (Flint and Flint, 2002). Particle density is required for sedimentation analysis; calculating soil volume or mass; and for mathematically correcting bulk soil samples containing significant amounts of rock fragments so as to determine fine-soil density, water content, or other soil properties affected by volume displacement of rock fragments (Flint and Childs, 1984).

2. Summary of Method

This method (3G1) determines particle density by the pycnometer gas displacement procedure. It employs the Archimedes’ principle of fluid displacement to determine the volume. The displaced fluid is a gas that can penetrate the finest pores, thereby assuring maximum accuracy (Quantachrome Instruments, 2003). Helium gas is the most commonly recommended gas because it has small atomic dimensions that assure penetration into crevices and
pores approaching one Angstrom ($10^{-10}$ m) in dimension. Also, its behavior as an ideal gas is also desirable.

3. Interferences
   Sample should be dry. Displacement gas evaporates water molecules and creates additional partial pressure (Flint and Flint, 2002). Temperature should be relatively constant because the method uses the ideal gas equation of state. Instrument should be calibrated when environmental conditions change.

4. Safety
   No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment
   5.1 Electronic balance, ±0.01-g sensitivity
   5.2 Oven, 110 °C
   5.3 Gas pycnometer, Penta-pycnometer, Quantachrome Instruments, Boynton Beach, FL

6. Reagents
   6.1 Helium gas

7. Procedure
   7.1 Oven-dry the soil sample at 110 °C overnight.
   7.2 Allow the instrument to warm up for at least 30 min prior to use.
   7.3 Set the regulator pressure to slightly over 20 PSIG.
   7.4 Validate the calibration by determining the volume of the calibration sphere.
   7.5 If volumes are outside specification range, recalibrate the instrument per instruction manual.
   7.6 Select the largest sample cell size of 135, 50, or 10 cc. The sample should fill at least half of the sample cell volume.
   7.7 Weigh the sample cell to the nearest 0.01 g. Place sample in the sample cell and reweigh to the nearest 0.01 g. The difference is the sample weight. Record the weight.
   7.8 Place the sample cells into the sample cell holder. Properly seal the cell holder with the cover.
   7.9 Define each cell with sample cell size, weight, and sample identification number.
7.10 Set the instrument to multi-run and enter the number of runs between 3 and 5.
7.11 Set the purge mode to 3 pulse cycles.
7.12 Start the sample run.
7.13 Record the average volume for each sample.
7.14 Calculate the particle density.

8. Calculations
Particle density (g cm\(^{-3}\)) = \(\frac{\text{Sample weight (g)}}{\text{Sample volume}}\)

9. Report
Report particle density (g cm\(^{-3}\)) to the nearest 0.01 unit on either the <2-mm or >2-mm particle-size fraction.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Atterberg Limits (3H)

<table>
<thead>
<tr>
<th>Liquid Limit (LL) (3H1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air-Dry, &lt;0.4 mm (3H1a1)</td>
</tr>
<tr>
<td>Field-Moist, &lt;0.4 mm (3H1b1)</td>
</tr>
</tbody>
</table>

Liquid Limit (LL) is the percent water content of a soil at the arbitrarily defined boundary between the liquid and plastic states. This water content is defined as the water content at which a pat of soil placed in a standard cup and cut by a groove of standard dimensions will flow together at the base of the groove for a distance of 13 mm (½ in) when subjected to 25 shocks from the cup being dropped 10 mm in a standard LL apparatus operated at a rate of 2 shocks s\(^{-1}\).
Refer to ASTM method D 4318 (ASTM, 2012). The LL is reported as percent water on a <0.4-mm basis (40-mesh) (method 3H1).

**Atterberg Limits (3H)**

**Plasticity Index (3H2)**

- **Air-Dry, <0.4 mm (3H2a1)**
- **Field-Moist, <0.4 mm (3H2b1)**

The plastic index (PI) is the range of water content over which a soil behaves plastically. Numerically, the PI is the difference in the water content between the LL and the plastic limit (PL). Refer to method 3H1 for the definition of LL. The PL is the percent water content of a soil at the boundary between the plastic and brittle states. The boundary is the water content at which a soil can no longer be deformed by rolling into 3.2-mm (⅛-in) threads without crumbling. Refer to ASTM method D 4318 (ASTM, 2012). The PI is reported as percent water on a <0.4-mm basis (method 3H2).

**References**


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**SOIL AND WATER CHEMICAL EXTRACTIONS AND ANALYSES (4)**

**Acid Standardization (4A)**

1. **Application**

   The process by which the concentration of a solution is accurately ascertained is known as standardization (Dean, 1995). Commonly, a solution is standardized by a titration in which it reacts with a weighed portion of a primary standard. The reaction between the titrant and the substance selected as a primary standard should fulfill the requirements for titrimetric analysis. In addition, a primary standard should be in a state of known high purity (typically <0.1 to 0.2% impurities), stable (easy to dry, not very hydroscopic, or does not lose weight upon exposure to air), and have a reasonably high equivalent weight in order to minimize the consequences of errors in weighing (Day and Underwood, 1980). For acid-base titrations, it is customary to prepare solutions of an acid and base.
of approximately the desired concentration and then to standardize one of the solutions against a primary standard. The solution thus standardized can be used as a secondary standard to obtain the normality of the other solution. Some widely used primary standards are as follows.

- **Benzoic acid**—($C_6H_5CO_2H$). Molecular weight = 122.123. Dissolve about 0.5 g in 50% ethanol and titrate to phenolphthalein end point.
- **Borax**—($Na_2B_4O_7\cdot10H_2O$). Molecular weight = 381.360. Use methyl red indicator. Dissolve in water. Borax forms a weak acid.
- **Mercuric oxide**—(HgO). Molecular weight = 216.599. Use bromthymol blue indicator. Dissolve 0.5 g HgO with 15 g of KBr in 25 mL of reverse osmosis deionized (RODI) water, excluding CO$_2$.
- **Potassium bicarbonate**—(KHCO$_3$). Molecular weight = 100.116. Use bromcresol green indicator. The first tint of green is the end point.
- **Potassium biphthalate**—(KHC$_8$H$_4$O$_4$). Molecular weight = 204.224. Use phenolphthalein indicator. Potassium biphthalate forms a weak acid.
- **Potassium bitartrate**—(KHC$_4$H$_4$O$_6$). Molecular weight = 188.178. Phenolphthalein indicator. Solutions of potassium bitartrate are susceptible to mold growth.
- **Sodium carbonate**—($Na_2CO_3$). Molecular weight = 105.988. Use bromocresol green indicator. The end point is the first green.

Refer to Table 1 for a list of some common acids and bases.

2. **Summary of Method**

Dissolve a known amount of the primary standard (e.g., $Na_2CO_3$) in reverse osmosis deionized (RODI) water. Prepare 10 of these $Na_2CO_3$ solutions plus 8 blanks and titrate with the acid to be standardized (e.g., HCl, H$_2$SO$_4$). Calculate the normality of the acid from the mean blank and titers. Report the normality and standard deviation for the acid standardization.

3. **Interferences**

Clean the glass electrode by rinsing with distilled water. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

Slow electrode response may cause the end point to be overshot. Cleaning the electrode with detergent solution may decrease the response time. If all else fails, change the electrode.

Dry primary standards and store in a desiccator to prevent hydration. Contamination of the primary standard may occur when drying, storing, or weighing the reagent. Use gloves when weighing the primary standard in the weighing vessel.
Table 1.—Common Commercial Strengths of Acids and Bases.

<table>
<thead>
<tr>
<th>Name</th>
<th>Molecular Weight</th>
<th>Moles per Liter</th>
<th>Grams per Liter</th>
<th>Percent by Weight</th>
<th>Specific Gravity</th>
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<tbody>
<tr>
<td>Acetic acid</td>
<td>60.05</td>
<td>17.4</td>
<td>1045</td>
<td>99.5</td>
<td>1.05</td>
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<tr>
<td>Glacial acetic acid</td>
<td>60.05</td>
<td>6.27</td>
<td>376</td>
<td>36</td>
<td>1.045</td>
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<td>Butyric acid</td>
<td>88.1</td>
<td>10.3</td>
<td>912</td>
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<td>Formic acid</td>
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<td>1080</td>
<td>90</td>
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<tr>
<td>Hydriodic acid</td>
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<td>1.18</td>
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<td>0.697</td>
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<td>642</td>
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<td>Hydrofluosilicic acid</td>
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<td>1008</td>
<td>71</td>
<td>1.42</td>
</tr>
<tr>
<td>Perchloric acid</td>
<td>100.5</td>
<td>11.65</td>
<td>1172</td>
<td>70</td>
<td>1.67</td>
</tr>
<tr>
<td>Phosphoric acid</td>
<td>98</td>
<td>14.7</td>
<td>1445</td>
<td>85</td>
<td>1.70</td>
</tr>
<tr>
<td>Sulfuric acid</td>
<td>98.1</td>
<td>18.0</td>
<td>1766</td>
<td>96</td>
<td>1.84</td>
</tr>
<tr>
<td>Sulfurous acid</td>
<td>82.1</td>
<td>0.74</td>
<td>61.2</td>
<td>6</td>
<td>1.02</td>
</tr>
<tr>
<td>Ammonia water</td>
<td>17.0</td>
<td>14.8</td>
<td>252</td>
<td>28</td>
<td>0.898</td>
</tr>
<tr>
<td>Potassium hydroxide</td>
<td>56.1</td>
<td>13.5</td>
<td>757</td>
<td>50</td>
<td>1.52</td>
</tr>
<tr>
<td>Sodium carbonate</td>
<td>106.0</td>
<td>1.04</td>
<td>110</td>
<td>10</td>
<td>1.10</td>
</tr>
<tr>
<td>Sodium hydroxide</td>
<td>40</td>
<td>19.1</td>
<td>763</td>
<td>50</td>
<td>1.53</td>
</tr>
</tbody>
</table>
4. Safety

Wear protective clothing and eye protection when preparing reagents. Dispense concentrated acids in a fume hood. Thoroughly wash hands after handling reagents. Be prepared to use safety showers and eyewash stations if necessary. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow the manufacturer’s safety precautions when using the automatic titrator. Review Material Safety Data Sheets (MSDS) for reagents.

5. Equipment

5.1 Electronic balance, ±0.10-mg sensitivity
5.2 Oven, 110 °C
5.3 Weighing vessel, 40 x 50 mm
5.4 Desiccator
5.5 Automatic titrator, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.
5.6 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.
5.7 Computer, with Titrino Workcell software, Metrohm Ltd., Brinkmann Instruments, Inc., and printer
5.8 Titration beakers, 250-mL, borosilicate, Metrohm Ltd., Brinkmann Instruments Inc.

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 pH buffers (4.00, 7.00, 9.18)
6.3 Sodium carbonate, anhydrous, 99.8% pure
6.4 Calcium sulfate (anhydrous) or equivalent desiccant

7. Procedure

7.1 Place a weighing vessel containing ≈10 g of the Na₂CO₃ in an oven at 110 °C overnight. Let cool in a desiccator to room temperature (≈30 min).
7.2 Refer to Acid Standardization Form. Weigh the weighing vessel plus the Na₂CO₃ designated in Section 7.1 to the nearest 0.1 mg (VW₁ₐ).
7.3 Tare a 250-mL titration beaker on the electronic balance. Add 0.15–0.55 g Na₂CO₃ to the beaker. Record the weight of the Na₂CO₃ to the nearest 0.1 mg (SW₁).
7.4 Verify the weighed SW₁ as follows:

7.4.1 Reweigh the weighing vessel plus Na₂CO₃ designated in Section 7.1 to the nearest 0.1 mg (VW₁b).

7.4.2 Calculate SWC₁ as follows: \( \text{VW₁a} - \text{VW₁b} = \text{SWC₁} \).

7.4.3 If SWC₁ calculated in Section 7.4b is within 0.5 mg of the weighed SW₁ determined in Section 7.3, proceed to Section 7.5; if not, return to Section 7.2 and begin again.

7.5 Prepare nine more Na₂CO₃ solutions (SW₂ ... SW₁₀) by this same procedure (Sections 7.2–7.4).

7.6 Add 100 mL of RODI water to each of the SW₁ ...... SW₁₀ and to eight empty beakers for blanks (B₁ ..... B₈) and gently swirl to facilitate dissolution of the salt.

7.7 Prepare the titration system according to the manufacturer’s instructions. Calibrate the pH electrode with 4.00, 7.00, and 9.18 pH buffers. Recalibrate electrode if \( R^2 < 0.950 \).

7.8 Set-up the automatic titrator to set end point titration mode. The “Set” pH parameters are listed as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measured value</td>
<td>pH</td>
</tr>
<tr>
<td>Titration rate</td>
<td>normal</td>
</tr>
<tr>
<td>Stop volume</td>
<td>100.0 mL</td>
</tr>
<tr>
<td>Stop end point</td>
<td>9</td>
</tr>
<tr>
<td>Stop potential</td>
<td>4.2 pH</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>High</td>
</tr>
<tr>
<td>Fixed end point</td>
<td>4.6 pH</td>
</tr>
<tr>
<td>Signal drift</td>
<td>25</td>
</tr>
<tr>
<td>Equilibration time</td>
<td>5 s</td>
</tr>
</tbody>
</table>

7.9 Titrate each of the SW₁ ...... SW₁₀ and B₁ ...... B₈ and record the titers.

8. Calculations

8.1 \( B_{\text{mean}} = \frac{B_{\text{sum}}}{n} \)

where:

\( B = \) Blank mean (mL)

\( B_{\text{sum}} = \) Sum of blank titers (mL)

\( n = \) Number of blanks (n=8)
8.2 \[ TCS_{1,...,10} = TS_{1,...,10} - B_m \]

where:
- \(TCS_{1,...,10}\) = Corrected sample titer (mL)
- \(TS_{1,...,10}\) = Sample titer (mL)
- \(B_m\) = Blank mean (mL)

8.3 \[ N_{1,...,10} = \left(\frac{SW_{1,...,10} \times 0.0188698 \text{ mole HCl/g Na}_2\text{CO}_3}{TCS_{1,...,10} \times 10^{-3} \text{ L/ml}}\right) = \frac{(SW_{1,...,10} \times 18.8698)}{TCS_{1,...,10}} \]

where:
- \(N_{1,...,10}\) = Normality calculated from titer (number of equivalents of solute per liter)
- \(SW_{1,...,10}\) = \(\text{Na}_2\text{CO}_3\) weight (g)

8.4 \[ N_{\text{mean}} = \frac{N_{\text{sum}}}{n} \]

where:
- \(N_{\text{mean}}\) = Normality mean
- \(N_{\text{sum}}\) = Sum of normalities
- \(n\) = Number of samples (n = 10)

8.5 Calculate standard deviation (Std) for \(N_{\text{mean}}\). If \(N_{\text{Std}} \leq 0.0005\), record \(N_{\text{mean}}\) to four decimals to right of decimal point. If \(N_{\text{Std}} > 0.0005\), discard no more than two suspicious values and return again to Sections 7.2 to 8.5 until 10 (n = 10) are achieved.

9. Report

Report the mean normality of acid \((N_{\text{mean}})\), standard deviation, and date of standardization.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


## Acid Standardization

<table>
<thead>
<tr>
<th>Acid:</th>
<th>Date:</th>
<th>Technician:</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Block</th>
<th>Vessel Weight</th>
<th>Sample Weight</th>
<th>Sample Titer</th>
<th>Blanks</th>
<th>Corrected Sample Titer</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1 Beginning Weight</td>
<td>( V_{WA1} )</td>
<td>( SW_1 )</td>
<td>( TS_1 )</td>
<td>( B_1 )</td>
<td>( TCS_1 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td>( B_2 )</td>
<td></td>
</tr>
<tr>
<td>#2 Beginning Weight</td>
<td>( V_{WA2} )</td>
<td>( SW_2 )</td>
<td>( TS_2 )</td>
<td>( B_3 )</td>
<td>( TCS_2 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td>( B_4 )</td>
<td></td>
</tr>
<tr>
<td>#3 Beginning Weight</td>
<td>( V_{WA3} )</td>
<td>( SW_3 )</td>
<td>( TS_3 )</td>
<td>( B_5 )</td>
<td>( TCS_3 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td>( B_6 )</td>
<td></td>
</tr>
<tr>
<td>#4 Beginning Weight</td>
<td>( V_{WA4} )</td>
<td>( SW_4 )</td>
<td>( TS_4 )</td>
<td>( B_7 )</td>
<td>( TCS_4 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td>( B_8 )</td>
<td></td>
</tr>
<tr>
<td>#5 Beginning Weight</td>
<td>( V_{WA5} )</td>
<td>( SW_5 )</td>
<td>( TS_5 )</td>
<td>( B_{sum} )</td>
<td>( TCS_5 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td>( B_{mean} )</td>
<td></td>
</tr>
<tr>
<td>#6 Beginning Weight</td>
<td>( V_{WA6} )</td>
<td>( SW_6 )</td>
<td>( TS_6 )</td>
<td></td>
<td>( TCS_6 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#7 Beginning Weight</td>
<td>( V_{WA7} )</td>
<td>( SW_7 )</td>
<td>( TS_7 )</td>
<td></td>
<td>( TCS_7 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#8 Beginning Weight</td>
<td>( V_{WA8} )</td>
<td>( SW_8 )</td>
<td>( TS_8 )</td>
<td></td>
<td>( TCS_8 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#9 Beginning Weight</td>
<td>( V_{WA9} )</td>
<td>( SW_9 )</td>
<td>( TS_9 )</td>
<td></td>
<td>( TCS_9 )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#10 Beginning Weight</td>
<td>( V_{WA10} )</td>
<td>( SW_{10} )</td>
<td>( TS_{10} )</td>
<td></td>
<td>( TCS_{10} )</td>
</tr>
<tr>
<td>Ending Weight</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[ N_{1-10} \]

\[ N_{Std} \]
Ion Exchange and Extractable Cations (4B)

Ion exchange is a reversible process by which one cation or anion held on the solid phase is exchanged with another cation or anion in the liquid phase. If two solid phases are in contact, ion exchange may also take place between two surfaces (Tisdale et al., 1985). In most agricultural soils, the cation exchange capacity (CEC) is generally considered to be more important than anion exchange (AEC). The anion molecular retention capacity of these soils is typically much smaller than the CEC (Tisdale et al., 1985). Some soils with abundant goethite and gibbsite, such as in some oxic horizons or subsoils of Oxisols (Soil Survey Staff, 2014), may have a CEC to AEC ratio approaching 1.0 (net charge of zero) or a small positive charge (Foth and Ellis, 1988).

Soil mineral and organic colloidal particles have negative valence charges that hold dissociable cations and thus are “colloidal electrolytes” (Jackson, 1958). The CEC is a measure of the quantity of readily exchangeable cations that neutralize negative charges in the soil (Rhoades, 1982). Cation exchange is a reversible reaction in soil solution, dependent upon negative charges of soil components arising from permanently charged or pH-dependent sites on organic matter and mineral colloid surfaces. The mechanisms for these negative charges are isomorphic substitution within layered silicate minerals; broken bonds at mineral edges and external surfaces; dissociation of acidic functional groups in organic compounds; and preferential adsorption of certain ions on particle surfaces (Rhoades, 1982). Isomorphic substitution produces permanent charge. The other charge mechanisms produce variable charge that is dependent on the soil solution phase as affected by soil pH, electrolyte level, valence of counter-ions, dielectric constant, and nature of anions (Rhoades, 1982). As a result of the variable charge in soils, the measurement of CEC is dependent on the method and conditions of determination. The method of determination is routinely reported with CEC data.

CEC is a measure of the total quantity of negative charges per unit weight of the material and is commonly expressed in units of milliequivalents per 100 g of soil (meq 100 g\(^{-1}\)) or centimoles per kg of soil (cmol(+) kg\(^{-1}\)). The KSSL reports cmol(+) kg\(^{-1}\) on a <2-mm basis. CEC can range from less than 1.0 to greater than 100 cmol(+) kg\(^{-1}\) soil. The term equivalent is defined as “1 gram atomic weight of hydrogen or the amount of any other ion that will combine with or displace this amount of hydrogen.” The milliequivalent weight of a substance is one thousandth of its atomic weight. Because the equivalent weight of hydrogen is about 1 gram, the term milliequivalent may be defined as “1 milligram of hydrogen or the amount of any other ion that will combine with or displace it” (Tisdale et al., 1985).

\[ S_{w} = W_{t4} - W_{t3} \]

Common CEC values for some soil components (NSSL Staff, 1975) are as follows:
<table>
<thead>
<tr>
<th>Soil Component</th>
<th>cmol(+) kg(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter</td>
<td>200 to 400</td>
</tr>
<tr>
<td>“Amorphous” clay</td>
<td>160 (at pH 8.2)</td>
</tr>
<tr>
<td>Vermiculite</td>
<td>100 to 150</td>
</tr>
<tr>
<td>Montmorillonite</td>
<td>60 to 100</td>
</tr>
<tr>
<td>Halloysite•4H(_2)O</td>
<td>40 to 50</td>
</tr>
<tr>
<td>Illite</td>
<td>20 to 40</td>
</tr>
<tr>
<td>Chlorite</td>
<td>10 to 40</td>
</tr>
<tr>
<td>Kaolinite</td>
<td>2 to 16</td>
</tr>
<tr>
<td>Halloysite•2H(_2)O</td>
<td>5 to 10</td>
</tr>
<tr>
<td>Sesquioxides</td>
<td>0</td>
</tr>
</tbody>
</table>

These very broad CEC ranges are intended only as general guidelines. More narrow groupings of CEC values are possible as data are continually collected and correlated. For example, the CEC of organic matter in Mollisols in the western United States ranges from 100 to 300 cmol (+) kg\(^{-1}\) (average 200), and the CEC of organic matter in Histosols ranges from 125 to 185 cmol (+) kg\(^{-1}\) and increases with decomposition of the organic matter (NSSL Staff, 1975).

Many procedures have been developed to determine CEC. These CEC measurements vary according to the nature of the cation employed, concentration of salt, and the equilibrium pH. The CEC measurement should not be thought of as highly exact but rather as an equilibrium measurement under the selected conditions (Jackson, 1958). Knowledge of the operational definition (procedure, pH, cation, and concentration) is necessary before evaluating the CEC measurement (Sumner and Miller, 1996). The more widely adopted methods of CEC determination are classified (Rhoades, 1982) as follows:

1. Cation summation
2. Direct displacement
3. Displacement after washing
4. Radioactive tracer

The KSSL performs a number of CEC methods, using several different reagents and pH levels. The most commonly reported methods by the KSSL are CEC–7 (4B1a1a1a1), CEC–8.2 (4B4b1), and effective cation exchange capacity (ECEC) (4B4b2). As a general rule, CEC–8.2 > CEC–7 > ECEC. Refer to Soil Survey Staff (2014) for the use and application of these CEC values in U.S. Soil Taxonomy.
Cation Exchange Capacity: \( \text{NH}_4\text{OAc, pH 7.0 (CEC–7)} \)

CEC–7 is a commonly used method (4B1a1a1a1) and has become a standard reference to which other methods are compared (Peech et al., 1947). Displacement after washing is the basis for this procedure. This CEC is determined by saturating the exchange sites with an index cation (\( \text{NH}_4^+ \)) by using a mechanical vacuum extractor (Holmgren et al., 1977); washing the soil free of excess saturated salt; displacing the index cation (\( \text{NH}_4^+ \)) adsorbed by the soil; and measuring the amount of the index cation (\( \text{NH}_4^+ \)). An advantage of using this method is that the extractant is highly buffered and therefore the extraction is performed at a constant and known pH (pH 7.0). In addition, the \( \text{NH}_4^+ \) on the exchange complex is easily determined. CEC–7 is an analytically determined value and is usually used for the calculation of CEC–7/clay ratios, although many Primary Characterization Data Sheets predating 1975 show CEC–8.2/clay.

Cation Exchange Capacity: Sum of Cations (CEC–8.2)

CEC–8.2 is calculated (4B4b1) by summing the \( \text{NH}_4\text{OAc} \) extractable bases (4B1a1b1-4) plus the \( \text{BaCl}_2\)-TEA extractable acidity (4B2b1a1). Cation summation is the basis for this procedure. CEC–8.2 minus CEC–7 is considered the pH dependent charge from pH 7.0 to pH 8.2. CEC–8.2 is not reported if carbonates, gypsum, or soluble salts are present in the soil because the \( \text{NH}_4\text{OAc} \) extracts cations from the dissolution of these soil constituents. CEC–8.2 is calculated as follows:

\[
\text{CEC–8.2} = \text{NH}_4\text{OAc extractable bases} + \text{Extractable acidity}
\]

Effective Cation Exchange Capacity: \( \text{NH}_4\text{OAc Extractable Bases} + \text{Aluminum} \)

CEC can be measured by extraction with an unbuffered salt. This measures the effective cation exchange capacity (ECEC), i.e., CEC at the normal soil pH (Coleman et al., 1958). Since the unbuffered salt solution, e.g., 1 N KCl, only affects the soil pH one unit or less, the extraction is determined at or near the soil pH and extracts only the cations held at active exchange sites at the particular pH of the soil. Neutral \( \text{NH}_4\text{OAc} \) extracts the same amounts of \( \text{Ca}^{2+}, \text{Mg}^{2+}, \text{Na}^+, \text{and K}^+ \) as KCl and therefore the method of extractable bases by \( \text{NH}_4\text{OAc} \) is used at the KSSL in place of KCl-extractable bases.

ECEC may be determined by extracting one soil sample with neutral normal \( \text{NH}_4\text{OAc} \) to determine the exchangeable basic cations (\( \text{Ca}^{2+}, \text{Mg}^{2+}, \text{Na}^+, \text{and K}^+ \)) and by extracting another sample of the same soil with 1.0 N KCl to determine the exchangeable Al. The 1 N KCl-extractable Al method approximates exchangeable Al and is a measure of “active” acidity present in soils with a 1:1 pH <5.5.
Aluminum is non-exchangeable at pH >5.5 due to hydrolysis, polymerization, and precipitation. For soils with pH <7.0, the ECEC should be less than the CEC measured with a buffered solution at pH 7.0. ECEC (4B4b2a) is calculated by summing the NH$_4$OAc bases (4B1a1b1-4) plus the KCl extractable Al (4B3a1a1) as follows:

$$ECEC = \text{NH}_4\text{OAc extractable bases} + \text{KCl-extractable Al}$$

**Effective Cation Exchange Capacity: NH$_4$Cl (ECEC)**

The CEC using a neutral unbuffered salt (NH$_4$Cl) is also an analytically determined value (4B1b1a1a1). The CEC by NH$_4$Cl provides an estimate of the ECEC of the soil (Peech et al., 1947). For a soil with a pH of <7.0, the ECEC value should be less than the CEC measured with a buffered solution at pH 7.0. The NH$_4$Cl CEC is about equal to the NH$_4$OAc extractable bases plus the KCl extractable Al for noncalcareous soils. This ECEC method is less commonly used at the KSSL.

**References**

National Soil Survey Laboratory Staff. 1975. Proposed tables for soil survey reports. RSSIU, USDA–SCS, Lincoln, NE.
Ion Exchange and Extractable Cations (4B)

Displacement after Washing, NH$_4$OAc, pH 7 (4B1a)

Automatic Extractor, 2 M KCl Rinse (4B1a1a)

Steam Distillation, HCl Titration (4B1a1a1a)

Cation Exchange Capacity (CEC–7) (4B1a1a1a1)

Air-dry or Field-Moist, <2 mm (4B1a1a1a1a-b1)

1. Application

The cation exchange capacity (CEC) determined with 1 N NH$_4$OAc buffered at pH 7.0 (CEC–7), is a commonly used method and has become a standard reference to which other methods are compared (Peech et al., 1947). The advantages of using this method are that the extractant is highly buffered and therefore the extraction is performed at a constant, known pH (7.0) and that the NH$_4^+$ on the exchange complex is easily determined.

2. Summary of Method

Displacement after washing is the basis for this procedure. The CEC is determined by saturating the exchange sites with an index cation (NH$_4^+$); washing the soil free of excess saturated salt; displacing the index cation (NH$_4^+$) adsorbed by the soil; and measuring the amount of the index cation (NH$_4^+$). A sample is leached using 1 N NH$_4$OAc and a mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed and saved for analyses of the cations. The NH$_4^+$ saturated soil is rinsed with ethanol to remove the NH$_4^+$ that was not adsorbed. The soil is then rinsed with 2 M KCl. This leachate is analyzed by steam distillation and titration to determine the NH$_4^+$ adsorbed on the soil exchange complex. The CEC by NH$_4$OAc, pH 7, is reported in meq 100 g$^{-1}$ or (cmol (+) kg$^{-1}$) soil in method 4B1a1a1a1.

3. Interferences

Incomplete saturation of the soil with NH$_4^+$ and insufficient removal of NH$_4^+$ are the greatest interferences to this method. Ethanol removes some adsorbed NH$_4^+$ from the exchange sites of some soils. Isopropanol rinses have been used for some soils in which ethanol removes adsorbed NH$_4^+$. Soils that contain large amounts of vermiculite can irreversibly “fix” NH$_4^+$. Soils that contain large amounts of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract. This method overestimates the “field” CEC of soils with pH <7 (Summer and Miller, 1996)

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents,
especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Nessler’s reagent contains mercury, which is toxic. Proper disposal of the Nessler’s reagent and clean-up of equipment in contact with the reagent are necessary.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjeltec Auto Analyzers.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity

5.2 Mechanical vacuum extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE (figs. 4B-1 and 4B-2)

5.3 Tubes, 60-mL, polypropylene, for extraction (0.45-µm filter), reservoir, and tared extraction tubes

Figure 4B-1.—Mechanical vacuum extractor, Sampletek, Mavco Industries, Lincoln, NE.
5.4 Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels
5.5 Kjeltec Auto 2300 Sampler System, Tecator, Perstorp Analytical
5.6 Digestion tubes, straight neck, 250 mL
5.7 Syringe filters, 0.45-μm, Whatman
5.8 Wash bottles
5.9 Vials, plastic
5.10 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Ammonium acetate solution (NH₄OAc), 1 N, pH 7.0. Add 1026 mL of glacial acetic acid (CH₃COOH) to 15 L RODI water. Add 1224 mL of concentrated
ammonium hydroxide (NH$_4$OH). Cool. Allow to stand one day to equilibrate to room temperature. Mix and adjust to pH 7.0 with CH$_3$COOH (typically, $\approx$40 mL) or NH$_4$OH and dilute with RODI water to 18 L.

6.3 Ethanol (CH$_3$CH$_2$OH), 95%, U.S.P.

6.4 Nessler's reagent. Add 4.56 g of potassium iodide (KI) to 30 mL RODI water. Add 5.68 g of mercuric iodide (Hgl$_2$). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL of RODI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with RODI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately. Store in brown bottle to protect from light.

6.5 Potassium chloride solution, 2 M. Add 1341.9 g of KCl reagent in 8 L RODI water. Allow solution to equilibrate to room temperature. Dilute to 9 L with RODI water.

6.6 Boric acid, 4% (w:v), with bromocresol green-methyl red indicator (0.075 % bromocresol green and 0.05% methyl red), Chempure

6.7 Hydrochloric acid (HCl), 0.1 N, standardized. Dilute 167 mL of concentrated HCl in 20 L of RODI water. Refer to procedure for standardization of acids.

6.8 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of RODI water. Dilute to 9 L with RODI water.

7. Procedure

**Extraction of Bases**

7.1 Weigh 2.5 g of <2-mm, air-dry soil to the nearest mg and place in a labeled extraction tube (ET). If sample is fine-grind, weigh 1 g to the nearest mg. If sample is moist, weigh enough soil to achieve $\approx$2.5 or 1 g, respectively, of air-dry soil. Prepare one quality-control check sample per 24 samples.

7.2 Place labeled ET on extractor and connect to corresponding tared extraction tube (TET$_{NH_4OAc}$) with rubber tubing.

7.3 Use wash bottle to rinse inside of ET with NH$_4$OAc. All soil should be wetted, and no air bubbles should be present. Shaking, swirling, or stirring may be required to wet organic samples. Fill ET to the 20-mL mark with NH$_4$OAc solution ($\approx$10 mL).

7.4 Secure reservoir tube (RT) to top of ET tube and let stand for 30 min. Extract the NH$_4$OAc solution at the 30-min rate until 2 mL of the solution remains above soil level. Turn off extractor. Do not let soil dry.

7.5 Add 40 mL of NH$_4$OAc solution to the RT. Set extractor for an overnight (12h) extraction. Extractor turns off automatically.
7.6 The next day, remove RT from top of extractor and place in a clean container. Carefully remove TET\textsubscript{NH\textsubscript{4}OAc}. Leave the rubber tubing on the ET. Weigh each TET\textsubscript{NH\textsubscript{4}OAc} containing the NH\textsubscript{4}OAc extract to the nearest mg.

7.7 Mix the extract in each TET\textsubscript{NH\textsubscript{4}OAc} by manually shaking. Fill a labeled plastic vial with extract solution and cap. Discard the excess properly. The solution in the vial is reserved for analyses of extracted cations (method 4B1a1b1-4) on the atomic absorption spectrophotometer (AAS). Some samples may be cloudy and need to be filtered prior to analysis on the AAS. If extracts are not to be determined immediately after collection, then store samples at 4 °C in plastic tubes.

7.8 Re-connect the TET\textsubscript{NH\textsubscript{4}OAc} with paired ET. Use a wash bottle to rinse the sides of the ET with ethanol to remove any remaining NH\textsubscript{4}OAc or soil particles adhering to the ET. All soil should be wetted, and no air bubbles should be present. Fill ET to the 20-mL mark with ethanol. Secure RT to top of ET tube and let stand for 30 min.

7.9 Extract the ethanol solution at the 30-min extraction rate until 2 mL of the solution remains above the soil level. Turn off the extractor. Do not let soil dry.

7.10 Add 45 mL of ethanol to the RT. Extract (≈45 min) the ethanol until 2 mL of this solution remains above the soil level. Turn off the extractor. Do not let soil dry. Disconnect the TET\textsubscript{NH\textsubscript{4}OAc} from the ET and discard the ethanol properly.

7.11 Re-connect the TET\textsubscript{NH\textsubscript{4}OAc} to the ET and add 55 mL of ethanol to the RT. Set the extractor for 45 min. Turn off the extractor. Remove the TET\textsubscript{NH\textsubscript{4}OAc}, leaving the tubing connected to the ET. Discard the ethanol properly.

7.14 After the final ethanol wash, collect a few drops of ethanol extract from the ET on a spot plate. Test for NH\textsubscript{4}\textsuperscript{+} by using Nessler’s reagent. A yellow, red, or reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler’s reagent. Repeat until a negative test is obtained.

7.15 Connect a new labeled extraction tube (ET\textsubscript{KCl}) with rubber tubing to ET on extractor.

7.16 Use a wash bottle to rinse inside of ET with 2 M KCl to remove any remaining ethanol or soil particles adhering to the ET. All soil should be wetted, and no air bubbles should be present. Fill ET to the 20-mL mark with KCl solution and let stand for 30.
7.17 Extract the KCl solution at the 30-min rate until 2 mL of the solution remains above soil level. Turn off extractor. Do not let soil dry.

7.18 Secure RT to top of ET tube. Add 40 mL KCl solution to RT and set the extract for 45 min. Remove the ET and ET\textsubscript{KCl} from the extractor.

**Steam Distillation: Setup, Operation, and Analysis**

7.19 Transfer the contents of the ET\textsubscript{KCl} to a 250-mL digestion tube. If extracts are not to be determined immediately after collection, then store samples at 4 °C.

7.20 Refer to the manufacturer’s manual for operation of the distillation unit. The following are only very general guidelines for instrument conditions.

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<thead>
<tr>
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<th></th>
</tr>
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</tr>
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<td>Time</td>
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</tr>
<tr>
<td>Distillation</td>
<td>Volume</td>
</tr>
<tr>
<td>Tube Drain</td>
<td>Yes</td>
</tr>
</tbody>
</table>

7.21 When using new reagents (e.g., boric acid), reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.22 Record the normality of standardized acid.

7.23 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

**8. Calculations**

\[ \text{CEC} = \frac{[\text{Titer} \times N \times 100 \times R]}{[\text{Sample Weight (g)}]} \]

where:

- CEC = Cation Exchange Capacity (meq 100 g\textsuperscript{−1})
- Titer = Titer of sample (mL)
- \(N\) = Normality of HCl titrant
- 100 = Conversion factor to 100-g basis
R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

9. Report
Report CEC–7 to the nearest 0.1 meq 100 g⁻¹ (cmol (+) kg⁻¹).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Ion Exchange and Extractable Cations (4B)
Displacement after Washing, NH₄Cl (4B1b)
   Automatic Extractor, 2 M KCl Rinse (4B1b1a)
   Steam Distillation, HCl Titration (4B1b1a1a)
   Cation Exchange Capacity (4B1b1a1a1)
   Air-dry or Field-Moist, <2 mm (4B1b1a1a1a-b1)

1. Application
The cation exchange capacity (CEC) determined with a neutral cation unbuffered salt, e.g., 1 N NH₄Cl, is an estimate of the "effective" CEC (ECEC) of the soil (Peech et al., 1947). For a soil with a pH of <7.0, the ECEC values should be <CEC measured with a buffered solution at pH 7.0. The NH₄Cl CEC is approximately equal to the NH₄OAc extractable bases plus the KCl extractable Al for noncalcareous soils.

2. Summary of Method
Displacement after washing is the basis for this procedure. The CEC is determined by saturating the exchange sites with an index cation (NH₄⁺); washing the soil free of excess saturated salt; displacing the index cation (NH₄⁺) adsorbed by the soil; and measuring the amount of the index cation (NH₄⁺). A sample is leached using 1 N NH₄Cl and a mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed and saved for analyses of the cations. The NH₄⁺ saturated soil is rinsed with ethanol to remove the NH₄⁺ that was not adsorbed.
The soil is then rinsed with 2 M KCl. This leachate is analyzed by steam distillation and titration to determine the NH$_4^+$ adsorbed on the soil exchange complex. The CEC by NH$_4$Cl is reported as meq 100 g$^{-1}$ or (cmol (+) kg$^{-1}$) soil in method 4B1b1a1a1.

3. Interferences

Incomplete saturation of the soil with NH$_4^+$ and insufficient removal of NH$_4^+$ are the greatest interferences to this method. Ethanol removes some adsorbed NH$_4^+$ from the exchange sites of some soils. Isopropanol rinses have been used for some soils in which ethanol removes adsorbed NH$_4^+$. Soils that contain large amounts of vermiculite can irreversibly “fix” NH$_4^+$. Soils that contain large amounts of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Nessler’s reagent contains mercury, which is toxic. Proper disposal of the Nessler’s reagent and clean-up of equipment in contact with the reagent is necessary.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjeltec Auto Analyzers.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical Vacuum Extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE (figs. 4B-1 and 4B-2)
5.3 Tubes, 60-mL, polypropylene, for extraction (0.45-µm filter), reservoir, and tared extraction tubes
5.4 Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels.
5.5 Kjeltec Auto 2300 Sampler System, Tecator, Perstorp Analytical
5.6 Digestion tubes, straight neck, 250 mL
5.7 Syringe filters, 0.45-µm, Whatman
5.8 Wash bottles
5.9 Vials, plastic
5.10 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Ammonium chloride solution (NH₄Cl), 1 N. Dissolve 535 g of NH₄Cl reagent in RODI water and dilute to 10 L.
6.3 Ethanol (CH₃CH₂OH), 95%, U.S.P.
6.4 Nessler’s reagent. Add 4.56 g of potassium iodide (KI) to 30 mL RODI water. Add 5.68 g of mercuric iodide (HgI₂). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL of RODI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with RODI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately. Store in brown bottle to protect from light.
6.5 Potassium chloride solution, 2 N. Add 1341.9 g of KCl reagent in 8 L RODI water. Allow solution to equilibrate to room temperature. Dilute to 9 L with RODI water.
6.6 Boric acid, 4% (w:v), with brom cresol green-methyl red indicator (0.075% brom cresol green and 0.05% methyl red), Chempure
6.7 Hydrochloric acid (HCl), 0.1 N, standardized. Dilute 167 mL of concentrated HCl in 20 L of RODI water. Refer to procedure for standardization of acids.
6.8 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of RODI water. Dilute to 9 L with RODI water.

7. Procedure

Extraction of Bases

7.1 Weigh 2.5 g of <2-mm, air-dry soil to the nearest mg and place in a labeled extraction tube (ET). If sample is fine-grind, weigh 1 g to the nearest mg. If sample is moist, weigh enough soil to achieve ≈2.5 or 1 g, respectively, of air-dry soil. Prepare one quality control check sample per 24 samples.
7.2 Place labeled ET on extractor and connect to corresponding tared extraction tube (TETNH₄Cl) with rubber tubing.
7.3 Use wash bottle to rinse inside of ET with NH₄Cl. All soil should be wetted, and no air bubbles should be present. Shaking, swirling, or stirring may be required to wet organic samples. Fill ET to the 20-mL mark with NH₄Cl solution (≈10 mL).
7.4 Secure reservoir tube (RT) to top of ET tube and let stand for 30 min. Extract the NH$_4$Cl solution at the 30-min rate until 2 mL of this solution remains above soil level. Turn off extractor. Do not let soil dry.

7.5 Add 40 mL of NH$_4$Cl solution to the RT. Set extractor for an overnight (12 h) extraction. Extractor turns off automatically.

7.6 The next day, remove RT from top of extractor and place in a clean container. Carefully remove TET$_{NH_4Cl}$. Leave the rubber tubing on the ET. Weigh each TET$_{NH_4Cl}$ containing the NH$_4$Cl extract to the nearest mg.

7.7 Mix the extract in each TET$_{NH_4Cl}$ by manually shaking. Fill a labeled plastic vial with extract solution and cap. Discard the excess properly. The solution in the vial is reserved for analyses of extracted cations (method 4B1b1b1-4) on the atomic absorption spectrophotometer (AAS). Some samples may be cloudy and need to be filtered prior to analysis on the AAS. If extracts are not to be determined immediately after collection, then store samples at 4 °C in plastic tubes.

**Removal of Excess Ammonium Chloride**

7.8 Re-connect the TET$_{NH_4Cl}$ with paired ET. Use a wash bottle to rinse the sides of the ET with ethanol to remove any remaining NH$_4$Cl or soil particles adhering to the ET. All soil should be wetted, and no air bubbles should be present. Fill ET to the 20-mL mark with ethanol. Secure RT to top of ET and let stand for 30 min.

7.9 Extract the ethanol solution at the 30-min rate until 2 mL of this solution remains above the soil level. Turn off the extractor. Do not let soil dry.

7.10 Add 45 mL of ethanol to the RT. Extract (≈45 min) the ethanol until 2 mL of this solution remains above the soil level. Turn off the extractor. Do not let soil dry. Disconnect the TET$_{NH_4Cl}$ from the ET and discard the ethanol properly.

7.11 Re-connect the TET$_{NH_4Cl}$ to the ET and add 55 mL of ethanol to the RT. Set the extractor for 45 min. Turn off the extractor. Remove the TET$_{NH_4Cl}$ leaving the tubing connected to the ET. Discard the ethanol properly.

7.12 After the final ethanol wash, collect a few drops of ethanol extract from the ET on a spot plate. Test for NH$_4^+$ by using Nessler’s reagent. A yellow, red, or reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler’s reagent. Repeat until a negative test is obtained.

**2 M KCl Rinse**

7.13 Connect a new labeled extraction tube (ET$_{KCl}$) with rubber tubing to ET on extractor.
7.14 Use a wash bottle to rinse inside of ET with 2 M KCl to remove any remaining ethanol or soil particles adhering to the ET. All soil should be wetted, and no air bubbles should be present. Fill ET to the 20-mL mark with KCl solution and let stand for 30 min.

7.15 Extract the KCl solution at the 30-min extraction rate until 2 mL of the solution remains above soil level. Turn off extractor. Do not let soil dry.

7.16 Secure RT to top of ET tube. Add 40 mL KCl solution to RT and set the extract for 45 min. Remove the ET and ET\text{KCl} from the extractor.

**Steam Distillation: Setup, Operation, and Analysis**

7.17 Transfer the contents of the ET\text{KCl} to a 250-mL digestion tube. If extracts are not to be determined immediately after collection, then store samples at 4 °C.

7.18 Refer to the manufacturer’s manual for operation of the distillation unit. The following are only very general guidelines for instrument conditions.

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<td>Time</td>
</tr>
<tr>
<td>Distillation</td>
</tr>
<tr>
<td>Tube Drain</td>
</tr>
</tbody>
</table>

7.19 When using new reagents (e.g., boric acid), reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.20 Record the normality of standardized acid.

7.21 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

\[
\text{CEC} = \frac{\text{Titer} \times \text{N} \times 100 \times \text{R}}{\text{Sample Weight (g)}}
\]

where:

\[
\text{CEC} = \text{Cation Exchange Capacity (meq 100 g}^{-1})
\]
Ion Exchange and Extractable Cations (4B)
Displacement after Washing, NH₄OAc, pH 7 (4B1a)
Automatic Extractor (4B1a1)
Atomic Absorption Spectrophotometer (4B1a1b)
Calcium, Magnesium, Potassium, and Sodium (4B1a1b1-4)
Air-dry or Field-Moist, <2-mm (4B1a1b1-4a-b1)

1. Application
The extractable bases (Ca²⁺, Mg²⁺, K⁺, and Na⁺) from the NH₄OAc extraction (method 4B1a1) are generally assumed to be those exchangeable bases on the cation exchange sites of the soil. The term “extractable” rather than “exchangeable” bases is used because any additional source of soluble bases influences the results. The abundance of these cations usually occurs in the sequence of Ca²⁺ > Mg²⁺ > K⁺ > Na⁺. Deviation from this usual order signals that some factor or factors, e.g., free CaCO₃ or gypsum, serpentine (high Mg²⁺), or natric material (high Na⁺), have altered the soil chemistry. The most doubtful cation extractions with this method are Ca²⁺ in the presence of free CaCO₃ and gypsum and K⁺ in soils that are dominated by mica or vermiculite (Thomas, 1982).

2. Summary of Method
The NH₄OAc extract from method 4B1a1 is diluted with an ionization suppressant (La₂O₃). The analytes are measured by an atomic absorption spectrophotometer (AAS). The analyte is measured by absorption of the light from
a hollow cathode lamp. An automatic sample changer is used to aspirate a series
of samples. The AAS converts absorption to analyte concentration. Data are
automatically recorded by a microcomputer and printer. The NH₄OAc extracted
cations, Ca²⁺, Mg²⁺, K⁺, and Na⁺, are reported in meq 100 g⁻¹ soil or (cmol (+) kg⁻¹)
in methods 4B1a1b1-4, respectively.

3. Interferences

Four types of interferences (matrix, spectral, chemical, and ionization) affect
the analyses of these cations. These interferences vary in importance, depending
upon the particular analyte selected. Do not use borosilicate tubes because of
potential leaching of analytes.

4. Safety

Wear protective clothing and safety glasses. Exercise special care when
preparing reagents. Restrict the use of concentrated HCl to a fume hood. Many
metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash
hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases.
Gas cylinders should be chained or bolted in an upright position. Acetylene
gas is highly flammable. Avoid open flames and sparks. Standard laboratory
equipment includes fire blankets and extinguishers for use if necessary. Follow
the manufacturer’s safety precautions when using the AAS.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Atomic absorption spectrophotometer (AAS), double-beam optical system,
AAAnalyst, 400, Perkin-Elmer Corp., Norwalk, CT
5.3 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.4 Computer, with AA WinLab software, Perkin-Elmer Corp., Norwalk, CT, and
printer
5.5 Single-stage regulator, acetylene
5.6 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight,
Microlab 500, Hamilton Co., Reno, NV
5.7 Plastic test tubes, 15-mL, 16 mm x 100, for sample dilution and sample
changer
5.8 Containers, polyethylene
5.9 Peristaltic pump

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent
water
6.2 Hydrochloric acid (HCl), concentrated 12 N

6.3 HCl, 1:1 HCl:RODI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part RODI water.

6.4 NH₄OH, reagent-grade, specific gravity 0.90

6.5 Glacial acetic acid, 99.5%

6.6 Ammonium acetate solution (NH₄OAc), 1 N, pH 7.0. Add 1026 mL of glacial acetic acid (CH₃COOH) to 15 L RODI water. Add 1224 mL of concentrated ammonium hydroxide (NH₄OH). Cool. Allow to stand one day to equilibrate to room temperature. Mix and adjust to pH 7.0 with CH₃COOH (typically, ≈40 mL) or NH₄OH and dilute with RODI water to 18 L. The NH₄OAc solution is used for extraction of cations (method 4B1a1).

6.7 NH₄OAc solution, 2.0 N, pH 7.0. Mix 228 mL of glacial acetic acid in 1200 mL of RODI water. While stirring, carefully add 272 mL of concentrated NH₄OH. Cool. Allow to stand one day to equilibrate to room temperature. Mix and adjust pH 7.0 using CH₃COOH or NH₄OH. Dilute to 2 L with RODI water.

6.8 Stock lanthanum ionization suppressant solution (SLISS), 65,000 mg L⁻¹. Wet 152.4 g lanthanum oxide (La₂O₃) with 100 mL RODI water. Slowly and cautiously add 500 mL of 6 N HCl to dissolve the La₂O₃. Cooling the solution is necessary. Dilute to 2 L with RODI water. Filter solution. Store in polyethylene container.

6.9 Working lanthanum ionization suppressant solution (WLISS), 2000 mg L⁻¹. Dilute 61.5 mL of SLISS with 1800 mL of RODI water (1:10). Dilute to 2 L with RODI water. Store in polyethylene container.

6.10 Primary stock standards solutions (PSSS), high purity, 1000 mg L⁻¹: Ca, Mg, K, and Na.

1.11 Working stock mixed standards solution (WSMSS), High, Medium, Low, Low/Low, and Blank. In five 500-mL volumetric flasks, add 250 mL of 2 N NH₄OAc and the following designated amounts of Ca PSSS, Mg PSSS, K PSSS, and Na PSSS. Dilute to volume with RODI. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use. Prepare WSMSS as follows:

6.11.1 High Standard WSMSS: 90 mL Ca PSSS, 7.5 mL Mg PSSS, 20.0 mL K PSSS, and 100.0 mL Na PSSS = 180 mg L⁻¹ Ca, 15 mg L⁻¹ Mg, 40 mg L⁻¹ K, and 200 mg L⁻¹ Na

6.11.2 Medium Standard WSMSS: 60 mL Ca PSSS, 5.0 mL Mg PSSS, 10.0 mL K PSSS, and 50.0 mL Na PSSS = 120 mg L⁻¹ Ca, 10 mg L⁻¹ Mg, 20 mg L⁻¹ K, and 100 mg L⁻¹ Na
6.11.3 Low Standard WSMSS: 30 mL Ca PSSS, 2.5 mL Mg PSSS, 5.0 mL K PSSS, and 10.0 mL Na PSSS = 60 mg L\(^{-1}\) Ca, 5 mg L\(^{-1}\) Mg, 10 mg L\(^{-1}\) K, and 20 mg L\(^{-1}\) Na

6.11.4 Low/Low Standard WSMSS: 12.5 mL Ca PSSS, 0.25 mL Mg PSSS, 0.30 mL K PSSS, and 5.0 mL Na PSSS = 25 mg L\(^{-1}\) Ca, 0.5 mg L\(^{-1}\) Mg, 0.60 mg L\(^{-1}\) K, and 10 mg L\(^{-1}\) Na

6.11.5 Blank WSMSS: 0 mL of Ca, Mg, K, and Na PSSS.

6.12 Mixed calibration standard solutions (MCSS), High, Medium, Low, Very Low, and Blank. Dilute 1 part WSMSS with 19 parts of WLISS (1:20) dilution with resulting concentrations for MCSS as follows:

6.12.1 MCSS High Standard: 9.0 mg L\(^{-1}\) Ca, 0.75 mg L\(^{-1}\) Mg, 2.0 mg L\(^{-1}\) K, and 10.0 mg L\(^{-1}\) Na

6.12.2 MCSS Medium Standard: 6.0 mg L\(^{-1}\) Ca, 0.5 mg L\(^{-1}\) Mg, 1.0 mg L\(^{-1}\) K, and 5.0 mg L\(^{-1}\) Na

6.12.3 MCSS Low Standard: 3.0 mg L\(^{-1}\) Ca, 0.25 mg L\(^{-1}\) Mg, 0.5 mg L\(^{-1}\) K, and 1.0 mg L\(^{-1}\) Na

6.12.4 MCSS Very Low Standard: 1.25 mg L\(^{-1}\) Ca, 0.025 mg L\(^{-1}\) Mg, 0.030 mg L\(^{-1}\) K, and 0.5 mg L\(^{-1}\) Na

6.12.5 Blank MCSS: 0 mg L\(^{-1}\) Ca, Mg, K, and Na

6.13 Compressed air with water and oil traps

6.14 Acetylene gas, purity 99.6%

7. Procedure

Dilution of Calibration Standards and Sample Extracts

7.1 The 10-mL syringe is for diluent (WLISS). The 1-mL syringe is for the MCSS and NH\(_4\)OAc extracts (method 4B1a1). Set the digital diluter at a 1:20 dilution. See reagents for preparation of the MCSS (High, Medium, Low, Very Low, and Blank). Dilute 1 part NH\(_4\)OAc sample extract with 19 parts of WLISS (1:20 dilution).

7.2 Dispense the diluted sample solutions into test tubes which have been placed in the sample holders of the sample changer.

AAS Set-up and Operation

7.3 Refer to the manufacturer’s manual for operation of the AAS. The following are only very general guidelines for instrument conditions for the various analytes.
7.4 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

**AAS Calibration and Analysis**

7.5 Calibrate the instrument by using the MCSS (High, Medium, Low, Very Low, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.6 If sample exceeds calibration standard, the sample is diluted 1:5, 1:20, 1:100, etc., with 1 N $\text{NH}_4\text{OAc}$ followed by 1:20 dilution with WLISS.

7.7 Perform one quality control (QC) (Low Standard MCSS) every 12 samples. If reading is not within 10%, the instrument is re-calibrated and QC re-analyzed.

7.8 Record analyte readings to 0.01 unit.

8. Calculations

The instrument readings for analyte concentration are in mg L$^{-1}$. These analyte concentrations are converted to meq 100 g$^{-1}$ as follows:

Soil Analyte Concentration (meq 100 g$^{-1}$) =
$$\frac{(A \times (B_1 - B_2)/B_3) \times C \times R \times 100}{(1000 \times E \times F)}$$

where:
- $A$ = Analyte (Ca, Mg, K, Na) concentration in extract (mg L$^{-1}$)
- $B_1$ = Weight of extraction syringe and extract (g)
- $B_2$ = Weight of tared extraction syringe (g)
- $B_3$ = Density of 1 N $\text{NH}_4\text{OAc}$ at 20 °C (1.0124 g cm$^{-3}$)
- $C$ = Dilution, if performed
- 100 = Conversion factor (100-g basis)
- $R$ = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
1000 = mL L⁻¹
E = Soil sample weight (g)
F = Equivalent weight (mg meq⁻¹)

where:
Ca⁺² = 20.04 mg meq⁻¹
Mg⁺² = 12.15 mg meq⁻¹
Na⁺¹ = 22.99 mg meq⁻¹
K⁺¹ = 39.10 mg meq⁻¹

9. Report
Report the extractable Ca²⁺, Mg²⁺, Na⁺, and K⁺ to the nearest 0.1 meq 100 g⁻¹ (cmol (+) kg⁻¹).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Ion Exchange and Extractable Cations (4B)
Displacement after Washing, NH₄Cl (4B1b)
Automatic Extractor (4B1b1)
Atomic Absorption Spectrophotometer (4B1b1b)
Calcium, Magnesium, Potassium, and Sodium (4B1b1b1-4)
Air-dry or Field-Moist, <2-mm (4B1b1b1-4a-b1)

1. Application
The extractable bases (Ca²⁺, Mg²⁺, Na⁺, and K⁺) from the NH₄Cl extraction (method 4B1b1) are generally assumed to be those exchangeable bases on the cation exchange sites of the soil. The abundance of these cations usually occurs in the sequence of Ca²⁺ > Mg²⁺ > K⁺ > Na⁺. Deviation from this usual order signals that some factor or factors, e.g., free CaCO₃ or gypsum, serpentine (high Mg²⁺), or natric material (high Na⁺), have altered the soil chemistry. The most doubtful cation extractions with this method are Ca²⁺ in the presence of free CaCO₃ or gypsum and K⁺ in soils that are dominated by mica or vermiculite (Thomas, 1982).
2. Summary of Method

The NH$_4$Cl extract from method 4B1b1 is diluted with an ionization suppressant (La$_2$O$_3$). The analytes are measured by an atomic absorption spectrophotometer (AAS). The analyte is measured by absorption of the light from a hollow cathode lamp. An automatic sample changer is used to aspirate a series of samples. The AAS converts absorption to analyte concentration. Data are automatically recorded by a microcomputer and printer. The NH$_4$Cl extracted cations, Ca$^{2+}$, Mg$^{2+}$, K$^+$, and Na$^+$, are reported in meq 100 g$^{-1}$ soil or (cmol (+) kg$^{-1}$) in methods 4B1b1b1-4, respectively.

3. Interferences

Four types of interferences (matrix, spectral, chemical, and ionization) affect the analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected. Do not use borosilicate tubes because of potential leaching of analytes.

4. Safety

Wear protective clothing and safety glasses. Exercise special care when preparing reagents. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory methods when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the AAS.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Atomic absorption spectrophotometer (AAS), double-beam optical system, Analyst, 400, Perkin-Elmer Corp., Norwalk, CT
5.3 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.4 Computer, with AA WinLab software, Perkin-Elmer Corp., Norwalk, CT, and printer
5.5 Single-stage regulator, acetylene
5.6 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.7 Plastic test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.8 Containers, polyethylene
5.9 Peristaltic pump
6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 Hydrochloric acid (HCl), concentrated 12 N

6.3 HCl, 1:1 HCl:RODI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part RODI water.

6.4 Ammonium chloride solution (NH\textsubscript{4}Cl), 1 N. Dissolve 535 g of NH\textsubscript{4}Cl reagent in RODI water and dilute to 10 L.

6.5 Stock lanthanum ionization suppressant solution (SLISS), 65,000 mg L\textsuperscript{-1}. Wet 152.4 g lanthanum oxide (La\textsubscript{2}O\textsubscript{3}) with 100 mL RODI water. Slowly and cautiously add 500 mL of 6 N HCl to dissolve the La\textsubscript{2}O\textsubscript{3}. Cooling the solution is necessary. Dilute to 2 L with RODI water. Filter solution. Store in polyethylene container.

6.6 Working lanthanum ionization suppressant solution (WLISS), 2000 mg L\textsuperscript{-1}. Dilute 61.5 mL of SLISS with 1800 mL of RODI water (1:10). Dilute to 2 L with RODI water. Store in polyethylene container.

6.7 Primary stock standards solutions (PSSS), high purity, 1000 mg L\textsuperscript{-1}: Ca, Mg, K, and Na.

6.8 Working stock mixed standards solution (WSMSS), High, Medium, Low, Very Low, and Blank. In five 500-mL volumetric flasks, add 250 mL of 2 N NH\textsubscript{4}Cl and the following designated amounts of Ca PSSS, Mg PSSS, K PSSS, and Na PSSS. Dilute to volume with RODI. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use. Prepare WSMSS as follows:

6.8.1 High Standard WSMSS: 90 mL Ca PSSS, 7.5 mL Mg PSSS, 20.0 mL K PSSS, and 100.0 mL Na PSSS = 180 mg L\textsuperscript{-1} Ca, 15 mg L\textsuperscript{-1} Mg, 40 mg L\textsuperscript{-1} K, and 200 mg L\textsuperscript{-1} Na

6.8.2 Medium Standard WSMSS: 60 mL Ca PSSS, 5.0 mL Mg PSSS, 10.0 mL K PSSS, and 50.0 mL Na PSSS = 120 mg L\textsuperscript{-1} Ca, 10 mg L\textsuperscript{-1} Mg, 20 mg L\textsuperscript{-1} K, and 100 mg L\textsuperscript{-1} Na

6.8.3 Low Standard WSMSS: 30 mL Ca PSSS, 2.5 mL Mg PSSS, 5.0 mL K PSSS, and 10.0 mL Na PSSS = 60 mg L\textsuperscript{-1} Ca, 5 mg L\textsuperscript{-1} Mg, 10 mg L\textsuperscript{-1} K, and 20 mg L\textsuperscript{-1} Na

6.8.4 Very Low Standard WSMSS: 12.5 mL Ca PSSS, 0.25 mL Mg PSSS, 0.30 mL K PSSS, and 5.0 mL Na PSSS = 25 mg L\textsuperscript{-1} Ca, 0.5 mg L\textsuperscript{-1} Mg, 0.60 mg L\textsuperscript{-1} K, and 10 mg L\textsuperscript{-1} Na

6.8.5 Blank WSMSS: 0 mL of Ca, Mg, K, and Na PSSS
6.9 Mixed calibration standard solutions (MCSS), High, Medium, Low, Very Low, and Blank. Dilute 1 part WSMSS with 19 parts of WLISS (1:20) dilution with resulting concentrations for MCSS as follows:

6.9.1 MCSS High Standard: 9.0 mg L\(^{-1}\) Ca, 0.75 mg L\(^{-1}\) Mg, 2.0 mg L\(^{-1}\) K, and 10.0 mg L\(^{-1}\) Na

6.9.2 MCSS Medium Standard: 6.0 mg L\(^{-1}\) Ca, 0.5 mg L\(^{-1}\) Mg, 1.0 mg L\(^{-1}\) K, and 5.0 mg L\(^{-1}\) Na

6.9.3 MCSS Low Standard: 3.0 mg L\(^{-1}\) Ca, 0.25 mg L\(^{-1}\) Mg, 0.5 mg L\(^{-1}\) K, and 1.0 mg L\(^{-1}\) Na

6.9.4 MCSS Very Low Standard: 1.25 mg L\(^{-1}\) Ca, 0.025 mg L\(^{-1}\) Mg, 0.030 K, and 0.5 mg L\(^{-1}\) Na

6.9.5 Blank MCSS: 0 mg L\(^{-1}\) Ca, Mg, K, and Na

6.10 Compressed air with water and oil traps

6.11 Acetylene gas, purity 99.6%

7. Procedure

Dilution of Calibration Standards and Sample Extracts

7.1 The 10-mL syringe is for diluent (WLISS). The 1-mL syringe is for the MCSS and NH\(_4\)Cl extracts (method 4B1b). Set the digital diluter at a 1:20 dilution. See reagents for preparation of the MCSS (High, Medium, Low, Very Low, and Blank). Dilute 1 part NH\(_4\)Cl sample extract with 19 parts of WLISS (1:20 dilution).

7.2 Dispense the diluted sample solutions into test tubes that have been placed in the sample holders of the sample changer.

AAS Set-up and Operation

7.3 Refer to the manufacturer’s manual for operation of the AAS. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Conc.</th>
<th>Burner &amp; angle</th>
<th>Wavelength</th>
<th>Slit</th>
<th>Fuel/Oxidant (C(_2)H(_2)/Air)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mg L(^{-1}))</td>
<td>(nm)</td>
<td>(mm)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca</td>
<td>9.0</td>
<td>10 cm @ 0°</td>
<td>422.7</td>
<td>0.7</td>
<td>1.5/10.0</td>
</tr>
<tr>
<td>Mg</td>
<td>0.75</td>
<td>10 cm @ 0°</td>
<td>285.2</td>
<td>0.7</td>
<td>1.5/10.0</td>
</tr>
<tr>
<td>K</td>
<td>2.0</td>
<td>10 cm @ 0°</td>
<td>766.5</td>
<td>0.7</td>
<td>1.5/10.0</td>
</tr>
<tr>
<td>Na</td>
<td>10.0</td>
<td>10 cm @ 30°</td>
<td>589.0</td>
<td>0.2</td>
<td>1.5/10.0</td>
</tr>
</tbody>
</table>
7.4 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

**AAS Calibration and Analysis**

7.5 Calibrate the instrument by using the MCSS (High, Medium, Low, Very Low, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.6 If sample exceeds calibration standard, the sample is diluted 1:5, 1:20, 1:100, etc., with 1 N NH$_4$Cl followed by 1:20 dilution with WLISS.

7.7 Perform one quality control (QC) (Low Standard MCSS) every 12 samples. If reading is not within 10%, the instrument is re-calibrated and QC re-analyzed.

7.8 Record analyte readings to 0.01 unit.

8. Calculations

The instrument readings for analyte concentration are in mg L$^{-1}$. These analyte concentrations are converted to meq 100 g$^{-1}$ as follows:

Soil Analyte Concentration (meq 100 g$^{-1}$) =

$$\frac{A \times \left(\frac{B_1 - B_2}{B_3}\right) \times C \times R \times 100}{1000 \times E \times F}$$

where:

- $A =$ Analyte (Ca, Mg, K, Na) concentration in extract (mg L$^{-1}$)
- $B_1 =$ Weight of extraction syringe and extract (g)
- $B_2 =$ Weight of tared extraction syringe (g)
- $B_3 =$ Density of 1 N NH$_4$Cl at 20 °C (1.0166 g cm$^{-3}$)
- $C =$ Dilution, if performed
- 100 = Conversion factor (100-g basis)
- $R =$ Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = mL L$^{-1}$
- $E =$ Soil sample weight (g)
- $F =$ Equivalent weight (mg meq$^{-1}$)

where:

- Ca$^{2+}$ = 20.04 mg meq$^{-1}$
- Mg$^{2+}$ = 12.15 mg meq$^{-1}$
- Na$^{+}$ = 22.99 mg meq$^{-1}$
- K$^{+}$ = 39.10 mg meq$^{-1}$
9. Report

Report the extractable Ca^{2+}, Mg^{2+}, K^{+}, and Na^{+} to the nearest 0.1 meq 100 g\(^{-1}\) (cmol (+) kg\(^{-1}\)).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Ion Exchange and Extractable Cations (4B)

BaCl\(_2\)-Triethanolamine, pH 8.2 Extraction (4B2)

Centrifuge (4B2b)

Automatic Titrator (4B2b1)

Back Titration with HCl (4B2b1a)

Extractable Acidity (4B2b1a1)

Air-Dry or Field-Moist, <2 mm (4B2b1a1-b1)

1. Application

Extractable acidity is the acidity released from the soil by a barium chloride-triethanolamine (BaCl\(_2\)-TEA) solution buffered at pH 8.2. Extractable acidity includes all the acidity generated by replacement of the H and Al from permanent exchange sites and pH-dependent exchange sites. Extractable acidity may be measured at any pH, and a variety of methods have been used to measure it. The Soil Conservation Service adopted a pH of 8.2, which approximates the calculated pH of a soil containing free CaCO\(_3\) in equilibrium with the normal CO\(_2\) content (0.03%) of the atmosphere. A pH of 8.2 also closely corresponds to the pH of complete neutralization of soil hydroxy-Al compounds. Although other pH values are valid for some types of soils and the BaCl\(_2\)-TEA, pH 8.2, method (4B2b1a1) may not always accurately reflect the nature of soils as they occur in the environment, this method has become a standard reference to which other methods are compared. Extractable acidity by BaCl\(_2\)-TEA, pH 8.2, is routinely determined by the KSSL for those samples that show very slight or no effervescence after treatment with 1 N HCl (method 1B1b2d4).

2. Summary of Method

A soil sample is leached with a BaCl\(_2\)-TEA solution buffered at pH 8.2. The sample is allowed to stand overnight, shaken, and centrifuged. The extract is back-
titrated with HCl. The difference between a blank and the extract is the extractable acidity. Extractable acidity is reported in meq 100 g$^{-1}$ soil or (cmol (+) kg$^{-1}$).

3. Interferences

No significant interferences are known to exist with this method. However, the buffer capacity of the BaCl$_2$-TEA solution may be exceeded for some very acid soils.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids in a fume hood. Thoroughly wash hands after handling reagents. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Pipettes or dispenser, adjustable volume to 40 mL
5.3 Vortexer, mini, Analog, VRW Scientific Products
5.4 Centrifuge tubes, 50-mL, polyethylene
5.5 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.6 Titration beakers, 250-mL, plastic, Metrohm Ltd., Brinkmann Instruments Inc.
5.7 Automatic titrator, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.
5.8 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.
5.9 Computer, with Titrino Workcell software, Metrohm Ltd., Brinkmann Instruments, Inc., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated, 12 N
6.3 HCl, 0.13 N, standardized. Dilute 193 mL of concentrated HCl to 16-L volume with RODI water.
6.4 Buffer solution (0.5 N BaCl$_2$, 0.2 N Triethanolamine (TEA), pH 8.2). Dissolve 977 g of BaCl$_2$$\cdot$2H$_2$O in 8 L of RODI water. Dissolve 477 g of TEA in 4 L of RODI water. Mix two solutions and bring to nearly 16-L volume
with RODI water. Adjust to pH 8.2 with ≈33 mL of concentrated HCl or barium hydroxide. Bring to 16-L volume with RODI water.

6.5 Replacement solution. Dissolve 977 g of BaCl\textsubscript{2}•2H\textsubscript{2}O in 8 L of RODI water. Add 80 mL of buffer solution and dilute to 16-L volume with RODI water.

7. Procedure

**Extraction of Acidity**

7.1 Weigh 5 g of <2-mm or fine-grind, air-dry soil to the nearest mg and place in a centrifuge tube. If sample is moist, weigh enough soil to achieve ≈5 g of air-dry soil. Prepare at least two reagent blanks (no sample in tube) and one quality control check sample per 21 samples.

7.2 Add 40.00 mL of BaCl\textsubscript{2}-TEA solution to sample. Cap the tube and shake to ensure all soil is wetted. Place tube in a rack.

7.3 Place tube rack on its side and gently shake to stratify the mixture lengthwise along the tube. Allow to stand overnight on its side.

7.4 Centrifuge sample at 2000 rpm for 5 min.

7.5 Decant extract into numbered titration beakers.

7.6 Add 40 mL of replacement solution to sample.

7.7 Cap tube and use a vortexer to loosen soil. Manually shake.

7.8 Repeat Sections 7.4–7.7.

7.9 Repeat Sections 7.4–7.5. Total volume in titration beaker should be ≈120 mL.

**Titration of BaCl\textsubscript{2}-TEA Extract**

7.10 Place titration beakers on automatic sample changer.

7.11 Refer to the manufacturer’s manual for operation of the automatic titrator.

7.12 Calibrate the titrator meter with pH 9.18, 7.00 and 4.00 buffers. Set-up the automatic titrator to “set end point” mode. The “Set” pH parameters are listed as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(E_p)</td>
<td>pH 4.60</td>
</tr>
<tr>
<td>Dyn change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s(^{-1})</td>
</tr>
<tr>
<td>Time delay</td>
<td>10 s</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s(^{-1})</td>
</tr>
<tr>
<td>Temp</td>
<td>25 °C</td>
</tr>
<tr>
<td>Stop volume</td>
<td>75 mL</td>
</tr>
</tbody>
</table>
7.13 If pre-titration pH is 0.3 units lower than the average pH of the blanks, re-run using a 0.5-g sample.

7.14 Record the titer to the nearest 0.01 mL. Record the normality of the HCl solution. Average the titer of the reagent blanks and record.

8. Calculations

\[
\text{Extractable acidity (meq 100 g}^{-1}\) = \frac{[(B-T) \times N \times R]}{C} \times 100
\]

where:
- \(B\) = Average reagent blank titer (mL)
- \(T\) = Sample titer (mL)
- \(N\) = Normality of HCl
- \(C\) = Sample Weight (g)
- 100 = Conversion factor (100-g basis)
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

9. Report

Report extractable acidity to the nearest 0.1 meq 100 g\(^{-1}\) (cmol (+) kg\(^{-1}\)).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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**Ion Exchange and Extractable Cations (4B)**

1. **1 \(N\) KCl Extraction (4B3)**
   - Automatic Extractor (4B3a)
     - Inductively Coupled Plasma Atomic Emission Spectrophotometer (4B3a1)
       - Radial Mode (4B3a1a)
         - Al and Mn (4B3a1a1-2)
         - Air-Dry or Field-Moist, <2 mm (4B3a1a1-2a-b1)

1. **Application**

   The Al extracted by 1 \(N\) KCl approximates exchangeable Al and is a measure of the “active” acidity present in soils with a 1:1 water pH <5.5. Above pH 5.5, precipitation of Al occurs during analysis. This method does not measure the acidity component of hydronium ions (H\(_3\)O\(^+\)). If Al is present in measurable
amounts, the hydronium is a minor component of the active acidity. Because the 1 $N$ KCl extractant is an unbuffered salt and usually affects the soil pH one unit or less, the extraction is determined at or near the soil pH. The KCl extractable Al is related to the immediate lime requirement and existing CEC of the soil. The “potential” acidity is better measured by the BaCl$_2$-TEA method (method 4B2b1a1) (Thomas, 1982). The use of NH$_4$Cl in place of KCl is useful where a single extractant for exchangeable bases and Al is preferred because NH$_4^+$ is as effective as K at displacing Al (Lee et al., 1985; Bertsch and Bloom, 1996). The Mn extracted by 1 $N$ KCl approximates exchangeable Mn. Mn is an essential trace metal for plant nutrition. Soil analysis for Mn is of interest from both deficiency and toxicity perspectives (Gambrell, 1996). Extractable Al is routinely determined by the KSSL when pH is <5.05 by 1:2 0.01 $M$ CaCl$_2$ (method 4C1a2a2).

2. Summary of Method

In this method (4B3a1a1-2), a soil sample is leached with 1 $N$ KCl using the mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed. The KCl extracted solution is diluted with 0.5 $N$ HCl. The analytes are measured by inductively coupled plasma atomic emission spectrophotometer (ICP–AES). The Mn and Al in the soil are reported in mg kg$^{-1}$ and cmol(+) kg$^{-1}$, respectively.

3. Interferences

Four types of interferences (matrix, spectral, chemical, and ionization) affect the ICP–AES analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

The soil to extractant ratio must remain constant. A soil to extractant ratio of 1:10 (w:v) for batch procedures is most commonly used. Using a leaching technique, a 1:20 (w:v) ratio gives comparable results. If the sample size is changed, the amount of extractable Al is changed. No other significant interferences have been identified for this procedure.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the ICP–AES.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical vacuum extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE (figs. 4B-1 and 4B-2)
5.3 Tubes, 60-mL, polypropylene, for extraction (0.45-µm filter), reservoir, and tared extraction tubes
5.4 Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels
5.5 Dispenser, 10-mL
5.6 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES), Perkin-Elmer Optima 7300 Dual View (DV), Perkin-Elmer Corp., Norwalk, CT
5.7 Scott spray chamber with end cap and gem cone x-flow nebulizer
5.8 Torch coupler at −3 position
5.9 RF generator, floor mounted power unit, 45 MHz free running, Perkin-Elmer Corp., Norwalk, CT
5.10 Computer, with WinLab software ver. 4.1, Perkin-Elmer Corp., Norwalk, CT, and printer
5.11 Recirculating chiller, Neslab, CFT Series
5.12 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.13 Single-stage regulator, high-purity, high-flow, argon
5.14 Pipettes, electronic digital, 10,000-µL and 1000-µL, with tips, 10,000-µL and 1000-µL
5.15 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.16 Containers, polyethylene
5.17 Vortexer, mini, MV1, VWR Scientific Products
5.18 Disposable tubes, glass, 10 mL

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Potassium chloride solution (KCl), 1.0 N. Dissolve 1341.9 g of KCl reagent in 16 L RODI water. Allow solution to equilibrate to room temperature. Dilute to 18 L with RODI water. Use 1.0 N KCl for Al and Mn extraction.
6.3 Potassium chloride solution (KCl), 2.0 N. Dissolve 298.2 g of KCl reagent in 1.5 L RODI water. Allow solution to equilibrate to room temperature. Dilute to 2 L with RODI water. Use 2.0 N KCl for standards.
6.5 HCl, 0.5 N. In a 1-L volumetric, add 41.67 mL concentrated HCl 12 N to RODI water and then dilute to volume with RODI water.
6.6 Primary Stock Standard Solution (PSSS), high purity, 1000 mg L⁻¹: Al and Mn
6.6.1 Mixed calibration standards solution (MCSS) for Al and Mn as follows:
6.6.1.1 MCSS High: In 1-L volumetric flask, mix 40 mL Al PSSS, 5 mL Mn PSSS, 14.91 g KCl, and 33.3 mL concentrated HCl and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 40 and 5 mg L⁻¹ Al and Mn, respectively. Store
in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use.

**6.6.2** MCSS Medium: In 1-L volumetric flask, mix 20 mL Al PSSS, 2 mL Mn PSSS, 14.91 g KCl, and 33.3 mL concentrated HCl and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 20 and 2 mg L\(^{-1}\) Al and Mn, respectively. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use.

**6.6.3** MCSS Low: In 1-L volumetric flask, mix 10 mL Al PSSS, 1 mL Mn PSSS, 14.91 g KCl and 33.3 mL concentrated HCl and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 10 and 1 mg L\(^{-1}\) Al and Mn, respectively. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use.

**6.6.4** MCSS Blank: In 1-L volumetric flask, mix 500 RODI water, 33.3 mL concentrated HCl, and 14.91 g KCl and dilute to volume with RODI water. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use.

**6.7** Argon gas, purity 99.9%

**7. Procedure**

**Extraction of Al and Mn**

**7.1** Weigh 2.5 g of <2-mm, air-dry soil to the nearest mg and place in a labeled extraction tube (ET). If sample is fine-grind, weigh 1.25 g to the nearest mg. If sample is moist, weigh enough soil to achieve ≈2.5 or 1.25 g, respectively, of air-dry soil. Prepare one quality control sample per 24 samples.

**7.2** Place labeled ET on extractor and connect to corresponding tared extraction tube (TET\(_{KCl}\)) with rubber tubing.

**7.3** Use a dispenser and add 10 mL of 1 \(N\) KCl to the ET. All soil should be wetted, and no air bubbles should be present. Shaking, swirling, or stirring may be required to wet organic samples.

**7.4** Secure reservoir tube (RT) to top of ET tube and let stand for 30 min. Extract the KCl solution at the 30-min rate until 2 mL of this solution remains above soil level. Turn off extractor. Do not let soil dry.

**7.5** Add 45 mL of KCl solution to the RT if sample weight is 2.5 g. Add 17.5 mL of KCl solution to the RT if sample weight is 1.25 g. Set extractor for 45-min extraction. Extractor turns off automatically.

**7.6** Remove RT from top of extractor. Carefully remove TET\(_{KCl}\). Leave the rubber tubing on the ET. Weigh each TET\(_{KCl}\) containing the KCl extract to the nearest mg.
7.7 Mix the extract in each TET$_{KCl}$ by manually shaking. Fill a disposable tube with extract solution and discard the excess properly. This solution is reserved for extractable Al and Mn analyses. If extracts are not be determined immediately after collection, then store samples at 4 °C.

Dilution of Extracts

7.8 Dilute samples (1:5 dilution). Dilute 1 part KCl sample extract with 4 parts 0.5 N HCl. Use Vortexer to mix sample.

7.9 Place the diluted sample solutions into test tubes and place in the sample holder of the sample changer.

ICP–AES Set-up and Operation

7.10 Refer to the manufacturer’s manual for operation of the ICP–AES. The following parameters are only very general guidelines for instrument conditions for the analytes.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>View</td>
<td>Radial</td>
</tr>
<tr>
<td>Wavelength Al</td>
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<tr>
<td>Wavelength Mn</td>
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<tr>
<td>Background correction</td>
<td>“ON” 2-Point</td>
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<tr>
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<tr>
<td>Gas Flow</td>
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<tr>
<td>Auxiliary gas flow</td>
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<tr>
<td>Nebulizer flow</td>
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<td>Peristaltic Pump</td>
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<tr>
<td>Sample flow rate</td>
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<tr>
<td>Flush rate</td>
<td>35 s</td>
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<tr>
<td>Relaxation time</td>
<td>15 s</td>
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<tr>
<td>Pump tubing type</td>
<td>Solvflex black/black, 2-stop, 0.030 ID for sample; Solvflex red/red, 2-stop, 0.045 ID for rinse</td>
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</tbody>
</table>
ICP–AES Calibration and Analysis

7.11 Calibrate the instrument by using the MCSS. The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.12 If sample exceeds calibration standard, the sample is diluted 1:10 with 1 N KCl followed by 1:5 0.5 N HCl.

7.13 Perform one quality control (QC) (Low MCSS) every 12 samples. If reading is not within 10%, the instrument is re-calibrated and QC re-analyzed.

7.14 Record analyte readings to 0.01 mg L$^{-1}$.

8. Calculations

The instrument readings are the analyte concentration (mg L$^{-1}$ Al and Mn). Use these values to calculate the analyte concentration in meq 100 g$^{-1}$ and mg kg$^{-1}$ for Al and Mn, respectively.

8.1 $\text{Al (meq 100 g}^{-1}) = \frac{A \times \left(\frac{B_1 - B_2}{B_3}\right) \times C_1 \times C_2 \times R \times 100}{1000 \times E \times F}$

where:
- $A$ = Al concentration in extract (mg L$^{-1}$)
- $B_1$ = Weight of extraction syringe and extract (g)
- $B_2$ = Weight of tared extraction syringe (g)
- $B_3$ = Density of 1 N KCl at 20 °C (1.0412 g mL$^{-1}$)
- $C_1$ = Dilution, required (1:5)
- $C_2$ = Dilution, if performed
- 100 = Conversion factor (100-g basis)
- $R$ = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = mL L$^{-1}$
- $E$ = Soil sample weight (g)
- $F$ = Equivalent weight ($\text{Al}^{3+} = 8.99$ mg meq$^{-1}$)

8.2 $\text{Mn (mg kg}^{-1}) = \frac{A \times \left(\frac{B_1 - B_2}{B_3}\right) \times C_1 \times C_2 \times R \times 1000}{1000 \times E}$

where:
- $A$ = Mn concentration in extract (mg L$^{-1}$)
- $B_1$ = Weight of extraction syringe and extract (g)
- $B_2$ = Weight of tared extraction syringe (g)
- $B_3$ = Density of 1 N KCl at 20 °C (1.0412 g mL$^{-1}$)
- $C_1$ = Dilution ratio, required (1:5)
- $C_2$ = Dilution ratio, if needed
R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
1000 = Conversion factor in numerator to kg-basis
1000 = Factor in denominator (mL L$^{-1}$)
E = Soil sample weight (g)

9. Report
   Report KCl extractable Al to the nearest 0.1 meq 100 g$^{-1}$ (cmol(+) kg$^{-1}$) and Mn to the nearest mg kg$^{-1}$.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

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Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Refer to Soil Survey Staff (2014) for the use and application of these ratios and estimates in U.S. Soil Taxonomy.
Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Sum of Extractable Bases (4B4a)
Sum of Extractable Bases by NH₄OAc, pH 7 (4B4a1)
Sum of Extractable Bases by NH₄OAc, pH 7, Calculated (4B4a1a)

Sum the NH₄OAc, pH 7, extractable bases (Ca²⁺, Mg²⁺, K⁺, and Na⁺) (4B4a1a) obtained by method 4B1a1 and analyzed in methods 4B1a1b1-4, respectively. This value is reported as meq 100 g⁻¹ (cmol (+) kg⁻¹).

Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Sum of Extractable Bases (4B4a)
Sum of Extractable Bases by NH₄Cl (4B4a2)
Sum of Extractable Bases by NH₄Cl, Calculated (4B4a2a)

Sum the NH₄Cl extractable bases (Ca²⁺, Mg²⁺, K⁺, and Na⁺) (4B4a2a) obtained by method 4B1b1 and analyzed in methods 4B1b1b1-4, respectively. This value is reported as meq 100 g⁻¹ (cmol (+) kg⁻¹).

Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Cation Exchange Capacity (CEC) (4B4b)
CEC–8.2 (Sum of Cations) (4B4b1)
CEC–8.2, Reported (4B4b1a)
CEC–8.2, Not Reported (4B4b1b)

Calculate the CEC–8.2 (4B4b1a) by adding the sum of the NH₄OAc extractable bases (4B4a1) plus the BaCl₂-TEA extractable acidity (method 4B2a1a1 or 4B2b1a1). This value is reported as meq 100 g⁻¹ (cmol (+) kg⁻¹). Cation summation is the basis for this procedure. The CEC–8.2 minus the CEC–7 is considered the pH dependent charge from pH 7.0 to pH 8.2. The CEC–8.2 is not reported (method 4B4b1b) if carbonates, gypsum, or soluble salts are present in the soil because the NH₄OAc extracts cations from the dissolution of these soil constituents. CEC–8.2 is calculated as follows:

\[ \text{CEC–8.2} = \text{NH}_4\text{OAc Bases} + \text{BaCl}_2\text{-TEA Acidity} \]
Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Cation Exchange Capacity (CEC) (4B4b)
Effective Cation Exchange Capacity (ECEC) (4B4b2)
Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Reported (4B4b2a)
Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Not Reported (4B4b2b)

Calculate the ECEC (method 4B4b2a) by adding the sum of the NH$_4$OAc extractable bases (4B4a1) plus the 1 N KCl extractable Al (method 4B3a1a1). This value is reported as meq 100 g$^{-1}$ (cmol (+) kg$^{-1}$). The ECEC is not reported (method 4B4b2b) if carbonates, gypsum, or significant quantities of soluble salts are present in the soil because the NH$_4$OAc extracts cations from the dissolution of these soil constituents. ECEC by NH$_4$OAc extractable bases and 1 N KCl Al is calculated as follows:

$$\text{ECEC} = \text{NH}_4\text{OAc Bases} + 1 \text{ N KCl Al}$$

References
Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Base Saturation (4B4c)

Base Saturation by NH$_4$Cl (4B4c2)

Base Saturation by NH$_4$Cl, Reported (4B4c2a)

Base Saturation by NH$_4$Cl, Set to 100% (4B4c2b)

Calculate the base saturation (method 4B4c2a) by dividing the sum of the NH$_4$Cl extractable (4B4a2) bases by CEC by NH$_4$Cl (method 4B1b1a1a1) and multiplying by 100. This value is reported as meq 100 g$^{-1}$ or cmol (+) kg$^{-1}$. If a soil has carbonates, gypsum, or soluble salts, this value is set to 100% (method 4B4c2b). Calculate base saturation by NH$_4$Cl as follows:

Base Saturation (%) = (NH$_4$Cl Bases/CEC by NH$_4$Cl) x 100

Base Saturation by CEC–8.2 (Sum of Cations) (4B4c3)

Base Saturation by CEC–8.2, Reported (4B4c3a)

Base Saturation by CEC–8.2, Set to 100% (4B4c3b)

Calculate the base saturation (method 4B4c3a) by dividing the sum of the NH$_4$OAc extractable bases (4B4a1) by CEC–8.2 (method 4B4b1a1a1) and multiplying by 100. This value is reported as meq 100 g$^{-1}$ or cmol (+) kg$^{-1}$. If a soil has carbonates, gypsum, or soluble salts, this value is set to 100% (method 4B4c3b). Calculate base saturation by CEC–8.2 (Sum of Cations) as follows:

Base Saturation (%) = [NH$_4$OAc Bases/(NH$_4$OAc Bases + BaCl$_2$-TEA Acidity)] x 100

References
Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Base Saturation (4B4c)

Base Saturation by Effective Cation Exchange Capacity (ECEC) (4B4c4)

Base Saturation by Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Reported (4B4c4a)
Base Saturation by Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Not Reported (4B4c4b)

Calculate the base saturation (method 4B4c4a) by dividing the sum of NH$_4$OAc extractable bases (method 4B4a1) by the ECEC (method 4B4b2a) and multiplying by 100. If a soil has carbonates, gypsum, or significant quantities of soluble salts, this value is not reported (4B4c4b). Calculate base saturation by ECEC as follows:

$$\text{Base Saturation (\%) = } \left[ \frac{\text{NH}_4\text{OAc Bases}}{\text{NH}_4\text{OAc Bases} + 1 \ N \ KCl \ Al} \right] \times 100$$

Ion Exchange and Extractable Cations (4B)
Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Aluminum Saturation (4B4d)

Aluminum Saturation by Effective Cation Exchange Capacity (ECEC) (4B4d1)

Aluminum Saturation by Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Reported (4B4d1a)
Aluminum Saturation by Sum of NH$_4$OAc Extractable Bases + 1 N KCl Extractable Aluminum, Not Reported (4B4d1b)

Calculate the Al saturation (method 4B4d1a) by dividing the 1 N KCl extractable Al (method 4B3a1a1) by the ECEC (method 4B4b2a) and multiplying by 100. If a soil has carbonates, gypsum, or significant quantities of soluble salts, this value is not reported (method 4B4d1b). Calculate Al saturation as follows:

$$\text{Al Saturation (\%) = } \left[ \frac{1 \ N \ KCl \ Al}{\text{NH}_4\text{OAc Bases} + 1 \ N \ KCl \ Al} \right] \times 100$$
Ion Exchange and Extractable Cations (4B)

Ratios and Estimates Related to Ion Exchange and Extractable Cations (4B4)

Activity (4B4e)

CEC–7/Clay (4B4e1)

Divide the CEC–7 (method 4B1a1a1a1) by the total clay (method 3A1a). This ratio is reported as a dimensionless value. In the past, the ratios of CEC to clay have been reported as meq g\(^{-1}\). For more detailed information on the application of this ratio, refer to Soil Survey Staff (2011, 2014).

References


Hydrogen-Ion Activity (4C)

Soil Suspensions (4C1)

Electrode (4C1a)

Standard Glass Body Combination (4C1a1)

Digital pH/Ion Meter (4C1a1a)

Soil pH is one of the most frequently performed determinations and one of the most indicative measurements of soil chemical properties (McLean, 1982). Soil pH tells more about a soil than merely indicating whether it is acidic or basic. The availability of essential nutrients and toxicity of other elements can be estimated because of their known relationship with pH (Thomas, 1996). Soil pH is affected by many factors, e.g., nature and type of inorganic and organic matter; amount and type of exchangeable cations and anions; soil to solution ratio; salt or electrolyte content; and CO\(_2\) content (McLean, 1982). The acidity, neutrality, or basicity of a soil influences the solubility of various compounds; the relative ion bonding to exchange sites; and microbial activities. Depending on the predominant clay type, the pH may be used as a relative indicator of base saturation (Mehlich, 1943). Soil pH is also a critical factor affecting the availability of most essential elements for plants.

The KSSL performs several pH determinations. These methods include, but are not limited to, NaF (1 N pH 7.5 to 7.8) (4C1a1a1); saturated paste pH (4C1a1a2); oxidized pH (4C1a1a3); 1:1 water and 1:2 CaCl\(_2\) (final solution: 0.01
An increase in the soil to water ratio or the presence of salts generally results in a decrease in the soil pH. The soluble salt content of the soil can be overcome by using dilute salt solutions, e.g., CaCl₂ or KCl, instead of distilled water. The use of dilute salt solutions is a popular method for masking seasonal variation in soil pH. The pH readings are usually less with dilute salt solutions than with distilled water but may be equal to or greater in highly weathered tropical soils, i.e., soils with a high anion exchange capacity. When the pH values of various soils are compared, determination by the same method is important (Foth and Ellis, 1988).

The pH as determined by the 1 N KCl method is an index of soil acidity. This method is more popular in those regions that have extremely acid soils and in which KCl is used as an extractant of exchangeable Al. The KCl pH indicates the pH at which Al is extracted. Similarly to the 1:2 CaCl₂ pH, the 1 N KCl pH readings tend to be uniform regardless of time of year.

The saturated paste pH method is popular in regions that have soils with soluble salts. The water content varies with the water storage characteristics of the soil. The saturated paste pH may be more indicative of the saturated, irrigated soil pH than is the soil pH measurement at a constant soil to water ratio. The saturated paste pH method is also the method from which the saturation extract is removed for salt analysis. The saturated paste pH is therefore the pH and the dilution at which the sodium adsorption ratio (SAR) is computed (method 4E4b).

The 1 N NaF pH may be used as an indicator that amorphous material dominates the soil exchange complex. The oxidized pH may be used to assess the activities of soil microorganisms. In soil taxonomy, the CaCl₂ pH is used to distinguish two family reaction classes in Histosols (Soil Survey Staff, 2014).

References


Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
   Electrode (4C1a)
   Standard Glass Body Combination (4C1a1)
   Digital pH/Ion Meter (4C1a1a)
   1 N NaF, pH 7.5–7.8 (4C1a1a1)
   Air-Dry or Field-Moist, <2 mm (4C1a1a1a-b1)

1. Application
The action of NaF upon noncrystalline (amorphous) soil material releases hydroxide ions (OH\(^{-}\)) to the soil solution and increases the pH of the solution. The amount of amorphous material in the soil controls the release of OH\(^{-}\) and the subsequent increase in pH (Fields and Perrott, 1966). The following reactions illustrate this action and form the basis of this procedure.

\[
\text{Al(OH)}_3 + 3 \text{F}^- \rightarrow \text{AlF}_3 + 3 \text{OH}^- \\
\text{Si(OH)}_4 + 4 \text{F}^- \rightarrow \text{SiF}_4 + 4 \text{OH}^- 
\]

Most soils contain components that react with NaF and release OH\(^{-}\). However, a NaF pH ≥9.4 is a strong indicator that amorphous material dominates the soil exchange complex. Amorphous material is usually an early product of weathering of pyroclastic materials in a humid climate. Amorphous material appears to form in spodic horizons in the absence of pyroclastics.

2. Summary of Method
A 1-g sample is mixed with 50 mL of 1 N NaF and stirred for 2 min. While the sample is being stirred, the pH is read at exactly 2 min in the upper ⅓ of the suspension (4C1a1a1).

3. Interferences
The difference in the sediment and supernatant pH is called the suspension effect (McLean, 1982). To maintain uniformity in pH determination, measure the pH just above the soil sediment. Clays may clog the KCl junction and slow the electrode response. Clean the electrode by rinsing with distilled water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization. Use high purity (99%) NaF.

Soils with a 1:1 water pH >8.2 do not give a reliable NaF pH. Free carbonates in a soil result in a high NaF pH. In general, soils with a 1:1 water pH <7.0 are not affected.
4. Safety

The NaF is poisonous. Avoid eye contact and ingestion. Skin penetration and irritation are moderately hazardous. Do not eat or drink while using NaF. Thoroughly wash hands after use. Wear protective clothing (coats, aprons, and gloves) and eye protection (safety glasses or goggles) when using NaF. Use the fume hood when using NaF. Follow standard laboratory safety practices.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Paper cup, 120-mL (4 fl. oz.), disposable, Solo Cup Co., no. 404
5.3 Titration beakers, polyethylene, 250 mL
5.4 Automatic titrator, Metrohm Titroprocessors, Control Units, Sample Changers, and Dosimats, Metrohm Ltd., Brinkmann Instruments, Inc.
5.5 Combination pH-reference electrode, Metrohm, Brinkmann Instruments, Inc.

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH buffers, pH 4.00, pH 7.00, pH 9.18, and pH 13.0 for electrode calibration, Beckman, Fullerton, CA
6.3 Phenolphthalein
6.4 Sodium fluoride (NaF), 99% purity, EM Science
6.5 Sodium fluoride (NaF), 1.0 N solution. In a plastic bottle, add 400 g NaF in 8 L of distilled water. Let stand for 3 days. On the third day, after excess NaF has settled, measure 50 mL of the solution and read pH. The pH should be between 7.5 and 7.8. Add 3 to 5 drops 0.25% phenolphthalein and titrate to pink end point (pH 8.2 to 8.3). If pH is outside the 7.5 and 7.8 range, then adjust pH with either HF or NaOH. If solution has a pH >8.2 or if the titratable acidity is >0.25 meq L⁻¹, use another source of NaF.

7. Procedure

7.1 Weigh 1 g of <2-mm or fine-grind, air-dry soil to the nearest 1 mg and place in a 120-mL (4-oz) paper cup. If sample is moist, weigh enough soil to achieve ≈1 g of air-dry soil.
7.2 Calibrate the titrator with pH 4.00, 7.00, and 9.18 buffer solutions.
7.3 The stirring of the sample, intervals for readings, addition of NaF solution, pH readings, and rinsing of electrode are controlled by computer.
7.4 The general sequence used by the automated system is as follows:

7.4.1 The sample is lifted so that the pH electrode is positioned above the soil sediment. Stirring begins immediately and is maintained during each sample cycle.
7.4.2 A 50-mL solution is added to sample.
7.4.3 After 2 min, NaF pH is read and recorded to the nearest 0.01 unit.
7.4.4 The sample is lowered, and the electrode and stirrer are rinsed with RO water.
7.4.5 The next sample is positioned for analysis.
7.4.6 The cycle is repeated until all samples have been analyzed.

7.5 Discard the solution and cup in safe containers. The paper cup with the NaF solution leaks in about 15 min.

8. Calculations
No calculations are required for this procedure.

9. Report
Report NaF pH to the nearest 0.1 pH unit.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
Electrode (4C1a)
   Standard Glass Body Combination (4C1a1)
      Digital pH/Ion Meter (4C1a1a)
         Saturated Paste pH (4C1a1a2)
            Air-Dry, <2 mm (4C1a1a2a1)

1. Application
When making interpretations about the soil, the saturated paste pH is usually compared to the 1:1 water pH and the 1:2 CaCl₂ pH. The usual pH sequence is as follows: 1:1 water pH > 1:2 CaCl₂ pH > saturated paste pH. If saturated paste
pH is > 1:2 CaCl₂ pH, the soil is not saline. If the saturated paste pH ≥ 1:1 water pH, the soil may be Na saturated and does not have free carbonates.

Because of the interrelations that exist among the various soil chemical determinations, the saturated paste pH value may be used as a means of cross-checking salinity data for internal consistency and reliability (U.S. Salinity Laboratory Staff, 1954). Some rules of thumb that apply to the saturated paste pH are as follows:

a. Soluble carbonates are present only if the pH is >9.
b. Soluble bicarbonate are seldom >3 or 4 meq L⁻¹, if the pH is ≤7.
c. Soluble Ca²⁺ and Mg²⁺ are seldom >2 meq L⁻¹, if the pH is >9.
d. Gypsiferous/gypseous soils seldom have a pH >8.2.

2. Summary of Method

The saturated paste is prepared (4F2), and the pH of paste is measured with a calibrated combination electrode/digital pH meter (4C1a1a2).

3. Interferences

The difference in the sediment and supernatant pH is called the suspension effect (McLean, 1982). To maintain uniformity in pH determination, measure the pH just beneath the surface of saturated paste. Clays may clog the KCl junction and slow the electrode response. Clean the electrode by rinsing with RO water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

4. Safety

No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Digital pH/ion meter, Accumet Model AR15, Fisher Scientific
5.2 Electrode, standard glass body combination, Accuflow, Fisher Scientific

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH Buffers, pH 4.00, pH 7.00 and pH 9.18, for electrode calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Prepare a saturated paste (method 4F2).
7.2 Calibrate the pH meter with pH 4.00, 7.00, and 9.18 buffer solutions.
7.3 After equipment calibration, gently wash the electrode with RO water. Dry the electrode. Do not wipe the electrode with a tissue as this may cause a static charge on the electrode.

7.4 Gently lower the electrode in the saturated paste until the KCl junction of the electrode is beneath the surface of saturated paste.

7.5 Allow the pH meter to stabilize before recording the pH. Record pH to the nearest 0.01 unit.

7.6 Gently raise the pH electrode from the paste and wash all particles adhering to the electrode with a stream of RO water.

8. Calculations
   No calculations are required for this procedure.

9. Report
   Report saturated paste pH to the nearest 0.1 pH unit.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
   Electrode (4C1a)
      Standard Glass Body Combination (4C1a1)
         Digital pH/Ion Meter (4C1a1a)
            Oxidized pH (4C1a1a3)

1. Application
   Sulfide material is waterlogged mineral, organic, or mixed soil material with a pH of 3.5 or higher, containing oxidizable sulfur compounds, and which if incubated as a 1-cm thick layer under moist, aerobic conditions (field capacity) at room temperature, shows a drop in pH of 0.5 or more units to a pH value of 4.0
or less (1:1 by weight in water or in a minimum of water to permit measurement) within 8 weeks (Van Breemen, 1982; Soil Survey Staff, 2014). The intent of the method described herein is to determine if known or suspected sulfidic materials will oxidize to form a sulfuric horizon (Soil Survey Staff, 2014). Identification of $\text{H}_2\text{S}$ in a soil by a “rotten-egg” smell or FeS in a saturated soil by its blue-black color indicates that sulfidic materials may be present. If such soils are drained and oxidized, the soil pH could drop to 3.5 or less, making the soil unsuitable for many uses. A field test for FeS is to add 1 N HCl and note the odor of $\text{H}_2\text{S}$.

2. Summary of Method

Transfer enough soil to fill a plastic cup one-half to two-thirds full. Add a little water if needed to make a slurry. Stir the slurry thoroughly to introduce air. Determine pH immediately. Place cup in a closed container with openings (inlet and outlet) providing humidified airflow. Keep at room temperature. After 24 h, open the container, stir the sample thoroughly, and determine the soil pH. Repeat the procedure for a minimum of 10 days until the pH reaches a steady state of $\leq 0.1$ units over a 2-day period. Record daily pH readings (4C1a1a3).

3. Interferences

Samples should be shipped in watertight containers completely filled with water from the ambient soil solution to prevent potential oxidation of sulfides and reduction in soil pH.

Extended time in stirring of sample and/or in reading the pH may result in the introduction of sufficient $\text{O}_2$ into the mixture to change the pH reading. This error can be minimized by quickly stirring the mixture and reading the pH.

Clean the electrode by rinsing with reverse osmosis (RO) water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

4. Safety

No significant hazard has been identified with this procedure. Follow standard laboratory safety precautions.

5. Equipment

5.1 Cups, plastic

5.2 Closed container, with openings (inlet and outlet) providing for humidified airflow. A tube from the inlet of this closed container is connected to the outlet of a stoppered 2.5-L container full of RO water. A pump supplies air to a bubbling stone placed near the bottom of the 2.5-L container (fig. 4C-1).

5.3 Digital pH/ion meter, Accumet Model AR15, Fisher Scientific
5.4 Electrode, standard glass body combination, Accuflow, Fisher Scientific

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH buffers, pH 4.00 and 7.00 for pH meter calibration, Beckman, Fullerton, CA

7. Procedure
7.1 Calibrate the pH meter with pH 4.00 and pH 7.00 buffer solutions.
7.2 After equipment calibration, gently wash the electrode with RO water. Dry the electrode. Do not wipe the electrode with a tissue as this may cause a static charge on the electrode.
7.3 Transfer enough soil to fill a small plastic cup one-half to two-thirds full. Add a little water if needed to make a slurry. Stir the slurry thoroughly to introduce air.
7.4 Immediately determine the pH of sample by carefully placing the electrode into the soil mixture. Ensure that the KCl junction and sensor membrane are in contact with the mixture.
7.5 Allow the pH meter to stabilize before recording the pH. Record the pH to the nearest 0.01 pH unit.
7.6 After pH determination, immediately place sample in a closed container with openings (inlet and outlet) providing for humidified airflow. A tube from the inlet of this closed container is connected to the outlet of a stoppered 2.5-L container full of RO water. A pump supplies air to a bubbling stone placed near the bottom of the 2.5-L container. Keep at room temperature (20 to 25 °C).
7.7 Stir the sample and record pH daily. Note any bubbling. After pH determination, immediately place sample back in closed container.

7.8 Record the pH for a minimum of 10 days until the change is ≤0.1 pH units for two days.

8. Calculations
No calculations are required for this procedure.

9. Report
Report the initial pH and the oxidized pH (end pH) to the nearest 0.1 pH unit.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
   Electrode (4C1a)
       Standard Glass Body Combination (4C1a1)
       Digital pH/Ion Meter (4C1a1a)
       Organic Materials CaCl₂ pH, Final Solution ≈0.01 M CaCl₂ (4C1a1a4)

1. Application
This method for determining pH is used in soil taxonomy to distinguish two family reaction classes in Histosols (Soil Survey Staff, 2014). Dysic families have a pH <4.5 in 0.01 M CaCl₂ in all parts of the organic materials in the control section. Euic families have a pH >4.5 in 0.01 M CaCl₂ in some part of the control section.

2. Summary of Method
Place 2.5 mL (2.5 cm³) of the prepared sample in a 30-mL plastic container and add 4 mL of 0.015 M CaCl₂, yielding a final concentration of ≈0.01 M CaCl₂ with most packed, moist organic materials. Mix, cover, and allow to equilibrate at least 1 h. Uncover and measure pH with pH paper or pH meter (4C1a1a4).
3. Interferences

This test of organic soil material can be used in field offices. Because it is not practical in the field to base a determination on a dry sample weight, moist soil is used. The specific volume of moist material depends on how it is packed. The packing of the material, therefore, must be standardized in order to obtain comparable results by different soil scientists (Soil Survey Staff, 2014).

Clean the pH electrode by rinsing with reverse osmosis (RO) water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

4. Safety

No significant hazard has been identified with this procedure. Follow standard laboratory safety precautions.

5. Equipment

5.1 Polycons, 30-mL
5.2 Digital pH/ion meter, Accumet Model AR15, Fisher Scientific
5.3 Electrode, standard glass body combination, Accuflow, Fisher Scientific
5.4 Half-syringe, 6-mL. Cut plastic syringe longitudinally to form a half-cylinder measuring device.
5.5 Metal spatula

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Calcium chloride (CaCl₂), 0.015 M. Dissolve 1.10 g of CaCl₂•2H₂O in RO water and dilute to 500 mL.
6.3 Borax pH buffers, pH 4.00 and 7.00, for pH meter calibration, Beckman, Fullerton, CA

7. Procedure

Sample Preparation

7.1 Prepare soil material. If the soil is dry, add water and let stand to saturate. Place 50 to 60 mL of a representative sample on a paper towel in a linear mound. Roll the towel around the sample and express water if necessary. Use additional paper towels as external blotters. Remove the sample and place on a fresh paper towel. The sample should be firm but saturated with water.

7.2 Use scissors to cut sample into 0.5- to 1.0-cm long segments.

7.3 Randomly select sample segments for determination of fiber (5C), solubility in pyrophosphate (5B), and pH (4C1a1a4).
pH Determination

7.4 Use a metal spatula to pack a half-syringe that is adjusted to the 5-mL mark or 2.5-mL (2.5-cm³) volume with the moist sample.

7.5 Place 2.5 mL (2.5 cm³) of the prepared sample in a 30-mL polycon and add 4 mL of 0.015 M CaCl₂, yielding a final concentration of approximately 0.01 M CaCl₂ with most packed moist organic materials.

7.6 Mix, cover, and allow to equilibrate at least 1 h.

7.7 Uncover, mix again, immerse electrode, and measure pH. Rinse electrode with RO water.

7.8 Alternatively, place pH strip on top of sample so that it wets from the bottom. Close cover and allow to equilibrate approximately 5 min. Remove pH strip with tweezers. Use a wash bottle to gently wash soil from bottom of strip. Compare color of active segment (center) with reference segments and with pH scale on box to determine pH.

8. Calculations
No calculations are required for this procedure.

9. Report
Report the 0.01 M CaCl₂ pH to the nearest 0.1 pH unit.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
Electrode (4C1a)

Combination pH-Reference Electrode (4C1a2)
Automatic Titrator (4C1a2a)

1:1 Water pH (4C1a2a1)
Air-Dry or Field-Moist, <2 mm (4C1a2a1a-b1)

1:2 0.01 M CaCl₂ pH (4C1a2a2)
Air-Dry or Field Moist, <2 mm (4C1a2a2a-b1)

1. Application
The 1:1 water pH (4C1a2a1) and 1:2 0.01 M CaCl₂ pH (4C1a2a2) determinations are two commonly performed soil pH measurements. The CaCl₂
soil pH generally reports lower values than the 1:1 water pH. The combination of exchange and hydrolysis in salt solutions (0.1 to 1 M) can lower the measured pH by 0.5 to 1.5 units compared to the pH measured in RO water (Foth and Ellis, 1988). These pH values are used as criteria for reaction classes (acid and nonacid) in some taxonomic families (Soil Survey Staff, 2014).

2. Summary of Method

The pH is measured in soil-water (1:1) and soil-salt (1:2 CaCl₂) solutions. For convenience, the pH is initially measured in water and then measured in CaCl₂. With the addition of an equal volume of 0.02 M CaCl₂ to the soil suspension that was prepared for the water pH, the final soil-solution ratio is 1:2 0.01 M CaCl₂.

A 20-g soil sample is mixed with 20 mL of reverse osmosis (RO) water (1:1 w:v) with occasional stirring. The sample is allowed to stand 1 h with occasional stirring. The sample is stirred for 30 s, and the 1:1 water pH is measured. The 0.02 M CaCl₂ (20 mL) is added to soil suspension, the sample is stirred, and the 1:2 0.01 M CaCl₂ pH is measured (4C1a2a2).

3. Interferences

The pH varies between the supernatant and soil sediment (McLean, 1982). Measure the pH just above the soil sediment to maintain uniformity. Clays may clog the KCl junction and slow the electrode response. Clean the electrode. Wiping the electrode dry with cloth, laboratory tissue, or similar material may cause electrode polarization. Rinse the electrode with distilled water and pat dry.

Atmospheric CO₂ affects the pH of the soil:water mixture. Closed containers and nonporous materials will not allow equilibration with CO₂. If critical work is being done, the partial pressure of CO₂ and the equilibrium point must be considered at the time of pH determination.

4. Safety

No significant hazards are associated with the procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Measuring scoop, handmade, ≈20-g capability
5.2 Paper cup, 120-mL (4 fl. oz.), disposable, Solo Cup Co., No. 404
5.3 Dispenser, 0 to 30 mL, Repipet or equivalent
5.4 Beverage stirring sticks, wood
5.5 Titration beakers, polyethylene, 250 mL
5.6 Automatic titrator, Metrohm Titroprocessors, Control Units, Sample Changers, and Dosimats, Metrohm Ltd., Brinkmann Instruments, Inc.
5.7 Combination pH-reference electrode, Metrohm part no. 6.0210.100, Brinkmann Instruments, Inc.

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH buffers, pH 4.00, 7.00, and 9.18, for electrode calibration, Beckman, Fullerton, CA
6.3 Calcium chloride (CaCl$_2$), 0.02 M. Dissolve 23.52 g of CaCl$_2$•2H$_2$O in RO water and dilute to 8 L.

7. Procedure
7.1 Use a calibrated scoop to measure ≈20 g of <2-mm or fine-grind, air-dry soil. If sample is moist, use calibrated scoop to achieve ≈20 g of air-dry soil.
7.2 Place the sample in a 120-mL (4-oz) paper cup.
7.3 Dispense 20 mL of RO water into sample and stir.
7.4 Place paper cup with sample in 250-mL titration beaker, allow to stand for 1 h, stirring occasionally.
7.5 Load beakers into sample changer.
7.6 Calibrate the pH meter using the pH 9.18, 7.00, and 4.00 buffer solutions.
7.7 The stirring of the sample, intervals for readings, addition of CaCl$_2$ solution, pH readings, and rinsing of electrode are controlled by computer.
7.8 The general sequence used by the automated system is as follows:
   7.8.1 The sample is lifted so that the pH electrode is positioned above the soil sediment.
   7.8.2 The sample is stirred for 30 s.
   7.8.3 After 1 min, 1:1 water pH is read and recorded to the nearest 0.01 unit.
   7.8.4 The 20 mL of 0.02 M CaCl$_2$ is added to sample. The sample is stirred for 30 s.
   7.8.5 After 1 min, the 1:2 CaCl$_2$ pH is read and recorded to the nearest 0.01 unit.
   7.8.6 The sample is lowered, and the electrode and stirrer are rinsed with RO water.
   7.8.7 The next sample is positioned for analysis.
   7.8.8 The cycle is repeated until all samples have been analyzed.

8. Calculations
   No calculations are required for this procedure.
9. Report
   Report the 1:1 water pH and the 1:2 0.01 M CaCl₂ pH to the nearest 0.1 pH unit.

10. Precision and Accuracy
   Precision and accuracy data are available from the KSSL upon request.

11. References
   Chemical and microbiological properties. 2nd ed. Agron. Monogr. 9. ASA
   and SSSA, Madison, WI.

Hydrogen-Ion Activity (4C)
Soil Suspensions (4C1)
   Electrode (4C1a)
      Combination pH-Reference Electrode (4C1a2)
      Automatic Titrator (4C1a2a)
      1 N KCl pH (4C1a2a3)
      Air-Dry or Field-Moist, <2 mm (4C1a2a3-b1)

1. Application
   The pH as determined by the 1 N KCl method is an index of soil acidity. If KCl
   pH is <5, significant amounts of Al are expected in the solution. If the pH is very
   much below 5, almost all the acidity is in the form of Al.

2. Summary of Method
   A 20-g soil sample is mixed with 20 mL of 1 N KCl. The sample is allowed to
   stand for 1 h with occasional stirring. The sample is stirred for 30 s, and after 1
   min, the KCl pH is read (4C1a2a3).

3. Interferences
   The difference in the sediment and supernatant pH is called the suspension
   effect (McLean, 1982). To maintain uniformity in determination, measure the
   pH just above the soil sediment. Clays may clog the KCl junction and slow
   the electrode response. Clean the electrode by rinsing with reverse osmosis
   (RO) water and patting it dry with tissue. Wiping the electrode dry with a cloth,
   laboratory tissue, or similar material may cause electrode polarization.
4. Safety

No significant hazards are associated with the procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Measuring scoop, handmade, ≈20 g
5.2 Paper cup, 120-mL (4 fl. oz.), disposable, Solo Cup Co., No. 404
5.3 Dispenser, 0 to 20 mL, Repipet or equivalent
5.4 Beverage stirring sticks, wood, FSN 7340-00-753-5565
5.5 Titration beakers, polyethylene, 250 mL
5.6 Automatic titrator, Metrohm Titroprocessors, Control Units, Sample Changers, and Dosimats, Metrohm Ltd., Brinkmann Instruments, Inc.
5.7 Combination pH-reference electrode, Brinkmann Instruments, Inc.

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH buffers, pH 4.00, 7.00, and 9.18 for electrode calibration, Beckman, Fullerton, CA
6.3 Potassium chloride (KCl), 1.0 N. Dissolve 74.56 g of KCl in RO water. Dilute to 1 L.

7. Procedure

7.1 Use a calibrated scoop to measure ≈20 g of <2-mm or fine-grind, air-dry soil. If sample is moist, use calibrated scoop to achieve ≈20 g of air-dry soil.
7.2 Place the sample in a 120-mL (4-oz) paper cup.
7.3 Dispense 20 mL of 1 N KCl into sample and stir with wooden beverage stirrer.
7.4 Place paper cup with sample in 250-mL titration beaker, allow to stand 1 h, stirring occasionally.
7.5 Load beakers into sample changer table.
7.6 Calibrate the pH meter using the pH 4.00, 7.00, and 9.18 buffer solutions.
7.7 The stirring of the sample, intervals for readings, pH reading, and rinsing of electrode are controlled by computer.
7.8 The general sequence used by the automated system is as follows:
   7.8.1 The sample is lifted so that pH electrode is positioned above the soil sediment.
   7.8.2 The sample is stirred for 30 s.
7.8.3 After 1 min, the KCl pH is read and recorded to the nearest 0.01 unit.
7.8.4 The sample is lowered, and the electrode and stirrer are rinsed with RO water.
7.8.5 The next sample is positioned for analysis.
7.8.6 The cycle is repeated until all samples have been analyzed.

8. Calculations
No calculations are required for this procedure.

9. Report
Report KCl pH to the nearest 0.1 pH unit.

10. Precision and Accuracy
Precision and accuracy data are available upon request from the KSSL.

11. References

Hydrogen-Ion Activity (4C)
Soil Extracts (4C2)
   Electrode (4C2a)
      Standard Glass Body Combination (4C2a1)
         Digital pH/Ion Meter (4C2a1a)
            Aqueous Extraction (4C2a1a1)
               Single-Point Extraction (4C2a1a1a)
                  1:5, 23-h, 1-h (4C2a1a1a1a)
                     Air-Dry or Field-Moist, <2 mm (4C2a1a1a1a-b1)

1. Application
   Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties (e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation
between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method

The extract is prepared (4D2a2), and the pH is measured with a calibrated, combination electrode/digital pH meter (4C2a1a1a1).

3. Interferences

The difference between the sediment pH and the supernatant pH is called the suspension effect (McLean, 1982). To maintain uniformity in pH determination, measure the pH just beneath the surface of saturated paste. Clays may clog the KCl junction and slow the electrode response. Clean the electrode by rinsing with RO water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material can cause electrode polarization.

4. Safety

No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Digital pH/ion meter, Accumet Model AR15, Fisher Scientific
5.2 Electrode, standard glass body combination, Accuflow, Fisher Scientific

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH Buffers, pH 4.00, 7.00, and 9.18, for electrode calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Prepare sample extract (method 4D2b1).
7.2 Calibrate the pH meter with pH 4.00, 7.00, and 9.18 buffer solutions.
7.3 After equipment calibration, gently wash the electrode with RO water. Dry the electrode. Do not wipe the electrode with a tissue as this may cause a static charge on the electrode.
7.4 Gently lower the electrode into extract until the KCl junction of the electrode is beneath the surface.
7.5 Allow the pH meter to stabilize before recording the pH. Record pH to the nearest 0.01 unit.
7.6 Gently raise the pH electrode from the paste and wash all particles adhering to the electrode with a stream of RO water.
8. Calculations

No calculations are required for this procedure.

9. Report

Report pH to the nearest 0.1 pH unit.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Test Analyses (4D)

For more than 30 years, soil testing has been widely used as a basis for determining lime and fertilizer needs (Soil and Plant Analysis Council, 1999). In more recent years, some of these tests have been employed in more diverse agronomic and environmental uses (SERA–IEG 17, 2000). It is for these reasons that the KSSL has expanded its suite of soil analyses to more completely characterize the inorganic and organic N fractions and to provide P analyses for a broad spectrum of soil applications.

Methods development in soil P characterization (Bray and Kurtz, 1945; Olsen et al., 1954; Chang and Jackson, 1957) has been instrumental in understanding the principles, nature, and behavior of P in soils (Olsen and Sommers, 1982). Amounts, forms, and distribution of soil P vary with soil-forming factors (Walker, 1974; Stewart and Tiessen, 1987); level and kind of added P (Barrow, 1974; Tisdale et al., 1985; Sharpley, 1996); other soil and land management factors (Haynes, 1982; Sharpley, 1985); and soil P-sorption characteristics (Goldberg
and Sposito, 1984; Van Riemsdijk et al., 1984; Polyzopoulos et al., 1985; Frossard et al., 1993). Knowledge of these factors and their impact on the fate and transport of soil P has been used in developing soil P interpretations for such broad and diverse applications as fertility, taxonomic classification, genesis and geomorphology models, and environmental studies (Burt et al., 2002).

To characterize the P in a soil system requires the selection of an appropriate method of determination. This selection is influenced by many factors, e.g., objectives of the study, soil properties, sample condition, sample environment, accuracy, and reproducibility (Olsen and Sommers, 1982). Most soil P determinations have two phases, i.e., preparation of a solution that contains the soil P or fraction thereof and the quantitative determination of P in the solution. Most P analyses of soil solutions have been colorimetric procedures because they are sensitive, reproducible, and lend themselves to automated analysis, accommodating water samples, digest solutions, and extracts (SERA–IEG 17, 2000). Inductively coupled plasma (ICP) spectrophotometry can also be used for P determination. The popularity of this procedure has increased due to the use of multi-element soil extractants (SERA–IEG 17, 2000). Results from colorimetric analyses are not always comparable to those from ICP because ICP estimates the total amount of P in solution while the colorimetric procedures measure P that can react with the color developing reagent (SERA–IEG 17, 2000).

The selected colorimetric method for P determination depends on the concentration of solution P, the concentration of interfering substances in the solution to be analyzed, and the particular acid system involved in the analytical procedure (Olsen and Sommers, 1982). The KSSL determines a number of P analyses. Most of these determinations are colorimetric, with the exception of the multi-element extractant Mehlich No. 3 (4D6b1b1-18). These P analyses include but are not limited to: anion-resin extractable (4D1a1a1a1-2); water soluble (4D2a1a1); Bray P-1 (4D3a1); Olsen sodium-bicarbonate (4D5a1); Mehlich No. 3 (4D6a1); citric acid soluble (4D7a1); and New Zealand P Retention (4D8a1). The methods for total P analysis (4H1a1b1-21, 4H1b1a1a1-12) are described in the section of this manual titled “Total Analysis.”

Nitrogen is ubiquitous in the environment because it is continually cycled among plants, soil organisms, soil organic matter, water, and the atmosphere (National Research Council, 1993). Nitrogen is one of the most important plant nutrients and forms some of the most mobile compounds in the soil-crop system. As such, it is commonly related to water-quality problems. Total N includes both organic and inorganic forms. The KSSL method for total N (4H2a2) is described in the section of this manual titled “Total Analysis.”

Inorganic N in soils is predominately NO$_3^-$ and NH$_4^+$. Nitrite is seldom found in detectable amounts, except in neutral to alkaline soils receiving NH$_4^+$.
or $\text{NH}_4^-$ producing fertilizers (Maynard and Kalra, 1993; Mulvaney, 1996). There is considerable diversity among laboratories in the extraction and determination of $\text{NO}_3^-$ and $\text{NH}_4^+$ (Maynard and Kalra, 1993). Nitrate is water soluble, and a number of soil solutions including water have been used as extractants. The most common is KCl. Refer to Maynard and Kalra (1993) and Mulvaney (1996) for a review of extractants. The KSSL determines inorganic $\text{N (NO}_3^-)$ by KCl extraction, with a flow injection automated ion analyzer used to measure the soluble inorganic nitrate ($\text{NO}_3^-$) (4D9a1a1-2). The nitrate is quantitatively reduced to nitrite by passage of the sample through a copperized cadmium column. The nitrite (reduced nitrate plus original nitrite) is then determined by diazotizing with sulfanilamide followed by coupling with N-1-naphthylethylenediamine dihydrochloride. The resulting water soluble dye has a magenta color that is read at 520 nm.

The concept of an organic N fraction that is readily mineralized has been used to assess soil N availability in cropland, forests, and waste-disposal sites (Campbell et al., 1993). Incubation-leaching techniques have been used to quantify the mineralizable pool of soil organic N. These techniques may be aerobic or anaerobic. The KSSL determines mineralizable $\text{N (N as NH}_3$) by anaerobic incubation (4D10a1a1).

References


Soil Test Analyses (4D)

Anion Resin Exchange Extraction (4D1)

Double-Point Extraction (4D1a)

- 1-h, 23-h, 1 M NaCl (4D1a1a1)
- UV-Visible Spectrophotometer, Dual-Beam (4D1a1a1a)
- Phosphorus, Two Points (4D1a1a1a1-2)
- Air-dry or Field-Moist, <2 mm (4D1a1a1a1-2a-b1)

1. Application

Anion resins remove P from soils without chemical alterations and with only minor pH changes. Amounts of P released from soil and adsorbed by resins have been used as a measure of available P; an assessment of the availability of residual phosphates; an estimation of release characteristics and runoff P for agricultural land (Elrashidi, et al., 2003); and the buffer capacity of soils (Olsen and Sommers, 1982).

Plotting g log of extraction periods (0.25, 0.50, 1, 2, 4, 8, 24, 48 h) against amounts of P released (mg kg\(^{-1}\)) showed a linear relationship in 24 U.S. benchmark soils (Elrashidi et al., 2003). Two extraction periods (1 and 24 h) are sufficient to develop linear equations that predict P release characteristics (PRC), describing the whole relationship between the extraction time (1 min to 48 h) and amount of P released (mg kg\(^{-1}\)) for soils (Elrashidi et al., 2003). The method describes a two-point measurement (1 and 24 h extraction). This method is called the double-point anion exchange resin (DP-AER) extraction method. It is applied to investigate P availability, capacity, and release characteristics in soils (Elrashidi et al., 2012).

2. Summary of Method

A 2-g soil sample and 4-g resin bag are shaken with 100 mL of reverse osmosis deionized water for 1 h. Soil suspension is shaken again with another 4-g resin bag for 23 h. Phosphorus released from soil during shaking is adsorbed by resin. To remove P from resin, resin bags are shaken for 1 h in 1 M NaCl solution. Concentrated HCl is added to sample extracts. A 1-mL aliquot is diluted with 4 mL of ascorbic acid molybdate solution. Absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg\(^{-1}\) soil (4D1a1a1a1-2).

3. Interferences

The Mo blue methods, which are very sensitive for P, are based on the principle that in an acid molybdate solution containing orthophosphate ions, a phosphomolybdate complex forms that can be reduced by ascorbic acid, SnCl\(_2\).
and other reducing agents to a Mo color. The intensity of blue varies with the P concentration but is also affected by other factors, such as acidity, arsenates, silicates, and substances that influence the oxidation-reduction conditions of the system (Olsen and Sommers, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H₂SO₄ and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 100 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Bottles, polyethylene, 250, 125, and 60 mL
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, 150-mm
5.6 Cups, plastic
5.7 Dispenser, 50-mL
5.8 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL
5.9 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.10 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.11 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Anion exchange resin (AER), DOWEX Marathon, type II, 510–610 µm spherical beads (DOW Chemical Company), converted to bicarbonate form by soaking bags overnight in 1.0 M NaHCO₃ solution and washing out excess salt with RODI water. Store in RODI water in refrigerator.
6.3 Nitex nylon fabric, 300-µm pores, and nylon thread, Sefar America, Inc., for making and sewing resin bags
6.4 Hydrochloric acid (HCl), concentrated, 12 N, trace pure grade

6.5 NaCl solution. Dissolve 58.4 g NaCl in 1000 mL RODI water.

6.6 Sulfuric-tartrate-molybdate solution (STMS). Dissolve 60 g of ammonium molybdate tetrahydrate \((\text{NH}_4\text{H}_2\text{Mo}_7\text{O}_{24}\cdot4\text{H}_2\text{O})\) in 200 mL of boiling RODI water. Allow to cool to room temperature. Dissolve 1.455 g of antimony potassium tartrate (potassium antimony tartrate hemihydrate \([\text{K(SbO)}\text{C}_4\text{H}_6\text{O}_6\cdot\frac{1}{2}\text{H}_2\text{O}]\)) in the ammonium molybdate solution. Slowly and carefully add 700 mL of concentrated \(\text{H}_2\text{SO}_4\). Cool and dilute to 1 L with RODI water. Store in a dark bottle in the refrigerator.

6.7 Ascorbic acid solution. Dissolve 13.2 g of ascorbic acid in RODI water and dilute to 100 mL with RODI water. Make fresh daily.

6.8 Working ascorbic acid molybdate solution (WAMS). Mix 25 mL of STMS solution with 800 mL of RODI water. Add 10 mL of ascorbic acid solution and dilute to 1 L with RODI water. Allow to stand at least 1 h before using. Prepare fresh daily.

6.9 Working stock standard P solution (WSSPS), 100.0 mg P L\(^{-1}\). In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate \((\text{KH}_2\text{PO}_4)\) (dried for 2 h at 110 °C) in about 800 mL 1.0 \(\text{M}\) NaCl. Dilute to volume with NaCl solution and invert to mix thoroughly. Store in a polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.10 Standard P calibration solutions (SPCS), or working standards, 4.0, 3.0, 2.0, 1.0, 0.5, 0.25, and 0.0 mg P L\(^{-1}\). Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. To seven 100-mL volumetric flasks, add as follows:

6.10.1 4.0 mg P L\(^{-1}\)=4.0 mL WSSPS
6.10.2 3.0 mg P L\(^{-1}\)=3.0 mL WSSPS
6.10.3 2.0 mg P L\(^{-1}\)=2.0 mL WSSPS
6.10.4 1.0 mg P L\(^{-1}\)=1.0 mL WSSPS
6.10.5 0.5 mg P L\(^{-1}\)=0.5 mL WSSPS
6.10.6 0.25 mg P L\(^{-1}\)=0.25 mL WSSPS
6.10.7 0.0 mg P L\(^{-1}\)=0.0 mL WSSPS

Add 70 mL of 1.0 \(\text{M}\) NaCl solution and 4.0 mL of concentrated HCl to each SPCS. Allow to cool and dilute to mark with 1.0 \(\text{M}\) NaCl solution. Invert to mix thoroughly.

6.11 Quality Control Samples: 0.1 mg L\(^{-1}\) solution made from SSPS; blanks; selected SPCS; and KSSL standard.
7. Procedure

7.1 Weigh 2 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place in a 250-mL polyethylene bottle. Add 100 mL RODI water to bottle. Use two replicates for each soil sample in addition to a control treatment where all steps of the extraction process are performed in the absence of soil.

7.2 Place 4-g resin bag in the soil suspension and control sample. Transfer sample to shaker. Shake for 1 h at 100 oscillations min\(^{-1}\) at room temperature (20 ±2 °C). After shaking, remove P from resin as described in Section 7.4 and then proceed with Sections 7.5–7.14.

7.3 Place another 4-g resin bag in the soil suspension and control sample. Transfer sample to shaker. Shake for 23 h at 100 oscillations min\(^{-1}\) at room temperature (20 ±2 °C). After shaking, remove P from resin as described in Section 7.4 and then proceed with Sections 7.5–7.14.

7.4 Remove P from resin by lifting resin bag out of soil suspension and rinsing with 5 mL RODI water to remove attached soil particles. Add RODI water to soil suspension. If necessary, keep soil suspension for the next extraction.

7.5 Place resin bag in a 125-mL polyethylene bottle containing 50 mL of 1.0 M NaCl solution.

7.6 Transfer bottle to shaker and shake for 1 h at 100 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.7 Transfer extracting solutions to 60-mL polyethylene bottles. Filter if soil particles are observed in the extract. Add 2 mL concentrated HCl to each bottle. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.8 Use the pipette to transfer a 1-mL aliquot of the sample to a plastic cup. Also transfer a 1-mL aliquot of each SPCS to a plastic cup. Use a clean pipette tip for each sample and SPCS.

7.9 Dispense 4 mL of the WAMS to sample aliquot and to each SPCS. Swirl to mix. The color reaction requires a minimum of 20 min before analyst records readings. Complete all readings within a 2-h period because blue color may fade after this period.

7.10 Transfer sample extract and SPCS to cuvettes.

7.11 Set the spectrophotometer to read at 882 nm. Autozero with calibration blank.

7.12 Calibrate the instrument using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is R\(^2\) <0.99.
7.13 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; or if KSSL standard varies by more than 20% from the accepted mean. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.14 If samples are outside the calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations

Convert extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:

\[
\text{Soil P (mg kg}^{-1}) = \frac{A \times B \times C \times R \times 1000}{E}
\]

where:

- \(A\) = Sample extract reading (mg L\(^{-1}\))
- \(B\) = Extract volume (L)
- \(C\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- \(E\) = Sample weight (g)

9. Report

Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

References


Soil Test Analyses (4D)
Aqueous Extraction (4D2)
  Single-Point Extraction (4D2a)
    1:10, 30-min (4D2a1)
      UV-Visible Spectrophotometer, Dual-Beam (4D2a1a)
        Phosphorus (4D2a1a1)
          Air-Dry or Field-Moist, <2 mm (4D2a1a1a-b1)

1. Application

Phosphorus occurs in soil in both the solution phase and solid phase. These forms are well documented, but questions still remain concerning the exact nature of the constituents and ionic forms found in water, soils, and sediments (National Research Council, 1993). These forms influence P availability in relation to root absorption and plant growth; runoff and water quality problems; and P loadings. Water soluble P has been defined as P measured in water, dilute salt extracts (e.g., O.01 M CaCl₂), displaced soil solutions, or saturation paste extracts (Olsen and Sommers, 1982). Even though the water soluble fraction principally consists of inorganic orthophosphate ions, evidence indicates that some organic P is also included (Rigler, 1968).

The water or dilute salt extracts represent an attempt to approximate the soil solution P concentration. As an index of P availability, the objectives of this method are (1) to determine the P concentration level in the soil extract that limits plant growth (Olsen and Sommers, 1982) and (2) to determine the composition of the soil solution so that the chemical environment of the plant roots may be defined in quantitative terms (Adams, 1974). The sum of water soluble P and pH 3 extractable P has also been defined as the available P in runoff (Jackson, 1958).

2. Summary of Method

A 2.5-g sample of <2-mm, air-dry soil is mechanically shaken for 30 min in 25-mL of reverse osmosis deionized water. The sample is centrifuged until solution is free of soil mineral particles and filtered until clear extracts are obtained. Absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg⁻¹ soil (4D2a1a1).

3. Interferences

The Mo blue methods, which are very sensitive for P, are based on the principle that in an acid molybdate solution containing orthophosphate ions, a phosphomolybdate complex forms that can be reduced by ascorbic acid, SnCl₂, and other reducing agents to a Mo color. The intensity of blue varies with the P concentration but is also affected by other factors, such as acidity, arsenates,
silicates, and substances that influence the oxidation-reduction conditions of the system (Olsen and Sommers, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated \( \text{H}_2\text{SO}_4 \) and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min\(^{-1}\), 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, polyethylene
5.4 Filter paper, Whatman 42, 150-mm
5.5 Funnel, 60° angle, long stem, 50-mm diameter
5.6 Centrifuge, Centra GP-8, Thermo IEC, Needham Heights, MA
5.7 Volumetric flasks, 2-L, 100-mL, and 25-mL
5.8 Bottles, plastic, dark, 2-L
5.9 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-µL
5.10 Cups, plastic
5.11 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.12 Dispenser, 30-mL or 10-mL
5.13 Spectrophotometer, UV-Visible, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.14 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Sulfuric acid (\( \text{H}_2\text{SO}_4 \)), concentrated, 36 \( \text{N} \), trace pure grade
6.3 Molybdate solution. Dissolve 12.0 g ammonium molybdate tetrahydrate \([\text{(NH}_4\text{)}_6\text{Mo}_7\text{O}_{24} \cdot 4\text{H}_2\text{O}]\) in approximately 250 mL RODI water. Dissolve 0.2908 potassium antimony tartrate \([\text{K(SbO)}\text{C}_4\text{H}_4\text{O}_6 \cdot \frac{1}{2}\text{H}_2\text{O}]\) in 100 mL RODI water. Add these two dissolved reagents to 1-L 5 \( \text{N} \) \( \text{H}_2\text{SO}_4 \) (141 mL conc.)
H₂SO₄ diluted to 1-L), mix thoroughly, and dilute with RODI water to 2 L. Store in dark bottle and refrigerate (Reagent A).

6.4 Ascorbic acid: Dissolve 2.112 g of ascorbic acid in 400 mL of reagent A and mix (Reagent B). Prepare fresh daily.

6.5 Stock standard P solution (SSPS), 100.0 mg P L⁻¹. In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate (KH₂PO₄) (dried for 2 h at 110 °C) in about 800 mL RODI water. Dilute to volume with RODI water and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.6 Working stock standard P solution (WSSPS), 10.0 mg P L⁻¹. In a 1-L volumetric flask, pipette 100 mL SSPS and add 700 mL RODI water. Dilute to volume with RODI water and invert to thoroughly mix. Make fresh daily.

6.7 Standard P calibration solutions (SPCS), or working standards, 0.8, 0.6, 0.4, 0.2, and 0.0 mg P L⁻¹. Make fresh daily. To five 25-mL volumetric flasks, add as follows:

6.7.1 0.8 mg P L⁻¹ = 2.0 mL WSSPS
6.7.2 0.6 mg P L⁻¹ = 1.5 mL WSSPS
6.7.3 0.4 mg P L⁻¹ = 1.0 mL WSSPS
6.7.4 0.2 mg P L⁻¹ = 0.5 mL WSSPS
6.7.5 0.0 mg P L⁻¹ = 0.0 mL WSSPS (blank)

Add 4 mL reagent B, dilute each SPCS to mark with RODI water (extracting solution), and invert to thoroughly mix.

6.8 Quality Control Samples: 0.1 mg L⁻¹ solution made from SSPS; blanks; and selected SPCS. In addition, KSSL soil standard is routinely included in a batch for quality control.

7. Procedure

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place into a 50-mL centrifuge flask. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Add 25.0 mL of RODI water to sample. Place the sample in shaker and shake for 30 min at 200 oscillations min⁻¹ at room temperature (20 ±2 °C).

7.3 Remove the sample from the shaker. Centrifuge at 3000 rpm for 20 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 24 h.

7.4 Pipette 2-mL of sample extract into plastic cup, add 4-mL reagent B and 19 mL RODI water. Color is stable for 24 h. Maximum intensity develops in 10 min.
7.5 Transfer SPCS and sample extracts into 4.5-mL cuvettes.

7.6 Set spectrophotometer to 882 nm. Autozero with calibration blank.

7.7 Calibrate the instrument using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is \( R^2 < 0.99 \).

7.8 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; or if KSSL standard varies by more than 20% from the accepted mean. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.9 If samples are outside the calibration range, dilute samples with extracting solution and re-run.

8. Calculations

Convert extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:

\[
\text{Soil P (mg kg}^{-1}\text{)} = \frac{(A \times B \times C_1 \times C_2 \times R \times 1000)}{E}
\]

where:

- A = Sample extract reading (mg kg\(^{-1}\))
- B = Extract volume (L)
- \( C_1 \) = Dilution, required
- \( C_2 \) = Dilution, if performed
- \( R \) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- E = Sample weight (g)

9. Report

Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Tests (4D)
Aqueous Extraction (4D2)
  Single-Point Extraction (4D2a)
  1:5, 23-h, 1-h (4D2a2)
    Air-Dry or Field-Moist, <2 mm (4D2a2a-b1)

1. Application
   Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties (e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method
   A 20.0-g sample of soil is added to 100 mL of water in a 250-mL polyethylene bottle. The soil/water suspension is maintained at room temperature for 23 h and then shaken on a reciprocating shaker for 1 h. The supernatant is filtered into a 100-mL polyethylene bottle. The extract obtained in procedure 4D2a2 is analyzed for Br\(^-\), Cl\(^-\), F\(^-\), NO\(_3\)\(^-\), NO\(_2\)\(^-\), PO\(_4\)\(^3-\), and SO\(_4\)\(^2-\) by ion chromatography (4D2a2a1a1-7, respectively); CO\(_3\)\(^2-\) and HCO\(_3\)\(^-\) by acid titration (4D2a2b1a1-2, respectively); concentration of all or selected elements (Al, As, B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, Pb, Se, Zn, Si, Sr, and Mo) by an inductively coupled plasma atomic emission spectrophotometer (ICP–MS) (4D2a2d1-22, respectively); and nitrate-nitrogen by flow-injection, automated ion analyzer method 4D2a2c1a1. In addition, the sample extract is analyzed for pH and EC by methods 4C2a1a1a1 and 4F1b1a1, respectively.
3. Interferences
   No interferences are known for this method.

4. Safety
   No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment
   5.1 Electronic balance, ±1.0-mg sensitivity
   5.2 Polyethylene bottles, 100 and 250-mL
   5.3 Mechanical Vacuum Extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE (figs. 4B1-1 and 4B1-2)
   5.4 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
   5.5 Membrane screen filters, Millipore Durapore, PVDF hydrophilic, 0.45-µm diameter, EMD Millipore, Billerica, MA
   5.6 Centrifuge, International No. 11, with No. 949 rotor head, International Equip. Co., Boston, MA
   5.7 Mechanical reciprocating shaker, 100 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI

6. Reagents
   6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

7. Procedure
   7.1 Weigh 7.0 g of <2-mm or fine-grind, air-dry soil to the nearest mg, place into a 50-mL tube, and add 35 mL RODI water. If sample was moist, weigh enough soil to achieve ≈7.0 g of air-dry soil.
   7.2 Maintain the soil/water suspension at room temperature for 23 h.
   7.3 Transfer the sample to a shaker. Shake for 1 h at 100 oscillations min⁻¹ at room temperature (20 ±2 °C).
   7.4 Centrifuge the suspension at 2000 rpm for 20 min. Use the automatic extractor to filter the supernatant into 100-mL polyethylene bottle. Cap bottle. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

8. Calculations
   No calculations are required for this procedure. Calculations are reported in methods 4D21b1a1-7 for Br⁻, Cl⁻, F⁻, NO₃⁻, NO₂⁻, PO₄³⁻, and SO₄²⁻, respectively; in 4D2b1c1a1-2 for CO₃²⁻ and HCO₃⁻, respectively; in 4D2b1d1-2a1-22 for Al, As,
B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, Pb, Se, Zn, Si, Sr, and Mo, respectively; in 4D2b1f1 or 4D9a1a1 for NO$_3^-$; and in 4C2a1a1a1 and 4F1b1a1 for pH and EC, respectively.

9. Report

Data are reported by methods 4D21b1a1-7 for Br$^-$, Cl$^-$, F$^-$, NO$_3^-$, NO$_2^-$, PO$_4^{3-}$, and SO$_4^{2-}$, respectively; 4D2b1c1a1-2 for CO$_3^{2-}$ and HCO$_3^-$, respectively; 4D2b1d1-2a1-22 for Al, As, B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, Pb, Se, Zn, Si, Sr, and Mo, respectively; 4D9a1a1 for NO$_3^-$; and 4C2a1a1a1 and 4F1b1a1 for pH and EC, respectively.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Tests (4D)
Aqueous Extraction (4D2)
Single-Point Extraction (4D2a)
  1:5, 23-h, 1-h (4D2a2)
  Ion Chromatograph (4D2a2a)
    Conductivity Detector (4D2a2a1)
      Self-Generation Suppressor (4D2a2a1a)
        Bromide, Chloride, Fluoride, Nitrate, Nitrite,
        Phosphate, Sulfate (4D2a2a1a1-7)
    Air-Dry or Field-Moist, <2 mm (4D2a2a1a1-7a-b1)

1. Application

Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties
(e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method

The soil extract (4D2a2) is diluted according to its electrical conductivity (4F1b1a1). The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to measure the anion species and content. Standard anion concentrations are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. A computer program automates these actions. The saturation extract anions, \( \text{Br}^- \), \( \text{Cl}^- \), \( \text{F}^- \), \( \text{NO}_3^- \), \( \text{NO}_2^- \), \( \text{PO}_4^{3-} \), and \( \text{SO}_4^{2-} \), are reported in meq L\(^{-1}\) (mmol (−) L\(^{-1}\)) in procedures 4D2a2b1a1-7, respectively.

3. Interferences

Some soil extracts contain suspended solids. Filtering after dilution removes the particles. Organic anions that have low molecular weight will co-elute with inorganic anions from the column.

4. Safety

Wear protective clothing and safety glasses. Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.

5. Equipment

5.1 Ion chromatograph, double-column, conductivity detection, Dionex CS200, Dionex Corp., Sunnyvale, CA
5.2 Guard column, IonPac AG-18, 4 x 50 mm, Dionex Corp., Sunnyvale, CA
5.3 Analytical column, IonPac AS-18, Dionex Corp., Sunnyvale, CA
5.4 Self-regeneration suppressor, ASRS–300, 4 mm, Dionex Corp., Sunnyvale, CA
5.5 Autosampler, AS40, Dionex Corp., Sunnyvale, CA
5.6 Computer with PeakNet software and printer
5.7 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.8 Poly-vials with caps, 5 mL, Dionex Corp., Sunnyvale, CA
6. Reagents

6.1 Reverse osmosis deionized filtered (RODI) water, ASTM Type I grade of reagent water

6.2 Helium gas

6.3 Primary stock standards solutions, (PSSS\textsubscript{1000}), high purity, 1000 mg L\textsuperscript{-1}: Cl\textsuperscript{-}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}.

6.4 Mixed calibration standards solutions (MCSS), A, B, C, and D and Blank as follows:

\textbf{6.4.1} MCSSA = In a 500-mL volumetric flask, add as follows:

\textbf{6.4.1.1} 32 mL Cl\textsuperscript{−} PSSS\textsubscript{1000} = 64 mg L\textsuperscript{-1}
\textbf{6.4.1.2} 32 mL SO\textsubscript{4}\textsuperscript{2−} PSSS\textsubscript{1000} = 64 mg L\textsuperscript{-1}
\textbf{6.4.1.3} 2 mL F\textsuperscript{−} PSSS\textsubscript{1000} = 4 mg L\textsuperscript{-1}
\textbf{6.4.1.4} 8 mL NO\textsubscript{3}\textsuperscript{−} PSSS\textsubscript{1000} = 16 mg L\textsuperscript{-1}
\textbf{6.5.1.1} 2 mL NO\textsubscript{2}\textsuperscript{−} PSSS\textsubscript{1000} = 4 mg L\textsuperscript{-1}
\textbf{6.5.1.1} 2 mL Br\textsuperscript{−} PSSS\textsubscript{1000} = 4 mg L\textsuperscript{-1}
\textbf{6.5.1.1} 2 mL PO\textsubscript{4}\textsuperscript{3−} PSSS\textsubscript{1000} = 4 mg L\textsuperscript{-1}

Dilute to volume with RODI water and invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

\textbf{6.4.2} MCSSB: In a 100-mL volumetric flask, add 50 mL MCSSA and dilute to volume with RODI water. Final concentrations are 32, 32, 2, 8, 2, 2, and 2, mg L\textsuperscript{-1} Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}, respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

\textbf{6.4.3} MCSSC: In a 100-mL volumetric flask, add 50 mL MCSSB and dilute to volume with RODI water. Final concentrations are 16, 16, 1, 4, 1, 1, and 1 mg L\textsuperscript{-1} Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}, respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

\textbf{6.4.4} MCSSD: In a 100-mL volumetric flask, add 50 mL MCSSC and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 8, 8, 0.5, 2, 0.5, 0.5, and 0.5 mg L\textsuperscript{-1} Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}, respectively. Store in plastic containers in the refrigerator. Prepare fresh weekly.

\textbf{6.4.5} MCSS Blank: 0 mL of Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}. Dilute RODI water to volume.
6.6 Quality Control: In a 1-L volumetric flask, add 2.0 mL from each PSSS\textsubscript{1000} Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−} and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 2 mg L\textsuperscript{−1} Cl\textsuperscript{−}, SO\textsubscript{4}\textsuperscript{2−}, F\textsuperscript{−}, NO\textsubscript{3}\textsuperscript{−}, NO\textsubscript{2}\textsuperscript{−}, Br\textsuperscript{−}, and PO\textsubscript{4}\textsuperscript{3−}. Store in plastic containers in the refrigerator. Prepare fresh weekly.

7. Procedure

**Dilution of Sample Extracts**

7.1 To estimate the total soluble anion concentration (meq L\textsuperscript{−1}), multiply the \( EC_s \) (procedure 4F2b1) by 10. Subtract the CO\textsubscript{3}\textsuperscript{2−} and HCO\textsubscript{3}\textsuperscript{−} concentrations (procedures 4F2c1c1a1-2) from the total anion concentration. The remainder is the estimated concentration (meq L\textsuperscript{−1}) of anions to be separated by ion chromatography.

\[
\text{Anion concentration (meq L}^{-1}) = \text{EC}_s \times 10 - (\text{HCO}_3^- + \text{CO}_3^{2-})
\]

7.2 Dilute the saturation extract with the RODI water as follows:

<table>
<thead>
<tr>
<th>( EC_s ) (dS cm\textsuperscript{−1})</th>
<th>Dilution factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00 to 0.54</td>
<td>1</td>
</tr>
<tr>
<td>0.55 to 0.66</td>
<td>2</td>
</tr>
<tr>
<td>0.70 to 1.13</td>
<td>5</td>
</tr>
<tr>
<td>1.14 to 1.47</td>
<td>10</td>
</tr>
<tr>
<td>1.48 to 2.10</td>
<td>20</td>
</tr>
<tr>
<td>2.11 to 4.00</td>
<td>60</td>
</tr>
<tr>
<td>4.01 to 8.83</td>
<td>100</td>
</tr>
<tr>
<td>8.84 to 11.8</td>
<td>150</td>
</tr>
<tr>
<td>11.9 to 26.5</td>
<td>250</td>
</tr>
<tr>
<td>26.6 to 38.7</td>
<td>400</td>
</tr>
<tr>
<td>38.8 to 80.6</td>
<td>1000</td>
</tr>
<tr>
<td>&gt;80.7</td>
<td>2000</td>
</tr>
</tbody>
</table>

7.3 Place the MCSS (A, B,C, D, and Blank) and diluted extract samples in the Poly-vials and cap with filter caps.
Set-up and Operation of Ion Chromatograph (IC)

7.4 Refer to the manufacturer’s manual for the operation of chromatograph. Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex ICS-2000 ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. Ranges and/or (typical settings) are as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range and/or (typical setting)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calibration</td>
<td>Peak height or (area)</td>
</tr>
<tr>
<td>Flow setting</td>
<td>0.5 to 4.5 mL min⁻¹ (1.00 mL min⁻¹)</td>
</tr>
<tr>
<td>Pressure</td>
<td>200 to 3000 psi (2200 to 2400 psi)</td>
</tr>
<tr>
<td>Detection</td>
<td>Suppressed conductivity</td>
</tr>
<tr>
<td>Total conductivity</td>
<td>0 to 999.9 µS</td>
</tr>
<tr>
<td>Injection volume</td>
<td>10 µL</td>
</tr>
<tr>
<td>Auto offset</td>
<td>-999.9 to 999.9 µS (On)</td>
</tr>
<tr>
<td>Cell temperature</td>
<td>35 ºC</td>
</tr>
<tr>
<td>Suppressor current</td>
<td>75 mA</td>
</tr>
</tbody>
</table>

7.5 Load the sample holder cassettes with the capped samples, standards, and check samples.

7.6 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

IC Calibration and Analysis

7.7 Calibrate the instrument by using the MCSS (A, B, C, D, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.8 If samples are outside calibration, dilute sample extracts with RODI water solution and re-analyze.

7.9 Perform one quality control (QC) (Low Standard MCSS, Standard C) for every 12 samples. If reading is not within tolerance limits (10 to 15%, based on analyte), the instrument is re-calibrated and QC re-analyzed.

7.10 Record analyte readings to 0.01 mg L⁻¹.

8. Calculations
   The instrument readings for analyte concentration are in mg L⁻¹. These analyte concentrations are converted to meq L⁻¹ as follows:
Analyte Concentration in Soil (meq L\(^{-1}\)) = (A \times B) / C

where:
A = Analyte (Br\(^-\), Cl\(^-\), F\(^-\), NO\(_3^-\), NO\(_2^-\), PO\(_4^{3-}\), SO\(_4^{2-}\)) concentration in extract (mg L\(^{-1}\))
B = Dilution ratio, if needed
C = Equivalent weight

where:
Cl\(^-\) = 35.45 mg meq\(^{-1}\)
SO\(_4^{2-}\) = 48.03 mg meq\(^{-1}\)
F\(^-\) = 19.00 mg meq\(^{-1}\)
NO\(_3^-\) = 62.00 mg meq\(^{-1}\)
NO\(_2^-\) = 46.00 mg meq\(^{-1}\)
Br\(^-\) = 79.90 mg meq\(^{-1}\)
PO\(_4^{3-}\) = 31.66 mg meq\(^{-1}\)

9. Report
Report the 1:5 aqueous extraction anions (Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\)) to the nearest 0.1 meq L\(^{-1}\) (mmol (−) L\(^{-1}\)).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Tests (4D)
Aqueous Extraction (4D2)
  Single-Point Extraction (4D2a)
    1:5, 23-h, 1-h (4D2a2)
  Automatic Titrator (4D2a2b)
    Combination pH-Reference Electrode (4D2a2b1)
      Acid Titration, H$_2$SO$_4$ (4D2a2b1a)
      Carbonate and Bicarbonate (4D2a2b1a1-2)
      Air-Dry or Field-Moist, <2 mm (4D2a2b1a1-2a-b1)

1. Application

Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties (e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method

An aliquot of the soil extract (procedure 4D2a2) is titrated on an automatic titrator to pH 8.25 and pH 4.60 end points. The carbonate and bicarbonate are calculated from the titers, aliquot volume, blank titer, and acid normality (4D2a2b1a1-2 procedures, respectively). Carbonate and bicarbonate are reported in meq L$^{-1}$ (mmol (+) L$^{-1}$). If the pH of the saturated paste extract $\leq$4.60, then carbonate and bicarbonate are not determined.

3. Interferences

Clean the electrode by rinsing with distilled water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

Slow electrode response time may cause the end point to be overshot. A combination of slowing the beret speed and increasing the time delay may help. Cleaning the electrode with detergent may decrease the response time. If all else fails, changing the electrode generally solves the problem. Blanks may not titrate properly because some sources of reverse osmosis (RO) water have a low pH.
4. Safety

Wear protective clothing and eye protection. Exercise care when preparing reagents. Thoroughly wash hands after handling reagents. Restrict the use of concentrated $\text{H}_2\text{SO}_4$ to the fume hood. Use showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow the manufacturer’s safety precautions when operating the automatic titrator.

5. Equipment

5.1 Automatic titrator, Metrohm 670 Titroprocessor, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.

5.2 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.

5.3 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

6.2 Helium gas

6.3 Sulfuric acid ($\text{H}_2\text{SO}_4$), concentrated, 36 N

6.4 $\text{H}_2\text{SO}_4$, 0.0240 N standardized. Carefully dilute 2.67 mL of concentrated $\text{H}_2\text{SO}_4$ in 4 L of RODI degassed water (≈15 min). Re-standardize the acid at regular intervals. Refer to the procedure for standardization of acids.

6.5 Borax pH buffers, pH 4.00, 7.00, and 9.18, for titrator calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Prepare an extract (procedure 4D2a2).

7.2 Pipette 3 mL of the fresh extract (procedure 4D2a2) into a 250-mL titration beaker.

7.3 Add 72 mL of RO water into a titration beaker. Final volume is 75 mL for blanks and samples. Run 8 to 12 blanks of RO water through the titration procedure.

7.4 Refer to manufacturer’s manual for operation of the automatic titrator.

7.5 Calibrate automatic titrator with 9.18, 7.00, and 4.00 pH buffers. Set-up the automatic titrator to set end point titration mode. The “Set” pH parameters are listed as follows:
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{p_1}$</td>
<td>pH 8.25</td>
</tr>
<tr>
<td>Dyn change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s$^{-1}$</td>
</tr>
<tr>
<td>Time delay</td>
<td>10 s</td>
</tr>
<tr>
<td>$E_{p_2}$</td>
<td>pH 4.60</td>
</tr>
<tr>
<td>Dyn change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s$^{-1}$</td>
</tr>
<tr>
<td>Temp</td>
<td>25 °C</td>
</tr>
<tr>
<td>Stop volume</td>
<td>35 mL</td>
</tr>
</tbody>
</table>

7.6 Place the 250-mL titration beakers in the sample changer.
7.7 Press “Start.”
7.8 If the titrator is operating properly, no other analyst intervention is required. The titers and other titration parameters are recorded on the Microprocessor printer.

8. Calculations
8.1 $\text{CO}_3^{2-}$ (meq L$^{-1}$) $= (2T_1 \times N \times 1000)/\text{Aliquot}$
8.2 $\text{HCO}_3^{-}$ (meq L$^{-1}$) $= \frac{T_2 + T_1 - \text{Blank} - (2 \times T_1 \times N \times 1000)}{\text{Aliquot}}$

where:
- $T_1$ = Titer of $\text{CO}_3^{2-}$ (mL)
- $T_2$ = Titer of $\text{HCO}_3^{-}$ (mL)
- $N$ = Normality of $\text{H}_2\text{SO}_4$
- Blank = Average titer of blank solutions (mL)
- Aliquot = Volume of saturation extract titrated (mL)
- 1000 = Conversion factor to meq L$^{-1}$

9. Report
Report extracted anions $\text{CO}_3^{2-}$ and $\text{HCO}_3^{-}$ to the nearest 0.1 meq L$^{-1}$ (mmol (−) L$^{-1}$).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Soil Test Analyses (4D)
Aqueous Extraction (4D2)
Single-Point Extraction (4D2a)
1:5, 23-h, 1-h (4D2a2)
  Cadmium-Copper Reduction (4D2a2c)
    Sulfanilamide N-1-Naphthylethylenediamine
    Dihydrochloride (4D2a1c1)
    Flow-Injection, Automated Ion Analyzer (4D2a2c1a)
      Nitrate (4D2a2c1a 1)
      Air-Dry or Field-Moist, <2 mm (4D2a2c1a1a-b1)

1. Application
The 1:5 aqueous extraction of nitrate is used in soil taxonomy as criteria for two taxa of the Gelisols order. The subgroups of Nitric Anhyorthels and Nitric Anhyturbels are defined by a minimum nitrate concentration of 118 mmol (−) /L in a horizon at least 15 cm thick. A additional part of the criterion uses a simple calculation of the nitrate content times the horizon thickness to connote a significant amount of nitrate accumulation. The product of nitrate concentration times thickness (cm) must be ≥3500. Based on this calculation, a horizon that is only 15 cm thick must have a nitrate concentration of 233 mmol (−)/L to qualify, whereas a horizon with a nitrate concentration of only 118 mmol (−)/L must be twice as thick (i.e., 30 cm) to meet the criterion (Soil Survey Staff, 2014).

2. Summary of Method
A 2.5-g soil sample is mechanically shaken for 30 min in 25 mL of reverse osmosis deionized water (RODI). The sample is then filtered through Whatman No. 42 filter paper. A flow injection automated ion analyzer is used to measure the soluble inorganic nitrate (NO₃⁻). The nitrate is quantitatively reduced to nitrite by passage of the sample through a copperized cadmium column. The nitrite (reduced nitrate plus original nitrite) is then determined by diazotizing with sulfanilamide followed by coupling with N-1-naphthylethylenediamine dihydrochloride. The resulting water soluble dye has a magenta color, which is read at 520 nm. Absorbance is proportional to the concentration of NO₃⁻ in the sample. Data are reported as mmol (−) L⁻¹ as NO₃⁻ (4D2a2c1a1).
3. Interferences

Nitrite is oxidized by air to nitrate in a few days. If analysis can be made within 24 h, the sample should be preserved by refrigeration at 4 ºC. When samples must be stored for more than 24 h, they should be preserved with sulfuric acid (2 mL concentrated H₂SO₄ per liter) and refrigerated (LACHAT, 2003). Low results can be obtained for samples that contain high concentration of Fe, Cu, or other metals. In this method, EDTA is added to the buffer to reduce this interference (LACHAT, 2003).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of NH₄OH and concentrated HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Cadmium is toxic and carcinogenic. Wear gloves and follow the precautions described on the Material Safety Data Sheet. If the cadmium-copper reduction column is repacked, all transfers should be done over a special tray or beaker dedicated to this purpose. Preferably, send the cadmium-copper column to LACHAT for repacking.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Centrifuge tubes, 25-mL, polyethylene, Oak Ridge
5.3 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Centrifuge, 50-mL, polyethylene
5.5 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL
5.6 Filter paper, Whatman 42, 150-mm
5.7 Funnel, 60° angle, long stem, 50-mm diameter
5.8 Volumetric flasks, 1-L and 250-mL
5.9 Bottles, plastic, dark, 1-L
5.10 Cups, plastic
5.11 Dispenser, 30-mL or 10-mL
5.12 Flow Injection Automated Ion Analyzer, QuikChem 8500, LACHAT Instruments, Loveland, CO, with computer and printer
5.13 Sampler, LACHAT Instruments, Loveland, CO
5.14 Reagent Pump, LACHAT Instruments, Loveland, CO
5.15 Automated Dilution Station, LACHAT Instruments, Loveland, CO
5.16 Sample Processing Module (SPM) or channel, QuikChem Method (12-107-04-1-B, nitrate in 1 M KCl 0.025 to 20.0 mg N L\(^{-1}\)), LACHAT Instruments, Loveland, CO
5.17 Computer, with QuikChem software, LACHAT Instruments, Loveland, CO, and printer
5.18 Vials, plastic, 25-mL (standards)
5.19 Culture tubes, glass, 10-mL (samples)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Helium, compressed gas
6.3 15 M NaOH. In a 500-mL container, add 250 mL RODI water. Slowly add 300 g NaOH. (CAUTION: The solution will get very hot!) Swirl until dissolved. Cool and store in a plastic bottle. Use to adjust ammonium chloride buffer to pH 8.5 (Reagent 6.4).
6.4 Ammonium chloride buffer, pH 8.5. In a hood, add 500 mL RODI to a 1-L volumetric flask. Add 105 mL concentrated HCl, 90 mL ammonium hydroxide (NH\(_4\)OH), and 1.0 g disodium EDTA. Dissolve and dilute to mark. Invert to mix. Degas with helium ≈5 min.
6.5 Sulfanilamide color reagent. To a 1-L volumetric flask, add 600 mL RODI H\(_2\)O followed by 100 mL 85 percent phosphoric acid (H\(_3\)PO\(_4\)), 40.0 g sulfanilamide, and 1.0 g N-1-naphthylethylenediamine dihydrochloride (NED). Shake to wet and stir to dissolve 20 min. Dilute to mark, invert to thoroughly mix. Degas with helium ≈5 min. Store in dark bottle and discard when pink.
6.6 1 M KCl extracting solution, carrier, and standards diluent. Dissolve 74.5 g potassium chloride (KCl) in 800 mL RODI water. Dilute to mark and invert to thoroughly mix. The extracting solution is used also as the carrier and a component of the N standards. Degas with helium ≈5 min.
6.7 The following are standards for a 1-channel system determining NO\(_2\)^− + NO\(_3\)^− or NO\(_2\)^− and a 2-channel system where one channel is used for determining NO\(_2\)^− + NO\(_3\)^− and the other channel is used for determining NO\(_2\)^−. For the 1-channel system, either NO\(_2\)^− or NO\(_3\)^− standards may be used. It is recommended that when running a 1 channel method for NO\(_2\)^− + NO\(_3\)^− that NO\(_2\)^− standards are used. For the 2-channel system, the use of both NO\(_2\)^− + NO\(_3\)^− standard sets is recommended.
6.7.1 Stock standard nitrate solution (SSNO\(_3\)S), 200.0 mg N L\(^{-1}\) as NO\(_3\)^− in 1 M KCl. In a 1-L volumetric flask, dissolve 1.444 g potassium nitrate (KNO\(_3\)) (dried in an oven for 2 h at 110 °C)
and 74.5 g KCl in 600 mL RODI water. Dilute to mark with RODI water and invert to thoroughly mix. Store in polyethylene containers in a refrigerator. Make fresh weekly.

6.7.2 Working stock standard nitrate solution (WSSNO₃S), 20.0 mg N L⁻¹ as NO₃⁻ in 1 M KCl. To a 1-L volumetric flask, add 100 mL SSNO₃S. Dilute to mark with 1 M KCl and invert to thoroughly mix. Make fresh daily.

6.7.3 Standard nitrate calibration standards (SNO₃CS), or working standards, 10.00, 1.00, 0.80, 0.08, and 0.00 mg N L⁻¹ as NO₃⁻ in 1 M KCl. Make fresh daily. To five 250-mL volumetric flasks, add as follows:

- 6.7.3.1 10.00 mg N L⁻¹ = 125.0 mL WSSNO₃S
- 6.7.3.2 1.00 mg N L⁻¹ = 12.5 mL WSSNO₃S
- 6.7.3.3 0.80 mg N L⁻¹ = 10.0 mL WSSNO₃S
- 6.7.3.4 0.08 mg N L⁻¹ = 1.00 mL WSSNO₃S
- 6.7.3.5 0.00 mg N L⁻¹ = 0.0 mL WSSNO₃S (blank)

Dilute each SNO₃CS to the mark with 1 M KCl and invert to thoroughly mix. Do not degas.

6.7.4 Stock standard nitrite solution (SSNO₂S), 200.0 mg N L⁻¹ as NO₂⁻ in 1 M KCl. In a 1-L volumetric flask, dissolve 74.5 g KCl and either 0.986 g sodium nitrite (NaNO₂) or 1.214 g potassium nitrite (KNO₂) in 800 mL RODI water. Dilute to mark with RODI water and invert to thoroughly mix. Store in polyethylene containers in a refrigerator. Make fresh weekly.

6.7.5 Working stock standard nitrite solution (WSSNO₂S), 20.0 mg N L⁻¹ as NO₂⁻ in 1 M KCl. To a 1-L volumetric flask, add 100 mL SSNO₂S. Dilute to mark with 1 M KCl and invert to thoroughly mix. Make fresh daily.

6.7.6 Standard nitrite calibration standards (SNO₂CS), or working standards, 10.00, 1.00, 0.80, 0.08, and 0.00 mg N L⁻¹ as NO₂⁻ in 1 M KCl. Make fresh daily. To five 250-mL volumetric flasks, add as follows:

- 6.7.6.1 10.00 mg N L⁻¹ = 125.0 mL WSSNO₂S
- 6.7.6.2 1.00 mg N L⁻¹ = 12.5 mL WSSNO₂S
- 6.7.6.3 0.80 mg N L⁻¹ = 10.0 mL WSSNO₂S
- 6.7.6.4 0.08 mg N L⁻¹ = 1.00 mL WSSNO₂S
- 6.7.6.5 0.00 mg N L⁻¹ = 0.0 mL WSSNO₂S (blank)

Dilute each SNO₂CS to the mark with 1 M KCl and invert to thoroughly mix. Do not degas.
7. Procedure

**Extraction**

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to the nearest mg and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Add ≈25 mL of RODI water to sample. Transfer the sample to a shaker. Shake for 30 min at 200 oscillations min⁻¹ at room temperature (20 ±2 °C).

7.3 Remove the sample from the shaker. Decant, filter, and collect extract in receiving cups. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 24 h.

**Flow Injection Set-up and Operation**

7.4 Transfer sample extracts into culture tubes and place in sample trays marked “Samples.”

7.5 Transfer working calibration standards into plastic vials and place in descending order in sample trays marked “Standards.”

7.6 Refer to the operating and software reference manuals for LACHAT for set-up and operation. Refer to LACHAT Method QuikChem Method 12-107-04-1-B for data system parameters, such as analyte and calibration data and sampler and valve timing.

7.7 Run samples using calibration curve. Sample concentration is calculated from the regression equations. Report results to the nearest 0.01 unit for the sample extract and each standard nitrogen calibration standard.

7.8 If samples are outside calibration range, dilute samples with extracting solution and re-analyze.

7.9 Upon completion of run, place the transmission lines into RODI water and pump for approximately 20 min before proceeding with the normal shut-down procedure.

7.10 KCl may accumulate and cause clogs in the manifold tubing and the fittings over time. The valves and fittings therefore need to be washed with RODI water upon completion of analysis. Some fittings may need to be soaked overnight or placed in a sonic bath for 10 to 15 min to remove KCl accumulations.

8. Calculations

Convert extract N (mg L⁻¹) to mmol (+) L⁻¹ as follows:

Soil N = (A x B) / C

where:

A = Sample extract reading (mg N L⁻¹)
B = Dilution, if performed  
C = Molecular weight (NO$_3^-$) = 62.0 mg mmol$^{-1}$

9. Report
Report data to the nearest 0.1 mmol (−) L$^{-1}$ as NO$_3^-$.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
LACHAT Instruments. 2003. QuikChem method 12-107-04-1-B, nitrate in 2 M KCl soil extracts by flow injection analysis, 0.025 to 20.0 mg N L$^{-1}$. LACHAT Instruments, Loveland, CO.

Soil Tests (4D)
Aqueous Extraction (4D2)
Single-Point Extraction (4D2a)
1:5, 23-h, 1-h (4D2a2)
Inductively Coupled Plasma Mass Spectrophotometer (4D2a2d)
Aluminum, Arsenic, Barium, Boron, Cadmium, Calcium, Chromium, Cobalt, Copper, Iron, Lead, Magnesium, Manganese, Molybdenum, Nickel, Phosphorus, Potassium, Selenium, Silicon, Sodium, Strontium, and Zinc (4D2a2d1-22)
Air-Dry or Field-Moist, <2-mm (4D2a2d1-22a-b1)

1. Application
Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties (e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method
The extract is prepared in procedure 4D2a1. Calibration standards plus a blank are prepared for elemental analysis. The concentration of all or selected
elements (Al, As, B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Mo, Na, Ni, P, Pb, Se, Sr, Si, and Zn) are determined by inductively coupled plasma mass spectrophotometer (ICP–MS). Data from this procedure (4D2a2d1-22) are reported as mg kg$^{-1}$ soil.

3. Interferences

Interferences are corrected or minimized by using an internal standard, collision/reaction cell technology, and careful selection of specific masses for data reporting. Interference corrections are made by ICP–MS software.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents.

5. Equipment

5.1 Pipettes, electronic digital, 250 µL and 10 mL, Rainin Instrument Co., Woburn, MA

5.2 Inductively coupled plasma mass spectrophotometer (ICP–MS), Agilent 7500cx, Agilent Technologies Inc. Wilmington, DE

5.3 Computer, with ICP–MS ChemStation software ver. B.03.07, Agilent Technologies Inc., Wilmington, DE

5.4 Heat Exchanger, G1879B, Agilent Technologies

5.5 Compressed gasses, argon (minimum purity 99.99%), hydrogen (minimum purity 99.999%) and helium (minimum purity 99.999%)

5.6 Autosampler, ASX-500 Series, Agilent Technologies Inc., Wilmington, DE

5.7 Quartz torch, for use with HMI, Part No. G3270-80027

5.8 Peristaltic pump (for automatic injection of internal standard)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 Concentrated hydrochloric acid (HCl), 12 N, trace pure grade

6.3 Concentrated nitric acid (HNO$_3$), 16 N, trace pure grade

6.4 Selenium Standard, commercially prepared solution containing 1000 mg/mL Se

6.5 Water Extractable Elements, commercially prepared solution containing 1000 µg/mL Ca, K, Mg; 150 µg/mL P, Sr; 100 µg/mL Al, Ba, Fe, Mn; 50 µg/mL Cu, V, Zn; 10 µg/mL Co, Cr, Ni, Pb; 5 µg/mL As, Cd; 1 µg/mL Mo
6.6 Phosphorus Standard, commercially prepared solution containing 1000 mg/mL P
6.7 Boron Standard, commercially prepared solution containing 1000 mg/mL B
6.8 Silicon Standard, commercially prepared solution containing 1000 mg/mL Si
6.9 Gold Standard, commercially prepared solution containing 1000 mg/mL Au
6.10 Lithium Standard, commercially prepared solution containing 1000 mg/mL Li
6.11 Scandium Standard, commercially prepared solution containing 1000 mg/mL Sc
6.12 Germanium Standard, commercially prepared solution containing 1000 mg/mL Ge
6.13 Yttrium Standard, commercially prepared solution containing 1000 mg/mL Y
6.14 Terbium Standard, commercially prepared solution containing 1000 mg/mL Tb
6.15 Bismuth Standard, commercially prepared solution containing 1000 mg/mL Bi
6.16 Agilent Stock Tuning Solution
6.17 Agilent 7500 Series PA Tuning 1 Solution

7. Procedure

**ICP–MS Calibration Standards, Set-up, and Operations**

7.1 Tuning Solution: In a 1L volumetric flask, add 300 mL RODI water, 1 mL commercially prepared Agilent Stock Tuning Solution (Reagent 6.16), and 18 mL concentrated HNO₃. Fill to volume with RODI water and mix well.

7.2 PA Tuning Solution: In 1-L volumetric, add 500 mL RODI water, 18 mL concentrated HNO₃, and 100 mL of commercially prepared Agilent 7500 Series PA Tuning 1 (Reagent 6.17). Fill to volume and mix well.

7.3 10 µg/mL Selenium Stock: In 500-mL flask, add 5 mL 1,000 µg/mL Se (Reagent 6.4) and 9 mL concentrated HNO₃. Fill to volume with RODI water and mix well.

7.4 Mixed Elements Stock: In 500-mL flask, add 300 mL RODI water, 9 mL concentrated HNO₃, 5 mL Water Extractable Elements (Reagent 6.5), 5 mL 10 µg/mL Se stock (Reagent 7.3), and 9 mL concentrated HNO₃. Fill to volume with RODI water and mix well.

7.5 Mixed Elements High: In 500-mL flask, add 300 mL RODI water and 50 mL Mixed Elements Stock (Reagent 7.4). Fill to volume with RODI water and mix well.
7.6 Mixed Elements Medium: In 500-mL flask, add 300 mL RODI water and 50 mL Mixed Elements High (Reagent 7.5). Fill to volume with RODI and mix well.

7.7 Mixed Elements Low: In a 500 mL flask, add 300 mL RODI water and 50 mL Mixed Elements Medium (Reagent 7.6). Fill to volume with RODI and mix well.

7.8 P1000: In 1-L volumetric flask, add 500 mL RODI water and 1 mL Phosphorus Standard (Reagent 6.6). Fill to volume with RODI water and mix well.

7.9 P100: In 1-L volumetric flask, add 500 mL RODI water and 100 mL of P1000 (Reagent 7.8). Fill to volume with RODI water and mix well.

7.10 B1000: In a 1-L volumetric flask, add 500 mL RODI water and 1 mL Boron Standard (Reagent 6.7). Fill to volume with RODI water and mix well. Transfer to 1-L polypropylene bottle.

7.11 B100: In 1-L volumetric flask, add 500 mL RODI water and 100 mL B1000 (Reagent 7.10). Fill to volume with RODI and mix well. Transfer to 1-L polypropylene bottle.

7.12 Si1000: In 1-L volumetric flask, add 500 mL RODI water and 1 mL Silicon Standard (Reagent 6.8). Fill to volume with RODI and mix well.

7.13 Si100: In 1-L volumetric flask, add 500 mL RODI and 100 mL Si1000 (Reagent 7.12). Fill to volume with RODI water and mix well.

7.14 Internal Standard (1 µg/mL Li⁺, Sc, Ge, Y, In, Tb, Bi): In 1-L flask, add 300 mL RODI water, 18 mL concentrated HNO₃, 6 mL concentrated HCl, 0.250 mL 1000 µg/mL Au (Reagent 6.9), and 1 mL each of 1000 µg/mL Li⁺, Sc, Ge, Y, In, Tb, and Bi (Reagents 6.10–6.15). Fill to volume with RODI water and mix well.

7.15 Rinse: In 2-L flask, add 300 mL RODI water and 58 mL concentrated HNO₃. Fill to volume with RODI and mix well.

7.16 Rinse #1: In 1-L flask, add 300 mL RODI water, 29 mL concentrated HNO₃, and 1 mL 1000 µg/mL Au (Reagent 6.9). Fill to volume with RODI water and mix well.

7.17 Rinse #2: In 1-L flask, add 300 mL RODI, 15 mL concentrated HNO₃, 45 mL concentrated HCl, and 1 mL of 1000 µg/mL Au (Reagent 6.9). Fill to volume with RODI water and mix well.

7.18 Rinse #3: RODI water.
### 7.20 Standard concentrations in \( \mu g/mL \) for each element.

<table>
<thead>
<tr>
<th>Element</th>
<th>Blank</th>
<th>Mixed elements low</th>
<th>Mixed elements mid</th>
<th>Mixed elements high</th>
<th>P100</th>
<th>P1000</th>
<th>B100</th>
<th>B1000</th>
<th>Si100</th>
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<td>---</td>
<td>---</td>
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### 7.21 Reporting m/z and tune step for each element analyzed.

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<tr>
<th>Element</th>
<th>m/z</th>
<th>Tune 1 ((H_2))</th>
<th>Tune 2 ((He))</th>
<th>Tune 3 (\text{no gas})</th>
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<tbody>
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<td>Element</td>
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<td>Tune 2 (He)</td>
<td>Tune 3 (no gas)</td>
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7.22 The ICP–MS is set up with an HMI system using a Burgener Mira mist nebulizer, quartz torch, and spray chamber to analyze samples. Internal standard is added via peristaltic pump using 0.19 mm i.d. pump tubing. Internal standard is mixed with the samples, or standards are added via coil prior to entering the nebulizer. Samples are diluted 1:10 or greater as necessary prior to analysis with sample diluent (Reagent 6.17). Perform instrument checks (tune for sensitivity, resolution axis, P/A factor, internal standard RSD, torch alignment, and EM tune) prior to analysis as discussed in operation manual of instrument. Check instrument gas pressures to ensure pressures are correct and in adequate supply.
7.23  Typical tune values for 1:5 aqueous extraction are as follows:

<table>
<thead>
<tr>
<th>Tune 1 (H₂)</th>
<th>Tune 1 (H₂)</th>
</tr>
</thead>
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<tr>
<td><strong>Plasma Parameters</strong></td>
<td><strong>Q-Pole Parameters</strong></td>
</tr>
<tr>
<td>RF power</td>
<td>1550 W</td>
</tr>
<tr>
<td>RF matching</td>
<td>1.78 V</td>
</tr>
<tr>
<td>Smpl depth</td>
<td>10.0 mm</td>
</tr>
<tr>
<td>Torch-H</td>
<td>0.4 mm</td>
</tr>
<tr>
<td>Torch-V</td>
<td>-0.4 mm</td>
</tr>
<tr>
<td>Carrier gas</td>
<td>0.50 L/min</td>
</tr>
<tr>
<td>Makeup gas</td>
<td>0.50 L/min</td>
</tr>
<tr>
<td>Optional gas</td>
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</tr>
<tr>
<td>Nebulizer pump</td>
<td>0.10 rps</td>
</tr>
<tr>
<td>Sample pump</td>
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</tr>
<tr>
<td>S/C temp</td>
<td>2 °C</td>
</tr>
<tr>
<td><strong>Ion Lenses</strong></td>
<td>He gas</td>
</tr>
<tr>
<td>Extract 1</td>
<td>0.0 V</td>
</tr>
<tr>
<td>Extract 2</td>
<td>-135.0 V</td>
</tr>
<tr>
<td>Omega Bias-ce</td>
<td>-24 V</td>
</tr>
<tr>
<td>Omega Lens-ce</td>
<td>-1.0 V</td>
</tr>
<tr>
<td>Cell entrance</td>
<td>-40 V</td>
</tr>
<tr>
<td>QP focus</td>
<td>-8 V</td>
</tr>
<tr>
<td>Cell exit</td>
<td>-40 V</td>
</tr>
<tr>
<td>Plasma Parameters</td>
<td>Tune 2 (He)</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>RF power</td>
<td>1550 W</td>
</tr>
<tr>
<td>RF matching</td>
<td>1.78 V</td>
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<tr>
<td>Smpl depth</td>
<td>10.0 mm</td>
</tr>
<tr>
<td>Torch-H</td>
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</tr>
<tr>
<td>Torch-V</td>
<td>-0.4 mm</td>
</tr>
<tr>
<td>Carrier gas</td>
<td>0.50 L/min</td>
</tr>
<tr>
<td>Makeup gas</td>
<td>0.50 L/min</td>
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<tr>
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<tr>
<td>S/C temp</td>
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<table>
<thead>
<tr>
<th>Ion Lenses</th>
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<tbody>
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<td>Extract 1</td>
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<tr>
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<tr>
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<tr>
<td>Omega Lens-ce</td>
<td>-1.0 V</td>
</tr>
<tr>
<td>Cell entrance</td>
<td>-40 V</td>
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<tr>
<td>Cell exit</td>
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<table>
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<tbody>
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<tr>
<td>He gas</td>
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<td>Tune 3 (No Gas)</td>
<td>Tune 3 (No Gas)</td>
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<td>----------------</td>
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<tr>
<td><strong>Plasma Parameters</strong></td>
<td><strong>Q-Pole Parameters</strong></td>
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<tr>
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<td>AMU gain</td>
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<td>Torch-H</td>
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</tr>
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<td>Torch-V</td>
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<td>Carrier gas</td>
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<tr>
<td>Makeup gas</td>
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</tr>
<tr>
<td>Optional gas</td>
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<tr>
<td>Nebulizer pump</td>
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<td>Sample pump</td>
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<tr>
<td>S/C temp</td>
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<td><strong>Ion Lenses</strong></td>
<td><strong>Octapole Parameters</strong></td>
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<td>Extract 2</td>
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<tr>
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<tr>
<td>Omega Lens-ce</td>
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<tr>
<td>QP focus</td>
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</tr>
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<td>Analog HV</td>
</tr>
<tr>
<td>He gas</td>
<td>Pulse HV</td>
</tr>
<tr>
<td>Optional gas</td>
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</tbody>
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320
7.24 Establish detection limits using the blank standard solution. The instrumental detection limits are calculated using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are reported as “ND” or non-detected.

8. Calculations
The calculation of mg kg\(^{-1}\) of an element in the soil from µg L\(^{-1}\) in solution is as follows:

\[
\text{Analyte concentration in soil (mg kg}\,^\text{−1}) = \left[\frac{A \times B \times C \times R \times 1000}{E}\right] \times 1000
\]

Where:
A = Sample extract reading (µg L\(^{-1}\))
B = Extract volume (L)
C = Dilution, if performed
R = Air-dry/oven-dry or field-moist/oven-dry ratio (method 3D1 and 3D2, respectively)
1000 = Conversion factor in numerator to kg-basis
E = Sample weight (g)
1000 = Factor in denominator (µg mg\(^{-1}\))

9. Report
Analysis is generally done on one mass per element. If more than one mass is analyzed, only the reporting mass is used for data reporting purposes. The particle-size fraction digested needs to be identified with each sample. Data are reported to the nearest 0.01 mg kg\(^{-1}\).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
Soil Tests (4D)
Aqueous Extraction (4D2)
  Single-Point Extraction (4D2a)
  1:500, 12-h (4D2a3)
    Ion Chromatograph (4D2a3a)
      Conductivity Detector (4D2a3a1)
        Self-Generation Suppressor (4D2a3a1a)
          Sulfate (4D2a3a1a1)
            Air-Dry or Field-Moist, <2 mm (4D2a3a1a1a-b1)

1. Application
   The content of water-soluble sulfate is used in soil taxonomy as a required characteristic for the sulfuric horizon (Soil Survey Staff, 2014). The sulfuric horizon connotes the presence of active acid sulfate soils and has pH values (1:1 by weight in water) of 3.5 or less. The sulfuric horizon is commonly accompanied by concentrations of iron and sulfur-bearing minerals, such as jarosite $\text{KFe}_3(\text{SO}_4)_2(\text{OH})_6$. These minerals are alteration products of pyrite that have been exposed to an oxidizing environment. Such minerals may be absent from some sulfuric horizons. The presence of 0.05 percent or more water-soluble sulfate is used as an additional criterion for evidence of a sulfuric horizon when pH ≤3.5 is caused by active acid sulfate effects and jarosite or similar minerals is not present (Soil Survey Staff, 2014).

2. Summary of Method
   The 1:500 soil extract is diluted according to its electrical conductivity. The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to measure the sulfate content. Standard anion concentrations are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. A computer program automates these actions. Data are reported as percent water soluble $\text{SO}_4^{2-}$ by method 4D2a3a1a1.

3. Interferences
   Some soil extracts contain suspended solids. Filtering after dilution removes the particles. Organic anions that have low molecular weight will co-elute with inorganic anions from the column.

4. Safety
   Wear protective clothing and safety glasses. Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if
ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.

5. Equipment

5.1 Bottles with caps, 500-mL, Nalgene
5.2 Syringe filters, 0.45-µm, Whatman
5.3 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1 ½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Ion chromatograph, double-column, conductivity detection, Dionex CS200, Dionex Corp., Sunnyvale, CA
5.5 Guard column, IonPac AG-18, 4 x 50 mm, Dionex Corp., Sunnyvale, CA
5.6 Analytical column, IonPac AS-18, Dionex Corp., Sunnyvale, CA
5.7 Self-regeneration suppressor, ASRS–300, 4 mm, Dionex Corp., Sunnyvale, CA
5.8 Autosampler, AS40, Dionex Corp., Sunnyvale, CA
5.9 Computer with PeakNet software and printer
5.10 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.11 Poly-vials with caps, 5 mL, Dionex Corp., Sunnyvale, CA

6. Reagents

6.1 Reverse osmosis deionized filtered (RODI) water, ASTM Type I grade of reagent water
6.2 Helium gas
6.3 Primary stock standards solutions, \((PSSS_{1000})\), high purity, 1000 mg L⁻¹: Cl⁻, \(SO_4^{2-}\), F⁻, NO₃⁻, NO₂⁻, Br⁻, and PO₄³⁻.
6.4 Mixed calibration standards solutions (MCSS), A, B, C, and D and Blank as follows:

6.4.1 MCSSA. In a 500-mL volumetric flask, add as follows:

- 6.4.1.1 32 mL Cl⁻ PSSS₁₀₀₀ = 64 mg L⁻¹
- 6.4.1.2 32 mL SO₄²⁻ PSSS₁₀₀₀ = 64 mg L⁻¹
- 6.4.1.3 2 mL F⁻ PSSS₁₀₀₀ = 4 mg L⁻¹
- 6.4.1.4 8 mL NO₃⁻ PSSS₁₀₀₀ = 16 mg L⁻¹
- 6.4.1.5 2 mL NO₂⁻ PSSS₁₀₀₀ = 4 mg L⁻¹
- 6.4.1.6 2 mL Br⁻ PSSS₁₀₀₀ = 4 mg L⁻¹
- 6.4.1.7 2 mL PO₄³⁻ PSSS₁₀₀₀ = 4 mg L⁻¹

Dilute to volume with RODI water and invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.
6.4.2 MCSSB: In a 100-mL volumetric flask, add 50 mL MCSSA and dilute to volume with RODI water. Final concentrations are 32, 32, 2, 8, 2, and 2 mg L\(^{-1}\) Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.3 MCSSC: In a 100-mL volumetric flask, add 50 mL MCSSB and dilute to volume with RODI water. Final concentrations are 16, 16, 1, 4, 1, 1, and 1 mg L\(^{-1}\) Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.4 MCSSD: In a 100-mL volumetric flask, add 50 mL MCSSC and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 8, 8, 0.5, 2, 0.5, 0.5, and 0.5 mg L\(^{-1}\) Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\), respectively. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.5 MCSS Blank: 0 mL of Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\). Dilute RODI water to volume.

6.5 Quality Control: In a 1-L volumetric flask, add 2.0 mL from each PSSS\(_{1000}\) Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\) and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 2 mg L\(^{-1}\) Cl\(^{-}\), SO\(_4^{2-}\), F\(^{-}\), NO\(_3^{-}\), NO\(_2^{-}\), Br\(^{-}\), and PO\(_4^{3-}\). Store in plastic containers in the refrigerator. Prepare fresh weekly.

7. Procedure

Sample Extraction

7.1 Weigh 1.0 g of <2-mm or fine-grind, air-dry soil to the nearest mg, place into a 500-mL bottle, and add 500 mL RODI water. If sample is moist, weigh enough to achieve ≈1.0 g of air-dry soil.

7.2 Transfer the sample to a shaker. Shake for 12 h at 100 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.3 Add 50 mL of filtered sample extract to 50-mL Falcon tube.

7.4 Measure and record EC of sample extract.

Dilution of Sample Extracts

7.5 Dilute the soil extract with the RODI water as follows:
### Set-up and Operation of Ion Chromatograph (IC)

**7.7** Refer to the manufacturer’s manual for the operation of chromatograph. Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex ICS-2000 ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. Ranges and/or typical settings are as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range and/or (Typical Setting)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calibration</td>
<td>Peak height or (area)</td>
</tr>
<tr>
<td>Flow setting</td>
<td>0.5 to 4.5 mL min⁻¹ (1.00 mL min⁻¹)</td>
</tr>
<tr>
<td>Pressure</td>
<td>200 to 3000 psi (2200 to 2400 psi)</td>
</tr>
<tr>
<td>Detection</td>
<td>Suppressed conductivity</td>
</tr>
<tr>
<td>Total conductivity</td>
<td>0 to 999.9 μS</td>
</tr>
<tr>
<td>Injection volume</td>
<td>10 μL</td>
</tr>
</tbody>
</table>

**7.6** Place the MCSS (A, B, C, D, and Blank) and diluted extract samples in the Poly-vials and cap with filter caps.
Parameter | Range and/or (Typical Setting)
--- | ---
Auto offset | -999.9 to 999.9 μS (On)
Cell temperature | 35 ºC
Suppressor current | 75 mA

7.8 Load the sample holder cassettes with the capped samples, standards, and check samples.

7.9 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

**IC Calibration and Analysis**

7.10 Calibrate the instrument by using the MCSS (A, B, C, D, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.11 If samples are outside calibration, dilute sample extracts with RODI water solution and re-analyze.

7.12 Perform one quality control (QC) (Low Standard MCSS, Standard C) for every 12 samples. If reading is not within tolerance limits (10 to 15%, based on analyte), the instrument is re-calibrated and QC re-analyzed.

7.13 Record analyte readings to 0.01 mg L$^{-1}$.

8. **Calculations**

The instrument readings for analyte concentration are in mg L$^{-1}$ and are converted to percent sulfate in the soil as follows:

\[
\text{Sulfate Concentration in Soil (\%) = } \left\{ \frac{(A \times B \times C)}{D \times E} \right\} \times 100
\]

where:

- $A =$ Analyte ($\text{SO}_4^{2-}$) concentration in extract (mg L$^{-1}$)
- $B =$ Sample extract volume (0.5 L)
- $C =$ Dilution ratio, if needed
- $D =$ Conversion factor (1000)
- $E =$ Sample weight (1 g)

9. **Report**

Report the water soluble $\text{SO}_4^{2-}$ to the nearest hundredths of a percent.

10. **Precision and Accuracy**

Precision and accuracy data are available from the KSSL upon request.
11. References

Soil Test Analyses (4D)
Bray P-1 Extraction (4D3)
   UV-Visible Spectrophotometer, Dual-Beam (4D3a)
   Phosphorus (4D3a1)
      Air-Dry or Field-Moist, <2 mm (4D3a1a-b1)

1. Application
The Bray P-1 procedure is widely used as an index of available P in the soil. Bray and Kurtz (1945) originally designed the Bray P-1 extractant to selectively remove a portion of the adsorbed form of P with the weak, acidified ammonium fluoride solution. Adsorbed phosphorus is the anion form adsorbed by different charged surface functional groups that have varying degrees of adsorption affinity. In general, this method has been most successful on acid soils (Olsen and Sommers, 1982). The acid solubilizes calcium and aluminum phosphates and partially extracts iron phosphates compounds. The NH₄F complexes the aluminum in solution and limits reabsorption of P on iron oxides (Kuo, 1996). The Bray P-1 has limited ability to extract P in calcareous soils due to the neutralization of the dilute acid by carbonates.

2. Summary of Method
A 2.5-g soil sample is shaken with 25 mL of Bray P-1 extracting solution for 15 min. The sample is centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained. A 2-mL aliquot is diluted with 8-mL of ascorbic acid molybdate solution. Absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg⁻¹ soil (4D3a1).

3. Interferences
Many procedures may be used to determine P. Studies have shown that incomplete or excessive extraction of P to be the most significant contributor to interlaboratory variation. The Bray P-1 procedure is sensitive to the soil/extractant ratio, shaking rate, and time. This extraction uses the ascorbic acid-potassium antimony-tartrate-molybdate method. The Fiske-Subbarrow method is less sensitive but has a wider range before dilution is required (North Central Regional Publication No. 221, 1988). For calcareous soils, the Olsen method is preferred. An alternative procedure for calcareous soils is to use the Bray P-1 extracting solution at a 1:50 ratio of soil to solution. This procedure has been shown to be satisfactory for some calcareous soils (Smith et al., 1957; North Central Regional Publication No. 221, 1988).
4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H₂SO₄ and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±0.10-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, polyethylene
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, ashless, 9-cm diameter
5.6 Centrifuge, Centra GP-8, Thermo IEC, Needham Heights, MA
5.7 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL
5.8 Cups, plastic
5.9 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.10 Dispenser, 30-mL or 10-mL
5.11 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.12 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated, 12 N, trace pure grade
6.3 HCl, 1 N. Carefully add 83.33 mL of concentrated HCl to RODI water and dilute to 1-L volume.
6.4 Sulfuric acid (H₂SO₄), concentrated, 36 N
6.5 Bray No. 1 extracting solution, 0.025 N HCl and 0.03 N NH₄F. Dissolve 8.88 g of NH₄F in 4 L RODI water. Add 200 mL of 1.0 N HCl and dilute to 8 L with RODI water. The solution pH should be 2.6 ±0.5. Store in a polyethylene bottle.
6.6 Sulfuric-tartrate-molybdate solution (STMS). Dissolve 60 g of ammonium molybdate tetrahydrate [(NH₄)₆Mo₇O₂₄•4H₂O] in 200 mL of boiling RODI water.
water. Allow to cool to room temperature. Dissolve 1.455 g of antimony potassium tartrate (potassium antimony tartrate hemihydrate [K(SbO)$_4$C$_4$H$_4$O$_6$•½H$_2$O]) in the ammonium molybdate solution. Slowly and carefully add 700 mL of concentrated H$_2$SO$_4$. Cool and dilute to 1 L with RODI water. Store in the dark in the refrigerator.

6.7 Ascorbic acid solution. Dissolve 6.6 g of ascorbic acid in RODI water and dilute to 50 mL with RODI water. Make fresh daily.

6.8 Working ascorbic acid molybdate solution (WAMS). Mix 25 mL of STSM solution with 800 mL of RODI water. Add 10 mL of ascorbic acid solution and dilute to 1 L with RODI water. Allow to stand at least 1 h before using. Prepare fresh daily.

6.9 Working stock standard P solution (WSSPS), 100.0 mg P L$^{-1}$. In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate (KH$_2$PO$_4$) (dried for 2 h at 110 °C) in about 800 mL extracting solution. Dilute to 1-L volume with extracting solution water and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.10 Standard P calibration solutions (SPCS), or working standards, 5.0, 4.0, 2.0, 0.8, 0.4, and 0.0 mg P L$^{-1}$. Make fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. To six 250-mL volumetric flasks, add as follows:

6.10.1 5.0 mg P L$^{-1}$ = 12.5 mL WSSPS
6.10.2 4.0 mg P L$^{-1}$ = 10.0 mL WSSPS
6.10.3 2.0 mg P L$^{-1}$ = 5.0 mL WSSPS
6.10.4 0.8 mg P L$^{-1}$ = 2.0 mL WSSPS
6.10.5 0.4 mg P L$^{-1}$ = 1.0 mL WSSPS
6.10.6 0.0 mg P L$^{-1}$ = 0 mL WSSPS (blank)

Dilute each SPCS to the mark with extracting solution and invert to thoroughly mix.

6.11 Quality Control Samples: 0.1 mg L$^{-1}$ solution made from SSPS; blanks; and selected SPCS. In addition, KSSL soil standard and WEPAL ISE's (Wageningen Evaluating Programmes for Analytical Laboratories, International Soil Exchange) from the Netherlands are routinely included in a batch for quality control.

7. Procedure

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Dispense 25.0 mL of extracting solution to tube.
7.3 Transfer the sample in the shaker. Shake for 15 min at 200 oscillations min\(^{-1}\) at room temperature (20 °C).

7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.5 Use the pipette to transfer a 2-mL aliquot of the sample to a plastic cup. Also transfer a 2-mL aliquot of each SPCS to a plastic cup. Use a clean pipette tip for each sample and SPCS.

7.6 Dispense 8 mL of the WAMS to sample aliquot and to each SPCS. Swirl to mix. The color reaction requires a minimum of 20 min before analyst records readings.

7.7 Transfer sample extract and SPCS to cuvettes.

7.8 Set the spectrophotometer to read at 882 nm. Autozero with calibration blank.

7.9 Calibrate the instrument using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is \(R^2 < 0.99\).

7.10 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; if KSSL standard varies by more than 20% from the accepted mean; or if ISE \(>(3 \times \text{MAD})\), where MAD=median of absolute deviations. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.11 If samples are outside the calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations
   Convert the extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:
   
   \[
   \text{Soil P (mg kg}^{-1}) = (A \times B \times C \times R \times 1000)/E
   \]

   where:
   
   \(A\) = Sample extract reading (mg L\(^{-1}\))

   \(B\) = Extract volume (L)

   \(C\) = Dilution, if performed

   \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

   1000 = Conversion factor to kg-basis

   \(E\) = Sample weight (g)
9. Report

Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


North Central Regional Publication No. 221. 1988. Recommended chemical soil test procedures for the North Central region. Agric. Exp. Stn. of IL, IN, IA, KS, MI, MN, MS, NE, ND, OH, SD, and WI and USDA cooperating.


Soil Test Analyses (4D)

Bray P-2 Extraction (4D4)

UV-Visible Spectrophotometer, Dual-Beam (4D4a)

Phosphorus (4D4a1)

Air-Dry or Field-Moist, <2 mm (4D4a1a-b1)

1. Application

The Bray P-2 procedure functions to extract a portion of the plant available P in the soil. It has a similar composition to Bray P-1 extraction solution. The difference is a slightly higher concentration of HCl (0.025 \(N\) to 0.1 \(N\)) in the Bray P-2. It was originally designated by Bray and Kurtz (1945) to extract the easily acid soluble P as well as a fraction of adsorbed phosphates. The HCl solubilizes calcium and aluminum phosphates and partially extracts iron phosphate compounds. The \(\text{NH}_4\text{F}\) complexes the aluminum in solution and limits re-adsorption of P on iron oxides (Kuo, 1996). The higher acid concentration of the Bray P-2 should allow greater extraction of P in calcareous soils compared to Bray P-1. Bray P-2 is not as widely used by soil testing laboratories as Bray P-1.
2. Summary of Method

A 2.5-g soil sample is shaken with 25 mL of Bray P-2 extracting solution for 15 min. The sample is centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained. A 2-mL aliquot is diluted with 8-mL of ascorbic acid molybdate solution. Absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg\(^{-1}\) soil (4D4a1).

3. Interferences

Many procedures may be used to determine P. Studies have shown that incomplete or excessive extraction of P to be the most significant contributor to interlaboratory variation. The Bray procedure is sensitive to the soil/extractant ratio, shaking rate, and time. This extraction uses the ascorbic acid-potassium antimony-tartrate-molybdate method. The Fiske-Subbarrow method is less sensitive but has a wider range before dilution is required (North Central Regional Publication No. 221, 1988). For calcareous soils, the Olsen method is preferred. An alternative procedure for calcareous soils is to use the Bray P-1 extracting solution at a 1:50 ratio of soil to solution. This procedure has been shown to be satisfactory for some calcareous soils (Smith et al., 1957; North Central Regional Publication No. 221, 1988). The higher acid concentration of the Bray P-2 should allow greater extraction of P in calcareous soils compared to Bray P-1. Bray P-2 is not as widely used by soil testing laboratories as Bray P-1.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H\(_2\)SO\(_4\) and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±0.10-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min\(^{-1}\), 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, polyethylene
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, ashless, 9-cm diameter
5.6 Centrifuge, Centra GP-8, Thermo IEC, Needham Heights, MA
5.7 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL
5.8 Cups, plastic
5.9 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.10 Dispenser, 30-mL or 10-mL
5.11 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.12 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated, 12 N, trace pure grade
6.3 HCl, 1 N. Carefully add 83.33 mL of concentrated HCl to RODI water and dilute to 1-L volume.
6.4 Sulfuric acid (H₂SO₄), concentrated, 36 N
6.5 Bray No. 2 Extracting solution, 0.1 N HCl and 0.03 N NH₄F. Dissolve 8.88 g of NH₄F in 4 L RODI water. Add 800 mL of 1.0 N HCl and dilute to 8 L with RODI water. Store in a polyethylene bottle.
6.6 Sulfuric-tartrate-molybdate solution (STMS). Dissolve 60 g of ammonium molybdate tetrahydrate [(NH₄)₆Mo₇O₂₄•4H₂O] in 200 mL of boiling RODI water. Allow to cool to room temperature. Dissolve 1.455 g of antimony potassium tartrate (potassium antimony tartrate hemihydrate [K(SbO)(C₄H₄O₆)•½H₂O]) in the ammonium molybdate solution. Slowly and carefully add 700 mL of concentrated H₂SO₄. Cool and dilute to 1 L with RODI water. Store in the dark in the refrigerator.
6.7 Ascorbic acid solution. Dissolve 6.6 g of ascorbic acid in RODI water and dilute to 50 mL with RODI water. Make fresh daily.
6.8 Working ascorbic acid molybdate solution (WAMS). Mix 25 mL of STMS solution with 800 mL of RODI water. Add 10 mL of ascorbic acid solution and dilute to 1 L with RODI water. Allow to stand at least 1 h before using. Prepare fresh daily.
6.9 Working stock standard P solution (WSSPS), 100.0 mg P L⁻¹. In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate (KH₂PO₄) (dried for 2 h at 110 °C) in about 800 mL extracting solution. Dilute to 1-L volume with extracting solution water and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.
6.10 Standard P calibration solutions (SPCS), or working standards, 5.0, 4.0, 2.0, 0.8, 0.4, 0.0 mg P L⁻¹. Make fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. To six 250-mL volumetric flasks, add as follows:
6.10.1  5.0 mg P L\(^{-1}\) = 12.5 mL WSSPS
6.10.2  4.0 mg P L\(^{-1}\) = 10.0 mL WSSPS
6.10.3  2.0 mg P L\(^{-1}\) = 5.0 mL WSSPS
6.10.4  0.8 mg P L\(^{-1}\) = 2.0 mL WSSPS
6.10.5  0.4 mg P L\(^{-1}\) = 1.0 mL WSSPS
6.10.6  0.0 mg P L\(^{-1}\) = 0 mL WSSPS (blank)

Dilute each SPCS to the mark with extracting solution and invert to thoroughly mix.

6.11 Quality Control Samples: 0.1 mg L\(^{-1}\) solution made from SSPS; blanks; and selected SPCS. In addition, KSSL soil standard and WEPAL ISE's (Wageningen Evaluating Programmes for Analytical Laboratories, International Soil Exchange) from the Netherlands are routinely included in a batch for quality control.

7. Procedure
7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.
7.2 Dispense 25.0 mL of extracting solution to tube.
7.3 Transfer the sample to the shaker. Shake for 15 min at 200 oscillations min\(^{-1}\) at room temperature (20 °C).
7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
7.5 Use the pipette to transfer a 2-mL aliquot of the sample to a plastic cup. Also transfer a 2-mL aliquot of each SPCS to a plastic cup. Use a clean pipette tip for each sample and SPCS.
7.6 Dispense 8 mL of the WAMS to sample aliquot and to each SPCS. Swirl to mix. The color reaction requires a minimum of 20 min before analyst records readings.
7.7 Transfer sample extract and SPCS to cuvettes.
7.8 Set the spectrophotometer to read at 882 nm. Autozero with calibration blank.
7.9 Calibrate the instrument using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is \(R^2 < 0.99\).
7.10 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; if
KSSL standard varies by more than 20% from the accepted mean; or if ISE > (3 x MAD), where MAD = median of absolute deviations. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.11 If samples are outside the calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations

Convert the extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:

\[
\text{Soil P (mg kg}^{-1}\) = (A \times B \times C \times R \times 1000) / E
\]

where:

- A = Sample extract reading (mg L\(^{-1}\))
- B = Extract volume (L)
- C = Dilution, if performed
- R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- E = Sample weight (g)

9. Report

Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


1. Application

Olsen extractant is 0.5 M sodium bicarbonate solution at pH 8.5. This extractant is most applicable to neutral to calcareous soils (Buurman et al., 1996). Solubility of Ca-phosphate in calcareous, alkaline, or neutral soils in increased because of the precipitation of Ca\(^{++}\) as CaCO\(_3\) (Soil and Plant Analysis Council, 1999). Olsen extractant correlates with Mehlich No. 3 on calcareous soils (R\(^2\) = 0.918), even though the quantity of Mehlich No. 3 extractable P is considerably higher (Soil and Plant Analysis Council, 1999). While Mehlich No. 3, Bray P-1, and Olsen sodium-bicarbonate are linearly related, relationships developed between some P tests (e.g., Olsen P, Mehlich No. 3) may have limited predictive capability with increasing soil P content (Burt et al., 2002).

2. Summary of Method

A 1.0-g soil sample is shaken with 20 mL of Olsen sodium-bicarbonate extracting solution for 30 min. The sample is centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained. Dilute 5-mL of sample extract with 5-mL of color reagent. The absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg\(^{-1}\) soil (4D5a1).

3. Interferences

The Mo blue methods, which are very sensitive for P, are based on the principle that in an acid molybdate solution containing orthophosphate ions, a phosphomolybdate complex forms that can be reduced by ascorbic acid, SnCl\(_2\), and other reducing agents to a Mo color. The intensity of blue varies with the P concentration but is also affected by other factors, such as acidity, arsenates, silicates, and substances that influence the oxidation-reduction conditions of the system (Olsen and Sommers, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H\(_2\)SO\(_4\) and HCl to a fume hood. Use safety showers and
eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1 ½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tube, 50-mL, polyethylene
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, 150-mm
5.6 Centrifuge, Centra, GP-8, Thermo IEC, Thermo IEC, Needham Heights, MA
5.7 Pipettes, electronic digital, 1000 µL and 10 mL, with tips, 1000 µL and 10 mL
5.8 Cups, plastic
5.9 Dispenser, 30-mL or 10-mL
5.10 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.11 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.12 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Sulfuric acid (H₂SO₄), concentrated, 36 N, trace pure grade
6.3 Sulfuric acid (H₂SO₄), 4 M. In a 250-mL volumetric, carefully add 56 mL concentrated H₂SO₄ to 150 mL RODI water. Allow to cool. Make to final volume with RODI water. Invert to thoroughly mix.
6.4 NaOH, 1 M. Dissolve 4 g NaOH in 100 mL RODI water.
6.5 Olsen Sodium Bicarbonate Extracting solution (0.5 M NaHCO₃). In a 6-L container, dissolve 252 g NaHCO₃ in RODI water. Adjust the pH to 8.5 with 1 M NaOH. Make to final volume. Mix thoroughly. Check pH every day.
6.6 Ammonium molybdate, 4%. Dissolve 4 g of [(NH₄)₆Mo₇O₂₄•4H₂O] in 100-mL volumetric with RODI water. Dilute to volume with RODI water. Store in the dark in the refrigerator.
6.7 Potassium antimony–(III) oxide tartrate, 0.275%. Dissolve 0.275 g [K(SbO)C₇H₆O₆•½H₂O] in 100 mL RODI water.
6.8 Ascorbic acid, 1.75%. Dissolve 1.75 g ascorbic acid in 100 mL RODI water. Prepare fresh daily.
6.9 Color developing reagent. In a 500-mL bottle, add 50 mL 4 M \( \text{H}_2\text{SO}_4 \), 15 mL 4% ammonium molybdate, 30 mL 1.75% ascorbic acid, 5 mL 0.275% potassium antimony–(III) oxide tartrate, and 200 mL RODI water. Mix well after each addition. Prepare fresh daily.

6.10 Stock standard P solution (SSPS), 100.0 mg P L\(^{-1}\). In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate (KH\(_2\)PO\(_4\)) (dried for 2 h at 110 °C) in about 800 mL extracting solution. Dilute to 1-L volume with extracting solution and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.11 Working stock standard P solution (WSSPS), 4.0 mg P L\(^{-1}\). Pipette 10 mL of 100 mg P L\(^{-1}\) SSPS to 250-mL volumetric flask. Dilute to 250-mL volume with extracting solution and invert to thoroughly mix. Make fresh weekly. Store in the refrigerator.

6.12 Standard P calibration solution (SPCS), or working standards, 2.0, 1.6, 1.2, 0.8, 0.4, and 0.0 mg P L\(^{-1}\). Make fresh weekly. Store in the refrigerator. Allow to equilibrate to room temperature before use. To six 50-mL volumetric flasks, add as follows:

6.12.1 2.0 mg P L\(^{-1}\)=25 mL WSSPS
6.12.2 1.6 mg P L\(^{-1}\)=20 mL WSSPS
6.12.3 1.2 mg P L\(^{-1}\)=15 mL WSSPS
6.12.4 0.8 mg P L\(^{-1}\)=10 mL WSSPS
6.12.5 0.4 mg P L\(^{-1}\)=5 mL WSSPS
6.12.6 0.0 mg P L\(^{-1}\)=0 mL WSSPS (blank)

Dilute each SPCS to mark with extracting solution and invert to thoroughly mix.

6.13 Quality Control Samples: 0.1 mg P L\(^{-1}\) solution made from SSPS; blanks; and selected SPCS. In addition, KSSL soil standard and WEPAL ISE’s (Wageningen Evaluating Programmes for Analytical Laboratories, International Soil Exchange) from the Netherlands are routinely included in a batch for quality control.

7. Procedure

7.1 Weigh 1.0 g of <2 mm or fine-grind, air-dry soil to the nearest mg into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈1.0 g of air-dry soil.

7.2 Dispense 20.0 mL of extracting solution to tube.

7.3 Transfer the sample to the shaker. Shake for 30 min at 200 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).
7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.5 Use the pipette to transfer a 5-mL aliquot of the sample to a plastic cup. Also transfer a 5-mL aliquot of each SPCS to a plastic cup. Use a clean pipette tip for each sample and SPCS.

7.6 Dispense 5 mL of color developing reagent to sample aliquot and to each SPCS. Swirl to mix. Do not place sample cups close together as carbon dioxide is released and solution will bubble. The color reaction requires a minimum of 20 min before analyst records readings. Allowing 1 h for color development usually improves results. Color will remain stable for 24 h.

7.7 Transfer sample extract and SPCS to cuvettes.

7.8 Set the spectrophotometer to read at 882 nm. Autozero with calibration blank.

7.9 Calibrate the instrument using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is $R^2 < 0.99$.

7.10 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are $>0.01$; if SPCS vary by more than 20% from calculated value; if KSSL standard varies by more than 20% from the accepted mean; or if ISE $(3 \times \text{MAD})$, where MAD = median of absolute deviations. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.11 If samples are outside calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations

8.1 Convert the extract P (mg L$^{-1}$) to soil P (mg kg$^{-1}$) as follows:

Soil P (mg kg$^{-1}$) = \( \frac{A \times B \times C \times R \times 1000}{E} \)

where:

- $A$ = Sample extract reading (mg L$^{-1}$)
- $B$ = Extract volume (L)
- $C$ = Dilution, if performed
- $R$ = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- $E$ = Sample weight (g)
9. Report

Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Soil Test Analyses (4D)
Mehlich No. 3 Extraction (4D6)
  UV-Visible Spectrophotometer, Dual-Beam (4D6a)
  Phosphorus (4D6a1)
    Air-Dry or Field-Moist, <2 mm (4D6a1a-b1)

1. Application

Mehlich No. 3 was developed by Mehlich (1984) as a multielement soil extraction (Ca, Mg, K, Na, P). In the Mehlich No. 3 procedure, P is extracted by reaction with acetic acid and F compounds. Mehlich No. 3 is used as an index of available P in the soil. Extraction of P by Mehlich No. 3 is designed to be applicable across a wide range of soil properties with reaction ranging from acid to basic (Mehlich, 1984). Mehlich No. 3 correlates well with Bray P-1 on acid to neutral (\(R^2=0.966\)) soils but does not correlate with Bray P-1 on calcareous soils (Soil and Plant Analysis Council, 1999). Mehlich No. 3 correlates with Olsen extractant on calcareous soils (\(R^2=0.918\)), even though the quantity of Mehlich No. 3 extractable P is considerably higher (Soil and Plant Analysis Council, 1999). The Mehlich No. 3 extractant is neutralized less by carbonate compounds in soil than the double acid (Mehlich No. 1) and the Bray P-1 extractants and is less aggressive towards apatite or other Ca-phosphate than the double acid and Bray P-2 extractants (Tran and Simard, 1993). Mehlich No. 3 can also be used to
extract Ca, Mg, K, and Na in a wide range of soils and correlates well with Mehlich No. 1, Mehlich No. 2, and NH₄OAc (Soil and Plant Analysis Council, 1999).

2. Summary of Method

A 2.5-g soil sample is shaken with 25 mL of Mehlich No. 3 extracting solution for 5 min. The sample is centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained. Dilute 0.5-mL of sample extract with 13.5-mL of working solution. Absorbance of the solution is read using a spectrophotometer at 882 nm. Data are reported as mg P kg⁻¹ soil (4D6a1).

3. Interferences

The Mo blue methods, which are very sensitive for P, are based on the principle that in an acid molybdate solution containing orthophosphate ions, a phosphomolybdate complex forms that can be reduced by ascorbic acid, SnCl₂, and other reducing agents to a Mo color. Ascorbic acid is the reducing agent most commonly used by agricultural laboratories (Murphy and Riley, 1962). The intensity of blue varies with the P concentration but is also affected by other factors, such as acidity, arsenates, silicates, and substances that influence the oxidation-reduction conditions of the system (Olsen and Sommers, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H₂SO₄ and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, polyethylene
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, 150-mm
5.6 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.7 Pipettes, electronic digital, 1000-µL and 10-mL, with tips, 1000-µL and 10-mL
5.8 Dispenser, 30-mL or 10-mL
5.9 Cups, plastic
5.10 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.11 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.12 Computer with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Sulfuric acid ($\text{H}_2\text{SO}_4$), concentrated, 36 N, trace pure grade
6.3 Mehlich No. 3 Stock Solution: Add 55.56 g NH₄F to 600 mL RODI water, followed by 29.23 g EDTA (FW 292.24), and bring to 1-L volume with RODI water.
6.4 Mehlich No. 3 Working Solution: Add 200.1 g NH₄NO₃, 100 mL Mehlich No. 3 Stock Solution, 115 mL CH₃COOH, and 82 mL 10% v/v HNO₃ (10 mL concentrated 70% HNO₃ in 100 mL RODI water). Bring to 10-L volume with RODI.
6.5 Sulfuric-tartrate-molybdate solution (STMS). Dissolve 100 g of ammonium molybdate tetrahydrate ($\left(\text{NH}_4\right)_6\text{Mo}_7\text{O}_{24}\cdot4\text{H}_2\text{O}$) in 500 mL of RODI water. Dissolve 2.425 g of antimony potassium tartrate (potassium antimony tartrate hemihydrate $\left[\text{K(SbO)}\text{C}_4\text{H}_4\text{O}_6\cdot\frac{1}{2}\text{H}_2\text{O}\right]$) in the ammonium molybdate solution. Slowly and carefully add 1400 mL of concentrated $\text{H}_2\text{SO}_4$ and mix well. Cool and dilute to 2 L with RODI water. Store in the dark in the refrigerator.
6.6 Ascorbic acid solution. Dissolve 8.8 g of ascorbic acid in RODI water and dilute to 100-mL with RODI water. Make fresh daily.
6.7 Working ascorbic acid molybdate solution (WAMS). Dilute 20 mL of STMS solution and 10 mL of the ascorbic acid solution with RODI water to make 1 L. Allow solution to come to room temperature before using. Prepare fresh daily.
6.8 Working stock standard P solution (WSSPS), 100.0 mg P L⁻¹. In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate ($\text{KH}_2\text{PO}_4$) (dried for 2 h at 110 °C) in about 800 mL extracting solution. Dilute to 1-L volume with extracting solution and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.
6.9 Standard P calibration solutions (SPCS), or working standards, 12.0, 10.0, 8.0, 4.0, 2.0, 1.0, 0.8, 0.4 and 0.0 mg P L⁻¹. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. In nine 250-mL volumetric flasks, add as follows:
6.9.1 12.0 mg P L⁻¹ $= 30$ mL WSSPS
6.9.2 10.0 mg P L⁻¹ $= 25$ mL WSSPS
6.9.3  8.0 mg P L\(^{-1}\)=20 mL WSSPS  
6.9.4  4.0 mg P L\(^{-1}\)=10 mL WSSPS  
6.9.5  2.0 mg P L\(^{-1}\)=5 mL WSSPS  
6.9.6  1.0 mg P L\(^{-1}\)=2.5 mL WSSPS  
6.9.7  0.8 mg P L\(^{-1}\)=2 mL WSSPS  
6.9.8  0.4 mg P L\(^{-1}\)=1 mL WSSPS  
6.9.9  0.0 mg P L\(^{-1}\)=0 mL WSSPS (blank)

Dilute each SPCS to the mark with extracting solution and invert to thoroughly mix.

6.10 Quality Control Samples: 0.1 mg P L\(^{-1}\) solution made from SSPS; blanks; and selected SPCS. In addition, KSSL soil standard and WEPAL ISE’s (Wageningen Evaluating Programmes for Analytical Laboratories, International Soil Exchange) from the Netherlands are routinely included in a batch for quality control.

7. Procedure

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to the nearest mg into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Dispense 25.0 mL of extracting solution to the tube.

7.3 Transfer the sample to the shaker. Shake for 5 min at 200 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cups. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.5 Use the pipette to transfer a 1 mL (or 0.5-mL) aliquot of the sample to a plastic cup. Also transfer a 1 mL (or 0.5-mL) aliquot of each SPCS to a plastic cup. Use a clean pipette tip for each sample and SPCS.

7.6 Dispense 27 mL (or 13.5 mL) of the WAMS to sample aliquot and to each SPCS. Swirl to mix. The color reaction requires a minimum of 20 min before analyst records readings. Allowing 1 h for color development usually improves results. Color will remain stable for 6 h.

7.7 Transfer sample extract and SPCS to cuvettes.

7.8 Set the spectrophotometer to read at 882 nm. Autozero with calibration blank.

7.9 Calibrate the instrument by using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is \(R^2 < 0.99\).
7.10 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; if KSSL standard varies by more than 20% from the accepted mean; or if ISE >\((3 \times \text{MAD})\), where MAD = median of absolute deviations. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.11 If samples are outside calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculation

8.1 Convert extract P \(\text{mg L}^{-1}\) to soil P \(\text{mg kg}^{-1}\) as follows:

\[
\text{Soil P \(\text{mg kg}^{-1}\)} = \left(\frac{A \times B \times C \times R \times 1000}{E}\right)
\]

where:
- \(A\) = Sample extract reading \(\text{mg L}^{-1}\)
- \(B\) = Extract volume (L)
- \(C\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- \(E\) = Sample weight (g)

9. Report

Report data to the nearest 0.1 mg P kg \(^{-1}\) soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Test Analyses (4D)

Mehlich No. 3 Extraction (4D6)

Inductively Coupled Plasma Atomic Emission Spectrophotometer (4D6b)

Axial Mode (4D6b1)

Cross-Flow Nebulizer (4D6b1b)

Aluminum, Arsenic, Barium, Calcium, Cadmium, Cobalt, Chromium, Copper, Iron, Potassium, Magnesium, Manganese, Sodium, Nickel, Phosphorus, Lead, Selenium, and Zinc (4D6b1b1-18)

Air-Dry or Field-Moist, >2 mm (4D6b1b1-18a-b1)

1. Application

Mehlich No. 3 was developed by Mehlich (1984) as a multielement soil extraction (Ca, Mg, K, Na, P). In the Mehlich No. 3 procedure, P is extracted by reaction with acetic acid and F compounds. Mehlich No. 3 is used as an index of available P in the soil. Extraction of P by Mehlich No. 3 is designed to be applicable across a wide range of soil properties with reaction ranging from acid to basic (Mehlich, 1984). Mehlich No. 3 correlates well with Bray P-1 on acid to neutral (R^2 = 0.966) soils but does not correlate with Bray P-1 on calcareous soils (Soil and Plant Analysis Council, 1999). Mehlich No. 3 correlates with Olsen extractant on calcareous soils (R^2 = 0.918), even though the quantity of Mehlich No. 3 extractable P is considerably higher (Soil and Plant Analysis Council, 1992). The Mehlich No. 3 extractant is neutralized less by carbonate compounds in soil than the double acid (Mehlich No. 1) and Bray P-1 extractants. It is also less aggressive towards apatite or other Ca-phosphate than the double acid and Bray P-2 extractants (Tran and Simard, 1993). Mehlich No. 3 can also be used to extract Ca, Mg, K, and Na in a wide range of soils and correlates well with Mehlich No. 1, Mehlich No. 2, and NH₄OAc (Soil and Plant Analysis Council, 1999). Additionally, Mehlich No. 3 can be used to extract Al, Cd, Cu, Fe, Mn, Ni, Pb, and Zn (Elrashidi et al., 2003).

2. Summary of Method

A 2.5-g soil sample is shaken with 25 mL of Mehlich No. 3 extracting solution for 5 min. The sample is centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained. Calibration standards are prepared for elemental analysis. A blank of Mehlich No. 3 is prepared. An inductively coupled plasma atomic emission spectrophotometer (ICP–AES) is used for analysis. The concentration of Al, As, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg,
Mn, Na, Ni, P, Pb, Se, and Zn are determined by an inductively coupled plasma atomic emission spectrophotometer (ICP–AES) in radial mode. Data by this method (4D6b1a1-18) are reported as mg kg\(^{-1}\) soil.

3. Interferences

Spectral and matrix interferences exist. Interferences are corrected or minimized by using both an internal standard and inter-elemental correction factors. Also, careful selection of specific wavelengths for data reporting is important. Background corrections are made by ICP–AES software. Samples and standards are matrix-matched to reduce interferences.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated acids to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronically balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min\(^{-1}\), 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, polyethylene
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter paper, Whatman 42, 150-mm
5.6 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.7 Pipettes, electronic digital, 1000-µL and 10-mL, with tips, 1000-µL and 10-mL
5.8 Dispenser, 30-mL or 10-mL
5.9 Cups, plastic
5.10 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES), Perkin-Elmer Optima 7300 Dual View (DV), Perkin-Elmer Corp., Norwalk, CT
5.11 Cross-flow nebulizer, Gem tip, Ryton double pass Scott type spray chamber, Perkin-Elmer Corp., Norwalk, CT
5.12 RF generator, floor mounted power unit, 45 MHz free running, Perkin-Elmer Corp., Norwalk, CT
5.13 Computer, with WinLab software ver. 4.1, Perkin-Elmer Corp., Norwalk, CT, and printer
5.14 Recirculating chiller, Neslab, CFT Series
5.15 Compressed gasses, Argon (minimum purity 99.996%) and Nitrogen (minimum purity 99.999%)
5.16 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.17 Quartz torch, Part No. N069-1662; alumina injector (2.0 mm id), Part No. N069-5362

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Mehlich No. 3 Extracting solution (0.2 \( N \) \( CH_3COOH \); 0.25 \( N \) \( NH_4NO_3 \); 0.015 \( N \) \( NH_4F \); 0.13 \( N \) \( HNO_3 \); 0.001 \( M \) EDTA). Premixed Mehlich No. 3 Extractant, Special-20, Hawk Creek Laboratory, Rural Route 1, Box 686, Simpson Road, Glen Rock, PA, 17327.
6.3 Primary standards, 1000 mg L\(^{-1}\): of Al, Fe, Ca, Mg, Na, K, Mn, P, As, Ba, Cd, Co, Cr, Cu, Ni, Pb, Se, and Zn, High Purity Standards, Charleston, SC
6.4 Mixed Standard High A, 100, 100, 100, 20, 40, 20, and 10 mg L\(^{-1}\) of Ca, Mg, Al, K, Fe, Na, Cr, respectively: To a 500-mL volumetric, add 50, 50, 50, 10, 20, 10, and 5 mL of the 1000 mg L\(^{-1}\) Ca, Mg, Al, K, Fe, and Na (Section 6.3) and dilute to volume with Mehlich No. 3 (Section 6.2) extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.
6.5 Mixed Standard Medium A, 50, 50, 50, 10, 20, 10, and 5 mg L\(^{-1}\) of Ca, Mg, Al, K, Fe, Na, and Cr, respectively: To a 100-mL volumetric, add 50 mL of Mixed Standard High A and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.
6.6 Mixed Standard Low A, 5, 5, 5, 1, 2, 1, and 0.5 mg L\(^{-1}\) of Ca, Mg, Al, K, Fe, Na, and Cr, respectively: To a 100-mL volumetric, add 10 mL of Mixed Standard Medium A and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.
6.7 Mixed Standard Very Low A, 0.5, 0.5, 0.5, 0.1, 0.2, 0.1 and 0.05 mg L\(^{-1}\) of Ca, Mg, Al, K, Fe, Na, and Cr, respectively: To a 100-mL volumetric, add 10 mL of Mixed Standard Low A and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.
6.8 Mixed Standard High B, 250, 10, 10, 10, 10, 10, 10, 10, 10, 10, 10 mg L\(^{-1}\) of P, As, Ba, Cd, Co, Cu, Mn, Ni, Se, Zn, and Pb, respectively: To a 100-mL volumetric, add 25, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1 mL of the 1000 mg L\(^{-1}\) of P, As, Ba, Cd, Co, Cu, Mn, Ni, Se, Zn, and Pb (Section 6.3) and dilute to volume with Mehlich No. 3 (Section 6.2) extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.

6.10 Mixed Standard Medium B, 25, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1 mg L\(^{-1}\) of P, As, Ba, Cd, Co, Cu, Mn, Ni, Pb, Se, and Zn, respectively: To a 100-mL volumetric, add 10 mL of Mixed Standard High B and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.

6.11 Mixed Standard Low B, 2.5, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1, 0.1 mg L\(^{-1}\) of P, As, Ba, Cd, Co, Cu, Mn, Ni, Pb, Se, and Zn, respectively: To a 100-mL volumetric, add 10 mL of Mixed Standard Medium B and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.

6.12 Mixed Standard Very Low B, 0.25, 0.01, 0.01, 0.01, 0.01, 0.01, 0.01, 0.01, 0.01, 0.01, 0.01 mg L\(^{-1}\) of P, As, Ba, Cd, Co, Cu, Mn, Ni, Pb, Se, and Zn, respectively: To a 100-mL volumetric, add 10 mL of Mixed Standard Low B and dilute to volume with Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.

6.13 Blanks: To a 100-mL volumetric, add Mehlich No. 3 extracting solution. Invert to thoroughly mix. Store in a polyethylene container in a refrigerator. Make fresh weekly.

7. Procedure

**Extraction of Al, As, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, Pb, Se, and Zn**

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to the nearest mg into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Dispense 25.0 mL of extracting solution to the tube.

7.3 Transfer the sample to the shaker. Shake for 5 min at 200 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cups. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
ICP–AES Set-up and Operation

7.5 Use the ICP–AES in axial mode to analyze elements. Use ultrasonic nebulization of sample. No initial dilutions of samples are necessary prior to analysis. Perform instrument checks (Hg alignment; BEC and %RSD of 1 mg L$^{-1}$ Mn solution) prior to analysis as discussed in operation manual of instrument. Check instrument alignment and gas pressures to obtain optimum readings with maximum signal to noise ratio. Analyses are generally performed at two or more wavelengths for each element. The selected wavelengths are as follows (reported wavelength listed first and in boldface):

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(nm)</td>
</tr>
<tr>
<td>Al</td>
<td>308.215, 396.153</td>
</tr>
<tr>
<td>Fe</td>
<td>259.939, 238.204</td>
</tr>
<tr>
<td>Ca</td>
<td>315.887, 317.932</td>
</tr>
<tr>
<td>Mg</td>
<td>280.271, 279.075</td>
</tr>
<tr>
<td>Na</td>
<td>589.592, 588.995</td>
</tr>
<tr>
<td>K</td>
<td>766.490</td>
</tr>
<tr>
<td>Mn</td>
<td>260.570, 257.608</td>
</tr>
<tr>
<td>P</td>
<td>178.221, 214.915</td>
</tr>
<tr>
<td>As</td>
<td>193.69</td>
</tr>
<tr>
<td>Ba</td>
<td>233.525, 455.507</td>
</tr>
<tr>
<td>Cd</td>
<td>226.501, 214.435</td>
</tr>
<tr>
<td>Co</td>
<td>228.614</td>
</tr>
<tr>
<td>Cr</td>
<td>267.710, 205.558</td>
</tr>
<tr>
<td>Cu</td>
<td>324.753, 327.396</td>
</tr>
<tr>
<td>Ni</td>
<td>232.003, 231.604,</td>
</tr>
<tr>
<td>Pb</td>
<td>220.353, 216.998</td>
</tr>
<tr>
<td>Se</td>
<td>196.026</td>
</tr>
<tr>
<td>Zn</td>
<td>213.857, 206.197</td>
</tr>
</tbody>
</table>

7.6 Use the Mehlich No. 3 extracting solution to dilute those samples with concentrations greater than the high standard. Rerun all elements and use only the data needed from the diluted analysis.
7.7 Establish detection limits using the blank standard solution. The instrumental detection limits are calculated by using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are reported as “ND” or non-detected. The digested fraction needs to be identified with each sample.

9. Calculations
The calculation of mg kg\(^{-1}\) of an element in the soil from mg L\(^{-1}\) in solution is as follows:

\[
\text{Analyte concentration in soil (mg kg}^{-1}\text{)} = \frac{(A \times B \times C \times R \times 1000)}{E}
\]

where:
- \(A\) = Sample extract reading (mg L\(^{-1}\))
- \(B\) = Extract volume (L)
- \(C\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (3D2)
- 1000 = Conversion factor to kg-basis
- \(E\) = Sample weight (g)

9. Report
Data are reported to the nearest 0.1 mg kg\(^{-1}\).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
Soil Test Analyses (4D)

Citric Acid Soluble (4D7)

UV-Visible Spectrophotometer, Dual-Beam (4D7a)

Phosphorus (4D7a1)

Air-Dry or Field-Moist, <2 mm (4D7a1a-b1)

1. Application

In soil taxonomy, citric acid soluble $P_2O_5$ is a criterion for distinguishing between mollic (<250 ppm $P_2O_5$) and anthropic epipedons (>250 ppm $P_2O_5$) (Soil Survey Staff, 2014). Additional data on anthropic epipedons from several parts of the world may permit improvements in this definition (Soil Survey Staff, 2014).

Phosphorus (citrate-soluble) and phosphorus (citrate-insoluble) are recognized methods (960.01 and 963.03, respectively) in the Official Methods of Analysis by the Association of Analytical Communities (AOAC) International (AOAC, 2000). The AOAC citrate-soluble P method considers the recovery of phosphite source materials as available phosphorus, even though the Association of American Plant Food Control Officials does not recognize phosphite as a source of available phosphorus. The procedure described herein is based on the method developed by Dyer (1894).

2. Summary of Method

A sample is checked for $CaCO_3$ equivalent. Sufficient citric acid is added to sample to neutralize the $CaCO_3$ plus bring the solution concentration of citric acid to 1%. A 1:10 ratio of soil to solution is maintained for all samples. The sample is shaken for 16 h and filtered. Ammonium molybdate and stannous chloride are added. Absorbance is read using a spectrophotometer at 660 nm. Data are reported as mg $P_2O_5$ kg$^{-1}$ soil (method 4D7a1).

3. Interferences

Unreacted carbonates interfere with the extraction of $P_2O_5$. Sufficient citric acid is added to sample to neutralize the $CaCO_3$. However, a high citrate level in a sample may interfere with the molybdate blue test. If this occurs, the method can be modified by evaporating the extract and ashing in a muffle furnace to destroy the citric acid.

Positive interferences in the analytical determination of $P_2O_5$ are silica and arsenic, if the sample is heated. Negative interferences in the $P_2O_5$ determination are arsenate, fluoride, thorium, bismuth, sulfide, thiosulfate, thiocyanate, or excess...
molybdate. A concentration of Fe >1000 ppm interferes with P$_2$O$_5$ determination. Refer to Snell and Snell (1949) and Metson (1956) for additional information on interferences in the citric acid extraction of P$_2$O$_5$.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow standard laboratory procedures.

5. Equipment

5.1 Electronic balance, ±0.10-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min$^{-1}$, 1½ strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL, graduated, polyethylene
5.4 Bottles, with gas release caps
5.5 Filter paper, Whatman 42, 150-mm
5.6 Funnel, 60° angle, long-stem, 50-mm diameter
5.7 Pipettes, electronic digital, 1000-µL and 10-mL, with tips, 1000-µL and 10-mL
5.8 Dispenser, 30-mL or 10-mL
5.9 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.10 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.11 Computer, with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water
6.2 Hydrochloric acid (HCl), concentrated, 12 N, trace pure grade
6.3 Citric acid solution, 10%. In a 1-L volumetric, dissolve 100 g of anhydrous citric acid (C$_6$H$_8$O$_7$) in RODI water and bring to volume with RODI water.
6.4 Citric acid solution, 1%. Dilute 100.0 mL of 10% citric acid solution to 1 L with RODI water.
6.5 Ammonium molybdate solution, 1.5%. Dissolve 15.0 g of ammonium molybdate [(NH$_4$)$_6$MO$_7$O$_{24}$•4H$_2$O] in 300 mL of distilled water. Transfer to a 1-L volumetric flask and carefully add 310 mL of concentrated HCl. Allow
to cool. Make to 1-L volume with RODI water. Store in brown bottle in the dark in a refrigerator. Solution is stable for ≈3 months.

6.6 Stock stannous chloride solution (SSCS). Dissolve 10 g of stannous chloride (SnCl₂·2H₂O) in 100 mL of concentrated HCl. Invert to mix thoroughly. Make fresh weekly. Store in a refrigerator.

6.7 Working stannous chloride solution (WSCS). Dilute 2 mL of SSCS with 100 mL of RODI water. Use immediately because solution is only stable for ≈4 h.

6.8 Stock standard P₂O₅ solution (SSPS), 250 mg L⁻¹ P. Dissolve 1.099 g of potassium dihydrogen orthophosphate (KH₂PO₄) (dried for 2 h at 110 °C) with RODI water in 1-L volumetric flask. Add 5 mL of 2 N HCl. Make to 1-L volume with RODI water. Invert to mix thoroughly. Store in polyethylene bottles. Make fresh weekly. Store in a refrigerator.

6.9 Working stock standard P₂O₅ solution (WSSPS), 2.5 mg L⁻¹ P. Pipette 10.0 mL of SSPS and dilute to 1 L in a volumetric flask with RODI water. Invert to mix thoroughly. Make fresh daily.

6.10 Standard P₂O₅ calibration solutions (SPCS). Pipette 0.0, 1.0, 2.0, 3.0, 4.0, and 5.0 mL of WSSPS into 50-mL tubes, add 1 mL of 1% citric acid solution, 4 mL of ammonium molybdate solution, and dilute to 25 mL with RODI water. Add 2 mL of the working stannous chloride solution (WSCS). Final concentrations are 0.0, 0.10, 0.20, 0.30, 0.40, and 0.50 mg L⁻¹, respectively. Invert to mix thoroughly. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use.

6.11 Quality Control Samples: 0.1 mg P L⁻¹ solution from SSPS; blanks; selected SPCS; and KSSL soil standard are routinely included in a batch for quality control.

7. Procedure

7.1 Weigh 3 g of <2-mm or fine-grind, air-dry soil to the nearest mg into a bottle with gas release tops. If sample is moist, weigh enough soil to achieve ≈3 g of air-dry soil. If the soil does not contain free carbonates, proceed to step 7.3.

7.2 If the soil contains free CaCO₃, refer to Table 1 to determine the amount of 10% citric acid solution required to neutralize the CaCO₃. Add required volume of 10% citric acid into a graduated cylinder and bring to a volume of 30 mL with RODI water. Add this solution to the soil. Place the bottle in a mechanical shaker and shake the bottle for 6 h at 200 oscillations min⁻¹ at room temperature (20 ±2 °C) to dissolve and neutralize the CaCO₃. Proceed to step 7.4.
Table 1.—Volume of 10% Citric Acid (mL) Required to Decompose CaCO₃ (%) and to Bring Solution Concentration to 1% in a Final Volume of 30 mL for 3-g Sample.

<table>
<thead>
<tr>
<th>% CC¹</th>
<th>mL CA²</th>
<th>% CC</th>
<th>mL CA</th>
<th>% CC</th>
<th>mL CA</th>
<th>% CC</th>
<th>mL CA</th>
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<tbody>
<tr>
<td>0</td>
<td>3.0</td>
<td>16</td>
<td>9.1</td>
<td>32</td>
<td>15.3</td>
<td>48</td>
<td>21.4</td>
</tr>
<tr>
<td>1</td>
<td>3.4</td>
<td>17</td>
<td>9.5</td>
<td>33</td>
<td>15.7</td>
<td>49</td>
<td>21.8</td>
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<tr>
<td>2</td>
<td>3.8</td>
<td>18</td>
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<td>10.3</td>
<td>35</td>
<td>16.4</td>
<td>51</td>
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<tr>
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<td>4.5</td>
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<td>10.7</td>
<td>36</td>
<td>16.8</td>
<td>52</td>
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<tr>
<td>5</td>
<td>4.9</td>
<td>21</td>
<td>11.1</td>
<td>37</td>
<td>17.2</td>
<td>53</td>
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<td>5.3</td>
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<td>11.4</td>
<td>38</td>
<td>17.6</td>
<td>54</td>
<td>23.7</td>
</tr>
<tr>
<td>7</td>
<td>5.7</td>
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<td>11.8</td>
<td>39</td>
<td>18.0</td>
<td>55</td>
<td>24.1</td>
</tr>
<tr>
<td>8</td>
<td>6.1</td>
<td>24</td>
<td>12.2</td>
<td>40</td>
<td>18.4</td>
<td>56</td>
<td>24.5</td>
</tr>
<tr>
<td>9</td>
<td>6.5</td>
<td>25</td>
<td>12.6</td>
<td>41</td>
<td>18.7</td>
<td>57</td>
<td>24.9</td>
</tr>
<tr>
<td>10</td>
<td>6.8</td>
<td>26</td>
<td>13.0</td>
<td>42</td>
<td>19.1</td>
<td>58</td>
<td>25.3</td>
</tr>
<tr>
<td>11</td>
<td>7.2</td>
<td>27</td>
<td>13.4</td>
<td>43</td>
<td>19.5</td>
<td>59</td>
<td>25.6</td>
</tr>
<tr>
<td>12</td>
<td>7.6</td>
<td>28</td>
<td>13.7</td>
<td>44</td>
<td>19.9</td>
<td>60</td>
<td>26.0</td>
</tr>
<tr>
<td>13</td>
<td>8.0</td>
<td>29</td>
<td>14.1</td>
<td>45</td>
<td>20.3</td>
<td>61</td>
<td>26.4</td>
</tr>
<tr>
<td>14</td>
<td>8.4</td>
<td>30</td>
<td>14.5</td>
<td>46</td>
<td>20.7</td>
<td>62</td>
<td>26.8</td>
</tr>
<tr>
<td>15</td>
<td>8.8</td>
<td>31</td>
<td>14.9</td>
<td>47</td>
<td>21.0</td>
<td>63</td>
<td>27.2</td>
</tr>
</tbody>
</table>

¹ %CC = percent calcium carbonate in a sample
² CA = mL of 10% citric acid needed to be diluted to 30-ml volume with RODI water and added to sample

7.3 If the soil contains no free CaCO₃, add 30 mL of 1% citric acid solution to the sample.

7.4 Cap the bottles, place in a mechanical shaker and shake for 16 h at 200 oscillations min⁻¹ at room temperature (20 ±2 °C).

7.5 Remove the sample from shaker and filter. Collect extract. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.6 Pipette 1 mL of sample extract into a plastic cup. Samples are treated the same as SPCS (Section 6.10). Add 4 mL of ammonium molybdate solution to all samples and SPCS and dilute to 25-mL with RODI water. Add 2 mL
working stannous chloride solution (WSCS). Swirl to mix and allow to stand 20 min for color development.

7.7 Transfer sample extract and SPCS to cuvettes.

7.8 Set the spectrophotometer to read at 660 nm. Autozero with calibration blank. A blank has all reagents contained in the sample extract except the soil.

7.9 Calibrate the instrument by using the SPCS. The data system then associates the concentrations with the instrument responses for each SPCS. Rejection criteria for SPCS is $R^2 < 0.99$.

7.10 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Rejection criteria for batch are as follows: if blanks are >0.01; if SPCS vary by more than 20% from calculated value; or if KSSL standard varies by more than 20% from the accepted mean. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.11 If samples are outside calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations

Convert the extract $P_2O_5$ (mg L$^{-1}$) to soil $P_2O_5$ (mg kg$^{-1}$) as follows:

$$\text{Soil } P_2O_5 \text{ (mg kg}^{-1}\text{)} = \left( A \times B \times C_1 \times C_2 \times R \times 1000 \times 2.29 \right) / E$$

where:

$A = P_2O_5$ in sample extract (mg L$^{-1}$)

$B =$ Extract volume (L)

$C_1 =$ Automatic dilution

$C_2 =$ Dilution, if necessary

$R =$ Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

1000 = Conversion factor to kg-basis

2.29 = Conversion factor $P$ to $P_2O_5$

$E =$ Sample weight (g)

9. Report

Report the 1% citrate acid extractable $P_2O_5$ in mg kg$^{-1}$ to nearest whole number.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.
11. References

Association of Analytical Communities (AOAC) International. 2000. Phosphorus (citrate-soluble) in fertilizers and phosphorus (citrate-insoluble) in fertilizers, methods 960.01 and 963.03, respectively. Official Methods of Analysis.


Soil Test Analyses (4D)
New Zealand P Retention (4D8)

UV-Visible Spectrophotometer, Dual-Beam (4D8a)
Phosphorus (4D8a1)
Air-Dry or Field-Moist, <2 mm (4D8a1a-b1)

1. Application

In soil taxonomy, the P retention of soil material is a criterion for andic soil properties (Soil Survey Staff, 2014). Andisols and other soils that contain large amounts of allophane and other amorphous minerals have capacities for binding P (Gebhardt and Coleman, 1984). The factors that affect soil P retention are not well understood. However, allophane and imogolite have been considered as major materials that contribute to P retention in Andisols (Wada, 1985). Phosphate retention is also called P adsorption, sorption, or fixation.

2. Summary of Method

A 5-g soil sample is shaken in a 25-mL aliquot of a 1000 mg L$^{-1}$ P solution for 24 h. The mixture is centrifuged at 2000 rpm for 15 min. An aliquot of the supernatant is transferred to a colorimetric tube. Nitric vanadomolybdate acid reagent (NVAR) is added. Absorbance of the solution is read using a spectrophotometer at 466 nm. This absorbance correlates to the concentration of the non–adsorbed P that remains in the sample solution. The New Zealand P retention (Blakemore et al., 1987) is the initial P concentration minus the P remaining in the sample solution and is reported as percent P retained (4D8a1).

3. Interferences

No significant problems are known to affect the P retention measurement.
4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated HNO₃ to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 100 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Digital diluter/dispenser with syringes, 10,000 and 1000 µL, gas tight, Microlab 500, Reno, NV
5.4 Centrifuge, Centra GP-8
5.5 Centrifuge tubes, 50-mL, polyethylene
5.6 Cups, plastic
5.7 Pipettes, electronic digital, 1000-µL and 10-mL, with tips, 1000-µL and 10-mL
5.8 Dispenser, 30-mL or 10-mL
5.9 Filter paper, Whatman 42, 150-mm
5.10 Cuvettes, plastic, 4.5-mL, 1-cm path, Daigger Scientific
5.11 Spectrophotometer, UV-Visible, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.12 Computer, with Cary WinUV software, Varian Australia Pty Ltd., with printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Superfloc 16, 0.2%, 2 g L⁻¹ in RODI water
6.3 Nitric acid (HNO₃), concentrated, 16 N
6.4 P retention solution (PRS), 1000 mg L⁻¹ P. Dissolve 35.2 g of KH₂PO₄ (dried for 2 h at 110 °C) and 217.6 g of sodium acetate trihydrate (CH₃COONa•3H₂O) in RODI water. Add 92 mL of glacial acetic acid. Dilute to 8 L with RODI water. The solution pH should range between 4.55 and 4.65.
6.5 Nitric acid solution. Carefully and slowly dilute 200 mL of concentrated HNO₃ to 2 L of RODI water. Add the acid to the water.
6.6 Molybdate solution. Dissolve 32 g of ammonium molybdate \( [(NH_4)_6Mo_7O_{24}\cdot4H_2O] \) in 50 °C RODI water. Allow the solution to cool to room temperature and dilute to 2 L with RODI water.

6.7 Nitric vanadomolybdate acid reagent (NVAR), vanadate solution. Dissolve 1.6 g of \( NH_4VO_3 \) in 500 mL of boiling RODI water. Allow the solution to cool to room temperature. Carefully and slowly add 12 mL of concentrated \( HNO_3 \). Dilute to 2 L with RODI water. Mix the nitric acid solution with the vanadate solution and then add the molybdate solution. Mix well. Note: This solution needs to be made in the order specified.

6.8 Diluent for Standard P Calibration Solutions (DSPCS). Add 54.4 g of sodium acetate trihydrate \( (CH_3COONa\cdot3H_2O) \) and 23 mL of glacial acetic acid and dilute to 2 L with RODI water.

6.9 Standard P calibration solutions (SPCS), 100, 80, 60, 40, 20, and 0% P retained. Make fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. To six 100-mL volumetric flasks, add the appropriate amount of PRS \( (1000 \text{ mg L}^{-1}) \) and bring to volume with DSPCS solution as follows:

6.9.1 100% SPCS = \( (0 \text{ mg L}^{-1}) \) = Bring to 100-mL volume with DSPCS solution.

6.9.2 80% SPCS = 1:20 (200 mg L\(^{-1}\)) = Add 20 mL PRS and bring to 100-mL volume with DSPCS solution.

6.9.3 60% SPCS = 1:10 (400 mg L\(^{-1}\)) = Add 40 mL PRS and bring to 100-mL volume with DSPCS solution.

6.9.4 40% SPCS = 3:20 (600 mg L\(^{-1}\)) = Add 60 mL PRS and bring to 100-mL volume with DSPCS solution.

6.9.5 20% SPCS = 1:5 (800 mg L\(^{-1}\)) = Add 80 mL PRS and bring to 100-mL volume with DSPCS solution.

6.9.6 0% SPCS = 1:4 (1000 mg L\(^{-1}\)) = Add 100 mL PRS to a 100-mL volumetric.

6.10 Quality Control: KSSL soil standard is routinely included in every batch of 24 samples for quality control.

7. Procedure

7.1 Weigh 5 g of <2-mm or fine-grind, air-dry soil to the nearest mg into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈5 g of air-dry soil. No fine-grind material is used for the moist procedure.

7.2 Use the dispenser to add 25.0 mL of P-retention solution to centrifuge tube.

7.3 Transfer the sample to the shaker. Shake for 24 h at 100 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).
7.4 Remove sample from the shaker. Add 2 or 3 drops of Superfloc, 0.02% w/v to each tube. Centrifuge at 2000 rpm for 15 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.5 Use the digital diluter to add the nitric vanadomolybdate acid reagent (NVAR) to each sample supernatant and to each SPCS. To fill a 4.5-mL cuvette, use a 1:20 sample dilution.

7.6 The color reaction requires a minimum of 30 min before the analyst records readings.

7.7 Set the spectrophotometer to read at 466 nm. Autozero with the calibration blank.

7.8 Calibrate the instrument using the SPCS. The data system then associates the percent P retained with the instrument responses. Rejection criteria for SPCS is $R^2 < 0.99$.

7.9 Run samples using calibration curve. Sample P retention is calculated from the regression equation, i.e., the absorbance is equated to the SPCS. Rejection criteria for batch are as follows: if blanks are $>0.01$; if SPCS vary by more than 20% from calculated value; or if KSSL standard varies by more than 20% from the accepted mean. Record results to the nearest 0.01 unit for the sample extract and each SPCS.

7.10 If samples are outside calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations
   None.

9. Report
   Report the percent New Zealand P retention to the nearest whole number.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

Soil Test Analyses (4D)
1 M KCl Extraction (4D9)
   Cadmium-Copper Reduction (4D9a)
   Sulfanilamide N-1-Naphthylethylenediamine Dihydrochloride (4D9a1)
   Flow-Injection, Automated Ion Analyzer (4D9a1a)
   Nitrate (4D9a1a1)
       Air-Dry or Field-Moist, <2 mm (4D9a1a1a-b1)

1. Application
   The inorganic combined N in soils is predominantly NH$_4^+$ and NO$_3^-$ (Keeney and Nelson, 1982). Nitrogen in the form of ammonium ions and nitrate are of particular concern because they are very mobile forms of nitrogen and are most likely to be lost to the environment (National Research Council, 1993). All forms of nitrogen, however, are subject to transformation to ammonium ions and nitrate as part of the nitrogen cycle in agro-ecosystems, and all can contribute to residual nitrogen and nitrogen losses to the environment (National Research Council, 1993).

2. Summary of Method
   A 2.5-g soil sample is mechanically shaken for 30 min in 25 mL of 1 M KCl solution. The sample is then filtered through Whatman No. 42 filter paper. A flow injection automated ion analyzer is used to measure the soluble inorganic nitrate (NO$_3^-$). The nitrate is quantitatively reduced to nitrite by passage of the sample through a copperized cadmium column. The nitrite (reduced nitrate plus original nitrite) is then determined by diazotizing with sulfanilamide followed by coupling with N-1-naphthylethylenediamine dihydrochloride. The resulting water soluble dye has a magenta color which is read at 520 nm. Absorbance is proportional to the concentration of NO$_3^-$ in the sample. Data are reported as mg N kg$^{-1}$ soil as NO$_3^-$ (4D9a1a1).

3. Interferences
   Nitrite is oxidized by air to nitrate in a few days. If analysis can be made within 24 h, the sample should be preserved by refrigeration at 4 ºC. When samples must be stored for more than 24 h, they should be preserved with sulfuric acid (2 mL concentrated H$_2$SO$_4$ per liter) and refrigerated (LACHAT, 2003). Low results can be obtained for samples that contain high concentration of Fe, Cu, or other metals. In this method, EDTA is added to the buffer to reduce this interference (LACHAT, 2003).
4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of NH₄OH and concentrated HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Cadmium is toxic and carcinogenic. Wear gloves and follow the precautions described on the Material Safety Data Sheet. If the cadmium-copper reduction column is repacked, all transfers should be done over a special tray or beaker dedicated to this purpose. Preferably, send the cadmium-copper column to LACHAT for repacking.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Centrifuge tubes, 25-mL, polyethylene, Oak Ridge
5.3 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Centrifuge, 50-mL, polyethylene
5.5 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL
5.6 Filter paper, Whatman 42, 150-mm
5.7 Funnel, 60° angle, long stem, 50-mm diameter
5.8 Volumetric flasks, 1-L and 250-mL
5.9 Bottles, plastic, dark, 1-L
5.10 Cups, plastic
5.11 Dispenser, 30-mL or 10-mL
5.12 Flow Injection Automated Ion Analyzer, QuikChem 8500, LACHAT Instruments, Loveland, CO, with computer and printer
5.13 Sampler, LACHAT Instruments, Loveland, CO
5.14 Reagent Pump, LACHAT Instruments, Loveland, CO
5.15 Automated Dilution Station, LACHAT Instruments, Loveland, CO
5.16 Sample Processing Module (SPM) or channel, QuikChem Method (12-107-04-1-B, nitrate in 2 M KCl soil extracts by flow injection analysis, 0.025 to 20.0 mg N L⁻¹), LACHAT Instruments, Loveland, CO
5.17 Computer, with QuikChem software, LACHAT Instruments, Loveland, CO, and printer
5.18 Vials, plastic, 25-mL (standards)
5.19 Culture tubes, glass, 10-mL (samples)
6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 Helium, compressed gas

6.3 15 M NaOH. In a 500-mL container, add 250 mL RODI water. Slowly add 300 g NaOH. (CAUTION: The solution will get very hot!) Swirl until dissolved. Cool and store in a plastic bottle. Use to adjust ammonium chloride buffer to pH 8.5 (Reagent 6.4).

6.4 Ammonium chloride buffer, pH 8.5. In a hood, add 500 mL RODI to a 1-L volumetric flask. Add 105 mL concentrated HCl, 90 mL ammonium hydroxide (NH₄OH), and 1.0 g disodium EDTA. Dissolve and dilute to mark. Invert to mix. Degas with helium ≈5 min.

6.5 Sulfanilamide color reagent. To a 1-L volumetric flask, add 600 mL RODI H₂O followed by 100 mL 85 percent phosphoric acid (H₃PO₄), 40.0 g sulfanilamide, and 1.0 g N-1-naphthylethylenediamine dihydrochloride (NED). Shake to wet and stir to dissolve 20 min. Dilute to mark and invert to thoroughly mix. Degas with helium ≈5 min. Store in dark bottle and discard when pink.

6.6 1 M KCl extracting solution, carrier and standards diluent. Dissolve 74.5 g potassium chloride (KCl) in 800 mL RODI water. Dilute to mark and invert to thoroughly mix. The extracting solution is used also as the carrier and a component of the N standards. Degas with helium ≈5 min.

6.7 The following are standards for a 1-channel system determining NO₂⁻ + NO₃⁻ or NO₂⁻ and a 2-channel system where one channel is used for NO₂⁻ + NO₃⁻ and the other channel is used for determining NO₂⁻. For the 1-channel system, either NO₂⁻ or NO₃⁻ standards may be used. It is recommended that when running a 1 channel method for NO₂⁻ + NO₃⁻ that NO₃⁻ standards are used. For the 2-channel system, the use of both NO₂⁻ + NO₃⁻ standard sets are recommended.

6.7.1 Stock Standard Nitrate Solution (SSNO₃S), 200.0 mg N L⁻¹ as NO₃⁻ in 1 M KCl. In a 1-L volumetric flask, dissolve 1.444 g potassium nitrate (KNO₃) (dried in an oven for 2 h at 110 °C) and 74.5 g KCl in 600 mL RODI water. Dilute to mark with RODI water and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.7.2 Working Stock Standard Nitrate Solution (WSSNO₃S), 20.0 mg N L⁻¹ as NO₃⁻ in 1 M KCl. To a 1-L volumetric flask, add 100 mL SSNO₃S. Dilute to mark with 1 M KCl and invert to thoroughly mix. Make fresh daily.

6.7.3 Standard Nitrate Calibration Standards (SNO₃CS), or working standards, 10.00, 1.00, 0.80, 0.08, and 0.00 mg N L⁻¹ as NO₃⁻ in
1 M KCl. Make fresh daily. To five 250-mL volumetric flasks, add as follows:

6.7.3.1 10.00 mg N L$^{-1}$ = 125.0 mL WSSNO$_3$S
6.7.3.2 1.00 mg N L$^{-1}$ = 12.5 mL WSSNO$_3$S
6.7.3.3 0.80 mg N L$^{-1}$ = 10.0 mL WSSNO$_3$S
6.7.3.4 0.08 mg N L$^{-1}$ = 1.00 mL WSSNO$_3$S
6.7.3.5 0.00 mg N L$^{-1}$ = 0.0 mL WSSNO$_3$S (blank)

Dilute each SNO$_3$CS to the mark with 1 M KCl and invert to thoroughly mix. Do not degas.

6.7.4 Stock Standard Nitrite Solution (SSNO$_2$S), 200.0 mg N L$^{-1}$ as NO$_2^-$ in 1 M KCl. In a 1-L volumetric flask, dissolve 74.5 g KCl and either 0.986 g sodium nitrite (NaNO$_2$) or 1.214 g potassium nitrite (KNO$_2$) in 800 mL RODI water. Dilute to mark with RODI water and invert to thoroughly mix. Store in polyethylene containers in a refrigerator. Make fresh weekly.

6.7.5 Working Stock Standard Nitrite Solution (WSSNO$_2$S), 20.0 mg N L$^{-1}$ as NO$_2^-$ in 1 M KCl. To a 1-L volumetric flask, add 100 mL SSNO$_2$S. Dilute to mark with 1 M KCl and invert to thoroughly mix. Make fresh daily.

6.7.6 Standard Nitrite Calibration Standards (SNO$_2$CS), or working standards, 10.00, 1.00, 0.80, 0.08, and 0.00 mg N L$^{-1}$ as NO$_3^-$ in 1 M KCl. Make fresh daily. To five 250-mL volumetric flasks, add as follows:

6.7.6.1 10.00 mg N L$^{-1}$ = 125.0 mL WSSNO$_2$S
6.7.6.2 1.00 mg N L$^{-1}$ = 12.5 mL WSSNO$_2$S
6.7.6.3 0.80 mg N L$^{-1}$ = 10.0 mL WSSNO$_2$S
6.7.6.4 0.08 mg N L$^{-1}$ = 1.00 mL WSSNO$_2$S
6.7.6.5 0.00 mg N L$^{-1}$ = 0.0 mL WSSNO$_2$S (blank)

Dilute each SNO$_2$CS to the mark with 1 M KCl and invert to thoroughly mix. Do not degas.

7. Procedure

**Extraction**

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to the nearest mg and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Add ≈25 mL of 1 M KCl to sample. Transfer the sample to a shaker. Shake for 30 min at 200 oscillations min$^{-1}$ at room temperature (20 ±2 °C).
7.3 Remove the sample from the shaker. Decant, filter, and collect extract in receiving cups. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 24 h.

**Flow Injection Set-up and Operation**

7.4 Transfer sample extracts into culture tubes and place in sample trays marked “Samples.”

7.5 Transfer WNCS standards into plastic vials and place in descending order in sample trays marked “Standards.”

7.6 Refer to the operating and software reference manuals for LACHAT for set-up and operation. Refer to LACHAT Method QuikChem Method 12-107-04-1-B for data system parameters, such as analyte and calibration data and sampler and valve timing.

7.7 Run samples using calibration curve. Sample concentration is calculated from the regression equations. Report results to the nearest 0.01 unit for the sample extract and each SPCS.

7.8 If samples are outside calibration range, dilute samples with extracting solution and re-analyze.

7.9 Upon completion of run, place the transmission lines into RODI water and pump for approximately 20 min before proceeding with the normal “Shut-down” procedure.

7.10 KCl may accumulate and cause clogs in the manifold tubing and the fittings over time. The valves and fittings therefore need to be washed with RODI water upon completion of analysis. Some fittings may need to be soaked overnight or placed in a sonic bath for 10 to 15 min to remove any KCl accumulations.

8. Calculations

Convert extract N (mg L⁻¹) to soil N (mg kg⁻¹) as follows:

\[
\text{Soil N} = \frac{(A \times B \times C \times R \times 1000)}{E}
\]

where:

- A = Sample extract reading (mg N L⁻¹)
- B = Extract volume (L)
- C = Dilution, if performed
- R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- E = Sample weight (g)
9. Report

Report data to the nearest 0.1 mg N kg\(^{-1}\) soil as NO\(_3^-\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


LACHAT Instruments. 2003. QuikChem method 12-107-04-1-B, nitrate in 2 \(M\) KCl soil extracts by flow injection analysis, 0.025 to 20.0 mg N L\(^{-1}\). LACHAT Instruments, Loveland, CO.


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Soil Test Analyses (4D)

Anaerobic Incubation (4D10)

2 \(M\) KCl Extraction (4D10a)

Ammonia–Salicylate (4D10a1)

Flow Injection, Automated Ion Analyzer (4D10a1a)

N as NH\(_3\) (Mineralizable N) (4D10a1a1)

Air-Dry or Field-Moist, <2 mm (4D10a1a1a-b1)

1. Application

The most satisfactory methods currently available for obtaining an index for the availability of soil N are those involving the estimation of the N formed when soil is incubated under conditions that promote mineralization of organic N by soil microorganisms (USEPA, 1992). The method described herein for estimating mineralizable N is an anaerobic incubation and is suitable for routine analysis of soils. This method involves estimation of the ammonium produced by a 1-week period of incubation of soil at 40 °C (Keeney and Bremner, 1966) under anaerobic conditions to provide an index of N availability.

2. Method Summary

An aliquot of air-dry, homogenized soil is placed in a test tube with water, stoppered, and incubated at 40 °C for 1 week. The contents are rinsed with 2 \(M\) KCl. A flow injection automated ion analyzer is used to measure the ammonium
produced in the soil after incubation. Absorbance of the solution is read at 660 nm. Data are reported as mg N kg\(^{-1}\) soil as NH\(_3\) by method 4D10a1a1.

3. Interferences

The temperature and incubation period must remain constant for all samples. The test can be performed on field-moist or air-dry soil samples. Ammonia is volatile and slowly leaves the sample even through polyethylene bottles. Samples should be run within 24 h of extraction. If this cannot be done, the samples should be adjusted to pH 3–5 with dilute sulfuric acid (LACHAT, 2003). The pH of the standards solutions should approximate that of the samples, i.e., if samples have been preserved with sulfuric acid, then the preservation acid should be added in standards preparation (LACHAT, 2003). Remove interfering turbidity by filtration. Soil extracts can contain sufficient concentrations of calcium and magnesium to cause precipitation during analysis. EDTA is added to eliminate this problem (Keeney and Nelson, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Test tubes, 16-mm x 150-mm
5.3 PVC stoppers
5.4 Incubator, Model 10-140, Quality Lab Inc., Chicago, IL
5.5 Centrifuge, 50-mL, polyethylene
5.6 Filter paper, Whatman 42, 150-mm
5.7 Funnel, 60° angle, long stem, 50-mm diameter
5.8 Flow Injection Automated Ion Analyzer, QuikChem 8500, LACHAT Instruments, Loveland, CO
5.9 Sampler, LACHAT Instruments, Loveland, CO
5.10 Reagent Pump, LACHAT Instruments, Loveland, CO
5.11 Automated Dilution Station, LACHAT Instruments, Loveland, CO
5.12 Sample Processing Module (SPM) or channel, QuikChem Method (12-107-06-2-A, ammonia (salicylate) in 2 M KCl soil extracts, 0.1 to 20.0 mg N L\(^{-1}\)), LACHAT Instruments, Loveland, CO
5.13 Computer, with QuikChem software, LACHAT Instruments, Loveland, CO, and printer

5.14 Vials, plastic, 25-mL (standards)

5.15 Culture tubes, glass, 10-mL (samples)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 Helium, compressed gas


6.4 EDTA (ethylene tetraacetic acid disodium salt dihydrate), 6% solution. Dissolve 66 g EDTA in 900 mL RODI water. Dilute to 1 L and invert to mix thoroughly. Degas with helium.

6.5 NaOH, buffer. Dissolve 28.0 g NaOH and 50.0 g sodium phosphate dibasic heptahydrate (Na₂HPO₄•7H₂O) in 900 mL RODI water. Dilute to 1 L and invert to mix thoroughly. Degas with helium.

6.6 Salicylate-Nitroprusside Color Reagent. Dissolve 150 g sodium salicylate [salicylic acid sodium salt (C₆H₄(OH)(COO)Na)] and 1.0 g sodium nitroprusside [sodium nitroferricyanide dihydrate (Na₂Fe(CN)₅NO•2H₂O)] in 800 mL RODI water. Dilute to 1 L and invert to mix thoroughly. Degas with helium. Store in dark bottle in a refrigerator.

6.7 Hypochlorite Reagent. In a 500-mL volumetric, dilute 250 mL of 5.25% sodium hypochlorite (NaOCl) to mark with RODI water. Invert to mix thoroughly. Degas with helium.

6.8 Stock Standard N Solution (SSNS), 100.0 mg N L⁻¹. In a 1-L volumetric flask, dissolve 150 g potassium chloride (KCl) and 0.3819 g of ammonium chloride (NH₄Cl) (dried for 2 h at 110 °C) in about 800 mL RODI water. Dilute to volume with RODI water and invert to thoroughly mix. Do not degas with helium. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.9 Standard N Calibration Solutions (SNCS), or working standards, 20.0, 8.00, 2.00, 0.50, 0.10, 0.00 mg N L⁻¹. Make fresh daily. To six 250-mL volumetric flasks, add as follows:

6.9.1 20.00 mg P L⁻¹=50.0 mL SSNS

6.9.2 8.00 mg P L⁻¹=20.0 mL SSNS

6.9.3 2.00 mg P L⁻¹=5.0 mL SSNS

6.9.4 0.50 mg P L⁻¹=1.25 mL SSNS
6.9.5 $0.10 \text{ mg P L}^{-1} = 0.25 \text{ mL SSNS}$
6.9.6 $0.00 \text{ mg P L}^{-1} = 0.0 \text{ mL SSNS (blank)}$

Dilute to mark with 2 M KCl. Invert to mix thoroughly.

7. Procedure

**Anaerobic Incubation of Soil Sample**

7.1 Weigh 5 g of <2 mm, air-dry soil to the nearest mg into a 16 mm x 150 mm test tube. If soil is fine-grind, weigh 1.25 g. If sample is moist, weigh enough soil to achieve ≈5 or 1.25 g, respectively, of air-dry soil.

7.2 Add 12.5 ±1 mL of RODI water. Do not add ethanol to overcome any wetting difficulties because ethanol interfere with microbial activity. Stopper the tube, shake, and place in a 40 °C constant-temperature incubator for 7 days. Refer to the manufacturer’s instructions for set-up and operation of the incubator.

7.3 At the end of 7 days, remove the tube and shake for 15 s.

7.4 Transfer the contents of the test tube to another test tube. Complete the transfer by rinsing the tube 3 times with 4 mL of 2 M KCl, using a total of 12.5 ±1 mL of the KCl. Filter contents into a centrifuge tube. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 24 h. Ammonia is volatile and slowly leaves the sample, even through the polyethylene bottles.

**Flow Injection Set-up and Operation**

7.5 Transfer sample extracts into culture tubes and place in XYZ sample trays marked “Samples.”

7.6 Transfer WNCS standards into plastic vials and place in descending order in XYZ sample trays marked “Standards.”

7.7 Refer to the operating and software reference manuals for LACHAT for set-up and operation. Refer to LACHAT Method QuikChem Method 12-107-06-2-A for data system parameters, such as analyte and calibration data and sampler and valve timing.

7.8 Run samples using calibration curve. Sample concentration is calculated from the regression equations. Report results to the nearest 0.01 unit for the sample extract and each SNCS.

7.9 If samples are outside calibration range, dilute samples with extracting solution and re-analyze.

7.10 Upon completion of run, place the transmission lines into the 1 M HCl. Pump the solution for approximately 5 min to remove any precipitated
reaction products, and then place the lines in RODI water and pump for an additional 5 min before proceeding with the normal “Shut-down” procedure.

8. Calculations
Convert extract N (mg L\(^{-1}\)) to soil N (mg kg\(^{-1}\)) as follows:

\[
\text{Soil N} = \frac{(A \times B \times C \times R \times 1000)}{E}
\]

where:
- \(A\) = Analyte reading (mg L\(^{-1}\))
- \(B\) = Extract volume (L)
- \(C\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis
- \(E\) = Sample weight (g)

9. Report
Report data to the nearest mg N kg\(^{-1}\) soil as NH\(_3\).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References


LACHAT Instruments. 2003. QuikChem method (12-107-06-2-A, ammonia (salicylate) in 2 M KCl soil extracts by flow injection analysis, 0.1 to 20.0 mg N L\(^{-1}\)), LACHAT Instruments, Loveland, CO.

Carbonate and Gypsum (4E)

3 N HCl Treatment (4E1)

CO₂ Analysis (4E1a)

Manometer, Electronic (4E1a1)

Calcium Carbonates (4E1a1a1)

Air-Dry, <2 mm or <20 mm (4E1a1a1a1-2)

1. Application

The distribution and amount of CaCO₃ are important factors affecting fertility, erosion, available water capacity, and genesis of the soil. Calcium carbonate provides a reactive surface for adsorption and precipitation reactions, e.g., phosphate, trace elements, and organic acids (Loeppert and Suarez, 1996; Amer et al., 1985; Talibudeen and Arambarr, 1964; Boischot et al., 1950). The determination of calcium carbonate (CaCO₃) equivalent is a criterion in soil taxonomy (Soil Survey Staff, 2014). Carbonate content of a soil is used to define carbonatic, particle-size, and calcareous soil classes and to define calcic and petrocalcic horizons (Soil Survey Staff, 2014). The formation of calcic and petrocalcic horizons has been related to a variety of processes, some of which include translocation and net accumulation of pedogenic carbonates from a variety of sources as well as the alteration of lithogenic (inherited) carbonate to pedogenic carbonate (soil-formed carbonate through in situ dissolution and re-precipitation of carbonates) (Rabenhorst et al., 1991). The CaCO₃ equivalent is most commonly reported on the <2-mm base. However, in some soils with hard carbonate concretions, carbonates are determined on both the <2-mm (4E1a1a1a1) and the 2- to 20-mm basis (4E1a1a1a2). The CaCO₃ equivalent is routinely determined by the KSSL if the CaCl₂ pH >6.95 (method 4C1a2a2) and/or effervescence after treatment with 1 N HCl is violent, strong, slight, or very slight (method 1B1b2d2).

2. Summary of Method

The amount of carbonate in the soil is measured by treating the samples with HCl. The evolved CO₂ is measured manometrically. The amount of carbonate is then calculated as percent CaCO₃.

3. Interferences

A chemical interference is the reaction by the acid with other carbonates, e.g., carbonates of Mg, Na, and K, that may be present in soil sample. The calculated CaCO₃ is only a semiquantitative measurement (Nelson, 1982).

Analytical interference may be caused by temperature changes within the reaction vessel. When sealing the vessel, the analyst should not hold the vessel any longer than necessary to tighten the cap. The internal pressure must be equalized with the atmosphere. After the septum has been pierced with a needle,
≈5 to 10 s are required to equalize the internal pressure of the bottle. With extensive use, the septa leak gas under pressure. The septa should be replaced at regular intervals. The analyst should not touch the glass of the vessel when reading the pressure.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when handling acids. Thoroughly wash hands after handling acids. Use the fume hood when diluting concentrated HCl. Use the safety showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

The gelatin capsule may leak acid while being filled. Keep other personnel away from the area when filling capsules.

High pressure may develop inside the bottle if there is a large amount of calcareous sample. Do not use more than 2 g of any sample in the bottles. If high pressure develops in the bottle, release the pressure by venting the gas with a syringe needle. Some bottles may break without shattering. Discard any bottle with hairline cracks or obvious defects.

5. Equipment

5.1 Electronic balance, ±0.10-mg sensitivity
5.2 Electronic balance, ±1-mg sensitivity
5.3 Threaded weighing bottles, wide-mouth, clear glass, standard, 120-mL (4 fl. oz.), 48-mm neck size. For best results, grind rim of bottle with 400–600 grit sandpaper on a flat glass plate.
5.4 Machined PVC caps for threaded 120 mL (4 fl. oz.) weighing bottles, 54-mm diameter with 12.7-mm diameter hole drilled in center, O-ring seal
5.5 O-rings, 3.2 x 50.8 x 57.2 mm (⅛ x 2 x 2⅛ in)
5.6 Flanged stopper No. 03-255-5, Fisher Scientific. Place in machined cap.
5.7 Manometer, hand-held gauge and differential pressure, PCL-200 Series, Omega Engineering, Stanford, CT
5.8 Hypodermic needle, 25.4-mm (1-in), 23-gauge. Connect needle to pressure tubing on transducer.
5.9 Mechanical rotating shaker, 140-rpm, Eberbach 6140, Eberbach Corp., Ann Arbor, MI.

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Methyl red indicator
6.3 Hydrochloric acid (HCl), concentrated, 12 N

6.4 HCl, 3 N. Dilute 500 mL of concentrated HCl with 1500 mL RODI water. Add a few crystals of methyl red indicator. Methyl red indicator turns yellow if HCl is consumed by sample. If this reaction occurs, adjust the sample size (smaller).

6.5 Gelatin capsule, 10-mL, size 11, Torpac Inc., Fairfield, NJ

6.6 Glycerin, USP. Put the glycerin in a small squeeze bottle and use as a lubricant for the O-rings.

6.7 CaCO$_3$, Ultrex, assay dried basis 100.01%

7. Procedure

Manometer Calibration

7.1 Calibrate the manometer quarterly or whenever equipment changes (e.g., old rubber septum replaced). Calibrate by weighing three replicates of CaCO$_3$ standards (0, 0.025, 0.05, 0.1, 0.2, 0.3, 0.4, 0.5, 0.75 g). Weigh to the nearest 0.1 mg. Dry the standard samples in the oven for 2 h at 110 °C. Remove samples from oven, place in desiccator, and cool to ambient temperature. Proceed as outlined in Sections 7.4 through 7.9.

<2-mm Basis

7.2 A CaCl$_2$ pH >6.95 is generally used as an indicator of the presence of carbonates. The presence of carbonates (effervescence with HCl) is also checked during lab preparation (method 1B1b2b5).

7.3 Weigh 0.5 to 2 g of fine-grind, air-dry soil sample to the nearest mg and place in a 120-mL, wide-mouth bottle. Run 3 blanks and a quality control check sample with every batch of 24 samples. The quality control check sample serves as a single point check. Vary the sample weight according to the CaCO$_3$ content based on effervescence of sample (method 1B1b2d2) as follows:

7.3.1 Use a 2-g sample weight if effervescence is none, very slight, or slight.

7.3.2 Use a 1-g sample weight if effervescence is strong.

7.3.3 Use a 0.5-g sample weight if effervescence is violent.

7.4 Lubricate the O-ring of bottle cap with glycerin from a squeeze bottle.

7.5 Dispense 10 mL of 3 N HCl into a gelatin capsule and carefully place the top on the capsule. The HCl may squirt or leak out of capsule. If this happens, discard the capsule.

7.6 Place the capsule in bottle and cap bottle immediately.
7.7 Release any pressure in the bottle by piercing the stopper with a hypodermic needle. Remove the needle after ≈5 to 10 s.

7.8 After 5 to 10 min, the HCl dissolves through the capsule. Shake the bottle at a rate of 140 rpm on the shaker for the first 10 min and the last 10 min of a 1-h interval at room temperature (20 ±2 °C). After this 1 h, measure the pressure in the bottle by piercing the stopper of the cap with a hypodermic needle connected to the manometer.

7.9 Autozero the manometer before taking readings. Record the manometer readings (mm Hg).

<20-mm Basis

7.10 Determine carbonate content of the 2- to 20-mm fraction on a fine-grind (<180 µm) air-dry sample by the above method.

7.11 The carbonate in the 2- to 20-mm fraction and in the <2-mm fraction are combined and converted to a <20-mm soil basis.

8. Calculations

8.1 Correct the manometer readings as follows:

\[
CR = (MR - BR)
\]

where:

CR = Corrected reading

MR = Manometer reading

BR = Blank reading

Three blanks are run with each batch of 24 samples. The average of three blanks is used as BR.

8.2 Calculate the regression equation for the corrected manometer readings. Use the CaCO₃ weights as the dependent variable (regressed or predicted values) and the corresponding manometer readings as the independent variable.

8.3 Use the corrected (CR) linear regression (slope, intercept) equation to estimate % CaCO₃ in the sample as follows:

\[
CCE = \left[ \frac{(CR \times \text{Slope} + \text{Intercept})}{\text{Sample Weight (g)}} \right] \times \text{AD/OD}
\]

where:

CCE = Calcium Carbonate Equivalent (%) in <2-mm fraction or 2- to 20-mm fraction

CR = Corrected manometer reading

AD/OD = Air-dry/oven-dry ratio (method 3D1)
8.5 Carbonate = \((A \times B) + [C \times (1-B)]\)

where:
Carbonate = Carbonate as CaCO\(_3\) on a <20-mm basis (%)
A = \(\text{CaCO}_3\) in <2-mm fraction (%)
B = Weight of the <20-mm fraction minus the weight of the 2- to 20-
mm fraction divided by the weight of <20-mm fraction (methods
1B2b2f and 3A2).
C = \(\text{CaCO}_3\) in 2- to 20-mm fraction (%)

9. Report
Report \(\text{CaCO}_3\) equivalent as a percentage of oven-dry soil to the nearest whole
number.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
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origin of calcium carbonates on the isotopically-exchangeable phosphate in
Carbonate and Gypsum (4E)
Aqueous Extraction (4E2)
  Precipitation in Acetone (4E2a)
  Conductivity Bridge (4E2a1)
  Electrical Conductivity (4E2a1a)
  Gypsum, Qualitative and Quantitative (4E2a1a1)
    Air-Dry, <2 mm or <20 mm (4E2a1a1a1-2)

1. Application

If the electrical conductivity of a soil sample is ≥0.50 dS cm\(^{-1}\) by method 4F1a1a1, gypsum content is routinely determined by the KSSL. Gypsum content of a soil is a criterion for gypsic and petrogypsic horizons and for mineralogical class at the family level (Soil Survey Staff, 2014). Soil subsidence through solution and removal of gypsum can crack building foundations, break irrigation canals, and make roads uneven. Failure can be a problem in soils with as little as 1.5% gypsum (Nelson, 1982). The gypsum content in the soil may be used to determine if reclamation of sodic soils requires chemical amendments. Corrosion of concrete is also associated with soil gypsum.

Gypsum formation by precipitation of calcium sulfate (CaSO\(_4\)) is typically greatest at the surface layers. Gypsum from deposits that have a high content of gypsum is typically greatest in the lower part of the soil profile. However, leaching may disrupt this sequence. Gypsum is reported on both a <2- and a <20-mm basis.

2. Summary of Method

A soil sample is mixed with water to dissolve gypsum. Acetone is added to a portion of the clear extract to precipitate the dissolved gypsum. After centrifuging, the gypsum is re-dissolved in water. The electrical conductivity (EC) of the solution is read. The EC reading is used to estimate the gypsum content in meq 100 g\(^{-1}\).

In method 4E2a1a1, gypsum content (meq 100 g\(^{-1}\)) is converted to percent gypsum (uncorrected). The percent gypsum (uncorrected) is used to calculate percent gypsum (corrected). The percent gypsum (corrected) is used to correct the AD/OD (air-dry/oven-dry ratio). The AD/OD and corrected AD/OD are determined in methods 3D1 and 3D3, respectively. The corrected AD/OD uses the correction for the crystal water of gypsum. Gypsum content on a <2-mm basis is reported in method 4E2a1a1a1.

Gypsum content may also be determined on the 2- to 20-mm fraction prepared by method 1B1b2f1a1. The gypsum determined on the 2- to 20-mm fraction and the gypsum determined on the fine earth (prepared by method 1B1b2d2) are
combined and converted to a <20-mm soil basis. Gypsum on a <20-mm basis is reported in method 4E2a1a1a2.

3. Interferences

Loss of the precipitated gypsum is the most significant potential error. Care in handling the precipitated gypsum is required. Incomplete dissolution of gypsum is also possible. In soils with large gypsum crystals, use fine-ground samples to reduce sampling errors.

When present in sufficiently high concentrations, the sulfates of Na and K are also precipitated by acetone. The concentration limits for sulfates of Na and K are 50 and 10 meq L\(^{-1}\), respectively.

4. Safety

Acetone is highly flammable. Avoid open flames and sparks. Use a nonsparking centrifuge. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Proper use and appropriate load balance of the centrifuge is required. Follow standard laboratory safety precautions.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Bottles, 250-mL, with caps, Wheaton
5.3 Mechanical reciprocating shaker, 200 oscillations min\(^{-1}\), Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Dispenser, 100-mL and 10-mL
5.5 Pipette, 20-mL, solvent, Pistolpet, Manostat
5.6 Pipette, 10-mL, electronic digital, with tips, polypropylene, 10-mL
5.7 Centrifuge, Sorvall, GLC-1, General Laboratory Centrifuge
5.8 Centrifuge tubes, 15-mL, plain, conical
5.9 Thermometer, 0 to 100 °C
5.10 Conductivity bridge and conductivity cell, Markson Model 1056, Amber Science, Eugene, Oregon
5.11 Filter paper, folded, 185-mm diameter, Whatman 2V
5.12 Funnel, 90-cm
5.13 Flask, Erlenmeyer, 250-mL
5.14 Vortexer, mini, MV1, VWR Scientific Products

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight or at least 4 h in
oven (105 °C). Dissolve 0.7456 g of dry reagent grade KCl in RO water and bring to 1-L volume. Conductivity at 25 °C is 1.41 mmhos cm⁻¹.

6.3 Acetone, purity 99%

7. Procedure
7.1 Weigh 5.0 g of fine-grind, air-dry soil into a 500-mL Wheaton bottle to the nearest mg. If a trace of gypsum is present, a 20-g sample size may be used. If the EC reading is >0.85 mmhos cm⁻¹, repeat the procedure using a smaller sample size (2.5, 1, 0.5, 0.25, or 0.1 g).

7.2 Use a dispenser to add 100 mL of RO water to sample and 1 blank.

7.3 Cap the bottle and shake at an oscillating rate of 200 oscillations min⁻¹ for 30 min at room temperature (20 ±2 °C).

7.4 Filter the suspension. The first few mL of filtrate is usually cloudy and should be discarded. Collect the clear filtrate in a 250-mL flask.

7.5 Pipette 5 mL of filtrate into 15-mL conical centrifuge tube.

7.6 Use a solvent dispenser to add 5 mL acetone.

7.7 Cap tube with a polyethylene stopper and mix.

7.8 Carefully release pressure within tube by loosening the stopper.

7.9 Let stand for at least 10 min to allow the precipitate to flocculate.

7.10 Use acetone (1 or 2 mL) to rinse stopper and inside rim of tube to prevent gypsum loss.

7.11 Remove stopper and centrifuge at 2200 rpm for 5 min.

7.12 Decant and discard supernatant. Invert and drain the tube on filter paper or on towel for 5 min.

7.13 Add 5 mL of acetone to the tube. Replace stopper. Use Vortexer to shake sample.

7.14 Carefully remove stopper and rinse it and the inside rim of tube with acetone (1 or 2 mL).

7.15 Centrifuge the sample tube at 2200 rpm for 5 min.

7.16 Discard supernatant. Drain tube upside down for 5 min.

7.17 Use a dispenser to add 10 mL of RO water to tube.

7.18 Stopper and shake with Vortexer until the precipitate dissolves.

7.19 Calibrate the EC meter and cell by drawing the 0.010 N KCl solution into the cell.

7.20 Flush the cell and fill with RO water. Digital reading should be 0.00.

7.21 Read the EC of dissolved precipitate by drawing up solution into cell and flush at least once.
7.22 If the EC reading is >0.85 mmhos cm\(^{-1}\), repeat the procedure using a smaller sample weight.

8. Calculations
8.1 Calculate % Gypsum\(_{uc}\) (gypsum, uncorrected) by using table 1 to convert EC reading (mmhos/cm) to gypsum content (meq/100 g) and proceeding with the following equation.

\[
% \text{Gypsum}_{uc} = \frac{\text{Gypsum} \times \text{Water} \times 0.08609 \times AD/OD}{\text{Sample Weight (g) \times 5}}
\]

where:
- % Gypsum\(_{uc}\) = % Gypsum in <2 mm fraction or 2- to 20-mm fraction
- Gypsum = Gypsum (meq L\(^{-1}\)). Refer to table 1.
- Water = Volume RO water (100 mL) to dissolve gypsum
- 0.08609 = Conversion factor (gypsum % = meq 100 g\(^{-1}\) x 0.08609)
- AD/OD = Air-dry/oven-dry ratio, (method 3D1)
- 5 = Filtrate (5 mL)

8.2 Table 1 converts EC (mmhos cm\(^{-1}\)) to gypsum (meq L\(^{-1}\)) for the above calculations. Use table 1 to determine gypsum content (meq L\(^{-1}\)) from the EC reading. Cross reference the EC reading using the y axis for tenths and the x axis for hundredths.

8.3 Alternatively to using table 1, calculate % Gypsum\(_{uc}\) from the following equation:

\[
\text{Result} = \frac{\{\exp(2.420384 + [1.1579713 \times \log(\text{EC} - \text{blank})]) \times \text{Water} \times 0.08609 \times \text{AD/OD}\}}{(\text{Sample Weight \times 5})}
\]

8.4 The following equation for calculation of % Gypsum\(_c\) (gypsum, corrected) assumes the crystal-water content of gypsum is 19.42% (Nelson et al., 1978) as opposed to the theoretical water content (20.21%).

\[
% \text{Gypsum}_{c} = \frac{(% \text{Gypsum}_{uc})}{[1 + (0.001942 \times % \text{Gypsum}_{uc})]}
\]

8.5 Use the % Gypsum\(_{uc}\) to recalculate the AD/OD (method 3D1). The corrected AD/OD (method 6F3) uses the correction for the crystal water of gypsum.

8.6 Calculate gypsum on <20-mm basis (method 4E2a1a1) as follows:

\[
(\%) \text{Gypsum} = AxB + [C \times (1-B)]
\]

where:
- A = Gypsum (%) in <2-mm fraction
B = Weight of the <20-mm fraction minus the 20- to 2-mm fraction divided by the weight of the <20-mm fraction
C = Gypsum (%) in the 20- to 2-mm fraction

9. Report
   Report gypsum as a percent to the nearest whole unit.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

Table 1.—Convert EC Reading (mmhos cm\(^{-1}\)) to Gypsum Content (meq L\(^{-1}\)).

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Carbonate and Gypsum (4E)
Aqueous Extraction (4E2)
   1:400, 24-h (4E2b)
Conductivity Bridge (4E2b1)
   Electrical Conductivity (4E2b1a)
   Equivalent Gypsum Content, Semiquantitative (4E2b1a1)
   Air-Dry, <2 mm or <20 mm (4E2b1a1a1-2)

1. Application
   Application of irrigated water on farmland in arid and semiarid areas poses engineering challenges related to gypsiferous/gypseous soils (Elrashidi et al., 2007). Subsidence and corrosion are also potential problems. Gypsum-related subsidence is attributed to the dissolution and removal of gypsum. Typically, gypsiferous/gypseous soils have a number of other water-soluble minerals associated with gypsum. Elrashidi et al. (2007) proposed that subsidence should not be estimated solely by gypsum content but also by other water-soluble minerals using Equivalent Gypsum Content (EGC). The EGC is defined as the quantity of both gypsum and other water-soluble minerals and is expressed as gypsum percentage (by weight) in soils. The method to estimate EGC is described herein. Refer to Elrashidi et al. (2007) for the application of EGC to estimate soil subsidence in gypsiferous/gypseous soils.

2. Summary of Method
   A 0.50-g sample is weighed and 200 mL water added. Sample is shaken for 24 h and allowed to settle for 15 min. A 20-mL sample is pipetted from the top 10-cm of the solution and filtered through a 0.45 µm disk. Electrical conductivity (EC) (1:400) is measured and recorded (dS m\(^{-1}\)).

3. Interferences
   A maximum of \(\approx 0.5\) g of gypsum can be dissolved completely in 200 mL of water, and the resulting system (2.5 g L\(^{-1}\)) is considered at a saturated state. Saturated aqueous solution of gypsum has 2.6 g L\(^{-1}\) at 25 °C (Smith and Robertson, 1962; Lagewerff et al., 1965; Van Alphen and Romero, 1971; Porta, 1998).

4. Safety
   Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents. Thoroughly wash hands after handling reagents. Refer to the Material Safety Data Sheets (MSDS) for information on the chemical make-up, use, storage,
emergency procedures, and potential health effects of the hazardous materials associated with this method.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Bottle, polyethylene, 250-mL
5.4 Tube, polyethylene, 50-mL
5.5 Pipette, 20-mL, solvent, Pistolpet, Manostat
5.6 Pipette, 10-mL, electronic digital, with tips, polypropylene, 10 mL
5.7 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.8 Conductivity bridge and conductivity cell, Markson Model 1056, Amber Science, Eugene, Oregon

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in distilled water and bring to 1-L volume. Conductivity at 25 °C is 1.4 dS m⁻¹.

7. Procedure

7.1 Weigh 0.50 g air-dry, <2-mm soil into 250-mL bottle.
7.2 Run all samples in duplicates.
7.3 Add 200 mL RO water to bottle.
7.4 Shake for 24 h at room temperature (23 ±1.0 °C).
7.5 Remove bottle from shaker and let bottle set upright 15 min, allowing soil to settle.
7.6 Pipette 20-mL sample from top 10-cm of solution and filter.
7.7 Calibrate conductivity meter using 0.010 N KCl solution.
7.8 Measure EC in filtrate.
7.9 If EC ≥ 1.0 dS m⁻¹, pipette 10 mL of soil solution and then add 20 mL distilled water into 50-mL polyethylene tube. Swirl, read, and record EC.
7.10 Rinse electrode with distilled water. Remove excess water by patting with tissue.
8. Calculations
The relationship between solution gypsum concentration (g/L) and EC of solution (dS m\(^{-1}\)) is as follows:

\[
\text{Gypsum (g L}^{-1}\text{)} = 0.998 \times \text{EC (dS m}^{-1}\text{)} = A
\]

The Soil Equivalent Gypsum Content (EGC) is calculated as follows:

\[
\text{EGC (%) = 100 \times A (g L}^{-1}\text{)} \times \text{DF} \times (200 \text{ mL/1000 mL/L}) / 0.5 \text{ g}
\]

where:
- \(\text{DF} = \text{Dilution factor. DF = 1 or 3, depending on whether dilution was necessary to determine “A.”}\)

Gypsum (%) is calculated as follows:

\[
\text{Gypsum (%) = 0.293 + [0.830 \times \text{EGC (%)}] - [0.144 \times \text{EC}_s (\text{dS m}^{-1})]}
\]

where:
- \(\text{EC}_s = \text{Electrical conductivity of saturation paste extract (dS m}^{-1}\text{) (method 4F2b1)}\)

If \(\text{EC}_s\) is unavailable, \(\text{EC}_{1:2}\) may be substituted as follows:

\[
\text{Gypsum (%) = 0.294 + [0.830 \times \text{EGC (%)}] - [0.318 \times \text{EC}_{1:2} (\text{dS m}^{-1})]}
\]

\(\text{EC}_{1:2} = \text{EC of 1:2 soil to water extract (method 4F1a1a1)}\)

9. Report
Report EC (1:400) to the nearest 0.1 dS m\(^{-1}\). Report gypsum (g L\(^{-1}\)), EGC (%), and gypsum (%).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
Electrical Conductivity and Soluble Salts (4F)
Aqueous Extraction (4F1)
  1:2 Extraction (4F1a)
    Conductivity Bridge (4F1a1)
      Electrical Conductivity (4F1a1a)
        Salt Prediction (4F1a1a1)
        Air-Dry, <2 mm (4F1a1a1a)

1. Application
   Salt prediction is used not only to predict which soils have measurable amounts of soluble salts but also to predict the quantity and appropriate dilutions for salt analyses of those soils. If salt prediction or conductivity is <0.25 mmhos cm$^{-1}$ (dS cm$^{-1}$) soils are considered nonsalty, and generally, no other salt analyses are performed on these soils by the KSSL.

2. Summary of Method
   A soil sample is mixed with water and allowed to stand overnight. The electrical conductivity (EC) of the mixture is measured using an electronic bridge. The EC by this method (4F1a1a1) is used to indicate the presence of soluble salts (U.S. Salinity Laboratory Staff, 1954).

3. Interferences
   Reverse osmosis deionized water is used to zero and flush the conductivity cell. The extract temperature is assumed to be 25 °C. If the temperature deviates significantly, a correction may be required.
   Provide airtight storage of KCl solution and samples to prevent soil release of alkali-earth cations. Exposure to air can cause gains and losses of water and dissolved gases, significantly affecting EC readings.

4. Safety
   No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment
  5.1 Electronic balance, ±1.0-mg sensitivity
  5.2 Conductivity bridge and conductivity cell, with automatic temperature adjustment, 25 ±0.1 °C, Markson Model 1056, Amber Science, Eugene, Oregon
  5.3 Plastic cups, 30-mL (1 oz), with lids, Sweetheart Cup Co. Inc., Owings Mills, MD
5.4 Dispenser, re-pipette or equivalent, 0 to 10 mL

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in RODI water and bring to 1-L volume. Conductivity at 25 °C is 1.412 mmhos cm⁻¹.

7. Procedure
7.1 Weigh 5.0 g of <2-mm, air-dry soil in a 30-mL (1 oz) condiment cup.
7.2 Add 10 mL of RO water to sample using a re-pipette dispenser.
7.3 Swirl to mix, cap, and allow to stand overnight.
7.4 Standardize the conductivity bridge using RO water (blank) and 0.010 N KCl (1.41 mmhos cm⁻¹).
7.5 Read conductance of supernatant solution directly from the bridge.
7.6 Record conductance to 0.01 mmhos cm⁻¹.

8. Calculations
8.1 No calculations are required for this procedure.
8.2 Use the following relationship to estimate the total soluble cation or anion concentration (meq L⁻¹) in the soil.
   EC (mmhos cm⁻¹) x 10 = Cation or Anion (meq L⁻¹)
8.3 Use the following relationship to estimate the total soluble cation or anion concentration (meq g⁻¹ oven-dry soil) in the soil.
   EC (mmhos cm⁻¹) x 20 = Cation (meq g⁻¹ soil)
   EC (mmhos cm⁻¹) x 20 = Anion (meq g⁻¹ soil)

9. Report
   Report prediction conductance to the nearest 0.01 mmhos cm⁻¹ (dS m⁻¹).

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References
Electrical Conductivity and Soluble Salts (4F)
Aqueous Extraction (4F1)
   Single-Point Extraction (4F1b)
      1:5, 23-h, 1-h Aqueous Extraction (4F1b1)
       Conductivity Bridge (4F1b1a)
       Electrical Conductivity (4F1b1a1)
       Air-Dry or Field-Moist, <2 mm (4F1b1a1a-b1)

1. Application
   Nutrients, particularly phosphorus and nitrogen, in runoff from agricultural land are leading causes of poor water quality in the United States (USEPA, 1996). When the environmental impact of agricultural land on natural water resources is evaluated, the amount of water-soluble elements and associated properties (e.g., pH, EC) should be measured in soil under conditions similar to those present during runoff events. In the laboratory, the soil:water system is allowed to equilibrate before extracting the soil solution. The pH, EC, and elements are then measured in the water extract. Studies at the KSSL reported a correlation between water-extractable elements for soils and their concentration in runoff from agricultural watersheds (Elrashidi et al., 2005a, 2005b).

2. Summary of Method
   The electrical conductivity (EC) of the extract (4D2a2) is measured using an electronic bridge (4F1b1a1).

3. Interferences
   Reverse osmosis deionized water is used to zero and flush the conductivity cell. The extract temperature is assumed to be 25 °C. If the temperature deviates significantly, a correction may be required.
   Provide airtight storage of KCl solution and samples to prevent soil release of alkali-earth cations. Exposure to air can cause gains and losses of water and dissolved gases, significantly affecting EC readings.

4. Safety
   No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment
   5.1 Conductivity bridge and conductivity cell, with automatic temperature adjustment, 25 ±0.1 °C, Markson Model 1056, Amber Science, Eugene, Oregon
5.2 Plastic cups, 30-mL (1 oz), with lids, Sweetheart Cup Co. Inc., Owings Mills, MD

5.3 Volumetric, 1-L

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in RODI water and bring to 1-L volume. Conductivity at 25 °C is 1.412 mmhos cm⁻¹.

7. Procedure

7.1 Prepare sample extract (procedure 4D2a2).

7.2 Standardize the conductivity bridge using RO water (blank) and 0.010 N KCl (1.41 mmhos cm⁻¹).

7.3 Read conductance of supernatant solution directly from the bridge.

7.4 Record conductance to 0.01 mmhos cm⁻¹.

8. Calculations

No calculations are required for this procedure.

9. Report

Report electrical conductance to the nearest 0.01 mmhos cm⁻¹ (dS m⁻¹).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Electrical Conductivity and Soluble Salts (4F)

Aqueous Extraction (4F1)

Subaqueous Soils (4F1c)

1:5 Aqueous Mixture by Volume (EC\(_{1:5\text{vol}}\)) (4F1c1)

Conductivity Bridge (4F1c1a)

Electrical Conductivity (4F1c1a1)

Field-Moist, Whole-Soil (4F1c1b2)

1. Application

Electrical conductivity (EC) measurements are quick, simple determinations for water-soluble salts in soils. For subaqueous soils, EC must be measured in a fresh, field-wet sample (moisture content at sample collection) that has been refrigerated or frozen because sulfides may oxidize during drying, forming sulfate salts, and thereby affecting (increasing) the EC value. The recommended EC method for subaqueous soils uses a soil to water ratio (volume) of 1:5 (EC\(_{1:5\text{vol}}\)) and is measured in the supernatant, not the extract (Soil Survey Staff, 2014). This method assumes that the salts in subaqueous soils are highly soluble chloride and sulfate salts in a dissolved state and that they include no important contributions from minerals, such as gypsum. Soil EC\(_{1:5\text{vol}}\) is used in the “Keys to Soil Taxonomy” (Soil Survey Staff, 2014) at the great group level to define freshwater subaqueous soils (Frasiwassents and Frasiwassists) from salt and brackish water subaqueous soils (Balduff, 2007; Payne, 2007). Salinity values for subaqueous soil interpretations are based on the pore-water salinity, which is what a plant root or aquatic organism experiences in situ.

2. Summary of Method

A fresh (or stored in the refrigerator) moist soil sample is mixed with 5 parts distilled water. The mixture is briefly stirred and left to equilibrate. EC\(_{1:5\text{vol}}\) is determined for unfiltered supernatant and reported as dS m\(^{-1}\). Optionally, another fresh moist sample is extracted using a vacuum pump and electrical conductivity is determined for extract and reported as dS m\(^{-1}\).

3. Interferences

Electrical conductivity increases at approximately 1.9% per degree Celsius increase in temperature (Rhoades et al., 1999). Therefore, EC needs to be expressed at a reference temperature for purposes of comparison and accurate salinity interpretations. The commonly used reference temperature is 25 °C. The best way to correct for the temperature effect on conductivity is to maintain the temperature of the sample and cell at 25 ±0.5 °C while EC is being measured. Alternatively, many EC conductivity meters correct to 25 °C.
Provide airtight storage of KCl calibration solutions. Exposure to air can cause gains and losses of water and dissolved gases, significantly affecting EC readings. Store calibration solutions in dry, dark, cool room.

Soil samples from salt water or brackish water may contain sulfides. The sulfides can oxidize to form sulfates if testing is not performed for several days and if the sample is not kept moist and either refrigerated or frozen. $EC_{1:5\text{vol}}$ is not directly comparable to EC determined by saturated paste or any other $EC_{1:5\text{vol}}$ measurement. At this time, $EC_{1:5\text{vol}}$ is only used for subaqueous soils.

4. Safety
No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment
5.1 Beakers, polypropylene, 100-mL
5.2 Stirring stick
5.3 Cylinder, graduated, 50-mL, polypropylene
5.4 Volumetric, 1-L
5.5 Conductivity bridge and conductivity cell, with automatic temperature adjustment, 25 ±0.1 °C, Markson Model 1056, Amber Science, Eugene, Oregon
5.6 Optional Equipment (if pore-water salinity determined) as follows:
   5.6.1 Vacuum pump, with tubing
   5.6.2 Buchner funnel, 56-mm
   5.6.3 Receiving tube, with 5-mL mark
   5.6.4 Filter flask, 125-mL
   5.6.5 Filter papers
   5.6.6 Spatula

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in RODI water and bring to 1-L volume. Conductivity at 25 °C is 1.412 mmhos cm$^{-1}$.

7. Procedure
7.1 Measure 10 mL of moist sample into a 100-mL beaker. Samples should be refrigerated prior to test if collected a day or two prior or should be frozen if stored longer.
7.2 Measure 50 mL RO water (5 times the volume of soil) into a second 100-ml beaker.
7.3 Pour the 50 mL RO water into the beaker containing 10 mL of soil and stir for 10 s.
7.4 Allow mixture to settle. Coarse textured samples settle in as little as 15 min. Fine-textured samples and those having a high content of organic matter may require overnight settling and should be refrigerated.
7.5 Standardize the conductivity bridge using RO water (blank) and 0.010 N KCl (1.41 mmhos cm$^{-1}$).
7.6 Allow readings to stabilize. Read conductance of supernatant solution directly from the bridge.
7.7 Record conductance of the unfiltered supernatant to 0.01 mmhos cm$^{-1}$.
7.8 Optionally, determine pore-water salinity as follows:
   7.8.1 Fill 100-mL plastic beaker approximately to 50-mL mark with soil.
   7.8.2 Connect Buchner funnel to receiving tube in beaker using the adapter.
   7.8.3 Moisten clean filter with water and place paper into Buchner funnel.
   7.8.4 Transfer moist sample into Buchner funnel. Carefully smooth sample over filter paper with spatula. Sample should cover bottom of Buchner funnel completely to depth of about $\frac{1}{2}$ in ($\approx$1.3 cm). Do not allow sample dry out.
   7.8.5 Connect vacuum pump to flask and pump to create vacuum in filter flask. Typically, about 10 pumps are sufficient to create vacuum. Pump frequently to maximize infiltration rate.
   7.8.6 Depending on soil type, drops of extract begin to collect in receiving tube. Obtain enough extract to determine the test. Filtering time can be reduced by increasing the moist sample amount in funnel.
   7.8.7 Disconnect apparatus and transfer contents in the receiving tube into another beaker. Read and record EC. Dilution of extract may be necessary.

8. Calculations
   None.

9. Report
   Report the EC$_{1:5vd}$ to the nearest 0.1 dS m$^{-1}$ (USDA–NRCS, 2013). Optionally, report EC for pore water to the nearest 0.1 dS m$^{-1}$. 
10. References


Electrical Conductivity and Soluble Salts (4F)
Saturated Paste (4F2)

Salt-affected soils, i.e., soils having excessive amounts of soluble salts and/or exchangeable sodium (ES), are common in, though not restricted to, arid and semi-arid regions. These soils are usually described and characterized in terms of the soluble salt concentrations, i.e., major dissolved inorganic solutes (Rhoades, 1982). Salt composition and distribution in the soil profile affect plant response, i.e., osmotic stress, specific ion effects, and nutritional imbalances. Soil texture and plant species also are factors in plant response to saline soils.

Traditionally, the classification of salt-affected soils has been based on the soluble salt concentrations in extracted soil solutions and on the exchangeable sodium percentage (ESP) in the associated soil (Bohn et al., 1979). In general, saline soils have been defined as having a salt content >0.1% or an EC ≥4 mmhos cm⁻¹; alkali soils have an ESP of ≥15%; and saline-alkali soils have properties of both saline and alkali soils (U.S. Salinity Laboratory Staff, 1954). In soil taxonomy, ESP and the sodium adsorption ratio (SAR) have been used as criteria for natric horizons (Soil Survey Staff, 2014). ESP and SAR are calculated in methods 4F3a1 and 4F3b, respectively.

The measurable absolute and relative amounts of various solutes are influenced by the soil to water ratio at which the soil solution extract is made. Therefore, this ratio is standardized to obtain results that can be applied and interpreted universally. Soil salinity is conventionally defined and measured on aqueous extracts of saturated soil pastes (U.S. Salinity Laboratory Staff, 1954). This soil to water ratio is used because it is the lowest reproducible ratio at which the extract for analysis can be readily removed from the soil with common laboratory equipment, i.e., pressure or vacuum, and because this soil to water ratio is often related in a predictable manner to field soil water contents (Rhoades,
Soil solutions obtained at lower soil moisture conditions are more labor intensive and require special equipment.

The KSSL measures salinity on aqueous extracts of saturated soil pastes. The saturated paste is prepared (4F2), and the saturation percentage (SP) determined (4F2a1). The saturated paste extract is obtained with an automatic extractor (4F2c1). Electrical conductivity and soil resistivity of saturated paste are measured in methods 4F2b1 and 4F2b2, respectively. The saturated paste pH is measured in method 4C1a1a2. The water-soluble cations of Ca\(^{2+}\), Mg\(^{2+}\), K\(^{+}\), and Na\(^{+}\) are measured by atomic absorption spectrophotometry in methods 4F2c1a1-4, respectively. The water-soluble anions of Br, Cl\(^{-}\), F\(^{-}\), NO\(_3\)^\(^{-}\), NO\(_2\)^\(^{-}\), PO\(_4\)^\(^{3-}\), and SO\(_4\)^\(^{2-}\) are measured by ion chromatography in methods 4F2c1b1a1-7, respectively. The carbonate and bicarbonate concentrations are determined by acid titration methods 4F2c1c1a1-2, respectively. Estimated total salt is calculated in method 4F3c. The KSSL also performs a salt prediction test (method 4F1a1a1), which is used not only to predict those soils that have measurable amounts of soluble salts but also to predict the quantity and the appropriate dilutions for salt analyses of those soils. If salt predictions or conductance are <0.25 mmhos cm\(^{-1}\), soils are considered nonsalty, and generally, no other salt analyses are performed on these soils by the KSSL.

The SP, i.e., the amount of moisture in the saturated paste, is an important measurement. An experienced analyst should be able to repeat the saturated paste preparation to an SP within 5%. The SP can be related directly to the field moisture range. Measurements on soils, over a considerable textural range (U.S. Salinity Laboratory Staff, 1954), indicate the following general rules of thumb.

\[
\begin{align*}
\text{SP} & \approx 4 \times \text{15-bar water} \\
\text{SP} & \approx 2 \times \text{upper-end field soil moisture content} \\
\text{AWC} & \approx \text{SP}/4
\end{align*}
\]

where:

- SP = Saturation percentage
- AWC = Available water capacity

Therefore, at the upper (saturated) and lower (dry) ends of the field moisture range, the salt concentration of the soil solution is \(\approx 4x\) and \(2x\) the concentration in the saturation extract, respectively.

If the soil texture is known and the 15-bar water content has been measured, the preceding SP relationships may be redefined (U.S. Salinity Laboratory Staff, 1954) as follows:
The electrical conductivity of the saturated paste (EC$_s$) is measured and is commonly reported as resistivity (R$_s$). Because of the preparation of the saturated soil paste, the EC$_s$ measurement requires more time than the R$_s$ measurement. However, the EC$_s$ is the easier measurement from which to make interpretations, i.e., EC$_s$ is more closely related to plant response (U.S. Salinity Laboratory Staff, 1954). Furthermore, there is a limited correlation between EC$_s$ and R$_s$, because the relationship is markedly influenced by variations in SP, salinity, and soil mineral conductivity. The EC$_s$ has been related to R$_s$ (U.S. Laboratory Staff, 1954) by the following equation:

$$EC_s \approx 0.25/R_s$$

where:

0.25 = Constant for Bureau of Soils electrode cup

Historically, the EC$_s$ is adjusted to a 60 °F (15.5 °C) basis before interpretative use. EC$_s$ and R$_s$ increase ≈2% per °C. The KSSL determines EC$_s$ and R$_s$ in methods 4F2b1 and 4F2b2, respectively. The unit EC x 10$^3$ is called mmhos cm$^{-1}$.

The EC$_s$ (mmhos cm$^{-1}$) may be used to estimate the salt percentage (P$_{sw}$) in solution (U.S. Salinity Laboratory Staff, 1954) as follows:

$$P_{sw} \approx 0.064 \times EC_s \text{ (mmhos cm}^{-1}\text{)}$$

The preceding equation may be used to estimate the salt percentage in the soil (P$_{ss}$) (U.S. Salinity Laboratory Staff, 1954) as follows:

$$P_{ss} = \frac{P_{sw} \times SP}{100}$$

The EC$_s$ (mmhos cm$^{-1}$) may be used to estimate the osmotic potential (OP) in atmospheres of a solution (U.S. Salinity Laboratory Staff, 1954) as follows:

$$OP \approx 0.36 \times EC_s \text{ (mmhos cm}^{-1}\text{)}$$

The EC$_s$ (mmhos cm$^{-1}$) may be used to estimate the total cation or anion concentration (meq L$^{-1}$) of the solution (U.S. Salinity Laboratory Staff, 1954) as follows:

$$\text{Total cations} \approx 10 \times EC_s \text{ (mmhos cm}^{-1}\text{)}$$

<table>
<thead>
<tr>
<th>15-Bar Water (%)</th>
<th>Texture</th>
<th>Relationship</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0 to 6.5</td>
<td>Coarse</td>
<td>SP $\approx$6⅔x15 bar</td>
</tr>
<tr>
<td>6.6 to 15</td>
<td>Medium</td>
<td>SP $\approx$4x15 bar</td>
</tr>
<tr>
<td>$&gt;$15</td>
<td>Fine</td>
<td>SP $\approx$3¼x15 bar</td>
</tr>
<tr>
<td>$&gt;$15</td>
<td>Organic</td>
<td>SP $\approx$3⅔x15 bar</td>
</tr>
</tbody>
</table>
Total anions \( \approx 10 \times EC_s \) (mmhos cm\(^{-1}\))

where:

\( EC_s \) at 25 °C

A means of cross-checking chemical analyses for consistency and reliability is provided by the interrelations that exist among the various soil chemical determinations (U.S. Salinity Laboratory Staff, 1954). The saturated paste pH is the apparent pH of the soil:water mixture and is a key indicator in many of these interrelations. The saturated paste pH is dependent upon the dissolved CO\(_2\) concentration; moisture content of the mixture; exchangeable cation composition; soluble salt composition and concentration; and the presence and amount of gypsum and alkaline-earth carbonates. Some general rules of thumb that apply to the saturated paste (U.S. Salinity Laboratory Staff, 1954) are as follows:

**Total Cation and Anion Concentrations**
- Total cations \( \approx \) Total anions, expressed on equivalent basis

**pH and Ca and Mg Concentrations**
- Concentrations of Ca\(^{2+}\) and Mg\(^{2+}\) are seldom \( >2 \) meq L\(^{-1}\) at pH >9.

**pH and Carbonate and Bicarbonate Concentrations**
- Carbonate concentration (meq L\(^{-1}\)) is measurable only if pH >9.
- Bicarbonate concentration is rarely \( >10 \) meq L\(^{-1}\) in absence of carbonates.
- Bicarbonate concentration is seldom \( >3 \) or \( 4 \) meq L\(^{-1}\) if pH <7.

**Gypsum**
- Gypsum is rarely present if pH >8.2.
- Gypsum has variable solubility in saline solutions (20 to 50 meq L\(^{-1}\)).
- Check for the presence of gypsum if Ca concentration \( >20 \) meq L\(^{-1}\) and pH \( \leq 8.2 \).

**pH, ESP, and Alkaline-Earth Carbonates**
- Alkaline-earth CO\(_3\)\(^{-}\) and ESP \( \geq 15 \) are indicated if pH \( \geq 8.5 \).
- ESP \( \leq 15 \) may or may not be indicated if pH <8.5.
- No alkaline-earth CO\(_3\)\(^{-}\) are indicated if pH <7.5.

**pH and Exchangeable Acidity**
- Significant amounts of exchangeable acidity are indicated if pH <7.0.

The commonly determined soluble cations and anions in the saturation extract include calcium, magnesium, sodium, potassium, chloride, sulfate, nitrate, fluoride, carbonate, bicarbonate, and nitrite. The less commonly analyzed cations and
anions include iron, aluminum, manganese, lithium, strontium, rubidium, cesium, hydronium, phosphate, borate, silicate, bromide, selenate, selenite, arsenate, and arsenite.

The effect of soluble cations upon the exchangeable cation determination is to increase the cation concentration in the extracting solution, i.e., \( \text{NH}_4\text{OAc} \), buffered at pH 7.0 (method 4B1a1b1-4). The dissolution of salts by the extractant necessitates an independent determination of soluble cations and a correction to the exchangeable cations. Therefore, in soils with soluble salts or carbonates, the soluble cations (meq L\(^{-1}\) solution) must be measured separately and the results subtracted from the extractable bases for determination of exchangeable bases as follows:

\[
\text{Exchangeable} = \text{Extractable} - \text{Soluble}
\]

The presence of alkaline-earth carbonates prevents accurate determination of exchangeable Ca and Mg.

**References**


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**Electrical Conductivity and Soluble Salts (4F)**

**Saturated Paste (4F2)**

- **Gravimetric (4F2a)**

- **Water Percentage (4F2a1)**

- **Air-Dry, <2 mm (4F2a1a1)**

**1. Application**

The saturated soil paste is a particular mixture of soil and water, i.e. the soil paste glistens as it reflects light, flows slightly when the container is tipped, and slides freely and cleanly from a spatula unless the soil has a high clay content. This soil to water ratio is used because it is the lowest reproducible ratio for which enough extract for analysis can be readily removed from the soil with pressure or vacuum, and because this ratio is often related in a predictable manner to the field soil water content (U.S. Salinity Laboratory Staff, 1954). Upon preparation
of a saturated paste, an aqueous extract is obtained, which is used in a series of chemical analyses, e.g., electrical conductivity and concentrations of the major solutes.

2. Summary of Method

A saturated paste is prepared (method 4F2) by adding water to a soil sample while stirring the mixture until the soil paste meets the saturation criteria, i.e. the soil paste glistens as it reflects light; flows slightly when the container is tipped; and slides freely and cleanly from a spatula unless the soil has a high clay content. The mixture is covered and allowed to stand overnight. The saturation criteria are then rechecked. If the mixture fails to meet these criteria, more water or soil is added until criteria are met. A saturated paste subsample is used to determine the moisture content, i.e., saturation percentage (SP) by method 4F2a1.

3. Interferences

Special precautions must be taken for peat and muck soils and very fine or very coarse-textured soils (Rhoades, 1982). Dry peat and muck soils, especially if coarse textured or woody, require an overnight wetting to obtain a definite end point for the saturated paste. After the first wetting, pastes of these soils usually stiffen and lose their glisten. However, upon adding water and remixing, the paste usually retains the saturated paste characteristics. With fine-textured soils, enough water should be added immediately, with a minimum of mixing, to bring the sample nearly to saturation. Care also should be taken not to overwet coarse-textured soils. The presence of free water on the surface of the paste after standing is an indication of oversaturation in the coarse-textured soils (Rhoades, 1982).

4. Safety

Use heat-resistant gloves to remove hot moisture cans from the oven. No other significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Aluminum cans, drying
5.2 Spatulas, stainless steel, hardwood handles
5.3 Electronic balance, ±1-mg sensitivity
5.4 Oven, thermostatically controlled, 110 °C
5.5 Thermometer, 0 to 200 °C
5.6 Plastic food containers, 1920-mL (16 oz), with recessed lids, Sweetheart Products Group, Owings Mills, MD
6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

7. Procedure

**Saturated Soil Paste Preparation (4F2)**

7.1 Place a <2-mm, air-dry, 250-g soil sample in the food container. This sample size is convenient to handle with the 1920-mL (16 oz) food containers and provides enough extract for most purposes. The sample size varies with the number of determinations to be made upon the paste or saturation extract.

7.2 Add enough RO water to bring the sample nearly to saturation. To reduce soil puddling and to obtain a more definite end point of the saturation criteria, mix with a minimum of stirring. Soils puddle easily when worked at a moisture content near field capacity. If the paste becomes too wet, add more dry soil.

7.3 Occasionally tap the container on the workbench to consolidate the soil:water mixture. At saturation, the soil paste glistens as it reflects light, flows slightly when the container is tipped, and slides freely and cleanly off the spatula unless the soil has a high clay content.

7.4 Cover the container and allow the sample to stand overnight.

7.5 Recheck saturation criteria, i.e., ordinarily, free water should not collect on the soil surface, paste should not stiffen markedly, and paste should not lose its glisten upon standing.

7.6 If the paste does not meet the saturation criteria, remix the paste with more RO water or dry soil. Allow to stand for at least 4 h and recheck the saturation criteria.

**Saturation Percentage Determination (4F2a1)**

7.7 Tare a moisture can and cover. Label each moisture can with the appropriate sample number.

7.8 Add ≈20 to 40 g of the saturated soil paste to the moisture can.

7.9 Cover the can, weigh the can plus sample, and record the weight to the nearest mg.

7.10 Remove the can cover, place the can in a vented drying oven at 110 °C, and leave in the oven overnight (12 to 16 h). A drying period of 24 h or longer is recommended. Do not place moist samples in the oven with other samples that are drying, unless these samples have been in the oven at least 12 to 16 h. Do not overcrowd the drying oven with samples.
7.11 Remove the cans from the oven and cover immediately. Allow the cans to cool for 1 h.

7.12 Weigh the oven-dry paste sample and record the weight. Before calculating the SP, subtract the tare weights from the saturated paste and oven-dry weights. Do not use the SP subsample for other analyses.

8. Calculations

\[ SP = \left( \frac{W_{SP} - W_{OD}}{W_{OD}} \right) \times 100 \]

where:

- \( SP \) = Saturation percentage
- \( W_{SP} \) = Weight of saturated paste
- \( W_{OD} \) = Weight of oven-dry soil

9. Report

Report the saturation percentage to the nearest 0.1%.

10. Precision and Accuracy

Precision and accuracy are available from the KSSL upon request.

11. References


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Electrical Conductivity and Soluble Salts (4F)

Saturated Paste (4F2)

Conductivity Bridge (4F2b)

Electrical Conductivity (4F2b1)

Air-Dry, <2 mm (4F2b1a1)

1. Application

The electrical conductivity of the saturation extract (EC<sub>s</sub>) is used as a criterion for classifying a soil as saline. Other uses of this measurement include the estimation of the total cation concentration in the extract, salt percentage in
solution ($P_{sw}$), salt percentage in soil ($P_{ss}$), and osmotic pressure (OP). The unit $EC \times 10^3$ is called the mmhos cm$^{-1}$. For solutions with a low $EC_s$, i.e., dilute solutions, the $EC_s$ (mmhos cm$^{-1}$) $\times 10 = \text{cation concentration (meq L}^{-1})$ (U.S. Salinity Laboratory Staff, 1954). The $EC_s$ (mmhos cm$^{-1}$) $\times 0.064 = (P_{sw})$; the ($P_{sw} \times \text{SP}$)/100 $= P_{ss}$; and the $EC_s$ (mmhos cm$^{-1}$) $\times 0.36 = \text{OP}$ in atmospheres (U.S. Salinity Laboratory Staff, 1954).

2. Summary of Method

The $EC_s$ of the saturation extract that is prepared in method 4F2 is measured using a conductivity cell and a direct reading digital bridge (method 4F2b1). The cell constant is set using a standard solution.

3. Interferences

Reverse osmosis deionized water is used to zero and flush the conductivity cell. The extract temperature is assumed to be 25 °C. If the temperature deviates significantly, a correction may be required. Provide airtight storage of KCl solution and samples to prevent soil release of alkali-earth cations. Exposure to air can cause gains and losses of water and dissolved gases, significantly affecting EC readings.

4. Safety

No significant hazards are associated with this procedure. Follow standard laboratory safety procedures.

5. Equipment

5.1 Conductivity bridge and conductivity cell, with automatic temperature adjustment, 25 ±0.1 °C, Markson Model 1096, Amber Science, Eugene, Oregon

6. Reagents

6.1 Reverse osmosis (RODI) water, ASTM Type I grade of reagent water, for rinse and KCl preparation

6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in RODI water and bring to 1-L volume. Conductivity at 25 °C is 1.412 mmhos cm$^{-1}$.

7. Procedure

7.1 Calibrate the conductivity meter and cell by drawing the 0.010 N KCl solution into the cell.
7.2 Set the meter to “D” scale and adjust the digital reading to “1.41”.
7.3 Flush the cell and fill with RODI water. Verify that digital reading is “0.00”.
7.4 Read the electrical conductivity of saturation extract (EC$_s$) by drawing up
the extract into the cell and flush at least once if the cell has not been dried.
Draw up extract a second time. Reading is started on the “C” scale. Higher
readings may require the use of the “D” or “E” scales.
7.5 When the reading has stabilized, record the EC$_s$. Rinse the cell with RODI
water and ensure that the conductivity reading falls to zero.

8. Calculations
   No calculations are required for this procedure.

9. Report
   Report EC$_s$ to the nearest 0.01 mmhos cm$^{-1}$ (dS m$^{-1}$).

10. Precision and Accuracy
    Precision and accuracy data are available from the Soil Survey upon request.

11. References
    U.S. Salinity Laboratory Staff. 1954. L.A. Richards (ed.) Diagnosis and
    improvement of saline and alkali soils. 160 p. USDA Handb. 60. U.S. Govt.
    Print. Office, Washington, DC.

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**Electrical Conductivity and Soluble Salts (4F)**

**Saturated Paste (4F2)**
- Conductivity Bridge (4F2b)
- Resistivity (4F2b2)
  - Air-Dry, <2 mm (4F2b2a1)

**1. Application**

The resistivity of the soil paste is mainly used to estimate the salt content in
the soil. The apparatus is simple and rugged, the measurements can be made
quickly, and the results are reproducible. Many agencies use the Bureau of Soils
electrode cup to estimate the soluble salt content in soils (Davis and Bryan, 1910;
Soil Survey Staff, 1951).

There is no simple method to convert saturation extract electrical conductivity
to soil paste resistivity or vice versa. There is a limited correlation between EC$_s$
and $R_s$ because the relationship is markedly influenced by variations in SP, salinity,
and soil mineral conductivity.
2. **Summary of Method**

A saturated paste that is prepared in method 4F2 is placed in an electrode cup. The resistance is measured (method 4F2b2). The temperature of the paste is measured. The resistance (ohms) is converted to a 60 °F (15.5 °C) basis using a fourth-order equation (Benham, 2003).

3. **Interferences**

No significant interferences are known to affect the saturated paste resistivity measurement.

4. **Safety**

No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. **Equipment**

5.1 Conductivity bridge, Standard Wheatstone, Model RC 16B2, Beckman Instruments, Inc.
5.2 Soil cup cell holder, soil cup Cel-M, Industrial Instruments, Inc.
5.3 Bureau of Soils electrode cup. Cell constant is defined as 0.25.
5.4 Thermometer, 0 to 100 °C

6. **Reagents**

No reagents or consumables are used in this procedure.

7. **Procedure**

7.1 Fill the electrode cup with the saturated paste that is prepared in method 4F2. Gently tap the cup to remove air bubbles. Level the soil paste by striking off the excess with a spatula.
7.2 Place the cup in the cell holder. Make sure that the surfaces of the cup and holder are clean and bright. Use steel wool or fine sandpaper to carefully clean the surfaces.
7.3 Set the conductivity meter to 1000 cycles s$^{-1}$ and adjust the multiplier range and the dial control to obtain the most distinct butterfly pattern on the fluorescent tube.
7.4 Measure resistivity. Adjustment of the sensitivity control also may be necessary.
7.5 Record the resistivity.
7.6 Place a thermometer in the saturated paste. When the temperature is stabilized, record the temperature.
8. Calculations

Use table 1 to convert measured resistance to specific resistance at 60 °F (15.5 °C).

Resistivity (ohms cm⁻¹) = ohms @ 60°F x electrode cup cell factor.

Alternatively, the following equation may be used reducing soil paste resistance readings to values at 60 °F with final results reported to 4 significant figures.

\[
A = (-0.013840786 + 0.028627073 B - 0.00037976971 B^2 + 3.7891593 e^{-06} B^3 - 1.2020657 e^{-08} B^4) \times C \times D \times E
\]

where:

A = Resistance (ohms) corrected to 60 °F
B = Temperature (°F) at which the resistance was measured
C = Resistance (ohms) measured at temperature B
D = Electrode cup cell factor
E = Scale (range multiplier)

9. Report

Report saturated paste resistivity in units of ohms at 60 °F (15.5 °C) to the nearest whole number.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Electrical Conductivity and Soluble Salts (4F)
Saturated Paste (4F2)
  Saturated Paste Extraction (4F2c)
  Automatic Extractor (4F2c1)

1. Application
   The saturated paste is operationally defined so that it may be reproduced by a
   trained analyst using limited equipment. The saturated paste extract derived from
   the saturated paste is an important aqueous solution because many soil properties
   have been related to the composition of the saturation extract, e.g., soluble salt
   composition and electrical conductivity. These soil properties or characteristics
   are related in turn to plant response to salinity (U.S. Salinity Laboratory Staff,
   1954).

2. Summary of Method
   The saturated paste (prepared in 4F2) is transferred to a plastic filter funnel
   fitted with filter paper. The funnel is placed on a mechanical vacuum extractor
   (Holmgren et al., 1977), and the saturated paste is extracted (4F2c1). The extract
   is used in subsequent chemical analyses, e.g., water-soluble cations (method
   4F2c1a1-4) and water-soluble anions (methods 4F2c1b1a1-7 and 4F2c1c1a1-2).

3. Interferences
   Some saturated pastes are difficult to extract because of soil dispersion and
   puddling. Repeated extractions may be necessary to obtain sufficient extract.
   High speed centrifuging or filtration of the extract also may be necessary. If the
   extract is to be stored for an extended period, sodium hexametaphosphate may
   be added to prevent calcium carbonate precipitation in the extract.

4. Safety
   No significant hazards are associated with this procedure. Follow standard
   laboratory procedures.

5. Equipment
   5.1 Mechanical vacuum extractor, 24-place, Sampletek, Mavco Industries,
       Lincoln, NE
   5.2 Paste extraction cups, 9-cm diameter, for mechanical vacuum extractors
   5.3 Syringes, disposable, 60-mL, polypropylene, for extraction
   5.4 Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm (⅛ ID x 1/16 OD x 1 in)
   5.5 Polycons, Richards Mfg. Co.
   5.6 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.7 Filter paper, 3- and 9-cm diameter, Whatman 40 or equivalent

6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

7. Procedure
7.1 Prepare the saturated paste extract cup to receive the saturated paste (method 4E3) by placing a 3-cm diameter filter paper circle over the center of the cup followed by two 9-cm diameter filter paper circles. Slightly moisten the filter paper to ensure that it remains in place. Ensure no air is trapped between the two 9-cm filter paper circles.
7.2 Place the extraction syringe on the lower disk of the mechanical vacuum extractor.
7.3 Use a clamp to close rubber tubing on the bottom of the paste extraction cup. Carefully transfer the saturated paste into the extraction cup. Gently tap the cup to remove entrapped air in the paste. Place cups on the extractor, connect the syringe, and remove the clamp.
7.4 When all cups are ready to extract, place a plastic cover over the extraction cup to retard evaporation.
7.5 Turn on the extractor. Set the extraction time to ≈1 h.
7.6 When the extractor stops, turn off the power.
7.7 If sufficient extract has been obtained, pull the plunger of the syringe down. Do not pull plunger from the barrel of the syringe. Carefully remove the syringe containing the extract. Leave the rubber tubing on the sample tube.
7.8 If insufficient extract has been obtained, re-extract by repositioning the extractor to its starting configuration. Remove excess air in the syringe and restart the extractor. The extraction may need to be slowed to an overnight extraction to obtain sufficient extract. Alternate methods are to extract any “unused” saturated paste in a new extraction cup or to re-extract by removing the top “moist” paste from the extraction cup, mixing with any “unused” paste, and re-extracting with a clean extraction cup or centrifuging.
7.9 Filtering the saturation extract is recommended to prevent the development of microorganisms. Connect the syringe to a 0.45-µm diameter syringe filter and express the extract into a polycon. If extracts are not to be determined immediately after collection, then store samples at 4 °C.

8. Calculations
   No calculations are required for this procedure.

9. Report
   None.
10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Electrical Conductivity and Soluble Salts (4F)
Saturated Paste (4F2)
  Saturated Paste Extraction (4F2c)
  Automatic Extractor (4F2c1)
    Atomic Absorption Spectrophotometry (4F2c1a)
      Calcium, Magnesium, Potassium, and Sodium (4F2c1a1-4)
        Air-Dry, <2 mm (4F2c1a1-4a1)

1. Application

The commonly determined soluble cations are Ca$^{2+}$, Mg$^{2+}$, K$^+$, and Na$^+$. In soils with a low saturation pH, measurable amounts of Fe and Al may be present. Determination of soluble cations is used to obtain the relations between total cation concentration and other properties of saline solutions, such as electrical conductivity and osmotic pressure (U.S. Salinity Laboratory Staff, 1954). The relative concentrations of the various cations in the soil-water extracts also provide information on the composition of the exchangeable cations in the soil. Complete analyses of the soluble ions provide a means to determine total salt content of the soils and salt content at field moisture conditions.

2. Summary of Method

The saturation extract from method 4F2c1 is diluted with an ionization suppressant (La$_2$O$_3$). The analytes are measured by an atomic absorption spectrophotometer (AAS). The data are automatically recorded by a computer and printer. The saturation extracted cations, Ca$^{2+}$, Mg$^{2+}$, K$^+$, and Na$^+$, are reported in meq L$^{-1}$ (mmol (+) L$^{-1}$) in methods 4F2c1a1-4, respectively.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the analysis of these cations. These interferences vary in importance, depending upon the particular analyte selected. Do not use borosilicate tubes because of potential leaching of analytes.
4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the AAS.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Atomic absorption spectrophotometer (AAS), double-beam, AAnalyist 400, Perkin-Elmer Corp., Norwalk, CT
5.3 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.4 Computer, with AA WinLab software, Perkin-Elmer Corp., Norwalk, CT, and printer
5.5 Single-stage regulator, acetylene
5.6 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.7 Plastic test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.8 Containers, polyethylene
5.9 Peristaltic pump

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated 12 N
6.3 HCl, 1:1 HCl:RODI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part RODI water.
6.4 Stock lanthanum ionization suppressant solution (SLISS), 65,000 mg L⁻¹. Wet 152.4 g of lanthanum oxide (La₂O₃) with 100 mL RODI water. Slowly and cautiously add 500 mL of 6 N HCl to dissolve the La₂O₃. Cooling the solution is necessary. Dilute to 2 L with RODI water. Filter solution. Store in polyethylene container.
6.5 Working lanthanum ionization suppressant solution (WLISS), 2000 mg L⁻¹. Dilute 61.5 mL of SLISS with 1800 mL of RODI water (1:10). Make up to
volume with RODI water. Invert to mix thoroughly. Store in polyethylene container.

6.6 Primary Stock Standards Solution (PSSS), high purity, 1000 mg L$^{-1}$: Ca, Mg, K, and Na.

6.7 Working Stock Mixed Standards Solution (WSMSS) for Ca, Mg, and K. In a 500-mL volumetric flask, add 250 mL Ca PSSS, 25 mL Mg PSSS, and 100 mL K PSSS = 500 mg L$^{-1}$ Ca, 50 mg L$^{-1}$ Mg, and 200 mg L$^{-1}$ K. Dilute to volume with RODI water. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator.

6.8 Mixed Calibration Standards Solution (MCSS), High, Medium, Low, Very Low, and Blank as follows:

6.8.1 MCSS High Standard (1:100): Dilute WSMSS 1:100 with WLISS. Invert to mix thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 5 mg L$^{-1}$ Ca, 0.5 mg L$^{-1}$ Mg, and 2 mg L$^{-1}$ K.

6.8.2 MCSS Medium Standard (1:200): To a 100-mL volumetric flask, add 50 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 2.5 mg L$^{-1}$ Ca, 0.25 mg L$^{-1}$ Mg, and 1 mg L$^{-1}$ K.

6.8.3 MCSS Low Standard (1:400): To a 100-mL volumetric flask, add 25 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 1.25 mg L$^{-1}$ Ca, 0.125 mg L$^{-1}$ Mg, and 0.5 mg L$^{-1}$ K.

6.8.4 MCSS Very Low Standard (1:600): To a 100-mL volumetric flask, add 16.65 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 0.83 mg L$^{-1}$ Ca, 0.08 mg L$^{-1}$ Mg, and 0.33 mg L$^{-1}$ K.

6.8.5 MCSS Blank: 0 mL of Ca, Mg, and K. Dilute RODI water 1:100 with WLISS.
6.9 Na Calibration Standards Solution (NaCSS), High, Medium, Low, and Very Low as follows:

6.9.1 NaCSS High Standard (1:100): Dilute Na PSMSS (1000 mg L\(^{-1}\)) 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 10 mg L\(^{-1}\) Na.

6.9.2 NaCSS Medium Standard (1:200): In a 50-mL volumetric, add 25 mL of Na PSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 5 mg L\(^{-1}\) Na.

6.9.3 NaCSS Low Standard (1:400): In a 50-mL volumetric flask, add 12.5 mL of PSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 2.5 mg L\(^{-1}\) Na.

6.9.4 NaCSS Very Low Standard (1:600) In a 50-mL volumetric flask, add 8.35 mL of PSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate before use. Final concentration is 1.67 mg L\(^{-1}\) Na.

6.9.5 NaCSS Blank: 0 mL Na PSMSS. Dilute RODI water 1:100 with WLISS.

6.10 Compressed air with water and oil traps

6.11 Acetylene gas, purity 99.6%

7. Procedure

**Dilution of Calibration Standards and Sample Extracts**

7.1 The 10-mL syringe is for diluent (WLISS). The 1-mL syringe is for the MCSS and saturation sample extracts (method 4F2c1). Set the digital diluter at a 1:100 dilution. See reagents for the preparation of the MCSS and the NaCSS. Dilute the saturation extract sample with 100 parts of WLISS (1:100).

7.2 Dispense the diluted sample solutions into test tubes that have been placed in the sample holders of the sample changer.
AAS Set-up and Operation

7.3 Refer to the manufacturer’s manual for operation of the AAS. The following are only very general guidelines for instrument conditions for the various analytes.

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<th>Slit $(mm)$</th>
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7.4 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

AAS Calibration and Analysis

7.5 Calibrate the instrument by using the MCSS and NaCSS. The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.6 If sample exceeds calibration standard, the sample is diluted 1:5, 1:20, 1:100, etc., with RODI water followed by 1:100 dilution with WLISS.

7.7 Perform one quality control (QC) (Low Standard) for every 12 samples. If reading is not within 10%, the instrument is re-calibrated and QC re-analyzed.

7.8 Record analyte readings to 0.01 mg $L^{-1}$.

8. Calculations

The instrument readings for analyte concentration are in mg $L^{-1}$. These analyte concentrations are converted to meq $L^{-1}$ as follows:

$$\text{Analyte Concentration in Soil (meq } L^{-1}) = \frac{(A \times B)}{C}$$

where:

$A =$ Analyte $(Ca^{2+}, Mg^{2+}, K^+, Na^+)$ concentration in extract $(mg \ L^{-1})$

$B =$ Dilution ratio, if needed

$C =$ Equivalent weight

where:

$Ca^{2+} = 20.04 \text{ mg meq}^{-1}$

$Mg^{2+} = 12.15 \text{ mg meq}^{-1}$
9. Report

Report the saturation extraction cations of Ca$^{2+}$, Mg$^{2+}$, Na$^{+}$, and K$^{+}$ to the nearest 0.1 meq L$^{-1}$ (mmol (+) L$^{-1}$).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Electrical Conductivity and Soluble Salts (4F)

Saturated Paste (4F2)

Saturated Paste Extraction (4F2c)

Automatic Extractor (4F2c1)

Ion Chromatograph 4F2c1b

Conductivity Detector (4F2c1b1)

Self-Regeneration Suppressor (4F2c1b1a)

Bromide, Chloride, Fluoride, Nitrate, Nitrite, Phosphate, and Sulfate (4F2c1b1a1-7)

Air-Dry, <2 mm (4F2c1b1a1-7a1)

1. Application

The soluble anions that are commonly determined in saline and alkali soils are carbonate, bicarbonate, sulfate, chloride, nitrate, nitrite, fluoride, phosphate, silicate, bromide, and borate (Khym, 1974; U.S. Salinity Laboratory Staff, 1954). Carbonate and bicarbonate are determined by titration. Phosphate, silicate, bromide, borate, and aluminate are found only occasionally in measurable amounts in soils. Chloride, sulfate, nitrate, fluoride, and nitrite are measured in solution by chromatography. In saline and alkali soils, carbonate, bicarbonate, sulfate, and chloride are the anions that are found in the greatest abundance. In general, soluble sulfate is usually more abundant than soluble chloride.

2. Summary of Method

The soil saturation extract is diluted according to its electrical conductivity (EC$\text{\textsubscript{s}}$). The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to measure the anion species...
and content. Standard anion concentrations are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. A computer program automates these actions. The saturation extract anions, Br\(^-\), Cl\(^-\), F\(^-\), NO\(_3^-\), NO\(_2^-\), PO\(_4^{3-}\), and SO\(_4^{2-}\), are reported in meq L\(^{-1}\) (mmol (−) L\(^{-1}\)) in procedures 4F2c1b1a1-7, respectively. This same method may also be used for water analysis.

3. Interferences

Some saturation extracts contain suspended solids. Filtering after dilution removes the particles. Saturation extracts of acid soils that contain Fe and/or Al may precipitate and clog the separator column. Saturation extracts of very high pH may contain organic material that may clog or poison the column. Organic anions that have low molecular weight will co-elute with inorganic anions from the column.

4. Safety

Wear protective clothing and safety glasses. Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.

5. Equipment

5.1 Ion chromatograph, double-column, conductivity detection, Dionex ICS2000, Dionex Corp., Sunnyvale, CA
5.2 Guard column, IonPac AG-18, 4 x 50 mm, Dionex Corp., Sunnyvale, CA
5.3 Analytical column, IonPac AG-18, Dionex Corp., Sunnyvale, CA
5.4 Self-regeneration suppressor, ASRS-300, 4-mm, Dionex Corp., Sunnyvale, CA
5.5 Autosampler, AS-40, Dionex Corp., Sunnyvale, CA
5.6 Computer with PeakNet software and printer
5.7 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.8 Poly-vials with caps, 5-mL, Dionex Corp., Sunnyvale, CA

6. Reagents

6.1 Reverse osmosis deionized filtered (RODI), ASTM Type I grade of reagent water
6.2 Helium gas
6.3 Primary stock standards solutions, (PSSS\(_{1000}\)), high purity, 1000 mg L\(^{-1}\): Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\).
6.4 Mixed calibration standards solutions (MCSS), A, B, C, and D and Blank as follows:

6.4.1 MCSSA: In a 500-mL volumetric flask, add as follows:

- 6.4.1.1 32 mL Cl\(^-\) PSSS\(_{1000}\) = 64 mg L\(^{-1}\)
- 6.4.1.2 32 mL SO\(_4^{2-}\) PSSS\(_{1000}\) = 64 mg L\(^{-1}\)
- 6.4.1.3 2 mL F\(^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
- 6.4.1.4 8 mL NO\(_3^-\) PSSS\(_{1000}\) = 16 mg L\(^{-1}\)
- 6.4.1.5 2 mL NO\(_2^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
- 6.4.1.6 2 mL Br\(^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
- 6.4.1.7 2 mL PO\(_4^{3-}\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)

Dilute to volume with RODI water and invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.2 MCSSB: In a 100-mL volumetric flask, add 50 mL MCSSA and dilute to volume with RODI water. Final concentrations are 32, 32, 2, 8, 2, 2, and 2 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.3 MCSSC: In a 100-mL volumetric flask, add 50 mL MCSSB and dilute to volume with RODI water. Final concentrations are 16, 16, 1, 4, 1, 1, and 1 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.4 MCSSD: In a 100-mL volumetric flask, add 50 mL MCSSC and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 8, 8, 0.5, 2, 0.5, 0.5, and 0.5 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.5 MCSS Blank: 0 mL of Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\). Dilute RODI water to volume.

7. Procedure

Dilution of sample extracts

7.1 To estimate the total soluble anion concentration (meq L\(^{-1}\)), multiply the EC\(_s\) (procedure 4F2b1) by 10. Subtract the CO\(_3^{2-}\) and HCO\(_3^-\) concentrations (procedures 4F2c1c1a1-2) from the total anion concentration. The remainder is the approximate concentration (meq L\(^{-1}\)) of anions to be separated by ion chromatography.

\[
\text{Anion concentration (meq L}^{-1}\text{)} = (\text{EC}_s \times 10) - (\text{HCO}_3^- + \text{CO}_3^{2-})
\]

7.2 Dilute the saturation extract with the RODI water as follows:
7.3 Place the MCSS (A, B, C, D and Blank) and diluted extract samples in the poly-vials and cap with filter caps.

**Set-up and Operation of Ion Chromatograph (IC)**

7.4 Refer to the manufacturer’s manual for the operation of chromatograph. Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex ICS-2000 ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. Ranges and/or (typical settings) are as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range and/or (Typical Setting)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calibration</td>
<td>Peak height or (area)</td>
</tr>
<tr>
<td>Flow setting</td>
<td>0.5 to 4.5 mL min⁻¹ (1.00 mL min⁻¹)</td>
</tr>
<tr>
<td>Pressure</td>
<td>200 to 3000 psi (2200 to 2400 psi)</td>
</tr>
<tr>
<td>Detection</td>
<td>Suppressed conductivity</td>
</tr>
<tr>
<td>Total conductivity</td>
<td>0 to 999.9 µS</td>
</tr>
<tr>
<td>Injection volume</td>
<td>10 µL</td>
</tr>
<tr>
<td>Auto offset</td>
<td>-999.9 to 999.9 µS (On)</td>
</tr>
<tr>
<td>Cell temperature</td>
<td>35 ºC</td>
</tr>
<tr>
<td>Suppressor current</td>
<td>75 mA</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ECₜ</th>
<th>Dilution Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>(dS cm⁻¹)</td>
<td></td>
</tr>
<tr>
<td>0.00 to 0.54</td>
<td>1</td>
</tr>
<tr>
<td>0.55 to 0.66</td>
<td>2</td>
</tr>
<tr>
<td>0.70 to 1.13</td>
<td>5</td>
</tr>
<tr>
<td>1.14 to 1.47</td>
<td>10</td>
</tr>
<tr>
<td>1.48 to 2.10</td>
<td>20</td>
</tr>
<tr>
<td>2.11 to 4.00</td>
<td>60</td>
</tr>
<tr>
<td>4.01 to 8.83</td>
<td>100</td>
</tr>
<tr>
<td>8.84 to 11.8</td>
<td>150</td>
</tr>
<tr>
<td>11.9 to 26.5</td>
<td>250</td>
</tr>
<tr>
<td>26.6 to 38.7</td>
<td>400</td>
</tr>
<tr>
<td>38.8 to 80.6</td>
<td>1000</td>
</tr>
<tr>
<td>&gt;80.7</td>
<td>2000</td>
</tr>
</tbody>
</table>
7.5 Load the sample holder cassettes with the capped samples, standards, and check samples.

7.6 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

IC Calibration and Analysis

7.7 Calibrate the instrument by using the MCSS (A, B, C, D, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.8 If samples are outside calibration, dilute sample extracts with RODI water solution and re-analyze.

7.9 Perform one quality control (QC) (Low Standard MCSS, Standard C) for every 12 samples. If reading is not within tolerance limits (10 to 15%, based on analyte), the instrument is re-calibrated and QC re-analyzed.

7.10 Record analyte readings to 0.01 mg L$^{-1}$.

8. Calculations

   The instrument readings for analyte concentration are in mg L$^{-1}$. These analyte concentrations are converted to meq L$^{-1}$ as follows:

   \[
   \text{Analyte Concentration in Soil (meq L}^{-1}) = \frac{(A \times B)}{C}
   \]

   where:

   \(A\) = Analyte (Br$^-$, Cl$^-$, F$^-$, NO$_3^-$, NO$_2^-$, PO$_4^{3-}$, and SO$_4^{2-}$) concentration in extract (mg L$^{-1}$)

   \(B\) = Dilution ratio, if needed

   \(C\) = Equivalent weight

   \[
   \begin{align*}
   \text{Cl}^- &= 35.45 \text{ mg meq}^{-1} \\
   \text{SO}_4^{2-} &= 48.03 \text{ mg meq}^{-1} \\
   \text{F}^- &= 19.00 \text{ mg meq}^{-1} \\
   \text{NO}_3^- &= 62.00 \text{ mg meq}^{-1} \\
   \text{NO}_2^- &= 46.00 \text{ mg meq}^{-1} \\
   \text{Br}^- &= 79.90 \text{ mg meq}^{-1} \\
   \text{PO}_4^{3-} &= 31.66 \text{ mg meq}^{-1}
   \end{align*}
   \]

9. Report

   Report the saturation extraction anions (Cl$^-$, SO$_4^{2-}$, F$^-$, NO$_3^-$, NO$_2^-$, Br$^-$, and PO$_4^{3-}$) to the nearest 0.1 meq L$^{-1}$ (mmol (−) L$^{-1}$).
10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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Electrical Conductivity and Soluble Salts (4F)

Saturated Paste (4F2)

Saturated Paste Extraction (4F2c)

Automatic Extractor (4F2c1)

Automatic Titrator (4F2c1c)

Combination pH-Reference Electrode (4F2c1c1)

Acid Titration, H₂SO₄ (4F2c1c1a)

Carbonate and Bicarbonate (4F2c1c1a1-2)

Air-Dry, <2 mm (4F2c1c1a1-2a)

1. Application

The water soluble anions that usually are determined in saturation extracts are carbonate, bicarbonate, sulfate, chloride, nitrate, nitrite, fluoride, phosphate, silicate, bromide, and borate. Carbonate and bicarbonate are analyzed by titration. In saturation extracts, carbonate is measurable if the pH >9 (U.S. Salinity Laboratory Staff, 1954). The bicarbonate concentration is seldom >10 meq L⁻¹ in the absence of carbonate anions (U.S. Salinity Laboratory Staff, 1954). The bicarbonate concentration at pH ≤7 seldom exceeds 3 or 4 meq L⁻¹ (U.S. Salinity Laboratory Staff, 1954).

The total dissolved ion amounts generally increase with increasing soil moisture content. While some ions increase, some ions may decrease. The carbonate and bicarbonate anions are among those ions that are most dependent upon soil moisture. Therefore, in making interpretations about carbonate and bicarbonate in soil solution, the chemistry of the soil and the soil solution must be carefully considered.

2. Summary of Method

An aliquot of the saturation extract (method 4F2c) is titrated on an automatic titrator to pH 8.25 and pH 4.60 end points. The carbonate and bicarbonate are calculated from the titers, aliquot volume, blank titer, and acid normality (methods 4F2c1c1a1-2, respectively). Carbonate and bicarbonate are reported in meq L⁻¹ (mmol (+) L⁻¹).
3. Interferences

Clean the electrode by rinsing with distilled water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

Slow electrode response time may cause over shooting the end point. A combination of slowing the buret speed and increasing the time delay may help. Cleaning the electrode with detergent may decrease the response time. If all else fails, changing the electrode generally solves the problem. Blanks may not titrate properly because some sources of reverse osmosis (RO) water have a low pH.

4. Safety

Wear protective clothing and eye protection. Exercise care when preparing reagents. Thoroughly wash hands after handling reagents. Restrict the use of concentrated H\textsubscript{2}SO\textsubscript{4} to the fume hood. Use showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow the manufacturer’s safety precautions when operating the automatic titrator.

5. Equipment

5.1 Automatic titrator, Metrohm 670 Titroprocessor, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.

5.2 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.

5.3 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

6.2 Helium gas

6.3 Sulfuric acid (H\textsubscript{2}SO\textsubscript{4}), concentrated, 36 \textit{N}

6.4 H\textsubscript{2}SO\textsubscript{4}, 0.0240 \textit{N} standardized. Carefully dilute 2.67 mL of concentrated H\textsubscript{2}SO\textsubscript{4} in 4 L of RODI degassed water (≈15 min). Re-standardize the acid at regular intervals. Refer to the procedure for standardization of acids.

6.5 Borax pH buffers, pH 4.00, 7.00, and 9.18, for titrator calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Pipette 3 mL of the fresh saturation extract (method 4E3c) into a 250-mL titration beaker.

7.2 Add 72 mL of RO water into a titration beaker. Final volume is 75 mL for blanks and samples. Run 8 to 12 blanks of RO water through the titration procedure.
7.3 Refer to manufacturer’s manual for operation of the automatic titrator.

7.4 Calibrate automatic titrator with pH 9.18, 7.00 and 4.00 buffers. Set-up the automatic titrator to set end point titration mode. The “Set” pH parameters are listed as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ep₁</td>
<td>pH 8.25</td>
</tr>
<tr>
<td>Dyn change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s⁻¹</td>
</tr>
<tr>
<td>Time delay</td>
<td>10 s</td>
</tr>
<tr>
<td>Ep₂</td>
<td>pH 4.60</td>
</tr>
<tr>
<td>Dyn change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s⁻¹</td>
</tr>
<tr>
<td>Temp</td>
<td>25 °C</td>
</tr>
<tr>
<td>Stop Volume</td>
<td>35 mL</td>
</tr>
</tbody>
</table>

7.5 Place the 250-mL titration beakers in the sample changer.

7.6 Press “Start.”

7.7 If the titrator is operating properly, no other analyst intervention is required. The titers and other titration parameters are recorded on the titroprocessor printer.

8. Calculations

8.1 \[ \text{CO}_3^{2-} \text{ (meq L}^{-1}\text{)} = (2T₁ x N x 1000)/\text{Aliquot} \]

8.2 \[ \text{HCO}_3^- \text{ (meq L}^{-1}\text{)} = [(T₂ + T₁) - \text{Blank} - (2 x T₁) x N x 1000]/\text{Aliquot} \]

where:
- \( T₁ \) = Titer of \( \text{CO}_3^{2-} \) (mL)
- \( T₂ \) = Titer of \( \text{HCO}_3^- \) (mL)
- \( N \) = Normality of \( \text{H}_2\text{SO}_4 \)
- \( \text{Blank} \) = Average titer of blank solutions (mL)
- \( \text{Aliquot} \) = Volume of saturation extract titrated (mL)
- 1000 = Conversion factor to meq L⁻¹

9. Report

Report saturation extract \( \text{CO}_3^{2-} \) and \( \text{HCO}_3^- \) to the nearest 0.1 meq L⁻¹ (mmol (−) L⁻¹).
10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Electrical Conductivity and Soluble Salts (4F)
Ratios and Estimates Related to Soluble Salts (4F3)
Exchangeable Sodium Percentage (ESP), NH$_4$OAc, pH 7.0 (4F3a)
ESP, Calculated without Saturated Paste Extraction (4F3a1)

Compute the exchangeable sodium percentage (ESP) by dividing the exchangeable sodium (ES) by the CEC by NH$_4$OAc, pH 7.0 (CEC–7) and multiplying by 100 (method 4F3a1). The ES is calculated by subtracting the water soluble Na$^+$ determined in method 4F2c1a4 from the NH$_4$OAc extractable Na$^+$ determined in method 4B1a1b4 (U.S. Salinity Laboratory Staff, 1954). The CEC–7 is determined in method 4B1a1a1a1. In soil taxonomy, an ESP ≥15% is a criterion for natric horizons (Soil Survey Staff, 2014). When the saturation extract is not prepared, the ESP is calculated as follows:

$$ESP = \frac{ES}{CEC–7} \times 100$$

where:

ESP=Exchangeable sodium percentage
ES=Extractable sodium (NH$_4$OAc extractable Na$^+$, (cmol (+) kg$^{-1}$)).
CEC–7=CEC by NH$_4$OAc, pH 7.0 (cmol (+) kg$^{-1}$).

Electrical Conductivity and Soluble Salts (4F)
Ratios and Estimates Related to Soluble Salts (4F3)
Exchangeable Sodium Percentage (ESP), NH$_4$OAc, pH 7.0 (4F3a)
ESP, Calculated with Saturated Paste Extraction (4F3a2)

Exchangeable Na is computed with acceptable accuracy unless salt contents >20 mmhos cm$^{-1}$ (dS m$^{-1}$) at 25˚C. Exchangeable Na equals extractable Na minus saturation-extract Na multiplied by saturation percentage. Saturation percentage is the water percentage in the saturated paste divided by 1000. Exchangeable Na can be determined with greater accuracy than the other cations in the presence of gypsum or carbonates. If exchangeable K is negligible compared to exchangeable Ca and Mg, then exchangeable Ca plus Mg equals CEC (NH$_4$OAc, pH 7.0) minus exchangeable Na. This approximation
is suitably reproducible for comparison between soils and for soil classification. Exchangeable Ca can be computed in the same manner as exchangeable Na. Results are not as satisfactory for exchangeable Ca when computed in the presence of carbonates or large amounts of gypsum.

When the saturation extract is prepared, the KSSL calculates the ESP by method 4F3a2 as follows:

$$ESP = 100 \times \frac{Na_{ex} - [Na_{ws} \times (H_2O_{ws}/1000)]}{CEC-7}$$

where:
- ESP = Exchangeable sodium percentage
- $Na_{ex}$ = Extractable Na (NH₄OAc extractable Na⁺, (cmol (+) kg⁻¹))
- $Na_{ws}$ = Water-soluble Na (mmol (+) L⁻¹)
- $H_2O_{ws}$ = Water saturation percentage
- CEC-7 = CEC by NH₄OAc, pH 7.0 (cmol (+) kg⁻¹)
- 1000 = Conversion factor to (cmol (+) kg⁻¹)
- 100 = Conversion factor to percent

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**Electrical Conductivity and Soluble Salts (4F)**

**Ratios and Estimates Related to Soluble Salts (4F3)**

**Sodium Adsorption Ratio (SAR) (4F3b)**

Compute the sodium adsorption ratio (SAR) by dividing the molar concentration of the monovalent cation Na⁺ by the square root of the molar concentration of the divalent cations Ca²⁺ and Mg²⁺ (U.S. Salinity Laboratory Staff, 1954). The water soluble Ca²⁺, Mg²⁺, and Na⁺ are determined in methods 4F2c1a1, 4F2c1a2, and 4F2c1a4, respectively. The SAR was developed as a measurement of the quality of irrigation water, particularly when the water is used for irrigating soils that are salt- or sodium-affected (U.S. Salinity Laboratory Staff, 1954). In soil taxonomy, a SAR ≥13 is a criterion for natric horizons (Soil Survey Staff, 2014). The KSSL calculates the SAR by method 4F3b. The SAR is calculated as follows:

$$SAR = \frac{[Na^+]}{\sqrt{[Ca^{2+}] + [Mg^{2+}]}}$$

where:
- SAR = Sodium Adsorption Ratio
- Na⁺ = Water soluble Na⁺ (mmol (+) L⁻¹)
- Ca²⁺ = Water soluble Ca²⁺ (mmol (+) L⁻¹)
- Mg²⁺ = Water soluble Mg²⁺ (mmol (+) L⁻¹)
References


Electrical Conductivity and Soluble Salts (4F)
Ratios and Estimates Related to Soluble Salts (4F3)
Estimated Total Salt (4F3c)

Use the charts and graphs available in U.S. Salinity Laboratory Staff (1954) to estimate total salt content (4F3c) from the electrical conductivity ($EC_s$) of the saturation extract (method 4F2b1). The essential relations are summarized in the equation as follows:

$$\text{Total Salt in Soil (ppm)} = [-4.2333 + (12.2347 \times EC_s) + (0.058 \times EC_s^2) - (0.0003 \times EC_s^3)] \times 0.000064 \times SP$$

where:
- $EC_s$ = Electrical conductivity of saturation extract
- $SP$ = Saturation percentage of saturation paste

Previous equations used to estimate total salt content are as follows:

$$\log \text{Total Salt in Soil (ppm)} = 0.81 + [1.08 \times \log EC_s \text{ (mmhos cm}^{-1})] + \log SP$$

where:
- $EC_s$ = Electrical conductivity of saturation extract
- $SP$ = Saturation percentage of saturation paste

$$\text{Total Salt in Soil } (%) = \text{Total salt (ppm)} \times 10^{-4}$$

These equations are applicable to saturation extracts with an $EC_s < 20$ mmhos cm$^{-1}$. Deviations occur at higher concentrations of salt.

References


Selective Dissolutions (4G)

Background: Over the years, various terms have been used to describe broad groupings of soil components, e.g., crystalline phyllosilicates, amorphous,
poorly crystalline, paracrystalline, noncrystalline, allophane, imogolite, short-
range-order minerals (SROMs), etc. (Jackson et al., 1986). These groupings
have been related, in part, to various laboratory analyses and have thereby been
operationally defined quantitatively and semiquantitatively by these analyses
(Jackson et al., 1986). Some of these analytical procedures include x-ray
diffraction analysis and selective chemical dissolutions, e.g., dithionite-citrate,
sodium pyrophosphate, and ammonium oxalate extractions. These terms have
not been used consistently in the literature. In addition, there is not always a
clear delineation between dissolution data, either conceptually or operationally.
For more detailed discussion of these various soil terms and the application of
selective chemical extractions, refer to Wada (1989) and Soil Survey Staff (2011).

Selective dissolution data have been used extensively in the study of the
noncrystalline material content of soils and sediments. There are, however,
limitations to using these data. In general, there exists a continuum of crystalline
order, ranging from no long-range order through paracrystalline and poorly
crystalline to well crystalline (Follet et al., 1965). Selective dissolution data are
necessary for independent determinations of various inorganic constituents of
soils. They are necessary because many physical analytical methods have
difficulty in estimating (or even recognizing) the presence of noncrystalline
and paracrystalline free oxides or aluminosilicates mixed with crystalline soil
components (Jackson et al., 1986). In general, the crystalline free oxides
and phyllosilicates of soils can be identified qualitatively and estimated
semiquantitatively by x-ray diffraction analysis. Those soils containing hydroxyls
(-OH groups), e.g., kaolinite, gibbsite, and goethite, can sometimes be determined
quantitatively by differential thermal analysis (DTA), differential scanning
colorimetry (DCS), and thermogravimetric analysis (TGA). Refer to additional
discussion on x-ray diffraction and thermal analysis in the mineralogy section (7)
of this manual.

Selective chemical dissolution data present difficulties in the adequate
assessment of the portion that is extracted by particular reagents, e.g., dithionite-
citrate, sodium pyrophosphate, and ammonium oxalate. In principle, it cannot be
expected that chemical methods are able to perfectly distinguish the degrees of
crystallinity, and some caution is required in the interpretation of these analytical
data (van Wambke, 1992). Refer to Wada (1989) regarding the dissolution of
Al, Fe, and Si in various clay constituents and organic complexes by treatment
with different reagents. The KSSL routinely performs three selective chemical
dissolutions: dithionite-citrate (4G1a1-3), ammonium oxalate (4G2a1a1-5), and
sodium pyrophosphate (4G3a1-3).

Dithionite-Citrate Extraction: The original objectives of the dithionite-citrate
extraction were to determine the free Fe oxides and to remove the amorphous
coatings and crystals of free Fe oxide, which were acting as cementing agents,
for subsequent physical and chemical analysis of soils, sediments, and clay
minerals (Weaver et al., 1968; Jackson, 1969; Jackson et al., 1986). Dithionite-
citrate extractable Fe ($\text{Fe}_{\text{d}}$) is considered a measure of “free iron” in soils and, as such, is pedogenically significant. Data regarding dithionite-citrate extractable Fe are of interest in studies of soil genesis and classification because of its increasing concentration with increasing weathering and its effect on soil colors (Schwertmann, 1992). “Free iron” is also considered an important factor in P-fixation and soil aggregate stability.

**Sodium Pyrophosphate Extraction:** Sodium pyrophosphate extracting solutions tend to selectively extract mainly Fe and Al associated with organic compounds. The dithionite-citrate extractions tend to extract these compounds plus the free oxides (McKeague et al., 1971). At one time, sodium pyrophosphate extractable Fe and Al in conjunction with dithionite-citrate data were used to help identify translocated Al and Fe humus complexes in spodic horizons (Soil Survey Staff, 1975). Numerous evaluations of pyrophosphate extracts have indicated that the pyrophosphate extraction does not necessarily correlate with organic-bound Fe and Al (Schuppli et al., 1983; Kassim et al., 1984; Parfitt and Childs, 1988; Birkeland et al., 1989) as commonly thought (Schwertmann and Taylor, 1977; Parfitt and Childs, 1988). Pyrophosphate not only extracts organic-bound Fe but also peptizes solid particles of ferrihydrite and in some instances even goethite (Yuan et al., 1993). The use of sodium pyrophosphate extract data in conjunction with dithionite-citrate data as chemical requirements for spodic horizons have been replaced by other criteria (Soil Survey Staff, 2014) and, at one time, were referred to as spodic horizon criteria on the KSSL data sheets.

**Ammonium Oxalate Extraction:** In general, ammonium oxalate allowed to react in darkness has been considered to be a selective dissolution for noncrystalline materials (McKeague and Day, 1966; Higashi and Ikeda, 1974; Fey and LeRoux, 1976; Schwertmann and Taylor, 1989; Hodges and Zelazny, 1980). The ammonium oxalate procedure removes most noncrystalline and paracrystalline materials (allophane and imogolite) from soils (Higashi and Ikeda, 1974; Hodges and Zelazny, 1980) as well as short-range-ordered oxides and hydroxides of Al, Fe, and Mn (Schwertmann, 1959, 1964; McKeague and Day, 1966; McKeague et al., 1971; Fey and LeRoux, 1976). In addition, this method is assumed to extract Al+Fe humus. Opaline Si is not dissolved by this method (Wada, 1977). This procedure has been reported to dissolve very little hematite and goethite and small amounts of magnetite (Baril and Bitton, 1969; McKeague et al., 1971; Walker, 1983). There have been conflicting data on the effect of this procedure on clay minerals, but in general, the ammonium oxalate treatment is considered to have very little effect on phyllosilicates (kaolinite, montmorillonite, and illite) or gibbsite. The ammonium oxalate extraction is assumed to dissolve selectively “active” Al and Fe components that are present in noncrystalline materials as well as associated or independent, poorly crystalline silica. The method also extracts allophane, imogolite, Al+Fe humus complexes, and amorphous or poorly crystallized oxides and hydroxides.
The intent of the ammonium oxalate procedure is to measure the quantities of poorly crystalline materials in the soil. At the present time, the ammonium oxalate extraction is considered the most precise chemical method for measuring these soil components. However, in principle it cannot be expected that chemical methods are able to perfectly distinguish degrees of crystallinity, and some caution is to be exercised in the interpretation of the analytical data (van Wambeke, 1992). A more reliable and accurate estimation of soil properties and a better understanding of noncrystallinity are provided when ammonium oxalate extraction is used in conjunction with other selective dissolution procedures, thermal techniques, and chemical tests (Jackson et al., 1986).

**Application, Ratios, and Estimates:** In a general way, the Fe\(_d\) is considered to be a measure of the total pedogenic Fe (e.g., goethite, hematite, lepidocrocite, and ferrihydrite), while the ammonium oxalate extractable Fe (Fe\(_o\)) (probably ferrihydrite) is a measure of the paracrystalline Fe (Birkeland et al., 1989). The Fe\(_o\)/Fe\(_d\) ratio is often calculated because it is considered an approximation of the relative proportion of ferrihydrite in soils (Schwertmann, 1985).

Mn\(_d\) is considered the “easily reducible Mn.” Al\(_d\) and Al\(_o\) are pedogenically significant. The Al\(_d\) represents the Al substituted in Fe oxides, which can have an upper limit of thirty-three percent mole substitution (Schwertmann et al., 1977; Schwertmann and Taylor, 1989). The Al\(_o\) is generally an estimate of the total pedogenic Al in soils dominated by allophane, imogolite, and organically-bound Al (Wada, 1977). Unlike Fe\(_d\), the Al\(_d\) extract is commonly less than the Al\(_o\) (Birkeland et al., 1989) and so does not necessarily represent the total pedogenic Al (Wada, 1977).

Allophane in soils can be estimated from the Al\(_o\) and Si\(_o\) and the pyrophosphate extractable Al (Al\(_p\)) (Parfitt and Henmi, 1982; Parfitt and Wilson, 1985; Parfitt, 1990). The Al\(_d\) represents the Al dissolved from allophane, imogolite, and Al-humus complexes, and the Al\(_p\) is the Al from the Al-humus complexes alone (Parfitt and Kimble, 1989). The Al\(_o\) minus the Al\(_p\) gives an estimate of the Al in allophane and imogolite, whereas the Si\(_o\) gives an estimate of the Si in allophane and imogolite. The (Al\(_o\) – Al\(_p\))/Si\(_o\) times the molar ratio (28/27) is an estimate of the Al/Si ratio of allophane and imogolite in the soil. The values of 28 and 27 represent the atomic weights of Si and Al, respectively.

Selective chemical dissolution data are used as taxonomic criteria for mineralogy classes, e.g., ammonium oxalate Fe and Si for the amorphous and ferrihydritic mineralogy classes and dithionite-citrate Fe for the ferritic mineralogy class. An optical density of ammonium oxalate extract (ODOE) of ≥0.25 is used as a chemical criterion for spodic materials (Soil Survey Staff, 2014). An increase in ODOE is used as an indicator of the accumulation of translocated organic materials in an illuvial horizon (Soil Survey Staff, 2014). Ammonium oxalate extractable Al\(_o\) plus 0.5 Fe\(_o\) is also used as a taxonomic criterion for andic soil properties (Soil Survey Staff, 2014). The weight of Fe atoms is approximately twice that of Al atoms. In evaluating the relative proportion of Fe and Al atoms
solubilized by ammonium oxalate, the weight percent of Fe must be divided by two, i.e., $\text{Al}_o + \frac{1}{2}\text{Fe}_o$. Refer to the Soil Survey Staff (2014, 2011) for more detailed discussion of these taxonomic criteria.

References


Selective Dissolutions (4G)
Dithionite-Citrate Extraction (4G1)
  Atomic Absorption Spectrophotometer (4G1a)
    Aluminum, Iron, and Manganese (4G1a1-3)
      Air-Dry or Field Moist, <2 mm (4G1a1-3a-b1)

1. Application

   Dithionite-citrate (CD) is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, and amorphous aluminosilicates (Wada, 1989). The CD solution is a poor extractant of crystalline hydrous oxides of Al, allophane, and imogolite. The CD solution does not extract opal, Si, or other constituents of crystalline silicate minerals (Wada, 1989).

   This extraction is also sometimes referred to as citrate-dithionite or sodium citrate-dithionite. The method (4G1a1-3) described herein is not the same extraction as described in the Soil Survey Investigations Report (SSIR) No. 1 (1972), method 6C3. This obsolete SSL method (6C3) incorporated sodium bicarbonate as a buffer (pH 7.3) in the dithionite-citrate method, resulting in a buffered neutral citrate-bicarbonate-dithionite (Aguilera and Jackson, 1953; Mehra and Jackson, 1960; Jackson, 1969), commonly referred to as the CBD method.

2. Summary of Method

   A soil sample is mixed with sodium dithionite, sodium citrate, and reverse osmosis deionized (RODI) water and shaken overnight. Solution is centrifuged, and a clear extract obtained. The CD extract is diluted with RODI water. The analytes are measured by an atomic absorption spectrophotometer (AAS). The data are automatically recorded by a computer and printer. The AAS converts absorption to analyte concentration. The percent CD extractable Al, Fe, and Mn are reported in methods 4G1a1-3, respectively.

3. Interferences

   There are four types of interferences (matrix, spectral, chemical, and ionization) in the AAS analyses of these elements. These interferences vary in importance, depending upon the particular analyte selected.

   The redox potential of the extractant is dependent upon the pH of the extracting solution and the soil system. Sodium citrate complexes the reduced Fe and usually buffers the system to a pH of 6.5 to 7.3. Some soils may lower the pH, resulting in the precipitation of Fe sulfides.

   Filtered extracts can yield different recoveries of Fe, Mn, and Al, relative to unfiltered extracts.
4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves); eye protection (face shields, goggles, or safety glasses); and a breathing filter when handling dry sodium dithionite. Sodium dithionite may spontaneously ignite if allowed to become moist, even by atmospheric moisture. Keep dithionite in a fume hood.

Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the AAS.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Atomic absorption spectrophotometer (AAS), double-beam optical system, AAnalyst, 400, Perkin-Elmer Corp., Norwalk, CT, with computer and printer
5.4 Autosampler, AS-93 Plus, Perkin-Elmer Corp., Norwalk, CT
5.5 Peristaltic pump
5.6 Single-stage regulators, acetylene and nitrous oxide
5.7 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.8 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.9 Dispenser, 30-mL
5.10 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.11 Containers, polypropylene
5.12 Volumetrics, Class A, 100-mL, 250-mL, and 1000-mL
5.13 Measuring scoop, handmade, 0.4 g calibrated
5.14 Centrifuge tubes, 50-mL

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Sodium dithionite (Na₂S₂O₄), purified powder
6.3 Sodium citrate dihydrate (Na₃C₆H₅O₇·2H₂O), crystal, reagent. Dissolve 336 g sodium citrate in approximately 1 L RODI water, followed by diluting to 2 L with RODI water. Final concentration is 0.57 M sodium citrate.
6.4 Sulfuric acid (H₂SO₄), concentrated
6.5 Phosphoric acid ($H_3PO_4$), concentrated (85%). For Fe analysis, samples are diluted 1:50 prior to analysis. A diluting solution for Fe analysis (for a final concentration of 0.5% $H_3PO_4$ in samples) may be made by adding 6.12 mL of concentrated $H_3PO_4$ to 500 mL volume of RODI water, diluting to 1000 mL, and mixing thoroughly. (Note: 1:5 sample dilutions for Al and Mn are in RODI water.)

6.6 Primary Stock Standard Solution (PSSS), high purity, 1000 mg L$^{-1}$: Fe, Mn, and Al.

6.7 Calibration standards for Fe (Section 6.8): To each 100 mL volume of blank, calibration, and quality control (QC) standards, add 25 mL of the following matrix matching mixture. This mixture is made by combining 20 mL Na citrate extracting solution, 0.21 mL $H_2SO_4$, and 6 mL $H_3PO_4$ and diluting to 250 mL volume with RODI water. Invert to mix thoroughly. (Note: Matrix of standards is prepared to match a 1:50 dilution of samples. Also, $H_2SO_4$ substitutes for dithionite).

6.8 Standard Fe Calibration Solutions (SFeCS), or working standards, (25.0, 20.0, 15.0, 10.0, 5.0, 1.0, and 0.0 mg Fe L$^{-1}$) and QC (12.5 mg L$^{-1}$). Prepare fresh weekly. In seven 100-mL volumetric flasks, add as follows:

- 6.8.1 25.0 mg Fe L$^{-1}$ = 2.5 mL PSSS$_{Fe}$
- 6.8.2 20.0 mg Fe L$^{-1}$ = 2.0 mL PSSS$_{Fe}$
- 6.8.3 15.0 mg Fe L$^{-1}$ = 1.5 mL PSSS$_{Fe}$
- 6.8.4 10.0 mg Fe L$^{-1}$ = 1.0 mL PSSS$_{Fe}$
- 6.8.5 5.0 mg Fe L$^{-1}$ = 0.5 mL PSSS$_{Fe}$
- 6.8.6 1.0 mg Fe L$^{-1}$ = 0.1 mL PSSS$_{Fe}$
- 6.8.7 0.0 mg Fe L$^{-1}$ = 0.0 mL PSSS$_{Fe}$ (blank)
- 6.8.8 12.5 mg Fe L$^{-1}$ = 1.25 mL PSSS$_{Fe}$ (QC)

Fill to volume with RODI water and invert to mix thoroughly. After dissolution, transfer solution to a plastic bottle.

6.9 Calibration standards for Mn (Section 6.10): To each 100 mL volume of blank, calibration, and quality control (QC) standards, add 25 mL of the following matrix matching mixture. This mixture is made by combining 200 mL Na citrate extracting solution, 2.1 mL $H_2SO_4$, and diluting to 250-mL volume with RODI water. Invert to mix thoroughly. (Note: Matrix of standards is prepared to match a 1:5 dilution of samples. Also, $H_2SO_4$ substitutes for dithionite).

6.10 Standard Mn Calibration Solutions (SMnCS), or working standards, (15.0, 10.0, 5.0, 2.5, 1.5, and 0.0 mg Mn L$^{-1}$) and QC (6.5 mg L$^{-1}$). Prepare fresh weekly. In six 100-mL volumetric flasks, add as follows:

- 6.10.1 15.0 mg Mn L$^{-1}$ = 1.5 mL PSSS$_{Mn}$
6.10.2 10.0 mg Mn L\(^{-1}\)=1.0 mL PSSS\(_{Mn}\)
6.10.3 5.0 mg Mn L\(^{-1}\)=0.5 mL PSSS\(_{Mn}\)
6.10.4 2.5 mg Mn L\(^{-1}\)=0.25 mL PSSS\(_{Mn}\)
6.10.5 1.5 mg Mn L\(^{-1}\)=0.15 mL PSSS\(_{Mn}\)
6.10.6 0.0 mg Mn L\(^{-1}\)=0.0 mL PSSS\(_{Mn}\) (blank)
6.10.7 6.5 mg Mn L\(^{-1}\)=0.65 mL PSSS\(_{Mn}\) (QC)

Fill to volume with RODI water and invert to mix thoroughly. After dissolution, transfer solution to a plastic bottle.

6.11 Calibration standards for Al (Section 6.12): To each 100 mL volume of blank, calibration, and quality control (QC) standards, add 25 mL of the following matrix matching mixture. This mixture is made by combining 200 mL Na citrate extracting solution, 2.1 mL H\(_2\)SO\(_4\), and then diluting to 250-mL volume with RODI water. Invert to mix thoroughly. (Note: Matrix of standards is prepared to match a 1:5 dilution of samples—same as with Mn. Also, H\(_2\)SO\(_4\) substitutes for dithionite).

6.12 Standard Al Calibration Solutions (SAICS), or working standards, (100.0, 80.0, 60.0, 40.0, 20.0, 10.0, and 0.0 mg Al L\(^{-1}\)) and QC (50.0 mg L\(^{-1}\)). Prepare fresh weekly. In seven 100-mL volumetric flasks, add as follows:

6.12.1 100.0 mg Al L\(^{-1}\)=10.0 mL PSSS\(_{Al}\)
6.12.2 80.0 mg Al L\(^{-1}\)=8.0 mL PSSS\(_{Al}\)
6.12.3 60.0 mg Al L\(^{-1}\)=6.0 mL PSSS\(_{Al}\)
6.12.4 40.0 mg Al L\(^{-1}\)=4.0 mL PSSS\(_{Al}\)
6.12.5 20.0 mg Al L\(^{-1}\)=2.0 mL PSSS\(_{Al}\)
6.12.6 10.0 mg Al L\(^{-1}\)=1.0 mL PSSS\(_{Al}\)
6.12.7 0.0 mg Al L\(^{-1}\)=0.0 mL PSSS\(_{Al}\) (blank)
6.12.8 50.0 mg Al L\(^{-1}\)=5.0 mL PSSS\(_{Al}\) (QC)

Fill to volume with RODI water and invert to mix thoroughly. After dissolution, transfer solution to a plastic bottle.

6.13 Acetylene gas, purity 99.6%
6.14 Nitrous oxide, USP
6.15 Compressed air with water and oil traps

7. Procedure

**Extraction of Al, Fe, and Mn**

7.1 Weigh 0.75 g of <2-mm or fine-grind, air-dry soil sample to the nearest mg and place in a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈0.75 g of air-dry soil.
7.2 Add 0.4 g of sodium dithionite (use one calibrated scoop) and 25 mL of sodium citrate solution.

7.3 Cap tubes and shake briefly by hand to dislodge soil from tube bottom. Place tubes in rack.

7.4 Place rack in shaker and shake overnight (12 to 16 h) at 200 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.5 Remove tubes from shaker and manually shake tubes to dislodge any soil from cap. Allow samples to sit overnight.

7.6 The following day, centrifuge at 4000 rpm for 15 min. The Fe, Mn, and Al are determined on the AAS from a clear aliquot of solution.

**Dilution of Sample Extracts**

7.7 No ionization suppressant is required because the Na in the extractant is sufficient in quantity. For a 1:50 dilution of samples for Fe analysis, use the H\(_3\)PO\(_4\) diluting solution (see Section 6.5). The dilution of Fe results in a final solution concentration of 0.5% H\(_3\)PO\(_4\). Dilute 1 part CD sample extract with 49 parts of H\(_3\)PO\(_4\) diluting solution (1:50 dilution).

7.8 A 1:5 dilution in RODI water is used for Al and Mn. Dilute 1 part CD sample extract with 4 parts RODI water.

7.9 Dispense the diluted sample solutions into test tubes that have been placed in the holders of the sample changer.

**AAS Set-up and Operation**

7.10 Refer to the manufacturer’s manual for operation of the AAS. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Concentration (mg L(^{-1}))</th>
<th>Wavelength (nm)</th>
<th>Burner head</th>
<th>Slit (mm)</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>25.0</td>
<td>248.8</td>
<td>10-cm parallel</td>
<td>0.2</td>
<td>3.0 C(_2)H(_2)/15.7 Air</td>
</tr>
<tr>
<td>Mn</td>
<td>15.0</td>
<td>279.8</td>
<td>10-cm parallel</td>
<td>0.2</td>
<td>3.0 C(_2)H(_2)/15.7 Air</td>
</tr>
<tr>
<td>Al</td>
<td>100.0</td>
<td>309.3</td>
<td>5-cm parallel</td>
<td>0.7</td>
<td>8.5 C(_2)H(_2)/15.7 N(_2)O</td>
</tr>
</tbody>
</table>

Typical read delay is 3 s, and integration time is 3 s but can vary depending on soil type. Three replicates are average for each sample.
7.11 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

7.12 The instrument readings are programmed to display analyte concentration in mg L⁻¹ (ppm).

### AAS Calibration and Analysis

7.13 Each element is analyzed during separate runs on the AAS. Use the calibration reagent blank and calibration standards to calibrate the AAS. Calibrations are linear with calculated intercept.

7.14 Use the QC after every 12th sample. It must pass within 15% to continue. If it fails, recalibrate and reread the QC. The QC is also read at the end of each run.

7.15 If samples are outside the calibration range, a serial dilution is performed. A 1:5 dilution of the sample using the calibration blank, followed by the typical dilution (1:5 dilution with RODI water for Al and Mn, and 1:50 dilution with the H₃PO₄ diluting solution for Fe). Maintain matrix match between standards and diluted samples by performing this extra dilution with calibration blank.

7.16 Record analyte readings to 0.01 unit.

### 8. Calculations

Convert analyte concentrations (mg L⁻¹) to percent in soil as follows:

\[
\text{Soil Fe, Al, Mn (\%) = } \frac{A \times B \times C \times R \times 100}{E \times 1000}
\]

where:

- \(A\) = Sample extract reading (mg L⁻¹)
- \(B\) = Extract Volume (L)
- \(C\) = Dilution, required
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- \(E\) = Sample weight (g)
- 100 = Conversion factor to 100-g basis
- 1000 = mg g⁻¹

### 9. Report

Report percent CD extractable Al, Fe, and Mn to the nearest 0.1 of a percent.

### 10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.
11. References


Selective Dissolutions (4G)
Ammonium Oxalate Extraction (4G2)
  Automatic Extractor (4G2a)
    Inductively Coupled Plasma Atomic Emission Spectrophotometer (4G2a1)
      Radial Mode (4G2a1a)
      Aluminum, Iron, Manganese, Phosphorus, and Silicon (4G2a1a1-5)
        Air-Dry or Field-Moist, <2 mm (4G2a1a1-5a-b1)
    UV-Visible Spectrophotometer, Dual Beam (4G2a2)
      Transmittance (4G2a2a)
      Optical Density (4G2a2a1)
        Air-Dry or Field-Moist, <2 mm (4G2a2a1-a-b1)

1. Application

Ammonium oxalate is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, allophane, and amorphous aluminosilicates (Wada, 1989). Ammonium oxalate is a poor extractant of imogolite and layer silicates and does not extract crystalline hydrous oxides of Fe and Al, opal, or crystalline silicate (Wada, 1989). A more reliable and accurate estimation of soil properties and a better understanding of soil exchange complex is provided when ammonium oxalate extraction is used in conjunction with other selective dissolution procedures, thermal techniques, and chemical tests. In soil taxonomy, ammonium oxalate extractable Fe and Al are criteria for andic soil properties (Soil Survey Staff, 2014). This extraction is also sometimes referred to as acid ammonium oxalate, acid oxalate, oxalate-oxallic acid, or oxalic acid-ammonium oxalate.
2. Summary of Method

A soil sample is extracted with a mechanical vacuum extractor (Holmgren et al., 1977) in a 0.2 M ammonium oxalate solution buffered at pH 3.0 under darkness. The ammonium oxalate extract is weighed. The ammonium oxalate extract is diluted with reverse osmosis deionized water. The analytes are measured by an inductively coupled plasma atomic emission spectrophotometer (ICP–AES). Data are automatically recorded by a computer and printer. The ammonium oxalate extractable Al, Fe, Mn, P, and Si are reported in methods 4G2a1a1-5, respectively. All these data are reported in percent, except Mn and P, which are reported in mg kg\textsuperscript{-1}. In method 4G2a2a1, the optical density of the extract is measured with a UV spectrophotometer at 430 nm.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the ICP analyses of these elements. These interferences vary in importance, depending on the analyte chosen.

The ammonium oxalate buffer extraction is sensitive to light, especially UV light. The exclusion of light reduces the dissolution effect of crystalline oxides and clay minerals. If the sample contains large amounts of amorphous material (>2% Al), an alternate method should be used, i.e., shaking with 0.275 M ammonium oxalate, pH 3.25, 1:100 soil:extractant.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the UV spectrophotometer and ICP–AES.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity

5.2 Mechanical vacuum extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE

5.3 Tubes, 60-mL, polypropylene, for extraction (0.45-µm filter), reservoir, and tared extraction tubes

5.4 Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm, (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels

5.5 Dispenser, 30-mL

5.6 Pipettes, electronic digital, 10,000-µL and 1000-µL, with tips, 10,000-µL and 1000-µL

5.7 Containers, polyethylene
5.8 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES),
dual-view, with high-solids nebulizer, alumina or quartz injector, Optima
7300 DV, Perkin-Elmer Corp., Norwalk, CT
5.9 Autosampler, S-10 Plus, Perkin-Elmer Corp., Norwalk, CT
5.10 Computer, with WinLab32™ software, and printer
5.11 Single-stage regulator, high-purity, high-flow, argon
5.12 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer,
Fisher Scientific
5.13 Vortexer, mini, MV1, VWR Scientific Products
5.14 Spectrophotometer, UV-Visible, Varian, Cary 50 Conc, Varian Australia Pty
Ltd.
5.15 Computer with Cary WinUV software, Varian Australia Pty Ltd., and printer
5.16 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent
water
6.2 Ammonium oxalate buffer solution, 0.2 M, pH 3.0. Solution A (base):
Dissolve 284 g of (NH$_4$)$_2$C$_2$O$_4$•2H$_2$O in 10 L of DDI water. Solution B (acid):
Dissolve 252 g of H$_2$C$_2$O$_4$•H$_2$O in 10 L of DDI water. Mix 4 parts solution A
with 3 parts solution B. Adjust ammonium oxalate solution pH by adding
either acid or base solution. Store in a polypropylene bottle.
6.3 Borax pH buffers, pH 4.00, 7.00, and 9.18 for electrode calibration,
Beckman, Fullerton, CA
6.4 Primary Fe standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher
Chemical Scientific Co., Fairlawn, NJ
6.5 Primary Al standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher
Chemical Scientific Co., Fairlawn, NJ
6.6 Primary Si standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher
Chemical Scientific Co., Fairlawn, NJ
6.7 Primary Mn standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher
Chemical Scientific Co., Fairlawn, NJ
6.8 Primary P standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher
Chemical Scientific Co., Fairlawn, NJ
6.9 High Mixed Calibration Standard. Mix 60 mL of each primary standard
(Si, Fe, and Al) with 10 mL of primary Mn standard and 20 mL of primary
P standard in 1-L volumetric flask. Add 100 mL of 0.2 M ammonium
oxalate solution and make to 1-L volume with RODI water. The elements
are added in the order (Si, Fe, Al, Mn, P) to avoid element precipitation.
Resulting solution contains $60 \text{ mg L}^{-1}$ each of Si, Fe, and Al, $10 \text{ mg L}^{-1}$ Mn, and $20 \text{ mg L}^{-1}$ P. Invert to mix thoroughly. Store in a polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.10 Medium Mixed Calibration Standard. Mix 30 mL of each primary standard (Si, Fe, and Al) with 5 mL of primary Mn standard and 10 mL of primary P standard in 1-L volumetric flask. Add 100 mL of 0.2 $M$ ammonium oxalate solution and make to 1-L volume with RODI water. The elements are added in the order (Si, Fe, Al, Mn, P) to avoid element precipitation. Resulting solution contains $30 \text{ mg L}^{-1}$ each of Si, Fe, and Al, $5 \text{ mg L}^{-1}$ Mn, and $10 \text{ mg L}^{-1}$ P. Invert to mix thoroughly. Store in a polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.11 Low Mixed Calibration Standard. Mix 10 mL of each primary standard (Si, Fe, and Al) with 2 mL of primary Mn standard, and 3 mL primary P standard in 1-L volumetric flask. Add 100 mL of 0.2 $M$ ammonium oxalate solution and make to 1-L volume with RODI water. The elements are added in the order (Si, Fe, Al, Mn, P) to avoid element precipitation. Resulting solution contains $10 \text{ mg L}^{-1}$ each of Si, Fe, and Al, $2 \text{ mg L}^{-1}$ Mn, and $3 \text{ mg L}^{-1}$ P. Invert to mix thoroughly. Store in a polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.12 Low Si Calibration Standard. Mix 5 mL of Si primary standard in 1-L volumetric flask. Add 100 mL of 0.2 $M$ ammonium oxalate solution and make to 1-L volume with RODI water. Resulting solution contains $5 \text{ mg L}^{-1}$ Si. Invert to mix thoroughly. Store in polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.13 Very Low Si Calibration Standard. Mix 2 mL of Si primary standard in 1-L volumetric flask. Add 100 mL of 0.2 $M$ ammonium oxalate solution and make to 1-L volume with RODI water. Resulting solution contains $2 \text{ mg L}^{-1}$ Si. Invert to mix thoroughly. Store in polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.14 Calibration reagent blank solution. Add 100 mL of 0.2 $M$ ammonium oxalate solution and make to 1-L volume with RODI water. Store in polyethylene bottle. Make fresh weekly. Store in a refrigerator.

6.15 Argon gas, purity 99.9%

6.16 Nitrogen, purity 99.9%

7. Procedure

**Extraction of Fe, Mn, Al, Si, and P**

7.1 Weigh 0.5 g of <2-mm, air-dry or fine-grind soil the nearest mg and place in sample tube. If sample is moist, weigh enough soil to achieve $\approx 0.5$ g. Prepare two reagent blanks (no sample in tube) per set of 48 samples.
7.2 Place labeled extraction tube (ET) on extractor and connect to corresponding tared extraction tube (TET\textsubscript{Oxalate}) with rubber tubing.

7.3 Use a dispenser to add 15.00 mL of ammonium oxalate buffer to the ET. Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min. Cover samples with black plastic bag to exclude light.

7.4 Secure reservoir tube (RT) to top of ET tube. Set extractor for 30-min extraction rate and extract until the ammonium oxalate buffer solution is at a 0.5 to 1.0-cm height above sample. Turn off extractor.

7.5 Add 35 mL of ammonium oxalate buffer to the RT.

7.6 Cover the extractor with a black plastic bag to exclude light. Adjust the extraction rate for a 12-h extraction.

7.7 After the extraction, shut off the extractor. Carefully remove TET\textsubscript{Oxalate}. Leave the rubber tubing on the ET.

7.8 Weigh each syringe containing ammonium oxalate extract to the nearest mg.

7.9 Mix extract in each TET\textsubscript{Oxalate} by manually shaking. Fill a disposable tube with extract solution. This solution is reserved for determinations of Fe, Mn, Al, Si, and P. If optical density is to be measured, fill a disposable cuvette with extract solution. Discard excess solution properly. If extracts are not to be determined immediately after collection, then store samples at 4 °C.

**Determination of Optical Density of Extract**

7.10 Place 4 mL of ammonium oxalate extract in disposable cuvette.

7.11 Place 4 mL of ammonium oxalate reagent blank in disposable cuvette.

7.12 On the spectrophotometer, select a 430-nm wavelength. Select normal slit width and height. Refer to manufacturer’s manual for operation of the spectrophotometer.

7.13 Use the ammonium oxalate reagent blank to zero spectrophotometer.

7.14 Record optical density of ammonium oxalate extract to nearest 0.001 unit.

**Dilution of Sample Extracts and Standards**

7.15 Dilute ammonium oxalate extracts (1:10) with RODI water. Add 1 part ammonium oxalate sample extract with 9 parts dilution solution. Pipette 0.7 mL of extract and 6.3 mL RODI water. Vortex. Calibration reagent blanks and calibration standards are not diluted.

7.16 Dispense the diluted solutions into test tubes that have been placed in the sample holder of the sample changer.
ICP–AES Set-up and Operation

7.17 Refer to the manufacturer’s manual for operation of the ICP–AES. The following parameters are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wash Parameters</td>
<td>Between samples</td>
</tr>
<tr>
<td>Wash rate</td>
<td>2.00 mL min(^{-1})</td>
</tr>
<tr>
<td>Wash time</td>
<td>30 sec</td>
</tr>
<tr>
<td>Background correction</td>
<td>2 point (all elements)</td>
</tr>
<tr>
<td>Read delay</td>
<td>2 sec</td>
</tr>
<tr>
<td>Replicates</td>
<td>2</td>
</tr>
<tr>
<td><strong>Plasma</strong></td>
<td></td>
</tr>
<tr>
<td>Source equilibration delay</td>
<td>20 sec</td>
</tr>
<tr>
<td>Plasma aerosol type</td>
<td>Wet</td>
</tr>
<tr>
<td>Nebulizer start-up conditions</td>
<td>Gradual</td>
</tr>
<tr>
<td><strong>Plasma</strong></td>
<td></td>
</tr>
<tr>
<td>Plasma</td>
<td>15 L min(^{-1})</td>
</tr>
<tr>
<td>Auxiliary</td>
<td>0.5 L min(^{-1})</td>
</tr>
<tr>
<td>Nebulizer</td>
<td>0.85 L min(^{-1})</td>
</tr>
<tr>
<td>Power</td>
<td>1450 Watts</td>
</tr>
<tr>
<td>View dist</td>
<td>15.0</td>
</tr>
<tr>
<td>Plasma view</td>
<td>Radial</td>
</tr>
<tr>
<td><strong>Peristaltic Pump</strong></td>
<td></td>
</tr>
<tr>
<td>Sample flow rate</td>
<td>2.00 L min(^{-1})</td>
</tr>
<tr>
<td>Sample flush time</td>
<td>35 sec</td>
</tr>
</tbody>
</table>

Nebulizer pressure depends on the type of nebulizer that is being used, i.e., low flow nebulizer requires a higher pressure whereas a higher flow nebulizer requires a lower pressure. To check for correct nebulizer pressure, aspirate with 1000.0 mg L\(^{-1}\) yttrium. Adjust pressure to correct yttrium bullet.

7.18 Analyte data are reported at the following wavelengths.
<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>259.94</td>
</tr>
<tr>
<td>Al</td>
<td>308.22</td>
</tr>
<tr>
<td>Si</td>
<td>251.61</td>
</tr>
<tr>
<td>Mn</td>
<td>257.61</td>
</tr>
<tr>
<td>P</td>
<td>213.61</td>
</tr>
</tbody>
</table>

7.19 Use the computer and printer to set instrument parameters and to collect and record instrument readings. The instrument readings are programmed in mg L\(^{-1}\).

**ICP–AES Calibration and Analysis**

7.20 Use a multipoint calibration for ICP–AES analysis of ammonium oxalate extracts. The ICP–AES calibrates the blank first, low standard, medium standard, followed by the high standard. Prepare a quality control (QC) standard with analyte concentration between the high and low calibration standards. The ICP–AES reads the QC after the high standard. If the QC falls within the range set by operator (±10%), the instrument proceeds to analyze the unknowns. If the QC is outside the range, the instrument restandardizes. The QC is analyzed approximately every 12 samples.

7.21 If sample exceeds calibration standard, dilute 1:5 (1 mL of sample extract with 4 mL 0.02 \(M\) ammonium oxalate extracting solution), followed by a 1:10 (1 mL of 1:5 solution with 9 mL RODI water). This makes for a 1:50 dilution.

7.22 Record analyte readings to the nearest 0.01 unit.

8. **Calculations (Al, Fe, Si)**

The instrument readings are the analyte concentration (mg L\(^{-1}\) Fe, Mn, Al, Si, and P). Use these values to calculate the analyte concentration in percent in the soil for Fe, Al, and Si and mg kg\(^{-1}\) for Mn and P as follows:

8.1 Soil Fe, Al, Si (%) = \(\frac{Ax[(B_1-B_2)/B_3]xC_1xC_2xR}{Ex1000x1000}\)

where:
- A = Sample extract reading (mg L\(^{-1}\))
- \(B_1\) = Weight of syringe + extract (g)
- \(B_2\) = Tare weight of syringe (g)
- \(B_3\) = Density of 0.2 \(M\) ammonium oxalate solution at 20 °C (1.007 g mL\(^{-1}\))
- \(C_1\) = Dilution, required
\[ C_2 = \text{Dilution, if performed} \]
\[ R = \text{Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)} \]
\[ E = \text{Sample weight (g)} \]
\[ 100 = \text{Conversion factor to 100-g basis} \]
\[ 1000 = \text{Factor in denominator (mL L}^{-1} \text{)} \]
\[ 1000 = \text{Factor in denominator (mg g}^{-1} \text{)} \]

8.2 Soil Mn, P (mg kg}^{-1} \text{soil}) = \{A \times \frac{(B_1 - B_2)}{B_3} \times C_1 \times C_2 \times R \times 1000\} / (E \times 1000)

where:
\[ A = \text{Sample extract reading (mg L}^{-1} \text{)} \]
\[ B_1 = \text{Weight of syringe + extract (g)} \]
\[ B_2 = \text{Tare weight of syringe (g)} \]
\[ B_3 = \text{Density of 0.2 M ammonium oxalate solution at 20 °C (1.007 g mL}^{-1} \text{)} \]
\[ C_1 = \text{Dilution, required} \]
\[ C_2 = \text{Dilution, if performed} \]
\[ R = \text{Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)} \]
\[ 1000 = \text{Conversion factor in numerator to kg-basis} \]
\[ 1000 = \text{Factor in denominator (mL L}^{-1} \text{)} \]
\[ E = \text{Sample weight (g)} \]

9. Report
Report the percent ammonium oxalate extractable Al, Fe, and Si to the nearest 0.01%. Report the concentration of ammonium oxalate extractable Mn and P to the nearest mg kg}^{-1} \text{soil}. Report the optical density of the ammonium oxalate extract to the nearest 0.01 unit.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
Jackson, M.L. 1979. Soil chemical analysis—Advanced course. 2nd ed., 12th printing. Published by author, Madison, WI.

Selective Dissolutions (4G)

Ammonium Oxalate (4G2)

Ratios and Estimates Related to Ammonium Oxalate Extraction (4G2b)

Al+½Fe (4G2b1)

The ratio using ammonium oxalate extractable Al plus ½ Fe determined in method 4G2a1a1-5 is used as taxonomic criterion for andic soil properties. Refer to Soil Survey Staff (2011, 2014) for more detailed information on the application of this ratio.

References


Selective Dissolutions (4G)

Sodium Pyrophosphate Extraction (4G3)

Atomic Absorption Spectrophotometry (4G3a)

Aluminum, Iron, and Manganese (4G3a1-3)

Air-Dry or Field-Moist, <2 mm (4G3a1-3a-b1)

1. Application

Sodium pyrophosphate (0.1 M Na₄P₂O₇) is used as a selective dissolution extractant for organically complexed Fe and Al (Wada, 1989). The Na₄P₂O₇ solution is a poor extractant for allophane, imogolite, amorphous aluminosilicates, and noncrystalline hydrous oxides of Fe and Al. The Na₄P₂O₇ solution does not extract opal, crystalline silicates, layer silicates, and crystalline hydrous oxides of Fe and Al (Wada, 1989). Sodium pyrophosphate extractable organic C, Fe, and Al were former criteria for spodic placement in soil taxonomy (Soil Survey Staff, 1975).
2. Summary of Method

The soil sample is mixed with 0.1 $M \text{Na}_4\text{P}_2\text{O}_7$ and shaken overnight. The solution is then allowed to settle overnight before centrifuging and filtering to obtain a clear extract. The analytes (Al, Fe, Mn) are measured by an atomic absorption spectrophotometer (AAS). The data are automatically recorded by a computer and printer. The AAS converts absorption to analyte concentration. Percent sodium pyrophosphate extractable Al, Fe, and Mn are reported in methods 4G3a1-3, respectively. The organic C in the sodium pyrophosphate extract is wet oxidized in a fume hood and gravimetrically measured in method 4G3b1a1.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the AAS analyses of these elements. These interferences vary in importance, depending upon the particular analyte selected.

There are several problems with this procedure, especially the peptization and dispersion of microcrystalline iron oxide by pyrophosphate (Jeanroy and Guilet, 1981). The quantity of Fe extracted with pyrophosphate decreases with increasing centrifugation (McKeague and Schuppli, 1982); therefore, uniform high-speed centrifugation or micropore filtration treatments are required (Schuppli et al., 1983; Loveland and Digby, 1984). Sodium pyrophosphate extraction works best at pH 10 (Loeppert and Inskeep, 1996). The concentration of Na$_4$P$_2$O$_7$ solution must be close to 0.1 $M$. Variable amounts of Fe, Al, Mn, and organic C may be extracted by varying the pyrophosphate concentration.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the atomic absorption spectrophotometer (AAS).

5. Equipment

5.1 Electronic balance, ±0.1-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min$^{-1}$, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Atomic absorption spectrophotometer (AAS), double-beam optical system, 
AAanalyst 400, Perkin-Elmer Corp., Norwalk, CT
5.4 Autosampler, AS-93 Plus, Perkin-Elmer Corp., Norwalk, CT
5.5 Peristaltic pump
5.6 Single-stage regulators, acetylene and nitrous oxide
5.7 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.8 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, 
Microlab 500, Hamilton Co., Reno, NV
5.9 Dispenser, 40 mL
5.10 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.11 Containers, polypropylene
5.12 Volumetrics, Class A, 100-mL, 250-mL, and 1000-mL
5.13 Centrifuge tubes, 50-mL
5.14 Filter paper, Whatman 42, 150-mm

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated, 12 N
6.3 Sodium pyrophosphate solution, 0.1 M. Dissolve 446.05 g of 
Na$_4$P$_2$O$_7$•10H$_2$O in 10 L of RODI water. Adjust pH of solution to 10.0 with 
either HCl or NaOH.
6.4 Primary Fe Standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher 
Chemical Scientific Co., Fairlawn, NJ
6.5 Primary Al Standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher 
Chemical Scientific Co., Fairlawn, NJ
6.6 Primary Mn Standard, 1000 mg L$^{-1}$. Certified Reference Solution, Fisher 
Chemical Scientific Co., Fairlawn, NJ
6.7 Mixed Calibration Standards (MCS), Fe, Al, Mn. To six 250-mL volumetrics, 
add the following amounts of Primary Standards (1000 mg L$^{-1}$) as follows. 
The elements are added in the order (Fe, Al, Mn) to avoid element 
precipitation.

6.7.1 25, 100, and 15 mg L$^{-1}$ Fe, Al, and Mn=6.25, 25.00, and 3.75 mL 
Fe, Al, and Mn, respectively.
6.7.2 20, 80, and 10 mg L$^{-1}$ Fe, Al, and Mn=5.00, 20.00, and 2.50 mL 
Fe, Al, and Mn, respectively.
6.7.3 10, 40, and 5 mg L$^{-1}$ Fe, Al, and Mn=2.50, 10.00, and 1.25 mL 
Fe, Al, and Mn, respectively.
6.7.4 5, 20, and 2.50 mg L\(^{-1}\) Fe, Al, and Mn = 1.25, 5.00, and 0.625 mL Fe, Al, and Mn, respectively.

6.7.5 1, 10, and 1.5 mg L\(^{-1}\) Fe, Al, and Mn = 0.25, 2.50, 0.375 mL Fe, Al, and Mn, respectively.

6.7.6 0, 0, and 0 mg L\(^{-1}\) Fe, Al, and Mn = 0.0, 0.0, and 0.0 mL Fe, Al, and Mn, respectively.

Add 50 mL of 0.1 \(M\) \(\text{Na}_4\text{P}_2\text{O}_7\) to each MCS and make to volume with RODI water. Final concentration of MCS is 0.02 \(M\) \(\text{Na}_4\text{P}_2\text{O}_7\). Quality control (QC) is the MCS with 10, 40, and 5 mg L\(^{-1}\) Fe, Al, and Mn, respectively.

6.8 Acetylene gas, purity 99.6%

6.9 Nitrous oxide gas, compressed

6.10 Compressed air with water and oil traps

7. Procedure

**Extraction of Al, Fe, and Mn**

7.1 Weigh 0.5 g <2-mm or fine-grind, air-dry soil to the nearest mg sample and place in a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve \(\approx 0.5\) g of air-dry soil.

7.2 Add 30-mL of 0.1 \(M\) \(\text{Na}_4\text{P}_2\text{O}_7\), pH 10.0 solution to centrifuge tube.

7.3 Cap tube and shake briefly by hand to dislodge soil from tube bottom. Place tube in rack.

7.4 Place rack in shaker and shake overnight (12 to 16 h) at 200 oscillations min\(^{-1}\) at room temperature (20 ±2 °C).

7.5 Remove tubes from shaker and manually shake tubes to dislodge any soil from cap. Allow samples to sit overnight.

7.6 The next day, centrifuge sample at 4000 rpm for 15 min. The Fe, Mn, and Al are determined from a clear aliquot of solution. Filter if necessary.

**Dilution of Sample Extracts and Standards**

7.7 No ionization suppressant is required as the Na in the extractant is present in sufficient quantity. Dilute samples 1:5 with RODI water. Samples have a final concentration of 0.02 \(M\) \(\text{Na}_4\text{P}_2\text{O}_7\).

7.8 Dispense the MCS and diluted sample solutions into test tubes that have been placed in the sample holder of the sample changer.

**AAS Set-up and Operation**

7.9 The following are only very general guidelines for instrument conditions for the various analytes.
<table>
<thead>
<tr>
<th>Analyte</th>
<th>Concentration</th>
<th>Wavelength</th>
<th>Burner head</th>
<th>Slit</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mg L(^{-1}))</td>
<td>(nm)</td>
<td></td>
<td>(mm)</td>
<td></td>
</tr>
<tr>
<td>Fe</td>
<td>25.0</td>
<td>248.8</td>
<td>10-cm parallel</td>
<td>0.2</td>
<td>3.0 C(_2)H(_2)/15.2 Air</td>
</tr>
<tr>
<td>Mn</td>
<td>15.0</td>
<td>279.8</td>
<td>10-cm parallel</td>
<td>0.2</td>
<td>3.0 C(_2)H(_2)/15.7 Air</td>
</tr>
<tr>
<td>Al</td>
<td>100.0</td>
<td>309.3</td>
<td>5-cm parallel</td>
<td>0.7</td>
<td>7.0 C(_2)H(_2)/3.5 N(_2)O</td>
</tr>
</tbody>
</table>

Typical read delay is 3 s, and integration time is 3 s but can vary depending on soil type. Three replicates are averaged for each sample.

7.10 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

7.11 If sample exceeds calibration standard, dilute the sample 1:10 with 0.1 M Na\(_4\)P\(_2\)O\(_7\) and then 1:5 with RODI water.

**AAS Calibration**

7.12 Each element is analyzed during separate runs on the AAS. Use the calibration reagent blank and calibration standards to calibrate the AAS. Calibrations are linear with calculated intercept.

7.13 Use the QC after every 12\(^{th}\) sample. It must pass within 15% to continue. If it fails, recalibrate and reread the QC. The QC is also read at the end of each run.

7.14 If samples are outside the calibration range, dilute.

7.15 Record analyte readings to 0.01 unit.

8. Calculations

8.1 Soil Fe, Al, Mn (%)= \((A \times B \times C_1 \times C_2 \times R \times 100) / (E \times 1000 \times 1000)\)

where:

- \(A\) = Analyte concentration reading (mg L\(^{-1}\))
- \(B\) = Extract volume (L)
- \(C_1\) = Dilution, required
- \(C_2\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- \(E\) = Sample weight (g)
- 100 = Conversion factor to 100-g basis
- 1000 = Factor in denominator (mg g\(^{-1}\))
- 1000 = Factor in denominator (mL L\(^{-1}\))

9. Report

Report sodium pyrophosphate extractable Fe, Mn, and Al to the nearest 0.1 of a percent.
10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


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**Total Analysis (4H)**

Acid Digestion (4H1)

HNO₃ + HCl Digestion (4H1a)

Microwave (4H1a1)

Inductively Coupled Plasma Mass Spectrophotometer (4H1a1b)

Antimony, Arsenic, Barium, Beryllium, Cadmium, Cobalt, Chromium, Copper, Lead, Manganese, Mercury, Molybdenum, Nickel, Phosphorus, Selenium, Silver, Strontium, Tin, Tungsten, Vanadium, and Zinc (4H1a1b1-21)

Air-Dry, <2 mm (4H1a1b1-21a1)

1. Application

The term “trace elements” is widely applied to a variety of elements that are generally present in plants, soils, and water in low concentrations, or
“background levels.” Knowledge of these levels is important in understanding the consequences of increasing levels of trace elements in ecosystems (Tiller, 1989; Holmgren et al., 1993). These elements may become elevated in concentration due to natural activities (e.g., magmatic activity, mineral weathering, and translocation through the soil or landscape) or human activities (e.g., pesticides, mining, smelting, and manufacturing). The relative reactivity or bioavailability of these elements in soils is governed by a variety of chemical factors, such as pH, redox potential, organic concentrations, and oxides (Pierzynski and Schwab, 1993; Gambrell, 1994; Keller and Vedy, 1994). Uses of elemental data in soil survey applications are broad and diverse, ranging from understanding natural distributions (Wilcke and Amelung, 1996; Jersak et al., 1997) to human-induced distributions (Wilcke et al., 1998). Knowledge of the amounts and distribution of elements in soils and their relationships with other soil properties can enhance the understanding of the fate and transport of anthropogenic elements. Such knowledge expands the utility and application of soil survey information in areas of environmental concern, such as urban soils, mine spoil reclamation, smelter emissions, and agricultural waste applications (Burt et al., 2002, 2003, 2011, 2013).

2. Summary of Method

The approach of this digestion methodology is to maximize the extractable concentration of elements in digested soils while minimizing the matrix interferences, such as those occurring in digestion procedures that use HF acid. This method (4H1a1) follows EPA method 3051A. A 500-mg <2-mm soil separate that has been air-dried and ground to <200 mesh (75 µm) is weighed into a 100-ml Teflon (PFA) sample digestion vessel. To the vessel, 9.0 mL HNO₃ and 3.0 mL HCl are added. The vessel is inserted into a protection shield, covered, and placed into a temperature-controlled rotor. Following microwave digestion, the rotor and samples are cooled, and the digestate is quantitatively transferred into a 50-ml glass volumetric with high purity reverse osmosis deionized water (RODI). The volumetrics are allowed to stand overnight and then are filled to volume. The samples are transferred to 50-mL Falcon tubes prior to analysis. The concentration of Ag, As, Ba, Be, Cd, Co, Cr, Cu, Hg, Mn, Mo, Ni, P, Pb, Sb, Se, Sn, Sr, V, W, and Zn are determined using an inductively coupled plasma mass spectrophotometer (ICP–MS) by methods 4H1a1b1-21a1, respectively.

3. Interferences

Organic constituents may contain metals and are difficult to digest if present in high concentrations. Certain elements are subject to volatile losses during digestion and transfer. Certain soil minerals (e.g., quartz, feldspars) are not soluble in HNO₃ + HCl.

Interferences are corrected or minimized by using both an internal standard and collision/reaction cell technology. Also, careful selection of specific masses
for data reporting is important. Background corrections are made by ICP–MS software. Samples and standards are matrix-matched to reduce interferences.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Restrict the use of concentrated acids to a fume hood. Wash hands thoroughly after handling reagents. Filling the digestion vessel to >25 percent of the free volume or adding organic reagents or oxidizing agents to the cup may result in explosion of the digestion microwave system.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Pipette, 3-mL and 9-mL, Omnifit Corp. manufacturers, variable volume (10 mL maximum) pipettes suitable for HNO₃ and HCl delivery from 2.5 L bottles
5.3 Volumetric flasks, class A glass, 50-mL
5.4 Polypropylene bottles, 60-mL, with cap
5.5 Electronic balance, (±0.1 mg sensitivity)
5.6 Microwave oven, CEM Mars 5, 14-position HP500 Plus vessel and rotor (vessels composed of PFA, sleeves composed of advanced composite)
5.7 Volumetrics, 500-, 250-, and 50-mL, class A glass
5.8 Containers, 500-mL, polypropylene, with screw caps
5.9 Pipettes, electronic digital, 250-µL and 10-mL, Rainin Instrument Co., Woburn, MA
5.10 Inductively coupled plasma mass spectrophotometer (ICP–MS), Agilent 7500cx, Agilent Technologies Inc., Wilmington, DE
5.11 Computer, with ICP–MS ChemStation software ver. B.03.07, Agilent Technologies Inc., Wilmington, DE
5.12 Heat exchanger, G1879B, Agilent Technologies
5.13 Compressed gasses, argon (minimum purity 99.99%), hydrogen (minimum purity 99.999%), and helium (minimum purity 99.999%)
5.14 Autosampler, ASX-500 Series, Agilent Technologies., Wilmington, DE
5.15 Quartz torch, for use with HIMI, Part No. G3270-80027
5.16 Peristaltic pump (for automatic injection of internal standard)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Concentrated hydrochloric acid (HCl), 12 N, trace pure grade
6.3 Concentrated nitric acid (HNO₃), 16 N, trace pure grade
6.4 Primary standards: 1000 mg L\(^{-1}\), from High Purity Standards, Charleston, SC. Single elemental standards are manufactured in dilute HNO\(_3\), HNO\(_3\) + HF, or H\(_2\)O.

7. Procedure

**Microwave Acid Digestion**

7.1 Weight about 500 mg of fine-earth (<2-mm) or a specific particle-size separate ground to <200-mesh (75 \(\mu\)m) to the nearest 0.1 mg in a 100-mL digestion vessel. For O horizons, weigh 250 mg of sample to the nearest 0.1 mg.

7.2 If sample is principally composed of organic materials (organic C >15%), perform a preliminary digestion in the muffle furnace in an digestion crucible: 250 °C for 15 min, 450 °C for 15 min, and then 550 °C for 1 h.

7.3 Pipette 9.0 mL HNO\(_3\) and 3.0 mL HCl into the sample and allow it to become completely wet. Add acids in a fume hood. If a strong reaction is observed, allow acids to react and vent in open vessels for 10 to 15 min.

7.4 Place covered vessels in protective sleeve, cover, and place into rotor.

7.5 Place digestion rotor in the microwave oven and insert the temperature probe into the reference vessel. Attach the probe cable into the fitting in the top of the microwave. Connect the pressure monitor to the vessel.

7.6 Microwave as follows: 1200 watts at 100 percent power for 5.5 min until 175 °C, maintain at 175 °C for 4.5 min, and then cool for 5 min.

7.7 After cooling, disconnect temperature probe and pressure sensor from microwave.

7.8 Remove rotor from oven and place in fume hood.

7.9 Open each vessel carefully. Quantitatively transfer contents of vessel to a 50-mL volumetric flask with RODI water.

7.10 Cap flask and mix well by inverting. Allow to stand overnight. Finish filling with RODI water.

7.11 Decant contents into a labeled 60-mL polypropylene container.

7.12 Prepare working standards of a blank, reference soil sample from the KSSL repository, NIST, or other standard reference material, and blank by the same digestion method. Run two of these standards or a blank with each set of 14 samples.

**ICP–MS Calibration Standards, Set-Up, and Operation**

7.13 Trace Method Stock C, commercially prepared solution containing 10 \(\mu\)g/mL Sb, As, Ba, Be, Cd, Cr, Co, Cu, Pb, Mo, Ni, Se, Sn, W, V, and Zn and 1 \(\mu\)g/mL Ag.
7.14 Trace Method Stock A, commercially prepared solution containing 10 µg/mL Mn, P, and Sr

7.15 Mercury Stock Standard, commercially prepared solution containing 1 µg/mL Hg

7.16 Tuning Solution: In a 1-L volumetric flask, add 300 mL RODI, followed by 1 mL commercially prepared Stock Tuning Solution and 18 mL concentrated HNO₃. Fill to volume with RODI and mix well.

7.17 PA Tuning Solution: In a 1-L volumetric flask, add 500 mL RODI, followed by 18 mL concentrated HNO₃ and 100 mL of commercially prepared 7500 Series PA Tuning. Fill to volume and mix well.

7.18 Standard Matrix: In a 1-L volumetric flask, add 500 mL RODI, followed by 0.250 mL 1000 µg/mL Au and 18 mL concentrated HNO₃. Fill to volume with RODI and mix well.

7.19 TM8: In a 1-L volumetric flask, add 300 mL RODI, followed by 18 mL concentrated HNO₃, 6 mL concentrated HCl, 0.250 mL 1000 µg/mL Au, and 1.0 mL of Mercury Stock standard (Reagent 7.15). Fill to volume with RODI and mix well.

7.20 TM7: In a 1-L volumetric flask, add 300 mL RODI, followed by 18 mL HNO₃, 6 mL concentrated HCl, 0.250 mL 1000 µg/mL Au, and 0.5 mL of Mercury Stock standard (Reagent 7.15). Fill to volume with RODI and mix well.

7.21 TM6: In a 500-mL volumetric flask, add 300 mL RODI and 50.0 mL Trace Method Stock A (Reagent 7.14). Fill to volume with RODI and mix well.

7.22 TM5: In a 500-mL volumetric flask, add 300 mL RODI and 12.5 mL Trace Method Stock A (Reagent 7.14). Fill to volume with RODI and mix well.

7.23 TM4: In a 500-mL volumetric flask, add 300 mL RODI and 1.25 mL Trace Method Stock A (Reagent 7.14). Fill to volume with RODI and mix well.

7.24 TM3: In a 500-mL volumetric flask, add 300 mL RODI, followed by 9 mL of concentrated HNO₃, 0.125 mL 1000 µg/mL Au, and 0.5 mL Trace Method Stock C (Reagent 7.13). Fill to volume with RODI, mix well.

7.25 TM2: In 500-mL volumetric flask, add 300 mL of Standard Matrix (Reagent 7.18) and 50 mL TM3 (Reagent 7.24). Fill to volume with Standard Matrix (Reagent 7.18) and mix well.

7.26 TM1: In a 500-mL volumetric flask, add 300 mL of Standard Matrix (Reagent 7.18) and 50 mL TM2 (Reagent 7.25). Fill to volume with Standard Matrix (Reagent 7.18) and mix well.

7.27 TM0: In a 500-mL volumetric flask, add 300 mL RODI, followed by 9 mL HNO₃, 3 mL concentrated HCl, and 0.125 mL 1000 µg/mL Au. Fill to volume with RODI and mix well.

7.28 Internal Standard (1 µg/mL Li⁶, Sc, Ge, Y, In, Tb, Bi): In a 1-L flask, add 300 mL RODI, followed by 18 mL concentrated HNO₃, 6 mL concentrated HCl,
0.250 mL 1000 µg/mL Au, and 1 mL each of 1000 µg/mL Li, Sc, Ge, Y, In, Tb, and Bi. Fill to volume with RODI and mix well.

7.29 Rinse: In a 2-L volumetric flask, add 300 mL RODI and 58 mL concentrated HNO₃. Fill to volume with RODI and mix well.

7.30 Rinse #1: In a 1-L volumetric flask, add 300 mL RODI, 29 mL concentrated HNO₃, and 1 mL 1000 µg/mL Au. Fill to volume with RODI and mix well.

7.31 Rinse #2: In a 1-L flask, add 300 mL RODI, followed by 15 mL concentrated HNO₃ acid, 45 mL concentrated HCl, and 1 mL of 1000 µg/mL Au. Fill to volume with RODI and mix well.

7.32 Sample diluent: In a 1-L volumetric flask, add 300 mL RODI, followed by 0.277 mL 1000 µg/mL Au, 18 mL concentrated HNO₃, and 6 mL concentrated HCl. Fill to volume with RODI and mix well.

7.33 Standard concentrations in µg/mL for each element.

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<th>TM1</th>
<th>TM2</th>
<th>TM3</th>
<th>TM4</th>
<th>TM5</th>
<th>TM6</th>
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### 7.34 Reporting m/z and tune step for each element analyzed.

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### 7.35 Use the ICP–MS with a micromist nebulizer, quartz torch for micro flow nebulizer, and quartz spray chamber to analyze samples. Internal standard is added via peristaltic pump using 0.19 mm i.d. pump tubing. Internal standard and samples or standards are mixed via coil prior to entering the nebulizer. Samples are diluted 1:10 or greater as necessary prior to analysis with sample diluent (7.32). Perform instrument checks (tune sensitivity, resolution axis, P/A factor, internal standard RSD, torch alignment, and EM tune) prior to analysis as outlined in operation manual for instrument. Check instrument gas pressures to ensure pressures are correct and in adequate supply.
Typical tune values for trace analysis method are as follows:

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<thead>
<tr>
<th>Tune 1 ($H_2$)</th>
<th>Tune 1 ($H_2$)</th>
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</thead>
<tbody>
<tr>
<td><strong>Plasma Parameters</strong></td>
<td><strong>Q-Pole Parameters</strong></td>
</tr>
<tr>
<td>RF power</td>
<td>AMU gain</td>
</tr>
<tr>
<td>RF matching</td>
<td>AMU offset</td>
</tr>
<tr>
<td>Smpl depth</td>
<td>Axis gain</td>
</tr>
<tr>
<td>Torch-H</td>
<td>Axis offset</td>
</tr>
<tr>
<td>Torch-V</td>
<td>QP bias</td>
</tr>
<tr>
<td>Carrier gas</td>
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</tr>
<tr>
<td>Makeup gas</td>
<td></td>
</tr>
<tr>
<td>Optional gas</td>
<td></td>
</tr>
<tr>
<td>Nebulizer pump</td>
<td></td>
</tr>
<tr>
<td>Sample pump</td>
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</tr>
<tr>
<td>S/C temp</td>
<td></td>
</tr>
<tr>
<td><strong>Ion Lenses</strong></td>
<td><strong>Octapole Parameters</strong></td>
</tr>
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<td>OctP RF</td>
</tr>
<tr>
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<td>OctP bias</td>
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<tr>
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<tr>
<td>Cell entrance</td>
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<tr>
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<tr>
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<tr>
<td>S/C temp</td>
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<td><strong>Detector Parameters</strong></td>
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<td>Discriminator</td>
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<td>He gas</td>
<td>Pulse HV</td>
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<td><strong>Q-Pole Parameters</strong></td>
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<td>Tune 3 (No Gas)</td>
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<td><strong>Detector Parameters</strong></td>
</tr>
<tr>
<td>Reaction mode</td>
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</tr>
<tr>
<td>H_2 gas</td>
<td></td>
</tr>
<tr>
<td>He gas</td>
<td></td>
</tr>
<tr>
<td>Optional gas</td>
<td></td>
</tr>
<tr>
<td><strong>Discriminator</strong></td>
<td><strong>Analog HV</strong></td>
</tr>
<tr>
<td>8.0 mV</td>
<td>17 V</td>
</tr>
<tr>
<td><strong>Pulse HV</strong></td>
<td>1260 V</td>
</tr>
</tbody>
</table>

7.37 Establish detection limits using the blank standard solution. Instrumental detection limits are calculated by using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Report analyzed values lower than the detection limits as “ND” or non-detected.

8. Calculations

The calculation of mg kg\(^{-1}\) of an element in the soil from µg L\(^{-1}\) in solution is as follows:
Analyte concentration in soil (mg kg⁻¹) = \((A \times B \times C \times R \times 1000) / (E \times 1000)\)

- \(A\) = Sample extract reading (µg L⁻¹)
- \(B\) = Extract volume (L)
- \(C\) = Dilution, if performed
- \(R\) = Air-dry/oven-dry ratio (method 3D1)
- \(1000\) = Conversion factor in numerator to kg-basis
- \(E\) = Sample weight (g)
- \(1000\) = Factor in denominator (µg mg⁻¹)

9. Report

Analysis is generally done on one mass per element. If more than one mass is analyzed, only the reporting mass is used for data reporting purposes. The digested particle-size fraction needs to be identified with each sample. Data are reported to the nearest 0.01 mg kg⁻¹.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Prior to the development of modern analytical techniques, e.g., x-ray diffraction and thermal analysis, identification of minerals was based on elemental analysis and optical properties (Washington, 1930; Bain and Smith, 1994). Chemical analysis is still essential to determine mineral structural formulas and to identify and quantify specific mineral species through elemental allocation to minerals. Many clay mineral groups are subdivided based on composition.

Analysis of the entire fine-earth (<2-mm) fraction or specific particle-size separates provides information on parent material uniformity, pedon development, and mineral weathering within or between pedons. This interpretation is determined from differences between horizons or pedons in elemental concentrations; elemental ratios, such as Si/Al, Si/Al + Fe, or Ti/Zr; or total...
elemental concentrations compared to concentrations determined by selective dissolution techniques.

The inherent fertility of a soil derived from its parent material can be examined by determination of the basic cations relative to the Si or Al content. Phosphorus fertility of a soil and potential water quality problems can be better understood by measurements of total P, especially when compared to other P measurements, such as water-soluble P or Bray-1 extractable P.

Hydrofluoric acid (HF) is efficient in the digestion and dissolution of silicate minerals for elemental dissolution (Bernas, 1968; Sawhney and Stilwell, 1994). HNO₃ + HCl aids in the digestion of soil components, especially the organic fraction. Method 4H1b1a is a digestion of 100 mg of dried clay suspension, the fine-earth (<2-mm) fraction, or other particle size separate with HF + HNO₃ + HCl. Samples are placed in Teflon digestion vessels and heated in a microwave. Elemental concentration of the digestate is determined using an inductively coupled plasma atomic emission spectrometer (ICP–AES). Method 4H1b1 follows EPA method 3052.

2. Summary of Method

A 250-mg sample of <2-mm or other particle-size soil separate that has been oven-dried and ground to <200 mesh (75 µm) is weighed into a 100-ml Teflon (PFA) sample digestion vessel. Dried clay (<0.002 mm) may be used, or a clay suspension (method 7A1b1) containing approximately 250 mg of clay material is pipetted into a digestion container and dried at 110 °C. An equal amount of suspension is pipetted into a tared, aluminum weighing dish and dried at 110 °C to obtain a dried sample weight. The P and Na content of the clay fraction is not measurable if the soil is dispersed in sodium hexametaphosphate (method 7A1b1).

To the vessel, 9.0 mL HNO₃, 3.0 mL HCl, and 4 mL HF are added. The vessel is inserted into a protection shield, covered, and placed into a rotor with temperature control. Following microwave digestion, the rotor and samples are cooled, and 20 mL of 4.5% boric acid solution is added (4H1b1a). The samples are then covered and heated in the microwave. The digestate is then quantitatively transferred onto a 100-ml polypropylene volumetric with 1.9% boric acid solution to achieve a final boric acid concentration of 2.1%. The volumetrics are allowed to stand overnight and filled to volume. Approximately 60 mL is saved for analysis. The concentration of Al, Ca, Fe, K, Mg, Mn, Na, P, Si, Sr, Ti, and Zr are determined by ICP–AES by methods 4H1b1a1a1-12, respectively.

3. Interferences

Insoluble fluorides of various metals may form. Formation of SiF₄ results in gaseous losses of Si, but additions of boric acid retards formation of this molecule as well as dissolves other metal fluorides. Spectral and matrix interferences exist. Careful selection of specific wavelengths for data reporting is important.
Background corrections are made by ICP–AES software. Samples and standards are matrix matched to reduce interferences where possible.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Restrict the use of concentrated acids to a fume hood. Keep HF acid refrigerated and avoid contact with skin of all acids. Wash hands thoroughly after handling reagents. Filling the digestion vessel to greater than 25 percent of the free volume or adding organic reagents or oxidizing agents to the cup may result in explosion of the digestion microwave system.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Pipette(s) capable of delivering 3, 4, and 9 mL, Omnifit Corp., variable volume (10-mL maximum) pipettes suitable for HNO₃ and HCl delivery from 2.5 L bottles
5.3 Volumetric flasks, Nalgene, 100-mL
5.4 Polypropylene bottles, 60-mL, with cap
5.5 Electronic balance, ±0.1 mg sensitivity
5.6 Microwave, CEM Mars 5, 14-position HP500 Plus vessel and rotor (vessels composed of PFA, sleeves composed of advanced composite)
5.7 Desiccator
5.8 Disposable aluminum weighing dishes
5.9 Volumetrics, 500-mL, polypropylene
5.10 Containers, 500-mL, polypropylene, with screw caps
5.11 Pipettes, electronic digital, 2500-µL and 10-mL, Rainin Instrument Co., Woburn, MA
5.12 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES), Perkin-Elmer Optima 7300 Dual View (DV), Perkin-Elmer Corp., Norwalk, CT
5.13 Computer, with WinLab 32 software, ver. 5.1.0.0527, Perkin-Elmer Corp., Norwalk, CT, and printer
5.14 Recirculating chiller, Polyscience
5.15 Compressed gasses, argon (minimum purity 99.996%) and nitrogen (minimum purity 99.999%)
5.16 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Calcium sulfate (anhydrous) or equivalent desiccant
6.3 Hydrofluoric acid (HF), 48%, low trace metal content
6.4 Concentrated hydrochloric acid (HCl), 12 N, trace pure grade
6.5 Concentrated nitric acid (HNO₃), 16 N, trace pure grade
6.6 Boric acid solution, 4.5 percent. Dissolve 45.0 g low trace metal, granular boric acid (H₃BO₃) in 1000 mL RODI water.
6.7 Boric acid solution, 1.9 percent. Dissolve 19.0 g low trace metal, granular boric acid (H₃BO₃) in 1000 mL RODI water.
6.8 Primary standards: 1000 mg L⁻¹, from High Purity Standards, Charleston, SC. Single elemental standards in dilute HNO₃, HNO₃ + HF, or H₂O

7. Procedure

**Microwave Acid Digestion**

7.1 Fine-earth (<2-mm) or a specific particle size separate ground to <200-mesh (75 µm) is used. A 250 mg sample is weighed to the nearest 0.1 mg in a 100-mL TFM vessel. If a clay suspension is used, proceed to Section 7.2 to 7.5. If using dried specimen, proceed to Section 7.6. Samples are typically prepared in sets of 14 because the digestion rotor accommodates that number of samples. If sample is principally composed of organic materials (organic C >15%), use a sample mass of 155 mg. Alternatively, perform a preliminary digestion in the muffle furnace in a digestion crucible: 250 °C for 15 min, 450 °C for 15 min, followed by 550 °C for 1 h.

7.2 Prepare Na-saturated clay as in method 7A1a1, (Preparation of Clay Suspension, Sections 7.8 to 7.19). Clay dispersion by this method eliminates quantitative analysis of Na and P in the clay due to dispersion by sodium hexametaphosphate. Digestion of the entire fine earth (<2-mm) fraction or any fraction not derived by dispersion with sodium hexametaphosphate (or other Na and P-containing dispersing agents) can be quantitatively analyzed for Na and P. Use RODI water for dispersion of clays and cleaning of test tubes and dishware.

7.3 Pipette a known aliquot of clay suspension containing approximately 250 mg clay into a 100-mL TFM vessel. The volume of required suspension depends on the clay concentration in the suspension but is generally 6 to 10 mL. More dilute suspensions should be partially evaporated under a fume hood to concentrate the clay prior to transfer to the Teflon container.

7.4 Pipette a duplicate aliquot of suspension (as used in Section 7.3) into a tared Al weighing dish, dry at 110 °C, cool in a desiccator, and weigh to the nearest 0.1 mg. Use this value as the sample weight in the calculations.

7.5 Dry the Teflon container and clay suspension in an oven for 4 h or until the aqueous portion of the suspension is completely evaporated. Remove from
oven and cool on the bench top or in a fume hood. Cooling in a desiccator is not required.

7.6 Pipette 9.0 mL HNO₃ and 3.0 mL HCl into the sample and allow to completely wet. Then pipette 4 mL HF into sample. Add acids in the fume hood.

7.7 Place covered Teflon digestion vessels in protective sleeve, cover, and place into rotor.

7.8 Place digestion rotor in the microwave oven and insert the temperature probe into the reference vessel. Attach the probe cable into the fitting in the top of the microwave. Connect the pressure monitor to the vessel.

7.9 Microwave settings are as follows:
- 1200 Watts, 100% power for 10 min (350 psi) to approximately 180 °C;
- 1200 Watts, 70% power for 9.5 min (350 psi), holding at 180 °C;
- cool (vent) for 15 min.

7.10 After venting, disconnect temperature probe from microwave.

7.11 Remove rotor from oven and place in fume hood.

7.12 Open each vessel carefully, add 20 mL 4.5 percent boric acid (H₃BO₃) solution, and then cover vessels and re-digest in microwave at:
- 1200 Watts, 100% power (350 psi) to 160 °C;
- maintain at 160 °C for 10 min;
- cool (vent) for 15 min.

7.13 Transfer contents of digestion vessel to a 100-mL Nalgene volumetric flask and adjust to near volume with 1.9% boric acid, achieving a final concentration of 2.1 percent H₃BO₃.

7.14 Cap flask and mix well by inverting. Allow to stand overnight to dissolve any metal fluorides. Finish filling to volume with 1.9% boric acid.

7.15 Invert the volumetric flask to mix.

7.16 Prepare working standards of a blank, reference soil sample from the KSSL repository, a National Institute of Standards and Technology (NIST) standard reference, or other reference material by the same digestion method. Run one of these standards with each set of 14 samples.

**ICP–AES Calibration Standards, Set-up, and Operation**

7.17 Instrument calibration standards for analysis are limited to specific combinations of elements because of chemical incompatibilities of certain elements. Each working standard is used in two concentrations, high and low. The concentrations of elements in the low standards (CALO, ALLO, and SILO) are 50 percent of the concentrations in the high standards.
(CAHI, ALHI, and SIHI). The amounts of primary standards (1000 mg L$^{-1}$) to make 500-mL volume of the low and high calibration standards, at the specified concentrations, for ICP–AES analysis are as follows:

**7.17.1** CALO is 75, 25, 20, and 10 mg L$^{-1}$ of Ca, K, Mg, and Mn, respectively. To a 500-mL volumetric flask, add 37.5, 12.5, 10.0, and 5.0 mL of the Ca, K, Mg, and Mn primary standards (1000 mg L$^{-1}$), respectively.

**7.17.2** CAHI is 150, 50, 40, and 20 mg L$^{-1}$ of Ca, K, Mg, and Mn, respectively. To a 500-mL volumetric flask, add 75.0, 25.0, 20.0, and 10.0 mL of the Ca, K, Mg, and Mn primary standards (1000 mg L$^{-1}$), respectively.

**7.17.3** ALLO is 100, 75, 5, 5, and 25 mg L$^{-1}$ of Al, Fe, Ti, Zr, and Na, respectively. To a 500-mL volumetric flask, add 50.0, 37.5, 2.5, 2.5, and 12.5 mL of the Al, Fe, Ti, Zr, and Na primary standards (1000 mg L$^{-1}$), respectively.

**7.17.4** ALHI is 200, 150, 10, 10, and 50 mg L$^{-1}$ of Al, Fe, Ti, Zr, and Na, respectively. To a 500-mL volumetric flask, add 100.0, 75.0, 5.0, 5.0, and 25.0 mL of the Al, Fe, Ti, Zr, and Na primary standards (1000 mg L$^{-1}$), respectively.

**7.17.5** SILO is 225, 5, and 1.5 mg L$^{-1}$ of Si, P, and Sr, respectively. To a 500-mL volumetric flask, add 112.5, 2.5, and 7.5 mL of the Si, P, and Sr primary standards (1000 mg L$^{-1}$), respectively.

**7.17.6** SIHI is 450, 10, and 10 mg L$^{-1}$ of Si, P, and Sr respectively. To a 500-mL volumetric flask, add 225.0, 5.0, and 5.0 mL of the Si, P, and Sr primary standards (1000 mg L$^{-1}$), respectively.

**7.18** To the calibration standards and a blank, also add the following chemicals: 20.0 mL HF, 45.0 mL HNO$_3$, 15.0 mL HCl, and 10.90 g granular boric acid. Make all standards and the blank to a final volume of 500-mL with RODI water.

**7.19** Use the ICP–AES in radial mode and analyze for the following elements: Fe, Mn, Al, Ca, Mg, Na, K, P, Si, Sr, Zr, and Ti. No initial dilutions of samples are necessary prior to analysis. Perform instrument checks (Hg alignment, BEC, %RSD of Mn solution) prior to analysis as discussed in operation manual of instrument. Check instrument alignment and gas pressures to obtain optimum readings with maximum signal to noise ratio.

**7.20** Analyses are generally performed at two or more wavelengths for each element. The selected wavelengths are as follows: (reported wavelength listed first and in bold):
<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(nm)</td>
</tr>
<tr>
<td>Al</td>
<td>308.215, 396.157</td>
</tr>
<tr>
<td>Ca</td>
<td>315.887, 317.932</td>
</tr>
<tr>
<td>Fe</td>
<td>259.939, 238.205</td>
</tr>
<tr>
<td>K</td>
<td>766.490</td>
</tr>
<tr>
<td>Mg</td>
<td>280.271, 279.075</td>
</tr>
<tr>
<td>Mn</td>
<td>257.610, 260.570</td>
</tr>
<tr>
<td>Na</td>
<td>589.592, 588.995</td>
</tr>
<tr>
<td>P</td>
<td>178.221, 213.620</td>
</tr>
<tr>
<td>Si</td>
<td>212.412, 251.612</td>
</tr>
<tr>
<td>Sr</td>
<td>407.747, 421.523</td>
</tr>
<tr>
<td>Ti</td>
<td>334.940, 368.522</td>
</tr>
<tr>
<td>Zr</td>
<td>339.197, 343.818</td>
</tr>
</tbody>
</table>

7.21 Use the blank standard solution to dilute those samples with concentrations greater than the high standard. Rerun all elements and use only the data needed from the diluted analysis.

7.22 Establish detection limits using the blank standard solution. These instrumental detection limits are calculated by using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are set equal to zero.

8. Calculations

8.1 The calculation of mg kg\(^{-1}\) of an element in the soil from mg L\(^{-1}\) in solution is as follows:

\[
\text{Analyte concentration in soil (mg kg}\,^\text{−1}) = \frac{(A \times B \times C \times R \times 1000)}{E}
\]

- A = Sample extract reading (mg L\(^{-1}\))
- B = Extract volume (L)
- C = Dilution, if performed
- R = Air-dry/oven-dry ratio (method 3D1)
- 1000 = Conversion factor in numerator to kg-basis
- E = Sample weight (g)

8.2 Data are recorded on an elemental basis. Often, users request data for an oxide form. The factor for converting from an elemental form to an
oxide form is based on the atomic weights of the element and oxygen. An example is as follows:

Atomic weight Si = 28.09
Atomic Weight O = 16.0
Molecular weight SiO$_2$ = 60.09

Calculate percent Si in SiO$_2$ as follows:

$$\text{Si} \text{ (%)} = \left(\frac{28.09}{60.09}\right) \times 100 = 46.7\%$$

There is 46.7 percent Si in SiO$_2$. To convert from mg/kg Si to percent Si oxide (SiO$_2$) in the soil, divide by 10,000 (to convert from mg kg$^{-1}$ to %), then divide the percent Si by 0.467 or multiply by the inverse of this value. The element, oxide form, and the elemental percent in the oxide form are as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Oxide Form</th>
<th>Elemental %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>SiO$_2$</td>
<td>46.7</td>
</tr>
<tr>
<td>Al</td>
<td>Al$_2$O$_3$</td>
<td>52.9</td>
</tr>
<tr>
<td>Fe</td>
<td>Fe$_2$O$_3$</td>
<td>69.9</td>
</tr>
<tr>
<td>Mg</td>
<td>MgO</td>
<td>60.3</td>
</tr>
<tr>
<td>Mn</td>
<td>MnO</td>
<td>77.4</td>
</tr>
<tr>
<td>K</td>
<td>K$_2$O</td>
<td>83.0</td>
</tr>
<tr>
<td>Ti</td>
<td>TiO$_2$</td>
<td>59.9</td>
</tr>
<tr>
<td>Ca</td>
<td>CaO</td>
<td>71.5</td>
</tr>
<tr>
<td>Zr</td>
<td>ZrO$_2$</td>
<td>74.0</td>
</tr>
<tr>
<td>P</td>
<td>P$_2$O$_5$</td>
<td>43.6</td>
</tr>
<tr>
<td>Na</td>
<td>Na$_2$O</td>
<td>74.2</td>
</tr>
</tbody>
</table>

9. Report

Analyses are generally performed at two or more wavelengths for each element, with one selected wavelength for reporting purposes. The particle-size fraction digested needs to be identified with each sample. Data are reported to the nearest mg kg$^{-1}$.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Total Analysis (4H)
Dry Combustion (4H2)
  Thermal Conductivity Detector (4H2a)
  Total Carbon, Nitrogen, and Sulfur (4H2a1-3)
  Air-Dry, <2 mm (4H2a1-3a1)

1. Application

Organic Matter and Organic Carbon: Soil organic matter has been defined as the organic fraction of the soil exclusive of undecayed plant and animal residues and has been used synonymously with “humus” (Soil Science Society of America, 1997). For laboratory analyses, however, the soil organic matter generally includes only those organic materials that accompany soil particles through a 2-mm sieve (Nelson and Sommers, 1982). The organic matter content influences many soil properties, including water retention capacity; extractable bases; capacity to supply N, P, and micronutrients; stability of soil aggregates; and soil aeration (Nelson and Sommers, 1996).

Organic C is a major component of soil organic matter. Organic C consists of the cells of microorganisms; plant and animal residues at various stages of decomposition; stable “humus” synthesized from residues; and nearly inert and highly carbonized compounds, such as charcoal, graphite, and coal (Nelson and Sommers, 1982). Because organic C is the major component of soil organic matter, a measurement of organic C can serve as an indirect determination of organic matter. Organic C determination is by either wet or dry combustion. In the past, the KSSL used the now-obsolete wet combustion method (6A1c): Walkley-Black modified acid-dichromate digestion, FeSO₄ titration, automatic titrator.

Values for organic C are multiplied by the “Van Bemmelen factor” (1.724) to calculate organic matter. This factor is based on the assumption that organic matter contains 58% organic C. The proportion of organic C in soil organic matter for a range of soils is highly variable. Any constant factor that is selected is only an approximation. Studies have indicated that subsoils have a higher factor than surface soils (Broadbent, 1953). Surface soils rarely have a factor <1.8 and usually range from 1.8 to 2.0. The subsoil factor may average ≈2.5. The
preference is to report organic C rather than to convert the organic C to organic matter through use of an approximate correction factor.

The KSSL also uses a direct determination of soil organic matter. In this determination, the organic matter is destroyed and then the loss in weight of the soil is taken as a measure of the organic matter content (5A). The percent organic matter lost on ignition (400 °C) can be used in place of organic matter estimates by the Walkley-Black organic C method.

Total Carbon: Total C is the sum of organic and inorganic C. Most of the organic C is associated with the organic matter fraction, and the inorganic C is generally found with carbonate minerals. The organic C in mineral soils generally ranges from 0 to 12% (Nelson and Sommers, 1996).

Total C is quantified by two basic methods, i.e., wet or dry combustion. The KSSL uses dry combustion (4H2a1). In total C determinations, all forms of C in a soil are converted to CO₂ followed by a quantification of the evolved CO₂. Total C can be used to estimate the organic C content of a soil. The difference between total and inorganic C is an estimate of the organic C. The inorganic C should be approximately equivalent to carbonate values measured by CO₂ evolution with strong acid (Nelson and Sommers, 1996). In KSSL method 4E1a1a1, the amount of carbonate in a soil is determined by treating a sample with HCl followed by a manometric measurement of the evolved CO₂. The amount of carbonate is then calculated as a CaCO₃ equivalent basis. Organic C defines mineral and organic soils. In soil taxonomy, organic C is also used at lower taxonomic levels, e.g., ustolic and fluventic subgroups (Soil Survey Staff, 2014).

Total Nitrogen: Total N includes organic and inorganic forms. The total N content of the soil may be <0.02% in subsoils, 2.5% in peats, and 0.06 to 0.5% in surface layers of many cultivated soil (Bremmer and Mulvaney, 1982). The total N data may be used to determine the soil C:N ratio, the soil potential to supply N for plant growth, and the N distribution in the soil profile. The C:N ratio generally ranges between 10 to 12. Variations in the C:N ratio may serve as an indicator of the amount of soil inorganic N. Uncultivated soils typically have higher C:N ratios than those of cultivated soils.

Soils that have large amounts of illites or vermiculites can “fix” significant amounts of N compared to those soils dominated by smectites or kaolinites (Young and Aldag, 1982; Nommik and Vahtras, 1982). Because the content of organic C of many soils diminishes with depth while the level of “fixed” N remains constant or increases, the C:N ratio narrows (Young and Aldag, 1982). The potential to “fix” N has important fertility implications as the “fixed” N is slowly available for plant growth.

Two methods of analysis of total N have gained acceptance for the determination of total N in soils. These are the Kjeldahl (1883) method, which is essentially a wet oxidation procedure, and the Dumas (1831) method, which is fundamentally a dry oxidation (i.e., combustion) procedure (Bremmer, 1996). The KSSL uses the combustion technique for analysis of total N (4H2a2).
**Total Sulfur:** Organic and inorganic S forms are found in soils. The organic fraction accounts for >95% of the total S in most soils from humid and semi-humid areas (Tabatabai, 1996). Mineralization of organic S and its conversion to sulfate by chemical and biological activity may serve as a source of plant-available S. Total S typically ranges from 0.01 to 0.05% in most mineral soils. In organic soils, total S may be >0.05%. The proportion of organic and inorganic S in a soil sample varies widely according to soil type and depth of sampling (Tabatabai, 1996).

In well drained, well aerated soils, most of the inorganic S normally occurs as sulfate. Marine tidal flats, other anaerobic marine sediments, and mine spoils usually have large amounts of reduced S compounds that oxidize to sulfuric acid upon exposure to the air. In arid regions, significant amounts of inorganic S are found as sulfates, such as gypsum and barite (Tabatabai, 1996).

The typical use of total S is an index of the total reserves of this element, which may be converted to plant-available S. The KSSL uses the combustion technique for analysis of total S (4H2a3). Extractable sulfate S (SO$_4^{2-}$-S) is an index of readily plant-available S. Reagents that have been used for measuring SO$_4^{2-}$-S include water, hot water, ammonium acetate, sodium carbonate and other carbonates, ammonium chloride and other chlorides, potassium phosphate and other phosphates, and ammonium fluoride (Bray-1). Extractable SO$_4^{2-}$-S does not include the labile fraction of soil organic S that is mineralized during the growing season (Tabatabai, 1996). Extraction reagents for organic S include hydrogen peroxide, sodium bicarbonate, sodium hydroxide, sodium oxalate, sodium peroxide, and sodium pyrophosphate. There are other methods available for determination of S, especially for total S and SO$_4^{2-}$-S. The investigator may refer to the review by Beaton et al. (1968). For detailed discussion of the application of total C, N, and S, refer to Soil Survey Staff (2011).

2. **Summary of Method**

An air-dry (80 mesh, <180 µm) sample is packed in a tin foil, weighed, and analyzed for total C, N, and S by an elemental analyzer (methods 4H2a1-3, respectively). The elemental analyzer works according to the principle of catalytic tube combustion in an oxygenated CO$_2$ atmosphere and high temperature. The combustion gases are freed from foreign gases. The desired measuring components (N$_2$, CO$_2$, and SO$_2$) are separated from each other with the help of specific adsorption columns and are determined in succession with a thermal conductivity detector. Helium is the flushing and carrier gas. Percent total C, N, and S are reported by methods 4H2a1-3a1, respectively.

3. **Interferences**

Contamination through body grease or perspiration must be avoided in sample packing. Substance loss after weighing should be avoided by exact folding of the sample into the tin foil. Air in the sample material (falsifying the N value) should be minimized by compressing the sample packing. Insufficient O$_2$ dosing
reduces the catalysts, decreasing their effectiveness and durability. Burnt sample substance that remains in ash finger falsifies the results of subsequent samples. WO$_3$ is used as sample additive and combustion filling to aid combustion or bind interfering substances (alkaline or earth-alkaline elements, non-volatile sulfates).

4. Safety

Exhaust gas pipes should lead into a ventilated fume hood. Aggressive combustible products should not be analyzed. Before working on electrical connections (adsorption columns) or before changing reaction tubes, the instrument must be cooled down and cooled off. Gloves and safety glasses should be worn at all times during operation and maintenance of instrument.

5. Equipment

5.1 Elemental analyzer with on-line electronic balance (0.1 ±mg sensitivity) and automatic sample feeder, Elementar vario EL Elementar vario EL III, and Elementar vario Cube, Elementar Analysensysteme GmbH, Hanau-Germany, and combustibles (Elementar Americas, Inc., Mt. Laurel, NJ; Alpha Resources Inc., Stevensville, MI) as follows:

5.1.1 Quartz ash finger, quartz
5.1.2 Quartz bridge
5.1.3 Combustion tube
5.1.4 Reduction tube
5.1.5 Gas purification (U-tube, GL 18)
5.1.6 Support tube (65 mm)
5.1.7 Protective tube
5.1.8 O$_2$ lance (150 mm rapid N)
5.1.9 Tin boats (4 x 4 x 11 mm)
5.1.10 Tin foil cups

5.2 Computer, with vario EL software, Elementar Analysensysteme GmbH, Hanau-Germany, and printer

6. Reagents

6.1 Sulfanilic acid, calibration standard, 41.6% C, 4.1% H, 8.1 % N, 27.7% O, and 18.5% S
6.2 Copper sticks
6.3 Corundum balls, high purity, alumina spheres, 3–5 mm
6.4 Cerium dioxide, 1–2 mm
6.5 Tungsten oxide powder, sample additive
6.6 Tungsten trioxide granulate, combustion tube filling
6.7 Quartz wool
6.8 Silver wool  
6.9 Phosphorus pentoxide, Sicapent, Elementar Americas, Inc., Mt. Laurel, NJ  
6.10 Helium, carrier gas, 99.996% purity  
6.11 Oxygen, combustion gas, 99.995% purity  

7. Procedure  

**Elemental Analyzer Set-up and Operation**  

7.1 Refer to the manufacturer’s manual for operation and maintenance of the elemental analyzer. Conditioning of the elemental analyzer and determination of factor and blank value limit are part of the daily measuring routine. The following are only very general guidelines for instrument parameters for the various analytes in the CNS mode.  

<table>
<thead>
<tr>
<th>Instrument Parameters</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Temperature</strong></td>
<td></td>
</tr>
<tr>
<td>Furnace 1</td>
<td>1140 °C</td>
</tr>
<tr>
<td>Furnace 2</td>
<td>850 °C</td>
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<tr>
<td>Furnace 3</td>
<td>0 °C</td>
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<tr>
<td>CO₂ column</td>
<td>85 °C</td>
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<tr>
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<tr>
<td>SO₂ col. standby</td>
<td>140 °C</td>
</tr>
<tr>
<td><strong>Timing</strong></td>
<td></td>
</tr>
<tr>
<td>Flush</td>
<td>5 s</td>
</tr>
<tr>
<td>Oxygen delay</td>
<td>10 s</td>
</tr>
<tr>
<td>Autozero delay</td>
<td>30 s</td>
</tr>
<tr>
<td>Integrator reset delay</td>
<td>50 s</td>
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<tr>
<td>Peak anticipation N</td>
<td>70 s</td>
</tr>
<tr>
<td>Peak anticipation C</td>
<td>125 s</td>
</tr>
<tr>
<td>Peak anticipation S</td>
<td>70 s</td>
</tr>
<tr>
<td><strong>Integrated Reset Delay for S</strong></td>
<td>60 s</td>
</tr>
</tbody>
</table>

<table>
<thead>
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<tr>
<td>N peak</td>
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<tr>
<td>C peak</td>
<td>3 mV</td>
</tr>
<tr>
<td>S peak</td>
<td>3 mV</td>
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</table>
Instrument Parameters

<table>
<thead>
<tr>
<th>O₂ Dosing</th>
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<tbody>
<tr>
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<tr>
<td>Index 4</td>
<td>120 s</td>
</tr>
<tr>
<td>Index 5</td>
<td>180 s</td>
</tr>
</tbody>
</table>

### Elemental Analyzer Calibration and Analysis

#### 7.2
A calibration that covers the desired working range of each element is performed. The calibration test analyzes sulfanilic acid with each given element content at different weights. The PC program automatically computes the calibration function (linear, polynomial, or mixed). Calibration typically remains stable for at least 6 months. Re-calibration is recommended when the daily factor is outside the range of 0.9 to 1.1 or if components that influence the results (e.g., detector or adsorption column) have been exchanged. Changing the desorption temperature of adsorption columns can also require a re-calibration.

#### 7.3
Add 0.100 g of tungsten oxide in tin foil and tare. A homogenized, fine-grind, air-dry soil sample is then packed in the tin foil, which is weighed (0.100 to 0.05 g) and placed into the carousel of the automatic sample feeder of the elemental analyzer. Sample weight is based on visual observation of the sample, related to element content, homogeneity, and combustion behavior of the sample. The sample weight is entered in the PC from an on-line electronic balance via an interface. A quality control (QC) sample is performed at a minimum of every 35 to 40 samples.

### 8. Calculations

\[
C(\%) = C_i \times AD/OD
\]

where:

\[
C(\%) = C(\%), \text{ oven-dry basis}
\]

\[
C_i = C(\%) \text{ instrument}
\]

\[
AD/OD = \text{Air-dry/oven-dry ratio (method 3D1)}
\]

\[
N(\%) = N_i \times AD/OD
\]

where:

\[
N(\%) = N(\%), \text{ oven-dry basis}
\]

\[
N_i = N(\%) \text{ instrument}
\]

\[
AD/OD = \text{Air-dry/oven-dry ratio (method 3D1)}
\]
\[ S(\%) = S_i \times \text{AD/OD} \]

where:
- \( S(\%) \) = \( S(\%) \) on oven-dry basis
- \( S_i \) = \( S(\%) \) instrument
- \( \text{AD/OD} \) = Air-dry/oven-dry ratio (method 3D1)

9. Report

Report total C and S percentage to the nearest 0.01% and total N to the nearest 0.001%.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Ground and Surface Water Analysis (4I)

Hydrogen-Ion Activity (4I1)

Electrode (4I1a)

Standard Glass Body Combination (4I1a1)

Digital pH/Ion Meter (4I1a1a)

pH (4I1a1a1)

1. Application

The pH of a water sample is a commonly performed determination and one of the most indicative measurements of water chemical properties. The acidity, neutrality, or basicity is a key factor in the evaluation of water quality.

2. Summary of Method

The pH of the water sample is measured with a calibrated combination electrode/digital pH meter (method 4I1a1a1).

3. Interferences

Water pH needs to be measured immediately upon arrival at the laboratory in order to maintain optimal preservation of sample (Velthorst, 1996).

4. Safety

No significant hazards are associated with the procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ

5.2 Tubes, 50-mL, with caps

5.3 Digital pH/ion meter, Accumet Model AR 15, Fisher Scientific

5.4 Electrode, standard glass body combination, Accuflow, Fisher Scientific

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Borax pH buffers, pH 4.00, 7.00, and 9.18, for electrode calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Water sample is filtered into a 50-mL tube. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.2 Calibrate the pH meter with pH 4.00, 7.00, and 9.18 buffer solutions.

7.3 After equipment calibration, gently wash the electrode with RO water. Dry the electrode. Do not wipe the electrode with a tissue as this may cause a static charge on the electrode.

7.4 Gently lower the electrode in the water sample until the KCl junction of the electrode is beneath the water surface.

7.5 Allow the pH meter to stabilize before recording the pH. Record pH to the nearest 0.01 unit.

7.6 Gently raise the pH electrode and wash the electrode with a stream of RO water.

8. Calculations

No calculations are required for this procedure.

9. Report

Report the pH of the water sample to the nearest 0.1 pH unit.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Ground and Surface Water Analyses (4l)
Electrical Conductivity and Salts (4l2)
Conductivity Bridge and Cup (4l2a)
Electrical Conductivity (4l2a1)

1. Application

Measuring electrical conductivity (EC) and total dissolved salts (TDS) is generally straightforward, but measuring these values in soils is not
straightforward because salinity is significantly affected by prevailing moisture content. A primary source of salts is chemical weathering of the minerals present in soils and rocks. The most important include dissolution, hydrolysis, carbonation, acidification, and oxidation-reduction (National Research Council, 1993). All of these reactions contribute to an increase in the dissolved mineral load in the soil solution and in waters.

2. Summary of Method

The electrical conductivity of the water sample is measured using an electronic bridge (4I2a1).

3. Interferences

Reverse osmosis water is used to zero and flush the conductivity cell. The extract temperature is assumed to be 25 °C. If the temperature deviates significantly, a correction may be required.

Provide airtight storage of KCl solution and samples to prevent soil release of alkali-earth cations. Exposure to air can cause gains and losses of water and dissolved gases, significantly affecting EC readings.

4. Safety

No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.2 Tubes, 50-mL, with caps
5.3 Conductivity bridge and conductivity cell, with automatic temperature adjustment, 25 ±0.1 °C, Markson Model 1056, Amber Science, Eugene, Oregon

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Potassium chloride (KCl), 0.010 N. Dry KCl overnight in oven (110 °C). Dissolve 0.7456 g of KCl in RODI water and bring to 1-L volume. Conductivity at 25 °C is 1.412 mhos cm⁻¹.

7. Procedure

7.1 Water sample is filtered into a 50-mL tube and capped. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
7.2 Standardize the conductivity bridge using RO water (blank) and 0.010 N KCl (1.41 mmhos cm\(^{-1}\)).

7.3 Read conductance of water sample directly from the bridge.

7.4 Record conductance to 0.01 mmhos cm\(^{-1}\).

8. Calculations

8.1 No calculations are required for this procedure.

8.2 Use the following relationship to estimate the total soluble cation or anion concentration (meq L\(^{-1}\)) in the water.

\[
\text{EC (mmhos cm}^{-1}\text{)} \times 10 = \text{Cation or Anion (meq L}^{-1}\text{)}
\]

9. Report

Report prediction conductance to the nearest 0.01 mmhos cm\(^{-1}\).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Ground and Surface Water Analyses (4I)

Electrical Conductivity and Salts (4I2)

Atomic Absorption Spectrophotometer (4I2b)

Calcium, Magnesium, Potassium, and Sodium (4I2b1-4)

1. Application

Nutrients (nitrogen and phosphorus), sediments, pesticides, salts, or trace elements in ground and surface water affect soil and water quality (National Research Council, 1993). This procedure is developed for the analysis of ground or surface water.

2. Summary of Method

The water sample is filtered and diluted with an ionization suppressant (La\(_2\)O\(_3\)). The analytes are measured by an atomic absorption spectrophotometer (AAS). The data are automatically recorded by a computer and printer. The saturation extracted cations Ca\(^{2+}\), Mg\(^{2+}\), K\(^+\), and Na\(^+\) are reported in meq L\(^{-1}\) (mmol (+) L\(^{-1}\)) in methods 4I2b1-4, respectively.
3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the analysis of these cations. These interferences vary in importance, depending upon the particular analyte selected. Do not use borosilicate tubes because of potential leaching of analytes.

4. Safety

Wear protective clothing and eye protection. Exercise special care when preparing reagents. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the AAS.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.3 Tubes, 50-mL, with caps
5.4 Atomic absorption spectrophotometer (AAS), double-beam, AAnalyst 400, Perkin-Elmer Corp., Norwalk, CT
5.5 Autosampler, S-10, Perkin-Elmer Corp., Norwalk, CT
5.6 Computer, with AA WinLab software, Perkin-Elmer Corp., Norwalk, CT, and printer
5.7 Single-stage regulator, acetylene
5.8 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.9 Plastic test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.10 Containers, polyethylene
5.11 Peristaltic pump

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Hydrochloric acid (HCl), concentrated 12 N
6.3 HCl, 1:1 HCl:RODI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part RODI water.
6.4 Stock Lanthanum Ionization Suppressant Solution (SLISS), 65,000 mg L\(^{-1}\). Wet 152.4 g of lanthanum oxide (La\(_2\)O\(_3\)) with 100 mL RODI water. Slowly and cautiously add 500 mL of 6 \(N\) HCl to dissolve the La\(_2\)O\(_3\). Cooling the solution is necessary. Dilute to 2 L with RODI water. Filter solution. Store in polyethylene container.

6.5 Working Lanthanum Ionization Suppressant Solution (WLISS), 2000 mg L\(^{-1}\). Dilute 61.5 mL of SLISS with 1800 mL of RODI water (1:10). Make to 2-L volume with RODI water. Invert to mix thoroughly. Store in polyethylene container.

6.6 Primary Stock Standards Solution (PSSS), high purity, 1000 mg L\(^{-1}\): Ca, Mg, K, and Na.

6.7 Working Stock Mixed Standards Solution (WSMSS) for Ca, Mg, and K. In a 500-mL volumetric flask, add 250 mL Ca PSSS, 25 mL Mg PSSS, and 100 mL K PSSS=500 mg L\(^{-1}\) Ca, 50 mg L\(^{-1}\) Mg, and 200 mg L\(^{-1}\) K. Dilute to volume with RODI water. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in the refrigerator.

6.8 Mixed Calibration Standards Solution (MCSS), High, Medium, Low, Very Low, and Blank as follows:

6.8.1 MCSS High Standard (1:100): Dilute WSMSS 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 5 mg L\(^{-1}\) Ca, 0.5 mg L\(^{-1}\) Mg, and 2 mg L\(^{-1}\) K.

6.8.2 MCSS Medium Standard (1:200): To a 100-mL volumetric flask, add 50 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 2.5 mg L\(^{-1}\) Ca, 0.25 mg L\(^{-1}\) Mg, and 1 mg L\(^{-1}\) K.

6.8.3 MCSS Low Standard (1:400): To a 100-mL volumetric flask, add 25 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 1.25 mg L\(^{-1}\) Ca, 0.125 mg L\(^{-1}\) Mg, and 0.5 mg L\(^{-1}\) K.

6.8.4 MCSS Very Low Standard (1:600): To a 100-mL volumetric flask, add 16.65 mL of WSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store
in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentrations are 0.83 mg L\(^{-1}\) Ca, 0.08 mg L\(^{-1}\) Mg, and 0.33 mg L\(^{-1}\) K.

**6.8.5** MCSS Blank: 0 mL of Ca, Mg, and K. Dilute RODI water 1:100 with WLISS.

**6.9** Na Calibration Standards Solution (NaCSS), High, Medium, Low, and Very Low as follows:

**6.9.1** NaCSS High Standard (1:100): Dilute Na PSMSS (1000 mg L\(^{-1}\)) 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 10 mg L\(^{-1}\) Na.

**6.9.2** NaCSS Medium Standard (1:200): In a 50-mL volumetric, add 25 mL of Na PSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to mix thoroughly. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 5 mg L\(^{-1}\) Na.

**6.9.3** NaCSS Low Standard (1:400): In a 50-mL volumetric flask, add 12.5 mL of PSMSS and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. Final concentration is 2.5 mg L\(^{-1}\) Na.

**6.9.4** NaCSS Very Low Standard (1:600): In a 50-mL volumetric flask, add 8.35 mL of PSMSS Na (1000 ppm) and bring to volume with RODI water. Dilute 1:100 with WLISS. Invert to thoroughly mix. Store in polyethylene containers. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate before use. Final concentration is 1.67 mg L\(^{-1}\) Na.

**6.9.5** NaCSS Blank = 0 mL Na PSMSS. Dilute RODI water 1:100 with WLISS.

**6.10** Compressed air with water and oil traps

**6.11** Acetylene gas, purity 99.6%

**7. Procedure**

**7.1** Water sample is filtered into a 50-mL tube and capped. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
Dilution of Calibration Standards and Sample Extracts

7.2 The 10-mL syringe is for diluent (WLISS). The 1-mL syringe is for the MCSS and water sample. Set the digital diluter at a 1:100 dilution. See Sections 6.8 and 6.9 for preparation of the MCSS and NaCSS. Dilute the saturation extract sample with 100 parts of WLISS (1:100).

7.3 Dispense the diluted sample solutions into test tubes that have been placed in the sample holders of the sample changer.

AAS Set-up and Operation

7.4 Refer to the manufacturer’s manual for operation of the AAS. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Conc.</th>
<th>Burner &amp; angle</th>
<th>Wavelength</th>
<th>Slit</th>
<th>Fuel/Oxidant (C\textsubscript{2}H\textsubscript{2}/Air)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mg L\textsuperscript{-1})</td>
<td>(nm)</td>
<td>(mm)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca</td>
<td>5.0</td>
<td>10 cm @ 0 °</td>
<td>422.7</td>
<td>0.7</td>
<td>1.5/0.0</td>
</tr>
<tr>
<td>Mg</td>
<td>0.5</td>
<td>10 cm @ 0 °</td>
<td>285.2</td>
<td>0.7</td>
<td>1.5/10.0</td>
</tr>
<tr>
<td>K</td>
<td>2.0</td>
<td>10 cm @ 0 °</td>
<td>766.5</td>
<td>0.7</td>
<td>1.5/10.0</td>
</tr>
<tr>
<td>Na</td>
<td>10.0</td>
<td>10 cm @ 30 °</td>
<td>589.0</td>
<td>0.2</td>
<td>1.5/10.0</td>
</tr>
</tbody>
</table>

7.5 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

AAS Calibration and Analysis

7.6 Calibrate the instrument by using the MCSS and NaCSS. The data system then associates the concentrations with the instrument responses for each MCSS and NaCSS. Rejection criteria for MCSS and NaCSS is R\textsuperscript{2} <0.99.

7.7 If sample exceeds calibration standard, the sample is diluted 1:5, 1:20, 1:100, etc., with RODI water followed by 1:100 dilution with WLISS.

7.8 Perform one quality control (QC) (Low Standard) for every 12 samples. If reading is not within 10%, the instrument is re-calibrated and QC re-analyzed.

7.9 Record analyte readings to 0.01 mg L\textsuperscript{-1}.

8. Calculations

The instrument readings for analyte concentration are in mg L\textsuperscript{-1}. These analyte concentrations are converted to meq L\textsuperscript{-1} as follows:

\[
\text{Analyte Concentration in Soil (meq L}^{-1}\text{)} = \frac{A \times B}{C}
\]
where:
\( A \) = Analyte (Ca, Mg, K, Na) concentration in extract (mg L\(^{-1}\))
\( B \) = Dilution ratio, if needed
\( C \) = Equivalent weight

where:
\( \text{Ca}^{2+} = 20.04 \text{ mg meq}^{-1} \)
\( \text{Mg}^{2+} = 12.15 \text{ mg meq}^{-1} \)
\( \text{K}^{+} = 39.10 \text{ mg meq}^{-1} \)
\( \text{Na}^{+} = 22.99 \text{ mg meq}^{-1} \)

9. Report
Report the saturation extraction cations \( \text{Ca}^{2+}, \text{Mg}^{2+}, \text{K}^{+}, \) and \( \text{Na}^{+} \) to the nearest 0.1 meq L\(^{-1}\) (mmol (+) L\(^{-1}\)).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Ground and Surface Water Analyses (4I)
Electrical Conductivity and Salts (4I2)
Ion Chromatograph (4I2c)
Conductivity Detector (4I2c1)
Self-Regeneration Suppressor (4I2c1a)
Bromide, Chloride, Fluoride, Nitrate, Nitrite, Phosphate, and Sulfate (4I2c1a1-7)

1. Application
Nutrients (nitrogen and phosphorus), sediments, pesticides, salts, or trace elements in ground water and surface water affect soil and water quality (National Research Council, 1993). This procedure is developed for the analysis of ground or surface water.

2. Summary of Method
The water sample is filtered and is diluted according to its electrical conductivity \( (\text{EC}_s) \). The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to measure the
anion species and content. Standard anion concentrations are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. A computer program automates these actions. The water anions Br<sup>−</sup>, Cl<sup>−</sup>, F<sup>−</sup>, NO<sub>3</sub><sup>−</sup>, NO<sub>2</sub><sup>−</sup>, PO<sub>4</sub><sup>3−</sup>, and SO<sub>4</sub><sup>2−</sup> are reported in meq L<sup>−1</sup> (mmol (−) L<sup>−1</sup>) in methods 4I2c1a1-7, respectively.

3. Interferences

Some water samples contain suspended solids and require filtering. Organic anions that have low molecular weight will co-elute with inorganic anions from the column.

4. Safety

Wear protective clothing and safety glasses. Exercise special care when preparing reagents. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.

5. Equipment

5.1 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.2 Tubes, 50-mL, with caps
5.3 Guard column, IonPac AG-18, 4 x 50 mm, Dionex Corp., Sunnyvale, CA
5.4 Analytical column, IonPac AS-18, Dionex Corp., Sunnyvale, CA
5.5 Self-regeneration suppressor, ASRS–300, 4-mm, Dionex Corp., Sunnyvale, CA
5.6 Autosampler, AS-40, Dionex Corp., Sunnyvale, CA
5.7 Computer with PeakNet software and printer
5.8 Digital diluter/dispenser, with syringes 10,000-µL and 1000-µL, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.9 Poly-vials with caps, 5-mL, Dionex Corp., Sunnyvale, CA

6. Reagents

6.1 Reverse osmosis deionized filtered (RODI), ASTM Type I grade of reagent water
6.2 Helium gas
6.3 Primary stock standards solutions, (PSSS<sub>1000</sub>), high purity, 1000 mg L<sup>−1</sup>: Cl<sup>−</sup>, SO<sub>4</sub><sup>2−</sup>, F<sup>−</sup>, NO<sub>3</sub><sup>−</sup>, NO<sub>2</sub><sup>−</sup>, Br<sup>−</sup>, and PO<sub>4</sub><sup>3−</sup>
6.4 Mixed Calibration Standards Solutions (MCSS), A, B, C, and D and Blank as follows:
   6.4.1 MCSSA=In a 500-mL volumetric flask, add as follows
6.4.1.1 32 mL Cl\(^-\) PSSS\(_{1000}\) = 64 mg L\(^{-1}\)
6.4.1.2 32 mL SO\(_4^{2-}\) PSSS\(_{1000}\) = 64 mg L\(^{-1}\)
6.4.1.3 2 mL F\(^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
6.4.1.4 8 mL NO\(_3^-\) PSSS\(_{1000}\) = 16 mg L\(^{-1}\)
6.4.1.5 2 mL NO\(_2^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
6.4.1.6 2 mL Br\(^-\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)
6.4.1.7 2 mL PO\(_4^{3-}\) PSSS\(_{1000}\) = 4 mg L\(^{-1}\)

Dilute to volume with RODI water and invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.2 MCSSB: In a 100-mL volumetric flask, add 50 mL MCSSA and dilute to volume with RODI water. Final concentrations are 32, 32, 2, 8, 2, 2, and 2 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.3 MCSSC: In a 100-mL volumetric flask, add 50 mL MCSSB and dilute to volume with RODI water. Final concentrations are 16, 16, 1, 4, 1, 1, and 1 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Invert to thoroughly mix. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.4 MCSSD: In a 100-mL volumetric flask, add 50 mL MCSSC and dilute to volume with RODI water. Invert to thoroughly mix. Final concentrations are 8, 8, 0.5, 2, 0.5, 0.5, and 0.5 mg L\(^{-1}\) Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\), respectively. Store in plastic containers in the refrigerator. Prepare fresh weekly.

6.4.5 MCSS Blank: 0 mL of Cl\(^-\), SO\(_4^{2-}\), F\(^-\), NO\(_3^-\), NO\(_2^-\), Br\(^-\), and PO\(_4^{3-}\).

Dilute RODI water to volume.

7. Procedure

7.1 Water sample is filtered into a 50-mL tube and capped. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

**Dilution of sample extracts**

7.2 To estimate the total soluble anion concentration (meq L\(^{-1}\)), multiply the EC\(_s\) (procedure 4F2b1) by 10. Subtract the CO\(_3^{2-}\) and HCO\(_3^-\) concentrations (procedures 4F2c1c1a1-2) from the total anion concentration. The remainder is the approximate concentration (meq L\(^{-1}\)) of anions to be separated by ion chromatography.

Anion concentration (meq L\(^{-1}\)) = EC\(_s\) x 10 - (HCO\(_3^-\)+CO\(_3^{2-}\))
7.3 Dilute the saturation extract with the RODI water as follows:

<table>
<thead>
<tr>
<th>ECs</th>
<th>Dilution Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>(dS cm(^{-1}))</td>
<td></td>
</tr>
<tr>
<td>0.00 to 0.54</td>
<td>1</td>
</tr>
<tr>
<td>0.55 to 0.66</td>
<td>2</td>
</tr>
<tr>
<td>0.70 to 1.13</td>
<td>5</td>
</tr>
<tr>
<td>1.14 to 1.47</td>
<td>10</td>
</tr>
<tr>
<td>1.48 to 2.10</td>
<td>20</td>
</tr>
<tr>
<td>2.11 to 4.00</td>
<td>60</td>
</tr>
<tr>
<td>4.01 to 8.83</td>
<td>100</td>
</tr>
<tr>
<td>8.84 to 11.8</td>
<td>150</td>
</tr>
<tr>
<td>11.9 to 26.5</td>
<td>250</td>
</tr>
<tr>
<td>26.6 to 38.7</td>
<td>400</td>
</tr>
<tr>
<td>38.8 to 80.6</td>
<td>1000</td>
</tr>
<tr>
<td>&gt;80.7</td>
<td>2000</td>
</tr>
</tbody>
</table>

7.4 Place the MCSS (A, B,C, D and Blank) and diluted extract samples in the Poly-vials and cap with filter caps.

**Set-up and Operation of Ion Chromatograph (IC)**

7.5 Refer to the manufacturer’s manual for the operation of chromatograph. Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex ICS-2000 ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. Ranges and/or (typical settings) are as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range and/or (Typical Setting)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calibration</td>
<td>Peak Height or (area)</td>
</tr>
<tr>
<td>Flow setting</td>
<td>0.5 to 4.5 mL min(^{-1}) (1.00 mL min(^{-1}))</td>
</tr>
<tr>
<td>Pressure</td>
<td>200 to 3000 psi (2200 to 2400 psi)</td>
</tr>
<tr>
<td>Detection</td>
<td>Suppressed conductivity</td>
</tr>
<tr>
<td>Total conductivity</td>
<td>0 to 999.9 µS</td>
</tr>
<tr>
<td>Injection volume</td>
<td>10 µL</td>
</tr>
<tr>
<td>Auto offset</td>
<td>-999.9 to 999.9 µS (On)</td>
</tr>
<tr>
<td>Cell temperature</td>
<td>35 ºC</td>
</tr>
<tr>
<td>Suppressor current</td>
<td>75 mA</td>
</tr>
</tbody>
</table>
7.6 Load the sample holder cassettes with the capped samples, standards, and check samples.

7.7 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

**IC Calibration and Analysis**

7.8 Calibrate the instrument by using the MCSS (A, B, C, D, and Blank). The data system then associates the concentrations with the instrument responses for each MCSS. Rejection criteria for MCSS is $R^2 < 0.99$.

7.9 If samples are outside calibration, dilute sample extracts with RODI water solution and re-analyze.

7.10 Perform one quality control (QC) (Low Standard MCSS, Standard C) for every 12 samples. If reading is not within tolerance limits (10 to 15%, based on analyte), the instrument is re-calibrated and QC re-analyzed.

7.11 Record analyte readings to 0.01 mg L$^{-1}$.

8. Calculations

The instrument readings for analyte concentration are in mg L$^{-1}$. These analyte concentrations are converted to meq L$^{-1}$ as follows:

$$\text{Analyte Concentration in Soil (meq L}^{-1}) = \frac{A \times B}{C}$$

where:

- $A =$ Analyte ($\text{Br}^-, \text{Cl}^-, \text{F}^-, \text{NO}_3^-, \text{NO}_2^-, \text{PO}_4^{3-}, \text{SO}_4^{2-}$) concentration in extract (mg L$^{-1}$)
- $B =$ Dilution ratio, if needed
- $C =$ Equivalent weight

where:

- $\text{Cl}^- = 35.45$ mg meq$^{-1}$
- $\text{SO}_4^{2-} = 48.03$ mg meq$^{-1}$
- $\text{F}^- = 19.00$ mg meq$^{-1}$
- $\text{NO}_3^- = 62.00$ mg meq$^{-1}$
- $\text{NO}_2^- = 46.00$ mg meq$^{-1}$
- $\text{Br}^- = 79.90$ mg meq$^{-1}$
- $\text{PO}_4^{3-} = 31.66$ mg meq$^{-1}$

9. Report

Report the water extraction anions ($\text{Br}^-, \text{Cl}^-, \text{F}^-, \text{NO}_3^-, \text{NO}_2^-, \text{PO}_4^{3-}, \text{SO}_4^{2-}$) to the nearest 0.1 meq L$^{-1}$ (mmol (−) L$^{-1}$).
10. **Precision and Accuracy**

   Precision and accuracy data are available from the KSSL upon request.

11. **References**


---

**Ground and Surface Water Analyses (4I)**

**Electrical Conductivity and Salts (4I2)**

- **Automatic Titrator (4I2d)**
  - **Combination pH-Reference Electrode (4I2d1)**
    - **Acid Titration, H₂SO₄ (4I2d1a)**
    - **Carbonate and Bicarbonate (4I2d1a1-2)**

1. **Application**

   Nutrients (nitrogen and phosphorus), sediments, pesticides, salts, or trace elements in ground water and surface water affect soil and water quality (National Research Council, 1993). This procedure is developed for the analysis of ground or surface water.

2. **Summary of Method**

   The water sample is filtered and aliquot titrated on an automatic titrator to pH 8.25 and pH 4.60 end points. The carbonate and bicarbonate are calculated from the titers, aliquot volume, blank titer, and acid normality. Carbonate and bicarbonate are reported in meq L\(^{-1}\) (mmol (\(\sim\)) L\(^{-1}\)) in methods 4I2d1a1-2, respectively.

3. **Interferences**

   Clean the electrode by rinsing with distilled water and patting it dry with tissue. Wiping the electrode dry with a cloth, laboratory tissue, or similar material may cause electrode polarization.

   Slow electrode response time may cause overshoot at the end point. A combination of slowing the buret speed and increasing the time delay may help. Cleaning the electrode with detergent may decrease the response time. If all else fails, changing the electrode generally solves the problem. Blanks may not titrate properly because some sources of reverse osmosis (RO) water have a low pH.

4. **Safety**

   Wear protective clothing and eye protection. Exercise care when preparing reagents. Thoroughly wash hands after handling reagents. Restrict the use of
concentrated $\text{H}_2\text{SO}_4$ to the fume hood. Use showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow the manufacturer’s safety precautions when operating the automatic titrator.

5. Equipment

5.1 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
5.2 Tubes, 50-mL, with caps
5.3 Automatic titrator, Metrohm 670 Titroprocessor, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.
5.4 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.
5.5 Pipettes, electronic digital, 2500-µL and 10-mL, with tips, 2500-µL and 10-mL

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Helium gas
6.3 Sulfuric acid ($\text{H}_2\text{SO}_4$), concentrated, 36 $N$
6.4 $\text{H}_2\text{SO}_4$, 0.0240 $N$ standardized. Carefully dilute 2.67 mL of concentrated $\text{H}_2\text{SO}_4$ in 4 L of RODI degassed water ($\approx$15 min). Re-standardize the acid at regular intervals. Refer to the procedure for standardization of acids.
6.5 Borax pH buffers, pH 4.00, 7.00, and 9.18, for titrator calibration, Beckman, Fullerton, CA

7. Procedure

7.1 Water sample is filtered into a 50-mL tube and capped. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
7.2 Pipette 3 mL of the water sample into a 250-mL titration beaker.
7.3 Add 72 mL of RO water into a titration beaker. Final volume is 75 mL for blanks and samples. Run 8 to 12 blanks of RO water through the titration procedure.
7.4 Refer to manufacturer’s manual for operation of the automatic titrator.
7.5 Calibrate automatic titrator with pH 9.18, 7.00 and 4.00 buffers. Set-up the automatic titrator to set end point titration mode. The “Set” pH parameters are listed as follows:
7.6 Place the 250-mL titration beakers in the sample changer.

7.7 Press “Start.”

7.8 If the titrator is operating properly, no other analyst intervention is required. The titers and other titration parameters are recorded on the Titroprocessor printer.

8. Calculations

8.1 \[ \text{CO}_3^{2-} (\text{meq L}^{-1}) = \frac{(2T_1 \times N \times 1000)}{\text{Aliquot}} \]

8.2 \[ \text{HCO}_3^{-} (\text{meq L}^{-1}) = \frac{[(T_2 + T_1) - \text{Blank} - (2 \times T_1) \times N \times 1000]}{\text{Aliquot}} \]

where:

- \( T_1 \) = Titer of \( \text{CO}_3^{2-} \) (mL)
- \( T_2 \) = Titer of \( \text{HCO}_3^{-} \) (mL)
- \( N \) = Normality of \( \text{H}_2\text{SO}_4 \)
- \( \text{Blank} \) = Average titer of blank solutions (mL)
- \( \text{Aliquot} \) = Volume of saturation extract titrated (mL)
- 1000 = Conversion factor to meq L\(^{-1}\)

9. Report

Report saturation extract \( \text{CO}_3^{2-} \) and \( \text{HCO}_3^{-} \) to the nearest 0.1 meq L\(^{-1}\) (mmol (−) L\(^{-1}\)).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References

Ground and Surface Water Analyses (4I)

Total Analysis (4I3)
  Inductively Coupled Plasma Mass Spectrophotometer (4I3b)
    Aluminum, Arsenic, Barium, Boron, Cadmium, Calcium, Chromium, Cobalt, Copper, Iron, Lead, Magnesium, Manganese, Molybdenum, Nickel, Phosphorus, Potassium, Selenium, Silicon, Sodium, Strontium, and Zinc (4I3b1-22)

1. Application
   Nutrients (nitrogen and phosphorus), sediments, pesticides, salts, or trace elements in ground water and surface water affect soil and water quality (National Research Council, 1993). This procedure is developed for the analysis of the elemental content of ground or surface water.

2. Summary of Method
   The water is filtered and acidified with HCl. Calibration standards plus a blank are prepared for elemental analysis. The concentration of Al, As, B, Ba, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Mo, Na, Ni, P, Pb, Se, Si, Sr, and Zn are determined by inductively coupled plasma mass spectrophotometer (mg L\(^{-1}\)) in methods 4I3b1-22, respectively.

3. Interferences
   Interferences are corrected or minimized by using an internal standard, collision/reaction cell technology, and careful selection of specific masses for data reporting. Interference corrections are made by ICP–MS software.

4. Safety
   Wear protective clothing and eye protection. Exercise special care when preparing reagents.

5. Equipment
  5.1 Pipettes, electronic digital, 250-µL and 10-mL, Rainin Instrument Co., Woburn, MA
  5.2 Inductively coupled plasma mass spectrophotometer (ICP–MS), Agilent 7500cx, Agilent Technologies Inc. Wilmington, DE
  5.3 Computer, with ICP–MS ChemStation software ver. B.03.07, Agilent Technologies Inc. Wilmington, DE
  5.4 Heat exchanger, G1879B, Agilent Technologies
  5.5 Compressed gasses, argon (minimum purity 99.99%), hydrogen (minimum purity 99.999%) and helium (minimum purity 99.999%)
5.6 Autosampler, ASX-500 Series, Agilent Technologies Inc. Wilmington, DE
5.7 Quartz torch, for use with HMI, Part No. G3270-80027
5.8 Peristaltic pump (for automatic injection of internal standard)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Concentrated hydrochloric acid (HCl), 12 \( N \), trace pure grade
6.3 Concentrated nitric acid (HNO\(_3\)), 16 \( N \), trace pure grade
6.4 Selenium Standard, commercially prepared solution containing 1000 mg/mL Se
6.5 Water Extractable Elements, commercially prepared solution containing 1000 \( \mu \)g/mL Ca, K, Mg; 150 \( \mu \)g/mL P, Sr; 100 \( \mu \)g/mL Al, Ba, Fe, Mn; 50 \( \mu \)g/mL Cu, V, Zn; 10 \( \mu \)g/mL Co, Cr, Ni, Pb; 5 \( \mu \)g/mL As, Cd; and 1 \( \mu \)g/mL Mo
6.6 Phosphorus Standard, commercially prepared solution containing 1000 mg/mL P
6.7 Boron Standard, commercially prepared solution containing 1000 mg/mL B
6.8 Silicon Standard, commercially prepared solution containing 1000 mg/mL Si
6.9 Gold Standard, commercially prepared solution containing 1000 mg/mL Au
6.10 Lithium\(^{6}\) Standard, commercially prepared solution containing 1000 mg/mL Li\(^{6}\)
6.11 Scandium Standard, commercially prepared solution containing 1000 mg/mL Sc
6.12 Germanium Standard, commercially prepared solution containing 1000 mg/mL Ge
6.13 Yttrium Standard, commercially prepared solution containing 1000 mg/mL Y
6.14 Terbium Standard, commercially prepared solution containing 1000 mg/mL Tb
6.15 Bismuth Standard, commercially prepared solution containing 1000 mg/mL Bi
6.16 Agilent Stock Tuning Solution
6.17 Agilent 7500 Series PA Tuning 1 Solution

7. Procedure

**ICP–MS Calibration Standards, Set-up, and Operations**

7.1 Tuning Solution: In a 1-L volumetric flask, add 300 mL RODI water, 1 mL
commercially prepared Agilent Stock Tuning Solution (Reagent 6.16), and 18 mL concentrated HNO₃. Fill to volume with RODI water and mix well.

7.2 PA Tuning Solution: In 1-L volumetric, add 500 mL RODI water, 18 mL concentrated HNO₃, and 100 mL of commercially prepared Agilent 7500 Series PA Tuning 1 (Reagent 6.17). Fill to volume and mix well.

7.3 10 µg/mL Selenium Stock: Add 5 mL 1,000 µg/mL Se (Reagent 6.4) and 9 mL concentrated HNO₃ to 500-mL flask. Fill to volume with RODI water and mix well.

7.4 Mixed Elements Stock: In 500-mL flask, add 300 mL RODI water, 9 mL concentrated HNO₃, 5 mL Water Extractable Elements (Reagent 6.5), 5 mL 10 µg/mL Se stock (Reagent 7.3), and 9 mL concentrated HNO₃. Fill to volume with RODI water and mix well.

7.5 Mixed Elements High: In 500-mL flask, add 300 mL RODI water and 50 mL Mixed Elements Stock (Reagent 7.4). Fill to volume with RODI water and mix well.

7.6 Mixed Elements Medium: In 500-mL flask, add 300 mL RODI water and 50 mL Mixed Elements High (Reagent 7.5). Fill to volume with RODI and mix well.

7.7 Mixed Elements Low: In a 500-mL flask, add 300 mL RODI water and 50 mL Mixed Elements Medium (Reagent 7.6). Fill to volume with RODI and mix well.

7.8 P1000: In 1-L volumetric flask, add 500 mL RODI water and 1 mL Phosphorus Standard (Reagent 6.6). Fill to volume with RODI water and mix well.

7.9 P100: In 1-L volumetric flask, add 500 mL RODI water and 100 mL of P1000 (Reagent 7.8). Fill to volume with RODI water and mix well.

7.10 B1000, In a 1-L volumetric flask add 500 mL DIRO and 1 mL Boron Standard (Reagent 6.7). Fill to volume with RODI water and mix well. Transfer to 1-L polypropylene bottle.

7.11 B100: In 1-L volumetric flask, add 500 mL RODI and 100 mL B1000 (Reagent 7.10). Fill to volume with RODI and mix well. Transfer to 1-L polypropylene bottle.

7.12 Si1000: In 1-L volumetric flask, add 500 mL RODI water and 1 mL Silicon Standard (Reagent 6.8). Fill to volume with RODI and mix well.

7.13 Si100: In 1-L volumetric flask, add 500 mL RODI and 100 mL Si1000 (Reagent 7.12). Fill to volume with RODI water and mix well.

7.14 Internal Standard (1 µg/mL Li⁶, Sc, Ge, Y, In, Tb, Bi): In 1-L flask, add 300 mL RODI water, 18 mL concentrated HNO₃, 6 mL concentrated HCl, 0.250
mL 1000 µg/mL Au (Reagent 6.9), and 1 mL each of 1000 µg/mL Li, Sc, Ge, Y, In, Tb, and Bi (Reagents 6.10–6.15). Fill to volume with RODI water and mix well.

7.15 Rinse: In 2-L flask, add 300 mL RODI water and 58 mL concentrated HNO₃. Fill to volume with RODI and mix well.

7.16 Rinse #1: In 1-L flask, add 300 mL RODI water, 29 mL concentrated HNO₃, and 1 mL 1000 µg/mL Au (Reagent 6.9). Fill to volume with RODI water and mix well.

7.17 Rinse #2: In 1-L flask, add 300 mL RODI, 15 mL concentrated HNO₃, 45 mL concentrated HCl, and 1 mL of 1000 µg/mL Au (Reagent 6.9). Fill to volume with RODI water and mix well.

7.18 Rinse #3 RODI water.

7.20 Standard concentrations in µg/mL for each element.

<table>
<thead>
<tr>
<th>Element</th>
<th>Blank</th>
<th>Mixed Elements Low</th>
<th>Mixed Elements Mid</th>
<th>Mixed Elements High</th>
<th>P100</th>
<th>P1000</th>
<th>B100</th>
<th>B1000</th>
<th>Si100</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>0</td>
<td>1</td>
<td>10</td>
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<tr>
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<tr>
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<td>---</td>
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<td>---</td>
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</tbody>
</table>
7.21 Reporting m/z and tune step for each element analyzed.

<table>
<thead>
<tr>
<th>Element</th>
<th>m/z</th>
<th>Tune 1 (H₂)</th>
<th>Tune 2 (He)</th>
<th>Tune 3 (No Gas)</th>
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</thead>
<tbody>
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<td>Zn</td>
<td>66</td>
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7.22 The ICP–MS is set up with an HMI system using a Burgener Mira mist nebulizer, quartz torch, and spray chamber to analyze samples. Internal standard is added via peristaltic pump using 0.19 mm id. pump tubing. Internal standard is mixed with the samples, or standards are added via coil prior to entering the nebulizer. Samples are diluted 1:10 or greater as necessary prior to analysis with sample diluent (Reagent 6.17). Perform instrument checks (tune for sensitivity, resolution axis, P/A factor, internal standard RSD, torch alignment, EM tune) prior to analysis as discussed in
operation manual of instrument. Check instrument gas pressures to ensure pressures are correct and of adequate supply.

7.23 Typical tune values for 1:5 aqueous extraction are as follows:

<table>
<thead>
<tr>
<th><strong>Plasma Parameters</strong></th>
<th><strong>Q-Pole Parameters</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>RF power 1550 W</td>
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</tr>
<tr>
<td>RF matching 1.78 V</td>
<td>AMU offset 125</td>
</tr>
<tr>
<td>Smpl depth 10.0 mm</td>
<td>Axis gain 0.9994</td>
</tr>
<tr>
<td>Torch-H 0.4 mm</td>
<td>Axis offset 0.01</td>
</tr>
<tr>
<td>Torch-V -0.4 mm</td>
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<tr>
<td>Carrier gas 0.50 L/min</td>
<td>Octapole Parameters</td>
</tr>
<tr>
<td>Makeup gas 0.50 L/min</td>
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<td>Optional gas 0.0</td>
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<tr>
<td>Extract 1 0.0 V</td>
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<tr>
<td>Extract 2 -135.0 V</td>
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</tr>
<tr>
<td>Omega Bias-ce -24 V</td>
<td>Pulse HV 1360 V</td>
</tr>
<tr>
<td>Omega Lens-ce -1.0 V</td>
<td></td>
</tr>
<tr>
<td>Cell entrance -40 V</td>
<td></td>
</tr>
<tr>
<td>QP focus -8 V</td>
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</tr>
<tr>
<td>Cell exit -40 V</td>
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<tr>
<td><strong>Plasma Parameters</strong></td>
<td><strong>Q-Pole Parameters</strong></td>
</tr>
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</tr>
<tr>
<td>Smpl depth 10.0 mm</td>
<td>Axis gain 0.9994</td>
</tr>
<tr>
<td>Torch-H 0.4 mm</td>
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<tr>
<td>Makeup gas 0.50 L/min</td>
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<tr>
<td>Optional gas 0.0</td>
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<td>Nebulizer pump 0.10 rps</td>
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</tr>
<tr>
<td>Sample pump 0.0</td>
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<tr>
<td>S/C temp 2 °C</td>
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<th><strong>Ion Lenses</strong></th>
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</thead>
<tbody>
<tr>
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<tr>
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<td>OctP bias -6.0 V</td>
</tr>
<tr>
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<tr>
<td>Omega Lens-ce -1.0 V</td>
<td></td>
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<tr>
<td>Cell entrance -40 V</td>
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<td>He gas 4.5 mL/min</td>
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<td>Plasma Parameters</td>
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<tr>
<td>Torch-H</td>
<td>0.4 mm</td>
</tr>
<tr>
<td>Torch-V</td>
<td>-0.4 mm</td>
</tr>
<tr>
<td>Carrier gas</td>
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</tr>
<tr>
<td>Makeup gas</td>
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<tr>
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<td>Cell entrance</td>
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<tr>
<td>He gas</td>
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<td>Pulse HV</td>
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7.24 Establish detection limits using the blank standard solution. The instrumental detection limits are calculated using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are reported as “ND” or non-detected.

8. Calculations
With the HCl treatment (0.5 mL per 10 mL water) in the calculations, the concentrations are then reported directly, unless additional dilutions are performed because of high analyte concentrations.

9. Report
Data are reported to the nearest 0.01 mg L$^{-1}$.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

ANALYSIS OF ORGANIC SOILS OR MATERIALS (5)

Mineral Content (5A)

1. Application
The mineral content consists of the plant ash and soil particles that remain after removal of organic matter. The percentage of organic matter lost on ignition can be used to define organic soils in place of estimates of organic matter by the Walkley-Black organic C method (6A1c, method obsolete). The determination of organic matter by loss on ignition is a taxonomic criterion for organic soil materials (Soil Survey Staff, 2014). Organic C data by Walkley-Black are generally considered invalid if organic C >8%.

2. Summary of Method
Dry sample overnight at 110 °C in moisture can. Cool and weigh. Place sample in a cold muffle furnace and raise the temperature to 400 °C. Heat sample
overnight (16 h), cool, and weigh. The ratio of the weights (400 °C/110 °C) is the mineral content percentage (method 5A).

3. Interferences
The sample must be placed in a cold muffle furnace to prevent rapid combustion and sample splattering.

4. Safety
Use caution when the muffle furnace is hot. Wear protective clothing and goggles. Handle the heated material with tongs.

5. Equipment
5.1 Metal weighing tins
5.2 Oven, 110 °C
5.3 Muffle furnace, 400 °C
5.4 Electronic Balance, ±0.01-g sensitivity

6. Reagents
None.

7. Procedure
7.1 Place a 10 to 15 g sample in a tared weighing tin.
7.2 Dry sample at 110 °C overnight.
7.3 Remove sample from oven, cap, and cool in a desiccator.
7.4 When cool, record weight to nearest 0.01 g.
7.5 Place sample and weighing tin in a cold muffle furnace. Raise temperature to 400 °C. Heat overnight (16 h).
7.6 Remove sample from oven, cap, and cool in a desiccator.
7.7 When cool, record sample weight to nearest 0.01 g.

8. Calculations
8.1 Mineral Content (%) = \((R_w/OD_w) \times 100\)
where:
\(R_w\) = Residue weight after ignition
\(OD_w\) = Oven-dry soil weight
Organic matter percent can then be calculated as follows:

8.2 Organic Content (%) = 100 − Mineral Content (%)
9. **Report**
   Report mineral content to the nearest whole percent.

10. **Precision and Accuracy**
    Precision and accuracy data are available from the KSSL upon request.

11. **References**

---

**Pyrophosphate Color (5B)**

1. **Application**
   Decomposed organic materials are soluble in sodium pyrophosphate. The combination of organic matter and sodium pyrophosphate forms a solution color that correlates with the decomposition state of the organic materials. Dark colors are associated with sapric materials, and light colors are associated with fibric materials (Soil Survey Staff, 2014).

2. **Summary of Method**
   Organic material is combined with sodium pyrophosphate. After standing, the color is evaluated by moistening a chromatographic strip in the solution and comparing the color with standard Munsell color charts (5B).

3. **Interferences**
   This test of organic soil material can be used in field offices. Because it is not practical in the field to base a determination on a dry sample weight, moist soil is used. The specific volume of moist material depends on how it is packed. The packing of the material, therefore, must be standardized in order to obtain comparable results by different soil scientists (Soil Survey Staff, 2014).

4. **Safety**
   Use caution when handling sodium pyrophosphate.

5. **Equipment**
   5.1 Polycons, 30-mL, Richards Mfg. Co.
   5.2 Chromatographic paper, Schleicher and Schuell no. 470 A-3
   5.3 Munsell Color Book, 10YR and 7.5YR pages
   5.4 Half-syringe, 6-mL. Cut plastic syringe longitudinally to form a half-cylinder measuring device.
   5.5 Scissors
5.6 Paper towel
5.7 Tweezers
5.8 Metal spatula

6. Reagents
6.1 Sodium pyrophosphate (Na₄P₂O₇•10H₂O)
6.2 Reverse osmosis (RO) water

7. Procedure

Sample Preparation

7.1 Prepare soil material. If the soil is dry, add water and let stand to saturate. Place 50 to 60 mL of a representative sample on a paper towel in a linear mound. Roll the towel around the sample and express water if necessary. Use additional paper towels as external blotters. Remove the sample and place on a fresh paper towel. The sample should be firm but saturated with water.

7.2 Use scissors to cut sample into 5- to 10-mm long segments.

7.3 Randomly select sample segments for determination of fiber (5C), solubility in pyrophosphate (5B), and pH (4C1a1a4).

Pyrophosphate

7.4 Dissolve 1 g (heaping 1/6 tsp) of sodium pyrophosphate in 4 mL of water in a 30-mL polycon container. Allow to equilibrate for 5 min.

7.5 Use a metal spatula to pack a half-syringe that is adjusted to the 5-mL mark or 2.5-mL (2.5-cm³) volume with the moist sample.

7.6 Transfer soil material cleanly into the container that holds the pyrophosphate solution.

7.7 Mix thoroughly using a wooden stirrer or metal spatula. Cover and let stand overnight.

7.8 Mix sample again next morning.

7.9 Use tweezers to insert a strip of chromatographic paper vertically into the sample to a 1-cm depth. Let stand until the paper strip has wetted to a 2-cm height above slurry surface. Generally, sample needs to stand ≈5 min but may stand longer if cover is closed. Remove the paper strip with tweezers. Cut strip and leave in the slurry that portion to which the soil adheres.

7.10 Place the strip on a piece of blotting paper and press gently with tweezers to make even contact.

7.11 Remove paper strip with tweezers and compare color of the strip to Munsell color charts.
8. Calculations
   No calculations.

9. Report
   Report color using Munsell color notation.

10. Precision and Accuracy
    Precision and accuracy data are not available for this procedure. Experienced analysts can usually reproduce results within ±1 color chip.

11. References

Fiber Volume (5C)

1. Application
   The water-dispersed fiber volume is a method to characterize the physical decomposition state of organic materials. The decomposition state of organic matter is used in soil taxonomy to define sapric, hemic, and fibric organic materials (Soil Survey Staff, 2014). Sapric material passes through a 100-mesh sieve (0.15-mm openings). Fibers are retained on the sieve. As defined in soil taxonomy, organic materials that are >2 mm in cross section and that are too firm to be readily crushed between thumb and fingers are excluded from fiber.

2. Summary of Method
   The sample is prepared to a standard water content. The unrubbed fiber content is determined in a series of three steps designed to remove the sapric material by increasingly vigorous treatments. The rubbed fiber content is determined by rubbing the sample between the thumb and fingers. The percent unrubbed fiber after each step and the final unrubbed and rubbed fiber are reported (5C).

3. Interferences
   This test of organic soil material can be used in field offices. Because it is not practical in the field to base a determination on a dry sample weight, moist soil is used. The specific volume of moist material depends on how it is packed. The packing of the material, therefore, must be standardized in order to obtain comparable results by different soil scientists (Soil Survey Staff, 2014).

4. Safety
   Use caution when using electrical equipment.
5. Equipment
5.1 Half-syringe, 6-mL. Cut plastic syringe longitudinally to form a half-cylinder measuring device.
5.2 Sieve, 100 mesh, 7.6-cm diameter
5.3 Eggbeater
5.4 Microscope or hand lens
5.5 Electric mixer, Hamilton Beach no. 35
5.6 Scissors
5.7 Paper towel
5.8 Metal spatula

6. Reagents
   Reverse osmosis (RO) water

7. Procedure

Sample Preparation

7.1 Prepare soil material. If the soil is dry, add water and allow to stand until saturated. Place 50 to 60 mL of a representative sample on a paper towel in a linear mound. Roll the towel around the sample and gently squeeze to express water if necessary. Use additional paper towels as external blotters. Remove the sample and place on a fresh paper towel. The sample should be firm but saturated with water.

7.2 Use scissors to cut sample into 0.5- to 1.0-cm length segments.

7.3 Randomly select sample segments for determination of fiber (method 5C), solubility in pyrophosphate (method 5B), and pH (method 4C1a1a4).

Unrubbed Fiber: Overview

7.4 The unrubbed fiber procedure involves a series of three steps designed to disperse sapric material by increasingly vigorous treatments. All three steps may not be necessary. Following each step that is performed, the percentage estimate of sapric material remaining is visually determined under a microscope or hand lens. Categories used to estimate the remaining sapric component are as follows:

7.4.1 Clean (<1% sapric)
7.4.2 Nearly clean (1 to 10% sapric)
7.4.3 Some sapric (10 to 30% sapric)
7.4.4 Sapric (>30% sapric)
Unrubbed Fiber: Part 1

7.5 Use a metal spatula to pack a half-syringe that is adjusted to the 5-mL mark or 2.5-mL (2.5 cm³) volume with the moist sample.

7.6 Transfer all the soil material to a 100-mesh sieve and wash under a stream of tapwater, adjusted to deliver 200 to 300 mL in 5 s. Wash sample until the water passing through the sieve appears clean. To more clearly determine the end point, catch the effluent in a white plastic container. Periodically empty the container until the effluent runs nearly clean.

7.7 Examine the sample under a microscope or hand lens to determine if sample is free of sapric material.

7.8 If sample is >10% sapric material, proceed to Unrubbed Fiber, Part 2. If sample is <10% sapric material, wash the residue to one side of the screen and blot from underneath with absorbent tissue to withdraw water and proceed on as follows with Unrubbed Fiber, Part 1.

7.9 Repack the residue into a half-syringe and blot again with absorbent tissue. The moisture content should be approximately that of the original sample.

7.10 Measure the volume by withdrawing the plunger and reading the value on the syringe scale. Record as a percentage of the initial 2.5-mL (2.5 cm³) volume.

7.11 Proceed with the Rubbed Fiber determination.

Unrubbed Fiber: Part 2

7.12 Transfer the residue obtained in Unrubbed Fiber, Part 1 to a 500-mL plastic container and fill about half full with water.

7.13 Stir vigorously with an eggbeater for 1 min.

7.14 Transfer to the 100-mesh sieve and repeat procedural steps in Unrubbed Fiber, Part 1 beginning with Section 7.9. If sample is >10% sapric material, proceed to Unrubbed Fiber, Part 3.

Unrubbed Fiber: Part 3

7.15 Transfer residue left from Unrubbed Fiber, Part 2 to an electric mixer container (malt mixer or blender) and fill to about two-thirds with water.

7.16 Mix for 1 min.

7.17 Transfer to a 100-mesh sieve and repeat Unrubbed Fiber Part 1 beginning with the washing procedure (Section 7.6).

7.18 Examine the residue under a microscope or hand lens and estimate the percentage of sapric material, if any.

7.19 Record the kind of fiber observed. Typical fibers are herbaceous, woody, and diatomaceous.

7.20 Blot the sample and measure the residue volume.
7.21 Proceed with the Rubbed Fiber determination.

**Rubbed Fiber**

7.22 Transfer the residue from the unrubbed fiber treatment to the 100-mesh sieve.

7.23 Rub sample between thumb and fingers under a stream of tapwater, adjusted to deliver 150 to 200 mL in 5 s, until water passing through the sieve is clean. Clean rubbed fibers should roll between the thumb and fingers rather than slide or smear.

7.24 Blot sample and measure volume in half-syringe.

8. **Calculations**

   Fiber volume (%) = Reading on half-syringe (mL) x 20

   where:

   Fiber volume = Rubbed + unrubbed fiber

9. **Report**

   Record the percentage of unrubbed fiber after each completed step. Report the final unrubbed fiber and the rubbed fiber to the nearest whole percent and report fiber type.

10. **Precision and Accuracy**

   No precision and accuracy data are available for this procedure.

11. **References**


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**Melanic Index (5D)**

1. **Application**

   Melanic and fulvic Andisols have high contents of humus, related to their soil color and reflecting pedogenic processes (Honna et al., 1988). Typically, Melanic Andisols are formed under grassland ecosystems, with humus dominated by A type humic acid (highest degree of humification); whereas Fulvic Andisols are found under forest ecosystems, with humus characterized by the high ratio of fulvic acid to humic acid (low degree of humification, e.g., P or B type humic acid) (Honna et al., 1988). The organic matter thought to result from large amounts of gramineous vegetation can be distinguished from organic matter formed under forest vegetation by the melanic index (Soil Survey Staff, 2014).
2. Summary of Method

A 0.5-g soil sample is mechanically shaken for 1 h in 25 mL of 0.5% NaOH solution. One drop of 0.2% Superfloc solution (flocculation aid) is added to sample and then mechanically shaken for 10 min. Either a 1 or 0.5 mL extract (<10% or >10% organic C, respectively) is pipetted into a test tube, followed by the addition of 20 mL of 0.1% NaOH solution and thorough mixing. Absorbance of the solution is read using a spectrophotometer at 450 and 520 nm, respectively, within 3 h after extraction. Melanic Index is calculated by dividing the absorbance at 450 nm by the absorbance at 520 nm (method 5D).

3. Interferences

No known interferences.

4. Safety

No significant hazards are associated with this procedure. Follow standard laboratory safety practices.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge tubes, 50-mL polypropylene
5.4 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.5 Pipettes, electronic digital, 1000-µL and 10-mL, with tips, 1000-µL and 10-mL
5.6 Dispenser, 30-mL or 10-mL
5.7 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.8 Spectrophotometer, UV-visible, Varian, Cary 50 Conc
5.9 Computer
5.10 Printer

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 NaOH, 0.5% and 0.1%
6.3 Superfloc 16, 0.2% (2 g L⁻¹) in RODI water

7. Procedure

7.1 Weigh 0.5 g of <2 mm or fine-grind air-dry soil to the nearest 1.0 mg into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈0.5 of air-dry soil.
7.2 Dispense 25 mL of 0.5% NaOH solution to the tube.

7.3 Transfer the sample to the shaker. Shake for 1 h at 200 oscillations min\(^{-1}\) at room temperature.

7.4 Remove the sample from the shaker. Add one drop of 0.2% Superfloc 16 solution and centrifuge at 4000 rpm for 10 min.

7.5 Use the pipette to transfer either 1 or 0.5 mL extract (<10% or >10% organic C, respectively) into test tube.

7.6 Add 20 mL of 0.1% NaOH solution and vortex.

7.7 Autozero the blank.

7.8 Set the spectrophotometer at 450 nm. Read absorbance.

7.9 Set the spectrophotometer at 520 nm. Read absorbance.

8. Calculations
   Melanic Index is calculated as follows:
   \[
   \text{Melanic Index} = \frac{\text{Absorbance at 450 nm}}{\text{Absorbance at 520 nm}}
   \]

9. Report
   Report Melanic Index.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References

Ratios and Estimates Related to Organic Matter (5E)
The KSSL Primary Characterization Data Sheets include several ratios and estimates associated with organic matter, using either estimated or measured C values. For more detailed information on these ratios, their calculations, and their applications, refer to the SSIR No. 45, “Soil Survey Laboratory Information Manual” (Soil Survey Staff, 2011). Additional information on these ratios and estimates can also be obtained from the KSSL upon request.

References
SOIL BIOLOGICAL AND PLANT ANALYSIS (6)

Soil is an ecosystem that contains a broad spectrum of biological components, representing many physiological types (Germida, 1993). The soil biota is critical to soil quality, affecting nutrient cycling, soil stability and erosion, water quality and quantity, and plant health (USDA–NRCS, 2004). Many components of this biota (e.g., fungi, bacteria, earthworms, protozoa, arthropods, and nematodes) and their relationship to soil health are discussed in the USDA–NRCS Soil Biology Primer (Tugel and Lewandowski, 2001). Also refer to Reeder et al. (2001) for information on root biomass and microbial biomass.

The KSSL performs several biological analyses, including but not limited to 0.02 M KMnO₄ extraction, oxidizable (POx C) or reactive carbon; particulate organic matter and mineral (HYPER-POM); and β-glucosidase assay.

References


Soil Analyses (6A)
0.02 M KMnO₄ Extraction (6A2)

UV-Visible Spectrophotometer, Dual Beam (6A2a)

Permanganate Oxidizable (POx C) or Reactive Carbon (6A2a1)

Air-Dry, <2 mm (6A2a1a1)

1. Application

This method, commonly called Weil Carbon (Weil et al., 2003), is designed to be a quick and easy field test for assessment of reactive soil organic C. Following the principle of bleaching chemistry, potassium permanganate (KMnO₄) is used to oxidize organic matter in soil. The oxidized organic matter is considered to be
associated with the reactive C pool (Blair et al., 1995). A reactive soil C index can be expressed as the quotient of reactive C to soil organic C (Blair et al., 2001). The stability of this index over time is considered to be a useful measure of soil quality (Islam and Weil, 1997).

2. Summary of Method

A 5-g sample is oxidized with 0.02 M potassium permanganate diluted with reverse osmosis water. Sample is vortexed and allowed to stand undisturbed for 10 minutes. A small aliquot of the supernatant is diluted with reverse osmosis water, and the absorbance of the solution is read at 550 nm using a spectrophotometer. Reactive carbon is reported in units of milligram POx C per kilogram oven dry soil (mg reactive carbon kg⁻¹).

The bleaching of the pink KMnO₄ color (reduction in absorbance) is proportional to the amount of POx C in soil, i.e., the KMnO₄ color loss (the lowering of the absorbance reading) is proportional to the amount of POx C in the soil (Weil et al., 2003). To estimate the amount of C oxidized, the method relies on the assumption of that 1 mol MnO₄⁻ is consumed (reduced from Mn⁷⁺ to Mn²⁺) in the oxidation of 0.75 moles (9000 mg) of C (Blair et al., 1995).

3. Interferences

Chemical oxidation methods for the determination of labile soil carbon have a number of limitations. Different soil samples may have variable amounts of readily oxidizable fractions, making standardization of any method difficult. Results are influenced by the amount of C in the sample, the concentration of MnO₄⁻, and the contact time (Blair et al., 1995).

4. Safety

Wear protective clothing (coats, aprons, and gloves) and eye protection (safety glasses and other devices as appropriate) while preparing reagents and performing procedure. Exercise special care when preparing reagents. Use a vented hood. Thoroughly wash hands after handling all chemicals.

Potassium permanganate is a strong oxidizer. Contact with other material may cause fire. Avoid contact with eyes, skin, and clothing. In case of fire, soak with water. In case of spill, sweep up and remove. Flush spill area with water. Inhalation of KMnO₄ dust may severely damage respiratory passages and/or lungs. Contact with skin or eyes may cause severe irritation or burns. Substance is readily absorbed through the skin. See the Material Safety Data Sheet (MSDS) for further information regarding KMnO₄.

5. Equipment

5.1 Centrifuge tubes, 50-mL, graduated, polyethylene, with screw tops
5.2 Dropper pipette, 1-mL, graduated, disposable
5.3 Squirt bottle
5.4 Electronic balance, 5-g (or electronic balance, ±0.01-g sensitivity)
5.5 Petri dish, for sun-drying crumbled soil, if necessary
5.6 Stopwatch, timer, or a watch with a second hand
5.7 Pipette, electronic digital, 10-mL
5.8 Volumetric flasks, 50-mL, 100-mL, 250-mL, and 2-L, with stoppers
5.9 Spectrophotometer, UV-Visible, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.10 Computer with Cary WinUV software, Varian Australia Pty Ltd., and printer
5.11 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.12 Vortexer

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 CaCl₂•2H₂O, 0.1 M solution. Dissolve 29.40 g of CaCl₂•2H₂O in a 2-L volumetric flask with 1L of RODI. Bring to 2-L volume. Store in a polyethylene bottle.
6.3 KOH, 0.1 M solution. Dissolve 0.561 g of potassium hydroxide with 50 mL of RODI water in a 100-mL volumetric flask. Bring to volume with RODI water. Invert to mix thoroughly.
6.4 Stock KMnO₄ solution, 0.2 M in 0.1 M CaCl₂ solution (pH 7.2). In a 250-mL volumetric, dissolve 7.90 g of potassium permanganate crystals in 100 mL of 0.1 M CaCl₂. Bring to volume with 0.1 M CaCl₂. Adjust solution pH to 7.2 with 0.1 M KOH (usually requires 1 or 2 drops of KOH). Solution is stable for approximately 3 days.
6.5 Working KMnO₄, 0.02 M. Add 200 mL of Stock KMnO₄ solution in a 2-L volumetric flask. Bring to volume with RODI and invert. Prepare fresh daily.
6.6 Standard KMnO₄ Working Solutions (SKMnO₄WS), 0.04, 0.02, 0.01, and 0.005 M KMnO₄. Prepare fresh weekly. Store in a refrigerator. Allow to equilibrate to room temperature before use. In four 50-mL volumetric flasks, add as follows: 10, 5, 2.5, and 1.25 mL of reagent 6.4 (0.2 M KMnO₄, in 0.1 M CaCl₂ solution, pH 7.2). Bring to volume with RODI water. Invert to mix thoroughly.
6.7 Standard KMnO₄ Calibration Solutions (SKMnO₄CS), 0.0004, 0.0002, 0.0001, 0.00005, and 0 M KMnO₄. To four 50-mL volumetric flasks, add 0.5-mL aliquots of the SKMnO₄WS. For the blank, add 5 mL 0.1 M CaCl₂. Make fresh daily. Bring to volume with RODI water. Invert to mix thoroughly.
7. Procedure

7.1 Weigh 5 g of <2-mm (sieved), air-dry soil to the nearest mg and add to centrifuge tube. Add 20 mL of Working KMnO₄, 0.2 M.

7.2 Vortex each sample and allow 10 min for soil to settle. Do not disturb during settling period.

7.3 After 10 min, centrifuge for 10 min at 2000 rpm.

7.4 Add 49.5 mL of RODI water to a clean, labeled centrifuge tube. Transfer 0.5 mL of the supernatant solution into the tube and mix.

7.5 Transfer sample extracts and SKMnO₄ CS to cuvettes.

7.6 Set the spectrophotometer to read at 550 nm. Autozero with calibration blank.

7.7 Calibrate the instrument by using the calibration solutions. The data system then associates the concentrations with the instrument responses for each calibration solution. Rejection criteria for calibration is $R^2 < 0.99$.

7.8 Run samples using calibration curve. Sample concentration is calculated from the regression equation. Record results to the nearest 0.01 unit for the sample extract and each calibration solution.

7.9 If samples have <0.00003 absorbance (A), reweigh smaller sample size (e.g., 2.50 g) and re-analyze. Samples that have low absorbance are those that have large amounts of reactive carbon.

8. Calculations

\[
\text{KMnO}_4 \text{ C (mg kg}^{-1} \text{)} = (0.02 \text{ mol L}^{-1} - A) \times (9000 \text{ mg C mol}^{-1}) \times \left(\frac{0.02 \text{ L solution}}{0.005 \text{ kg}}\right) \times \frac{\text{AD/OD}}{}
\]

where:

- 0.02 mol L\(^{-1}\) = initial solution concentration
- A = analyte reading (mol L\(^{-1}\))
- 9000 = mg C (0.75 mole) oxidized by 1 mole MnO\(_4\), changing from Mn\(^{7+}\) to Mn\(^{2+}\)
- 0.02 L = volume of KMnO\(_4\) solution reacted
- 0.005 = kg of soil used
- AD/OD = air-dry/oven-dry ratio (method 3D1)

9. Report

Report reactive C (mg kg\(^{-1}\)) as oxidizable C, potassium permanganate (POx C).

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.
11. References


Soil Analyses (6A)
Particulate Organic Matter and C-Mineral (6A4)

Total Analysis (6A4a)
   Dry Combustion (6A4a1)
      Thermal Conductivity Detector (6A4a1a)
      Carbon, Nitrogen, Sulfur (6A4a1a1-3)
      Air-Dry (6A4a1a1-3a)
         ≥53 µm, Particulate Organic Matter (6A4a1a1-3a1)
         <53 µm, C-Mineral, Calculated (6A4a1a1-3a3)

1. Application

Particulate organic matter (POM) is soil organic matter (SOM) associated with a physical fraction of soil, operationally defined at >53 µm in diameter (Elliott and Cambardella, 1991; Cambardella and Elliott 1992; Follett and Pruessner, 1997). Researchers are not in agreement on the exact dynamic character of POM. Some considering it to be a relatively labile component of SOM, whereas others have described it as slowly decomposable, or stabilized, SOM (Cambardella and Elliott, 1992). This fraction is similar to various sieved and physical fractions, such as resistant plant material (Jenkinson and Rayner, 1977), sand-size organic matter (Tiessen and Stewart, 1983), and size fractions (Gregorich et al., 1988). Although “light” fraction OM (LF) (Strickland and Sollins, 1987; Hassink, 1995) is often confused with POM, LF and POM are not the same. Particulate organic matter is a size-defined fraction, whereas LF is defined by density.

The quantity of POM measured in soils is sensitive to management practices. Under tillage, the POM fraction becomes depleted (Jenkinson and Rayner, 1977;
Cambardella and Elliott, 1992). Reductions of more than 50% have been found in long-term tillage plots (20-year). Measurable reductions are believed to occur in the range of 1 to 5 years (Cambardella and Elliott, 1992). When paired sampling is done between tillage practices or repeated sampling of the same plots through time, the impact of the management upon POM becomes more apparent (Marriot and Wander, 2006).

As an intermediately labile pool of soil carbon (C) and a tenable soil quality indicator, POM can be used in SOM modeling. Several models have been developed to estimate the dynamics of SOM, and all have incorporated at least two SOM components based on susceptibility to degradation—slow and rapid. Physical fractions, such as POM, have been found to be more useful in modeling applications than chemical fractions (e.g., fulvic and humic acids) (Hassink, 1995). A minimum data set for modeling soil organic carbon proposed by Gregorich (1994) includes POM as one of the primary parameters. Examples of models that incorporate these parameters can be found in Jenkinson and Rayner (1977) and in tests of the CENTURY Soil Organic Model (Parton et al., 1994; Metherell et al., 1993; Montavalli et al., 1994).

2. Summary of Method

The Hyper POM (HPOM) procedure is primarily the physical separation (method 1B3b2a) of <2 mm sieved soil into the ≥53-µm fraction and the <53-µm fraction (Cambardella and Elliott, 1992; Follett and Pruessner, 1997). HPOM-C and HPOM-N are measured directly by dry combustion analysis of the ≥53-µm fraction, whereas MIN-C and MIN-N (<53-µm fraction) are calculated by subtracting the HPOM-C and HPOM-N values from the direct measurement by combustion of total C and N of the <2 mm soil (method 4H2a1-3).

3. Interferences

Interferences include inorganic carbonates, which are, however, not commonly expected in surface horizons. If carbonates are present, the calculations need to be adjusted to account for them. Highly organic samples tend to clump or ball up during shaking. To break up these clumps, tubes are vortexed a few times during the 15 h shaking.

4. Safety

Always wear safety glasses when working with soils and glass containers.

5. Equipment

5.1 Pressure regulator for water, with stop cock attached to tubing
5.2 Centrifuge tube, 50-mL
5.3 Sieve, 270-mesh, 53-µm
5.4 Mechanical reciprocating shaker, 200 oscillations min$^{-1}$, 1½ strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI

5.5 Evaporating crucibles

5.6 Oven, 110 °C

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 0.5% sodium hexametaphosphate solution: Dissolve 20 g sodium hexametaphosphate in 4 liters RODI water, ASTM Type I grade of reagent water

7. Procedure

**Physical Separation of ≥53-µm and <53-µm Fractions**

7.1 Weigh 10 g of <2-mm (sieved), air-dry soil to the nearest mg into a 50-mL centrifuge tube. Add 30 mL 0.5% sodium hexametaphosphate solution to the tube. Highly organic samples may require an increased volume of sodium hexametaphosphate solution.

7.2 Stopper sample tightly and shake for 15 h (overnight) at 200 oscillations min$^{-1}$ at room temperature (25 ±3 °C).

7.3 Sieve slurry through a 53-µm sieve to retain the ≥53-µm fraction. Rinse the material retained on the sieve with RODI water until no further fine material comes through and the water that passes the sieve is clear.

7.4 Transfer the ≥53-µm fraction into a preweighed evaporating crucible. Rinse the sieve, including sides, into the crucible with a small amount of RODI water.

7.5 Dry the ≥53-µm fraction in an oven at 110 °C for 15 h or until fully evaporated.

7.6 Cool to room temperature and record the weight of crucible plus ≥53-µm fraction to the nearest 0.1 mg.

7.7 Transfer the entire oven-dried ≥53-µm fraction into a labeled scintillation vial.

**Total Carbon and Nitrogen Analysis**

7.8 Determine total C and N for the ≥53-µm fraction as derived in Section 7.1–7.7, analyzing fine-ground (≈180 µm) sample. Refer to method 4H2a1-3 for the remaining procedural steps for 6A4a1a1-3 as related total C and N by dry combustion analysis of the ≥53-µm fraction and the <2-mm fraction.
8. Calculations

8.1 HPOM-C = \( \frac{C_{\geq 53-\mu m}}{AD/OD} \times \left( \frac{Wt_1}{Wt_2} \right) \times AD/OD \)

8.2 HPOM-N = \( \frac{N_{\geq 53-\mu m}}{AD/OD} \times \left( \frac{Wt_1}{Wt_2} \right) \times AD/OD \)

where:
- HPOM-C = POM-Carbon in oven-dry, <2-mm fraction (%)
- HPOM-N = POM-Nitrogen in oven-dry, <2-mm fraction (%)
- \( C_{\geq 53-\mu m} \) = Carbon in oven-dry, ≥53-µm fraction (%) (method 4H1a1-3)
- \( N_{\geq 53-\mu m} \) = Nitrogen in oven-dry, ≥53-µm fraction (%) (method 4H1a1-3)
- \( Wt_1 \) = Weight of oven-dry, ≥53-µm fraction (g)
- \( Wt_2 \) = Weight of air-dry, <2-mm fraction (g)
- AD/OD = Air dry/oven dry ratio (method 3D1)

8.3 MIN-C = \( C_T - \text{POM-C} \)

8.4 MIN-N = \( N_T - \text{POM-N} \)

where:
- MIN-C = MIN-Carbon in oven-dry, <2-mm fraction (%)
- MIN-N = MIN-Nitrogen in oven-dry, <2-mm fraction (%)
- \( C_T \) = Total Carbon in oven-dry, <2-mm fraction (%) (method 4H2a1-3)
- \( N_T \) = Total Nitrogen in oven-dry, <2-mm fraction (%) (method 4H2a1-3)

9. Report

Report HPOM-C, MIN-C, HPOM-N, and MIN-N as percent of <2 mm soil.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Soil Analyses (6A)
Soil Enzymes (6A5)
p-Nitrophenol (6A5a)
ß-Glucosidase (6A5a1)
Air-Dry, <2 mm (6A5a1a1)
Field-Moist, <2 mm (6A5a1a2)

1. Application

Soil enzymes are important to the biochemical functions of organic matter decomposition and serve as catalysts in reactions necessary for the life processes of organic wastes, organic matter formation, and nutrient cycling (Das and Varma, 2011). These enzymes include amylase, arylsulphatases, ß-glucosidase, cellulose, chitinase, dehydrogenase, phosphatase, protease, and urease released from plants, animals, organic compounds, microorganisms, and soils. ß-glucosidase is a common and predominant enzyme in soils (Eivazi and Tabatabai, 1988; Tabatabai, 1994). Glucosidase is involved in the hydrolysis and biodegradation of various ß-glucosidase present in plant debris decomposing.
in the ecosystem, with its final product being glucose, which is an important C energy source of life to microbes in the soil (Esen, 1993). ß-glucosidase enzyme is sensitive to changes in pH and soil management practices, and as such it has proved to be useful as an indicator of soil quality (Acosta-Martinez and Tabatabai, 2000; Madejon et al., 2001; Das and Varma, 2011). Enzyme assays, such as ß-glucosidase, reflect potential activity, do not represent true in situ activity levels, and should be viewed as an index.

2. Summary of Method

A Modified Universal Buffer is added to a 3-g sample (control and treatment), followed by the addition of 0.05 $M$ p-nitrophenyl-ß-glucosidase (PNG) and incubated for 1 h at 37 ºC. After incubation, 0.5 $M$ CaCl$_2$ and 0.1 $M$ 2-amino-2 (hydroxymethyl)-1-3-propanediol (THAM) (pH 12) are added. To the control sample only, PNG is added and allowed to settle for 5 min. A 10-mL sample is then pipetted and filtered. A calibration curve is prepared plotting absorbance at 410 nm versus amount of p-nitrophenol. If readings exceed high standard, then dilute sample with THAM (pH 10). Data are reported as mg p-nitrophenol per kg oven-dry soil per h.

3. Interferences

The ß-glucosidase assay can be determined on both air-dry and field-moist samples. Air-drying makes handling easier on a routine basis. Additionally, immediate analysis is not required for air-dry samples. However, ß-glucosidase activity declines with air-drying, but the ranking of soil treatments within a soil type or the ranking across different soil types stays the same (Bandick and Dick, 1999). Samples must be read as soon as possible after filtration because p-nitrophenol is light sensitive.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Always work under the fume hood when handling the p-Nitrophenol solutions. Standard laboratory equipment includes fire blankets and extinguishers for use if necessary. Follow the manufacturer’s safety precautions when using the spectrophotometer.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Volumetric flasks, acid washed, 50-mL, 100-mL, 1000-mL
5.3 Plastic bottle, amber, 1000-mL
5.4 Funnel, 60° angle, long stem, 50-mm diameter
5.5 Filter, 0.45-µm
5.6 Pipettes, electronic digital, 2500-µL and 10-mL, with 2500-µL and 10-mL tips
5.7 Syringe filters, 0.45-µm, Whatman
5.8 Centrifuge tubes, 50-mL, polypropylene
5.9 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.10 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.11 Disposable pipettes
5.12 Incubator or water bath, 37 ºC
5.13 Spectrophotometer, UV-Visible, Dual-View, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.14 Vortexer, mini, Analog, VWR Scientific Products
5.15 Syringes, 1-mL

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Modified Universal Buffer (MUB) stock solution: Dissolve 12.10 g THAM, 11.60 g maleic acid, 14.00 g citric acid, and 6.30 g boric acid in 488 mL 1 M NaOH (40.00 g NaOH in 1 L RODI in a volumetric flask) in 1-L volumetric flask and dilute to 1 L with RODI. Store in refrigerator.
6.3 MUB working solution (pH 6.0): Place 200 mL MUB stock solution in a 500 mL beaker with a magnetic stir bar and place the beaker on a magnetic stirrer. Slowly, with stirring, add 0.1 M HCl to the MUB stock solution until the pH reaches 6.0. Transfer acidified solution to 1-L volumetric flask and adjust the volume to 1 L with RODI.
6.4 0.05 M p-nitrophenyl-β-D-glucoside (PNG): Prepare and use same week as analysis. Dissolve 1.308-g PNG in about 80 mL MUB working solution (pH 6.0) in 100-mL volumetric flask. After the PNG is dissolved, bring the solution to a volume of 100 mL with MUB working solution (pH 6.0). Store in the refrigerator (stable for several days at 4°C). Reagent is sufficient for approximately 50 samples in duplicate for “control” and “treatment.”
6.5 0.5 M CaCl₂•2H₂O: Dissolve 73.5 g CaCl₂•2H₂O in about 700 mL RODI in 1-L volumetric flask. After the salt is dissolved, bring to volume with RODI.
6.6 0.5 M NaOH: Dissolve 20 g NaOH in about 700 mL RODI in 1-L volumetric flask. After the NaOH is dissolved, bring to volume with RODI.
6.7 0.1 \( M \) 2-amino-2-(hydroxymethyl)-1,3-propanediol (THAM) (pH ≈10): Dissolve 12.20-g THAM in about 800 mL RODI in 1-L volumetric flask. After the THAM is dissolved, bring to volume with RODI.

6.8 0.1 \( M \) THAM (pH 12): Dissolve 12.20-g THAM in about 800 mL RODI. After the THAM is dissolved, adjust the pH to 12 by slowly adding, with stirring, 0.5 \( M \) NaOH. Bring to volume with RODI.

6.9 p-Nitrophenol Stock Standard Solution: Work in a fume hood. Prepare and use the same month as analysis. Dissolve 1.00 g p-nitrophenol in about 700 mL RODI in 1-L volumetric flask. After p-nitrophenol is dissolved, bring to volume with RODI. Store in amber bottle at 4°C.

6.10 p-Nitrophenol Working Standard Solution: Work in a fume hood. Prepare and use same week as end analysis. Add 1 mL p-nitrophenol Stock Standard Solution to 100 mL-volumetric flask and bring to volume with RODI. Mix well. Store in amber bottle at 4°C.

6.11 Standard p-nitrophenol calibration solutions: 50, 40, 30, 20, 10, 0 \( \mu \)g p-nitrophenol. To six 50-mL centrifuge tubes, mix the following amounts of working standard and RODI:

6.11.1 50.0 \( \mu \)g p-nitrophenol = 5.0 mL p-nitrophenol Working Standard Solution

6.11.2 40.0 \( \mu \)g p-nitrophenol = 4.0 mL p-nitrophenol Working Standard Solution and 1.0 mL RODI

6.11.3 30.0 \( \mu \)g p-nitrophenol = 3.0 mL p-nitrophenol Working Standard Solution and 2.0 mL RODI

6.11.4 20.0 \( \mu \)g p-nitrophenol = 2.0 mL p-nitrophenol Working Standard Solution and 3.0 mL RODI

6.11.5 10.0 \( \mu \)g p-nitrophenol = 1.0 mL p-nitrophenol Working Standard Solution and 4.0 mL RODI

6.11.6 0.0 \( \mu \)g p-nitrophenol = 0 mL p-nitrophenol Working Standard Solution and 5.0 mL RODI

Add 1.0 mL 0.5 \( M \) CaCl\(_2\) and 4.0 mL of THAM (pH 12) to tube. Cap tube and shake. Standards are not incubated.

7. Procedure

7.1 Weigh 1 g (±0.03 g) of <2 mm air-dry soil into Falcon tubes. Label odd-numbered tubes “control” and even-numbered tubes “treatment.”

7.2 Work in a fume hood. Add 4 mL MUB (pH 6.0) working solution to all tubes (controls and treatments).

7.3 Add 1 mL of PNG solution to the “treatment” tubes only. Cap and vortex all tubes. Start the 60 min timer immediately after adding the PNG and before mixing by vortex.
7.4 Incubate all samples at 37 °C for 1 h.

7.5 Add 1 mL of 0.5 M CaCl₂ and 4 mL 0.1 THAM (pH 12) to all tubes and swirl. This reagent may be combined and added to each tube.

7.6 Add 1 mL PNG solution to “control” tubes and vortex. Remove caps and let samples settle 5 minutes before filtering.

7.7 Draw up sample into a 10 mL syringe and attach a 0.45 µm filter. Filter sample directly into cuvette. Samples must be read as soon as possible after filtration because p-nitrophenol is light sensitive. Cap the samples waiting to be read.

7.8 Prepare calibration curve plotting absorbance at 410 nm versus amount of p-nitrophenol.

7.9 Record amounts of p-nitrophenol. Read the samples in control/treatment pairs.

7.10 If readings exceed high standard, dilute with THAM (pH 10). Cap and swirl. Transfer to cuvettes and record the amount of p-nitrophenol.

8. Calculations

\[ \text{mg p-nitrophenol/ kg oven-dry soil/h} = B \times \left[ (D_T \times A_T / W_{T_t}) - (D_C \times A_C / W_{C_t}) \right] / T \]

where:
- \( A_T \) = Amount of p-nitrophenol in treatment sample (µg)
- \( A_C \) = Amount of p-nitrophenol in control sample (µg)
- \( W_{T_t} \) = Weight of treatment sample (g)
- \( W_{C_t} \) = Weight of control sample (g)
- \( D_C \) = Dilution factor for control sample
- \( D_T \) = Dilution factor for treatment sample
- \( T \) = Incubation time (h)
- \( B \) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

Note: µg p-nitrophenol/g oven-dry soil/hour = mg p-nitrophenol/kg oven-dry soil/hour

9. Report

Report Beta-glucosidase activity as mg p-nitrophenol/kg oven-dry soil/h.

10. Precision and Accuracy

Precision and accuracy data are available from the KSSL upon request.

11. References


Ratios and Estimates Related to Biological Analyses (6D)
Estimates or calculated values associated with soil biological and plant analyses are described within the respective methods. Additional information on the reporting of these calculated values can be obtained from the KSSL upon request.

MINERALOGY (7)

Instrumental Analyses (7A)
The physical and chemical properties of a soil are controlled to a very large degree by the soil minerals, especially by those minerals constituting the clay fraction (McBride, 1989; Whittig and Allardice, 1986). Positive identification of mineral species and quantitative estimation of their proportions in soils usually require the application of several complementary qualitative and quantitative analyses (Whittig and Allardice, 1986). Some of the semiquantitative and quantitative procedures that have been performed by the KSSL include x-ray diffraction (method 7A1a1) and thermal analysis (7A2a, 7A3a, and 7A4a). Other indirect, ancillary procedures to infer mineral composition include linear extensibility, elemental analysis, and CEC/clay ratios.
Analysis by x-ray diffraction facilitates identification of crystalline mineral components of soil and semiquantitative estimates of relative amounts. It is commonly applied to the clay fraction in soils and to layer-silicate (phyllosilicates) minerals in particular. Identification is by d-spacings (spatial distance between
repeating planes of atoms) characteristic of a mineral, according to Bragg’s Law. Because layer-silicate structures are very similar from one mineral to another, except in the direction perpendicular to the layers, several treatments (cation saturation and heating) must be used to correctly identify the several minerals. In x-ray analysis of soils or clay samples, there are difficulties in evaluation of and compensation for the variations in chemical composition, crystal perfection, amorphous substances, and particle size (Whittig and Allardice, 1986; Hughes et al., 1994). A more reliable and accurate estimation of mineral percentages is provided when x-ray diffraction analysis is used in conjunction with other methods, e.g., differential-thermal, surface-area, elemental analysis, and other species-specific chemical methods (Alexiades and Jackson, 1966; Karathanasis and Hajek, 1982).

Upon heating, many soil constituents undergo thermal reactions that serve as diagnostic properties for qualitative and quantitative identification of these substances (Tan et al., 1986; Karathanasis and Harris, 1994). Thermogravimetric analysis (TGA) (method 7A2a) is a technique for determining weight loss of a sample when it is heated at a constant rate. TGA is an outgrowth of dehydration curves that were used in early studies of various phyllosilicate clay minerals (Jackson, 1956). The TGA sample weight, however, is monitored continuously rather than measured at discrete intervals after periods of heating at a constant temperature (Wendlandt, 1986). TGA measures only reactions that involve weight loss of the sample.

Differential scanning calorimetry (DSC) (7A4a, obsolete method) is a calorimetric technique that theoretically measures the amount of energy required to establish zero temperature difference between sample and reference material as the two are heated side by side at a controlled rate (Tan et al., 1986; Karathanasis and Harris, 1994). Most common DCS instruments have sample and reference pans heated in a single furnace, and the difference in temperature is measured during various endothermic and exothermic reactions. This difference in temperature is then converted to a value equivalent to an enthalpy change (expressed in calories) using instrumental calibrations (Karathanasis and Harris, 1994).

TGA and DSC are complementary methods available to the analyst. Many of the same clay mineral reactions, e.g., dehydroxylation, loss of surface adsorbed water, decomposition of carbonates, and oxidation, that are studied by DSC can also be studied by TGA. However, some transformation reactions, e.g., melting or structural reorganization (quartz alpha-beta transition), cannot be measured by TGA because no weight loss is involved (Karathanasis and Harris, 1994). The DSC procedures provide information about energy relationships in the structures and reactions of the solid phase, whereas TGA provides quantitative information about quantities of substances gained or lost by the solid phase during certain thermally driven reactions.
References

Jackson, M.L. 1956. Soil chemical analysis. Advan. course. SSSA, Madison, WI.

Instrumental Analyses (7A)

X-Ray Diffractometer (7A1)

Filter Peel on Glass (7A1b)

Mg Room Temperature, Mg Glycerol Solvated, K 300 °C, K 500 °C (7A1b1)

1. Application

Clay fractions of soils are commonly composed of mixtures of one or more phyllosilicate minerals together with primary minerals inherited directly from the parent material (Olson et al., 1999). Positive identification of mineral species
and quantitative estimation of their proportions in these polycomponent systems usually require the application of several complementary qualitative and quantitative analyses (Whittig and Allardice, 1986; Amonette and Zelazny, 1994; Wilson, 1994; Moore and Reynolds, 1997). One of the most useful methods to identify and to make semiquantitative estimates of the crystalline mineral components of soil is x-ray diffraction analysis (Hughes et al., 1994; Kahle et al., 2002). Quantification of a mineral by x-ray diffraction requires attention to many details, including sample (slide) size relative to the incident x-ray beam, thickness and particle size uniformity of sample, and beam-sample orientation (Moore and Reynolds, 1997). More complex quantification procedures include using standard additions, full pattern fitting, and determining mineral intensity factors (Kahle et. al., 2002). At best, quantification can approach a precision of ±5% and an accuracy of ±10 to 20% (Moore and Reynolds, 1997).

The operational strategy at the KSSL and the former Lincoln SSL has been to base mineral quantification on first order peak intensities. Semiquantitative interpretations have been held consistent over time (1964 to the present) by adjusting instrumental parameters (e.g., scan speed) to maintain a constant peak intensity for an in-house reference clay standard and subsequently soil samples. The intent is to keep interpretations consistent from sample to sample.

2. Summary of Method

Soils are dispersed and separated into fractions of interest. Sands and silts are mounted for analysis on glass slides as slurries, on a smear of Vaseline, or on double sticky tape. Clay suspensions are placed on glass slides to dry and to preferentially orient clay minerals. Most samples of soil clays contain fewer than 7 minerals that require identification. The soil clay minerals of greatest interest are phyllosilicates, e.g., kaolinite, mica (illite), smectite, vermiculite, hydroxy-interlayered vermiculite, smectite, hydroxy-interlayered smectite, and chlorite.

Diffraction maxima (peaks) develop from the interaction of x-rays with planes of elements that repeat at a constant distance (d-spacing) through the crystal structure. Generally, no two minerals have exactly the same d-spacings in three dimensions and the angles at which diffraction occurs are distinctive for a particular mineral (Whittig and Allardice, 1986; Moore and Reynolds, 1997). Phyllosilicates (or layer-silicate minerals) have very similar structures, except in the direction perpendicular to the layers (c-dimension). Several treatments are needed to sort out which minerals are present. Glycerol is added to expand smectites. Ionic saturation and/or heat treatments are used to collapse some 2:1 layer silicates and to dehydroxylate kaolinite, gibbsite, and goethite, eliminating characteristic peaks.

The crystal “d” spacings of minerals, i.e., the interval between repeating planes of atoms, can be calculated by Bragg’s Law as follows:
\[ n\lambda = 2d \sin \theta \]

where:
\( n \) = integer that denotes order of diffraction
\( \lambda \) = x-radiation wavelength (Angstroms, Å)
\( d \) = crystal “d” spacing (Å)
\( \theta \) = angle of incidence

When \( n = 1 \), diffraction is of the first order. The wavelength of radiation from an x-ray tube is constant and characteristic for the target metal in the tube. Copper radiation (CuK\( \alpha \)) with a wavelength of 1.54 Å (0.154 nm) is used at the KSSL. Because of the similar structure of layer silicates commonly present in soil clays, several treatments that characteristically affect the “d” spacings are necessary to identify the clay components. At the KSSL, four treatments are used. They are: Mg\( ^{2+} \) (room temperature); Mg\( ^{2+} \)-glycerol (room temperature); K\( ^+ \) (300 °C); and K\( ^+ \) (500 °C).

Standard tables to convert \( \theta \) or 2\( \theta \) angles to crystal d-spacings are published in the U.S. Geological Survey Circular 29 (Switzer et al., 1948) and in other publications (Brown, 1980). Through the years, hardware has been updated and the recording of data has evolved from a strip chart recorder through several kinds of electronic software. X-ray by this method (7A1a1) is semiquantitative.

3. Interferences

Interstratification of phyllosilicate minerals causes problems in identification. These interstratified mixtures, differences in crystal size, purity, chemical composition, atomic unit cell positions, and background or matrix interferences affect quantification (Moore and Reynolds, 1997; Kahle et al., 2002). No pretreatments other than ionic saturation and dispersion with sodium hexametaphosphate are used for separation and isolation of the clay fraction in the routine procedure. Impurities, such as organic matter, carbonates, and iron oxides, may act as matrix interferences, causing peak attenuation during x-ray analysis, or may interfere with clay dispersion and separation. Pretreatments to remove these impurities serve to concentrate the crystalline clay fraction and may increase accuracy but also potentially result in degradation of certain mineral species (e.g., smectites) and loss of precision (Hughes et al., 1994).

The separation (centrifuge) procedure used to isolate the clay fraction from the other size fractions of the soil skews the <2-μm clay suspension toward the fine clay, but it minimizes the inclusion of fine silt in the fraction. Sedimentation of the clay slurry on a glass slide tends to cause differential settling by particle size (i.e., increasing the relative intensity of finer clay minerals).

Dried clay may peel from the XRD slide. One remedy is to rewet the peeled clay on the slide with 1 drop of glue-water mixture (1:7). An optimum amount of glycerol on the slides is required to solvate the clay, i.e., to expand smectites to
18 Å. X-ray analysis should be performed 1 to 2 days after glycerol addition. If excess glycerol is applied to the slide and free glycerol remains on the surface, XRD peaks are attenuated. Using a chamber (such as a desiccator with no desiccant) to dry the slide, especially when the clay is thin, is suggested to achieve optimum glycerol solvation.

4. Safety

Operate the centrifuge with caution. Keep the centrifuge lid closed when in operation. Ensure that all rotors and tubes are seated firmly in their proper location. Use tongs and appropriate thermal protection when operating the muffle furnace. The diffraction unit presents an electrical and radiation hazard. Analysts must receive radiation safety training before operating the equipment. Area radiation monitors must be used in the vicinity of the x-ray diffractometer.

5. Equipment

5.1 Dispenser, for sodium carbonate solution
5.2 Centrifuge, International No. 2, with No. 240 head and carriers for centrifuge tubes, International Equip. Co., Boston, MA
5.3 Centrifuge tubes, plastic, 100-mL, on which 10-cm solution depth is marked
5.4 Rubber stoppers, No. 6, for centrifuge tubes
5.5 Burrell Wrist Action™ Shaker Model 75
5.6 Sieve, 80-mesh, copper
5.7 Spray bottle, plastic, 30-mL (1 oz), for a 1:7 glycerol:water mixture
5.8 Muffle furnace, NeyTech Vulcan™ 3-1750A
5.9 X-ray diffractometer, Bruker D-4 Endeavor, with X-Y autosampler that accommodates 66 samples or standards, Bruker AXS Inc., Madison, WI
5.10 EVA 16 software, release 2010, Bruker AXS Inc., Madison, WI, and printer
5.11 XRD Commander 2.6 software, release 2009 Bruker AXS Inc., Madison, WI
5.12 XRD slides, glass, 25.4 x 25.4 mm
5.13 66 slide holders
5.14 Reference slides: quartz and clay from reference soil
5.15 Check standard soil
5.16 Vortex mixer
5.17 Pall membrane filters, Nylaflo membrane, diameter 47 mm, pore size 0.2 µm.
5.18 Fisherbrand reusable heavy wall filter flask
5.19 Four Millipore filter holder vacuum manifolds, PVC, three-place
5.20 Twelve 47-mm filter holders for flasks or manifolds, glass
5.21 Slide roller (fig. 7A1-1)
5.22 Vacuum chuck (fig. 7A1-1)

Figure 7A1-1.—Vacuum chuck (top) and slide roller (bottom).

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Sodium carbonate (Na$_2$CO$_3$), 0.25 N. Dissolve 26.50 g Na$_2$CO$_3$ in 1 L RO water or 132.50 g in 5 L RO water.
6.3 Potassium chloride (KCl), 1.0 N. Dissolve 74.60 g KCl in 1 L RO water or 671.40 g KCl in 9 L RO water.
6.4 Magnesium chloride (MgCl$_2$), 1.0 N. Dissolve 47.61 g MgCl$_2$ in 1 L RO water or 428.49 g MgCl$_2$ in 9 L RO water.
6.5 Glycerol:water mixture (1:7). Add 4 mL of glycerol to 28 mL RO water plus 2 drops of toluene.

7. Procedure

Preparation of Clay Suspension

7.1 In the laboratory information system (LIMS), create a batch file. The batch will consist of 11 samples plus the check standard soil. Data in the file is transferred to a job program on the x-ray computer software for data
analysis. These data include project and sample identification. Include both the quartz and the soil standard with each run.

7.2 Each sample has two filter treatments (Mg$^{2+}$ and K$^+$) so that two tubes are prepared for each sample.

7.3 Label each 100 mL plastic centrifuge tube with a sample number from the run sheet and the treatment. There are 24 tubes per batch.

7.4 Weigh out enough soil to obtain 150 mg of clay (use particle size distribution data to calculate the weight) into the 100 mL tube.

7.5 Fill tube to 9.5-cm height with sodium carbonate solution and add stopper.

7.6 Place the tubes on Wrist Action™ shaker and shake for 2 hours.

7.7 Remove stopper from tube and rinse stopper and sides of tube with enough solution to bring the volume to the 10-cm mark. Vortex the clay suspension.

7.8 Balance the pairs of tubes and place in centrifuge. Centrifuge at 750 rpm for 3.0 min.

7.9 If the clay is dispersed, carefully decant 30 mL of suspension into a labeled, 200 mL beaker, making sure to stop pouring before the silt reaches the mouth of the tube. You can usually pour off 5–7 cm of solution.

7.10 Fill the tubes to the 10 cm mark solution again. Vortex the tubes.

7.11 Repeat Sections 7.8 to 7.10 three times.

7.12 Clay suspension in the beaker is used for x-ray diffraction and TGA analysis.

**Thin Film on Glass, Filter Peel Preparation**

7.13 Squirt a few drops of water on the base of the vacuum filter holder and then place the filter, being careful to center it. Clamp the funnel down over the base, being careful not to displace the filter paper.

7.14 Turn vacuum valve below the base of the filter holder.

7.15 Label glass slides with sample number and treatment.

7.16 Pour half of the clay suspension on the filter.

7.17 Wait for the water to pull through the filter. The required amount of time varies, depending on clay type. When the very center of the filter paper begins to dry, swirl the remaining suspension to disperse the clay particles once again, then pour the other half of the clay suspension into the funnel. Wash any settled clay from the bottom of the beaker into the funnel using a squirt bottle filled with RO water.
7.18 When the filter begins drying again, use a squirt bottle to cover the filter with desired treatment solution (MgCl₂ or KCl). Add approximately 5 mL.

7.19 When the filter begins drying once again, rinse the sides of the filter with approximately 5 mL RO water.

7.20 Wait until the filter begins drying. Within 15 seconds, peel the filter from the base and turn the vacuum valve to the “off” position.

7.21 Place the glass slide (labeled-side down) on the vacuum chuck (fig. 7A1-1). Apply vacuum.

7.22 Place the filter (clay-side down) on the slide and then (using the roller, fig. 7A1-1) run across all four sides of the slide. Then roll across the top of the filter gently. Peel up the filter paper, leaving the clay film on the slide.

7.23 If extra clay is needed (such as for TGA analysis), roll the remaining clay on another slide. This clay is the residual area on the circular filter that extended past the square slide in step 7.23.

7.24 Allow slides to dry in air for at least 4 h.

7.25 For each sample number on the bench sheet, there should be two slides, magnesium-treated and potassium-treated. Each slide is analyzed twice so that there are four treatments in total. Treatments are as follows:

Mg²⁺-room temperature
Mg²⁺-glycerol
K⁺-300 °C (heated 2 h)
K⁺-500 °C (heated 2 h)

7.26 Transfer the K⁺ treated slides to a 300 °C muffle furnace for 2 hours. Analyze the Mg²⁺-room temperature and K⁺-300 °C slides at the same time.

7.27 After analyzing, transfer K⁺-300 °C slides to 500 °C muffle furnace for 2 h. Transfer Mg²⁺-room temperature to a paper lined tray and spray with glycerol solution, making sure to coat all slides evenly. Allow to stand for at least 8 h. Analyze the Mg²⁺-glycerol and the K⁺-500 °C slides at the same time.

X-Ray Diffraction Operation

7.28 Complete x-ray analysis of the glycerol treatment within 1 day after the slide dries. If not possible, spray additional glycerol prior to run.

7.29 Place the slides in the autosampler grid of the XRD. Slides should be placed in the holders inside the grid, making sure each slide fits snugly.

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Parameter Setting

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuKα radiation, λ</td>
<td>1.54 Å (0.154 nm)</td>
</tr>
<tr>
<td>Scan range</td>
<td>2° to 35° 2θ</td>
</tr>
<tr>
<td>Generator settings</td>
<td>40 kv, 30 ma</td>
</tr>
<tr>
<td>Divergence slit</td>
<td>1°</td>
</tr>
<tr>
<td>Receiving slit</td>
<td>0.2 mm</td>
</tr>
</tbody>
</table>

Step-size and scan-speed are varied depending on intensity of x-rays generated. Adjust settings to maintain the same peak intensities on the standard reference clay and quartz standard over the long term regardless of tube intensities.

7.30 Activate the sample information file (SIF) for analysis in LIMS. The SIF can be imported into XRD Commander. There are two SIF files for each batch. The first SIF file is for the Mg²⁺ room temperature and the K⁺ 300 °C. The second SIF file is for the Mg²⁺ glycerol and the K⁺ 500 °C. The SIF file stores raw data on the hard disk under the subdirectory designated by year, project type, project name.

7.31 Prepare and print a 4-color graphics chart. Colors are as follows: blue (Mg²⁺); green (Mg²⁺-glycerol); pink (K⁺ 300 °C); and red (K⁺ 500 °C). File hard copies of detected peaks and graphics chart in pasteboard binders by LIMS batch number.

7.32 Compare quartz and soil standard patterns electronically with previous runs to ensure peak intensity and positions have remained constant.

Interpretation of X-Ray Diffraction Data

7.33 The angle in degrees two theta (2θ) measured in x-ray diffraction analyses is converted to angstroms (Å) using tables compiled according to Bragg’s Law. Refer to summary of method. Angstroms convert to nanometers (nm) by a factor of 0.1, e.g., 14 Å = 1.4 nm.

7.34 Use the following x-ray diffraction criteria to identify some common crystalline minerals. The reported “d” values are for 00I basal spacings. The Miller index (hkl) specifies a plane or crystal face that has some orientation to the three crystallographic axes of a, b, and c. The Miller index (00I) indicates a crystal face that is perpendicular to the a and b axes (Schultz, 1989). The following x-ray diffraction criteria also have some questions (Q) that may aid the analyst in interpreting the diffraction patterns. These questions are a suggested procedural approach to help the analyst identify the relative locations of a few peaks and to confirm
key criteria. For a more complete list of d-spacings for confirmation or identification of a mineral, consult the Mineral Powder Diffraction File–Data Book (JCPDS, 1980).

**X-Ray Diffraction Criteria**

7.34.1 Kaolinite and Halloysite
   a. Crystal structure missing at 500 °C
   b. 7 Å (7.2 to 7.5 Å) with all other treatments
   Q. Is there a 7 Å peak? Is it destroyed at 500 °C? If so, the mineral is kaolinite or halloysite.
   Q. Is the peak sharp and at ≈7.1 Å (but absent at 500 °C)? If so, the mineral is kaolinite.
   Q. Is the peak broad and at 7.2 to 7.5 Å (but absent at 500 °C)? If so, the mineral is halloysite.

7.34.2 Mica (Illite)
   a. 10 Å with all treatments
   b. 10 Å with Mg²⁺-saturation
   Q. Is there a 10 Å peak with Mg²⁺-saturation? If so, the mineral is mica (illite).

7.34.3 Chlorite
   a. Crystal structure of Fe-chlorites destroyed at 650 to 700 °C
   b. 14 Å with all other treatments
   c. 14 Å at 500 °C
   d. Generally, also has a strong 7 Å peak
   Q. Is there a 14 Å peak when heated to 500 °C? If so, the mineral is chlorite.

7.34.4 Vermiculite
   a. 14 Å with Mg²⁺-saturation
   b. 14 Å with Mg²⁺-glycerol solvation
   c. Nearly 10 Å with K⁺ saturation
   d. 10 Å when K⁺-saturated and heated to 300 °C
   Q. Is there an enhanced 10 Å peak with K⁺-saturation in comparison to Mg²⁺-saturation that cannot be attributed to smectite? If so, the mineral is vermiculite.

7.34.5 Smectite
   a. 14 Å with Mg²⁺-saturation
   b. 12 to 12.5 Å with K⁺- or Na⁺-saturation
   c. 17 to 18 Å with Mg²⁺-glycerol solvation
   d. 10 Å with K⁺-saturation and heating to 300 °C
   Q. Is there a 17 to 18 Å peak upon solvation? If so, the mineral is smectite.
7.34.6  Gibbsite  
a. Peak at 4.83 to 4.85 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol but destroyed when heated to 300 °C

7.34.7  Goethite  
a. Peak at 4.16 to 4.18 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol but destroyed when heated to 300 °C

7.34.8  Hydroxy-interlayered Vermiculite or Smectite  
a. Failure to completely collapse to 10 Å of smectite or vermiculite when K$^+$-saturated and heated to 300 °C

7.34.9  Quartz  
a. Peaks at 4.27 Å and 3.34 Å with all treatments (only 3.34 if small amounts)

7.34.10  Lepidocrocite  
a. Peak at 6.2 to 6.4 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol but destroyed when heated to 300 °C

7.34.11  Potassium Feldspar  
a. Peak at 3.24 Å with all treatments

7.34.12  Plagioclase Feldspar  
a. Twin peaks between 3.16 and 3.21 with all treatments

7.34.13  Calcite  
a. Peak at 3.035 Å with all treatments

7.34.14  Dolomite  
a. Peak at 2.88 to 2.89 Å with all treatments

7.34.15  Gypsum  
a. Peak at 7.56 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol but destroyed when heated to 300 °C

7.34.16  Mixed Layer Vermiculite-Mica  
a. Randomly interstratified: Peak between 10 and 14 Å with Mg$^{2+}$ which does not expand with Mg$^{2+}$-glycerol; peak collapses to 10 Å with K$^+$-saturation and heating to 300 °C.  
b. Regularly interstratified: A 24 Å peak (and higher orders); no change with Mg$^{2+}$-glycerol treatment; K$^+$ saturation and heating collapses vermiculite and a produces a 10 Å peak.

7.34.17  Mixed Layer Smectite-Mica  
a. Randomly interstratified: Peak between 10 and 14 Å with Mg$^{2+}$ which expands to 14–16 Å with Mg$^{2+}$-glycerol; Peak collapses to 10 Å with K$^+$-saturation and heating to 300 °C.  
b. Regularly interstratified: A small 24 Å peak and large peak at 12 Å with Mg$^{2+}$-saturation; expands to 28 Å with Mg$^{2+}$-glycerol treatment; K$^+$-saturation and heating collapses smectite, then produces a 10 Å peak.
7.34.18 Mixed Layer Chlorite-Vermiculite
   a. Randomly Interstratified: Peak at 14 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol; Peak collapses incompletely to between 10 and 14 Å with K$^+$-saturation and heating.
   b. Regularly interstratified: A 28 Å peak (and higher orders) with Mg-saturation; no expansion with Mg$^{2+}$-glycerol treatment; K$^+$-saturation and heating to 500 °C collapses vermiculite and a produces a 24 Å peak.

7.34.19 Mixed Layer Chlorite-Smectite
   a. Randomly interstratified: Peak at 14 Å with Mg$^{2+}$-saturation; expands to higher spacings (≈16 Å) with Mg$^{2+}$-glycerol treatment; Peak collapses incompletely to between 10 and 14 Å with K$^+$-saturation and heating.

7.35 Use the x-ray diffraction criteria, i.e., diagnostic basal 00$l$ spacings (Å), in table 1 for identification and ready reference of some common crystalline minerals as affected by differentiating sample treatments.

7.36 Preferential orientation of clay mineral samples enhances diffraction from the basal (00$l$) spacing and tends to minimize the number and intensity of peaks from diffraction by other $hkl$ planes. With preferential orientation, second, third, and fourth order peaks may be recorded in addition to the basal first order peaks. Groups of associated peaks that differ by order of diffraction are as follows:

7.36.1 Smectite (Mg$^{2+}$-glycerol)
   a. 17 to 18 Å
   b. 8.5 to 9 Å (weak)

7.36.2 Chlorite, vermiculite, and smectite
   a. 14, 7, 4.7, and 3.5 Å
   b. 7, 4.7, and 3.5 Å weak for smectite
   (Note: High Fe substitution in the chlorite structure results in a decrease in the peak intensity of odd numbered orders (e.g., 14 and 4.7 Å) and increase in peak intensity of even number orders (7 and 3.5 Å).

7.36.3 Mica
   a. 10, 5 (weak in biotites and moderate in muscovites), and 3.3 Å

7.36.4 Kaolinite
   a. 7 and 3.5 Å

7.37 The differentiation of kaolinite and halloysite in a sample can be aided by the use of formamide (Churchman et al., 1984). The intercalation and expansion of halloysite to a d-spacing of ≈10.4 Å is relatively rapid (20 to 30 min), whereas kaolinite expansion requires ≈4 h upon treatment. The procedure is as follows:
7.37.1 Lightly spray formamide as an aerosol on the dried Mg\(^{2+}\)-saturated slide.

7.37.2 Wait 15 min but not more than 1 h and x-ray approximately 7.6 to 13.5 ° 2θ (d = 11.6 to 6.55 Å).

7.37.3 Halloysite will expand to ≈10.4 Å, whereas kaolinite will remain unchanged.

7.37.4 Heating the sample to 110 °C for 15 min will collapse the halloysite to ≈7 Å.

7.37.5 The total amount of kaolinite and halloysite can be determined by thermal analysis. The intensity ratio of the 10.4 to 7.2 Å peaks of the formamide-treated sample can be used to determine the relative percentage of halloysite and kaolinite.

8. Calculations

X-ray diffraction produces peaks on a chart that correspond to 2θ angle on a goniometer as well as detected x-ray intensity from the detector. Standard tables to convert θ or 2θ to crystal “d” spacings are published in the U.S. Geological Survey Circular 29 (Switzer et al., 1948) and in other publications (Brown, 1980). The crystal “d” spacings of minerals, i.e., the interval between repeating planes of atoms, can be calculated by Bragg’s Law. Refer to summary of method.

9. Report

From the “Detected Peaks File” and graphics chart, identify the minerals present according to the registered “d” spacings. As a first approximation, use the following peak intensities, i.e., peak heights above background in counts s\(^{-1}\), to assign each layer silicate mineral to one of the 5 semiquantitative classes.

<table>
<thead>
<tr>
<th>Class</th>
<th>Peak Height above Background (counts sec(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 (Very large)</td>
<td>&gt;1800</td>
</tr>
<tr>
<td>4 (Large)</td>
<td>1120 to 1800</td>
</tr>
<tr>
<td>3 (Medium)</td>
<td>360 to 1120</td>
</tr>
<tr>
<td>2 (Small)</td>
<td>110 to 360</td>
</tr>
<tr>
<td>1 (Very small)</td>
<td>&lt;110</td>
</tr>
</tbody>
</table>

Adjust class placement to reflect area under the curve if peak is broad relative to peak height or if thermal, elemental, clay activity data, or other evidence warrant class adjustment. If there are no peaks or no evidence of crystalline components,
place the sample in NX class (non-crystalline). If there are only 1 to 3 very small (class 1) peaks, also indicate NX to infer a major non-crystalline component. Indicate the exact counts of each identified peak in the LIMS database.

10. Precision and Accuracy

X-ray by method 7A1b1 is semiquantitative. Precision and accuracy data are available from the KSSL upon request.

11. References


Table 1.—X-Ray Diffraction Parameters of Common Soil Minerals

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Treatment</th>
<th>Na⁺</th>
<th>Mg²⁺</th>
<th>Mg²⁺ Gly</th>
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<th>K⁺ 500 °C</th>
<th>K⁺ 700 °C</th>
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<td>7B²/</td>
<td>7B</td>
<td>7B</td>
<td>7B</td>
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<td>14*</td>
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<td>14*</td>
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</table>

¹/ LD = Lattice destroyed.
²/ B = Broad peak is common.
³/ * = Sometimes <14 Å.
⁴/ T = Temperature of decomposition varies with chemical composition, particle-size, and heating conditions.

Instrumental Analyses (7A)
Thermogravimetric Analysis (7A2)
Thermal Analyzer (7A2a)

1. Application
Thermal analysis defines a group of analyses that determine some physical parameter, e.g., energy, weight, or evolved substances, as a dynamic function of temperature (Tan et al., 1986; Karathanasis and Harris, 1994). Thermogravimetric
analysis (TGA) is a technique for determining weight loss of a sample as it is being heated at a controlled rate. The weight changes are recorded as a function of temperature, i.e., a thermogravimetric curve, and provide quantitative information about substances under investigation, e.g., gibbsite (Al(OH)₃), kaolinite (Al₂Si₂O₅(OH)₄), goethite (FeOOH), and 2:1 expandable minerals (smectite and vermiculite).

2. Summary of Method

A 5- to 10-mg sample of soil clay or fine earth (finely ground) is placed in a platinum sample pan, and the pan is placed in the TGA balance. The instrument records the initial sample weight. The sample is then heated from a temperature of 30 °C to 900 °C at a rate of 20 °C min⁻¹ in a flowing N₂ atmosphere. The computer collects weight changes as a function of temperature and records a thermogravimetric curve. Gibbsite and kaolinite are quantified by calculating the weight loss between approximately 250 to 350 °C and 450 to 550 °C, respectively, and then relating these data to the theoretical weight loss of pure gibbsite or kaolinite. The weight loss is due to dehydroxylation, i.e., loss of crystal lattice hydroxyl ions. Although not presently performed by the KSSL, quantification of the 2:1 expandable minerals (smectite + vermiculite) is related to weight loss at <250 °C, i.e., loss of adsorbed water (Karathanasis and Hajek, 1982a, 1982b; Tan et al., 1986). At this low temperature, adsorbed water is proportional to the specific surface area of the sample (Jackson, 1956; Mackenzie, 1970; Tan and Hajek, 1977; Karathanasis and Hajek, 1982b). In the absence of gibbsite, goethite (a Fe oxyhydroxide) can be quantified based on the characteristic weight loss of 10.1 to 11.2% between 300 and 400 °C (Karathanasis and Harris, 1994). Recent work in the KSSL has found good agreement between gypsum quantification using dissolution procedures and thermal analysis. Gypsum has a weight loss of 20.9% between 100 and 350 °C (Karathanasis and Harris, 1994). The TGA method (7A2a) is especially useful for soils with a large percentage (>20%) of gypsum. Burt et al. (2001) had good agreement between total Mg analysis and TGA quantification (12.9% weight loss between 600 and 650 °C) of serpentine minerals in ultramafic-derived soils in Oregon.

3. Interferences

Organic matter is objectionable because it has a weight loss by dehydrogenation and by oxidation to CO₂ between 300 and 900 °C (Tan, et al., 1986). Analysis in an inert N₂ atmosphere helps to alleviate this problem, but samples with significant organic matter should be pretreated with H₂O₂ (method 3A1a1). Mineral salts that contain water of crystallization may also interfere. Samples should be washed free of any soluble salts. In some cases, weight loss from gibbsite and goethite overlap and prevent quantitative interpretation. These samples can be deferrated (method 4F1a1) to eliminate goethite.
A representative soil sample is important because sample size is small (<10 mg). Avoid large aggregates in sample, the presence of which may cause thermal interferences, i.e., differential kinetics of gas diffusion through the sample and physical movement of sample in a reaction.

In general, the same reactions that interfere with DSC/DTA also interfere with TGA determinations of kaolinite, gibbsite, and 2:1 expandable minerals. However, TGA is more sensitive to small water losses at slow rates, whereas DSC/DTA is more sensitive to large water losses at rapid rates (Tan, et al., 1986). This difference in sensitivity may help to explain why quantification of kaolinite and gibbsite in TGA and DSC/DTA are often not equivalent, i.e., TGA estimates tend to be greater than the corresponding DSC/DTA estimates. In TGA, there is a greater probability of measuring water losses in specific temperature regimes that are not specifically associated with dehydroxylation reactions of interest. This problem is particularly apparent with illitic samples, which characteristically contain more "structural" water than ideal structural formulae would indicate (Rouston, et al., 1972; Weaver and Pollard, 1973).

Even though it is well established that various minerals lose the major portion of their crystal lattice water in different temperature ranges (Tan et al., 1986), the weight loss regions (WLR) of minerals have overlaps that interfere in the identification and measurement of the minerals of interest. The goethite WLR (250 to 400 °C) overlaps the gibbsite WLR (250 to 350 °C) (Mackenzie and Berggen, 1970). The illite WLR (550 to 600 °C) overlaps the high end of the kaolinite WLR (450 to 550 °C) (Mackenzie and Caillere, 1975). The WLR of hydroxy-Al interlayers in hydroxy-Al interlayered vermiculite (HIV) (400 to 450 °C) overlaps the low end of the kaolinite WLR (450 to 550 °C), especially in the poorly crystalline kaolinites (Mackenzie and Caillere, 1975). Similarly, the dehydroxylation of nontronites, Fe-rich dioctahedral smectites (450 to 500 °C), may interfere with kaolinite identification and measurement (Mackenzie and Caillere, 1975).

4. Safety

Secure high pressure N₂ tanks and handle with care. When changing the tanks, protect valves with covers. Do not program the analyzer for >950 °C because it may present a safety hazard during sample analysis and cleaning cycles. Always use high quality purge gases with the TGA. Minimum purity of 99.9% is recommended. Handle hot furnace with care.

5. Equipment

5.1 Thermal analyzer, TGA Q5000 IR, TA Instruments, New Castle, DE
5.2 Thermal analyzer operating system software, TA Instrument Explorer 2000, TA Instruments, New Castle, DE
5.3 Computer Data analysis software, TA Instruments Universal Analysis 2000, TA Instruments, New Castle, DE
5.5 N₂ gas, 99.99% purity
5.7 Two-stage gas regulator 20 psi maximum outlet pressure
5.8 Forceps, flat-tipped
5.9 Weighing spatula
5.10 Desiccator, glass
5.11 Mortar and pestle
5.12 Sieve, 80-mesh
5.13 Razor blade
5.14 Gibbsite, standard, Surinam Gibbsite, SSL, 67L022

6. Reagents
6.1 Magnesium nitrate saturated solution Mg(NO₃)₂•6H₂O

7. Procedure

Derive <2 µm Clay Fractions

7.1 Create LIMS batch for TGA. Include 1–25 samples plus the kaolinite standard.
7.2 Prepare Mg-saturated clay film on slide (created during dispersion and filtration of clay fraction). Refer to method 7A1a1, Sections 7.1 to 7.24.
7.3 Scrape clay from slide using a razor blade.
7.4 Lightly grind sample with pestle to make a homogeneous powder.
7.5 Sieve sample through an 80-mesh sieve. Equilibrate sample for 4 hours over a saturated magnesium nitrate solution (55% relative humidity) in a glass desiccator.

TGA Operation

7.6 Load empty pans into autosampler, making sure that they are lined up properly and that the bails are in line with each other.
7.7 Access Instrument Explorer. Input batch information, including sample numbers, project information, and layer key.
7.8 Run the tare procedure on the empty pans. This stores the pan weights.
7.9 Take autosampler tray from machine. Use spatula to place approximately 10 mg (estimated by volume) of sample into each pan. Load the pans back
into the autosampler tray, making sure that they are seated correctly and that their bails are lined up with one another.

7.10  Run “Preweigh” procedure on filled pans. This stores sample weights.

7.11  Before starting the run, make sure that batch information is still correct and that the right pan number is associated with the right sample. Start the run.

7.12  Access Universal Analysis software. Display file and calculate weight loss in specific regions for selected minerals.

8. Calculations

The thermogravimetric curve is displayed on the computer monitor. The ordinate (Y) is expressed in a relative weight percentage, i.e., the initial sample weight is 100.0%. Use the computer to calculate the total change in sample weight (ΔY), within the predetermined temperature range, as a sample weight percent.

8.1  % Kaolinite = \[\frac{(Δ \text{ sample weight} \% 450–550 \, °C)}{14}\] x 100

or

\[(Δ \text{ sample weight} \% 450–550 \, °C) \times 7.14\]

where:

Δ sample weight = total change in sample weight expressed as relative percent

14 = percent weight of hydroxyl water lost from pure kaolinite during dehydroxylation

8.2  % Gibbsite = \[\frac{(Δ \text{ sample weight} \% 250–350 \, °C)}{34.6}\] x 100

or

\[(Δ \text{ sample weight} \% 250–350 \, °C) \times 2.89\]

where:

Δ sample weight = total change in sample weight expressed as relative percent of the 110 °C base weight.

34.6 = percent weight of hydroxyl water lost from pure gibbsite during dehydroxylation

If Fe oxides are removed prior to analysis to prevent the interference with gibbsite determination, the calculation is modified to account for weight loss due to deferration as follows:
8.3 % Gibbsite = {[(Δ Sample weight % 250–350 °C x (Wt₂/Wt₁))/34.6] x 100
where:
Wt₁ = Weight before deferration
Wt₂ = Weight after deferration

The percent weights of hydroxyl water lost from kaolinite and gibsite are derived from the following assumed dehydroxylation reactions.

8.4 \( \text{Si}_2\text{Al}_2\text{O}_5(\text{OH})_4 \rightarrow 2\text{SiO}_2 + \text{Al}_2\text{O}_3 + 2\text{H}_2\text{O} \) (kaolinite)

8.5 \( 2\text{Al} (\text{OH})_3 \rightarrow \text{Al}_2\text{O}_3 + 3\text{H}_2\text{O} \) (gibbsite)

Using kaolinite as an example, percent weight of hydroxyl water lost is calculated from the following formula weights.

8.6 \( \text{Si}_2\text{Al}_2\text{O}_5(\text{OH})_4 = 258 \text{ g mol}^{-1} \)
\( 2\text{H}_2\text{O} = 36 \text{ g mol}^{-1} \)
Percent weight of hydroxyl water lost = \( \frac{36}{258} \times 100 = 14\% \)

If serpentine minerals are present in the sample, TGA can be used to quantify these minerals (Burt et al., 2001) based on an onset temperature of 600 to 650 °C (Karathanasis and Harris, 1994) and a weight loss from 600 to 900 °C (12.9%) based on the mineral structure \( \text{Mg}_3\text{Si}_2\text{O}_5(\text{OH})_4 \).

8.7 % Serpentine minerals = \[\frac{\Delta \text{ sample weight % 600–900 °C}}{12.9}\] x 100
or
\( \Delta \text{ sample weight % 600–900 °C} \times 7.75 \)

Gypsum can be quantified based on a loss of 20.9% (Karathanasis and Harris, 1994) based on the weight loss in the region of 100 to 350 °C.

8.8 % Gypsum = \[\frac{\Delta \text{ sample weight % 100–350 °C}}{20.9}\] x 100
or
\( \Delta \text{ sample weight % 100–350 °C} \times 4.78 \)

9. Report
Report percent gibbsite, kaolinite, gypsum, or antigorite to nearest whole number.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.
11. References


Instrumental Analysis (7A)
Surface Area (7A5)
   \( \text{N}_2 \) Adsorption (7A5a)
  Brunauer, Emmett, and Teller (BET) Theory (7A5a1)
   Vacuum Degassing (7A5a1a)
      Multi-point (7A5a1a1)
         Air-Dry, <2 mm (7A5a1a1a1)
   Single Point (7A5a1a2)
      Air-Dry, <2-mm (7A5a1a2a1)

1. Application

Surface area influences many physical and chemical properties of materials, e.g., physical adsorption of molecules and the heat loss or gain that results from this adsorption, shrink-swell capacity, water retention and movement, cation exchange capacity, pesticide adsorption, and soil aggregation (Carter et al., 1986). In addition, many biological processes are closely related to specific surfaces. Soils vary widely in their relative surface area because of differences in mineralogical and organic composition and differences in particle-size distribution. Specific surface area (SSA) is an operationally defined concept, dependent upon the measurement technique and sample preparation (Pennell, 2002).

The most common approach used to measure SSA is considered indirect. It is based on measurements of the adsorption or retention of probe molecules on a solid surface at monolayer coverage (Pennell, 2002). Two common methods of measuring SSA are by ethylene glycol monoethyl ether (EGME) and \( \text{N}_2 \)-sorption, using the theory of Brunauer, Emmett, and Teller (\( \text{N}_2 \)-BET). \( \text{N}_2 \) is a nonpolar gas and does not interact with, or have access to, interlayer crystallographic planes of expandable clay minerals. \( \text{N}_2 \) is therefore considered to provide a measure of external surface area. Polar molecules, such as EGME, are known to penetrate the interlayer surfaces of expandable clay minerals and therefore have been used to provide a measure of total surface areas (internal + external surface area) (Pennell, 2002). Significant differences between these methods are most apparent in soils containing expandable clay minerals and soil organic matter (Chlou and Rutherford, 1993; Pennell et al., 1995; de Jong, 1999; Quirk and Murray, 1999). In the past, the KSSL determined surface area by glycerol retention (7D1, method obsolete) or EGME retention (7D2, method obsolete). The current method, which is described herein, is \( \text{N}_2 \)-BET (multi-point).

2. Summary of Method

A <2-mm, air-dry soil sample is ground to pass a 0.25 mm (60 mesh) sieve and oven-dried (24 h, 110 °C). Enough soil (typically 0.5 to 1 g) is added to weighed sample cell to achieve 2- to 50- \( \text{m}^2 \) total area. Soil is cleaned of contaminants,
e.g., water and oils, by vacuum degassing at 10 millitorr for a minimum of 3 h at 110 °C and then reweighed to obtain degassed sample weight. The sample is brought to a constant temperature by means of an external bath (77 °K), and then small amounts of gas (N₂), called the “absorbate,” are admitted in steps to evacuate sample chamber. Gas molecules that stick to the surface of the solid (absorbent) are said to be adsorbed and tend to form a thin layer that covers the entire adsorbate surface. The number of molecules required to cover the adsorbent surface with a monolayer of adsorbed molecules, Nₘ, can be estimated based on the BET theory. Multiplying Nₘ by the cross-sectional area of an adsorbate molecule yields the sample's surface area. Specific surface area is reported in m² g⁻¹ by method 7A5a1a1 or 7A5a1a2.

3. Interferences

Organic material can coat or cover mineral surfaces, generally reducing SSA as determined by N₂-BET. Removal of organic matter prior to analysis typically increases these values. Freeze-drying may provide SSA values that are more representative of field conditions because air-drying may result in the collapse and shrinkage of soil humic acid whereas freeze-drying maintains an intricate structural network more characteristic of a natural state (Pennell, 2002). Sample size will vary depending on the SSA of the solid.

4. Safety

The outer surfaces of the heating mantle may become hot during use. Do not hold hot heating mantles without wearing protective gloves. Never insert fingers inside the pocket to determine if the mantle is heating up. Do not place a hot heating mantle on a surface that is not heat-resistant. Switch off the heating mantle when not in use. Refer to the manufacturer’s manual for safe operation of the surface area analyzer. Handle liquid nitrogen with care. Skin and eye contact with liquid nitrogen is hazardous due to the low temperatures. Use liquid nitrogen in well-ventilated areas; high concentrations of nitrogen gas in an enclosed area can cause asphyxiation. Because liquid nitrogen boils and splashes when added to a warm container, liquid nitrogen should be added slowly to minimize these reactions.

5. Equipment

5.1 Surface area analyzer, with vacuum degassing, Quantachrome, Nova 3000 Series, Boynton Beach, FL
5.2 Computer, with Nova, Enhanced Data Reduction Program, Version 2.13, Quantachrome, Boynton Beach, FL, and printer
5.3 Vacuum pump, capable of achieving 50 millitorr required, 10 millitorr recommended
5.4 Dewar flask with insulated lid for liquid N₂
5.5 Oven, 110 °C
5.6 Sample cells, glass, with outside stem diameters of 6, 9, and 12 mm and internal diameters of 4, 7, and 10 mm, respectively
5.7 Glass filler rods, 6 mm x 268.5 mm, 6 mm x 131.5 mm
5.8 Mortar and pestle

6. Reagents
6.1 N₂, high purity, 99.9%
6.2 Liquid N₂ (77 °K)
6.3 Standard reference material, 31.16 m² g⁻¹ SSA, with ±2.03 m² g⁻¹ reproducibility, Quantachrome, Boynton Beach, FL

7. Procedure

   Surface Area Analyzer Set-up and Operation

7.1 Set N₂-regulator at 10 PSIG (70 kPa).
7.2 Refer to the manufacturer’s manual for the routine operation, manifold calibration, and sample cell calibration.
7.3 The dosing manifold is factory calibrated, and there is no need to repeat this calibration before every analysis. The operator should check this calibration periodically (e.g. once every month), or if changes to the system may have altered the manifold volume.
7.4 A sample cell calibration needs to be performed for each combination of sample cell + filter rod + station and for each combination of adsorbate + coolant. Once a calibration is complete, further calibration for that particular combination is unnecessary. For most users, all standard (bulbless) cells can be considered equivalent for each diameter, i.e., one cell + rod + station calibration suffices for all cells of the same diameter with the same rod in the same station.

   Sample Preparation and Vacuum Degas

7.5 Pulverize a <2-mm, air-dry soil sample with a mortar and pestle to break up any aggregates.
7.6 Weigh sample cell to the nearest 0.1 mg. Weigh enough soil (typically 0.5 to 1.0 g) into sample cell to achieve 2- to 50-m² total area (sample size will vary depending on the SSA of the soil). Oven-dry at 110 °C for 24 h.
7.7 Remove sample cell with soil from oven and seal immediately. Allow to cool to touch.
7.8 Place the sample cell in the pouch of the heating mantle, set clamp in place, insert cell into fitting, tighten fitting, and loop elastic cords over hooks provided. Set the degas temperature at 110 °C for a minimum of 3 h.
7.9 Upon completion of degassing, switch the mantle off. Allow sample to cool. Unload degasser when ready to analyze sample.

7.10 Remove cell and reweigh to obtain dry, degassed soil sample weight to the nearest 0.1 mg.

7.11 Using the “Preset Analysis” option on surface area analyzer allows the user to preset and save the following: User ID, stations for analysis, setup files, cell numbers, sample ID numbers, and comments.

Analysis

7.12 Ensure Dewar flask is filled to the red line (approximately 1 in from top) with liquid \( N_2 \). Allow 5 min for liquid \( N_2 \) to equilibrate for best results. If the liquid \( N_2 \) is still boiling heavily, then the Dewar needs to be cleaned before filling with liquid \( N_2 \). If boiling continues after a dry clean Dewar is filled for 5 min, then replace the Dewar. Residual boiling will require that the Dewar be topped-off again. Ensure proper alignment of sample cells with Dewar mouth. Foam cap is not required for BET analysis only. If, however, a long isotherm is required, then use the foam cap. Ensure that there is no condensed water visible in the stem of the sample cell.

7.13 Follow set-up instructions on the surface area analyzer (e.g., select cell for Stations, enter sample ID for Stations, enter dry degassed sample weight by measuring sample volume, etc.) until prompted to proceed with analysis. Analytical data include Multi Point BET (adsorption), slope, intercept, correlation coefficient, BET C, total surface area in cell \( (m^2) \), and specific surface area \( (m^2 g^{-1}) \). Set-up and analytical data are automatically recorded by computer and printer.

8. Calculations

The BET equation for determination of the surface area of solids (Quantachrome Corp., 2000) is as follows:

\[
1 / \{W[(P_0/P)−1)]\} = [1/(W_m C)] + \{[(C−1)/W_m C]\}x(P/P_0)
\]

where:

- \( W \) = weight of gas adsorbed at a relative pressure of \( P/P_0 \)
- \( W_m \) = weight of adsorbate constituting a monolayer of surface coverage
- \( C \) = BET C constant, related to energy of adsorption in first adsorbed layer and indicative of the magnitude of absorbed/adsorbate interactions (typically, 50 to 250 for most solid surfaces)

The BET equation requires a linear plot of \( 1 / \{W[(P_0/P)−1)]\} \) versus \( P/P_0 \) which for most solids, using nitrogen as adsorbate, is restricted to a limited region of the adsorption isotherm, usually in the \( P/P_0 \) range of 0.05 to 0.35 (Quantachrome
The standard multi-point BET procedure requires a minimum of three points in the appropriate relative pressure range. The weight of the monolayer of adsorbate \( W_m \) can then be obtained from the slope \( s \) and intercept \( i \) of the BET plot.

\[
\begin{align*}
8.2 & \quad \text{From the BET equation,} \\
8.2.1 & \quad s = \frac{(C - 1)}{W_m C} \\
8.2.2 & \quad i = \frac{1}{W_m C}
\end{align*}
\]

Thus, the weight of the monolayer \( W_m \) can be obtained by combining these two equations as follows:

\[
W_m = \frac{1}{(s + i)}
\]

Total surface area \( S_t \) of the sample is calculated as follows:

\[
S_t = \frac{W_m N A_{cs}}{M}
\]

where:

\[
\begin{align*}
N &= \text{Avogadro's number (6.023 x 10^{23})} \\
M &= \text{Molecular weight of adsorbate (28.02 g mol}^{-1}) \\
A_{cs} &= \text{Close-packed nitrogen monolayer at 77 °K, the cross-sectional area for nitrogen = 16.2 Angstroms}^{2}
\end{align*}
\]

Specific surface area \( S \) is calculated as follows:

\[
S = \frac{S_t}{w}
\]

where:

\[
w = \text{Degassed sample weight}
\]

9. **Report**

Report specific surface area in \( \text{m}^2\text{g}^{-1} \) to the nearest 0.01 unit.

10. **Precision and Accuracy**

    Precision and accuracy data are available from the KSSL upon request.

11. **References**


Instrumental Analyses (7A)
Visible and Near-Infrared Diffuse Reflectance Spectroscopy (VNIR–DRS) (7A6)
   Air-Dry, <2 mm (7A6a1)

1. Application
   VNIR–DRS is the measurement of diffuse reflected spectra of a sample after exposure to visible near-infrared radiation (350–2500 nm) (McWhirt, 2012; Workman and Springsteen, 1998; Sparks, 1996). The spectrometer contains a dispersive energy source that enables the intensity at different wavelengths to be detected and recorded (McWhirt, 2012; Workman and Springsteen, 1998). The resulting spectrum of DRS does not necessarily produce a directly proportional relationship between wavelength intensity and analyte concentration; therefore, corrections and statistical analysis must be used to interpret the resulting spectra and to build models (McWhirt, 2012). Refer to Workman and Springsteen (1998) for multivariate regression techniques often used for this statistical application.

   VNIR–DRS provides nondestructive, noninvasive, and nearly instantaneous measurements. Small samples that require minimal sample preparation are adequate for VNIR–DRS analysis. VNIR analysis has been used in the evaluation and prediction of a wide range of soil properties, including, but not limited to, exchangeable cations, cation-exchange capacity, pH, organic C, free Fe, particle-size separates, gravimetric water content, and relative kaolinite and smectite (Ben-Dor and Banin, 1995; Shepherd and Walsh, 2002; Islam et al., 2003; Brown et al., 2006). VNIR–DRS has been used in the USDA–NRCS Rapid Carbon Assessment (RaCA) project, providing information on U.S. soil carbon (USDA–NRCS, 2013a, 2013b).

2. Summary of Method
   An air-dry, <2-mm soil sample is placed on sample holder and pressed at approximately 46 psi. The sample holder is placed onto Muglight, and scan is performed. The resulting VNIR pattern is stored for future data analysis.
3. Interferences
Calibration may be less accurate than other, more convention-based chemistry. An increased set of samples may therefore be required compared to these other methods. Measurements outside the range of calibration samples are invalid, and small calibration sample sizes can lead to overconfidence. Pure compounds can be positively identified only if a library of compounds is developed. Due to the IR signature of water, moist samples are typically not scanned by VNIR–DRS analysis.

4. Safety
Only trained personnel should use VNIR–DRS equipment. Follow the manufacturer’s safety precautions when using the VNIR–DRS.

5. Equipment
5.1 Visible and near-infrared diffuse reflectance spectrometer (VNIR–DRS) and sample holders (pucks), ASD Inc., LabSpec 2500, Analytical Spectral Devices, Boulder, CO
5.2 Computer with Indico Pro Software, ASD Inc., Boulder, CO
5.3 Press, approximately 46 psi
5.4 Pancake air compressor, 3-gal, 100 psi, oil-less, Central Pneumatic, item 95275, Harbor Freight Tools
5.5 Microfiber cloth

6. Reagents
6.1 Water, distilled
6.2 Isopropyl alcohol wipes, 70%
6.3 QC standards (high and low), appropriate reference materials with data models

7. Procedure
7.1 Clean lens and window:
7.1.1 To minimize dust coatings, use a clean, dry microfiber cloth to clean Muglight lens, inner and outer Muglight sample holder windows, and ASD White Reference window and ASD Wave Cal Puck window.
7.1.2 If smudges remain, slightly dampen microfiber cloth with distilled water and wipe lens or window. Thoroughly dry with a clean, dry microfiber cloth.
7.1.3 If smudges remain, use an isopropyl alcohol wipe and thoroughly dry with a clean, dry microfiber cloth.
7.2 Prepare spectrometer:

7.2.1 Make sure the Muglight is connected to spectrometer and that both are energized and properly warmed. Warm up Muglight for 3 h. Warm up spectrometer for 20 min after power is on. Alternatively, leave Muglight and spectrometer on continuously during times of constant or semi-constant usage.

7.2.2 Run Wavelength Analyzer program daily before scanning samples.

7.2.3 Check window on ASD White Reference Puck (#1) and lens on Muglight to ensure that they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.2.4 Load ASD White Reference Puck (#1) onto Muglight; push downward and inward to ensure it is in place. (For Pucks that have ASD White Reference on one side and ASD Wave Cal on the other, place the #1 side face down onto the Muglight).

7.2.5 Select Run and wait for completion of scan (approximately 30 s). ASD White Reference, ASD Wave Cal Puck, white reference, or soil samples should be scanned immediately upon loading onto Muglight. The exposure time of material to the intense light (heat) source affects the IR signature.

7.2.6 Remove ASD White Reference Puck (#1) from Muglight and place in clean plastic bag.

7.2.7 Check window on ASD Wave Cal Puck (#2) and lens on Muglight to ensure that they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.2.8 Load ASD Wave Cal Puck (#2) onto Muglight with #2 facing downward.

7.2.9 Select Run and wait for completion of scan (approximately 15 s). Print report and store.

7.2.10 Remove ASD Wave Cal Puck (#2) from Muglight, clean if necessary, and place in clean plastic bag.

7.3 Perform daily QC sample check:

7.3.1 Go to Start menu, select ASD Programs, and then select Indico Pro.

7.3.2 On the toolbar, select Open. Under project name, select QC and then OK.

7.3.3 On the toolbar, select Spectrum, Sample Count, Average, Averaging, None, Set Instrument Sample Count to 100, Select AB Even, and OK. This sequence is only necessary once per QC project. It is conducted at the time that the QC project is
initially set-up and the scans made. All future accesses to QC project will have spectra settings in place. Select set-up mode Reflectance.

7.3.4 Check window on Muglight white reference and lens on Muglight to ensure they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3. This white reference is not the same as the ASD White Reference Puck used for Wavelength Analyzer program.

7.3.5 Load white reference onto Muglight. On the toolbar, select Baseline, check Optimize First, and select Yes. Wait until the scan is complete (approximately 30 s). Result should be 1.00 on graph. Baseline should be taken at least every 15 to 20 min to avoid “drift.”

7.3.6 Remove white reference from Muglight. Clean if necessary and place in plastic bag or on clean cloth. Cover until next use.

7.3.7 Prepare QC labeled “Low.”

7.3.8 Check outside window of sample holder and Muglight lens to ensure they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.3.9 Place sample holder with “Low” QC onto Muglight.

7.3.10 On the toolbar, select Scan. Wait until scan is complete (approximately 15 s). Select Log 1/R mode.

7.3.11 Go to Chemometrics and Set-up. Under Project Predictors, select Galactic PLS PLS/IQ. Under selected predictor, select Galactic PLS/IQ. Click Add button. At the dialogue box, click on Desktop. Find and select the appropriate “.cal” file (QC.cal). Click Open. At the Chemometrics dialogue box, this file should show up under “Selected Model” files. Click Close.

7.3.12 On the toolbar, select Predict to view prediction and M-distance of QC sample labeled “Low” and record. If prediction or M-distance is outside of acceptable ranges, check fiber optic cable for proper connection, ensure Muglight and spectrometer are properly energized and warmed-up, and then repeat steps.

7.3.13 Prepare QC labeled “High.”

7.3.14 Check outside window of sample holder and Muglight lens to ensure that they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.3.15 Place sample holder with “High” QC onto Muglight.

7.3.16 On the toolbar, select Scan. Wait until scan is complete (approximately 15 s).
7.3.17 On the toolbar, select **Predict** to view prediction and M-distance of QC sample labeled “High” and record. If prediction or M-distance is outside of acceptable ranges, check fiber optic cable for proper connection, ensure Muglight and spectrometer are properly energized and warmed-up, and then repeat steps.

7.4 Clean sample holder after scanning:

7.4.1 Work over waste container and in front of dust collector. Turn sample holder upside down and gently tap with wooden dowel or knife handle until compacted soil falls out.

7.4.2 Use compressed air from an oil-less air compressor to blow out remaining residue from inside of sample holder. Set output regulator on compressor to 25 psi.

7.5 Prepare air-dry samples:

7.5.1 Prepare air-dry, <2-mm samples.

7.5.2 Check inner window of Muglight sample holder to ensure it is clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.5.3 Heap prepared sample into sample holder to slightly overflowing capacity.

7.5.4 Level sample to the top of sample holder by striking off excess sample with a straight edge.

7.5.5 Before packing the sample, support the underside of the sample holder window with a rubber stopper, preventing the window from being pushed out or broken. Properly align piston so it does not catch as it presses into the well of the sample holder. Brush away any excess from top and sides of sample holder while the sample is being packed, preventing the soil from spilling onto the Muglight lens.

7.5.6 Pack sample with approximately 46 psi pressure using press/penetrometer and hold for approximately 10 s.

7.5.7 After the sample is packed, wipe the bottom of the sample holder free of extraneous soil. Place the sample holder on a clean towel.

7.5.8 Load the sample holder onto the Muglight before scanning.

7.5.9 During compaction process, some soil particles may cling to the piston surface. Wipe face of piston before packing the next sample.

7.6 Scanning air-dry samples:

7.6.1 On the **Start** menu, select **ASD Programs** and then **Indico PRO**.
7.6.2 On the toolbar, select Open for appropriate project. Select set-up mode Reflectance.

7.6.3 On the toolbar, select Spectrum, Sample Count, Average, Averaging, None, Set Instrument Sample Count to 100, Select AB Even, and OK.

7.6.4 Check window on white reference and Muglight lens to ensure that they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.6.5 Load white reference onto Muglight.

7.6.6 On the toolbar, select Baseline. Ensure Optimize First is checked: Yes. Wait until scan is complete (approximately 30 s). Result should be straight line at 1.00 on the graph.

7.6.7 Check outside window of filled sample holder and on Muglight to ensure they are clean. Clean if necessary. Refer to Sections 7.1.1 to 7.1.3.

7.6.8 On the toolbar, select Scan. Wait until scan is complete (approximately 15 s).

7.6.9 On the toolbar, select Save.

7.6.10 Enter file name and save as type ASD Spectrum files (*.ADS), and click OK.

7.6.11 To clean sample holder after scanning, refer to Sections 7.4.1 and 7.4.2.

8. Calculations
   Not applicable.

9. Report
   The process of recording the spectral information is called a scan, the result of which is saved as a file in binary form for future corrections and statistical analysis to interpret spectra and to build models.

10. Precision and Accuracy
    Precision and accuracy data are available from the KSSL upon request.

11. References
Instrumental Analyses (7A)
Mid-Infrared Diffuse Reflectance Spectroscopy (MIR–DRS) (7A7)
Air-Dry, <2 mm (7A7a1)

1. Application

Mid-infrared diffuse reflectance spectroscopy (MIR–DRS) is a combination of spectroscopy and chemometric (multivariate analysis) methods used to identify and/or quantify chemical species. The resulting MIR spectrum is dependent on the composition of the soil, particularly on specific vibrational signatures for organic matter, quartz, kaolinite, smectite, carbonates, gypsum, and Fe and Al oxides (CSIRO, 2013). MIR–DRS can predict a wide range of chemical and physical properties closely associated with the soil bulk properties (e.g., clay, organic matter, moisture content, CEC, and mineralogy). It does not, however, directly measure any of these properties (CSIRO, 2013). The advantages of MIR spectral analysis for measurement of soil carbon are that the method is nondestructive, consumes no reagents, and permits simultaneous measurements of organic and inorganic carbon, thereby simplifying analysis relative to the traditional chemical methods (McCarty et al., 2002).
2. Summary of Method

A sample of <2-mm, air-dry soil is processed to ≈80 mesh (180 µm). Processed sample is placed in a microplate and pressed. Microplate is placed in microplate module. MIR–DRS analysis is performed, and the resulting MIR pattern is stored for future analysis.

3. Interferences

Because of the small area sampled with MIR–DRS, the homogeneity and particle-size of the sample are important to examine when the spectral results are interpreted (Baldock and Hawke, 2010). MIR–DRS can be used to predict total soil N, surface adsorption properties, and other total element concentrations in soils. MIR–DRS has difficulty with commonly used soil analyses based on the soil solution rather the soil matrix (e.g., extractable P, S, and N) because of the generally low concentration in the soil environment (Merry and Janik, 2013). MIR–DRS relies on calibration against recognized laboratory procedures for both chemical and physical properties and is therefore only as accurate as the data against which it is calibrated.

4. Safety

Only trained personnel should use MIR–DRS equipment. Follow the manufacturer’s safety precautions. Handle liquid nitrogen with care. Skin and eye contact with liquid nitrogen is hazardous due to the low temperatures. Use liquid nitrogen in well-ventilated areas; high concentrations of nitrogen gas in an enclosed area can cause asphyxiation. Because liquid nitrogen boils and splashes when added to a warm container, liquid nitrogen should be added slowly to minimize these reactions.

5. Equipment

5.1 Spectrometer, Vertex 70, FT-IR, Bruker Optics
5.2 Microplate module, HTS-XT, Bruker Optics
5.3 Microplates, 96-well sample plate, aluminum (fig. 7A7-1)
5.4 Opus 6.5 Software, Bruker Optics
5.5 Computer
5.6 Sticks, wooden
5.7 Steel rod, for packing sample plates (fig. 7A7-2)
5.8 Glass plate (fig. 7A7-1)
5.9 Weight, placed on glass plate to hold it in place (fig. 7A7-1)
5.10 Vacuum cleaner (fig. 7A7-3)
5.11 Funnel
5.12 Dewar flask with insulated lid for liquid N₂
Figure 7A7-1.—Filled sample plate with glass plate covering leftmost two columns and with weight holding glass plate in place.

Figure 7A7-2.—Steel rod for packing sample plates.

Figure 7A7-3.—Vacuum cleaner to clean sample plates.
6. Reagents
6.1 Reverse osmosis (RO) water, ASTM Type III grade reagent water
6.2 Isopropyl alcohol wipes, 70%
6.3 Liquid N₂ (77 °K)
6.4 Molecular sieves, enclosed in removable cartridges
6.5 QC standards, appropriate reference materials with data models

7. Procedure

Loading 96-Well Sample Plate with Air-Dry, Fine-Grind Soil

7.1 Position a clean plate with well A1 at the upper left corner.
7.2 Place cello tape over wells A1–D1 (blank cells) to prevent contamination.
7.3 Using a wooden spatula, place the standard sample material in wells E1–H1. Fill the wells with loose soil.
7.4 Pack the sample in each well with a steel rod. The surface of the sample needs to be flat. The height is less important, provided the sample is at least 1 mm thick in the well. Clean the rod after each sample.
7.5 Vacuum any excess soil from the top of the plate, using a plastic tubing fitting attached to rubber tubing attached to a vacuum cleaner hose. If necessary, wipe the surface of the plate around the wells with a cotton swab.
7.6 Place another 96-well plate upside down to the left of the plate being filled. Place a glass plate on the 96-well plates, positioning the glass so it covers the left column of the plate being filled. Place a weight on the glass to hold it in place.
7.7 Place 80-mesh material for the first unknown sample in wells B2–D2.
7.8 Repeat Sections 7.4 through 7.5.
7.9 Place 80-mesh material for the second unknown sample in wells E2–H2.
7.10 Slide the glass plate to the right so it covers the leftmost two columns on the plate.
7.11 Fill the rest of the plate in a similar manner.
7.12 Clean aluminum plate by soaking in soapy water and then using a soft brush and clear water. Rinse with RO water and then with ethanol before setting out to dry overnight.

Instrument Start-Up, Validation, Operation, and Calibration

7.13 Instrument remains on all the time. If instrument is switched on or off, the following sequence is used:
7.13.1 **Powering On**: Vertex 70, Twister, Computer, and Software
7.13.2 **Powering Off**: Software, Computer, Vertex 70, and Twister
7.13.3 If the instrument is powered off, it requires 1 h warm-up at restart before performing any test.

7.14 Open Opus software and provide information for User ID, Assigned Workspaces, and Password. Click *Login*.

7.15 Perform instrument performance qualification once a week and operational qualification once a year.

7.16 To calibrate the sample stage in the HTS-XT, go to menu item *Open Drawer*.

7.17 Load the 96-well plate in the HTS-XT with well A1 positioned near red dot of plate holder.

7.18 Go to menu item *Close Drawer*.

7.19 Go to menu item *Measure* and select *Motorized Stage Control*.

7.20 Click *Calibrate Stage* and then *Exit*.

7.21 Using Plate Positioning Utility on menu: If stage was not previously calibrated, go to *Calibrating Stage* and select *Continue*. Select sample well by clicking on the corresponding plate position (e.g., A1, D3, and H6). When position is selected, either click *Exit* or go to tab *Display, SSL.ows*.

### Sample Batch Set-Up

7.22 On the menu, select Run OPUS/Lab.

7.23 Select Measure and then HTS-XT.

7.24 For plate format, select *96-well plate*. Click *Initialize Plate*.

7.25 For Background Position, select a clean, empty position on the plate. Select *Background as Before Each Sample*.

7.26 Select sample positions to be scanned. Samples are analyzed in letter order first and then in number order.

7.27 Click *Create Sample Table*.

7.28 Enter microplate number (1,2,3, etc.).

7.29 Enter batch number and sample number.

7.30 Select and highlight each position of samples to be analyzed.

7.31 Click on the *Assign Selected Positions* bar. Repeat batch and sample number assignment process to assign batch and sample numbers for all remaining positions.

7.32 Click *OK* and *Return*.

7.33 Click *Measure*. Insert sample tray. Click *OK*. 

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**555**
7.34 Well positions that are orange have not been scanned. Well positions that are green have been scanned. Well positions that are white will not be scanned.

7.35 Upon completion of scanning, remove the plate and click OK.

**Instrument Maintenance**

7.36 Before any measurements are performed, cool the HTS-XT MCT detector with liquid nitrogen. Use a funnel to fill the MCT detector with liquid nitrogen. Wait 30 min before making spectral measurements. A single charge of liquid nitrogen lasts approximately 6 h before a refill is required. The hold-time of the cooled MCT depends less on environmental conditions than on the degree to which the Dewar is filled. To continuously keep the MCT detector cool during measurements, top off the detector with liquid nitrogen every 2 h. If the MCT is refilled by hand while still-cooled, the wait for stabilization is less than the wait when cooling down the warm detector. A minimum delay of 2 min before starting the next measurement is recommended.

7.37 Keep the air in the interferometer and detector chamber dry by using desiccant (molecular sieve) enclosed in removable cartridges. Replace or regenerate the desiccant cartridge approximately every month and whenever the cartridge indicator changes from blue to pink. The desiccant cartridges are located in the detector and interferometer compartments. Pour molecular sieves into clean beakers and heat to 230 °C for 4 h.

8. **Calculations**
   
   Not applicable.

9. **Report**
   
   The process of recording the spectral information is called a scan, the result of which is saved as a file in binary form for future corrections and statistical analysis to interpret spectra and to build models.

10. **Precision and Accuracy**
    
    Precision and accuracy data are available from the KSSL upon request.

11. **References**
    
Optical Analyses^2 (7B)
Grain Studies (7B1)
Analysis and Interpretation (7B1a)

Minerals

Identification criteria: Important properties in grain identification are listed below in approximate order of ease and convenience of determination. Because estimates of several of these properties commonly allow identification of a grain, detailed or extremely accurate measurements are seldom necessary. In the finer soil separates, grain identification may be impossible because the grains may be too small or not in the right position to permit measurement of some properties, e.g., optic angle (2V) or optic sign. The estimation of properties can be practiced by crushing, sieving, and mounting a set of known minerals and comparing these known standards to unknowns.

Refractive index is the ratio of the speed of light in the medium (mineral) to the speed of light in a vacuum. It can be estimated by relief or can be accurately determined by using calibrated immersion liquids. When relief is used to estimate refractive index, the grain shape, color, and surface texture are considered, i.e., thin platy grains may be estimated low, whereas colored grains and grains with rough, hackly surface texture may be estimated high. Estimation is aided by comparing an unknown to known minerals.

Relief is an expression of the difference in refractive index between the grain and the mounting medium. The greater the difference, the greater the relief. The analogy is to topographic relief. When viewed through the microscope, grains with high relief are distinct, whereas grains with low relief tend to fade into the background. The KSSL selects a mounting medium with an index

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^2 The discussion of identification and significance of minerals, microcrystalline aggregates, and amorphous substances in optical studies of grain mounts was from material after John G. Cady (1965), with permission, and modified by Warren C. Lynn, Research Soil Scientist, NRCS, Lincoln, NE.
of refraction close to quartz, which has low relief. Most other minerals are identified by comparison.

**Becke line** is a bright halo of light that forms near the contact of the grain and the mounting medium because of the difference in refractive index between the two. As the plane of focus is moved upward through the grain, the Becke line appears to move into the component with the higher refractive index. In Petropoxy 154™, the Becke line moves away from potassium feldspar (index of refraction <1.54) but moves into mica (index of refraction >1.54).

**Birefringence** is the difference between the highest and lowest refractive index of the mineral. Accounting for grain thickness and orientation, the birefringence is estimated by interference color. Interference color is observed when an anisotropic mineral is viewed between cross-polarized light. Several grains of the same species must be observed because the grains may not all lie in positions that show the extremes of refractive index. For example, the birefringence of mica is high but appears low when the platy mineral grain is perpendicular to the microscope axis because the refractive indices of the two crystallographic directions in the plane are similar. However, a mica grain viewed on edge in a thin section shows a high interference color. The carbonate minerals have extremely high birefringence (0.17 to 0.24). Most of the ferromagnesian minerals are intermediate (0.015 to 0.08). Orthoclase feldspar and apatite are low (0.008) and very low (0.005), respectively.

**Color** helps to discriminate among the heavy minerals. Pleochroism is the change in color or light absorption with stage rotation when the polarizer is inserted. Pleochroism is a good diagnostic characteristic for many colored minerals. Tourmaline, green beryl, and staurolite are examples of pleochroic minerals.

**Shape, cleavage, and crystal form** are characteristic or possibly unique for many minerals. Cleavage may be reflected in the external form of the grain or may appear as cracks within the grain that show as regularly repeated straight parallel lines or as sets of lines that intersect at definite repeated angles. The crystal shape may be different from the shape of the cleavage fragment. Plagioclase feldspars, kyanite, and the pyroxenes have strong cleavage. Zircon and rutile usually appear in crystal forms.

**Extinction angle and character of extinction** observed between cross-polarized light are important criteria for some groups of minerals. For extinction angles to be measure, the grain must show its cleavage or crystal form. These angles may be different along different crystallographic axes. Some minerals have sharp, quick total extinction, whereas other minerals have more gradual extinction. In some minerals with high light dispersion, the interference color dims and changes at the extinction position.

**Optic sign, optic angle, and sign of elongation** are useful, if not essential, determinations but are often difficult, unless grains are large or in favorable orientation. Determination of optic sign requires that the grains show dim, low-
order interference colors or show no extinction. Grains with bright colors and with sharp, quick extinction rarely provide usable interference figures.

**Particular mineral species:** The following are the outstanding diagnostic characteristics of the most commonly occurring minerals and single-particle grains in the sand and silt fractions of soils. The refractive indices that are provided are the intermediate values.

**Quartz** has irregular shapes. The refractive index of quartz (1.54) approximates that of the epoxy (Petropoxy 154™) mounting medium. The Becke line may be split into yellow and blue components. The interference colors are low order but are bright and warm. There is sharp extinction with a small angle of rotation, i.e., “blink extinction.” Crystal forms are sometimes observed and usually indicate derivation from limestone or other low-temperature secondary origin.

**Potassium feldspars:** *Orthoclase* may resemble quartz, but the refractive index (1.52) and birefringence are lower than that of quartz. In addition, orthoclase may show cleavage. *Microcline* has a refractive index of 1.53. The Becke line moves away from the grain with upward focus. A twinning intergrowth produces a plaid or grid effect between cross-polarized light that is characteristic of microcline. *Sanidine* has the same refractive index and birefringence as other potassium feldspars. Grains are usually clear, and twinning is not evident. In sanidine, the 2V angle is low (12°) and characteristic. The 2V angle is the acute angle between two optic axes, or more simply, the optical axial angle.

**Plagioclase feldspars** have refractive indices that increase with an increase in the proportion of calcium. The refractive index of the sodium end-member albite (1.53) is lower than that of quartz, but the refractive index of the calcium end-member anorthite (1.58) is noticeably higher than that of quartz. Some *oligoclase* has the same refractive index as quartz; thus distinctions cannot be made by the Becke line. Plagioclase feldspars often show a type of twinning (defined as albite twinning) that appears as multiple alternating dark and light bands in cross-polarized light. Cleavage is good in two directions parallel to (001) and (010), often producing lathlike or prismatic shapes.

**Micas** occur as platy grains that are commonly very thin. The plate view shows very low birefringence, whereas the edge view shows a very high birefringence. Plates are commonly equidimensional and may appear as hexagons or may have some 60° angles. *Biotite* is green to dark brown. Green grains may be confused with chlorite. Paler colors, a lowering of refractive index, and a distortion of the extinction and interference figure indicate weathering to *hydrobiotite*, *kaolinite*, or *vermiculite*. *Muscovite* is colorless. Muscovite has a moderate refractive index (1.59) in the plate view and an interference figure that shows a characteristic 2V angle of 30° to 40°, which can be used as a standard for comparing 2V angles of other minerals.

**Amphiboles** are fibrous to platy or prismatic minerals that typically have slightly inclined extinction but occasionally have parallel extinction. Color and
refractive index increase as the Fe content increases. Amphiboles have good cleavage at angles of ≈56° and 124°. Refractive index of the group ranges from 1.61 to 1.73. Hornblende is the most common member of the amphiboles. It is slightly pleochroic, usually has a distinctive color close to olive-green, has inclined extinction, and is often used as an indicator of weathering.

Pyroxenes: Enstatite and aegirine-augite are prismatic and have parallel extinction. Aegirine-augite has unique and striking green-pink pleochroism. Augite and diopside have good cleavage at angles close to 90° and large extinction angles. Colors usually are shades of green, with interference colors of reds and blues. Refractive indices in the pyroxenes (1.65 to 1.79) are higher than those for amphiboles.

Olivine is colorless to very pale green and usually irregular in shape (weak cleavage). It has vivid, warm interference colors. It is an easily weathered mineral and may have cracks or seams filled with serpentine or goethite. It is seldom identified in soils, but has been observed in certain soils from Hawaii.

Staurolite is pleochroic yellow to pale brown and sometimes contains holes, i.e., the “Swiss cheese” effect. The refractive index is ≈1.74. Grains may have a foggy or milky appearance, which may be caused by colloidal inclusions.

Epidote is a common heavy mineral, but the forms that occur in soils may be difficult to identify positively. Typical epidote is unmistakable with its high refractive index (1.72 to 1.76), strong birefringence, and a pleochroism that includes pistachio-green color. The typical interference colors are reds and yellows. Commonly, grains show an optic axis interference figure with a 2V angle that is nearly 90°. However, epidote is modified by weathering or metamorphism to colorless forms with lower birefringence and refractive index. Zoisite and clinozoisite in the epidote group are more common than some of the literature indicates. These minerals of the epidote group commonly appear as colorless, pale-green, or bluish-green, irregularly shaped or roughly platy grains with high refractive index (1.70 to 1.73). Most of these minerals show anomalous interference colors (bright pale blue) and no complete extinction and can be confused with several other minerals, e.g., kyanite and diopside. Zoisite has a distinctive deep blue interference color. Identification usually depends on determination of properties for many grains.

Kyanite is a common mineral but is seldom abundant. A pale blue color, large cleavage angles, large extinction angles (30° extinction), and platy, angular cleavage flakes usually can be observed and make identification easy.

Sillimanite and andalusite resemble each other. These minerals are fibrous to prismatic with parallel extinction. However, their signs of elongation are different. In addition, sillimanite is colorless, and andalusite commonly is pink.

Garnet occurs in irregularly shaped, equidimensional grains that are isotropic and have high refractive index (≥1.77). Garnet the size of fine sand and silt is commonly colorless. Pale pink or green colors are diagnostic in the larger grains.
**Tourmaline** has a refractive index of 1.62 to 1.66. Prismatic shape, strong pleochroism, and parallel extinction are characteristic. Some tourmaline is almost opaque when at right angles to the vibration plate of the polarizer.

**Zircon** occurs as tetragonal prisms with pyramidal ends. Zircon has very high refractive index (>1.9), parallel extinction, and bright, strong interference colors. Broken and rounded crystals frequently occur. Zircon crystals and grains are almost always clear and fresh appearing.

**Sphene**, in some forms, resembles zircon, but the crystal forms have oblique extinction. The common form of sphene, a rounded or subrounded grain, has a color change through ultrablue with crossed polarizers instead of extinction because of its high dispersion. Sphene is the only pale-colored or colorless high-index mineral that provides this effect. It is amber colored in reflected light. The refractive index of sphene is slightly lower than that of zircon, and the grains are commonly cloudy or rough-surfaced.

**Rutile** grains have prismatic shape. The refractive index and birefringence are extremely high (2.6 to 2.9). The interference colors typically are obscured by the brown, reddish-brown, or yellow colors of the mineral. Other TiO₂ minerals, *anatase* and *brookite*, also have very high refractive indices and brown colors and may be difficult to distinguish in small grains. The anatase and brookite typically occur as tabular or equidimensional grains.

**Apatite** is common in youthful soil materials. Apatite has a refractive index slightly <1.63 and a very low birefringence. Crystal shapes are common, may appear as prisms, and are often bullet shaped. Rounding by solution produces ovoid forms. Apatite is easily attacked by acid and may be lost in pretreatments.

**Carbonates:** *Calcite*, *dolomite*, and *siderite*, in their typical rhombohedral cleavage forms, are easily identified by their extremely high birefringence. In soils, these minerals have other forms, e.g., scales and chips; cements in aggregates; microcrystalline coatings or aggregates; and other fine-grained masses that are often mixed with clay and other minerals. The extreme birefringence is always the identification clue and is shown by the bright colors between cross-polarized light and by the marked change in relief when the stage is rotated with one polarizer in. The microcrystalline aggregates produce a twinkling effect when rotated between cross-polarized light. These three minerals have differences in their refractive indices, which may be used to distinguish them. Siderite is the only one with both indices greater than those of Petropoxy 154™. It is more difficult to distinguish calcite from dolomite, and additional techniques, such as staining or x-ray diffraction, may be used.

**Gypsum** occurs in platy or prismatic, flat grains with refractive index approximately equal to orthoclase. It usually has a brushed or “dirty” surface.

**Opaque** minerals, of which *magnetite* and *ilmenite* are the most common, are difficult to identify, especially when they are worn by transportation or otherwise affected by weathering. Observations of color and luster by reflected light,
aided by crystal form if visible, are the best procedures. Magnetic separations help to confirm the presence of magnetite and ilmenite. Many grains that appear opaque by plain light can appear translucent if viewed between strong cross-polarized light. Most grains that behave in this way are altered grains or aggregates and are not opaque minerals.

**Microcrystalline Aggregates and Amorphous Substances**

**Identification criteria:** Most microcrystalline aggregates have one striking characteristic feature, i.e., they show birefringence but do not have definite, sharp, complete extinction in cross-polarized light. Extinction may occur as dark bands that sweep through the grain or parts of the grain when the stage is turned or may occur in patches of irregular size and shape. With a few exceptions, e.g., well-oriented mineral pseudomorphs and certain clay-skin fragments, some part of the grain is bright in all positions. Aggregates and altered grains should be examined with a variety of combinations of illumination and magnification in both plain and polarized lights. Following is a discussion of the principal properties that can be used to identify or at least characterize aggregates.

**Color,** if brown to bright red, is usually related to Fe content and oxidation. Organic matter and Mn may contribute black and grayish-brown colors.

**Refractive index** is influenced by a number of factors, including elemental composition, atom packing, water content, porosity, and crystallinity. Amorphous (noncrystalline) substances have a single index of refraction, which may vary depending on chemical composition. For example, allophane has a refractive index of 1.47 to 1.49, but the apparent refractive index increases with increasing inclusion of ferrihydrite (noncrystalline Fe oxide) in the mineral.

**Strength of birefringence** is a clue to the identity of the minerals. Even though the individual units of the aggregate are small, birefringence can be estimated by interference color and brightness. Amorphous substances, having only a single index of refraction, exhibit no birefringence and are isotropic between cross-polarized light.

**Morphology** may provide clues to the composition or origin of the aggregate. Some aggregates are pseudomorphs of primary mineral grains. Characteristics of the original minerals, i.e., cleavage traces, twining, or crystal form can still be observed. Morphology can sometimes be observed in completely altered grains, even in volcanic ash shards and basalt fragments. Other morphological characteristics may be observed in the individual units or in the overall structure. For example, the units may be plates or needles, or there may be banding.

**Particular species of microcrystalline aggregates and amorphous substances:** For purposes of soil genesis studies, the aggregates that are present in sand or silt fractions are not of equal significance. Some are nuisances but must be accounted for, and others are particles with important
diagnostic value. Useful differentiating criteria for some of the commonly occurring aggregate types are discussed below.

*Rock fragments* include chips of shale, schist, and fine-gained igneous rocks, e.g., rhyolite. Identification depends on the recognition of structure and individual components and the consideration of possible sources. Rock fragments are common in mountainous regions and are often hydrothermally-altered in the western United States.

*Clay aggregates* may be present in a wide variety of forms. Silt and sand that are bound together into larger grains by a nearly isotropic brownish material usually indicate incomplete dispersion. Clay skins may resist dispersion and consequently may appear as fragments in grain mounts. Such fragments are usually brown or red and translucent with wavy extinction bands. Care is required to distinguish these fragments from weathered biotite. Clay aggregates may be mineral pseudomorphs. Kaolin pseudomorphs of feldspar are common. Montmorillonite aggregates, pseudomorphic of basic rock minerals, have been observed. In this form, montmorillonite shows high birefringence and an extinction that is mottled or patchy on a small scale. Coarse kaolinite flakes, books, and vermicular aggregates resist dispersion and may be abundant in sand and silt. These particles may resemble muscovite but are cloudy. They show no definite extinction and have very low birefringence. Many cases of anomalously high cation exchange capacity (CEC) of sand and silt fractions that are calculated from whole soil CEC and from clay CEC and percent content can be accounted for by the occurrence of these aggregates in the sand and silt fractions.

*Volcanic glass* is isotropic and has a low refractive index, lower than most of the silicate minerals. The refractive index ranges from 1.48 in the colorless siliceous glasses to as high as 1.56 in the green or brown glasses of basalt composition. Shapes vary, but the elongated, curved shard forms, often with bubbles, are common. This glassy material may adhere to or envelop other minerals. Particles may contain small crystals of feldspar or incipient crystals with needles and dendritic forms. The colorless siliceous types (acidic, pumiceous) are more common in soils, as the basic glasses weather easily. Acidic glasses are more commonly part of “ash falls”, as the magma usually is gaseous and explosive when pressure is released. Basic glasses are more commonly associated with volcanic flow rocks, which are usually not gaseous.

*Allophane* is present in many soils that are derived from volcanic ash. Allophane seldom can be identified directly, but its presence can be inferred when sand and silt are cemented into aggregates by isotropic material with low refractive index, especially if volcanic ash shards are also present.

*Opal*, an isotropic material, occurs as a cementing material and in separate grains, some of which are of organic origin, i.e., plant opal, sponge spicules, and diatoms. The refractive index is very low (<1.45), which is lower than the value for volcanic ash. Identification may depend in part on form and occurrence.
Iron oxides may occur as separate grains or as coatings, cementing agents, and mixtures with other minerals. Iron oxides impart brown and red colors and raise the refractive index in the mixtures. Goethite is yellow to brown. Associated red areas may be hematite. These red varieties have a refractive index and birefringence that are higher and seem to be better crystallized, often having a prismatic or fibrous habit. Aggregates have parallel extinction. In oriented aggregates, the interference colors often have a greenish cast. Hematite has higher refractive index than goethite and is granular rather than prismatic. Large grains of hematite are nearly opaque.

Gibbsite often occurs as separate, pure, crystal aggregates, either alone or inside altered mineral grains. The grains may appear to be well-crystallized single crystals, but close inspection in cross-polarized light shows patchy, banded extinction, indicating intergrown aggregates. Gibbsite is colorless. The refractive index (1.56 to 1.58) and the birefringence are higher for gibbsite than the corresponding values for quartz. The bright interference colors and aggregate extinction are characteristic of gibbsite.

Chalcedony is a microcrystalline form of quartz that was formerly considered a distinct species. Chalcedony occurs as minute quartz crystals and exhibits aggregate structure with patchy extinction between cross-polarized light. It may occur in nodules of limestone deposits and may be a pseudomorphic replacement in calcareous fossils. The refractive index is slightly lower than that of quartz, and the birefringence is lower than that of gibbsite. Chert is a massive form of chalcedony.

Glaucnite occurs in aggregates of small micaceous grains with high birefringence. When fresh, glauconite is dark green and almost opaque, but it weathers to brown and more translucent forms. Glaucnite is difficult to identify on optical evidence alone. Knowledge of source area or history is helpful in identification.

Titanium oxide aggregates have been tentatively identified in the heavy mineral separates of many soils. These bodies have an extremely high refractive index and high birefringence and thus are similar to rutile. Their yellow to gray colors are similar to those of anatase. The TiO₂ aggregates are granular and have a rough surface. This growth habit with the little spurs and projections suggests that TiO₂ aggregates may be secondary.

References
Optical Analyses (7B)
Grain Studies (7B1)
Separation by Heavy Liquids (7B1a1)

1. Application

The sand and silt fractions of most soils are dominated by quartz or by quartz and feldspars (Cady, 1965). These minerals have a relatively low specific gravity (2.57 to 2.76). The large numbers of “heavy” mineral grains (i.e., having specific gravity >2.8 or 2.9) with a wide range of weatherability and diagnostic significance may be only a small percentage of the grains (Cady, 1965). However, these “heavy” minerals are commonly indicative of provenance, weathering intensity, and parent material uniformity (Cady et al., 1986).

2. Procedure

To study “heavy” minerals, a common practice is to concentrate these grains by specific-gravity separations in a heavy liquid. This method (7B1a1) is rarely used at the KSSL, generally only for special studies.

Micas are difficult to separate because of their shape and because a little weathering, especially in biotite, significantly decreases the specific gravity. These differences in density in biotite may be used to concentrate weathered biotite in its various stages of alteration.

Separation of grains by heavy liquids is most effective when grains are clean. Organic matter may prevent wetting and cause grains to clump or raft together. Light coatings may cause heavy grains to float, and iron-oxide coatings may increase specific gravity. For some kinds of materials, an additional technique is to separate and weigh the magnetic fraction, either before or after the heavy-liquid separation.

Concentrate the “heavy” minerals, i.e., those with specific gravity >2.8 or 2.9, by specific-gravity separations in a heavy liquid. The reagent of choice is sodium polytungstate (density 2.8 g⁻¹ mL). Dilute the sodium polytungstate with distilled water to obtain required densities <2.8 g⁻¹ mL. Use a specific gravity ≈2.5, to concentrate volcanic glass, plant opal, or sponge spicules. When using this liquid, avoid contact with skin and work in a well-ventilated area.

Separation by specific gravity alone in separatory funnels, cylinders, or various kinds of tubes is usually adequate for grains >0.10 mm. Separation by centrifuging is required for grains <0.10 mm. Use pointed, 15-ml centrifuge tubes for these separations.

Decant the light minerals after inserting a smooth-bulb glass rod to stop off the tapered end of centrifuge tube. Alternatively, remove the heavy minerals by gravity flow using a lower stopcock or maintain the heavy minerals in place by freezing the lower part of tube.
3. References

Optical Analysis (7B)
Grain Studies (7B1)
Grain Mounts, Epoxy (7B1a2)

1. Application
Grain counts are used to identify and quantify minerals in the coarse silt and sand fractions of soils. The results are used to classify soil pedons in mineralogy families of soil taxonomy (Soil Survey Staff, 2014), to help determine substrate provenance of source materials, and to support or identify lithologic discontinuities.

2. Summary of Method
In particle-size analysis, soils are dispersed so that material <20 µm in diameter is separated by settling and decanting and the sand and coarse silt fractions are separated by sieving. Refer to the procedure for the separation by heavy liquids of the less abundant minerals with a specific gravity >2.8 or 2.9 (method 7B1a1).
Following sample selection, permanent mounts are prepared for the two most abundant particle-size fractions among the fine sand, very fine sand, and coarse silt. The grains are mounted in a thermo-setting epoxy cement with a refractive index of 1.54. The grains are then identified and counted under a petrographic microscope.

A mineralogical analysis of a sand or silt fraction may be entirely qualitative, or it may be quantitative to different degrees (Cady, 1965). The KSSL performs a quantitative analysis (7B1a2). Data are reported as a list of minerals and an estimated quantity of each mineral as a percentage of the grains counted in the designated fraction. The percentages of minerals are obtained by identifying and counting a minimum of 300 grains on regularly spaced line traverses that are 2 mm apart.

The identification procedures and reference data on minerals are described in references on sedimentary petrography (Krumbein and Pettijohn, 1938; Durell,
1948; Milner, 1962; Kerr, 1977; Deer et al., 1992) and optical crystallography (Bloss, 1961; Stoiber and Morse, 1972; Shelley, 1978; Klein and Hurlbut, 1985; Drees and Ransom, 1994).

3. Interferences
The sample must be thoroughly mixed because the subsample on the slide is small. If grains are coated with clay or if aggregates of finer material remain in the fraction that is counted, the results may be skewed. Variations in the time or temperature of heating the epoxy may result in either matrix stress or variation in the refractive index of the epoxy. Do not use steel needles or spatulas because magnetic minerals may adhere to steel, resulting in uneven distribution of grains on the slide.

4. Safety
Heat the epoxy in a fume hood. Use caution in handling hot glass slides. Use heat resistant gloves as needed. Immediately wash or remove any epoxy that comes in contact with the skin. Carefully handle slides and cover slips to avoid cuts.

5. Equipment
5.1 Petrographic microscope slides, precleaned, 27 x 46 mm
5.2 Cover slips, glass, 25 x 25 mm
5.3 Hot plate
5.4 Micro-spatula
5.5 Dissecting needle
5.6 Plywood covered with Formica (6 x 8 x 1.25 cm)
5.7 Timer
5.8 Polarizing petrographic microscope
5.9 Tally counter
5.10 Set of 76-mm (3 in) sieves, square-weave phosphor bronze wire cloth, except 300 mesh, which is twilled weave. U.S. series and Tyler Screen Scale equivalent designations are as follows:

<table>
<thead>
<tr>
<th>Sand Size</th>
<th>Opening (mm)</th>
<th>U.S. No.</th>
<th>Tyler Mesh Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very coarse sand (VCS)</td>
<td>1.0</td>
<td>18</td>
<td>16</td>
</tr>
<tr>
<td>Coarse sand (CS)</td>
<td>0.5</td>
<td>35</td>
<td>32</td>
</tr>
<tr>
<td>Medium sand (MS)</td>
<td>0.25</td>
<td>60</td>
<td>60</td>
</tr>
<tr>
<td>Fine sand (FS)</td>
<td>0.105</td>
<td>140</td>
<td>150</td>
</tr>
<tr>
<td>Very fine sand (VFS)</td>
<td>0.047</td>
<td>300</td>
<td>300</td>
</tr>
</tbody>
</table>
5.11 Oven, 110°C

6. Reagents

6.1 Petropoxy 154™ Resin and Curing Agent, Palouse Petro Products, 425 Sand Rd., Palouse, WA 99163

6.2 Index immersion oils

6.3 Reverse osmosis (RO) water, ASTM Type III grade of reagent water

7. Procedure

**Sample Selection and Grain Mount Preparation**

7.1 Refer to the analysis request sheets. Record optical mineralogy requests in the LIMS Report “Optical Priority List.” Note any special instructions by soil scientists. Sample selection depends on the purpose of analysis. In most work, e.g., checks on discontinuities or estimation of degree of weathering in different soil horizons, the study of those fractions that comprise a significant quantitative part of the soil is important. The KSSL convention is to count the most abundant fraction, i.e., coarse silt (CSI), very fine sand (VFS), or fine sand (FS), especially if the fraction is clearly larger. This procedure works well in the establishment of mineralogy families for soil taxonomy (Soil Survey Staff, 2014). This procedure may result in different size fractions being counted for different horizons within a single pedon. If fractions are rather equal in abundance, the VFS is selected because it provides the widest range of information. The KSSL does not count multiple fractions for a single sample, does not count combined fractions, and does not present the data as weighted averages. If it is appropriate to count the same size fraction for each horizon within a pedon or project, such as for a study of soil lithology, this request must be specified by the project coordinator.

7.2 Refer to the laboratory information system (LIMS) report “Optical Priority List” for daily work or check the sand box lids to determine which sands have been fractionated. Sands are fractionated during particle-size distribution analysis (PSDA) (method 3A1a). Fine sand and very fine sand fractions are placed in gelatin capsules and stored in a labeled vial. Coarse silts are stored in aluminum pans. For projects, the LIMS report provides the following:

a. Sample numbers by project

b. Type of count requested (full grain count or glass count)

c. Percent of <2-mm fraction for CSI, VFS, or FS as determined by KSSL PSDA
d. Protocol for glass counts are as follows: Count predominant fraction. If CSI is not the predominant fraction in sample then count the CSI fraction as well, resulting in two glass counts per slide.

7.3 If the particle-size section does not provide a sand and coarse silt separate, derive these fractions by repeated gravity sedimentation at 20 µm and sieving the 20-µm to 2.0-mm material as follows:

7.3.1 Disperse the sample in sodium hexametaphosphate as described in method 7A1a1.

7.3.2 Pour the soil suspension into a 200-mL beaker that has a line marked 5 cm above the bottom.

7.3.3 Add RO water to the beaker up to the 5-cm mark.

7.3.4 Stir the suspension and allow to settle 2.0 min. Use a stopwatch.

7.3.5 Decant and discard the suspension containing the clay and fine silt.

7.3.6 Repeat Steps 7.3.3 to 7.3.5 until the supernatant is clear or reasonably clear.

7.3.7 Transfer the sediment to a drying dish and dry at 105–110 °C.

7.3.8 Sieve the dried sample to isolate the individual fractions.

7.4 Review the PSDA data and select samples. Make grain mounts from the one or two most abundant fractions, preferably from the CSI, VFS, or FS. Record sample numbers and respective PSDA data.

7.5 Mix a small amount of Petropoxy 154™ resin and curing agent (1:10 ratio of resin to curing agent) in a clean graduated plastic beaker that is provided with the reagents.

7.6 Prepare epoxy at least 1 day prior to use and refrigerate until needed.

7.7 Turn on hot plate and allow to equilibrate at 125 °C for ≈1 h.

7.8 Remove mixture from refrigerator at least 40 min prior to use. If the petropoxy crystallizes, gently warm mixture until crystals dissolve.

7.9 At the base of the glass slides, record the grain size fraction (CSI, VFS, FS, etc.) and KSSL LIMS sample number; for example, VFS 35126.

7.10 Obtain sand vials and/or silt dishes. Arrange in an orderly manner. Work with 4–6 slides and samples at a time.

7.11 Remove lids from sand vials and place upside down in front of respective vials. Remove gelatin capsules (VFS or FS) from vial. Rotate capsule to mix contents and place in lid. Stir with a micro-spatula to mix coarse silts.

7.12 Use a small, rounded glass or plastic rod to drop petropoxy mixture on the upper middle of each slide. Use ½ drop petropoxy for CSI, one drop VFS, and 1½ drops for FS.
Use a micro-spatula to add the mixed grains to petropoxy. Use larger amounts for smaller fractions. The analyst’s technique for adding the appropriate amount of petropoxy and making grain counts on prepared slides develops with experience. Use a dissecting needle to slowly and carefully stir the grains into the petropoxy. Avoid introduction of air bubbles. Obvious air bubbles can be popped with the dissecting needle.

Gently place one (check to be certain) cover slip on the petropoxy. Avoid leaving fingerprints. Allow the petropoxy to spread under the cover slip. Center the cover slip at top center of glass microscope slide so that there is a parallel, equidimensional border around the top and sides of slide.

To ensure the uniform distribution of grains and the removal of air bubbles, use a dissecting needle to gently tap or press down cover slip. If necessary, the analyst may need to re-center the cover slip. Be careful not to crack the cover slip.

Align a batch of 4–6 slides in two rows on center of hot plate. Set timer and heat slides at 125 °C for 8 min. Time can be adjusted by experience. As a rule, when epoxy is set, it has cured to yield a refractive index of 1.540. Longer heating may result in a distortion of the optical characteristics of the petropoxy and a refractive index differing from 1.540.

As one batch of slides heats, prepare the next batch. Heat the slides for 8 min and then slide them off the hot plate onto the Formica block. Allow to cool.

Examine the grain mount for quality. The epoxy medium should be isotropic. The presence of anisotropic stress lines around grains under X-Nicols may interfere with observation of optical properties. Remake any unsatisfactory grain mounts. Place satisfactory mounts in a microscope slide file box. Neatly record project number and grain mount positions on interior of box lid. Record box number(s) in the LIMS Sample Disposition files.

Return the petropoxy mixture to the refrigerator in order to extend shelf life of the mixture.

**Observations of Grain Mount**

Record raw grain-count data in a logbook. Most grain counts are made with a 10X magnification ocular and either a 10X (for very fine or fine sand) or 25X (for coarse silt) magnification objective lens.

The first step is to seat the grain mount in the mechanical stage of the microscope and to survey the slide with a low-power magnification power (10X). This step is intended to familiarize the analyst with the grain assemblage and to make a rough estimate of the relative abundance of minerals and other grains.
7.23 Initially, identify the most abundant minerals. They are probably the easiest to identify, and their elimination decreases the number of possibilities to consider in identifying the less common minerals. Furthermore, there are certain likely and unlikely assemblages of minerals, and an awareness of the overall types that are present gives clues to the minor species that may be expected.

7.24 Note the observed minerals by a two-letter code, e.g., QZ for quartz. Refer to the list of mineralogy codes, provided at the end of the mineralogy section (7).

7.25 Make grain counts in horizontal traverses across the grain mount. A 10X magnification objective is appropriate for FS and VFS. A 25X objective is appropriate for CSI.

7.26 To make a grain count, move the slide via the mechanical stage so that the left border of cover slip is in view and in the proximity of, but not in, the upper left corner. Place vertical scale on mechanical stage on an even number, e.g., 72 or 74 mm.

7.27 Set the rotating stage so that the horizontal movement of a grain, via the mechanical stage, parallels the horizontal cross-hair in the ocular.

7.28 List the most abundant grains and associated counter number in logbook. Mineral identification is facilitated by the familiarity with a few striking features and by the process of elimination.

7.29 Set counters to zero. Move the slide laterally one field width at a time. Identify and tally each grain that touches the horizontal cross-hair in each field of view until the right margin of cover slip is in view.

7.30 Translate the slide vertically a distance of 2 mm and run another traverse in the reverse direction.

7.31 Repeat process until the end of the traverse in which 300 grains have been tallied. If there are only a few species, a counting of 300 grains provides a good indication of composition. As the number of species increases, the count should increase within limits of practicability. To count more than 1000 grains is seldom necessary.

7.32 The counting of complete traverses minimizes the effects of nonrandom distribution of grains on the slide. This nonrandom distribution of grains is usually most pronounced near the edges of the cover slip. If the entire slide has been traversed and the total grain count is <300, reverse the direction of vertical translation and count traverses on odd-numbered settings, e.g., 81 or 79 mm.

7.33 Counting isotropic grains only (e.g., volcanic glass) can be done more rapidly using either of the following microscope configurations:

7.33.1 Positioning the polarizer slightly off the extinction or “blackout” position.
7.33.2 With crossed Nicols and a gypsum plate, the outline of the grains is visible; the color of the grain is the same as the epoxy background.

7.34 When the count is complete, enter the raw data (project, sample number, fraction(s), minerals, and counts) into the KSSL LI MS data base.

8. Calculations
Calculations are made by a computer program that facilitates data entry and manipulation. Required inputs are as follows: project number, sample number, grain-size fractions, mineral identification, and number of grains counted per mineral.
Percentage of minerals (frequency per 100 grains) is calculated formula as follows:

8.1 Mineral frequency (%) = \( \frac{\text{Number of grains for a mineral} \times 100}{\text{Total number of grains counted}} \)

9. Report
Report mineral contents to the nearest whole percentage of grains counted. These data are accurate number percentages for the size fraction analyzed but may need to be recomputed to convert to weight percentages (Harris and Zelazny, 1985). Grain counts can deviate significantly from weight percentage due to platy grains and density variations. These data are reported on the mineralogy data page of the primary characterization data set. For each grain size counted, the mineral type and amount are recorded. For example, quartz, 87% of fraction, is recorded as QZ87. The percentage of resistant minerals in each fraction is reported on the KSSL datasheet. Refer to 7C1 for more information on resistant minerals.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References


Optical Analysis (7B)
Platy Grains (7B2)
Magnetic Separation (7B2a)

1. Application
A magnetic separator is used to separate magnetic or paramagnetic minerals from nonmagnetic mineral grains in fine-earth fractions that range from 0.02 to 0.5 mm in size. At the KSSL, the common application of this procedure is to quantify the amount of platy grains (phyllosilicates) in micaceous or paramicaceous soils. Ferrimagnetic magnetic grains, such as magnetite and ilmenite, typically are separated first with a hand magnet. The separator then concentrates grains of paramagnetic minerals, such as biotite and muscovite, away from nonmagnetic minerals, such quartz and feldspar. This method (7B2a) has been often used in combination with static tube separation (obsolete method 7B2b) or froth flotation (obsolete method 7B2c). Grains in each separate can be further analyzed by optical microscopy or x-ray diffraction.

2. Summary of Method
A magnetic separator applies a strong magnetic field along a shallow trough that slopes down from an entry point to an exit point. Grains travel the path
under the force of gravity. The trough is also tilted perpendicular to the travel path. The magnetic field draws paramagnetic grains up the tilt slope. A divider at midslope along the path separates the paramagnetic grains from the nonmagnetic grains. Grains exit the path into separate containers, and the two components are weighed to obtain a relative percentage. Percent platy minerals of specific analyzed fraction are reported (7B2a).

3. Interferences

Some mafic minerals are paramagnetic and exit with the platy grains. The two groups may be separated by the static tube method (obsolete method7B2b). Mafic minerals commonly are heavy minerals and may be separated from platy grains by density separation (7B1a2).

4. Safety

The magnetic field is not a direct biological health hazard. Keep mechanical watches away from the magnetic field. Keep iron/steel objects away from the magnetic field because of the strength of field. The separator will need to be turned off to retrieve them.

5. Equipment

5.1 300-mL fleaker, with a fill line marked at 5 cm above bottom
5.2 Glass stirring rod, with a rubber policeman
5.3 Stop watch
5.4 Sonic probe, Vibra Cell Model V1A, Sonics and Materials, Inc., Danbury, CT
5.5 Magnetic separator; Frantz Isodynamic Magnetic Separator, Model L-1, S.G. Frantz, Inc., Trenton, NJ (fig. 7B2-1)
5.6 Centrifuge tube, plastic, 100-mL
5.7 Beaker, Pyrex, 150–200 mL, marked with a line 5-cm above bottom
5.8 Sieves:
   5.8.1 0.5–2.0 mm = U.S. 35 mesh sieve or Tyler 32 mesh sieve
   5.8.2 0.1–0.5 mm = U.S. 140 mesh sieve or Tyler 150 mesh sieve
   5.8.3 0.05–0.1 mm = U.S. 300 mesh sieve or Tyler 300 mesh sieve
   5.8.4 0.02–0.05 mm = Material that passes through the 300 mesh sieve
5.9 Mechanical reciprocating shaker, 100 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.10 Oven, 110°C

6. Reagents

6.1 Reverse Osmosis (RO) water (ASTM type III grade)
6.2 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate ($\text{NaPO}_3$)$_6$ and 7.94 g of sodium carbonate ($\text{Na}_2\text{CO}_3$) in 1 L of RO water.

7. Procedure

Separation of Target Fraction (0.02 to 2.0 mm)  
(for magnetic separation, static tube separation, and/or froth flotation)

7.1 For magnetic separation alone or in combination with static tube separation, 5 grams of <2-mm soil is sufficient. If froth floatation is used in combination with magnetic separation, use 20 to 30 g of <2.0-mm sample.

7.2 Place sample into a 100-mL centrifuge tube for magnetic or tube methods or in a 300-mL fleaker for froth flotation. Add 10 to 60 mL of dispersing agent, as appropriate for sample weight.

7.3 Fill the centrifuge tube half full or the fleaker to the 5-cm mark with distilled water. Stopper and shake the solution overnight at 100 oscillations min$^{-1}$.

7.4 Transfer contents of the centrifuge tube to a 150 to 200 mL beaker. Add RO to the fleaker to the 5-cm mark, if needed.

7.5 Using the stirring rod, stir the sample for 15 seconds and allow to stand for 2 minutes. Decant and discard the dispersant.

7.6 Repeat the settling process, usually 4 or 5 times, until the dispersant is clear. Sediment is the 0.02–2 mm fraction.
Sonic Cleaning of Sample

7.8 Add approximately 50 mL of RO water to the fleaker or beaker containing the 0.02–2.0 mm sample.

7.9 Turn on and tune the sonic cleaner for optimal performance. Set the % Duty cycle knob to 50; output control knob to 10; and run time to 360 s. Turn on the pulser switch.

7.10 Lower the sonic probe horn 0.5 to 1.0 cm below the water level of the fleaker containing the washed sample and turn on the sonic probe.

7.11 Repeat the decanting process (see separation of working fraction) to remove fines (<0.02 mm) produced by the sonic probe treatment.

7.1 Oven dry the sample at 110 °C.

Particle-Size Separation

7.13 Dry sieve the .02–2 mm fraction cleaned by sonication into the following subfractions: 0.5–2.0, 0.1–0.5, 0.05–0.1, and 0.02–0.05 mm.

7.14 Weigh and record dry weigh of each of the separates.

Procedure Options for Each Size Fraction

7.15 0.5–2 mm fraction: Separate platy grains via magnetic separator only (fig. 7B2-1). This fraction seems unsuited for static tube or froth floatation separation.

7.16 Separate 0.1–0.5, 0.05–0.1, and 0.02–0.05 mm fractions by magnetic separation.

Magnetic Separation

7.17 Turn on the magnetic separator; adjust the machine to the high-high, positive position; and set the amp meter to 1.25 amperes.

7.18 Set the slope to about 20 degrees and the tilt to 15 degrees.

7.19 Introduce sample into the funnel and set the vibrator adjustment knob between 5 and 8. Observe the flow of the grains down the separator trough and adjust the vibrator setting accordingly.

7.20 Repeat the process using the nonmagnetic portion 2 to 4 times or until the nonmagnetic fraction is free of platy minerals.

7.21 Weigh and record the weight of the magnetic fraction as platy grains and the nonmagnetic fraction as residual grains. Calculate amount of platy grains as percent by weight of analyzed fraction.

8. Calculations

8.1 Percent platy grains = \[100 \times (\text{weight of platy grains})/(\text{sample weight})\]
8.2 Percent residual grains = \[\frac{100 \times \text{(weight of residual grains)}}{\text{(sample weight)}}\]

8.3 Recovery = \(\frac{\text{(weight of platy and residual grains)}}{\text{(sample weight)}}\)

9. Report

Report platy grains as a percent of the specific particle size fraction analyzed, oven-dried soil weight.

10. Precision and Accuracy

Precision and accuracy data are not available for this method.

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Ratios and Estimates Related to Optical Analysis (7C)
Total Resistant Minerals (7C1)

The sum of the grain-count percentages of resistant minerals are reported (7C1). For more detailed information on total resistant minerals, refer to the Soil Survey Staff (2014, 2011). Also refer to the list of mineralogy codes for resistant and weatherable minerals, following method 7C1.

References


### MINERALOGY CODES

**Resistant Minerals**

<table>
<thead>
<tr>
<th>Code</th>
<th>Mineral</th>
</tr>
</thead>
<tbody>
<tr>
<td>AE</td>
<td>Anatase</td>
</tr>
<tr>
<td>AN</td>
<td>Andalusite</td>
</tr>
<tr>
<td>BY</td>
<td>Beryl</td>
</tr>
<tr>
<td>CD</td>
<td>Chalcedony (Chert, Flint, Jasper, Agate, Onyx)</td>
</tr>
<tr>
<td>CE</td>
<td>Cobaltite</td>
</tr>
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Glass Count Minerals and Mineraloids

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<th>Organic Origin Grains&lt;sup&gt;4&lt;/sup&gt;</th>
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<tr>
<td>BG=Basic Glass</td>
<td>DI=Diatoms</td>
<td>OT=Other</td>
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<tr>
<td>FG=Glass-Coated Feldspar</td>
<td>PO=Plant Opal</td>
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<tr>
<td>GA=Glass Aggregates</td>
<td>SS=Sponge Spicule</td>
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<tr>
<td>GC=Glass-Coated Grain</td>
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<td>GM=Glassy Materials</td>
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<tr>
<td>GS=Glass</td>
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<td></td>
</tr>
<tr>
<td>HG=Glass-Coated Hornblende</td>
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<tr>
<td>OG=Glass-Coated Opaque</td>
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<td>PM=Pumice</td>
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<tr>
<td>QG=Glass-Coated Quartz</td>
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</table>

<sup>1</sup>Minerals not included as “weatherable minerals” as defined by “Keys to Soil Taxonomy” (Soil Survey Staff, 2014): “The intent is to include... only those weatherable minerals that are unstable in a humid climate compared to other minerals such as quartz and 1:1 lattice clays, but are more resistant to weathering than calcite.” This group of minerals is not part of the calculation for percent resistant minerals used in the siliceous family mineralogy class or percent weatherable minerals used as criteria for oxic horizon but are included in the calculation of “total resistant minerals” on the Kellogg Soil Survey Laboratory (KSSL) mineralogy data sheet. Therefore, the value on the data sheet should be recalculated for strict use in “Soil Taxonomy” criteria if these minerals (e.g., calcite) are in the grain count of a selected horizon.

<sup>2</sup>Minerals on this list are identified during the “glass count” procedure by the KSSL during the quantification of particle size-separates in the sand-silt fraction. Minerals in the “OT” category are other weatherable or resistant minerals that would be quantified during a “full grain count.”

<sup>3</sup>Minerals and mineraloids in this column are all considered weatherable according to the KSSL and are defined in the “Keys to Soil Taxonomy,” Twelfth Edition, 2014, as being “volcanic glass.” The percentages of these minerals are summed as “volcanic glass” and used in the criteria for andic soil properties, subgroups with the formative element “vitr(i)”, families with “ashy” substitutes for particle-size class, and the glass mineralogy class in “Soil Taxonomy.”

<sup>4</sup>Mineraloids included in this list are regarded as resistant minerals according to the KSSL and included in the calculation of “total resistant minerals” as shown on the KSSL mineralogy data sheet.
OBSELETE METHODS OF ANALYSIS

This section describes the methods that are no longer used at the Kellogg Soil Survey Laboratory (KSSL). These methods were described in earlier versions of Soil Survey Investigations Report (SSIR) No. 42 (1989, 1992, 1996, and 2004) and in SSIR No. 1, “Procedures for Collecting Soil Samples and Methods of Analysis for Soil Survey” (1972, 1982, and 1984). Some of these procedures are in the old format. Information is not available to describe some of these obsolete procedures in the same detail as used to describe the current methods in the laboratory. Also included in this manual are examples of earlier versions of the KSSL data sheet (appendix fig. 1) and pedon description (appendix fig. 2).

Since the publication of SSIR No. 42 (1996), the number and kinds of methods performed at the KSSL have increased significantly, resulting in a re-structuring of the laboratory method codes. This re-structuring is reflected in this version of SSIR No. 42 (2014) as well as in SSIR No. 42, version 4.0 (2004). Some of the methods described in SSIR No. 1, 1972, 1982, and 1984, as well as in SSIR No. 42, 1989, 1992, and 1996, carry the old method codes which may not necessarily be the same as current method codes. These older method codes have a maximum of four characters, e.g., 6A2b.

It is important to document and archive these obsolete methods because many older SSIRs and scientific publications report these methods. The intent of this documentation is to provide a historical linkage for the KSSL core methods. The following section of this manual documents the obsolete methods and is divided into parts as follows:

Part III: SSIR No. 42, Soil Survey Laboratory Methods Manual, Versions 1.0 and 2.0 (1989, 1992, respectively)
# INDEX TO OBSOLETE METHODS

Obsolete Methods that have the New Method Code Structure

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SAMPLE COLLECTION AND PREPARATION (1)

Laboratory Sample Collection and Preparation (1B)
Soils (1B1)
  Soil Bulk Sample Preparation (1B1b)
    Air-Dry Preparation (1B1b2)
      <2-mm Fraction (1B1b2b)
        Presence of Carbonates (1B1b2b5)

7.32 Use a sub-sample of the ADOD sample (method 1B1b2b4) and check for the presence of carbonates. Reference samples (knowns) are available for comparisons. Place 1 g of the air-dry fine-earth fraction in porcelain spot plate, add reverse osmosis water, and stir to remove entrapped air. Add 1 N HCl to soil, observe amount of effervescence, and record as follows:

None.—No visual effervescence.

Very Slight.—Bubbles rise at a few points in the sample and consistently appear at the same point in either a steady stream of tiny bubbles or in a slower stream of larger bubbles. Do not mistake trapped air bubbles for a positive test. Generally, these air bubbles appear immediately after the addition of 1 N HCl.

Slight.—More small bubbles, and possibly a few larger bubbles, appear throughout the sample than with a very slight reaction.

Strong.—More large bubbles are evident than with a slight reaction. Often the reaction is violent at first and then quickly decreases to a reaction that produces many small bubbles.

Violent.—The sample effervescences violently. Many large bubbles appear to burst from the spot plate.
SOIL AND WATER CHEMICAL EXTRACTIONS AND ANALYSES (4)

Ion Exchange and Extractable Cations (4B)

BaCl₂-Triethanolamine, pH 8.2 Extraction (4B2)
   Automatic Extractor (4B2a)
   Automatic Titrator (4B2a1)
   Back Titration with HCl (4B2a1a)
   Extractable Acidity (4B2a1a1)
       Air-Dry or Field-Moist, <2 mm (4B2a1a1a-b1)

1. Application
   The extractable acidity is the acidity released from the soil by a barium chloride-triethanolamine (BaCl₂-TEA) solution buffered at pH 8.2 and includes all the acidity generated by replacement of the H and Al from permanent and pH dependent exchange sites. Extractable acidity may be measured at any pH, and a variety of methods have been used to measure it. The Soil Conservation Service adopted a pH of 8.2 because it approximates the calculated pH of a soil containing free CaCO₃ in equilibrium with the normal CO₂ content (0.03%) of the atmosphere. A pH of 8.2 also closely corresponds to the pH of complete neutralization of soil hydroxy-Al compounds. Although other pH values are valid for some types of soils, and the BaCl₂-TEA, pH 8.2 method (4B2a1a1) may not always accurately reflect the nature of soils as they occur in the environment, this method has become a standard reference to which other methods are compared.

2. Summary of Method
   A soil sample is leached with a BaCl₂-TEA solution buffered at pH 8.2. Sample is allowed to stand overnight and extracted using a mechanical vacuum extractor (Holmgren et al., 1977). The extract is back-titrated with HCl. The difference between a blank and the extract is the extractable acidity. Extractable acidity is reported in meq 100 g⁻¹ soil or (cmol (+) kg⁻¹).

3. Interferences
   No significant interferences are known to exist with this method. However, for some very acid soils, the buffer capacity of the BaCl₂-TEA solution may be exceeded.

4. Safety
   Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids in a fume hood. Thoroughly wash hands after handling reagents. Use the safety showers.
and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Sampletek, Mavco Industries, Lincoln, NE
5.3 Pipettes or dispenser, adjustable volume to 20 mL
5.4 Titration beakers, 250-mL, plastic, Metrohm Ltd., Brinkmann Instruments Inc.
5.5 Automatic titrator, with control unit, sample changer, and dispenser, Metrohm Ltd., Brinkmann Instruments, Inc.
5.6 Combination pH-reference electrode, Metrohm Ltd., Brinkmann Instruments, Inc.
5.7 Computer, with Titrino Workcell software, Metrohm Ltd., Brinkmann Instruments, Inc., and printer
5.8 Titration beakers, 250-mL, plastic, Metrohm Ltd., Brinkmann Instruments Inc.
5.9 Tubes, 60-mL, polypropylene, for extraction, with 0.45-µm filter
5.10 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm, (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels

6. Reagents

6.1 Reverse osmosis deionized (RODI) water
6.2 Reverse osmosis (RO) water
6.3 Hydrochloric acid (HCl), concentrated, 12 N
6.4 HCl, 0.13 N, standardized. Dilute 193 mL of concentrated HCl to 16-L volume with RODI water.
6.5 Buffer solution [0.5 N BaCl₂, 0.2 N Triethanolamine (TEA), pH 8.2]. Dissolve 977 g of BaCl₂•2H₂O in 8 L of RODI water. Dissolve 477 g of TEA in 4 L of RODI water. Mix two solutions and bring to nearly 16-L volume with RODI water. Adjust to pH 8.2 with ≈33 mL of concentrated HCl or barium hydroxide. Bring to 16-L volume with RODI water.
6.6 Replacement solution. Dissolve 977 g of BaCl₂•2H₂O in 8 L of RODI water. Add 80 mL of buffer solution and dilute 16-L volume with RODI water.

7. Procedure

Extraction of Acidity

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to the nearest mg and place in a labeled extraction (ET₁). If sample is moist, weigh enough soil
to achieve ≈2.5 g of air-dry soil. Prepare at least one reagent blank (no sample in syringe) and one quality control check sample per 24 samples.

7.2 Place labeled ET on extractor and connect to corresponding extraction tube \((ET_{\text{Acidity}})\) with rubber tubing.

7.3 Use a dispenser to add 20.00 mL of \(\text{BaCl}_2\)-TEA solution to the \(ET_1\). During the addition, wash the sides of the tube and wet the sample. For organic soils, shaking, swirling, or stirring may be required to wet the sample.

7.4 Let \(ET_1\) tube stand overnight.

7.5 Set the extractor for a 30-min extraction rate. Extract solution to a 0.5- to 1.0-cm height above the sample. Turn off the extractor. Do not allow the sample to become dry.

7.6 Use a dispenser to add 20.00 mL of replacement solution to \(ET_1\). Extract the sample at 30-min rate, pulling the solution almost completely through the sample.

7.7 Add a second 20.00-mL aliquot of replacement solution to \(ET_1\). Extract at 30-min rate until all the solution has been drawn through the sample.

7.8 Carefully remove \(ET_{\text{Acidity}}\). Leave rubber tubing on the \(ET_1\).

**Titration of \(\text{BaCl}_2\)-TEA Extract**

7.9 Transfer the \(\text{BaCl}_2\)-TEA extract from the \(ET_{\text{Acidity}}\) to a 250-mL polyethylene titration beaker.

7.10 Add 100 mL of RO water to the beaker. The solution is ready to be titrated.

7.11 Refer to manufacturer’s manual for operation of the automatic titrator.

7.12 Calibrate automatic titrator with 9.18, 7.00, and 4.00 pH buffers. Set-up the automatic titrator to set end point titration mode. The “Set” pH parameters are listed as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(E_{p1})</td>
<td>pH 4.60</td>
</tr>
<tr>
<td>Dynamic change pH</td>
<td>1.5 units</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s(^{-1})</td>
</tr>
<tr>
<td>Time delay</td>
<td>10 s</td>
</tr>
<tr>
<td>Drift</td>
<td>0.4 mV s(^{-1})</td>
</tr>
<tr>
<td>Temp</td>
<td>25 °C</td>
</tr>
<tr>
<td>Stop volume</td>
<td>75 mL</td>
</tr>
</tbody>
</table>

7.13 If pre-titration pH is 0.3 units lower than the average pH of the blanks, re-run using a 0.25 g sample.
7.14 Record the titer to the nearest 0.01 mL. Record the normality of the HCl solution. Average the titer of the reagent blanks and record.

8. Calculations

Extractable acidity (meq 100 g⁻¹) = [(B − T) x N x R] / C x 100

where:
B = Average reagent blank titer (mL)
T = Sample titer (mL)
N = Normality of HCl
C = Sample Weight (g)
100 = Conversion factor (100-g basis)
R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)

9. Report
Report extractable acidity to the nearest 0.1 meq 100 g⁻¹ (cmol (+) kg⁻¹).

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

Soil Test Analyses (4D)
Aqueous Extraction (4D2)
Single-Point Extraction (4D2a)
1:10, 30 min (4D2a1)
Flow Injection, Automated Ion Analyzer (4D2a1b)
Phosphorus (4D2a1b1)
Air-Dry or Field-Moist, <2 mm (4D2a1b1a-b1)

1. Application
Phosphorus occurs in soil in both the solution and solid phase. These forms are well documented, but questions still remain concerning the exact nature of the constituents and ionic forms found in water, soils, and sediments (National Research Council, 1993). These forms influence P availability in relation to root absorption and plant growth; runoff and water quality problems; and P loadings.
Water soluble P has been defined as P measured in water, dilute salt extracts (e.g., 0.01 $M \text{CaCl}_2$), displaced soil solutions, or saturation paste extracts (Olsen and Sommers, 1982). Even though the water soluble fraction principally consists of inorganic orthophosphate ions, there is evidence that some organic P is also included (Rigler, 1968).

The water or dilute salt extracts represent an attempt to approximate the soil solution P concentration. As an index of P availability, the objectives of this method are (1) to determine the P concentration level in the soil extract that limits plant growth (Olsen and Sommers, 1982) and (2) to determine the composition of the soil solution so that the chemical environment of the plant roots may be defined in quantitative terms (Adams, 1974). The sum of water soluble P and pH 3 extractable P has also been defined as the available P in runoff (Jackson, 1958).

2. Summary of Method

A 2.5-g sample of <2-mm, air-dry soil is mechanically shaken for 30 min in 25-mL of reverse osmosis deionized water (RODI). The sample is then centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained.

A flow injection automated ion analyzer is used to measure the orthophosphate ion ($\text{PO}_4^{3-}$). This ion reacts with ammonium molybdate and antimony potassium tartrate under acidic conditions to form a complex. This complex is reduced with ascorbic acid to form a blue complex which absorbs light at 880 nm. Absorbance is proportional to the concentration of $\text{PO}_4^{3-}$ in the sample. Data are reported as mg P kg$^{-1}$ soil (4D2a1b1).

3. Interferences

Silica forms a pale blue complex which also absorbs at 880 nm. This interference is generally insignificant as a Si concentration of approximately 30 mg SiO$_2$ L$^{-1}$ would be required to produce a 0.005 mg P L$^{-1}$ positive error in orthophosphate (LACHAT, 1993).

Glassware contamination is a problem in low-level P determinations. Glassware should be washed with 1:1 HCl and rinsed with deionized water. Commercial detergents should rarely be needed but, if they are used, use P-free preparation for lab glassware (LACHAT, 1993).

Concentrations of ferric ion >50 mg L$^{-1}$ will cause a negative error due to competition with the complex for the reducing agent ascorbic acid. Samples high in Fe can be pretreated with sodium bisulfite to eliminate this interference. Treatment with bisulfite will also remove the interference due to arsenates (LACHAT, 1993).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). When preparing reagents,
exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated H$_2$SO$_4$ and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Centrifuge tubes, 50-mL, polyethylene
5.3 Mechanical reciprocating shaker, 200 oscillations min$^{-1}$, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Centrifuge, Centra GP-8, Thermo IEC, Needham Heights, MA
5.5 Filter paper, Whatman No. 42, 150 mm
5.6 Funnel, 60° angle, long stem, 50-mm diameter
5.7 Volumetric flasks, 1-L and 250-mL
5.8 Bottles, plastic, dark, 1-L
5.9 Cups, plastic
5.10 Flow Injection Automated Ion Analyzer, QuikChem AE, LACHAT Instruments, Milwaukee, WI, with computer and printer
5.11 XYZ Sampler, LACHAT Instruments, Milwaukee, WI
5.12 Reagent Pump, LACHAT Instruments, Milwaukee, WI
5.13 Automated Dilution Station, LACHAT Instruments, Milwaukee, WI
5.14 Sample Processing Module (SPM) or channel, QuikChem Method (10-115-01-1-A, orthophosphate in waters, 0.01 to 2.0 mg P L$^{-1}$), LACHAT Instruments, Milwaukee, WI
5.15 Computer, with QuikChem software, LACHAT Instruments, Milwaukee, WI, and printer
5.16 Pipettes, electronic digital, 2500 µL and 10 mL, with tips, 2500 µL and 10 mL
5.17 Vials, plastic, 25-mL (standards)
5.18 Culture tubes, glass, 10-mL (samples)
5.19 Dispenser, 30 mL or 10 mL

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Helium, compressed gas
6.3 Sulfuric acid (H$_2$SO$_4$), concentrated, 36 N, trace pure grade
6.4 Stock ammonium molybdate solution. In 1-L volumetric flask dissolve 40.0 g ammonium molybdate tetrahydrate \( \left[ \text{(NH}_4\right)_{6}\text{Mo}_7\text{O}_{24}\cdot4\text{H}_2\text{O} \] in approximately 800 mL RODI water. Dilute to the mark with RODI water and invert to thoroughly mix. Stir for 4 h. Store in plastic and refrigerate.

6.5 Stock antimony potassium tartrate solution. In 1-L flask, dissolve 3.0 g antimony potassium tartrate (potassium antimony tartrate hemihydrate \( \text{K(SbO)}\text{C}_4\text{H}_4\text{O}\cdot\frac{1}{2}\text{H}_2\text{O} \] in approximately 800 mL RODI water. Dilute to the mark and invert to thoroughly mix. Store in dark bottle and refrigerate.

6.6 Molybdate color reagent. In 1 L volumetric flask, add about 500 mL RODI water, then add 35.0 mL concentrated sulfuric acid (CAUTION: The solution will get very hot!). Swirl to mix. When it can be comfortably handled, add 72 mL stock antimony potassium tartrate solution and 213 mL stock ammonium molybdate solution. Dilute to volume with RODI water and invert three times. Degas with helium ≈5 min.

6.7 Ascorbic acid reducing solution. In 1-L volumetric flask, dissolve 60.0 g ascorbic acid in about 700 mL RODI water. Dilute to volume with RODI water and invert three times. Degas with helium ≈5 min. Optional: After dilution to volume and degassing, dissolve 1.0 g dodecyl sulfate \( \text{CH}_3\text{(CH}_2\right)_{11}\text{OSO}_3\text{Na} \). Prepare fresh weekly.

6.8 Sodium hydroxide-EDTA rinse. Dissolve 65 g sodium hydroxide (NaOH) and 6 g tetrasodium ethylenediamine tetraacetic acid (Na\textsubscript{4}EDTA) in 1.0 L RODI water.

6.9 Stock standard P solution (SSPS), 100.0 mg P L\textsuperscript{-1} (ppm). In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate \( \text{KH}_2\text{PO}_4 \) that has been dried for 2 h at 110 °C in about 800 mL RODI water. Dilute to volume and invert to thoroughly mix. Do not degas. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.10 Working stock standard P solution (WSSPS), 10.0 mg P L\textsuperscript{-1}. In a 1-L volumetric flask, dilute 100.0 mL SSPS to mark with RODI water. Invert to thoroughly mix. Make fresh daily.

6.11 Standard P calibration solutions (SPCS) or working standards, 2.0, 1.0, 0.5, 0.20, 0.05, 0.01, and 0.00 mg P L\textsuperscript{-1}. Make fresh daily. To seven 250-mL volumetric flasks add as follows:

6.11.1 2.0 mg P L\textsuperscript{-1}=50.0 mL WSSPS
6.11.2 1.0 mg P L\textsuperscript{-1}=25.0 mL WSSPS
6.11.3 0.5 mg P L\textsuperscript{-1}=12.5 mL WSSPS
6.11.4 0.20 mg P L\textsuperscript{-1}=5.0 mL WSSPS
6.11.5 0.05 mg P L\textsuperscript{-1}=1.25 mL WSSPS
6.11.6 0.01 mg P L\textsuperscript{-1}=0.25 mL WSSPS
6.11.7 0.00 mg P L\(^{-1}\) = 0 mL WSSPS (blank)
Dilute each SPCS to the mark with RODI water and invert to thoroughly mix. Do not degas.

7. Procedure

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Add 25.0 mL of RODI water to sample. Transfer the sample to the shaker. Shake for 30 min at 200 oscillations min\(^{-1}\) at room temperature (20 °C ±2 °C).

7.3 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min decant, filter, and collect extract in receiving cup.

7.4 Transfer sample extracts into culture tubes and place in XYZ sample trays marked “Samples”. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 24 h.

7.5 Transfer SPCS standards into plastic vials and place in descending order in XYZ sample trays marked “Standards”.

7.6 Refer to the operating and software reference manuals for LACHAT set-up and operation.

7.7 Turn main power switch “ON” and allow 15 min for heater module to warm up to 37 °C.

7.8 On reagent pump, set speed to 35. Pump RODI water through system for 20 min.

7.9 On computer main menu, select “Methods” and then “Analysis Select and Download.” On method list, select water soluble P method. System unit receives the downloaded method and initializes it.

7.10 Pump reagents into manifold. Continue this step and observe baseline. A good baseline needs to be smooth and at zero absorbance. Scatter is indicative of air bubbles and irregular reagent flow. Also observe for any back-pressure in manifold tubing.

7.11 On computer main menu, select “Samples,” “Tray Definition and Submit,” and then “Edit” to create new sample tray followed by “Submit” to run new sample tray.

7.12 Method parameters specific to water soluble P are defined within the “Method Definition” menu. Some of these parameters have been modified from the QuikChem Method 10-115-01-1-A, orthophosphate in waters (U.S. Environmental Protection Agency, 1983; LACHAT Instruments, 1993; U.S. Department of Interior, Geological Survey 1993). Modifications are
primarily related to the criteria and strategies for calibration standards and to injection timing.

7.13 Some of the method parameters as they relate to calibration standards are as follows:

**7.13.1** There are 7 calibration standards (2.00, 1.00, 0.50, 0.20, 0.05, 0.01, and 0.00 mg P L\(^{-1}\)) with a data format of ####.###, i.e., data rounded to 3 places.

**7.13.2** The segments/boundaries for the calibration standards are A–D (2.0 to 0.20 mg P L\(^{-1}\)) and D–G (0.20 to 0.00 mg P L\(^{-1}\)).

**7.13.3** The protocol (replications) for the calibration standards is as follows: AA BB CCC DDDD EEEE FFFF GG

**7.13.4** The check standard is 2.0 mg P L\(^{-1}\). Maximum number of consecutive trays between check standard is one; maximum number of consecutive samples between check standard is 60; and maximum elapse time between check standards is 2 h.

**7.13.5** Calibration strategy for segments A–D and D–G are normal. The normal strategy requires a minimum correlation coefficient of 0.99. Both segments require a maximum standard deviation in slope of 50%. A calibration passes only when both criteria are met. Strategies are user designated. In addition, calibration strategies are based on the full chord. Chord 0 is full chord, and chord 1–5 are sections of peak from start of peak to end of peak.

**7.13.6** The instrument is calibrated with the injection of SPCS. The data system then associates the concentrations with the instrument response for each SPCS.

7.14 Method parameters in relation to timing are as follows:

**7.14.1** Cycle period: 40 s

**7.14.2** Inject to start of peak period: 18 s. To see if peaks are being timed correctly, scan across correlation coefficients for all chords 1–5. The most peak area should be between chords 2–4 with the most signal-to-noise ratio in chords 1 and 5.

**7.14.3** Inject to end of peak period: 52 s

**7.14.4** Automatic timing, where standard assumptions are in effect; no manual timing

7.15 Method parameters in relation to data presentation are as follows:

**7.15.1** Top Scale Response: 0.50 abs

**7.15.2** Bottom Scale Response: 0.00 abs

7.16 Method parameters in relation to data results are as follows:
Set Default Chord to 3. This change must be made to both the sample and the calibration RDF’s.

Refer to the “Method Definition” for water soluble P for other method parameters not discussed here.

Run samples using calibration curve. Sample concentration is calculated from the regression equations. Report results to the nearest 0.01 unit for the sample extract and each SPCS.

If samples are outside calibration range, dilute samples with extracting solution and re-analyze.

Upon completion of run, place the transmission lines into the NaOH-EDTA solution. Pump the solution for approximately 5 min to remove any precipitated reaction products. Then place these lines in RODI water and pump for an additional 5 min and proceed with the normal “Shut-down” procedure.

8. Calculations
Convert extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:

\[
\text{Soil P (mg kg}^{-1}) = \frac{(A \times B \times C \times R \times 1000)}{E}
\]

where:
A = Sample extract reading (mg L\(^{-1}\))
B = Extract volume (L)
D = Dilution, if performed
R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (procedure 3D2)
1000 = Conversion factor to kg-basis
E = Sample weight (g)

9. Report
Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
Soil Test Analyses (4D)
Bray P-1 Extraction (4D3)
   Flow-Injection, Automated Ion-Analyzer (4D3b)
      Phosphorus (4D3b1)
         Air-Dry or Field-Moist, <2 mm (4D3b1a-b1)

1. Application
   The Bray P-1 procedure is widely used as an index of available P in the soil. Bray and Krutz (1945) originally designed the Bray P-1 extractant to selectively remove a portion of the adsorbed form of P with the weak, acidified ammonium fluoride solution. Adsorbed phosphorus is in the anion form adsorbed by different charged surface functional groups that have varying degrees of adsorption affinity. In general, this method has been most successful on acid soils (Olsen and Sommers, 1982). The acid solubilizes calcium and aluminum phosphates and partially extracts iron phosphates compounds. The NH₄F complexes the aluminum in solution and limits re-adsorption of P on iron oxides (Kuo, 1996). The Bray P-1 has limited ability to extract P in calcareous soils due to the neutralization of the dilute acid by carbonates. For most soils, Bray P-1 and Mehlich No. 3 are nearly comparable in their abilities to extract native P but exceed Olsen sodium-bicarbonate method by two- to three-fold, indicating that predictive models for Bray
P-1, Mehlich No. 3, and Olsen sodium-bicarbonate are closely associated with pH buffering of extractant (acid versus alkaline) (Burt et al., 2002).

2. Summary of Method

A 2.5-g soil sample is mechanically shaken for 15 min in 25-mL of Bray P-1 extracting solution. The sample is then centrifuged until solution is free of soil mineral particles and then filtered until clear extracts are obtained.

A flow injection automated ion analyzer is used to measure the orthophosphate ion ($PO_{4}^{3-}$). This ion reacts with ammonium molybdate and antimony potassium tartrate under acidic conditions to form a complex. This complex is reduced with ascorbic acid to form a blue complex which absorbs light at 660 nm. Absorbance is proportional to the concentration of $PO_{4}^{3-}$ in the sample. Data are reported as mg P kg$^{-1}$ soil (4D3b1).

3. Interferences

Silica forms a pale blue complex which also absorbs at 660 nm. This interference is generally insignificant as a silica concentration of approximately 4000 mg L$^{-1}$ would be required to produce a 1 mg L$^{-1}$ positive error in orthophosphate (LACHAT Instruments, 1989).

Concentrations of ferric iron greater than 50 mg L$^{-1}$ will cause a negative error due to competition with the complex for the reducing agent ascorbic acid. Samples high in iron can be pretreated with sodium bisulfite to eliminate this interference. Treatment with bisulfite will also remove the interference due to arsenates (LACHAT Instruments, 1989).

The determination of phosphorus is sensitive to variations in acid concentrations in the sample since there is no buffer. With increasing acidity, the sensitivity of the method is reduced. Samples, standards, and blanks should be prepared in a similar matrix.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated HCl, $NH_{4}F$, and $H_{2}SO_{4}$ to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity

5.2 Centrifuge tubes, 50-mL, polyethylene
5.3 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.4 Centrifuge, Centra GP-8, Thermo IEC, Needham Heights, MA
5.5 Filter paper, Whatman No. 42, 150 mm
5.6 Funnel, 60° angle, long stem, 50-mm diameter
5.7 Volumetric flasks, 1-L and 250-mL
5.8 Bottles, plastic, dark, 1-L
5.9 Cups, plastic
5.10 Dispenser, 30 mL or 10 mL
5.11 Flow Injection Automated Ion Analyzer, QuikChem AE, LACHAT Instruments, Milwaukee, WI, with computer and printer
5.12 XYZ Sampler, LACHAT Instruments, Milwaukee, WI
5.13 Reagent Pump, LACHAT Instruments, Milwaukee, WI
5.14 Automated Dilution Station, LACHAT Instruments, Milwaukee, WI
5.15 Sample Processing Module (SPM) or channel, QuikChem Method (12-115-01-1-A, orthophosphate in waters, 0.4 to 20 mg P L⁻¹), LACHAT Instruments, Milwaukee, WI
5.16 Computer, with QuikChem software, LACHAT Instruments, Milwaukee, WI, and printer
5.17 Pipettes, electronic digital, 2500 µL and 10 mL, with tips 2500 µL and 10 mL
5.18 Vials, plastic, 25-mL (standards)
5.19 Culture tubes, glass, 10-mL (samples)

6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Helium, compressed gas
6.3 Hydrochloric acid (HCl), concentrated, 12 N, trace pure grade
6.4 Sulfuric acid (H₂SO₄), concentrated, 36 N, trace pure grade
6.5 HCl, 1 N. Carefully add 83.33 mL of concentrated HCl to RODI water and dilute to 1-L volume.
6.6 Bray No. 1 Extracting Solution. 0.025 M HCl, and 0.03 M NH₄F. Dissolve 8.88 g of NH₄F in 4 L RODI water. Add 200 mL of 1.0 N HCl and dilute to 8 L with RODI water. The solution pH should be 2.6 ±0.05. Store in a polyethylene bottle.
6.7 Stock ammonium molybdate solution. In 1-L volumetric flask dissolve 40.0 g ammonium molybdate tetrahydrate [(NH₄)₆Mo₇O₂₄•4H₂O] in approximately
800 mL RODI water. Dilute to the mark with RODI water and invert to thoroughly mix. Stir for 4 h. Store in plastic and refrigerate.

6.8 Stock antimony potassium tartrate solution. In 1-L flask, dissolve 3.0 g antimony potassium tartrate (potassium antimony tartrate hemihydrate \([K(SbO)C_4H_4O_6\cdot\frac{1}{2}H_2O]\) in approximately 800 mL RODI water. Dilute to the mark and invert to thoroughly mix. Store in dark bottle and refrigerate.

6.9 Molybdate color reagent. In 1 L volumetric flask, add 72 mL stock antimony potassium tartrate solution and 213 mL stock ammonium molybdate solution. Dilute to volume with RODI water and invert to thoroughly mix. Degas with helium ≈15 min.

6.10 Ascorbic acid reducing solution. In 1-L volumetric flask, dissolve 60.0 g ascorbic acid in about 700 mL RODI. Dilute to volume with RODI water and invert to thoroughly mix. Degas with helium ≈5 min. After dilution to volume and degassing, dissolve 1.0 g dodecyl sulfate \((CH_3(CH_2)_11OSO_3Na)\). Prepare fresh daily.

6.11 0.8 \(M \) \(H_2SO_4\) Carrier. To 1-L container, add 44.4 mL concentrated \(H_2SO_4\) and bring volume with RODI water. (CAUTION: The solution will get very hot!) Invert to thoroughly mix. Degas with helium ≈5 min.

6.12 Sodium hydroxide-EDTA rinse. Dissolve 65 g sodium hydroxide (NaOH) and 6 g tetrasodium ethylenediamine tetraacetic acid \((Na_4EDTA)\) in 1.0 L RODI water.

6.13 Working stock standard P solution (WSSPS), 100.0 mg P L\(^{-1}\). In a 1-L volumetric flask, dissolve 0.4394 g primary standard grade anhydrous potassium dihydrogen phosphate \((KH_2PO_4)\) that has been dried for 2 h at 110 °C in about 800 mL extracting solution. Dilute to 1-L volume with extracting solution and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.

6.14 Standard P calibration solutions (SPCS) or working standards, 20.00, 12.00, 4.00, 0.800, and 0.000 mg P L\(^{-1}\) as PO\(_4^{3-}\). Make fresh weekly. Store in refrigerator. Allow to equilibrate to room temperature before use. To five 250-mL volumetric flasks add as follows:

6.14.1 20.00 mg P L\(^{-1}\)=50 mL WSSPS
6.14.2 12.00 mg P L\(^{-1}\)=30 mL WSSPS
6.14.3 4.00 mg P L\(^{-1}\)=10 ml WSSPS
6.14.4 0.80 mg P L\(^{-1}\)=2 mL WSSPS
6.14.5 0.00 mg P L\(^{-1}\)=0 mL WSSPS (blank)

Dilute each SPCS to the mark with extracting solution and invert to thoroughly mix. Do not degas.
7. Procedure

7.1 Weigh 2.5 g of <2-mm or fine-grind, air-dry soil to nearest mg on an electronic balance and place into a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈2.5 g of air-dry soil.

7.2 Dispense 25.0 mL of extracting solution to tube.

7.3 Transfer the sample to the shaker. Shake for 15 min at 200 oscillations min⁻¹ at room temperature (20 °C ±2 °C).

7.4 Remove the sample from the shaker. Centrifuge at 2000 rpm for 10 min, decant, filter, and collect extract in receiving cup. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.

7.5 Transfer sample extracts into culture tubes and place in XYZ sample trays marked “Samples”.

7.6 Transfer SPCS standards into plastic vials and place in descending order in XYZ sample trays marked “Standards”.

7.7 Refer to the operating and software reference manuals for LACHAT set-up and operation.

7.8 Turn main power switch “ON” and allow 15 min for heater module to warm up to 60 °C.

7.9 On reagent pump, set speed to 35.

7.10 On computer main menu, select “Methods” and then “Analysis Select and Download”. On method list, select Bray P-1 Method. System unit receives the downloaded method and initializes it.

7.11 Pump reagents into appropriate chambers of the manifold. Continue this step and observe baseline. A good baseline needs to be smooth and at zero absorbance. Scatter is indicative of air bubbles and irregular reagent flow. Also observe for any back-pressure in manifold tubing.

7.12 On computer main menu, select “Samples,” “Tray Definition and Submit,” and then “Edit” to create new sample tray followed by “Submit” to run new sample tray.

7.13 Method parameters specific to Bray P-1 are defined within the “Method Definition” menu. Some of these parameters have been modified from the QuikChem Method 12-115-01-1-A, orthophosphate in soils (U.S. Environmental Protection Agency, 1983; LACHAT Instruments, 1989; U.S. Department of Interior, Geological Survey, 1993). Modifications are primarily related to the criteria and strategies for calibration standards and to injection timing.

7.14 Some of the method parameters as they relate to calibration standards are as follows:
7.14.1 There are 5 calibration standards (20.00, 12.00, 4.00, 0.80, and 0.00 mg P L\(^{-1}\)) with a data format of ####.###, i.e., data rounded to 3 places.

7.14.2 The segments/boundaries for the calibration standards are A–C (20.0 to 4.0 mg P L\(^{-1}\)); C–E (4.0 to 0.0 mg P L\(^{-1}\)).

7.14.3 The protocol (replications) for the calibration standards is as follows: AA BB CC DDD EEE.

7.14.4 The check standard is 20.0 mg P L\(^{-1}\). Maximum number of consecutive trays between check standard is one; maximum number of consecutive samples between check standard is 60; and maximum elapse time between check standards is 2 h.

7.14.5 Calibration strategy for segments A–C and C–E are normal. The normal strategy requires a minimum correlation coefficient of 0.99. Both segments require a maximum standard deviation in slope of 50%. A calibration passes only when both criteria are met. Strategies are user designated. In addition, calibration strategies are based on the full chord. Chord 0 is full chord, and chord 1–5 are sections of peak from start of peak to end of peak.

7.14.6 The instrument is calibrated with the injection of SPCS. The data system then associates the concentrations with the instrument response for each SPCS.

7.15 Method parameters in relation to timing are as follows:

7.15.1 Cycle period: 40 s

7.15.2 Inject to start of peak period: 18 s. To see if peaks are being timed correctly, scan across correlation coefficients for all chords 1–5. The most peak area should be between chords 2–4 with the most signal-to-noise ratio in chords 1 and 5.

7.15.3 Inject to end of peak period: 46 s

7.15.4 Automatic timing, where standard assumptions are in effect. Manual timing may be helpful in this method.

7.16 Method parameters in relation to data presentation are as follows:

7.16.1 Top Scale Response: 0.50 abs

7.16.2 Bottom Scale Response: 0.00 abs

7.17 Refer to the “Method Definition” for Bray P-1 for other method parameters not discussed here.

7.18 Run samples using calibration curve. Sample concentration is calculated from the regression equations. Report results to the nearest 0.01 unit for the sample extract and each SPCS.
7.19 If samples are outside calibration range, dilute samples with extracting solution and re-analyze.

7.20 Upon completion of run, place the transmission lines into the NaOH-EDTA solution. Pump the solution for approximately 5 min to remove any precipitated reaction products. Then place these lines in RODI water and pump for an additional 5 min and proceed with the normal “Shut-down” procedure.

8. Calculations
Convert extract P (mg L\(^{-1}\)) to soil P (mg kg\(^{-1}\)) as follows:

\[\text{Soil P (mg kg}^{-1}\text{)} = \left(\frac{A \times B \times C \times R \times 1000}{E}\right)\]

where:
A = Sample extract reading (mg L\(^{-1}\))
B = Extract volume (L)
C = Dilution, if performed
R = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (procedure 3D2)
1000 = Conversion factor to kg-basis
E = Sample weight (g)

9. Report
Report data to the nearest 0.1 mg P kg\(^{-1}\) soil.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References
LACHAT Instruments. 1989. QuikChem method 12-115-01-1-A, phosphorus as orthophosphate, 0.4 to 20 mg P L\(^{-1}\). LACHAT Instruments, 6645 West Mill Rd., Milwaukee, WI.
Selective Dissolutions (4G)

Sodium Pyrophosphate Extraction (4G3)

Acid Digestion (4G3b)

\[ K_2Cr_2O_7 + (H_2SO_4 + H_3PO_4 \text{ Digestion}) \] (4G3b1)

\[ CO_2 \text{ Evolution, Gravimetric (4G3b1a)} \]

Organic Carbon (4G3b1a1)

Air-Dry or Field-Moist, <2 mm (4G3b1a1-a-b1)

1. Application

Sodium pyrophosphate (0.1 \( M \) \( Na_4P_2O_7 \)) is used as a selective dissolution extractant for organically complexed Fe and Al (Wada, 1989). The \( Na_4P_2O_7 \) solution is a poor extractant for allophane, imogolite, amorphous aluminosilicates, and noncrystalline hydrous oxides of Fe and Al. The \( Na_4P_2O_7 \) solution does not extract opal, crystalline silicates, layer silicates, and crystalline hydrous oxides of Fe and Al (Wada, 1989). Sodium pyrophosphate extractable organic C, Fe, and Al were former criteria for spodic placement in soil taxonomy (Soil Survey Staff, 1975). Sodium pyrophosphate extractable Al, Fe, and Mn are currently determined by method (4G3a1-3).

2. Summary of Method

The soil sample is mixed with 0.1 \( M \) \( Na_4P_2O_7 \) and shaken overnight. The solution is then allowed to settle overnight before centrifuging and filtering to obtain a clear extract. The organic C in the sodium pyrophosphate extract is wet oxidized in a fume hood and gravimetrically measured in method 4G3b1a1.

3. Interferences

There are several interferences with this procedure, especially the peptization and dispersion of microcrystalline iron oxide by pyrophosphate (Jeanroy and Guilet, 1981). The quantity of Fe extracted with pyrophosphate decreases with increasing centrifugation (McKeague and Schuppli, 1982); therefore uniform high-speed centrifugation or micropore filtration treatments are required (Schuppli et al., 1983; Loveland and Digby, 1984). Sodium pyrophosphate extraction works best at pH 10 (Loeppert and Inskeep, 1996). The concentration of \( Na_4P_2O_7 \)
solution must be close to 0.1 \( M \). Variable amounts of organic C may be extracted by varying the pyrophosphate concentration.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

5. Equipment

5.1 Electronic balance, ±0.1-mg sensitivity
5.2 Mechanical reciprocating shaker, 200 oscillations min\(^{-1}\), 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.3 Centrifuge, Centra, GP-8, Thermo IEC, Needham Heights, MA
5.4 Digital diluter/dispenser, with syringes 10,000 and 1000 \( \mu \)L, gas tight, Microlab 500, Hamilton Co., Reno, NV
5.5 Dispenser, 40 mL
5.6 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer
5.7 Containers, polypropylene
5.8 Volumetrics, Class A, 100, 250, and 1000-mL
5.9 Centrifuge tubes, 50-mL
5.10 Funnel, 60° angle, long stem, 50-mm diameter
5.11 Filter paper, Whatman 42, 150 mm
5.12 Absorption bulb, Nesbitt with stopper
5.13 Absorption bulb, Stetser-Norton
5.14 Flask, boiling, round bottom, short neck
5.15 Condenser, Allihn
5.16 Funnel, separatory, cylindrical, open top, with stopcock
5.17 Tube, drying, Schwartz

6. Reagents

6.1 Reverse osmosis deionized (RODI) water
6.2 Hydrochloric acid (HCl), concentrated, 12 \( N \)
6.3 Sodium pyrophosphate solution, 0.1 \( M \). Dissolve 446.05 g of \( \text{Na}_4\text{P}_2\text{O}_7 \cdot 10\text{H}_2\text{O} \) in 10 L of RODI water. pH solution to 10.0 with either HCl or NaOH.
6.4 Potassium dichromate \( (\text{K}_2\text{Cr}_2\text{O}_7) \) reagent.
6.5 Potassium iodide solution. Dissolve 100 g of KI in 100 mL of RODI water.
6.6 Silver sulfate, saturate aqueous solution
6.7 Digestion acid mixture: Mix 600 mL of concentrated $\text{H}_2\text{SO}_4$ and 400 mL of 85% $\text{H}_3\text{PO}_4$
6.8 Indicarb or Mikohibite
6.9 Soda lime
6.10 Zinc granules, 300 mesh
6.11 Anhydride
6.12 Acetylene gas, purity 99.6%
6.13 Nitrous oxide gas, compressed
6.14 Compressed air with water and oil traps

7. Procedure

**Extraction of Al, Fe, and Mn**

7.1 Weigh 0.5 g <2-mm or fine-grind, air-dry soil to the nearest mg sample and place in a 50-mL centrifuge tube. If sample is moist, weigh enough soil to achieve ≈0.5 g of air-dry soil.
7.2 Add 30-mL of $0.1 \text{ M Na}_4\text{P}_2\text{O}_7$, pH 10.0 solution to centrifuge tube.
7.3 Cap tube and shake briefly by hand to dislodge soil from tube bottom. Place tube in rack.
7.4 Place rack in shaker and shake overnight (12 to 16 h) at 200 oscillations min$^{-1}$ at room temperature (20 °C ±2°C).
7.5 Remove tubes from shaker and manually shake tubes to dislodge any soil from cap. Allow samples to sit overnight.
7.6 Next day centrifuge sample at 4000 rpm for 15 min. Filter if necessary.

**Organic C Determination**

7.7 Pipet 100 mL of the extract into a 100-ml flask.
7.8 Evaporate the extract to near dryness using a 50 °C water bath and a gentle stream of clean, filtered air.
7.9 Construct the wet combustion apparatus. Refer to figure 4G3-1 for the apparatus for gravimetric organic C determination.
7.10 Add 1 to 2 g of potassium dichromate.
7.11 Wash the neck of the flask with 3 mL of RODI $\text{H}_2\text{O}$ and connect to condenser.
7.12 Attach a weighed Nesbitt bulb to the system and open the valve at the top.
7.13 Pour 25 mL of digestion-acid mixture into the funnel. Add the mixture to the flask and immediately close the stopcock. Use the digestion-acid mixture to lubricate the stopcock.
7.14 The tip of the air-delivery tube should be ≈0.5 cm below the digestion-acid mixture. Adjust the flow of the “carrier stream” to maintain 1 to 2 bubble s⁻¹ rate throughout the digestion. Apply suction on the outlet side of the Nesbitt bulb. Gentle air pressure and needle valve on the air-pressure line aids flow-adjustment.

7.15 With a gas flame or a variable power-heating mantle, gently heat the flask until the mixture boils (≈3 to 4 min). Continue a gentle boiling for 10 min. Heating is too rapid if white fumes of SO₂ are visible above the second bulb of the reflux condenser.

7.16 Remove the heat and allow to aerate for 10 additional min at a rate of 6 to 8 bubbles s⁻¹.

7.17 Close the stopcock on the Nesbitt bulb, disconnect the bulb from the system, and weigh to the nearest 0.0001g.
8. Calculations

8.1 Organic C(%) = \[(W_t - W_i) \times 27.3 \times Volume \times R\] / \[(Sample Weight (g) \times 236.6)\]

where:

- \(W_t\) = Nesbitt bulb weight after digestion (g)
- \(W_i\) = Nesbitt bulb weight before digestion (g)
- Volume = Extract volume digested (mL)
- \(R\) = Air-dry/oven-dry ratio (method 3D1) or field-moist/oven-dry ratio (method 3D2)
- 27.3 = Conversion factor
- 236.6 = Total extract volume (mL)

9. Report

Report organic C the nearest 0.1 of a percent.

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References


Total Analysis (4H)
Acid Digestion (4H1)
  HNO₃ + HCl Digestion (4H1a)
  Microwave (4H1a1)
    Inductively Coupled Plasma Atomic Emission Spectrophotometer (4H1a1a)
      Axial Mode (4H1a1a1)
        Ultrasonic Nebulizer (4H1a1a1a)
          Silver, Arsenic, Barium, Beryllium, Cadmium, Cobalt, Chromium, Copper, Manganese, Molybdenum, Nickel, Phosphorus, Lead, Antimony, Tin, Strontium, Thallium, Vanadium, Tungsten, and Zinc (4H1a1a1a1-20)
      Air-Dry, <2 mm (4H1a1a1a1-20a1)

1. Application

The term of trace elements is widely applied to a variety of elements that are generally present in plants, soils, and water in low concentrations or what is termed background levels. Knowledge of these levels is important in understanding the consequences of increasing levels of trace elements in ecosystems (Tiller, 1989; Holmgren et al., 1993). These elements may become elevated in concentration due to natural (e.g., magmatic activity, mineral weathering, translocation through the soil or landscape) or through human-induced activities (e.g., pesticides, mining, smelting, manufacturing). The relative reactivity or bioavailability of these elements in soils is governed by a variety of chemical factors such as pH, redox potential, organic concentrations, and oxides (Pierzynski and Schwab, 1993; Gambrell, 1994; Keller and Vedy, 1994; Burt et al., 2002). Uses of elemental data in soil survey applications are broad and diverse, ranging from understanding natural (Wilcke and Amelung, 1996; Jersak et al., 1997) to human-induced distributions (Wilcke et al., 1998). Knowledge of the elemental amounts and distribution in soils and their relationships with other soil properties can enhance the understanding of the fate and transport of anthropogenic elements, thereby expanding the utility and application of soil survey knowledge in areas of environmental concern such as urban, mine spoil reclamation, smelter emissions, and agricultural waste applications (Burt et al., 2003).

2. Summary of Method

The approach of this digestion methodology is to maximize the extractable concentration of elements in digested soils while minimizing the matrix interferences such as found in digestion procedures that use HF acid. This method (4H1a1) follows EPA Method 3051A. A 500-mg <2-mm soil separate
which has been air-dried and ground to <200 mesh (75 µm) is weighed into a 100-ml Teflon (PFA) sample digestion vessel. To the vessel, 9.0 mL HNO₃ and 3.0 mL HCl are added. The vessel is inserted into a protection shield and covered, and placed into a rotor with temperature control. Following microwave digestion, the rotor and samples are cooled, and digestate quantitatively transferred into a 50-ml glass volumetric high purity reverse osmosis deionized water. The volumetrics are allowed to stand overnight, filled to volume, and samples transferred into appropriate acid-washed polypropylene containers for analysis. The concentration of Ag, As, Ba, Be, Cd, Co, Cr, Cu, Mn, Mo, Ni, P, Pb, Sb, Sn, Sr, TI, V, W, and Zn are determined using an inductively coupled plasma atomic emission spectrophotometer (ICP–AES) in axial mode by methods 4H1a1a1a1-20, respectively. Mercury is analyzed by a cold-vapor atomic absorption spectrophotometer (CVASS) (4H1a1c1), and As and Se are determined by flow through hydride-generation and atomic absorption spectrophotometer (HGAAS) (4H1a1b1a1-2), respectively.

3. Interferences

Organic constituents may contain metals and are difficult to digest if present in high concentrations. Certain elements are subject to volatile losses during digestion and transfer. Certain soil minerals (e.g., quartz, feldspars) are not soluble in HNO₃ + HCl.

Spectral and matrix interferences exist. Interferences are corrected or minimized by using both an internal standard and inter-elemental correction factors. Also, careful selection of specific wavelengths for data reporting is important. Background corrections are made by ICP software. Samples and standards are matrix-matched to help reduce interferences.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated acids to the fume hood. Wash hands thoroughly after handling reagents. Filling the digestion vessel to greater than 25 percent of the free volume or adding organic reagents or oxidizing agents to the cup may result in explosion of the digestion microwave system.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Pipette(s) capable of delivering 3 and 9 mL, Omnifit Corp. manufacturers variable volume, (10-ml maximum) pipettes suitable for HNO₃ and HCl delivery from 2.5 L bottles
5.3 Volumetric flasks, class A glass, 50 mL
5.4 Polypropylene bottles, 60 mL, with cap
5.5 Electronic balance, (±0.1 mg sensitivity)
5.6 Microwave oven, CEM Mars 5, 14 position-HP500 Plus vessel and rotor (vessels composed of PFA, sleeves composed of advanced composite)
5.7 Volumetrics, 500, 250, and 50-mL class A glass
5.8 Containers, 500-mL, polypropylene, with screw caps
5.9 Pipettes, electronic digital, 250 µL and 10 mL, Rainin Instrument Co., Woburn, MA
5.10 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES), Perkin-Elmer Optima 3300 Dual View (DV), Perkin-Elmer Corp., Norwalk, CT
5.11 RF generator, floor mounted power unit, 45 MHz free running, Perkin-Elmer Corp., Norwalk, CT.
5.12 Computer, with WinLab software ver. 4.1, Perkin-Elmer Corp., Norwalk, CT, and printer
5.13 Recirculating chiller, Neslab, CFT Series
5.14 Compressed gasses, argon (minimum purity = 99.996%) and nitrogen (minimum purity = 99.999%)
5.15 Autosampler, AS-90, Perkin-Elmer Corp., Norwalk, CT.
5.16 Quartz torch, Part No. N069-1662; alumina injector (2.0 mm id), Part No. N069-5362
5.17 Ultrasonic nebulizer, Model U-5000AT+, CETAC Corp., Omaha, NE
5.18 Peristaltic pump (for automatic injection of internal standard)

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Concentrated hydrochloric acid (HCl), 12 N, trace pure grade
6.3 Concentrated nitric acid (HNO₃), 16 N, trace pure grade
6.4 Primary standards: 1000 mg L⁻¹, from High Purity Standards, Charleston, SC. Single elemental standards are manufactured in dilute HNO₃, HNO₃ + HF, or H₂O

7. Procedure

Microwave Acid Digestion

7.1 About 500 mg of fine-earth (<2-mm) or a specific particle size separate ground to <200-mesh (75 µm) is weighed to the nearest 0.1 mg in a 100-mL digestion vessel.
7.2 Note: If sample is principally composed of organic materials (organic C > 15%), perform a preliminary digestion in the muffle furnace in an digestion crucible: 250°C for 15 min, 450 °C for 15 min, followed by 550 °C for 1 h.

7.3 Pipette 9.0 mL HNO$_3$ and 3.0 mL HCl into the sample and allow to completely wet. Add acids in the fume hood. Allow acids to react and vent in uncovered vessels for about 30 min.

7.4 Place covered vessels in protective sleeve, cover and place into rotor.

7.5 Place digestion rotor in the microwave oven and insert the temperature probe into the reference vessel. Attach the probe cable into the fitting in the top of the microwave. Connect the pressure monitor to the vessel.

7.6 Microwave settings are as follows:
- 1200 watts at 100% power for 5.5 min until 175 °C
- Hold at 175 °C for 4.5 min
- Cool for 5 min

7.7 After cooling, disconnect temperature probe and pressure sensor from microwave.

7.8 Remove rotor from oven, and place in fume hood.

7.9 Open each vessel carefully and then quantitatively transfer contents of vessel to a 50-mL volumetric flask with RODI water.

7.10 Cap flask and mix well by inverting. Allow to stand overnight. Finish filling to volume with RODI water.

7.11 Decant contents into a labeled 60-mL polypropylene container.

7.12 Prepare working standards of a blank, reference soil sample from the SSL repository, NIST or other standard reference material, and blank by the same digestion method. Run two of these standards or blank with each set of 14 samples.

**ICP–AES Calibration Standards, Set-Up, and Operation**

7.13 A primary mixed calibration standard (PMCS) is prepared from the respective primary elemental standards (1000 mg L$^{-1}$) to a 500-mL final volume. From this PMCS, three working calibration standards (WCS) are prepared. In addition, a single element standard for Tl (STl$_1$) is prepared separately from a 1000 µg L$^{-1}$ stock standard to a 50-mL final volume. Also, prior to diluting to volume, add 90 mL HNO$_3$ and 30 mL HCl to the PMCS and 9 mL HNO$_3$ and 3 mL HCl to the STl$_1$. Use RODI water to dilute to final volume for PMCS and STl$_1$. Invert to mix thoroughly. Store in polyethylene container in refrigerator. Make fresh on a routine basis. The amount of the primary standards (1000 mg L$^{-1}$) to make the PMCS, amount of the1000 µg L$^{-1}$ Tl stock standard to make the STl$_1$ and the final elemental concentrations of the PMCS and STl$_1$ are as follows:
<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration</th>
<th>Primary Standard Required</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(µg L(^{-1}))</td>
<td>(mL)</td>
</tr>
<tr>
<td>As</td>
<td>2,000</td>
<td>1</td>
</tr>
<tr>
<td>Ni</td>
<td>4,000</td>
<td>2</td>
</tr>
<tr>
<td>P</td>
<td>60,000</td>
<td>30</td>
</tr>
<tr>
<td>Cr</td>
<td>4,000</td>
<td>2</td>
</tr>
<tr>
<td>M</td>
<td>40,000</td>
<td>20</td>
</tr>
<tr>
<td>Cu</td>
<td>20,000</td>
<td>10</td>
</tr>
<tr>
<td>Zn</td>
<td>20,000</td>
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<tr>
<td>C</td>
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<td>1</td>
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<tr>
<td>Pb</td>
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<tr>
<td>Co</td>
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<tr>
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<td>Sb</td>
<td>400</td>
<td>0.2</td>
</tr>
<tr>
<td>Sr</td>
<td>60,000</td>
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<tr>
<td>M</td>
<td>400</td>
<td>0.2</td>
</tr>
<tr>
<td>V</td>
<td>20,000</td>
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<td>8,000</td>
<td>4</td>
</tr>
<tr>
<td>Ti</td>
<td>500</td>
<td>25</td>
</tr>
<tr>
<td>W</td>
<td>2,000</td>
<td>1</td>
</tr>
</tbody>
</table>

7.14 The WCS are made from dilution of the PMCS with the exception of single element Ti (STI\(_2\)), which is made up separately. The three WCS (Low, Medium, and High) require 0.625, 6.25, and 62.5 mL PMCS diluted to 250-mL final volume, respectively. Also, prior to diluting to volume, add 44.89, 43.88, and 33.75 mL HNO\(_3\) and 14.96, 14.63, and 11.25 mL HCl to the WCS (Low, Medium, and High, respectively). The STI\(_2\) requires 0.5, 5, and 50 mL of the STI\(_1\) diluted to a 50-mL final volume. Also, prior to diluting to volume, add 8.91 and 8.1 mL HNO\(_3\) and 2.97 and 2.7 mL HCl for the Low and Medium STI\(_2\), respectively. Use RODI water to dilute to final volume for WCS and STI\(_2\). Invert to mix thoroughly. Store in polyethylene container in a refrigerator. Make fresh on a routine basis. The elemental
concentrations of the Low, Medium, and High WCS and the ST1₂ are as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low (µg L⁻¹)</td>
<td>Medium</td>
<td>High</td>
<td></td>
</tr>
<tr>
<td>Ni</td>
<td>10</td>
<td>100</td>
<td>1000</td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>150</td>
<td>1500</td>
<td>15,000</td>
<td></td>
</tr>
<tr>
<td>Cr</td>
<td>10</td>
<td>100</td>
<td>1000</td>
<td></td>
</tr>
<tr>
<td>Mn</td>
<td>100</td>
<td>1000</td>
<td>10,000</td>
<td></td>
</tr>
<tr>
<td>Cu</td>
<td>50</td>
<td>500</td>
<td>5000</td>
<td></td>
</tr>
<tr>
<td>Zn</td>
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</tr>
<tr>
<td>Cd</td>
<td>5</td>
<td>50</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>Pb</td>
<td>10</td>
<td>100</td>
<td>1000</td>
<td></td>
</tr>
<tr>
<td>Co</td>
<td>10</td>
<td>100</td>
<td>1000</td>
<td></td>
</tr>
<tr>
<td>Ag</td>
<td>1</td>
<td>10</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>Ba</td>
<td>300</td>
<td>300</td>
<td>3000</td>
<td></td>
</tr>
<tr>
<td>Be</td>
<td>1.5</td>
<td>15</td>
<td>150</td>
<td></td>
</tr>
<tr>
<td>Sb</td>
<td>1.0</td>
<td>10</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>Sr</td>
<td>150</td>
<td>1500</td>
<td>15,000</td>
<td></td>
</tr>
<tr>
<td>Mo</td>
<td>1</td>
<td>10</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>50</td>
<td>500</td>
<td>5000</td>
<td></td>
</tr>
<tr>
<td>Sn</td>
<td>20</td>
<td>200</td>
<td>2000</td>
<td></td>
</tr>
<tr>
<td>Tl</td>
<td>5</td>
<td>50</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>W</td>
<td>5</td>
<td>50</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>As</td>
<td>5</td>
<td>50</td>
<td>500</td>
<td></td>
</tr>
</tbody>
</table>

7.15 Single element primary standards (1000 mg L⁻¹, Al, Fe, Mo, V, Mn) are required to create the inter-elemental correction (IEC) factors. These are prepared in the matrix of the digests and are combined into one solution for routine calibration. The single element IEC standards (SEIECS) are required to determine the IEC's. The mixed IEC standard (MIECS) is required for routine calibration. The SEIECS is based on a 50-mL final volume and the MIECS is based a 250-mL final volume. Use RODI water to dilute to final volume for SEIECS and MIECS. Invert to mix thoroughly.
Store in polyethylene container in a refrigerator. Make fresh on routine basis. The amount of the primary standard (1000 mg L\(^{-1}\)) to make the SEIECS and MIECS solutions and the final elemental concentration of these IEC solutions are as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>IEC Solution</th>
<th>Primary Standard Required</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Concentration</td>
<td>MIECS</td>
</tr>
<tr>
<td></td>
<td>mg L(^{-1})</td>
<td>mL</td>
</tr>
<tr>
<td>Al</td>
<td>100</td>
<td>25.00</td>
</tr>
<tr>
<td>Fe</td>
<td>100</td>
<td>25.00</td>
</tr>
<tr>
<td>Mo</td>
<td>5</td>
<td>1.25</td>
</tr>
<tr>
<td>V</td>
<td>10</td>
<td>2.50</td>
</tr>
<tr>
<td>Mn</td>
<td>10</td>
<td>2.50</td>
</tr>
</tbody>
</table>

7.16 To MIECS, add 45 and 15 mL HNO\(_3\) and HCl, respectively. To SEIECS, add 9 and 3 mL HNO\(_3\) and HCl, respectively. Use RODI water to dilute to final volume for MIECS and SEIECS.

7.17 The elements chosen for IEC factors are based on established spectral interferences with chosen analyte wavelengths. The SEIECS should initially be prepared in separate 50-mL volumetrics for establishment of IEC factors and then prepared (MIECS) in a single 250-mL volumetric for routine analysis. IEC factors are established via a procedure in the WinLab software in which the amount of interference on the analyte (in µg L\(^{-1}\)) is measured for each mg L\(^{-1}\) of interferent concentration in the digest.

7.18 A 10 mg L\(^{-1}\) Lu internal standard (read at 291.138 nm) is added to the blank, all calibration standards, and samples. It is prepared by adding 5.0 mL Lu primary standard (1000 mg L\(^{-1}\)) and 10 ml conc. HNO\(_3\) to 500-mL volumetric flask, and diluting to volume with RODI water. Internal standard is automatically injected via the peristaltic pump and mixing block.

7.19 Use the ICP–AES in axial mode and ultrasonic nebulization to analyze sample. Internal standard is added via an external peristaltic pump at 15% pump speed using 0.44 mm id. pump tubing. Internal standard and samples or standards are mixed via a mixing block and coil prior to entering the ultrasonic nebulizer. No initial dilutions of samples are necessary prior to analysis. Perform instrument checks (Hg alignment; BEC and %RSD of 1 mg L\(^{-1}\) Mn solution) prior to analysis as discussed in operation manual of instrument. Check instrument alignment and gas pressures to obtain optimum readings with maximum signal to noise ratio.
7.20 Analyses are generally performed at two or more wavelengths for each element. The selected wavelengths are as follows: (reported wavelength listed first and in boldface):

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>237.312 308.215</td>
</tr>
<tr>
<td>Fe</td>
<td>302.107 238.203</td>
</tr>
<tr>
<td>Mn</td>
<td>260.570 203.844</td>
</tr>
<tr>
<td>P</td>
<td>178.221, 213.620</td>
</tr>
<tr>
<td>Cr</td>
<td>267.710, 205.558</td>
</tr>
<tr>
<td>Cu</td>
<td>324.753, 327.396</td>
</tr>
<tr>
<td>Ni</td>
<td>232.003, 231.604,</td>
</tr>
<tr>
<td>Zn</td>
<td>213.857, 206.197</td>
</tr>
<tr>
<td>Cd</td>
<td>228.802, 226.501</td>
</tr>
<tr>
<td>Pb</td>
<td>220.353, 216.998</td>
</tr>
<tr>
<td>Co</td>
<td>228.614</td>
</tr>
<tr>
<td>Sb</td>
<td>217.582, 206.833</td>
</tr>
<tr>
<td>Sr</td>
<td>460.733, 407.771</td>
</tr>
<tr>
<td>Ba</td>
<td>233.525, 455.507</td>
</tr>
<tr>
<td>Be</td>
<td>313.104, 313.046</td>
</tr>
<tr>
<td>As</td>
<td>188.979</td>
</tr>
<tr>
<td>Ag</td>
<td>328.068, 338.287</td>
</tr>
<tr>
<td>Mo</td>
<td>202.031, 203.845</td>
</tr>
<tr>
<td>V</td>
<td>292.402, 310.230</td>
</tr>
<tr>
<td>Sn</td>
<td>189.927, 235.485</td>
</tr>
<tr>
<td>W</td>
<td>207.912, 224.876</td>
</tr>
<tr>
<td>Ti</td>
<td>190.801, 276.787</td>
</tr>
<tr>
<td>Lu</td>
<td>291.138 (Internal Standard)</td>
</tr>
<tr>
<td>Al (IEC)</td>
<td>237.312</td>
</tr>
<tr>
<td>Fe (IEC)</td>
<td>302.107</td>
</tr>
<tr>
<td>Mo (IEC)</td>
<td>202.031</td>
</tr>
<tr>
<td>V (IEC)</td>
<td>292.402</td>
</tr>
<tr>
<td>Mn (IEC)</td>
<td>260.568</td>
</tr>
</tbody>
</table>
7.21 Use the blank standard solution to dilute those samples with concentrations greater than the high standard. Rerun all elements and use only the data needed from the diluted analysis.

7.22 Establish detection limits using the blank standard solution. The instrumental detection limits are calculated by using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are reported as “ND” or non-detected.

7.23 The extract obtained in this method (4H1a1) is used in method 4H1a1c1 for Hg analysis and in methods 4H1a1b1a1-2 for As and Se analysis, respectively.

8. Calculations
The calculation of mg kg\(^{-1}\) of an element in the soil from \(\mu g\ L^{-1}\) in solution is as follows:

\[
\text{Analyte concentration in soil (mg kg}^{-1}) = \frac{A \times B \times C \times R \times 1000}{E \times 1000}
\]

- A = Sample extract reading (\(\mu g\ L^{-1}\))
- B = Extract volume (L)
- C = Dilution, if performed
- R = Air-dry/oven-dry ratio (method 3D1)
- 1000 = Conversion factor in numerator to kg-basis
- E = Sample weight (g)
- 1000 = Factor in denominator (\(\mu g\ mg^{-1}\))

9. Report
Analyses are generally performed at two or more wavelengths for each element, with the one selected wavelength for reporting purposes. The particle-size fraction digested needs to be identified with each sample. Data are reported to the nearest 0.01 mg kg\(^{-1}\).

10. Precision and Accuracy
Precision and accuracy data are available from the SSL upon request.

11. References


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**Total Analysis (4H)**

**Acid Digestion (4H1)**

HNO₃ + HCl Digestion (4H1a)

Microwave (4H1a1)

HCl Digestion (4H1a1b)

Water Bath (4H1a1b1)

Flow Through Hydride-Generation and Atomic Absorption Spectrophotometer (4H1a1b1)

Arsenic and Selenium (4H1a1b1a1-2)

Air-Dry, <2 mm (4H1a1b1a1-2a1)

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**1. Application**

Arsenic is an extremely toxic element that is occurs in both organic and inorganic forms in soils. It is typically found in low concentrations, but has been widely applied to soils as a component in pesticides and herbicides and also via industrial pollution and smelting operations. The element is used in drugs, soaps, dyes, and metals, though 90% of industrial As in the U.S. is used in wood preservatives (Pinsker, 2001). Concerns exist for both short-term (acute) and long-term (chronic) soil exposure. Primary route for exposure is via soil ingestion or inhalation of air-borne particles. Data from As measurements in groundwater
by U.S. Geological Survey and Environmental Protection Agency has suggested that most As in groundwater is related to natural sources (Ryker, 2001), i.e., from mineral dissolution from minerals in geologic formations and soils. For example, As released via pyrite oxidation is in part responsible for groundwater As levels in Bangladesh ranging between 50 and 2,500 µg/L in many wells. Soil applied As is generally immobile, with soil chemistry similar to phosphorus. The element occurs as arsenate (As$^{5+}$) and arsenite (As$^{3+}$), and in soils, is in the form of the oxyanion, AsO$_4$$^{3-}$. The weathering of limestone and biological accumulation of the element by aquatic organisms is responsible for the high levels in wetland soils of Florida (Chen et. al., 2002).

Selenium is a naturally occurring element in rocks, but is especially concentrated in certain geologic formations, such as Mancos Shale in Colorado and Wyoming and in the shales of the Moreno and Kreyenhagen Formations of California (Martens and Suarez, 1997). Selenium occurs in four species (related to valance states): selenate (Se$^{6+}$), selenite (Se$^{4+}$), elemental Se (Se$^0$), and selenide (Se$^{2-}$). The bioavailability and toxicity is related to speciation. The oxidized species are more commonly found in soils and water. The element is important due to both deficiency (forages for animals) and toxicity (bioaccumulation) concerns (Huang and Fujii, 1996).

2. Summary of Method

A soil sample is digested with HNO$_3$ and HCl in a microwave oven (method 4H1a1). Following extraction, samples are diluted with water to a final 50-mL volume. A 6-mL aliquot of the digestate is combined with 6 mL concentrated H$_2$SO$_4$ and heated at 180 °C for 5 min in the microwave oven to eliminate the HNO$_3$. Then, the extract is combined with 14 mL of water and 20 mL of concentrated HCl and boiled for 30 min. Sample extracts are allowed to cool. Potassium iodide is added as a pre-reduction step for As analysis, with a final concentration of 1% in analysis solutions. Solutions are allowed to stand 1 h before analyzed for total As and/or Se, using flow through hydride-generation and atomic absorption spectrophotometer (HGAAS). Under acidic conditions, sodium borohydride (NaBH$_4$) reduces As and Se to form gaseous products that can be detected by atomic adsorption. For example:

$$3\text{NaBH}_4 + 4\text{H}_2\text{SeO}_3 \rightarrow 4\text{H}_2\text{Se}_9 + 3\text{H}_3\text{BO}_4 + 3\text{NaOH}$$

The data are automatically recorded by a computer and printer. The As and Se concentrations are reported as mg kg$^{-1}$ in the soil by method 4H1a1b1a1-2, respectively.

3. Interferences

Oxidizing acids (e.g., nitric) can produce interferences, and inter-element interferences (e.g., Cu, Sn, Ni, Fe, Cr, Pb, Co) can affect determinations. Even
small amounts of nitric acid produce suppressed, erratic absorbance signals and low recoveries, more so for As than Se. This interference can be effectively eliminated with either (1) addition of urea before the potassium iodide pre-reduction step, or (2) boiling the samples with $\text{H}_2\text{SO}_4$. Alternatively, if samples are boiled in $\text{HCl}$, the quantity of nitric acid can be sufficiently reduced to eliminate the need for urea. Hydride-forming elements may exist in more than one oxidation state, affecting the signal. In general, $\text{As}^{5+}$ methods produce a signal that may be 20 to 50% of that produced by $\text{As}^{3+}$. There are typically more inter-element interferences for $\text{As}^{5+}$ methods than for $\text{As}^{3+}$. Inter-element interferences can be reduced by using the lowest possible concentration of sodium borohydride. Best results can be obtained for difficult samples containing high concentrations of metals if the sodium borohydride concentration is reduced to 0.3% w/v.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves); eye protection (face shields, goggles, or safety glasses); and a breathing filter when handling As and Se solutions. These elements are extremely toxic.

Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±0.1-mg sensitivity
5.2 Atomic absorption spectrophotometer (AAS), PE Model Analyst 300, Perkin-Elmer Corp., Norwalk, CT
5.3 System 2 Electrodeless Discharge Lamp (EDL) Power Supply, with lamps for As and Se, Perkin-Elmer Corp., Norwalk, CT
5.4 Autosampler, Model 90A, Perkin-Elmer Corp., Norwalk, CT
5.5 WinLab Software, Ver. 4.1, Perkin-Elmer Corp., Norwalk, CT
5.6 Computer, Dell Optiplex GXM 333 MHz Pentium, Dell Computer Corp., 17 in color monitor
5.7 Printer, Hewlett-Packard LaserJet 880A
5.8 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.9 Double-stage regulator, argon service
5.10 Varian Vapor Generation Accessory, Model VGA-77
5.11 Tubes, 50-mL for calibration standards
5.12 Test tubes, 50-mL, Corning Pyrex, for sample digestion
5.13 Test tubes, 25-mL, 16 mm x 100, for sample dilution and autosampler
5.14 100, 250, and 1000-mL volumetrics, class A.
5.15 Containers, polypropylene and glass
5.16 Water bath, 95 °C-capability

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Sulfuric acid (H$_2$SO$_4$), 36 M, trace metal purity
6.3 Sodium hydroxide (NaOH•3H$_2$O)
6.4 Sodium borohydride (NaBH$_4$) pellets
6.5 Hydrochloric acid (HCl), 12 M, trace metal purity, assay 35–38%
6.6 Acetylene gas, purity 99.6%.
6.7 Compressed air with water and oil traps
6.8 Compressed Argon
6.9 Primary standards, 1000 mg L$^{-1}$ As and 1000 mg L$^{-1}$ Se, High Purity Standards, Inc.
6.10 Primary mixed working standard (PMWS), 1000 µg L$^{-1}$ As and Se: Add 500 mL RODI water to a 1-L volumetric. Add 1 mL of 1000 mg L$^{-1}$ As, 1 mL of 1000 mg L$^{-1}$ Se, and 10 mL concentrated HCl. Fill to volume with RODI water. Invert to mix thoroughly. Final concentration is ≈1% HCl. Store in polyethylene container in a refrigerator. Make fresh weekly.
6.11 Mixed Calibration Standards (MCS), 150, 100, 75, 50, 25, 12, and 0 µg L$^{-1}$: To seven 250-mL volumetrics, add PMWS and concentrated HCl as follows:
   6.11.1 150 µg L$^{-1}$ = 37.5 mL PMWS + 106 mL HCl
   6.11.2 100 µg L$^{-1}$ = 25.0 mL PMWS + 113 mL HCl
   6.11.3 75 µg L$^{-1}$ = 18.75 mL PMWS + 116 mL HCl
   6.11.4 50 µg L$^{-1}$ = 12.5 mL PMWS + 119 mL HCl
   6.11.5 25 µg L$^{-1}$ = 6.25 mL PMWS + 122 mL HCl
   6.11.6 12 µg L$^{-1}$ = 3.0 mL PMWS + 124 mL HCl
   6.11.7 0 µg L$^{-1}$ = 0 mL PMWS + 125 mL HCl
6.12 Bring to volume with RODI water and invert to mix thoroughly. Store in polyethylene container in a refrigerator. Standards will keep for 2 to 3 days with refrigeration. Quality Control (QC) check is MCS 100 µg L$^{-1}$. Final concentration of MCS and QC check is ≈6 M HCl.
6.13 Acid Carrier, 6 M HCl: Add 200 mL RODI water to a 500-mL volumetric. Carefully add 250 mL concentrated HCl and fill to volume with RODI water. Invert to mix thoroughly.
6.14 Reductant, 0.5% NaOH-0.3% NaBH₄: Add 200 mL RODI water to a 500-mL volumetric. Always stabilize the solution by first adding the NaOH. Add 2.5 g NaOH and mix until dissolved. Add 1.5 g of NaBH₄, mix until dissolved, and fill to volume. Invert to mix thoroughly. Degas for 10 min. Make fresh daily.

6.15 Potassium iodide (KI) (10%) – ascorbic acid solution (10%): To a 500-mL volumetric, add 250 mL of RODI water. Dissolve 50 g of KI and 50 g of ascorbic acid and dilute to volume with RODI water. Invert to mix thoroughly. Make fresh daily. (Procedure requires 1-mL per sample.)

6.16 Diluent, 6 M HCl, for samples: Add 400 mL RODI water to a 1-L volumetric. Carefully add 500 mL concentrated HCl and fill to volume with RODI water. Invert to mix thoroughly.

6.17 QC Soil Standards: Loam C (certified reference material, purchased from High purity Standards) and NIST SRM 2710 (National Institute of Standards and Technology, Standard Reference Material), all prepared to <200 mesh (75 µm).

7. Procedure

Digestion of acid (HNO₃ + HCl) extract

7.1 Prepare soil samples, soil standards, blanks, and spikes prepared in method 4H1a1 as follows: Pipette 6mL of acid (HNO₃ + HCl) extract and 6mL concentrated H₂SO₄ into a 50-mL Pyrex tube. Place in a glass beaker (4 tubes into each 250-mL beaker; maximum of 3 beakers, i.e., 12 samples, in microwave at one time). Heating program for microwave is as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage</td>
<td>1</td>
</tr>
<tr>
<td>Max. Power (watts)</td>
<td>600</td>
</tr>
<tr>
<td>Heating Power (%)</td>
<td>100</td>
</tr>
<tr>
<td>Program Ramp (min)</td>
<td>5:00</td>
</tr>
<tr>
<td>Pressure (psi)</td>
<td>800</td>
</tr>
<tr>
<td>Temperature (°C)</td>
<td>180</td>
</tr>
<tr>
<td>Hold (min)</td>
<td>30:00</td>
</tr>
</tbody>
</table>

Place thermocouple (in thermowell) into one sample. Heat microwave until 180 °C is reached for 5 minutes. At that time, fuming (loss of HNO₃) should cease. Allow samples to cool and vent in the microwave for 5 minutes prior to removal.

7.2 To the microwave digested soil samples and MCS, pipette 15 mL of RODI water+20 mL of concentrated HCl. Include one QC soil standard and blank
in each sample rack. Soil samples and QC soil standards have a final concentration of \( \approx 6 \, M \) HCl.

7.3 Digested the open tubes (do not cover) in a sample rack by submerging up to the neck of the tubes in a water bath (95 °C). Heat for 30 min and remove to cool.

7.4 Remove tubes, cool, fill to volume and cap. Mix well by inverting. (Note: digested soil samples and QC soil standards can be analyzed the same day, or placed into the refrigerator overnight for subsequent analysis the following day.) For As analysis, proceed to Section 7.5. For Se analysis, proceed to Section 7.7.

7.5 Arsenic (MCS): Pipette 36 mL of digested MCS + 4 mL of KI-ascorbic acid solution into test tubes that have been placed in the sample holder of the sample changer. Allow to stand 1 h before As analysis. Do not allow MCS to stand greater than 2 h before As analysis. For every sample extract rack, do a set of MCS.

7.6 Arsenic (sample extracts): Pipette 9 mL of sample extract + 1 mL of KI-ascorbic acid solution into test tubes that have been placed in the sample holder of the sample changer. Allow to stand 1 h before As analysis. Do not allow samples to stand longer than 2 h before As analysis.

7.7 Selenium (sample extracts and MCS): Pour sample extract and MCS (without KI-ascorbic acid solution) into test tubes that have been placed in the sample holder of the sample changer.

**HGAAS Set-up and Operation**

7.8 Each element is analyzed separately on the atomic absorption spectrometer. Refer to manufacturer’s manual for operation of AAS. Connect EDL power supply to AAS. Allow EDL to warm up 20 to 30 minutes. Follow the manufacturer’s operating procedures for AA EDL settings, warm-up, and adjustments to settings. Connect vapor generation accessory to AA. Follow the manufacturer’s operating procedures for appropriate gas and liquid flow. Instrumental parameters for each element are as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength (nm)</th>
<th>Slit Width (mA)</th>
<th>EDL Current (mA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenic</td>
<td>193.7</td>
<td>0.7</td>
<td>380</td>
</tr>
<tr>
<td>Selenium</td>
<td>196.0</td>
<td>2.0</td>
<td>250</td>
</tr>
</tbody>
</table>

The flame is required only for heating the gas flow tube and is maintained as a low temperature as possible. Use an air/C\(_2\)H\(_2\) mixture of 6.5 and 0.2 L min\(^{-1}\).
7.9 For automated analysis (using the autosampler), the analysis is performed using atomic absorption with background correction, time average mode, with a read delay of 48 s, BOC time of 5 s, and read time of 2 s. Rinse for 30 s between samples. Reported values are the average of five replications.

7.10 Use the computer and printer to set instrument parameters and to collect and record instrument readings.

7.11 The instrument readings are programmed to display analyte concentration in $\mu$g L$^{-1}$ (ppb).

**HGAAS Calibration**

7.12 Each element is analyzed during separate runs on the AA. Use the calibration reagent blank and calibration standards to calibrate the AAS. Detection limits are 5 and 10 $\mu$g L$^{-1}$ for As and Se, respectively.

7.13 Use the QC standards after every 12th sample. The QC is 100 $\mu$g L$^{-1}$ for As and Se, respectively, with $\pm$30% rejection criteria. If QC fails after three attempts, recalculate and reread the QC. The QC is read at the end of each run.

7.14 If samples are outside the calibration range, a 1:5 serial dilution is performed using 6 M HCl. The QC soil standards (Loam C and SRM 2710) have As contents of 47±3 and 626±38, mg kg$^{-1}$ soil, respectively, as consensus/certified values. These QC soil standards have automatic dilutions of 1:10 and 1:100, respectively.

**Clean-up and Maintenance**

7.15 Soak the absorption cell in dilute HNO$_3$ acid (0.1% w/v) for 30 min, rinse thoroughly with RODI water, and allow to dry.

7.16 If gas/liquid separator and tubing (including autosampler sipper) has been exposed to contamination with KI, pump a freshly prepared 1% sodium thiosulfate solution through the system for 10 min. The sodium thiosulfate solution must be removed by pumping RODI water through the system for 10 min.

7.17 If gas/liquid separator and tubing have not been exposed to contamination with KI, pump RODI water through the system for 10 min.

**8. Calculations**

Convert extract As and Se ($\mu$g L$^{-1}$) to soil As and Se (mg kg$^{-1}$) as follows:

Soil As (mg kg$^{-1}$) = \[\text{A} \times \text{B} \times \text{C} \times \text{R} \times 1000\] / E \times 1000

where:

A = Sample extract reading ($\mu$g L$^{-1}$)
B = Extract volume (L) (0.05)
C = Dilution, if performed
1000 = Conversion factor in numerator to kg-basis
1000 = Factor in denominator (µg mg⁻¹)
R = Air-dry/oven-dry ratio (method 3D1)
E = Sample weight (g) (0.5)

Soil Se (mg kg⁻¹) = [A x B x C x R x 1000] / E x 1000

where:
A = Sample extract reading (µg L⁻¹)
B = Extract volume (L) (0.05)
C = Dilution, if performed
1000 = Conversion factor in numerator to kg-basis
1000 = Factor in denominator (µg mg⁻¹)
R = Air-dry/oven-dry ratio (method 3D1)
E = Sample weight (g) (0.5)

9. Report
Report As and Se to the nearest 0.1 mg kg⁻¹ soil.

10. Precision and Accuracy
Precision and accuracy data are available from the SSL upon request.

11. References

Total Analysis (4H)
Acid Digestion (4H1)
   HNO₃ + HCl Digestion (4H1a)
   Microwave (4H1a1)
      Cold Vapor Atomic Absorption Spectrophotometer (4H1a1c)
         Mercury (4H1a1c1)
            Air-Dry, <2-mm (4H1a1c1a1)

1. Application
   Mercury is highly toxic to both plants and animals, and enters the food chain primarily through atmospheric deposition (smelting, coal combustion, volcanic activity) and pesticide usage (Pais and Jones, 1997). Due to the absorption of Hg by both organic and inorganic soil components, many studies have been performed which have examined soil-Hg interactions (MacNaughton and James, 1974; Barrow and Cox, 1992; Yin et al., 1996) and ecosystem distributions (Hall et al., 1987; Inacio et al., 1998).

2. Summary of Method
   Soil digests (HNO₃ + HCl) from method 4H1a1 are analyzed for Hg using cold-vapor atomic absorption spectroscopy (CVAAS). This method is based on absorption of radiation at 253.7 nm wavelength by Hg vapor. The digest is mixed with stannous chloride to reduce Hg to the elemental state. Using argon as a carrier gas, the solution is passed over a gas-liquid separator in a closed system to separate the gaseous Hg from solution. The Hg vapor passes through a cell positioned in the light path of an atomic absorption spectrophotometer. Absorbance (peak height) is measured as a function of mercury concentration. Mercury data are reported as µg kg⁻¹ soil by method 4H1a1c1.

3. Interferences
   Copper, chlorides, and certain volatile organic materials may be interferences.

4. Safety
   Soil digests contain acid and must be handled appropriately. Procedure uses a primary Hg standard that is diluted for working standards. Use gloves and avoid skin contact. Gasses exhausting from the Hg analyzer cabinet, prior to passing through the Hg vapor trap may contain Hg vapor. Do not run the instrument unless the exhaust gas is properly scrubbed or removed.
5. Equipment
5.1 Cold-vapor atomic absorption spectrophotometer (CVAAS), CETAC M-6000A Mercury Analyzer, CETAC Corp., Omaha, NE
5.2 Autosampler, CETAC ASX-500 Model 510, CETAC Corp., Omaha, NE
5.3 Autodilutor Accessory, CETAC ADX-500, CETAC Corp., Omaha, NE
5.4 Nafion drying tube, CETAC Corp., Omaha, NE
5.5 Peristaltic Pump, CETAC Corp., Omaha, NE
5.6 Computer, Microsoft Windows 97, CETAC M-6000A Software, CETAC, Corp., Omaha, NE, and printer
5.7 Pipettors, electronic digital, Rainin Instrument Co., Woburn, MA, 2500 µL and 10 mL
5.8 Compressed argon gas
5.9 Calcium sulfate (anhydrous) or equivalent desiccant

6. Reagents
6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water
6.2 Concentrated hydrochloric acid (HCl), 12 N. Use trace pure HCl.
6.3 Concentrated nitric acid (HNO₃), 16 N. Use trace pure HNO₃.
6.4 Primary standard: 1000 mg L⁻¹ Hg, High-Purity Standards, Charleston, SC.
6.5 Stock standard, 500 µg L⁻¹: Add 0.25 mL primary standard to 500 mL volumetric and dilute to mark with RODI water and invert to mix thoroughly.
6.6 Standard Hg calibration solutions or working standards are 4.0, 3.0, 2.5, 1.0, 0.5, and 0.0 µg L⁻¹. To six 50-mL volumetric flasks add as follows:

<table>
<thead>
<tr>
<th>Concentration (µg L⁻¹)</th>
<th>Reagent Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>9 mL HNO₃ + 3 mL HCl</td>
</tr>
<tr>
<td>0.5</td>
<td>0.05 mL stock standard + 9 mL HNO₃ + 3 mL HCl</td>
</tr>
<tr>
<td>1.0</td>
<td>0.1 mL stock standard + 9 mL HNO₃ + 3 mL HCl</td>
</tr>
<tr>
<td>2.5</td>
<td>0.25 mL stock standard + 9 mL HNO₃ + 3 mL HCl</td>
</tr>
<tr>
<td>3.0</td>
<td>0.3 mL stock standard + 9 mL HNO₃ + 3 mL HCl</td>
</tr>
<tr>
<td>4.0</td>
<td>0.4 mL stock standard + 9 mL HNO₃ + 3 mL HCl</td>
</tr>
</tbody>
</table>

Dilute each working standard to mark with RODI water and invert to mix thoroughly. These working standards are used for Normal and High Throughput Ranges.

6.7 Stannous chloride, reducing agent 10% stannous chloride solution (SnCl₂ in 7% HCl). Add 50 g of stannous chloride and 97.2 mL concentrated HCl
to 500 mL volumetric and dilute to mark with RODI water. Invert to mix thoroughly.

6.8 Diluent: Add 90 mL concentrated HNO$_3$ and 30 mL concentrated HCl to 500 mL volumetric and dilute to mark with RODI water. Invert to mix thoroughly.

6.9 Rinse (5% HNO$_3$): Add 71.4 mL HNO$_3$ to 1-L volumetric and dilute to mark with RODI water. Invert to mix thoroughly.

6.10 Potassium permanganate, solid, crystalline, fills safety trap for Hg vapor exhaust.

6.11 Glass wool. Fine glass wool only.

7. Procedure

7.1 Oven radiator temperature must be at 125 °C to maintain the actual gas temperature of 50 °C. Argon gas carrier must be supplied at 100 psig (6.9 bar). Liquid flow is always set at fixed flow of 4.0 mL min$^{-1}$ (sample) and 0.8 mL min$^{-1}$ (reagent).

7.2 Turn mercury analyzer and mercury lamp for warm-up (90 min) prior to analysis. Ensure integrity of lamp (there is some loss of performance at 13 mA but replace at 15 mA).

7.3 Turn on computer and choose Worksheet Template appropriate to range of analysis. Three ranges of analysis (Highest Sensitivity, Normal, and High Throughput) have been developed on the CETAC Mercury Analyzer. Method parameters for each of these ranges are saved on a different Worksheet Template. General parameters are as follows:

<table>
<thead>
<tr>
<th>High Throughput Range</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sampling Times</strong></td>
</tr>
<tr>
<td>Integration</td>
</tr>
<tr>
<td>Read Delay</td>
</tr>
<tr>
<td><strong>Auto-Adjust Integration</strong></td>
</tr>
<tr>
<td>Replicates</td>
</tr>
<tr>
<td><strong>Instrument Control</strong></td>
</tr>
<tr>
<td>Gas Flow</td>
</tr>
<tr>
<td><strong>Autosampler Setup</strong></td>
</tr>
<tr>
<td>Sip Duration</td>
</tr>
</tbody>
</table>
### High Throughput Range

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rinse Time</td>
<td>20 s</td>
</tr>
<tr>
<td>Repeats</td>
<td>1</td>
</tr>
<tr>
<td><strong>Sample Matrix</strong></td>
<td>Liquid</td>
</tr>
<tr>
<td><strong>Reslope Frequency</strong></td>
<td>0</td>
</tr>
<tr>
<td><strong>Reslope Standard</strong></td>
<td>Standard No. 3 (check standard)</td>
</tr>
<tr>
<td><strong>Detection Limit</strong></td>
<td>0.050 µg L(^{-1})</td>
</tr>
<tr>
<td><strong>Baseline Correction</strong></td>
<td>1 point</td>
</tr>
</tbody>
</table>

### Normal Range

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sampling Times</strong></td>
<td></td>
</tr>
<tr>
<td>Integration</td>
<td>1 s</td>
</tr>
<tr>
<td>Read Delay</td>
<td>50 s</td>
</tr>
<tr>
<td><strong>Auto-Adjust Integration</strong></td>
<td></td>
</tr>
<tr>
<td>Replicates</td>
<td>4</td>
</tr>
<tr>
<td><strong>Instrument Control</strong></td>
<td></td>
</tr>
<tr>
<td>Gas Flow</td>
<td>85 mL min(^{-1})</td>
</tr>
<tr>
<td><strong>Autosampler Setup</strong></td>
<td></td>
</tr>
<tr>
<td>Sip Duration</td>
<td>30 s</td>
</tr>
<tr>
<td>Rinse Time</td>
<td>45 s</td>
</tr>
<tr>
<td>Repeats</td>
<td>1</td>
</tr>
<tr>
<td><strong>Sample Matrix</strong></td>
<td>Liquid</td>
</tr>
<tr>
<td><strong>Reslope Frequency</strong></td>
<td>0</td>
</tr>
<tr>
<td><strong>Reslope Standard</strong></td>
<td>Standard No. 3 (check standard)</td>
</tr>
<tr>
<td><strong>Detection Limit</strong></td>
<td>0.015 µg L(^{-1})</td>
</tr>
<tr>
<td><strong>Baseline Correction</strong></td>
<td>1 point</td>
</tr>
</tbody>
</table>
### Highest Sensitivity Range

<table>
<thead>
<tr>
<th>Sampling Times</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Integration</td>
<td>1</td>
</tr>
<tr>
<td>Read Delay</td>
<td>50</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Auto-Adjust Integration</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Replicates</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Instrument Control</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas Flow</td>
<td>40 mL min⁻¹</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Autosampler Setup</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sip Duration</td>
<td>60 s</td>
</tr>
<tr>
<td>Rinse Time</td>
<td>140 s</td>
</tr>
<tr>
<td>Repeats</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample Matrix</th>
<th>Liquid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reslope Frequency</td>
<td>0</td>
</tr>
<tr>
<td>Reslope Standard</td>
<td>Standard No. 3 (check standard)</td>
</tr>
<tr>
<td>Detection Limit</td>
<td>0.001 µg L⁻¹</td>
</tr>
</tbody>
</table>

| Baseline Correction     | 2 point          |

### 7.4

Prior to analysis ensure that the gas-liquid separator (GLS) post is fully wetted as follows:

#### 7.4.1
Check the bottle supplying the ASX-500 rinse station is full of 5% HNO₃.

#### 7.4.2
Use quick release mechanism and fully release clamp tension on lower two tube channels of peristaltic pump (drain channels).

#### 7.4.3
Use sample and reagent tubes and pump 5% HNO₃ and stannous chloride reagent, respectively.

#### 7.4.4
With drain pump tubes unclamped, GLS should begin to fill with 5% HNO₃.

#### 7.4.5
Allow GLS to fill until liquid level reaches top of GLS center post or until gas bubble propels a meniscus upward to wet post all along its length, including apex.
7.4.6 Upon wetting, immediately reengage quick-release clamps on drain pump tubes.
7.4.7 Do not let liquid level overflow GLS into Nafion drying tube.
7.4.8 With drain tube clamps properly reengaged and pump running, liquid level normally stops rising and goes back down
7.4.9 Once GLS has emptied, leave pump running (keep liquid flowing)
7.5 Zero analyzer to compensate for baseline offsets (due to microscopic dust buildup on the optics, dirty sample windows, and thermal drift).
7.6 Define Time Profile using Signal Profile Chart (sample time, and 1st and 2nd baseline correction points) as follows:
7.6.1 Current sample time on chart will start at dark green vertical line and stop at light red vertical line.
7.6.2 Using mouse, define sample time by clicking on chart and dragging to right.
7.6.3 Integration times will automatically be recalculated based on new total sample time.
7.6.4 Current 1st baseline correction point will start at light blue vertical line and stop at dark red line.
7.6.5 Use mouse and shift key and change point times by clicking on chart and dragging to right.
7.6.6 To do this, press and hold shift key.
7.6.7 Click with left mouse button on left limit of part of signal to be used on baseline correction point.
7.6.8 Light blue vertical cursor line will appear.
7.6.9 Without releasing shift key or left mouse button, drag mouse to right until portion of signal to be used as baseline correction point is between light blue and dark red cursor lines.
7.6.10 Release shift and mouse key and chart will update baseline times in worksheet.
7.6.11 Current 2nd baseline correction point on chart will start at light green vertical line and stop at purple line.
7.6.12 Use mouse and control key, change point times by clicking on chart and dragging to right.
7.7 Enter sample numbers, final volume, and weights in “Labels”.
7.8 Run calibration and analysis in “Analysis”.
7.9 After completion of analytical run, run RODI water through sample sipper tube and stannous chloride reagent lines. Pump all lines dry after rinsing.
7.10 Perform shutdown: turn off mercury lamp, argon gas, ASX-500, pump, M-6000 main power, close the M-6000A Software, and turn off the computer.

8. Calculations
Analytical data is reported by the instrument in µg mL\(^{-1}\) in solution. It is converted to µg kg\(^{-1}\) in soil as follows:

\[
\text{Hg in soil (µg kg}^{-1}\text{)} = \frac{[A \times B \times C \times R \times 1000]}{E}
\]

where:
- \(A\) = Sample extract reading (µg L\(^{-1}\))
- \(B\) = Extract volume (mL)
- \(C\) = Dilution, if performed
- 1000 = Conversion factor in numerator to kg-basis
- \(R\) = Air-dry/oven-dry ratio (method 3D1)
- \(E\) = Sample weight (g)

9. Report
Hg data are reported to the nearest 0.1 µg kg\(^{-1}\).

10. Precision and Accuracy
Precision and accuracy data are available from the SSL upon request.

11. References
Ground and Surface Water Analyses (4I)
Total Analysis (4I3)
  Inductively Coupled Plasma Atomic Emission Spectrophotometer (4I3a)
    Axial Mode (4I3a1)
      Ultrasonic Nebulizer (4I3a1a)
        Aluminum, Iron, Manganese, Phosphorus, and Silicon
(4I3d1a1-5)

1. Application
   Nutrients (nitrogen and phosphorus), sediments, pesticides, salts, or trace elements in ground and surface water affect soil and water quality (National Research Council, 1993). This procedure is developed for the analysis of ground or surface water. This procedure is developed for the analysis of the elemental content of ground or surface water.

2. Summary of Method
   The water is filtered and acidified with HCl. Two calibration standards plus a blank are prepared for elemental analysis. An inductively coupled plasma atomic emission spectrophotometer (ICP–AES) in axial mode is used to determine the concentration of Al, Fe, Mn, P, and Si (mg L⁻¹) by methods 4I3d1a1-5, respectively.

3. Interferences
   Spectral and matrix interferences exist. Interferences are corrected or minimized by using an internal standard. Also, careful selection of specific wavelengths for data reporting is important. Samples and standards are matrix matched to help reduce interferences.

4. Safety
   Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated acids to the fume hood. Wash hands thoroughly after handling reagents.

5. Equipment
   5.1 Syringe filters, 0.45-µm diameter, Whatman, Clifton, NJ
   5.2 Tubes, 50-mL, with caps
   5.3 Volumetrics, 500-mL and 200 mL, class A glass
   5.4 Containers, 500-mL, polypropylene, with screw caps
   5.5 Pipettors, electronic digital, Rainin Instrument Co., Woburn, MA, 2500 µL and 10 mL
5.6 Inductively coupled plasma atomic emission spectrophotometer (ICP–AES), Perkin-Elmer Optima 3300 Dual View (DV), Perkin-Elmer Corp., Norwalk, CT.
5.7 Computer, with WinLab software, ver. 4.1, Perkin-Elmer Corp., Norwalk, CT, and printer
5.8 Compressed gasses, Argon (minimum purity = 99.996%) and Nitrogen (minimum purity = 99.999%)
5.9 Autosampler, AS-90, Perkin-Elmer Corp., Norwalk, CT.
5.10 Quartz torch, alumina injector (2.0 mm id), Ultrasonic Nebulizer, Model U-5000AT+, CETAC Corp., Omaha, NE

6. Reagents
6.1 Deionized, reverse osmosis (RODI) water, ASTM Type 1 grade of reagent water
6.2 Concentrated hydrochloric acid (HCl), 12 N. Use trace-pure grade that contains low levels of impurities.
6.3 Primary standards: 1000 mg L\(^{-1}\), High Purity Standards, Charleston, SC. Single elemental standards are manufactured in dilute HNO\(_3\), HNO\(_3\)+HF, or H\(_2\)O
6.4 Internal standard: HNO\(_3\), trace pure

7. Procedure
7.1 Water sample is filtered into a 50-mL tube and capped. If extracts are not to be determined immediately after collection, then store samples at 4 °C. Analyze samples within 72 h.
7.2 The working calibration standards (WCS) are made from dilution of the primary standards. The low and high WCS for Al and Si are prepared as follows:
   7.2.1 ALLO is 0.5, 2.5, and 2.5 mg L\(^{-1}\) of Al, Fe, and Mn, respectively. To a 500-mL volumetric flask, add 0.25, 1.25, and 1.25 mL of the Al, Fe, and Mn primary standard (1000 mg L\(^{-1}\)), respectively.
   7.2.1 ALHI is 1.0, 5.0, and 5.0 mg L\(^{-1}\) of Al, Fe, and Mn, respectively. To a 500-mL volumetric flask, add 0.50, 2.50, and 2.50 mL of the Al, Fe, and Mn primary standard (1000 mg L\(^{-1}\)), respectively.
   7.2.1 SILO is 0.5 and 0.5 mg L\(^{-1}\) of Si and P, respectively. To a 500-mL volumetric flask, add 0.25 and 0.25 mL of the Si and P primary standard (1000 mg L\(^{-1}\)), respectively.
   7.2.1 SIHI is 1.0 and 1.0 mg L\(^{-1}\) of Si and P, respectively. To a 500-mL volumetric flask, add 0.50 and 0.50 mL of the Si and P primary standard (1000 mg L\(^{-1}\)), respectively.
7.3 Samples are treated with 0.5 mL HCl for each 10 mL water.
7.4 A 10 mg L$^{-1}$ Lu internal standard (read at 291.138 nm) is added to the blank, all calibration standards, and samples. It is prepared by adding 5.0 mL Lu primary standard (1000 mg L$^{-1}$) and 10 mL conc. HNO$_3$ to a 500 ml volumetric flask, and diluting to volume with RODI water.

7.5 Use the ICP–AES spectrophotometer in axial mode to analyze elements. Use ultrasonic nebulization of sample. Internal standard is added via an external peristaltic pump at 15% pump speed using 0.44 mm id. pump tubing. Internal standard and samples or standards are mixed via a mixing block and coil prior to entering the ultrasonic nebulizer. Typically, no initial dilutions of samples because of high concentrations are necessary prior to analysis. Perform instrument checks (Hg alignment; BEC and %RSD of 1 mg L$^{-1}$ Mn solution) prior to analysis as discussed in operation manual of instrument. Check instrument alignment and gas pressures to obtain optimum readings with maximum signal to noise ratio.

7.6 Analyses are generally performed at two or more wavelengths for each element. The selected wavelengths are as follows: (reported wavelength listed first and in boldface):

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mn</td>
<td>260.570, 257.610, 403.075</td>
</tr>
<tr>
<td>P</td>
<td>178.221, 213.620, 214.910</td>
</tr>
<tr>
<td>Al</td>
<td>308.215, 167.022, 396.153</td>
</tr>
<tr>
<td>Fe</td>
<td>238.203, 239.562, 259.939</td>
</tr>
<tr>
<td>Si</td>
<td>251.611, 212.412, 288.158</td>
</tr>
<tr>
<td>Lu</td>
<td>291.138 (Internal Standard)</td>
</tr>
</tbody>
</table>

7.7 Use the blank standard solution to dilute those samples with concentrations greater than the high standard. Rerun all elements and use only the data needed from the diluted analysis.

7.8 Establish detection limits using the blank standard solution. The instrumental detection limits are calculated by using 3 times the standard deviation of 10 readings of the blank. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are reported as “ND” or non-detected.

8. Calculations

With the HCl treatment (0.5 mL per 10 mL water) in the calculations, the concentrations are then reported directly, unless additional dilutions are performed because of high analyte concentrations.
9. Report
Data are reported to the nearest 0.01 mg L$^{-1}$.

10. Precision and Accuracy
Precision and accuracy data are available from the KSSL upon request.

11. References

SOIL BIOLOGICAL AND PLANT ANALYSIS (6)

Soil Analyses (6A)
Sample Preparation (1B3b2b1)
0.5 M $K_2SO_4$ Extraction + Heating with Disodium Bicinchoninic Reagent (6A1)
UV-Visible Spectrophotometer, Dual Beam (6A1a)
Hot Water Extractable Organic Carbon (6A1a1)
Air-Dry, <2 mm (6A1a1a1)

1. Application
Hot water soluble soil carbohydrates are thought to be primarily extra-cellular polysaccharides of microbial origin. They help bind soil particles together into stable aggregates. Water stable aggregates reduce soil loss through erosion, increase organic matter, and nutrient content. They also occur as part of the fast or labile organic carbon pool in soils. This labile pool contains the most available carbon for plant, animal and microbial use. The hot water soluble organic C makes up from 4 to 10% of the microbial biomass C determined by chloroform fumigation. It also makes up about 6 to 8% of the total carbohydrate content in the soil. This pool is the most easily depleted of the three organic C pools (Joergensen et al., 1996; Haynes and Francis, 1993).

2. Summary of Method
Water is added to a 10-g soil sample and autoclaved at 1 h at 121 °C. Extractable carbohydrates are measured by adding disodium bicinchoninic (BCA) reagent to a 0.5 M $K_2SO_4$ soil extract, heating to 60 °C for 2 h, cooling, and reading the absorbance at 562 nm using a spectrophotometer. Glucose is used as a standard and results expressed as glucose-C. Data are reported as mg glucose equivalent-carbon kg$^{-1}$ soil by method 6A1a1.
3. Interferences

Carbohydrates from non-microbial sources can be avoided by using the BCA reagent that is selective for microbial carbohydrates. 0.5 M $K_2SO_4$ extracts of soil are usually supersaturated with $CaSO_4$, with the excess $CaSO_4$ precipitating during storage, especially if samples are frozen and during heating of the extract with BCA reagent (Joergensen et al., 1996). Sodium hexametaphosphate is added to buffer to prevent $CaSO_4$ precipitation (Joergensen et al., 1996). Aspartic acid is used to chelate $Cu^{2+}$, preventing undesirable oxidation and so improving precision (Sinner and Puls, 1978).

4. Safety

Wear safety glasses when preparing solutions and handling soil extracts. Use oven mitts, tongs, and other devices to avoid contact with hot water and instruments used.

5. Equipment

5.1 Electronic balance, ±1.0-mg sensitivity
5.2 Steam sterilizer, 121 °C-capability
5.3 Erlenmeyer flasks, 125-mL
5.4 Filter paper, Whatman 42, 150 mm
5.5 Pipette, electronic digital, 10,000 µL, with tips, 10,000 µL
5.6 Test tubes, 10-mL
5.7 Hot water bath, 60 °C capability
5.8 Cuvettes, plastic, 4.5-mL, 1-cm light path, Daigger Scientific
5.9 Spectrophotometer, UV-Visible, Varian, Cary 50 Conc, Varian Australia Pty Ltd.
5.10 Computer with Cary WinUV software, Varian Australia Pty Ltd., and printer

6. Reagents

6.1 Reverse osmosis deionized water (RODI), ASTM Type I grade of reagent water
6.2 0.5 $M K_2SO_4$ solution. In a 1-L volumetric, dissolve 87.135 g $K_2SO_4$ (dried for 2 h at 110 °C) in RODI water. Dilute to volume and invert to mix thoroughly.
6.3 Stock standard glucose solution (SSGS), 20.0 mg glucose L⁻¹. In 1-L of 0.5 $M K_2SO_4$ solution, dissolve 0.02 g glucose (dextrose). Dilute to volume and invert to thoroughly mix. Store in polyethylene containers. Make fresh weekly. Store in a refrigerator.
6.4 Solution 1: Add 4 g Na$_2$CO$_3$, 4 g sodium hexametaphosphate [(NaPO$_3$)$_6$], and 0.2 g DL-aspartic acid in 100 mL RODI water. pH solution to 11.25 with NaOH.
6.5 Solution 2: Dissolve 0.48 g bicinchoninic acid in 12 mL RODI water (0.1 M).
6.6 Solution 3: Dissolve 1 g CuSO$_4$ in 25 mL RODI water (0.25 M).
6.7 Disodium bicinchoninic (BCA) reagent: Mix 100 mL Solution 1, 12 mL Solution 2, and 1.8 mL of Solution 3 = BCA reagent. Store in polyethylene containers. Make fresh daily. Store in a refrigerator.
6.8 Standard glucose working solutions (SGWS), 10.0, 5.0, 2.5, 1.25, 0.75, and 0.375 mg glucose L$^{-1}$. In six test tubes, add 5 mL RODI water. Perform six serial dilutions. Begin as follows: add 5 mL of SSGS to Tube 1 and shake (10.0 mg glucose L$^{-1}$) and extract 5 mL from Tube 1, add to Tube 2, and shake (5.0 mg glucose L$^{-1}$). Proceed to make all six SGWS.
6.9 Standard glucose calibration solutions (SGCS). Add 2 mL of each SGWS to a separate test tube, followed by 2 mL BCA reagent. Blank = 2 mL RODI water and 2 mL BCA reagent.

7. Procedure
7.1 Weigh 10 g of <2-mm (sieved), air-dry soil to the nearest mg and place into a 125-mL Erlenmeyer flask. If soil is highly organic, weigh 2 g of fine-grind material to the nearest mg.
7.2 Add RODI water to soil at a 1:4 ratio (10 g to 40 mL water or 2 g soil to 8 mL water).
7.3 Autoclave 1 h at 121 °C and 15 psi. Cool and filter.
7.4 Pipette 2 mL of each sample extract, 0.5 mL K$_2$SO$_4$, and 2 mL disodium BCA reagent into test tubes.
7.5 Place all tubes (SGCS, blank, and samples) in hot water bath for 2 h at 60 °C.
7.6 Allow to cool and transfer sample extract and SGCS to cuvettes.
7.7 Set the spectrophotometer to read at 562 nm. Autozero with calibration blank.
7.8 Calibrate the instrument by using the SGCS. The data system will then associate the concentrations with the instrument responses for each SGCS. Rejection criteria for SPCS, if $R^2 <0.99$.
7.9 Run samples using calibration curve. Sample concentration is calculated from the regression equation.
7.10 If samples are outside calibration range, dilute sample extracts with extracting solution and re-analyze.

8. Calculations
Convert extract glucose equivalent (mg L$^{-1}$) to glucose equivalent-carbon in the soil (mg kg$^{-1}$) as follows:

$$\text{Soil glucose-C (mg kg}^{-1}) = \frac{[(AxBxCxRx0.40 \times 1000)]}{E}$$
where:
A = Sample reading (mg L$^{-1}$)
B = Extract volume (L)
C = Dilution, if performed
R = Air-dry/oven dry ratio (method 3D1)
0.40 = Mass fraction C in glucose
1000 = Conversion factor to kg-basis
E = Sample weight (g)

9. Report
Report data as to the nearest 0.1 mg glucose equivalent-carbon kg$^{-1}$ soil.

10. Precision and Accuracy
Precision and accuracy data are available from the SSL upon request.

11. References

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Soil Analyses (6A)
Sample Preparation (1B3b2b1)

Acid Dissolution (6A3)
1 N HCl + FeCl$_2$ (6A3a)
CO$_2$ Analysis (6A3a1)
Gas Chromatography (6A3a1a)
Carbonates (6A3a1a1)
Air-Dry, <2 mm (6A3a1a1a)

1. Application
Methods involving determination of CO$_2$ have usually been preferred for measuring soil carbonate (Loeppert and Suarez, 1996). CO$_2$ released can be measured gravimetrically (Allison, 1960; Allison and Moodie, 1965), titrimetrically
(Bundy and Bremmer, 1972), manometrically (Martin and Reeve, 1955; Presley, 1975), volumetrically (Dreimanis, 1962), spectrophotometrically by infrared spectroscopy, or by gas chromatography (Loeppert and Suarez, 1996). The SSL routinely determines the amount of carbonate in the soil by treating the CaCO$_3$ with HCl, with the evolved CO$_2$ measured manometrically (methods 4E1a1a1a1-2 for <2-mm and 2- to 20-mm bases, respectively). The method herein describes soil carbonate by acid decomposition and CO$_2$ analysis by gas chromatography. This method is more commonly used in soil biochemical and biology studies, where organic C in soils with carbonates may be more precisely determined by subtracting the total carbonates (inorganic C) from total C (method 4H2a1).

2. Summary of Method

Soil carbonate is determined by chromatographic analysis of CO$_2$ evolved upon acidification of soil in a closed system of known headspace. Ferrous iron (FeCl$_2$) is added to the acid as an anti-oxidant, and the dilute acid solution (1N HCl) is chilled before addition to soil to minimize the decarboxylation of organic matter by the acid. Data are reported as mg CO$_2$-C per g of soil to the nearest 0.1 g (6A3a1a1). These data can be used to estimate soil organic carbon by subtracting (CO$_2$-C x 0.2727) from total carbon (4H2a1).

3. Interferences

It is essential that precautions be taken to ensure that there is no interference from organic matter oxidation (Loeppert and Suarez, 1996). This procedure may be more appropriate for soils with relatively low amounts of carbonates (<15%).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when handling acids. Thoroughly wash hands after handling acids. Use the fume hood when diluting concentrated HCl. Use the safety showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Canning jars, 1-qt (0.984-L), with lids fitted with gas-sampling septa
5.2 Syringe, 20-mL, with 18-gauge needle
5.3 Disposable syringe, 1-mL, for gas sampling
5.4 Needle, 18 or 20 gauge, for venting jars
5.5 Electronic balance, ±0.01-g sensitivity
5.6 Gas chromatograph (GC) with thermal conductivity detector (TCD)
5.7 Beaker, glass, 600-mL
5.8 Filter paper, Whatman 42, 150 mm
5.9 Stirrer
6. Reagents

6.1 Reverse osmosis deionized (RODI) water, ASTM Type I grade of reagent water

6.2 1N HCl with 3.1 g FeCl₂ added per 100 mL of solution, chilled to 4 to 5 °C. Prepare 500 mL of solution of as follows: weigh 15.5 g FeCl₂ into 600-mL beaker; add 20-mL RODI water; stir (medium speed) until all crystals have dissolved; filter solution to remove insoluble ferric iron particulates; transfer to 500-mL volumetric; add 41.7 mL concentrated HCl to aqueous FeCl₂; bring to volume with RODI water; and chill solution to 4 to 5 °C.

7. Procedure

7.1 Determine water content of soil sample so results may be expressed on an oven-dry soil basis. Refer to water retention methods (3C).

7.2 Weigh 5g of <2-mm (sieved), air-dry soil to the nearest 0.01 g into clean 1-qt jar.

7.3 Tightly seal jar with lid fitted with gas sampling septum. Also include one or more jars with no soil added to serve as reagent controls and CO₂ background indicators.

7.4 Add 20 ml of RODI water (room temperature) to jar through septum with 20-mL syringe, venting jar with 18 or 20 gauge needle.

7.5 Add 20 mL of chilled 1N HCl+FeCl₂ solution through septum as in Section 7.4 (do not vent), let stand 2 h, swirling soil-acid solution occasionally.

7.6 Take gas sample for gas chromatograph analysis using 1-mL syringe. Purge syringe several times before withdrawing final sample for analysis.

8. Calculations

Calculation of Soil Carbonate mass from CO₂% volumetric concentrations (Kettler and Doran, 1995).

8.1 Reaction equation:

\[
\text{CaCO}_3 + 2\text{HCl (aq)} \rightarrow \text{CO}_2 \uparrow + \text{H}_2\text{O} + \text{CaCl}_2 \text{(aq)}
\]

\[
\text{mg CO}_2\text{-C produced} = \text{mg CO}_3\text{−C dissolved by acid in soil}
\]

\[
\text{mg CO}_2\text{-C produced/OD} = \text{mg CO}_3\text{-C/OD}
\]

where:

OD = Oven-dry soil (g)

8.2 Net jar headspace

Volume of Empty 1-qt. jar = 978 cm³ (measured by H₂O volume displacement).
Soil Solid Volume (cm³) = \[\frac{\text{g Moist Soil}}{1 + \% \text{H}_2\text{O}/100}\] / (Particle Density)

Soil Solid Volume (cm³) = \(\frac{5.00 \text{ g sample}}{1 + \% \text{H}_2\text{O}/100}\) / (2.65 g cm⁻³)

where:
2.65 = assumed particle density (g cm⁻³)

Net jar Headspace (cm³)

= Empty Jar – Soil Solid Volume – Soil H₂O Volume – Liquid Volume
= 978 cm³ – (Soil Solid Volume, cm³) – (g Oven-dry Soil/1 + % H₂O/100) – (20 cm³ H₂O + 20 cm³ Acid)
= 978 cm³ – 1.89 cm³ – 40 cm³
= 936.1 cm³ (936 to 935.4 cm³ for 3–30% H₂O)

where:
5.00 = g soil oven-dry basis, assumed

8.3 Gaseous CO₂-Carbon produced:
This step is important for soils with >15% carbonates.

mg CO₂-Carbon produced by soil carbonate decomposition and detected as CO₂ in vapor space
= 4.594 mg CO₂-Carbon/atm \times ((\text{Mole % CO}_2s \times \text{P}_{ts}, \text{atm}) – (\text{Mole % CO}_2b \times \text{P}_{tb}, \text{atm}))

where:
\text{P}_{tb} = \text{Total pressure in blank jar by electronic manometer}
\text{P}_{ts} = \text{Total pressure in sample jar by electronic manometer}
\text{Mole % CO}_2b = \text{Mole % CO}_2 in blank jar by GC
\text{Mole % CO}_2s = \text{Mole % CO}_2 in sample jar by GC

8.4 CO₂-Carbon produced but dissolved in solution:

mg CO₂-Carbon produced by soil carbonate decomposition but dissolved in solution
= (0.163 mg CO₂-C/atm) \times ((\text{Mole % CO}_2s \times \text{P}_{ts}, \text{atm}) – (\text{Mole % CO}_2b \times \text{P}_{tb}, \text{atm}))

Total mg CO₃-Carbon/OD = Total mg CO₂-Carbon/OD
= \{[[0.163 mg CO₂-C/atm] \times ((\text{Mole % CO}_2s \times \text{P}_{ts}, \text{atm}) – (\text{Mole % CO}_2b \times \text{P}_{tb}, \text{atm}))]+[4.594 mg CO₂-Carbon/atm \times ((\text{Mole % CO}_2s \times \text{P}_{ts}, \text{atm}) – (\text{Mole % CO}_2b \times \text{P}_{tb}, \text{atm}))]/\{FM/FMOD}\}
\[= \frac{[4.757 \text{ mg CO}_2\text{-C/FM/atm}][(\text{Mole }\%\ CO_2\text{s} \times \text{Pts, atm}) - (\text{Mole }\%\ CO_2\text{b} \times \text{P}_{tb}, \text{ atm})]} \times (\text{FMOD})}{\text{OD}=\text{Oven-dry soil (g)}} \]

where:

\[\text{FM}=\text{Field-moist soil (g)}\]
\[\text{FMOD}=\text{Field-moist soil/oven-dry ratio (g/g) (method 3D2)}\]

8.5 Convert mg CO\text{2-C} per g of soil to percent CO\text{2-C} in soil as follows:

\[(\text{mg CO}_2\text{-C/g soil}) \times \left(\frac{1\text{g}}{1000 \text{ mg}}\right) \times (100 \text{ g soil}) = \text{Percent CO}_2\text{-C in soil}\]

9. Report

Report % CO\text{2-C} in soil to the nearest 0.1%. These data can be used to estimate soil organic carbon by subtracting (CO\text{2-C} \times 0.2727) from total carbon (4H2a1).

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References


Kettler, T., and J. Doran. 1995. Determination of soil carbonate concentration by acid decomposition and GC CO\text{2 analysis. USDA–ARS, University of Nebraska, Lincoln (unpublished).}


Particulate Organic Matter and C-Mineral (6A4)
Sample Preparation (1B3b2a1)
Total Analysis (6A4a)
Dry Combustion (6A4a1)
Thermal Conductivity Detector (6A4a1a)
Carbon, Nitrogen, Sulfur (6A4a1a1-3)
Air-Dry (6A4a1a1-3a)
≥53 µm, Particulate Organic Matter (6A4a1a1-3a1)
<53 µm, C-Mineral, Analyzed (6A4a1a1-3a2)

1. Application

Particulate organic matter (POM) is a physical fraction of the soil >53 µm in diameter (Elliott and Cambardella, 1991; Cambardella and Elliott 1992; Follett and Pruessner, 1997). Some researchers combine this fraction with the fast or labile pool. Others have described this pool as slow, decomposable, or stabilized organic matter (Cambardella and Elliott, 1992). To avoid confusion, this fraction may best described as representing an intermediate pool with regards to decomposition. This fraction is similar to various sieved and physical fractions such as the resistant plant material (RPM) (Jenkinson and Rayner, 1977), and size fractions (Gregorich et al., 1988), and variously determined light fractions of the soil organic matter (Strickland and Sollins, 1987; Hassink, 1995).

Under tillage, the POM fraction becomes depleted (Jenkinson and Rayner, 1977; Cambardella and Elliott, 1992). Reductions of more than 50% have been found in long-term tillage plots (20 yr.). Measurable reductions are believed to occur in the range of 1 to 5 years (Cambardella and Elliott, 1992).

When paired samples are selected either in time or between two tillage treatments a comparison can be made to determine the impact of the tillage practice. POM can be used in soil organic matter modeling, as a soil quality indicator and as an indicator of the SOM that can move into the active C pool.

Since the late 1970’s several models have been developed to estimate the dynamics of organic matter in the soil. All of these models have at least two phases, slow and rapid. In measuring these two phases chemical fractionation (humic and fulvic acids) has been found to be less useful than physical fractionation (Hassink, 1995). Examples of some of these models can be found in Jenkinson and Rayner (1977), tests of the CENTURY Soil Organic Model, (Parton et al., 1994; Metherell et al., 1993; and Montavalli et al., 1994). A minimum data set for soil organic carbon is proposed by Gregorich et al. (1994) that includes POM as one of the primary parameters.

2. Summary of Method

The procedural steps described herein encompasses the physical separation (1B3b2a) of the soil organic matter (<2 mm) into two fractions: (1) ≥53-µm, POM
and (2) <53 µm, C-Mineral (C-Min) (Cambardella and Elliot, 1992; Follett and Pruessner, 1997) and the analysis of these two fractions for total C, N, and S by method 6A4a1a1-3, respectively. Typically, this procedure is determined on the A horizons (Soil Survey Staff, 1999) because detectable levels of both C and N are most likely to occur in this horizon.

3. Interferences

In some weathered soils there is approximately the same amount of C in both fractions. To date, no research has been done to establish the interpretation of this result. Charcoal in native sod that has been historically burned if residence time was to be determined from the two fractions, does not affect the POM determination and C and N analysis themselves.

4. Safety

Always wear safety glasses when working with glass containers.

5. Equipment

5.1 Pressure regulator for water, with stop cock attached to tubing
5.2 Sieve, 10 mesh, 2 mm
5.3 Sieve, 270 mesh, 53 µm
5.4 Mechanical reciprocating shaker, 200 oscillations min⁻¹, 1½ strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI
5.5 Glass, Pyrex, pie or round cake pans
5.6 Evaporating crucibles
5.7 Drying Oven (110 °C)

6. Reagents

6.1 Reverse osmosis deionized water (RODI), ASTM Type I grade of reagent water

7. Procedure

Physical Separation of Organic Matter (POM and C-Min)

7.1 Remove large pieces of organic matter, roots, plant residue, gravel, wood material from an air dry soil sample. Do not shake. Save the large fraction and weigh.
7.2 Weigh 10 g of <2-mm (sieved), air-dry soil to the nearest mg into a 125-mL Erlenmeyer flask. Add 30 mL of RODI water to sample.
7.3 If the soil has carbonates, take a sub-sample and measure the inorganic carbon following the gas chromatograph method (6A3a1a1).
7.4 Stopper sample tightly and shake for 15 h (overnight) at 200 oscillations min\(^{-1}\) at room temperature (20 °C ±2 °C).

7.5 Sieve the soil through a 53-µm sieve. The POM fraction will remain on top of the sieve and the C-min will be a slurry that is collected in a pie pan underneath the sieve. Use the regulator that is attached to the RODI water to rinse the POM with a steady gentle stream of water. Keep rinsing until the water that comes through the sieve is clear. Capture all the soil slurry and water (C-Min fraction) that passes the sieve.

7.6 Label and tare a glass pie pan and an evaporating crucible to the nearest 0.01 g.

7.7 Transfer the POM into an evaporating crucible by rinsing the sieve, including the sides, with a small amount of RODI water.

7.8 Transfer the C-Min slurry into a glass pie pan for drying. Rinse the contents of the pie pan into the labeled glass pie pan.

7.9 Dry the two fractions (POM, C-Min) in an oven at 110 °C oven. The C-Min fraction may require 48 h to dry, depending on how much water was used to rinse the sample.

7.10 Once the samples are dry, let them cool briefly and record the weight to the nearest 0.1 mg.

7.11 Transfer the entire contents for each fraction to an appropriately labeled scintillating vial.

**Total Carbon, Nitrogen, and Sulfur Analysis**

7.12 Determine total C, N, and S for POM and C-min fractions (6A4a1a1-3), using fine-grind samples (=180 µm). Refer to 4H2a1-3 for the remaining procedural steps for 6A4a1a1-3.

8. Calculations

Calculate POM-C and C-Min using soil bulk density values determined by methods 4A or ASTM method D-2167 (American Society of Testing and Materials, 2004) and total C values determined by methods 6A4a1a1.

8.1 \(\frac{\% \text{Total C of POM Fraction}}{100} \times \frac{\text{POM (g)}}{10} \times \text{FMOD} = \text{POM-C g/g soil}\),

where:

\[\text{FMOD} = \text{Field-moist/oven-dry ratio (method 3D2)}\]

8.2 \(\frac{\% \text{Total C of C-Min Fraction}}{100} \times \frac{\text{C-Min (g)}}{10} \times \text{FMOD} = \text{C-Min g/g soil}\)

8.3 \(\frac{\% \text{Total C of POM fraction}}{100} \times \frac{\text{POM (g)}}{10} \times \text{Bulk Density} \times 100,000 \times \text{Depth} = \text{POM-C kg ha}^{-1} \text{ at given depth interval}\)

8.4 \(\frac{\% \text{Total C of C-Min fraction}}{100} \times \frac{\text{C-Min (g)}}{10} \times \text{Bulk Density} \times 100,000 \times \text{Depth} = \text{C-Min kg ha}^{-1} \text{ at given interval depth}\).
Use the above equations for similar computations of $N$ and $S$ (6A4a1a2-3, respectively).

9. Report

Report POM-C and C-Min in kg ha$^{-1}$ at a given depth interval (cm). Report separately similarly calculated values for $N$ and $S$. Report the percent >2-mm fraction.

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References


Fumigation Incubation (6B)
Sample Preparation (1B3b1a1)
Gas Chromatography (6B1)
CO₂ Analysis (6B1a)
Microbial Biomass (6B1a1)
Field-Moist, <2 mm (6B1a1a1)

2 M KCl Extraction (6B2)
Automatic Extractor (6B2a)
Ammonia—Salicylate (6B2a1)
Flow Injection, Automated Ion Analyzer (6B2a1a)
N as NH₃ (Mineralizable Nitrogen) (6B2a1a1)
Field-Moist, <2 mm Fumigated and <2 mm Non-Fumigated (6B2a1a1a1-2)

1. Application
Soil microorganisms are an important component of soil organic matter. One of their functions is to breakdown non-living organic matter in the soil. A variety of methods exist to measure the biomass of living soil microbes. The method described herein by the chloroform fumigation incubation (Jenkinson and Powlson, 1976) with modifications, and CO₂ evolution measurement by gas chromatography. Mineralizable N may also be determined on microbial biomass.

2. Summary of Method
A freshly collected soil sample is weighed into two separate vials. One sample is fumigated using chloroform and the other is used as a control (non-fumigated). After fumigation both the fumigated and non-fumigated samples are brought up to 55% water filled pore space (WFPS) (Horwath and Paul, 1994). Both samples are placed in a sealed container and aerobically incubated for 20 days. During this incubation period it is assumed that normal respiration occurs in the control sample container. The fumigated sample having a large carbon source for food, supplied from the dead microorganisms, has a higher CO₂ production. At the end of 10 days respiration readings are taken on both the control and the fumigated
sample to determine the amount of CO₂ evolved by gas chromatography. The CO₂ level of the control sample is also measured at the end of 20 days. CO₂ produced by biomass flush (g CO₂-C/g of soil) and soil microbial biomass (kg C/ha for a given depth interval) are reported by method 6B1a1. Mineralizable N by 2 M KCl extraction may also be determined on microbial biomass using a flow injection automated ion analyzer by method 6B2a1a. Mineralizable N is reported as mg N kg⁻¹ soil as NH₃.

3. Interferences
The determination of CO₂ evolution by gas chromatography gives a rapid and accurate measurement and can be used in acidic soils. However, this technique is prone to error in neutral and alkaline soils (Martens, 1987), as accumulation of carbonate species in the soil solution can lead to lowered CO₂ determinations (Horwath and Paul, 1994).

4. Safety
All fumigation work needs to be conducted in an adequate fume hood because chloroform has carcinogenic-volatile properties. Never determine residual chloroform by sense of smell. Make sure the vacuum pump is maintained to ensure proper operations.

5. Equipment
5.1 Face Shield
5.2 Goggles
5.3 Rubber Apron
5.4 Rubber Gloves
5.4 Chloroform spill kit
5.6 Fume hood, 100-fpm face velocity
5.7 Vacuum chambers, fiberglass
5.8 Incubator, 25 °C
5.9 Vacuum pump, 26 and 14 in Hg, organic/oil free
5.10 Mason jars, 1-qt, with lids and septa
5.11 Vials, 60-mL glass, with snap caps, for samples
5.12 Beakers, 100-mL
5.13 Refrigerator, for sample and titrate storage
5.14 Sieve, 10 mesh, 2 mm
5.15 Electronic balance, ±0.01-g sensitivity
5.16 Electronic balance, ±1.0-mg sensitivity
5.17 Oven, 110 °C
5.18  Permanent marker
5.19  Paper Towels
5.20  Tongs, 12 in
5.21  Mechanical vacuum extractor, 24-place, Sampletek, Mavco Industries, Lincoln, NE
5.22  Tubes, 60-mL polypropylene, for extraction tubes
5.23  Rubber tubing, 3.2 ID x 1.6 OD x 6.4 mm (⅛ ID x ⅛ OD x 1 in) for connecting syringe barrels
5.24  Containers, polycon
5.25  Aluminum weighing pans, 60 mm diameter x 15 mm depth
5.26  Gas chromatograph (GC) with thermal conductivity detector (TCD)
5.27  GC syringes, 1-mL

6. Reagents
6.1  Reverse osmosis deionized (RODI), ASTM Type I grade of reagent water
6.2  Chloroform stabilized in amylene. Use purified chloroform within 3 weeks.
6.3  Helium, compressed gas

7. Procedure
7.1  Until sample preparation and analysis, keep soils moist and refrigerated.
7.2  Weigh soil to the nearest 0.01 g for bulk density determination. Remove a 15 to 20 g sample for water content. Sieve moist soil to <2 mm. Sub-sample (10 to 15 g) for post-sieve water content. Refrigerate samples until analyses can be performed. Dry post-sieve water content samples at 110 °C overnight. Weigh samples the following day.
7.3  Prepare 1 sample for the Fumigated Day 10 (F 10) and 1 for the Non-Fumigated Day 20 (NF 20). Prepare replicates for each sample.
7.4  Mark the volume on the glass sample container corresponding to 20 mL (or more if a larger quantity of soil is needed).
7.5  Label the non-fumigated container clearly with permanent marker.
7.6  Use etched containers for the fumigated set, as chloroform can dissolve written labels.
7.7  Weigh enough moist soil (nearest mg) to achieve approximately 25 g (or 50 g) oven-dry soil into 60-mL glass vial. Use soil moisture content conversion. Adjust soil, by gently tapping against the counter, so that it is leveled off at the bulk density line. Carefully add RODI water with a dropper to bring the moisture up to 55% WFPS, using the moisture content conversion. Make the surface as uniformly moist as possible.
7.8  Cap the vials and refrigerate samples overnight to equilibrate.
Fumigated Samples

7.9 Line the vacuum chambers with wet paper towels to prevent desiccation.
7.10 Place the vacuum chambers in the fume hood.
7.11 Place beaker with 30 to 40 mL of stabilized chloroform in the chambers. Evacuate at 14” Hg to drive off the amylene stabilizer. A volume change will be visible, approximately 5 to 10 mL.
7.12 Place the sample vials in the vacuum chambers.
7.13 Place the pure chloroform into the pan of the vacuum chambers.
7.14 Fumigate samples for 24 h.
7.15 Evacuate the fumigated samples 4 times for approximately 15 min at 27” Hg to drive off the chloroform.

Fumigated and Non-fumigated Samples

7.16 Add 5 to 10 mL of RODI water to the bottom of the mason jar to prevent desiccation.
7.17 Seal mason jars securely with rings and lids. Lids must be airtight during the incubation. Incubate samples at 25 °C for 10 days.

Day 10 Samples

7.18 Remove mason jars from incubator. Proceed to Section 7.25 for analysis.
7.19 Remove all F 10 samples from the mason jars.
7.20 Cap F 10 samples and store at 4 °C until they can be extracted for mineralizable N (2 M KCl). If a microbial inhibitor is used, samples can be stored in the refrigerator for up to two weeks before analysis. For longer periods they should be frozen. For extraction, proceed to Section 7.28.
7.21 Incubate NF 20 samples at 25 °C for 10 days. Make sure the mason jar lids are still sealing. If not, replace with new lids

Day 20 Samples

7.22 Remove mason jars from incubator. Proceed to Section 7.25 for analysis.
7.23 Remove all NF 20 samples from the mason jars.
7.24 Cap samples and store at 4 °C until they can be extracted for mineralizable N (method 6B3a1a1). If a microbial inhibitor is used, samples can be stored in the refrigerator for up to two weeks before analysis. For longer periods they should be frozen. For extraction, proceed to Section 7.28.

Gas Chromatography

7.25 Measure the CO₂ accumulated in the headspace of the mason jars by gas chromatography.
7.26 Refer to manufacturer’s manual for operation of the gas chromatograph.

7.27 Calibration curves and retention times for gas under analysis is established by analyzing the certified standard gas mixture (1% CO\textsubscript{2}) by the procedure used for analysis of the sample. Flow rate is 30 mL min\textsuperscript{-1}. Monitor the baseline prior to analysis.

2 M KCl Extraction

7.28 Mix fumigated replicates together. Mix non-fumigated replicates together.

7.29 Weigh 5 g of moist soil to the nearest mg into 60-mL polypropylene extraction tubes. Tube will need to be tapped and rinsed with 2 M KCl in order to get the moist soil to bottom of tube.

7.30 Set-up vacuum extractors. Add 25 mL of 2 M KCl to extraction tubes. Extract for 1 h.

7.31 Following extraction, transfer contents of tubes into polycon containers. Proceed with determining mineralizable N. Also analyze N in reagent RODI water as blanks. Refer to 4D10a1a1 for the remaining procedural steps for 6B2a1a1.

8. Calculations

8.1 Calculate the bulk density of 25 g of <2-mm, field-moist soil in a 100-mL beaker manually compressed to 20 cm\textsuperscript{3} volume.

\[
Db_1 (g/cm^3) = \frac{25 g}{(1 + H_2O_f)} / 20 cm^3
\]

where:

- \(Db_1\) = Bulk density (g cm\textsuperscript{-3})
- \(H_2O_f\) = Field water content (g g\textsuperscript{-1})
- \(H_2O_f\) is determined by methods 4B.

8.2 Calculate the gravimetric water content [Gravimetric H\textsubscript{2}O\textsubscript{0.55} (g/g)] required for soil to be at 55% water filled pore space (WFPS), using calculated \(Db_1\) in Section 8.1 and an assumed particle density of 2.65 g cm\textsuperscript{-3}:

\[
Gravimetric H_2O_{0.55} (g/g) = \frac{0.55 \times [1 - (Db_1 / 2.65)]}{Db_1}
\]

where:

- Gravimetric H\textsubscript{2}O\textsubscript{0.55} (g/g) = 55% water filled pore space (WFPS)
- 2.65 = Assumed particle density (g cm\textsuperscript{-3})

8.3 Calculate the additional gravimetric water needed for soil to reach 55% WFPS:

\[
H_2O_{add} = ([Gravimetric H_2O 0.55 (g/g)] - \text{(H}_2\text{O}_f\text{)}) \times [W / (1 + H_2O_f)]
\]
8.4 Determine the CO$_2$-Carbon produced by biomass flush during 10 day incubation.

8.4.1 At 10 days, determine the mole % concentration of CO$_2$ in head space of jar by GC analysis.

8.4.2 Calculate the CO$_2$-Carbon produced by biomass flush as follows:

\[
\text{CO}_2\text{-Carbon produced by biomass flush, g BioCO}_2\text{-Carbon/g FM Soil} = \\
[(\text{mole } %\text{CO}_2, 10 \text{ days, fumigated} - \text{mole } %\text{CO}_2, 10 \text{ days, non-fumigated}) \times (0.47 \text{ g CO}_2\text{-C})] / W
\]

where:
- FM = Field-moist soil (g)

8.5 Determine the difference between fumigated CO$_2$-Carbon produced during the 10 to 20 day incubation period and the non-fumigated CO$_2$-Carbon produced during the same period.

8.5.1 At 20 days, determine the mole % concentration of CO$_2$ in head space of jar by GC analysis.

8.5.2 Calculate the (fumigated – non-fumigated) CO$_2$-Carbon produced during the 10 to 20 day incubation period as follows:

\[
\text{(fumigated – non-fumigated) CO}_2\text{ -Carbon produced during the 10 to 20 day incubation period, g BioCO}_2\text{-Carbon/g FM Soil} = \\
\{(\text{mole } %\text{CO}_2, 20 \text{ days, fumigated} - \text{mole } %\text{CO}_2, 10 \text{ days, fumigated}) \\
- (\text{mole } %\text{CO}_2, 20 \text{ days, non-fumigated} - \text{mole } %\text{CO}_2, 10 \text{ days, non-fumigated})\} \times (0.47 \text{ g CO}_2\text{-C})] / W
\]

8.6 Calculate the Soil Biomass Flush (kg CO$_2$-C/ha), using the bulk density value (Db$_2$) determined by method 4A or ASTM method D-2167:

\[
\text{Soil biomass flush (kg CO}_2\text{-Carbon/ha)} = (\text{g BioCO}_2\text{-Carbon/g FM Soil}) \times (\text{g FM Soil/g OD soil}) \times (\text{Db}_2 \times \text{g OD soil/cm}^3 \text{ FM Soil}) \times (1 \text{ kg CO}_2\text{-Carbon/1000 g CO}_2\text{-Carbon}) \times (100,000,000 \text{ cm}^2/\text{ha}) \times (\text{layer thickness, cm})
\]

8.7 Calculate Soil Microbial Biomass (kg C/ha for a given depth interval):

\[
\text{Soil Biomass Flush (kg C/ha)/0.41}
\]

0.41 = $K_c$, fraction of biomass C mineralized to CO$_2$ (Anderson and Domsch, 1978)
8.8 Calculate (fumigated – non-fumigated) mineralizable N (mg kg\(^{-1}\)):

\[
\text{Fumigated N – Non-fumigated N} = \frac{[(F_1 - F_2) \times A \times B \times E \times 1000]}{C} - \frac{[(N_1 - N_2) \times D \times F \times E \times 1000]}{G}
\]

where:

- \(F_1\) = Analyte reading, fumigated (mg L\(^{-1}\))
- \(F_2\) = Blank reading, reagent RODI Water, fumigated (mg L\(^{-1}\))
- \(A\) = Extract volume, fumigated (L)
- \(B\) = Dilution, fumigated (if performed)
- \(C\) = Sample weight, fumigated (g)
- \(N_1\) = Analyte reading non-fumigated sample extract (mg L\(^{-1}\))
- \(N_2\) = Blank reading, reagent RODI water, non-fumigated (mg L\(^{-1}\))
- \(D\) = Extract volume, non-fumigated (L)
- \(F\) = Dilution, non-fumigated (if performed)
- \(G\) = Sample weight, non-fumigated (g)
- \(E\) = Field-moist/oven-dry ratio (method 3D2)
- 1000 = Conversion factor to kg-basis

9. Report

Report CO\(_2\) produced by biomass flush (g CO\(_2\)-C/g of soil) and soil microbial biomass (kg C ha\(^{-1}\) for a given depth interval). Report the difference between mineralizable N of fumigated and non-fumigated to the nearest mg N kg\(^{-1}\) soil as NH\(_3\).

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References


Plant Analyses (6C)
   Sample Preparation (1B3b3a1a1, 1B3b3b1a1)
Root Biomass (6C1)
   Plant (Above-Ground) Biomass (6C2)
Plant Nutrition (6C3)
   Total Analysis (6C3a)
      Dry Combustion (6C3a1)
         Thermal Conductivity Detector (6C3a1a)
            Carbon, Nitrogen, and Sulfur (6C3a1a1-3)
            Dry (50°C), Roots (6C3a1a1-3a1)
            Dry (50°C), Plant Material (Above-Ground) (6C3a1a1-3a2)

1. Application
   Root biomass in the upper 4 inches of the soil is an input value for the Revised Universal Soil Loss Equation (RUSLE) (Renard et al., 1997). The mass, size, and distribution of roots in the near surface are among the most important factors in determining the resistance of the topsoil to water and wind erosion. Root biomass is also one of the major Carbon pools found in soil. Commonly, root mass and plant residue in the soil form between 3,000 (annual crop) and 15,000 (perennial grasses) lbs/ac/yr soil biomass (Harwood et al., 1998). Above-ground biomass (production) represents annual yield and can be measured following the protocols found in the National Range and Pasture Handbook (USDA–NRCS, 1997). Root biomass represents biomass from more than one year.

   The development of new roots and ultimately the decomposition of roots within the soil is a major contributor to the Soil Organic Carbon (SOC) pool. In this way, plant roots also contribute to the fertility of soils by slowly releasing macro- and micro-nutrients back into the soil.

   Root biomass and SOC help bind the soil together by forming aggregates and granular structure. This improves the tilth as well as the erosion resistance of soil. Depending upon the root turnover rate (known for some species), climate, and residue decomposition rate (known for some areas, based on climate and soil moisture status) the amount of Carbon stored in the soil can be determined from the root biomass, plant residue, and SOC.
Root biomass is frequently used to calculate root/shoot ratios in order to evaluate the health and vigor of plants, and determine the success of establishment of seeded plants at the 4-leaf stage.

Dried roots can be fine-grind, and total C, N, P, and S can be determined. The C/N ratio can also be determined, which is typically different from the C/N ratio of the above ground plant material. Low levels of N in the soil will promote root growth over top growth (Bedunah and Sosebee, 1995). The C/N ratio of roots, plant residue in the soil, and SOC each contribute to the residue decomposition rate for soils. Low C/N values lead to more rapid decomposition, high C/N levels slow decomposition. The C/N ratio required for decomposition of plant residue, without a net tie-up of N, is approximately 25:1. Plant residue from young legumes commonly has a C/N ratio of 15:1. Plant residue from woody materials commonly is 400:1 (Harwood et al., 1998). The C/N ratio of soil microbes is quite variable but commonly falls between 15:1 and 3:1 (Paul and Clark, 1989).

Root biomass/horizon can be paired with the description of roots in each soil horizon (i.e. few fine, many very fine, etc.) in the pedon description and thus a qualitative estimate can be made of the mass in each size fraction of roots. This automated method for determining root biomass also includes some plant residue. Woody material is removed and weighed separately. Because root biomass determined in this manner includes plant residue, it can be used to estimate the soil plant residue pool in most models (Jenkinson and Rayner, 1977; Metherell et al., 1993).

2. Summary of Method

The procedural steps described herein encompass the physical separation of roots and plant residue from a soil sample using an automated root washer (1B3b3); these weights recorded for root (6C1) and plant biomass (6C2); and these fractions analyzed for total C, N, and S by method 6C3a1a1-3, respectively.

3. Interferences

The soil must be dispersed for successful separation of the roots and plant residue from the soil sample. Tap water rather than distilled water should be used to help avoid puddling and dispersion problems.

4. Safety

Do not touch moving parts of the root washer when it is in operation. Avoid electrical shock by ensuring that the electrical cord is dry, and prevent the formation of pools of water near the cord.

5. Equipment

5.1 Automated root washer (after Brown and Thilenius, 1976)
5.1.1 Root cages, basket sieves, with No. 30 mesh and 0.5 mm-diameter openings
5.1.2 Garden hose
5.1.3 Sediment tank

5.2 Buckets
5.3 Analytical balance, ±0.01 g sensitivity
5.4 Drying oven (60 °C capability)
5.5 Weighing dishes
5.6 Scintillating vials
5.7 Tweezers
5.8 Drying trays

6. Reagents
6.1 Tap water
6.2 Algaecide, Bath Clear

7. Procedure

Sample Preparation

7.1 Weigh approximately 200 g of field-moist soil to the nearest 0.01 g and record the weight.
7.2 Pour all of the weighed soil into a root cage and cap it.
7.3 Immerse cage in tap water until soil disperses (overnight if samples are cloddy).

Root Washing

7.4 Make sure that machine is level and that the sediment tank is under the drain.
7.5 Load the root cages containing the soil and root slurry into the rotation bars. Be sure to load them evenly. If not using all of the rotation bar slots, load into every other slot.
7.6 Fill the washing tank with water to the top of the bottom cage.
7.7 Add 10 drops of algaecide to the washing tank. Attach machine to water source.
7.8 Turn on the water at the faucet then turn on the machine’s spray nozzle. Do not start the machine with the lid open. Once the rotator has started, turn on spray nozzles.
7.9 Depending upon the number of samples, let the machine run from 40 to 90 min. (Ex: 12 samples usually take about 60 min.)
Clean Up and Maintenance

7.10 Upon completion of sample washing, shut down the sprayer first then the rotator. Drain the machine first by opening the bottom plug. Make sure the sediment tank is under the drain. After the machine is drained, let the water in the sediment tank settle. Replace plug in the machine.

7.11 Drain the sediment tank water off. Collect the sediment out of the machine and the sediment tank and properly dispose of it.

7.12 Flush out all of the sediment in the machine over the sediment tank. Repeat procedure until the machine is completely clean.

7.13 Clean the entire area. Run water down the drain for about 30 min after everything is clean.

Root/Plant Material Separation and Drying

7.14 Air-dry roots and plant material at room temperature overnight while still in the sieve cages.

7.15 Remove the roots/plant residue in the cage by tapping them. Brush out any roots/plant residue that clings to the side of the sieve cages.

7.16 Add water to a tray of roots/plant material. Float off as much of the organic matter as possible by adding water to a tray roots/plant residue. Much of the organic fraction will be less dense than the sand particles that are not removed during root washing. Pour floating matter into root cage to trap roots/plant residue; avoid introduction of inorganic portion into cage.

7.17 If roots/plant material remain in the inorganic fraction, use tweezers to remove as much of it as possible and return it to the cage.

7.18 Air dry at room temperature overnight all material in cage. Next day, tap and brush the air-dry material into a tray.

7.19 Remove the woody material, dry at 50 °C in an oven overnight, and record weight of woody material.

7.20 Separate plant residue from roots, dry at 50 °C in an oven overnight, and record weights of plant residue and roots.

7.21 Place the roots and plant residue into separate scintillation vials.

Total Carbon, Nitrogen, and Sulfur Analysis

7.22 Determine total C, N, and S for roots and plant material (6C3a1a1-3), using fine-grind samples (≈180 µm). Refer to 4H2a1-3 for the remaining procedural steps for 6C3a1a1-3.
Separating Roots and Organic Matter Residue (picking)

7.23 Following initial air-drying, use tweezers and separate organic matter residue from roots using tweezers. Roots are usually light colored, and organic residue is usually darker colored.

7.24 Place the organic residue and roots on separate tared watch glasses and re-dry and weigh.

7.25 Record each individual weight for plant residue and roots. Subtract the tare weights and record the total weight of air-dry roots and the total weight of air-dry plant residue. Report separately root biomass and plant residue rather than just roots including some organic residue.

8. Calculations

Calculate root biomass using soil bulk density values determined by methods (3B) described in this manual or ASTM method D-2167 (American Society for Testing and Materials, 2004).

\[
\text{Root biomass/ha for soil layer of given thickness (kg ha}^{-1}\) = \\
\left[\frac{\text{Dry Roots (g)}}{\text{Total sample weight (g)}} \times \text{(Bulk density: g OD soil/ cm}^3\text{ FM soil)} \times (\text{g FM soil/g OD soil)} \times (1 \text{ kg/1000 g)} \times (100,000,000 \text{ cm}^2/ \text{ha}) \times (\text{Layer thickness, cm})\right]
\]

where:
OD=Oven-dry
FM=Field-moist

9. Report

Report root biomass as kg ha\(^{-1}\) at a given depth interval (cm). If plant residue was separated from roots, report each separately.

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References


MINERALOGY (7)

Instrumental Analyses (7A)
X-Ray Diffractometer (7A1)
  Thin Film on Glass, Resin Pretreatment II (7A1a)
    Mg Room Temperature, Mg Glycerol Solvated, K 300°, K 500° C (7A1a1)

1. Application

Clay fractions of soils are commonly composed of mixtures of one or more phyllosilicate minerals together with primary minerals inherited directly from the parent material (Olson et al., 1999). Positive identification of mineral species
and quantitative estimation of their proportions in these polycomponent systems usually require the application of several complementary qualitative and quantitative analyses (Whittig and Allardice, 1986; Amonette and Zelazny, 1994; Wilson, 1994; Moore and Reynolds, 1997). One of the most useful methods to identify and to make semiquantitative estimates of the crystalline mineral components of soil is x-ray diffraction analysis (Hughes et al., 1994; Kahle et al., 2002). Quantification of a mineral by x-ray diffraction requires attention to many details, including sample (slide) size relative to the incident x-ray beam, thickness and particle size uniformity of sample, and beam-sample orientation (Moore and Reynolds, 1997). More complex quantification procedures include using standard additions, full pattern fitting, and determining mineral intensity factors (Kahle et al., 2002). At best, quantification can approach a precision of ±5% and an accuracy of ±10 to 20% (Moore and Reynolds, 1997).

The operational strategy at the SSL and the preceding Lincoln SSL has been to base mineral quantification on first order peak intensities. Semi-quantitative interpretations have been held consistent over time (1964 to the present) by adjusting instrumental parameters (e.g., scan speed) to maintain a constant peak intensity for an in-house reference clay standard and subsequently soil samples. The intent is to keep interpretations consistent from sample to sample.

2. Summary of Method

Soils are dispersed and separated into fractions of interest. Sands and silts are mounted on glass slides as slurries, on a smear of Vaseline, or on double sticky tape for analysis. Clay suspensions are placed on glass slides to dry and to preferentially orient clay minerals. Most samples of soil clays contain fewer than 7 minerals that require identification. The soil clay minerals of greatest interest are phyllosilicates, e.g., kaolinite, mica (illite), smectite, vermiculite, hydroxy-interlayered vermiculite, smectite, hydroxy-interlayered smectite and chlorite.

Diffraction maxima (peaks) develop from the interaction of x-rays with planes of elements that repeat at a constant distance (d-spacing) through the crystal structure. Generally, no two minerals have exactly the same d-spacings in three dimensions and the angles at which diffraction occurs are distinctive for a particular mineral (Whittig and Allardice, 1986; Moore and Reynolds, 1997). Phyllosilicates (or layer silicate minerals) have very similar structures except in the direction perpendicular to the layers (c-dimension). Several treatments are needed to sort out which minerals are present. Glycerol is added to expand smectites. Ionic saturation and/or heat treatments are used to collapse some 2:1 layer silicates and dehydroxylate kaolinite, gibbsite, and goethite, eliminating characteristic peaks.

The crystal “d” spacings of minerals, i.e., the interval between repeating planes of atoms, can be calculated by Bragg’s Law as follows:
\[ n\lambda = 2d \sin \theta \]

Where:
- \( n \) = integer that denotes order of diffraction
- \( \lambda \) = x-radiation wavelength (Angstroms, Å)
- \( d \) = crystal "d" spacing (Å)
- \( \theta \) = angle of incidence

When \( n = 1 \), diffraction is of the first order. The wavelength of radiation from an X-ray tube is constant and characteristic for the target metal in the tube. Copper radiation (CuK\( \alpha \)) with a wavelength of 1.54 Å (0.154 nm) is used at the SSL. Because of the similar structure of layer silicates commonly present in soil clays, several treatments that characteristically affect the "d" spacings are necessary to identify the clay components. At the SSL, four treatments are used, i.e., Mg\(^{2+}\) (room temperature); Mg\(^{2+}\)-glycerol (room temperature); K\(^+\) (300 °C); and K\(^+\) (500 °C).

Standard tables to convert \( \theta \) or 2\( \theta \) angles to crystal d-spacings are published in the U.S. Geological Survey Circular 29 (Switzer et al., 1948) and in other publications (Brown, 1980). Through the years hardware has been updated and the recording of data has evolved from a strip chart recorder through several kinds of electronic software. X-ray by this method (7A1a1) is semiquantitative.

### 3. Interferences

Interstratification of phyllosilicate minerals causes problems in identification. These interstratified mixtures, differences in crystal size, purity, chemical composition, atomic unit cell positions, and background or matrix interferences affect quantification (Moore and Reynolds, 1997; Kahle et al., 2002). No pretreatments other than ionic saturation and dispersion with sodium hexametaphosphate are used for separation and isolation of the clay fraction in the routine procedure. Impurities, such as organic matter, carbonates, and iron oxides, may act as matrix interferences causing peak attenuation during X-ray analysis or may interfere with clay dispersion and separation. Pretreatments to remove these impurities serve to concentrate the crystalline clay fraction and may increase accuracy, but also potentially result in degradation of certain mineral species (e.g., smectites) as well as loss of precision (Hughes et al., 1994).

The separation (centrifuge) procedure used to isolate the clay fraction from the other size fractions of the soil skews the <2-µm clay suspension toward the fine clay, but it minimizes the inclusion of fine silt in the fraction. Sedimentation of the clay slurry on a glass slide tends to cause differential settling by particle size (i.e., increasing the relative intensity of finer clay minerals).

Dried clay may peel from the XRD slide. One remedy is to rewet the peeled clay on the slide with 1 drop of glue-water mixture (1:7). Other remedies are:
a) Place double sticky tape on the slide prior to re-wetting the dried clay with the glue-water mixture.

b) Dilute the suspension if thick.

c) Crush with ethanol and dry, and then add water to make a slurry slide.

d) Roughen the slide surface with a fine-grit sandpaper.

An optimum amount of glycerol on the slides is required to solvate the clay, i.e., to expand smectites to 18 Å. X-ray analysis should be performed 1 to 2 days after glycerol addition. If excess glycerol is applied to the slide and free glycerol remains on the surface, XRD peaks are attenuated. Some suggestions to dry the slides and achieve optimum glycerol solvation are as follows:

a) Use a chamber such as a desiccator (with no desiccant) to dry slide, especially when the clay is thin.

b) If the center of slide is whitish and dry, usually with thick clay, brush slide with glycerol or add an additional drop of glycerol.

4. Safety

Operate the centrifuge with caution. Keep the centrifuge lid closed when in operation. Ensure that all rotors and tubes are seated firmly in proper location. Use tongs and appropriate thermal protection when operating the muffle furnace. The diffraction unit presents an electrical and radiation hazard. Analysts must receive radiation safety training before operating the equipment. Employees must wear a radiation film badge while in the room when the diffraction unit is in operation.

5. Equipment

5.1 Teaspoon (5 g)

5.2 Dispenser, 5 mL, for sodium hexametaphosphate solution

5.3 Centrifuge, International No. 2, with No. 240 head and carriers for centrifuge tubes, International Equip. Co., Boston, MA

5.4 Centrifuge tubes, plastic, 100 mL, on which 10-cm solution depth is marked

5.5 Rubber stoppers, No. 6, for centrifuge tubes

5.6 Mechanical reciprocating shaker, 100 oscillations min⁻¹, 1½ in strokes, Eberbach 6000, Eberbach Corp., Ann Arbor, MI

5.7 Plastic cups, 60 mL (2 fl. oz.) with lids

5.8 Label printer

5.9 Hypodermic syringes, plastic, 12 mL, with tip caps

5.10 Screen, 80 mesh, copper

5.11 Dropper bottle, plastic, 30 mL (1 fl. oz.), for a 1:7 glycerol:water mixture

5.12 Muffle furnace
5.13 X-ray diffractometer, Bruker 5000-Dmatic, with X-Y autosampler that accommodates 66 samples or standards, Bruker AXS Inc., Madison, WI
5.14 Computer, Diffract\textsuperscript{plus} EVA software, release 2000, Bruker AXS Inc., Madison, WI, and printer
5.15 XRD slides, glass, 2.54 X 2.54 mm (frosted glass slides used for K-treated samples)
5.16 XRD sample preparation board, wood, with 32 places for glass XRD slides
5.17 Slide holder
5.18 Reference slides: quartz and clay from reference soil

6. Reagents
6.1 Reverse Osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate (NaPO\textsubscript{3})\textsubscript{6} and 7.94 g of sodium carbonate (Na\textsubscript{2}CO\textsubscript{3}) in 1 L RO water.
6.3 Potassium chloride (KCl), 1.0 N. Dissolve 74.60 g KCl in 1 L RO water or 671.40 g KCl in 9 L RO water.
6.4 Magnesium chloride (MgCl\textsubscript{2}), 1.0 N. Dissolve 47.61 g MgCl\textsubscript{2} in 1 L RO water or 428.49 g MgCl\textsubscript{2} in 9 L RO water.
6.5 Glycerol:water mixture (1:7). Add 4 mL of glycerol to 28 mL RO water plus 2 drops of toluene.
6.6 Exchange resin, Rexyn 101 (H), analytical grade. Pretreatment of resin as follows:
6.6.1 Divide equally Rexyn 101 (H), approximately 250-g portions, into two 600-mL beakers labeled K and Mg and add appropriate salt solution (1.0 N KCl or 1.0 N MgCl\textsubscript{2}). Cover resin with salt solution.
6.6.2 Stir, let settle for 10 min, decant clear solution, and add salt solution. Repeat 3 times. Leave resin covered in salt solution for 8 to 12 h.
6.6.3 Repeat step 6.6.2 on a second day. Resin is ready for syringes. Saturated resin not used initially for syringes can be saved for future use.
6.7 White glue, diluted 1:7 with RO water

7. Procedure

**Preparation (Recharge) of Resin-Loaded Syringes**

7.1 Place a small circle of 80-mesh screen in a 12-mL syringe and add 4 cm\textsuperscript{3} of exchange resin from which salt solution has been drained. The procedure
requires 2 Mg and 2 K slides for each sample, so two sets of syringes are prepared.

7.2 Saturate the resin in each of the syringes with 4 mL of the appropriate 1.0 N salt solution (MgCl₂ or KCl) and expel. Repeat saturation of resin. Individual steps follow.

7.3 Fill syringe completely with the salt solution and allow to equilibrate for 4 to 20 h.

7.4 Rinse syringe twice with 4 mL of RO water and rinse tip cap.

7.5 Completely fill syringe with RO water and allow to equilibrate for 4 to 20 h.

7.6 Rinse syringe twice with RO water.

7.7 Expel water, cap syringe, and store.

**Preparation of Clay Suspension**

7.8 Print run sheets using LIMS. Each run consists of 8 samples with 4 treatments for each sample.

7.9 Label each 100 ml plastic centrifuge tube with a sample number from the run sheet.

7.10 Place ≈5 g (1 tsp) of air-dry <2-mm soil in a 100-mL plastic centrifuge tube. If the sample appears to be primarily sand, use 10 g (2 tsp) of <2-mm soil to obtain sufficient clay.

7.11 Add 5 mL of sodium hexametaphosphate dispersion agent. If the soil contains gypsum or is primarily calcium carbonate, use 10 mL of sodium hexametaphosphate dispersing agent.

7.12 Fill tube to 9.5-cm height with RO water and close with a stopper.

7.13 Place the tubes in a mechanical shaker and shake overnight (at least 4 hours).

7.14 Remove stopper from tube and rinse stopper and sides of tube with enough water to bring the volume to the 10-cm mark.

7.15 Balance the pairs of tubes and place in centrifuge. Centrifuge at 750 rpm for 3.0 min.

7.16 If the clay is dispersed, carefully decant 30 mL of suspension into a labeled, 60-mL, plastic cup and cover with lid.

7.17 If the clay did not disperse after being shaken overnight, decant and discard the clear supernatant. Then add an additional 10 mL sodium hexametaphosphate and sufficient RO water to bring the level up to 9.5 cm depth. Repeat Sections 7.13 to 7.16.

7.18 Clay suspension is used for X-ray diffraction analysis. It can be dried and used for elemental or thermal analysis.
Thin Film on Glass, Resin Pretreatment

7.19 The SSL uses sample boards that hold 32 slides each, i.e., 8 samples x 4 treatments. Place run number on the sample board. Prepare the sample board with glass XRD slides to receive the following 4 treatments per clay suspension sample.

- Mg$^{2+}$-room temperature
- Mg$^{2+}$-glycerol (room temperature)
- K$^{+}$-300 °C (heated 2 h)
- K$^{+}$-500 °C (heated 2 h)

7.20 Use a hypodermic syringe to place 6 drops of the glycerol:water mixture (1:7) on each Mg$^{2+}$-glycerol slide.

7.21 Draw 3 to 4 mL of the clay suspension into the Mg syringe and invert back and forth to facilitate cation exchange.

7.22 Dispense 3 drops to clear the tip.

7.23 Dispense ≈0.3 mL (6 to 10 drops) to cover the Mg and mg-glycerol XRD slides. Similarly, use the K syringes to apply clay suspension to the frosted glass K-300 and K-500 slides. Draw RO water into each syringe and expel 3 times to remove all of the clay suspension, cap and store syringes. Recharge all syringes after 10 run boards.

7.24 When the clay suspension has dried, transfer the slides with the K$^{+}$-saturated clays to the muffle furnace. Heat for a minimum of 2 hours at 300 °C, remove the K-300 batch of slides. Set the temperature to 500 °C and heat slides for a minimum of 2 hours at 500 °C. After slides are cool, return them to the run board.

X-Ray Diffraction Operation

7.25 Complete X-ray analysis of the glycerol slide within 1 to 2 days after the slide dries. If this is not possible, add additional glycerol prior to run (e.g., add 6 drops of glycerol:water mixture to dry slide 24 h prior to x-ray analysis).

7.26 Place the tray with filled sample holders in the autosampler and execute the run. Use the following parameters:

- CuK$\alpha$ radiation, $\lambda=1.54$ Å (0.154 nm)
- Scan range=2° to 35°2$\theta$
- Generator settings=40 kv, 30 ma
- Divergence slit=1°
- Receiving slit=0.2 mm
Step-size and scan-speed vary depending on intensity of X-rays generated. Settings should be adjusted to maintain the same peak intensities on the standard reference clay and quartz standard over the long term regardless of tube intensities.

7.27 In the laboratory information system (LIMS), create a batch file. Data in file is transferred to a job program on the x-ray computer software for data analysis. These data include project and sample identification. Include both the quartz and soil standard with each run.

7.28 Activate job program for analysis. The job stores raw data on the hard disk under the subdirectory designated by year, project type, project name.

7.29 Prepare and print a 4-color graphics chart. The four colors are blue (Mg$^{2+}$); green (Mg$^{2+}$-glycerol); pink (K$^+$ 300 °C); and red (K$^+$ 500 °C). File hard copies of detected peaks and graphics chart in pasteboard binders by state, county, and chronology.

7.30 Compare quartz and soil standard patterns electronically with previous runs to ensure peak intensity and positions have remained constant.

**Interpretation of X-Ray Diffraction Data**

7.31 The angle in degrees two theta ($2\theta$) measured in X-ray diffraction analyses is converted to angstroms (Å) using tables complied according to Bragg’s Law. Refer to summary of method. Angstroms convert to nanometers (nm) by a factor of 0.1, e.g., 14 Å = 1.4 nm.

7.32 Use the following X-ray diffraction criteria to identify some common crystalline minerals. The reported “d” values are for 00l basal spacings. The Miller index ($hkl$) specifies a plane or crystal face which has some orientation to the three crystallographic axes of a, b, and c. The Miller index (00l) indicates a crystal face that is perpendicular to the a and b axes (Schulze, 1989). The following X-ray diffraction criteria also have some questions (Q) that may aid the analyst in interpreting the diffraction patterns. These questions are a suggested procedural approach to help the analyst identify the relative locations of a few peaks and to confirm key criteria. For a more complete list of d-spacings for confirmation or identification of a mineral consult the “Mineral Powder Diffraction File – Data Book” (JCPDS, 1980).

**X-Ray Diffraction Criteria**

7.32.1 Kaolinite and Halloysite
- a. Crystal structure missing at 500 °C.
- b. 7 Å (7.2 to 7.5 Å) with all other treatments.
Q. Is there a 7 Å peak? Is it destroyed at 500 °C? Kaolinite or Halloysite.
Q. Is the peak sharp and at ~ 7.1 Å (but absent at 500 °C)? Kaolinite.
Q. Is the peak broad and at 7.2 to 7.5 Å (but absent at 500 °C)? Halloysite.

7.32.2 Mica (Illite)
a. 10 Å with all treatments.
b. 10 Å with Mg²⁺-saturation.
Q. Is there a 10 Å peak with Mg²⁺-saturation? Mica (Illite).

7.32.3 Chlorite
a. Crystal structure of Fe-chlorites destroyed at 650 to 700 °C.
b. 14 Å with all other treatments.
c. 14 Å at 500 °C.
d. Generally also has strong 7 Å peak.
Q. Is there a 14 Å peak when heated to 500 °C? Chlorite.

7.32.4 Vermiculite
a. 14 Å with Mg²⁺-saturation.
b. 14 Å with Mg²⁺-glycerol solvation.
c. Nearly 10 Å with K⁺ saturation.
d. 10 Å when K⁺-saturated and heated to 300 °C.
Q. Is there an enhanced 10 Å peak with K⁺-saturation in comparison to Mg²⁺-saturation that cannot be attributed to smectite? Vermiculite.

7.32.5 Smectite
a. 14 Å with Mg²⁺-saturation.
b. 12 to 12.5 Å with K⁺- or Na⁺-saturation.
c. 17 to 18 Å with Mg²⁺-glycerol solvation.
d. 10 Å with K⁺-saturation and heating to 300 °C.
Q. Is there a 17 to 18 Å peak upon solvation? Smectite.

7.32.6 Gibbsite
a. Peak at 4.83 to 4.85 Å with Mg²⁺ and Mg²⁺-glycerol but destroyed when heated to 300 °C.

7.32.7 Goethite
a. Peak at 4.16 to 4.18 Å with Mg²⁺ and Mg²⁺-glycerol but destroyed when heated to 300 °C.
7.32.8 Hydroxy-interlayered Vermiculite or Smectite
   a. Failure to completely collapse to 10 Å of smectite or
      vermiculite when K⁺-saturated and heated to 300 °C.

7.32.9 Quartz
   a. Peaks at 4.27 Å and 3.34 Å with all treatments (only 3.34 if
      small amounts).

7.32.10 Lepidocrocite
   a. Peak at 6.2 to 6.4 Å with Mg²⁺ and Mg²⁺-glycerol but destroyed
      when heated to 300 °C.

7.32.11 Potassium Feldspar
   a. Peak at 3.24 Å with all treatments.

7.32.12 Plagioclase Feldspar
   a. Twin peaks between 3.16 and 3.21 with all treatments.

7.32.13 Calcite
   a. Peak at 3.035 Å with all treatments.

7.32.14 Dolomite
   a. Peak at 2.88 to 2.89 Å with all treatments.

7.32.15 Gypsum
   a. Peak at 7.56 Å with Mg²⁺ and Mg²⁺-glycerol, but destroyed
      when heated to 300 °C.

7.32.16 Mixed Layer Vermiculite-Mica
   a. Randomly interstratified: Peak between 10 and 14 Å with Mg²⁺
      that does not expand with Mg²⁺-glycerol; peak collapses to
      10 Å with K⁺-saturation and heating to 300 °C.
   b. Regularly interstratified: A 24 Å peak (and higher orders); no
      change with Mg²⁺-glycerol treatment; K⁺ saturation and
      heating collapses vermiculite and produces a 10 Å peak.

7.32.17 Mixed Layer Smectite-Mica
   a. Randomly interstratified: Peak between 10 and 14 Å with
      Mg²⁺ that expands to 14–16 Å with Mg²⁺-glycerol; Peak
      collapses to 10 Å with K⁺-saturation and heating to 300 °C.
   b. Regularly interstratified: A small 24 Å peak and large peak
      at 12 Å with Mg²⁺-saturation; expands to 28 Å with Mg²⁺-
      glycerol treatment; K⁺-saturation and heating collapses
      smectite, then produces a 10 Å peak.

7.32.18 Mixed Layer Chlorite-Vermiculite
   a. Randomly Interstratified: Peak at 14 Å with Mg²⁺ and Mg²⁺-
      glycerol; Peak collapses incompletely to between 10 and
      14 Å with K⁺-saturation and heating.
b. Regularly interstratified: A 28 Å peak (and higher orders) with Mg-saturation; no expansion with Mg$^{2+}$-glycerol treatment; K$^+$-saturation and heating to 500 °C collapses vermiculite and a produces a 24 Å peak.

**7.32.19** Mixed Layer Chlorite-Smectite

a. Randomly interstratified: Peak at 14 Å with Mg$^{2+}$-saturation; expands to higher spacings (≈16 Å) with Mg$^{2+}$-glycerol treatment; Peak collapses incompletely to between 10 and 14 Å with K$^+$-saturation and heating.

**7.33** Use the X-ray diffraction criteria, i.e., diagnostic basal 00l spacings (Å), in Table 1 for identification and ready reference of some common crystalline minerals as affected by differentiating sample treatments.

**7.34** Preferential orientation of clay mineral samples enhances diffraction from the basal (00l) spacing and tends to minimize the number and intensity of peaks from diffraction by other hkl planes. With preferential orientation, second, third, and fourth order peaks may be recorded in addition to the basal first order peaks. Groups of associated peaks that differ by order of diffraction are as follows:

**7.34.1** Smectite (Mg$^{2+}$-glycerol):

a. 17 to 18 Å.

b. 8.5 to 9 Å (weak).

**7.34.2** Chlorite, vermiculite, and smectite:

a. 14, 7, 4.7, and 3.5 Å.

b. 7, 4.7, and 3.5 Å weak for smectite.

(Note: High Fe substitution in the chlorite structure results in a decrease in the peak intensity of odd numbered orders (e.g., 14 and 4.7 Å) and increase in peak intensity of even number orders (7 and 3.5 Å)).

**7.34.3** Mica:

a. 10, 5 (weak in biotites and moderate in muscovites), and 3.3 Å.

**7.34.4** Kaolinite:

a. 7 and 3.5 Å.

**7.35** The differentiation of kaolinite and halloysite in a sample can be aided by the use of formamide (Churchman et al., 1984). The intercalation and expansion of halloysite to a d-spacing of ≈10.4 Å is relatively rapid (20 to 30 min), whereas kaolinite expansion requires ≈4 h upon treatment. The procedure is as follows:

**7.35.1** Lightly spray formamide as an aerosol on the dried Mg$^{2+}$-saturated slide.
7.35.2 Wait 15 min but not more than 1 h and X-ray approximately 7.6 to 13.5° 2θ (d=11.6 to 6.55 Å).

7.35.3 Halloysite will expand to ≈10.4 Å, whereas kaolinite will remain unchanged.

7.35.4 Heating the sample to 110 °C for 15 min will collapse the halloysite to ≈7 Å.

7.35.5 The total amount of kaolinite and halloysite can be determined by thermal analysis. The intensity ratio of the 10.4 to 7.2 Å peaks of the formamide-treated sample can be used to determine the relative percentage of halloysite and kaolinite.

7.35.6 The total amount of kaolinite and halloysite can be determined by thermal analysis. The intensity ratio of the 10.4 to 7.2 Å peaks of the formamide-treated sample can be used to determine the relative percentage of halloysite and kaolinite.

8. Calculations

X-ray diffraction produces peaks on a chart that corresponds to 2θ angle on a goniometer. Standard tables to convert θ or 2θ to crystal “d” spacings are published in the U.S. Geological Survey Circular 29 (Switzer et al., 1948) and in other publications (Brown, 1980). The crystal “d” spacings of minerals, i.e., the interval between repeating planes of atoms, can be calculated by Bragg’s Law. Refer to summary of method.

9. Report

From the “Detected Peaks File” and graphics chart, identify the minerals present according to the registered “d” spacings. As a first approximation, use the following peak intensities, i.e., peak heights above background in counts s⁻¹, to assign each layer silicate mineral to one of the 5 semiquantitative classes.

<table>
<thead>
<tr>
<th>Class</th>
<th>Peak Height above background</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 (Very large)</td>
<td>&gt;1800</td>
</tr>
<tr>
<td>4 (Large)</td>
<td>1120 to 1800</td>
</tr>
<tr>
<td>3 (Medium)</td>
<td>360 to 1120</td>
</tr>
<tr>
<td>2 (Small)</td>
<td>110 to 360</td>
</tr>
<tr>
<td>1 (Very small)</td>
<td>&lt;110</td>
</tr>
</tbody>
</table>

Adjust class placement to reflect area under the curve if peak is broad relative to peak height or if thermal, elemental, clay activity data, or other evidence warrant.
class adjustment. If there are no peaks or no evidence of crystalline components, place the sample in NX class (noncrystalline). If there are only 1 to 3 very small (class1) peaks, also indicate NX to infer a major noncrystalline component.

10. Precision and Accuracy

X-ray by method 7A1a1 is semi-quantitative. Precision and accuracy data are available from the SSL upon request.

11. References


Instrumental Analyses (7A)
Differential Scanning Calorimetry (7A4)
Thermal Analyzer (7A4a)

1. Application
Calorimetry measures specific heat or thermal capacity of a substance. Two separate types of differential scanning calorimetry (DSC) instruments have evolved over time. The term “DSC” is most appropriate for the power-compensated-type instrument in which the difference in the rate of heat flow between a sample and a reference pan is measured as materials are held isothermal to one another using separate furnaces (Karathanasis and Harris, 1994). The DSC therefore directly measures the magnitude of an energy change ($\Delta H$, enthalpy or heat content) in a material undergoing an exothermic or endothermic reaction. Heat flow-type DSC instruments are more common and are similar in principal to differential thermal analyzers (DTA). The heat flow instruments have the sample and reference pans in a single furnace and monitor pan temperature from the conducting base. The difference in pan temperatures ($\Delta T$) results from clay mineral decomposition reactions in the sample as the furnace temperature is increased. The configuration of this instrument results in a signal that is independent of the thermal properties of the sample and $\Delta T$ can be converted to a calorimetric value via instrument calibration (Karathanasis and Harris, 1994). DSC is commonly used to quantify gibbsite ($\text{Al(OH)}_3$) and kaolinite ($\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$) in soils and clays by measuring the magnitude of their dehydroxylation endotherms which are between approximately 250 to 350 °C and 450 to 550 °C, respectively (Jackson, 1956; Mackenzie, 1970; Mackenzie and Berggen, 1970; Karathanasis and Hajek, 1982).

2. Summary of Method
An 8-mg sample of soil clay is weighed into an aluminum sample pan and placed in the DSC sample holder. The sample and reference pans are heated under flowing $\text{N}_2$ atmosphere from a temperature of 30 to 600 °C at a rate of 10 °C min$^{-1}$. Data are collected by the computer and a thermograph is plotted. Gibbsite and kaolinite are quantified by measuring the peak area of any endothermic reactions between 250 to 350 °C and 450 to 550 °C, respectively, and by calculating the $\Delta H$ of the reaction. These values are related to the measured enthalpies of standard mineral specimens (gibbsite and kaolinite). Percent kaolinite and gibbsite are reported by method 7A4a.

3. Interferences
Organic matter is objectionable because it produces irregular exothermic peaks in air or $\text{O}_2$, commonly between 300 to 500 °C, which may obscure
important reactions from the inorganic components of interest (Schnitzer and Kodama, 1977). Analysis in an inert N\textsubscript{2} atmosphere helps to alleviate this problem although thermal decomposition of organic matter is still observed. Pretreatment with H\textsubscript{2}O\textsubscript{2} may be necessary for soils with significant amounts of organic matter. Mineral salts that contain water of crystallization also may be interferences. Samples should be washed free of any soluble salts.

Use a representative soil sample as sample size is small (<10 mg). Avoid large aggregates in sample, the presence of which may cause thermal interferences because of differential kinetics of gas diffusion through the sample and physical movement of sample in a reaction.

The dehydroxylation of goethite is between 250 to 400 °C and may interfere with the identification and integration of the gibbsite endotherm (250 to 350 °C) (Mackenzie and Berggen, 1970). The dehydroxylation of illite is between 550 to 600 °C and partially overlaps the high end of the kaolinite endotherm (450 to 550 °C), resulting in possible peak integrations (Mackenzie and Caillere, 1975). The dehydroxylation of hydroxy-Al interlayers in hydroxy-Al interlayered vermiculite (HIV) is between 400 to 450 °C and may interfere with the low end of the kaolinite endotherm (450 to 550 °C), especially in the poorly crystalline kaolinites (Mackenzie and Caillere, 1975). Similarly, the dehydroxylation of nontronites, Fe-rich dioctahedral smectites is between 450 to 500 °C and may interfere with kaolinite identification and measurement (Mackenzie and Caillere, 1975).

4. Safety

Secure high pressure N\textsubscript{2} tanks and handle with care. When changing the tanks, valves should be protected with covers. Do not heat aluminum sample pans >600 °C. Aluminum melts at 660 °C, and the sample pans alloy with and destroy the DSC cell. Always use high quality purge gases with the DSC. Minimum purity of 99.9% is recommended.

5. Equipment

5.1 Thermal analyzer, DSC 910S, TA Instruments, New Castle, DE
5.2 Thermal analyzer operating system software, Thermal Analyst 2100, Version 8.10B, TA Instruments, New Castle, DE
5.3 Data analysis software, TGA Standard Data Analysis Version 4.0, TA Instruments, New Castle, DE
5.4 Computer, IBM-PC 386, TA Instruments Operating System, Version 8.10B
5.5 Thermal analyzer instrument controller (MIM), TA Instruments, New Castle, DE
5.6 Autosampler, 920 Auto DSC, TA Instruments, New Castle, DE
5.7 Printer, Hewlett Packard, HP-7440, 8-pen plotter
5.8 Two-stage gas regulators, 50 psi maximum outlet pressure
5.9   Electronic balance, ±0.1-mg sensitivity, Mettler AE160
5.10  Forceps, flat-tipped
5.11  Weighing spatula
5.12  Desiccator
5.13  Mortar and pestle
5.14  Sieve, 80 mesh
5.15  N$_2$ gas, 99.99% purity
5.17  Gibbsite, standard, Surinam Gibbsite, SSL 67L022.

6. Reagents
6.1   Magnesium nitrate saturated solution [Mg(NO$_3$)$_2$•6H$_2$O]
6.2   Ethanol

7. Procedure

   Derive <2-µm Clay Fractions

7.1   Prepare Na-saturated clay as in method 7A1a1, preparation of clay suspension, sections 7.8 to 7.19.
7.2   Dry the clay suspension and transfer to mortar. Moisten sample with ethanol and grind with pestle to make an homogeneous slurry.
7.3   Air-dry sample using flowing air in hood. Lightly grind sample with pestle to make a homogeneous powder. Transfer to original container for storage until use.
7.4   Prior to analysis, sieve sample with 80-mesh screen. Equilibrate sample overnight over a saturated magnesium nitrate solution (55% relative humidity) in a glass desiccator.

   DSC Operation

7.5   Set-up the instrument and calibrate. Refer to the manufacturer’s manual for operation of the DSC. Samples can be analyzed singly or with the autosampler for multiple samples.
7.6   Weigh ≈8 mg of sample, i.e., <80-mesh fine-earth (<2 mm) soil fraction or derived <2-µm clay fraction, into tared aluminum sample pan. Refer to section on derived <2-µm clay fractions, Steps 7.1 to 7.4.
7.7   Use flat-tipped forceps to remove aluminum sample pan from balance. Drop sample from a 4- to 5-mm height to uniformly distribute sample in pan. Return the sample pan with sample to the balance and record weight to nearest ±0.1 mg. This weight is entered into computer in appropriate menu.
7.8 Carefully place the aluminum sample pan in the center of DSC platinum sample side (front section) of sample holder.

7.9 Place empty aluminum sample pan in reference side (back section) of sample holder.

7.10 Cover the DSC cell.

7.11 The standard sample run heating program has a heating rate of 10 °C min⁻¹, a starting temperature of 30 °C, and an ending temperature of 600 °C.

7.12 Start the “Run” program.

7.13 When the run is complete, data are analyzed by entering the Data Analysis 2100 System and selecting the DSC Standard Data Analysis Program.

7.14 Display file and calculate joules g⁻¹ for the mineral endotherm.

8. Calculations

The area under a curve representing an endothermic dehydroxylation reaction is proportional to the enthalpy (ΔH) of the reaction. The enthalpy is calculated with the DSC software per g of kaolinite or gibbsite (joules g⁻¹) as appropriate.

Analyze each of the standard clays on the DSC. Calculate the enthalpy per g for the endothermic reactions of the standard kaolinite and gibbsite (joules g⁻¹).

The purity of the standard clays is evaluated via TGA (7A2a). Adjust the DSC results of the standards using the purity measurements from TGA.

Determine the amount of kaolinite and gibbsite in soil samples by dividing the enthalpy of the sample (joules g⁻¹) by the enthalpy of the standard (joules g⁻¹). Multiply this result by 100 to express as a percentage.

9. Report

Report percent kaolinite and/or gibbsite to the nearest whole number.

10. Precision and Accuracy

Precision and accuracy data are available from the SSL upon request.

11. References

Jackson, M.L. 1956. Soil chemical analysis. Advan. course. M. L. Jackson, Madison, WI.


Optical Analyses (7B)

Platy Grains (7B2)

Static Tube Separation (7B2b)

1. Application

Static charge of mineral grains to glass and a magnetic separator are used to separate platy grains from non-platy grains in the 0.02–2 mm fraction of soil. The separates are weighed to determine the quantity of platy minerals. The platy separates are examined by optical microscope and analyzed by X-ray diffraction to determine the kinds of minerals present.

2. Summary of Method

A sample of <2-mm soil is prepared according to the procedure described in 7B2a. A small portion of sample is introduced into the top of an inclined glass tube mounted on a vibrator. As the tube is rotated and vibrated, the platy grains adhere
to the tube and the non-platy grains (residue) roll or slide through. The residue is run through a magnetic separator to separate the coarser platy grains that did not adhere to the glass tube. Percent platy minerals of specific analyzed fraction are reported (7B2b).

3. Interferences

Large platy grains tend to slide through the tube into the residue, especially if the plates are stacked into a book.

4. Safety

No known hazards exist.

5. Equipment

5.1 Glass tube, 1.5 cm inside diameter, 30 cm long vibrating mechanism
5.2 Receptacles to hold grains
5.3 Funnel or glassine paper or aluminum weighing dish
5.4 Camel's hair brush
5.5 Gelatin capsules
5.6 Mechanical Vibrator

6. Reagents

None.

7. Procedure

7.1 Prepare sample (disperse, fractionate, and dry sample as described in Method 7B2a).

**Static Charge Separation by Glass Tube**

7.2 Set up vibrator as shown in figure 7B2b-1.
7.3 Weigh 0.1500 g of 0.02-2 mm material or particle-size separate onto a square of glassine paper and introduce into the upper end of the glass tube.
7.4 Turn on the vibrator until the material begins to flow. Rotate the tube slowly so the platy grains adhere to the tube wall. Adjust vibrator intensity and rotation to achieve a slow, smooth flow rate.
7.5 When rounded grains have passed through the tube, remove the tube and hold it vertically over a tared weighing dish. Tap the tube to remove the platy grains. If grains remain, wash them out with RO water, dry, and weigh.

Note: The glass tube separation can be done by hand, without mechanical vibration in the field office to obtain a fair approximation of the platy grain component.
**8. Calculations**

8.1 Percent platy grains = \( \frac{100 \times \text{(weight of platy grains)}}{\text{(sample weight)}} \)

8.2 Percent residual grains = \( \frac{100 \times \text{(weight of residual grains)}}{\text{(sample weight)}} \)

8.3 Recovery = \( \frac{\text{(weight of platy and residual grains)}}{\text{(sample weight)}} \)

**9. Report**

Report platy grains as a percent of the specific particle size fraction analyzed, oven-dried soil weight.

**10. Precision and Accuracy**

Precision and accuracy data are not available for this method.

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**Optical Analyses (7B)**

**Platy Grains (7B2)**

**Froth Flotation (7B2c)**

**1. Application**

This method used with coarse silt, very fine, fine, and medium sand fractions of soils with significant amounts of platy minerals (mica, vermiculite, chlorite, and their pseudomorphically altered weathering products). It provides weight percent data on each of the fractions. Combined use of froth flotation and magnetic
separation improve separation of platy and non-platy grains to better estimate weight percentages of components.

2. Summary of Method

Platy minerals (muscovite, biotite, vermiculite, and kaolinite) are floated off over the top of a container in an agitated aqueous suspension by action of a complexer and frother, adapted from procedure provided by Louis Schlesinger of the Minerals Research Laboratory School of Engineering, North Carolina State University, in Asheville, NC. Percent platy minerals of specific analyzed fraction are reported (7B2c).

3. Interferences

There are no known interferences.

4. Safety

There are no known safety hazards.

5. Equipment

5.1 Modified 800 mL glass beaker for mixing container
5.2 Plastic bucket, 5 qt, for catch container
5.3 Mechanical mixer–1 laboratory reagent mixer or a magnetic bar stirrer
5.4 Manual mixer–1 glass rod pH meter
5.5 Oven, 110 °C
5.6 Wood tongue depressors or a similar spatula-like device
5.7 Syringe, 1 mL
5.8 Beaker, 800 mL aerator
5.9 Ring stand (to hold Aerator assembly)
5.10 Funnel and stand to hold funnel 300-mesh sieve
5.11 Glass rod with rubber policeman

6. Reagents

6.1 Reverse osmosis (RO) water, ASTM Type III grade of reagent water
6.2 Frother: Econofroth 910 frother, 100% solution (Mfg: Nottingham Company; P.O. Box 250049; Station N; 1303 Boyd Ave. NW; Atlanta, GA 303025; phone: 404-351-3501)
6.3 Promoter: Econofloat A-50 promoter, 5% solution (1 mL/20 mL or 50 mL/1000 mL); (Mfg: Nottingham Company)
6.4 NaOH (2.5% solution or 12.5 g per 500 g) 25 g/liter
6.5 \( H_2SO_4\), 0.9 \( N \) (2.5% solution or 12.5 g conc. \( H_2SO_4 \) per 500 g water)

6.6 Ethyl alcohol in wash bottle

7. Procedure

7.1 Prepare sample (disperse, fractionate, and dry sample as described in method 7B2a.

7.2 Set up apparatus as shown in figure 7B2c-1.

![Figure 7B2c-1.—Apparatus for froth flotation.](image)

7.3 Prepare 700 mL RO water, pH 2.5: Put 800-mL glass beaker on mechanical mixer and fill to 700 mL level. Adjust to pH 2.5 (not over 3.0) with 0.9 \( N \) \( H_2SO_4 \) solution and set the full beaker aside for use later.

7.4 Place 800 mL mixing container on mechanical mixer, and fill \( \frac{3}{4} \) full with RO water.

7.5 Add 5 g of sample to water and start a strong mixing action.

7.6 Adjust pH of sample and water to 2.5 with 0.9 \( N \) \( H_2SO_4 \) solution.

7.7 Add 0.25ml of Promoter.
7.8 Add 0.2ml of frother.
7.9 Continue to mix the sample solution for 2–5 minutes.
7.10 Check and readjust the pH level as necessary.
7.11 Place mixing container in plastic 5-qt catch bucket, lower aerator into solution, fill mixing container almost to top, using the water from section 7.3.
7.12 Turn off the mechanical mixer and turn on the air to the aerator to start the frothing action. The frothing foam should build to a point where the foam pours out of the modified mixing container. Add water from section 7.3 as needed to maintain foam overflow from the mixing container.
7.13 Use a glass stirring rod with a rubber policeman to mix the sample on the bottom of the mixing chamber in a grid like motion. As the platy minerals froth to the surface use the wood tongue depressor to rake them over the side of the mixing chamber.
7.14 After 1–5 minutes the amount of platy minerals floating to the surface should diminish. At this time turn off the air, thoroughly remix the sample with the glass stirring rod, and repeat the procedure starting at section 7.12.
7.15 After 2–5 minutes during the second run through section 7.12, carefully inspect the foam to see if platy minerals are still frothing to the surface. Continue until little or no platy minerals are frothing to the surface.
7.16 Transfer contents of the catch bucket with ethyl alcohol to the 300-mesh sieve and rinse the sample.
7.17 Transfer the rinsed grains to an aluminum dish, dry in an oven at 110 °C, weigh and record the weight as platy grains.
7.18 Repeat steps 7.16 and 7.17 for the contents in the mixing container and record the weight as residual grains. Note: Delay weighing samples if magnetic separation (method 7B2a) will be done next.

8. Calculations
8.1 Percent platy grains=[100 x (weight of platy grains)]/(sample weight)
8.2 Percent residual grains=[100 x (weight of residual grains)]/(sample weight)
8.3 Recovery=(weight of platy and residual grains)/(sample weight)

9. Report
Report platy grains as a percent of the specific particle-size fraction analyzed, oven-dried soil weight.

10. Precision and Accuracy
Precision and accuracy data are not available for this method.
Table 1.—X-Ray Diffraction Parameters of Common Soil Minerals.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Treatment</th>
<th>Na⁺</th>
<th>Mg²⁺</th>
<th>Mg²⁺</th>
<th>K⁺ 300 °C</th>
<th>K⁺ 500 °C</th>
<th>K⁺ 700 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kaolinite</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>LD¹</td>
<td>LD</td>
</tr>
<tr>
<td>Halloysite</td>
<td>7B²</td>
<td>7B</td>
<td>7B</td>
<td>7B</td>
<td>7B</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Mica (Illite)</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Chlorite</td>
<td>14*³</td>
<td>14*</td>
<td>14*</td>
<td>14*</td>
<td>14*</td>
<td>14*</td>
<td>T⁴</td>
</tr>
<tr>
<td>Vermiculite</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>10</td>
<td>10</td>
<td>10</td>
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<tr>
<td>Smectite</td>
<td>12.5</td>
<td>14</td>
<td>18</td>
<td>12.5</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Gibbsite</td>
<td>4.85</td>
<td>4.85</td>
<td>4.85</td>
<td>4.85</td>
<td>LD</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Goethite</td>
<td>4.18</td>
<td>4.18</td>
<td>4.18</td>
<td>4.18</td>
<td>LD</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Interlayer</td>
<td>10-14</td>
<td>10-14</td>
<td>10-18</td>
<td>10-14</td>
<td>10-14</td>
<td>10-14</td>
<td>10-14</td>
</tr>
<tr>
<td>Quartz</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3.34 and 4.27</td>
<td>for all treatments</td>
<td></td>
</tr>
<tr>
<td>Calcite</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3.035</td>
<td>for all treatments</td>
<td></td>
</tr>
<tr>
<td>Dolomite</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2.886</td>
<td>for all treatments</td>
<td></td>
</tr>
</tbody>
</table>

¹ LD = Lattice destroyed.
² B = Broad peak is common.
³ * = Sometimes <14 Å.
⁴ T = Temperature of decomposition varies with chemical composition, particle-size, and heating conditions.
ION ANALYSES (5)

Cation Exchange Capacity (5A)
NH₄OAc, pH 7.0 (5A8)
Automatic Extractor (CEC-7)
Steam Distillation
   Kjeltec Auto 1035 Analyzer (5A8c)

1. Application

The CEC determined with 1 N NH₄OAc buffered at pH 7.0, is a commonly
used method and has become a standard reference to which other methods are
compared (Peech et al., 1947). The advantages of using this method are that
the extractant is highly buffered so that the extraction is performed at a constant,
known pH (7.0) and that the NH₄⁺ on the exchange complex is easily determined.

2. Summary of Method

Displacement after washing is the basis for this procedure. The CEC is
determined by saturating the exchange sites with an index cation (NH₄⁺); washing
the soil free of excess saturated salt; displacing the index cation (NH₄⁺) adsorbed
by the soil; and measuring the amount of the index cation (NH₄⁺). A sample is
leached using 1 N NH₄OAc and a mechanical vacuum extractor (Holmgren et al.,
1977). The extract is weighed and saved for analyses of the cations. The NH₄⁺
saturated soil is rinsed with ethanol to remove the NH₄⁺ that was not adsorbed.
Steam distillation and titration are used to determine the NH₄⁺ adsorbed on the soil
exchange complex. The CEC by NH₄OAc, pH 7 is reported in meq 100 g⁻¹ oven-
dry soil in method 5A8c.

3. Interferences

Incomplete saturation of the soil with NH₄⁺ and insufficient removal of NH₄⁺
are the greatest interferences to this method. Ethanol removes some adsorbed
NH₄⁺ from the exchange sites of some soils. Isopropanol rinses have been used
for some soils in which ethanol removes adsorbed NH₄⁺. Soils that contain large
amounts of vermiculite can irreversibly “fix” NH₄⁺. Soils that contain large amounts
of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Nessler’s reagent contains mercury, which is toxic. Proper disposal of the Nessler’s reagent and clean-up of equipment in contact with the reagent is necessary.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjeltec Auto 1035 Analyzer.

5. Equipment

5.1 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.2 Mechanical vacuum extractor, Mavco Sampletek, 5300 N. 57th St., Lincoln, NE
5.3 Syringes, polypropylene, disposable, 60 mL, for extraction vessel, extractant reservoir and tared extraction syringe
5.4 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels
5.5 Polycons, Richards Mfg. Co.
5.6 Kjeltec Auto 1035/1038 Sampler System, Tecator, Perstorp Analytical Inc.
5.7 Digestion tubes, straight neck, 250 mL
5.8 Analytical filter pulp, ash-free, Schleicher and Schuell, No. 289
5.9 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.10 Electronic balance, ±1-mg sensitivity

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Ammonium acetate solution (NH₄OAc), 1 N, pH 7.0. Add 1026 mL of glacial acetic acid (CH₃COOH) to 15 L DDI water. Add 1224 mL of concentrated
ammonium hydroxide (NH₄OH). Mix and cool. Dilute with DDI water to 18 L and adjust to pH 7.0 with CH₃COOH or NH₄OH.

6.3 Ethanol (CH₃CH₂OH), 95%, U.S.P.

6.4 Nessler’s reagent. Add 4.56 g of potassium iodide (KI) to 30 mL DDI water. Add 5.68 g of mercuric iodide (HgI₂). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL of DDI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with DDI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately. Store in brown bottle to protect from light.

6.5 Sodium chloride (NaCl), reagent, crystal.

6.6 Antifoam agent, slipicone release spray, Dow Chemical Corp. Alternatively, use n-octyl alcohol.

6.7 Boric acid, 4% (w:v), with bromcresol green-methyl red indicator (0.075 % bromcresol green and 0.05% methyl red), Chempure Brand

6.8 Hydrochloric acid (HCl), 0.05 N, standardized. Dilute 83 mL of concentrated HCl in 20 L of DDI water.

6.9 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of DDI water. Dilute to 9 L with DDI water.

7. Procedure

Extraction of Bases

7.1 Prepare extraction vessel by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh 2.50 g of <2-mm, air-dry soil and place in an extraction vessel. Weigh a smaller amount of sample, if the soil is highly organic. Prepare one quality control check sample per 48 samples.

7.3 Place extraction vessel on upper disk of the extractor and connect a tared extraction syringe. Use a 25.4-mm (1 in) length rubber tubing and insert the plunger in the slot of the stationary disk of the extractor.

7.4 Use a squeeze bottle to fill extraction vessel to the 20-mL mark with NH₄OAc solution (≈10 mL). Thoroughly wet the sample. Let stand for at least 20 min.

7.5 Put reservoir tube on top of the extraction vessel. Rapidly extract the NH₄OAc solution to a 0.5- to 1.0-cm height above sample. Turn off extractor. Add ≈45 mL of NH₄OAc solution to the reservoir tube. Set extractor for an overnight (12 to 16 h) extraction.

7.6 Next morning turn off the extractor. Pull the plunger of the syringe down. Do not pull plunger from the barrel of the syringe. Carefully remove the
syringe containing the extract. Leave the rubber tubing on the extraction vessel. Weigh each syringe containing the NH₄OAc extract to the nearest 0.01 g.

7.7 Mix the extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. The solution in the polycon is reserved for analyses of extracted cations (methods 6N2, 6O2, 6P2, and 6Q2).

**Removal of Excess Ammonium Acetate**

7.8 Return the extractor to starting position. Attach syringe to the extraction vessel and rinse the sides of the extraction vessel with ethanol from a wash bottle. Fill the extraction vessel to the 20-mL mark with ethanol and let stand for 15 to 20 min.

7.9 Place reservoir tube on the extraction vessel. Rapidly extract the ethanol level to a 0.5- to 1.0-cm height above the sample. Turn off the extractor and add 55 to 60 mL of ethanol to the reservoir. Extract at a 45-min rate.

7.10 After the extractor has stopped, turn off the switch. Pull the plunger of the syringe down. Do not pull the plunger from the syringe barrel. Remove the syringe and discard the ethanol.

7.11 Repeat the ethanol wash.

7.12 After the second wash, collect a few drops of ethanol extract from the extraction vessel on a spot plate. Test for NH₄⁺ by using Nessler's reagent. A yellow, red to reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler's reagent. Repeat until a negative test is obtained.

**Steam Distillation: Samples and Reagent Blanks**

7.13 Remove the extraction vessel and transfer the sample to a 250-mL digestion tube. Add 6 to 7 g of NaCl to the digestion tube.

7.14 Perform the same transfer and addition of reagents for blanks as for samples.

7.15 Spray silicone antifoam agent (or 2 drops of n-octyl alcohol solution) into the digestion tubes for each of the samples and reagent blanks.

7.16 When using new reagents, e.g., boric acid, reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.17 On bench worksheet, record the normality of standardized acid, i.e., ≈0.05 N HCl.
7.18 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

$$\text{CEC} = \frac{[\text{Titer} \times N \times 100 \times \text{AD/OD}]}{[\text{Sample Weight (g)-tooltip}]}$$

where:

- CEC = Cation Exchange Capacity (meq 100 g⁻¹)
- Titer = Titer of sample (mL)
- N = Normality of HCl titrant
- 100 = Conversion factor to 100-g basis
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report CEC-7 in units of meq 100 g⁻¹ of oven-dry soil to the nearest 0.1 meq 100 g⁻¹.

10. Precision

Precision data are not available for this procedure.

11. References


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Cation Exchange Capacity (5A)

$\text{NH}_4\text{Cl (5A9)}$

**Automatic Extractor**

- Steam Distillation (5A9c)
- Kjeltec Auto 1035 Analyzer (5A9c)

1. Application

The CEC determined with a neutral unbuffered salt, e.g., 1 N $\text{NH}_4\text{Cl}$, is an estimate of the “effective” CEC (ECEC) of the soil (Peech et al., 1947). For a soil with a pH of <7.0, the ECEC value should be < CEC measured with a buffered solution at pH 7.0. The $\text{NH}_4\text{Cl}$ CEC is ≈ equal to the $\text{NH}_4\text{OAc}$ extractable bases plus the KCI extractable Al for noncalcareous soils.
2. Summary of Method

Displacement after washing is the basis for this procedure. The CEC is determined by saturating the exchange sites with an index cation (NH₄⁺); washing the soil free of excess saturated salt; displacing the index cation (NH₄⁺) adsorbed by the soil; and measuring the amount of the index cation (NH₄⁺). A sample is leached using 1 N NH₄Cl and a mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed and saved for analyses of the cations. The NH₄⁺ saturated soil is rinsed with ethanol to remove the NH₄⁺ that was not adsorbed. Steam distillation and titration are used to determine the NH₄⁺ adsorbed on the soil exchange complex. The CEC by NH₄Cl is reported in meq 100 g⁻¹ oven-dry soil in method 5A9c.

3. Interferences

Incomplete saturation of the soil with NH₄⁺ and insufficient removal of NH₄⁺ are the greatest interferences to this method. Ethanol removes some adsorbed NH₄⁺ from the exchange sites of some soils. Isopropanol rinses have been used for some soils in which ethanol removes adsorbed NH₄⁺. Soils that contain large amounts of vermiculite can irreversibly “fix” NH₄⁺. Soils that contain large amounts of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Nessler’s reagent contains mercury, which is toxic. Proper disposal of the Nessler’s reagent and clean-up of equipment in contact with the reagent is necessary. Thoroughly wash hands after handling reagents. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjelteck Auto 1030 Analyzer.

5. Equipment

5.1 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE

5.2 Mechanical vacuum extractor, Mavco Samultiplek, 5300 N. 57th St., Lincoln, NE
5.3 Syringes, polypropylene, disposable, 60 mL, for extraction vessel, extractant reservoir, and tared extraction syringe
5.4 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in), for connecting syringe barrels
5.5 Polycons, Richards Mfg. Co.
5.6 Kjeltec Auto 1035/1038 Sampler System, Tecator, Perstorp Analytical Inc.
5.7 Digestion tubes, straight neck, 250 mL
5.8 Analytical filter pulp, ash-free, Schleicher and Schuell, No. 289
5.9 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.10 Electronic balance, ±1-mg sensitivity

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Ammonium chloride solution (NH₄Cl), 1 N. Dissolve 535 g of NH₄Cl reagent in DDI water and dilute to 10 L.
6.3 Ethanol (CH₃CH₂OH), 95%, U.S.P.
6.4 Nessler’s reagent. Add 4.56 g of potassium iodide (KI) to 30 mL DDI water. Add 5.68 g of mercuric iodide (Hgl₂). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL DDI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with DDI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately. Store the reagent in a brown bottle to protect from light.
6.5 Sodium chloride (NaCl), reagent, crystal
6.6 Antifoam agent, slipicone release spray, Dow Chemical Corp. Alternatively, use n-octyl alcohol.
6.7 Boric acid, 4% (w:v), with bromcresol green-methyl red indicator (0.075 % bromcresol green and 0.05% methyl red), Chempure Brand
6.8 Hydrochloric acid (HCl), 0.05 N, standardized. Dilute 83 mL of concentrated HCl in 16 L of DDI water.
6.9 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of DDI water. Dilute to 9 L with DDI water.

7. Procedure

**Extraction of Bases**

7.1 Prepare extraction vessel by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.
7.2 Weigh 2.50 g of <2-mm, air-dry soil and place in an extraction vessel. Weigh a smaller amount of sample, if the soil is highly organic. Prepare one quality control check sample per 48 samples.

7.3 Place extraction vessel on upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1 in) length rubber tubing and insert the plunger in the slot of the stationary disk of the extractor.

7.4 Use a squeeze bottle to fill extraction vessel to the 20-mL mark with NH$_4$Cl solution (≈10 mL). Thoroughly wet the sample. Let stand for at least 20 min.

7.5 Put reservoir tube on top of the extraction vessel. Rapidly extract the NH$_4$Cl solution to a 0.5- to 1.0-cm height above sample. Turn off extractor. Add ≈45 mL of NH$_4$Cl solution to the reservoir tube. Set extractor for an overnight (12 to 16 h) extraction.

7.6 Next morning turn off the extractor. Pull the plunger of the syringe down. Do not pull plunger from the barrel of the syringe. Carefully remove the syringe containing the extract. Leave the rubber tubing on the extraction vessel. Weigh each syringe containing the NH$_4$Cl extract to the nearest 0.01 g.

7.7 Mix the extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. The solution in the polycon is reserved for analyses of extracted cations (methods 6N2, 6O2, 6P2, and 6Q2).

7.8 Return the extractor to starting position. Attach syringe to the extraction vessel and rinse the sides of the extraction vessel with ethanol from a wash bottle. Fill the extraction vessel to the 20-mL mark with ethanol and let stand for 15 to 20 min.

7.9 Place reservoir tube on the extraction vessel. Rapidly extract the ethanol level to a 0.5- to 1.0-cm height above the sample. Turn off the extractor and add 55 to 60 mL of ethanol to the reservoir. Extract at a 45-min rate.

7.10 After the extractor has stopped, turn off the switch. Pull the plunger of the syringe down. Do not pull the plunger from the syringe barrel. Remove the syringe and discard the ethanol.

7.11 Repeat the ethanol wash.

7.12 After the second wash, collect a few drops of ethanol extract from the extraction vessel on a spot plate. Test for NH$_4^+$ by using Nessler’s reagent. A yellow, red to reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler’s reagent. Repeat until a negative test is obtained.
Steam Distillation: Samples and Reagent Blanks

7.13 Remove the extraction vessel and transfer the sample to a 250-mL digestion tube. Add 6 to 7 g of NaCl to the sample.

7.14 Perform the same transfer and addition of reagents for blanks as for samples.

7.15 Spray silicone antifoam agent (or 2 drops of n-octyl alcohol solution) into the digestion tubes for each of the samples and reagent blanks.

7.16 When using new reagents, e.g., boric acid, reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.17 On bench worksheet, record the normality of standardized acid, i.e., ≈0.05 N HCl.

7.18 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

\[
\text{CEC} = \frac{\text{Titer} \times N \times 100 \times \text{AD/OD}}{\text{Sample Weight (g)}}
\]

where:

- CEC = Cation Exchange Capacity (meq 100 g\(^{-1}\))
- Titer = Titer of sample (mL)
- \(N\) = Normality of HCl titrant
- 100 = Conversion factor to 100-g basis
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report neutral salts CEC in units of meq 100 g\(^{-1}\) of oven-dry soil to the nearest 0.1 meq 100 g\(^{-1}\).

10. Precision

Precision data are not available for this procedure.

11. References

CHEMICAL ANALYSES (6)

Organic Carbon (6A)
Walkley-Black Modified Acid-Dichromate Organic Carbon (6A1)
FeSO₄ Titration, Automatic Titrator
Metrohm 686 Titroprocessor (6A1C)

1. Application
Organic C by the Walkley-Black method is a wet combustion technique to estimate organic C. A correction factor is used to convert the Walkley-Black value to an organic matter content. A common value for the factor is 1.724 based upon the assumption that soil organic matter contains 58% organic C. A review of the literature reveals that the factor is highly variable, not only among soils but also between horizons in the same soil (Broadbent, 1953). In addition, a recovery factor is used because the Walkley-Black method does not completely oxidize all the organic C.

2. Summary of Method
The SSL uses the Walkley-Black modified acid-dichromate FeSO₄ titration organic carbon procedure. A sample is oxidized with 1 N potassium dichromate and concentrated sulfuric acid (1:2 volume ratio). After 30 min, the reaction is halted by dilution with water. The excess dichromate is potentiometrically back-titrated with ferrous sulfate. A blank is carried throughout the procedure to standardize the ferrous sulfate. Percent organic C is reported on an oven-dry soil basis.

3. Interferences
Dichromate methods that do not use additional heating do not give complete oxidation of organic matter. Even with heating, the recovery may not be complete. Walkley and Black (1934) determined an average recovery factor of 76%. Other studies have found recovery factors ranging from 60% to 86%. Thus, an average correction factor yields erroneous values for many soils. The Walkley-Black method is only an approximate or semiquantitative estimate of organic C.

Maintain the ratio of dichromate solution to concentrated H₂SO₄ at 1:2 to help maintain uniform heating of the mixture.

The presence of significant amounts of chloride in the soil results in a positive error. If the chloride in the soil is known, use the following correction factor (Walkley, 1947) for the organic C.

\[ \text{Organic C (\%) = Apparent soil C \% - (Soil Cl⁻ \%)/12} \]

The presence of significant amounts of ferrous ions results in a positive error (Walkley, 1947). The dichromate oxidizes ferrous to ferric iron.
\[ Cr_2O_7^{2-} + 6 Fe^{2+} + 14 H^+ = 2 Cr^{3+} + 6 Fe^{3+} + 7 H_2O \]

The presence of manganese dioxide results in a negative error (Walkley, 1947). When heated in an acidic medium, the higher oxides of manganese, e.g., MnO_2, compete with dichromate for oxidizable substances.

\[ 2 MnO_2 + C^0 + 4 H^+ = CO_2^+ + 2 Mn^{2+} + 2 H_2O \]

All dichromate methods assume that the organic C in the soil has an average oxidation state of zero and an equivalent weight of 3 g per equivalent when reacting with dichromate. When the soil has carbonized material, e.g., charcoal, graphite, coal and soot, the Walkley-Black method gives low recovery of this material, i.e., recovery range is from 2 to 36%.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing acids and dichromate. Toxic chromyl chloride may be released from the sample, if high concentrations of chloride are present. Use the fume hood to contain the gases released by this procedure. Use the safety showers and eyewash stations to dilute spilled acids. Use sodium bicarbonate and water to neutralize and dilute spilled acids and dichromate. Follow the manufacturer’s safety precautions when using the automatic titrator.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Titration beakers, borosilicate glass, 250 mL
5.3 Automatic dispenser, 5 to 20 mL, Oxford no. 470 or equivalent, for K_2Cr_2O_7, capable of volume adjustment to 10.00 ±0.01 mL, 0.5% reproducibility.
5.4 Dispenser, Zippette 30 mL or equivalent, for concentrated H_2SO_4, Brinkmann Instruments, Inc.
5.5 Shaker, Eberbach 6000 power unit, fitted with spring holders for titration beakers, reciprocating speed of 60 to 260 epm, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
5.6 Automatic titrator, Metrohm 686 Titroprocessor Series 04, 664 Control Unit, 674 Sample Changer Series 5, and 665 Dosimat Series 14, Metrohm Ltd., Brinkmann Instruments, Inc.
5.7 Platinum electrode, Metrohm part no. 6.0412.000

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Potassium dichromate, 1.000 N, primary standard. Dissolve 49.035 g of K₂Cr₂O₇ reagent, dried @ 105 °C, in 1-L volumetric flask with DDI water.

6.3 Sulfuric acid (H₂SO₄), concentrated, reagent

6.4 Ferrous sulfate, 1 N, acidic. Dissolve 1 kg of FeSO₄·7H₂O in 6 L of DDI water. Carefully add 640 mL of concentrated H₂SO₄ with stirring. Cool and dilute to 8 L with DDI water.

7. Procedure

**Digestion of Organic C**

7.1 Weigh 1.000 g air-dry soil and place in a titration beaker. If the sample contains >3% of organic C, use a smaller sample size. Refer to Table 1 for sample weight guide. If sample size is <0.5 g, use <80-mesh soil. If sample size is >0.5 g, use <2-mm soil.

7.2 With automatic dispenser, add 10.00 mL of K₂Cr₂O₇ solution to the titration beaker. Mix by swirling the sample.

7.3 Use the dispenser to carefully add 20 mL of concentrated H₂SO₄ to the beaker. Mix by swirling solution. Adjustment in the amount of K₂Cr₂O₇ added to sample requires appropriate adjustment in the amount of H₂SO₄ so that a 1:2 volume is maintained.

7.4 Place titration beaker on the reciprocating shaker and shake 1 min. If the dichromate-acid mixture turns a blue-green color, all the dichromate has been reduced. Add more dichromate and acid to maintain a 1:2 volume. Refer to Table 1 for dichromate:acid volumes.

<table>
<thead>
<tr>
<th>OC (%)</th>
<th>Sample (g)</th>
<th>K₂Cr₂O₇ (mL)</th>
<th>H₂SO₄ (mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-3</td>
<td>1.000</td>
<td>10.00</td>
<td>20</td>
</tr>
<tr>
<td>3-6</td>
<td>0.500</td>
<td>10.00</td>
<td>20</td>
</tr>
<tr>
<td>3-6</td>
<td>1.000</td>
<td>20.00</td>
<td>40</td>
</tr>
<tr>
<td>6-12</td>
<td>0.500</td>
<td>20.00</td>
<td>40</td>
</tr>
<tr>
<td>12-24</td>
<td>0.250</td>
<td>20.00</td>
<td>40</td>
</tr>
<tr>
<td>24-50</td>
<td>0.100</td>
<td>30.00</td>
<td>60</td>
</tr>
</tbody>
</table>

7.5 Place the beaker on a heat resistant surface for 30 min.

7.6 Add ≈180 mL DDI water to the beaker to stop the reaction.
Titration of Excess Dichromate

7.7 Titrate eight reagent blanks at the start of each batch to determine the normality of the ferrous sulfate. A blank is 10.00 mL K$_2$Cr$_2$O$_7$ plus H$_2$SO$_4$ without soil. The average titer is used for the blank titer value.

7.8 Place the appropriate blanks and samples in the sample holder magazines and place on the sample changer.

7.9 Refer to the manufacturer’s instruction manual for operation of automatic titrator.

7.10 Set the endpoint to 700 mV. Set the controls of the 664 Control Unit to the appropriate settings.

7.11 Prime the burette with 50 mL of ferrous sulfate solution before starting the titrations.

7.12 When a long series of samples are being titrated, intersperse blank samples throughout the titrations. The blank titer drifts over time, mainly because of the temperature change of the solution. Any sample with a titer of less one milliliter and/or endpoint of less than 620 millivolts should be reanalyzed.

7.13 Press “Start” on the titrator.

8. Calculations

\[
OC \, (\%) = \frac{[(\text{Blank} \times \text{Volume}) - (10 \times \text{Titer}) \times 3 \times 100 \times \text{AD/OD}]}{\text{Blank} \times \text{Sample Weight (g)} \times 0.77 \times 1000}
\]

where:
- OC (\%) = Organic C (\%)
- Blank = Average titer of reagent blanks (mL)
- Volume = Volume of 1 N K$_2$Cr$_2$O$_7$ (mL)
- Titer = Titer of FeSO$_4$ (mL)
- AD/OD = Air-dry/oven-dry ratio (method 4B5)
- 3 = Equivalents per C (assumed)
- 1000 = Meq eq$^{-1}$
- 100 = Convert to 100-g basis
- 0.77 = Assumed C oxidation factor

9. Report

Report organic C percentage to two decimal places, e.g., 0.95% OC, on an oven-dry basis.
10. Precision

Precision data are not available for this procedure. A quality control check sample is run in every batch of 20 samples. With 251 observations of the quality control check sample, the mean, standard deviation, and C.V. for organic carbon are 1.47, 0.025, and 1.7%, respectively.

11. References


Total Carbon (6A)
Dry Combustion (6A2)
LECO SC-444 Carbon Analyzer (6A2e)

1. Application

Total C in soils is the sum of organic and inorganic C. Most of the organic C is associated with the organic matter fraction, and the inorganic C is generally found with carbonate minerals. The organic C in mineral soils generally ranges from 0 to 12 percent.

Total C is quantified by two basic methods, i.e., wet or dry combustion. The SSL uses dry combustion. In total C determinations, all forms of C in a soil are converted to CO$_2$ followed by a quantification of the evolved CO$_2$. Total C can be used to estimate the organic C content of a soil. The difference between total and inorganic C is an estimate of the organic C. Organic C also can be determined directly (method 6A1c). The inorganic C should be equivalent to carbonate values measured by CO$_2$ evolution with strong acid (Nelson and Sommers, 1982).

Organic C defines mineral and organic soils. In Soil Taxonomy, organic C is also used at lower taxonomic levels, e.g., ustolic and fluventic subgroups (Soil Survey Staff, 1975).

2. Summary of Method

A fine-ground (<80-mesh) soil sample is oxidized at high temperatures. The released gases are scrubbed, and the CO$_2$ in the combustion gases is measured by using an infrared detector. The microprocessor formulates the analytical results (C$_i$) by combining the outputs of the infrared detector and the system...
ambient sensors with pre-programmed calibration, linearization, and weight compensation factors. Percent total C is reported on an oven-dry soil basis.

3. Interferences

This procedure simultaneously measures inorganic and organic C. A high rate of combustion can oversaturate the carbon detection cell. The rate of combustion can be retarded by adding a solid/powder combustion controller.

4. Safety

Wear protective clothing and safety glasses. Magnesium perchlorate may form explosive mixtures. Magnesium perchlorate may contain traces of perchloric acid, which remain from manufacturer’s operations. This acid is anhydrous because of the strong desiccating capability of the salt. Avoid prolonged contact with oxidizable material or material capable of forming unstable perchlorate esters or salts. Remove magnesium perchlorate by using an excess of water to thoroughly dilute the material.

The use of high temperatures in the oxidation of samples requires that extreme caution be used to prevent burns and fires. Follow standard laboratory procedures when handling compressed gases. Oxygen is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the carbon analyzer.

5. Equipment

5.1 Carbon analyzer, Leco Model SC-444, Sulfur and Carbon Analyzers, Leco Corp., St. Joseph, MI
5.2 Combustion boats, part no. 529-203, Leco Corp., St. Joseph, MI
5.3 Single-stage regulator, oxygen service, part no. E11-W-N115Box, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.4 Electronic balance, ±1-mg sensitivity

6. Reagents

6.1 Anhydrous magnesium perchlorate, granular
6.2 Glass wool
6.3 Compressed oxygen, >99.5% @ 30 psi
6.4 Calcium carbonate, CaCO₃, reagent grade.
6.6 Soil Calibration Sample, part no. 502-062, Leco Corp., St. Joseph, MI
7. Procedure

7.1 Use a fine-ground 80-mesh, air-dry soil.

7.2 Prepare instrument as outlined in the operator’s instruction manual (Leco, 1994; Leco, 1993).

7.3 Methods are created with the method menu and stored in the instrument memory. System parameters are set as follows:
- Furnace operating temperature: 1450 °C
- Lance delay: 20 s
- Analysis time settings: 70 to 180 s
- Comparator level settings: 0.1%

7.3 Condition instrument by analyzing a few soil samples, until readings are stable.

7.4 Calibrate instrument by analyzing at least three replicates of each calibration standard. Use the soil calibration standard for samples with less than 3 to 4 percent total carbon and calcium carbonate for samples with more than 4 percent total carbon. Weigh standards in a range from 0.2 to 0.7 g.

7.5 Load samples on autoload rack, place in the analyzer, and press analyze key.

7.6 Weigh 0.2 to 0.5 g sample in a tared combustion boat.

7.7 Load samples on autoload rack, place in the analyzer, and press analyze key.

7.8 If results exceed calibration range, reduced weight of sample. If carbon detection cell is saturated, add approximately 1 g of solid/powder combustion controller to sample.

7.9 Repack the reagent (anhydrous magnesium perchlorate) tubes whenever the reagent becomes caked or moist or the warning alarm displays.

8. Calculations

\[ C(\%) = C_i \times \frac{AD}{OD} \]

where:
- \( C(\%) = C(\%), \) oven-dry basis
- \( C_i = C(\%) \) instrument
- \( \frac{AD}{OD} = \text{air-dry/oven-dry ratio (method 4B5)} \)

9. Report

Report total C percentage on an oven-dry basis to the nearest 0.1%.
10. Precision

A quality control check sample is included in every batch of ten samples. For 191 observations of calcium carbonate (actual total C=12%), the mean, standard deviation, and C.V. for total carbon are 12.04, 0.31, and 2.5%, respectively. For 86 observations of soil calibration standard (reported total C=0.77%), the mean, standard deviation, and C.V. for total carbon are 0.79, 0.02, and 2.2%, respectively.

11. References


Total Nitrogen (6B)
Dry Combustion (6B4)
LECO FP-428 Analyzer (6B4a)

1. Application

The total N content of the soil may range from <0.02% in subsoils, 2.5% in peats, and 0.06 to 0.5% in surface layers of many cultivated soils (Bremmer and Mulvaney, 1982). The total N data may be used to determine the soil C:N ratio, the soil potential to supply N for plant growth, and the N distribution in the soil profile. The C:N ratio generally ranges between 10 to 12. Variations in the C:N ratio may serve as an indicator of the amount of soil inorganic N. Uncultivated soils usually have higher C:N ratios than do cultivated soils.

Soils with large amounts of illites or vermiculites can “fix” significant amounts of N compared to those soils dominated by smectites or kaolinites (Young and Aldag, 1982; Nommik and Vahtras, 1982). Since the organic C of many soils diminishes with depth while the level of “fixed” N remains constant or increases, the C:N ratio narrows (Young and Aldag, 1982). The potential to “fix” N has important fertility implications as the “fixed” N is slowly available for plant growth.
2. **Summary of Method**

A soil sample is combusted at high temperature with oxygen to release NO\(_x\). The gases released are scrubbed to remove interferences (e.g., CO\(_2\) and H\(_2\)O), and the NO\(_x\) is reduced to N\(_2\). The N\(_2\) is measured by thermal conductivity detection and reported as percent N.

3. **Interferences**

The total N that is measured by the combustion method does not distinguish among the types of N that are present in the soil. The purity of the helium and oxygen gases used in the instrument may affect the results of the analysis. The highest purity gases available are required to assure low detection limits and consistent results.

4. **Safety**

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (goggles or safety glasses) when handling hot crucibles. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary.

5. **Equipment**

5.1  
Electronic balance, ±0.001-g sensitivity

6. **Reagents**

None.

7. **Procedure**

7.1  
Weigh 0.200 g of 80-mesh, air-dry soil into a tin foil cup.

7.2  
Close the tin foil cup by twisting the top closed as to fit the sample holder.

7.3  
Place the enclosed sample in the sample holder.

7.4  
When all the samples are in the sample holder, place the sample holder on the instrument.

7.5  
Refer to manufacturer’s manual for operation and calibration of the LECO FP-438 Analyzer.

7.6  
On the bench worksheet, record the percent N for the samples.

8. **Calculations**

\[
N (\%) = \text{Instrument Reading} \times \frac{\text{AD}}{\text{OD}}
\]

where:

\[
\frac{\text{AD}}{\text{OD}} = \text{Air-dry/Oven-dry ratio (method 4B5)}
\]
9. Report
   Report total N as a dimensionless value to the nearest 0.001 unit on an oven-dry basis.

10. Precision
   Precision data are not available for this procedure. For 105 observations of the quality control check sample for total N, the mean, standard deviation, and C.V. are 0.143, 0.004, and 2.7 percent, respectively.

11. References

Mineralizable Nitrogen (6B)
Steam Distillation (6B5)
Kjeltec Auto 1035 Sampler (6B5a)

1. Application
   The most satisfactory methods currently available for obtaining an index for the availability of soil N are those involving the estimation of the N formed when soil is incubated under conditions which promote mineralization of organic N by soil microorganisms (Environmental Protection Agency, 1992). The method described herein for estimating mineralizable N is one of anaerobic incubation and is suitable for routine analysis of soils. This method involves estimation of the ammonium produced by a 1-week period of incubation of soil at 40 °C (Keeney and Bremner, 1966) under anaerobic conditions to provide an index of N availability.

2. Method Summary
   An aliquot of air-dry homogenized soil is placed in a test tube with water, stoppered, and incubated at 40 °C for 1 week. The contents are transferred to a steam distillation, rinsed with 4 N KCl. The amount of ammonium-N is determined by steam distillation and titration for the KCl:soil mixture.
3. Interferences

There are no known interferences. The temperature and incubation period must remain constant for all samples. The test can be performed on field-moist or air-dry soil samples.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow the manufacturer’s safety precautions when using the incubator and Kjeltec Auto 1035 Analyzer.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Test tubes, 16-mm x 150-mm
5.3 PVC stoppers
5.4 Incubator, Model 10-140, Quality Lab Inc., Chicago, IL
5.5 Digestion tubes, straight neck, 250 mL
5.6 Kjeltec Auto 1035/1038 Sampler System, Tecator, Perstorp Analytical Inc.

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Potassium chloride (KCl), 4 N. Dissolve 298.24 g KCl in DDI water and dilute to 1-L volume.
6.3 Hydrochloric acid (HCl), 0.05 N, standardized. Dilute 83 mL of concentrated HCl in 20 L of DDI water.
6.4 Antifoam agent, slipicone release spray, Dow Chemical Corp. Alternatively, use n-octyl alcohol.
6.5 Boric acid, 4% (w:v), with brom cresol green-methyl red indicator (0.075 % brom cresol green and 0.05% methyl red), Chempure Brand

7. Procedure

**Anaerobic Incubation of Soil Sample**

7.1 Place 5.00 g of mineral soil (or 1.25 g of organic soil) into a 16-mm x 150-mm test tube. Record the soil sample weight to the nearest 0.00 g.
7.2 Add 12.5 ±1 mL of DDI water. Do not add ethanol to overcome any wetting difficulties as ethanol may act as an interference with microbial activity. Stopper the tube, shake, and place in a 40 °C constant-temperature incubator for 7 days. Refer to the manufacturer's instructions for set-up and operation of the incubator.

7.3 At the end of 7 days, remove the tube and shake for 15 s.

7.4 Transfer the contents of the test tube to a 250-mL digestion tube. Complete the transfer by rinsing the tube with 3 times with 4 ml of 4 N KCl, using a total of 12.5 ±1 mL of the KCl.

7.5 Remove the extraction vessel and transfer the sample to a 250-mL digestion tube.

7.6 Perform the same transfer and addition of reagents for blanks as for samples.

7.7 Spray silicone antifoam agent (or 2 drops of n-octyl alcohol solution) into the digestion tubes for each of the samples and reagent blanks.

7.8 When using new reagents, e.g., boric acid, reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.9 On bench worksheet, record the normality of standardized acid, i.e., ≈0.0500 N HCl.

7.10 Load samples in racks of 20. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

\[ N = \frac{(\text{Titer} \times N \times 100 \times \text{AD/OD})}{\text{Sample Weight (g)}} \]

where:

- \( N \) = Mineralizable N (meq 100 g\(^{-1}\))
- \( \text{Titer} \) = Titer of sample (mL)
- \( N \) = Normality of HCl titrant
- 100 = Conversion factor to 100-g basis
- \( \text{AD/OD} \) = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report mineralizable N in units of meq 100 g\(^{-1}\) of oven-dry soil to the nearest 0.001 meq 100 g\(^{-1}\).
Iron, Manganese, and Aluminum (6C, 6D, and 6G)
Dithionite-Citrate Extraction (6C2, 6D2, and 6G7)
Atomic Absorption Perkin-Elmer AA 5000 (6C2b, 6D2a, and 6G7a)

1. Application
Dithionite-citrate (CD) is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, and amorphous aluminosilicates (Wada, 1989). The CD solution is a poor extractant of crystalline hydrous oxides of Al, allophane, and imogolite. The CD solution does not extract opal, Si, or other constituents of crystalline silicate minerals (Wada, 1989). In Soil Taxonomy, the CD extractable Fe and Al are criteria for spodic placement (Soil Survey Staff, 1975).

2. Summary of Method
A soil sample is mixed with sodium dithionite, sodium citrate, and distilled deionized water, and shaken overnight. Superfloc 16 is added, and the mixture is made to volume. Solution is allowed to settle, and a clear extract is obtained. The CD extract is diluted with distilled deionized (DDI) water. The analytes are by an atomic absorption spectrophotometer (AA). The data are automatically recorded by a microcomputer and printer. The percent CD extractable Fe, Mn, and Al are reported in methods 6C2b, 6D2a, and 6G7a, respectively.

3. Interferences
There are four types of interferences (matrix, spectral, chemical, and ionization) in the AA analyses of these elements. These interferences vary in importance, depending upon the particular analyze selected.

The redo potential of the extractant is dependent upon the pH of the extracting solution and the soil system. Sodium citrate complexes the reduced Fe and
usually buffers the system to a pH of 6.5 to 7.3. Some soils may lower the pH, resulting in the precipitation of Fe sulfides. The SSL has not had significant problems with this interference.

Filtered extracts can yield different recoveries of Fe, Mn, and Al, relative to unfiltered extracts.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves); eye protection (face shields, goggles, or safety glasses); and a breathing filter when handling dry sodium dithionite. Sodium dithionite may spontaneously ignite if allowed to become moist, even by atmospheric moisture. Keep dithionite in a fume hood.

Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Filter paper, pre-pleated, 185-mm diameter, Schleicher and Schuell
5.3 Atomic absorption spectrophotometer (AA), model 5000, Perkin-Elmer Corp., Norwalk, CT
5.4 Automatic burner control, model 5000, Perkin-Elmer Corp., Norwalk, CT
5.5 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.6 Dot matrix printer, P-132, Interdigital Data Systems, Inc.
5.7 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.8 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.9 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.10 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.11 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Sodium dithionite (Na₂S₂O₄), purified powder
6.3 Sodium citrate dihydrate (Na₃C₆H₅O₇•2H₂O), crystal, reagent
6.4 Hydrochloric acid (HCl), concentrated 12 N
6.5 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.
6.6 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.
6.7 Superfloc 16, 0.2% solution (w:v). Dissolve 2 g of Superfloc 16 in 1 liter of DDI water. Do not shake the mixture as this breaks the polymer chains of the Superfloc. Gently swirl the mixture occasionally over the several days that the solution requires to completely dissolve the Superfloc. Suggested source is American Cyanamid Co., P.O. Box 32787, Charlotte, NC.
6.8 Primary mixed standard, 4000 mg L\(^{-1}\) (4000 ppm) Fe, 600 mg L\(^{-1}\) (600 ppm) Mn, and 3000 mg L\(^{-1}\) (3000 ppm) Al. Dissolve 4.0 g of Fe wire, 0.6 g of Mn metal powder, and 3.0 g of Al wire with 1:1 HCl in a glass beaker. When dissolved, transfer to a 1-L volumetric flask and make to volume with 1% HCl solution. Store in a polypropylene bottle.
6.9 High calibration standard, 240 mg/8 oz (1012 ppm) Fe; 36 mg/8 oz (75 ppm) Mn; and 180 mg/8 oz (76 ppm) Al. Pipette 60 mL of primary mixed standard into 8 oz bottle. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H\(_2\)SO\(_4\), and 2 mL of Superfloc 16 solution. In standards, the H\(_2\)SO\(_4\) substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.
6.10 Low calibration standard, 120 mg/8 oz (506 ppm) Fe; 18 mg/8 oz (76 ppm) Mn; and 90 mg/8 oz (380 ppm) Al. Pipette 30 mL of primary mixed standard into 8 oz bottle. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H\(_2\)SO\(_4\), and 2 mL of Superfloc 16 solution. In standards and reagent blanks, the H\(_2\)SO\(_4\) substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.
6.11 Calibration reagent blank solution. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H\(_2\)SO\(_4\), and 2 mL of Superfloc 16 solution. In standards and reagent blanks, the H\(_2\)SO\(_4\) substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.
6.12 Acetylene gas, purity 99.6%
6.13 Compressed air with water and oil traps

7. Procedure

**Extraction of Fe, Mn, and Al**

7.1 Weigh 4.0 g of <2-mm, air-dry soil sample and place in an 8-oz nursing bottle.
7.2 Add 2 g of sodium dithionite and 20 to 25 g of sodium citrate dihydrate.

7.3 Add DDI water to 4-oz level on bottle and securely stopper bottle.

7.4 Shake overnight (12 to 16 h) in a reciprocating shaker. After shaking, use a dispenser to add 2 ml of Superfloc 16 solution.

7.5 Fill bottle to 8-oz volume with DDI water. Stopper and shake thoroughly for ~15 s.

7.6 Allow to settle for at least 3 day (3 to 5 days typical). The Fe, Mn, and Al are determined from a clear aliquot of solution.

**Dilution of Sample Extracts and Standards**

7.7 No ionization suppressant is required as the Na in the extractant is present in sufficient quantity. Set the digital diluter at 66 for diluent and 35 for CD extracts, calibration reagent blanks, and calibration standards for a 1:20 dilution as follows:

7.8 Dilute 1 part CD sample extract with 19 parts of DDI water (1:20 dilution).

7.9 Dilute 1 part calibration reagent blank with 19 parts of DDI water (1:20 dilution).

7.10 Dilute 1 part low calibration standard with 19 parts of DDI water (1:20 dilution).

7.11 Dilute 1 part high calibration standard with 19 parts of DDI water (1:20 dilution).

7.12 Dispense the reagent blanks and calibration standards in polycons from which the solutions are transferred to test tubes. Dispense the diluted sample solutions into test tubes which have been placed in the sample holders of the sample changer.

**AA Calibration**

7.13 Use the calibration reagent blank and high calibration standard to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

7.14 Use the low calibration standard (120 mg/8 oz Fe; 18 mg/8 oz Mn; and 90 mg/8 oz Al) as a check sample. Use high calibration standard for Fe check sample and low calibration standard for Mn and Al check sample.

**AA Set-up and Operation**

7.15 Refer to the manufacturer’s manual for operation of the AA. The following are only very general guidelines for instrument conditions for the various analytes.
<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength (nm)</th>
<th>Burner Head &amp; Angle</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>248.5</td>
<td>5-cm, parallel</td>
<td>10 C₂H₂/25 Air</td>
</tr>
<tr>
<td>Al</td>
<td>309.35</td>
<td>5-cm, parallel</td>
<td>30 C₂H₂/17 N₂O</td>
</tr>
<tr>
<td>Mn</td>
<td>280.15</td>
<td>5-cm, parallel</td>
<td>10 C₂H₂/25 Air</td>
</tr>
</tbody>
</table>

Typical read delay is 6 s, and integration time is 8 s.

7.16 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.17 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with DDI water (1:20 dilution).

7.18 The instrument readings are usually programmed to display analyte concentration in mg/8 oz.

8. Calculations

\[
\text{Fe} (\%) = \frac{\text{Fe} \times \text{DR} \times 100 \times \text{AD}/\text{OD}}{\text{Sample} \times 1000}
\]

\[
\text{Fe}_2\text{O}_3 (\%) = \frac{\text{Fe} \times \text{DR} \times 1.43 \times 100 \times \text{AD}/\text{OD}}{\text{Sample} \times 1000}
\]

\[
\text{Mn (\%)} = \frac{\text{Mn} \times \text{DR} \times 100 \times \text{AD}/\text{OD}}{\text{Sample} \times 1000}
\]

\[
\text{Al (\%)} = \frac{\text{Al} \times \text{DR} \times 100 \times \text{AD}/\text{OD}}{\text{Sample} \times 1000}
\]

where:

- Fe = mg/8 oz
- Mn = mg/8 oz
- Al = mg/8 oz
- DR = Dilution Ratio
- Sample = Sample weight (g)
- 1.43 = Conversion factor from Fe to Fe₂O₃
- 100 = Conversion factor to percent
- AD/OD = Air-dry/oven-dry ratio (method 4B5)
- 1000 = Conversion factor (mg g⁻¹)

9. Report

Report percent CD extractable Fe, Mn, and Al on oven-dry soil basis to the nearest whole number.
10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of samples. For the quality control check sample, the mean, standard deviation, and C.V. for Fe, Mn, and Al are as follows:

<table>
<thead>
<tr>
<th>Element</th>
<th>Mean</th>
<th>n</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>2.5</td>
<td>35</td>
<td>0.05</td>
<td>2.2%</td>
</tr>
<tr>
<td>Mn</td>
<td>0.01</td>
<td>19</td>
<td>0.00</td>
<td>0.0%</td>
</tr>
<tr>
<td>Al</td>
<td>0.26</td>
<td>33</td>
<td>0.01</td>
<td>5.4%</td>
</tr>
</tbody>
</table>

11. References


Iron, Manganese, and Aluminum (6C, 6D, and 6G)
Dithionite-Citrate Extraction (6C2, 6D2, and 6G7)
Atomic Absorption
Thermo Jarrell Ash, Smith-Hieftje 4000 (6C2c, 6D2b, and 6G7b)

1. Application

Dithionite-citrate (CD) is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, and amorphous aluminosilicates (Wada, 1989). The CD solution is a poor extractant of crystalline hydrous oxides of Al, allophane, and imogolite. The CD solution does not extract opal, Si, or other constituents of crystalline silicate minerals (Wada, 1989). In Soil Taxonomy, the CD extractable Fe and Al are criteria for spodic placement (Soil Survey Staff, 1975).

2. Summary of Method

A soil sample is mixed with sodium dithionite, sodium citrate, and distilled deionized water, and shaken overnight. Superfloc 16 is added, and the mixture is made to volume. Solution is allowed to settle, and a clear extract is obtained. The CD extract is diluted with distilled deionized (DDI) water. The analytes are measured by an atomic absorption spectrophotometer (AA). The data are automatically recorded by a microcomputer and printer. The percent CD
extractable Fe, Mn, and Al are reported in methods 6C2c, 6D2b, and 6G7b, respectively.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the AA analyses of these elements. These interferences vary in importance, depending upon the particular analyte selected.

The redox potential of the extractant is dependent upon the pH of the extracting solution and the soil system. Sodium citrate complexes the reduced Fe and usually buffers the system to a pH of 6.5 to 7.3. Some soils may lower the pH, resulting in the precipitation of Fe sulfides. The SSL has not had significant problems with this interference.

Filtered extracts can yield different recoveries of Fe, Mn, and Al, relative to unfiltered extracts.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves); eye protection (face shields, goggles, or safety glasses); and a breathing filter when handling dry sodium dithionite. Sodium dithionite may spontaneously ignite if allowed to become moist, even by atmospheric moisture. Keep dithionite in a fume hood.

Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Filter paper, pre-pleated, 185-mm diameter, Schleicher and Schuell
5.3 Atomic absorption spectrophotometer (AA), Smith-Hieftje Model 4000, Thermo Jarrell Ash Corp., Franklin, MA
5.4 Autosampler, Model 150, Thermo Jarrell Ash Corp., Franklin, MA
5.5 ThermoSpec software, Version 3.01, Enable 4.0, DOS 5.0, Thermo Jarrell Ash Corp., Franklin, MA
5.6 Computer, CUi Advantage 486, Thermo Jarrell Ash Corp., Franklin, MA
5.7 Printer, NEC Pinwriter P3200
5.8 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.9 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.10 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.11 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.12 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Sodium dithionite (Na₂S₂O₄), purified powder
6.3 Sodium citrate dihydrate (Na₃C₆H₅O₇•2H₂O), crystal, reagent
6.4 Hydrochloric acid (HCl), concentrated 12 N
6.5 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.
6.6 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.
6.7 Superfloc 16, 0.2% solution (w:v). Dissolve 2 g of Superfloc 16 in 1 liter of DDI water. Do not shake the mixture as this breaks the polymer chains of the Superfloc. Gently swirl the mixture occasionally over the several days that the solution requires to completely dissolve the Superfloc. Suggested source is American Cyanamid Co., P.O. Box 32787, Charlotte, NC.
6.8 Primary mixed standard, 4000 mg L⁻¹ (4000 ppm) Fe, 600 mg L⁻¹ (600 ppm) Mn, and 3000 mg L⁻¹ (3000 ppm) Al. Dissolve 4.000 g of Fe wire, 0.6000 g of Mn metal powder, and 3.000 g of Al wire with 1:1 HCl in a glass beaker. When dissolved, transfer to a 1-L volumetric flask and make to volume with 1% HCl solution. Store in a polypropylene bottle.
6.9 High calibration standard, 240 mg/8 oz (1012 ppm) Fe; 36 mg/8 oz (152 ppm) Mn; and 180 mg/8 oz (759 ppm) Al. Pipette 60 mL of primary mixed standard into 8 oz bottle. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H₂SO₄, and 2 mL of Superfloc 16 solution. In standards, the H₂SO₄ substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.
6.10 Low calibration standard, 120 mg/8 oz (506 ppm) Fe; 18 mg/8 oz (76 ppm) Mn; and 90 mg/8 oz (380 ppm) Al. Pipette 30 mL of primary mixed standard into 8 oz bottle. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H₂SO₄, and 2 mL of Superfloc 16 solution. In standards and reagent blanks, the H₂SO₄ substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.
6.11 Calibration reagent blank solution. Add 20 g of sodium citrate dihydrate, 1.24 mL of concentrated H₂SO₄, and 2 mL of Superfloc 16 solution. In
standards and reagent blanks, the H$_2$SO$_4$ substitutes for the dithionite. Fill to 8-oz volume with DDI water and mix thoroughly. After dissolution, transfer solution to a plastic bottle.

6.12 Acetylene gas, purity 99.6%
6.13 Compressed air with water and oil traps

7. Procedure

Extraction of Fe, Mn, and Al

7.1 Weigh 4.00 g of <2-mm, air-dry soil sample and place in an 8-oz nursing bottle.
7.2 Add 2 g of sodium dithionite and 20 to 25 g of sodium citrate dihydrate.
7.3 Add DDI water to 4-oz level on bottle and securely stopper bottle.
7.4 Shake overnight (12 to 16 h) in a reciprocating shaker. After shaking, use a dispenser to add 2 ml of Superfloc 16 solution.
7.5 Fill bottle to 8-oz volume with DDI water. Stopper and shake thoroughly for ~15 s.
7.6 Allow to settle for at least 3 day (3 to 5 days typical). The Fe, Mn, and Al are determined from a clear aliquot of solution.

Dilution of Sample Extracts and Standards

7.7 No ionization suppressant is required as the Na in the extractant is present in sufficient quantity. Set the digital diluter at 66 for diluent and 35 for CD extracts, calibration reagent blanks, and calibration standards for a 1:20 dilution as follows:
7.8 Dilute 1 part CD sample extract with 19 parts of DDI water (1:20 dilution).
7.9 Dilute 1 part calibration reagent blank with 19 parts of DDI water (1:20 dilution).
7.10 Dilute 1 part low calibration standard with 19 parts of DDI water (1:20 dilution).
7.11 Dilute 1 part high calibration standard with 19 parts of DDI water (1:20 dilution).
7.12 Dispense the reagent blanks and calibration standards in polycons from which the solutions are transferred to test tubes. Dispense the diluted sample solutions into test tubes which have been placed in the sample holders of the sample changer.

AA Calibration

7.13 Use the calibration reagent blank and high calibration standard to calibrate the AA. The AA program requires a blank and a standard, in that order,
to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

7.14 Use the low calibration standard (120 mg/8 oz Fe; 18 mg/8 oz Mn; and 90 mg/8 oz Al) as a check sample. Use high calibration standard for Fe check sample and low calibration standard for Mn and Al check sample.

**AA Set-up and Operation**

7.15 Refer to manufacturer’s manual for operation of the AA. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
<th>Burner Head &amp; Angle</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>248.5</td>
<td>5-cm, parallel</td>
<td>4 C₂H₂/16 Air</td>
</tr>
<tr>
<td>Al</td>
<td>309.35</td>
<td>5-cm, parallel</td>
<td>20 C₂H₂/10 N₂O</td>
</tr>
<tr>
<td>Mn</td>
<td>280.15</td>
<td>5-cm, parallel</td>
<td>4 C₂H₂/10 Air</td>
</tr>
</tbody>
</table>

Typical read delay is 6 s, and integration time is 8 s.

7.16 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.17 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with DDI water (1:20 dilution).

7.18 The instrument readings are usually programmed to display analyte concentration in mg/8 oz.

**8. Calculations**

\[
\text{Fe} \text{ (%) } = \frac{(\text{Fe} \times \text{DR} \times 100 \times \text{AD/OD})}{(\text{Sample} \times 1000)} \\
\text{Fe₂O₃} \text{ (%) } = \frac{(\text{Fe} \times \text{DR} \times 1.43 \times 100 \times \text{AD/OD})}{(\text{Sample} \times 1000)} \\
\text{Mn} \text{ (%) } = \frac{(\text{Mn} \times \text{DR} \times 100 \times \text{AD/OD})}{(\text{Sample} \times 1000)} \\
\text{Al} \text{ (%) } = \frac{(\text{Al} \times \text{DR} \times 100 \times \text{AD/OD})}{(\text{Sample} \times 1000)}
\]

where:
- Fe = mg/8 oz
- Mn = mg/8 oz
- Al = mg/8 oz
DR = Dilution Ratio
Sample = Sample weight (g)
1.43 = Conversion factor from Fe to Fe$_2$O$_3$
100 = Conversion factor to percent
AD/OD = Air-dry/oven-dry ratio (method 4B5)
1000 = Conversion factor (mg g$^{-1}$)

9. Report
   Report percent CD extractable Fe, Mn, and Al on oven-dry soil basis to the nearest whole number.

10. Precision
   Precision data are not available for this procedure. A quality control check sample is run with every batch of samples.

11. References

Iron, Manganese, Aluminum, Calcium, Magnesium, Sodium, Potassium, Phosphorus, Silicon, Zirconium, Copper, Zinc, Titanium, Cadmium, Lead, Nickel, Chromium, and Cobalt
   HF Plus Aqua Regia (HF + HNO$_3$ + HCl) Dissolution
   Inductively Coupled Plasma Spectrometry
   Thermo Jarrell Ash ICAP 61E Optima 3300 DV
   (6C7b, 6D6a, 6G11b, 6N5b, 6O5b, 6P3b, 6Q3b, 6S6a, 6V1b, 8K1a, 8L1a, 8M1a, 8N1a, 8O1a, 8P1a, 8Q1a, and 8R1a)

1. Application
   This procedure is an integral part of total analysis (7C4a) and represents the spectroscopic analysis of elements in the digestate.

2. Summary of Method
   High and low calibration standards are prepared for Ca, K, Mg, Mn, Cu, Zn, Cd, Pb, Co (mixed standards CALO and CAHI); Al, Fe, Ti, Zr, Na, (mixed
standards ALLO and ALHI); and Si, P, Se, As (mixed standards SILO and SIHI). A blank of HF, HNO$_3$, HCl, and H$_3$BO$_3$ is prepared. A Thermo Jarrell Ash ICAP 61E spectrometer is used for analysis. The concentration of Fe, Mn, Al, Ca, Mg, Na, K, P, Si, Zr, Cu, Zn, Ti, Cd, Pb, Cr, and Co are determined by ICP analysis by methods 6C7b, 6D6a, 6G11b, 6N5b, 6P3b, 6Q3b, 6S6a, 6V1b, 8K1a, 8L1a, 8M1a, 8N1a, 8O1a, 8P1a, 8Q1a, and 8R1a.

3. Interferences
None.

4. Safety
Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated acids to the fume hood. Keep HF acid refrigerated and avoid contact with skin of all acids. Wash hands thoroughly after handling reagents.

5. Equipment
5.1 Volumetrics, 500-mL, polypropylene
5.2 Containers, 500-mL, polypropylene, with screw caps
5.3 Pipettors, electronic digital, Rainin Instrument Co., Woburn, MA, 2500 µL and 10 mL
5.4 Inductively coupled plasma spectrometer, ICAP-61E, Thermo Jarrell Ash Corp., Franklin, MA
5.5 RF generator, floor mounted power unit, 45 MHz free running, Perkin-Elmer Corp., Norwalk, CT
5.6 Computer, AT&T 386 Starstation, Model CPU-G72, and printer, NEC Pinwriter, P2200XE, Dot Matrix
5.7 ThermoSpec software, Thermo Jarrell Ash Corp., Franklin, MA
5.8 Line conditioner, Unity/1, Model UT8K, Best Power Technology, Inc., Necedah, WI
5.9 Compressed argon gas
5.10 Autosampler, Thermo Jarrell Ash Corp., Franklin, MA
5.11 High flow torch, Part No. 126440-01; Saffire (HF-resistant) tip, Part No. 127190-00; Polypropylene spray chamber, Part 131129-00, Thermo Jarrell Ash Corp., Franklin, MA

6. Reagents
6.1 Deionized distilled (DDI) water
6.2 Hydrofluoric acid (HF), 48%, low trace metal content
6.3 Concentrated hydrochloric acid (HCl), 12 N. Use instrumental grade which contains low levels of impurities.

6.4 Concentrated nitric acid (HNO₃), 16 N. Use instrumental grade which contains low levels of impurities.

6.5 Boric acid solution. Dissolve 25.0 g low trace metal, granular boric acid (H₃BO₃) in 1000 mL DDI water.

6.6 Standards, 1000 ppm, suitable for atomic absorption spectroscopy for all elements

7. Procedure

7.1 Instrument calibration standards for analysis are limited to specific combinations of elements because of chemical incompatibilities of certain elements. Specific combinations of elements in calibration standards are based on suggestion by Thermo Jarrell Ash (TJA), Inc. Each working standard is used in two concentrations, high and low. The concentrations of elements in the low standards (CALO, ALLO, and SILO) are 50 percent of the concentrations of elements in the low standards (CALO, ALLO, and SILO). Refer to Tables 1-3 for the amounts of primary standards (1000 ppm) to make 500-mL volume of the low and high calibration standards, at the specified concentrations, for ICP analysis.

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration</th>
<th>Concentration</th>
<th>Primary Std. Required For</th>
<th>Primary Std. Required For</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CALO  (ppm)</td>
<td>CAHI  (ppm)</td>
<td>CALO (mL)</td>
<td>CAHI (mL)</td>
</tr>
<tr>
<td>Ca</td>
<td>75</td>
<td>150</td>
<td>37.5</td>
<td>75.0</td>
</tr>
<tr>
<td>K</td>
<td>25</td>
<td>50</td>
<td>12.5</td>
<td>25.0</td>
</tr>
<tr>
<td>Mg</td>
<td>20</td>
<td>40</td>
<td>10.0</td>
<td>20.0</td>
</tr>
<tr>
<td>Mn</td>
<td>10</td>
<td>20</td>
<td>5.0</td>
<td>10.0</td>
</tr>
<tr>
<td>Cu</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
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<tr>
<td>Zn</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
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<tr>
<td>Cd</td>
<td>5</td>
<td>10</td>
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<tr>
<td>Pb</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
</tbody>
</table>

1All calibration standards based on 500-ml final volume.
Table 2.—Calibration standards for ALLO and ALHI¹.

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration</th>
<th>Concentration</th>
<th>Primary Std. Required For</th>
<th>Primary Std. Required For</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ALLO</td>
<td>AlHI</td>
<td>ALLO</td>
<td>AlHI</td>
</tr>
<tr>
<td></td>
<td>(ppm)</td>
<td>(ppm)</td>
<td>(mL)</td>
<td>(mL)</td>
</tr>
<tr>
<td>Al</td>
<td>100</td>
<td>200</td>
<td>50.0</td>
<td>100.0</td>
</tr>
<tr>
<td>Fe</td>
<td>75</td>
<td>150</td>
<td>37.5</td>
<td>75.0</td>
</tr>
<tr>
<td>Ti</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
<tr>
<td>Zr</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
<tr>
<td>Na</td>
<td>25</td>
<td>50</td>
<td>12.5</td>
<td>25.0</td>
</tr>
</tbody>
</table>

¹All calibration standards based on 500-ml final volume

Table 3.—Calibration standards for SILO and SIHI¹.

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration</th>
<th>Concentration</th>
<th>Primary Std. Required For</th>
<th>Primary Std. Required For</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SILO</td>
<td>SIHI</td>
<td>SILO</td>
<td>SIHI</td>
</tr>
<tr>
<td></td>
<td>(ppm)</td>
<td>(ppm)</td>
<td>(mL)</td>
<td>(mL)</td>
</tr>
<tr>
<td>Si</td>
<td>225</td>
<td>450</td>
<td>112.5</td>
<td>225.0</td>
</tr>
<tr>
<td>P</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
<tr>
<td>Ti</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
<tr>
<td>Se</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
<tr>
<td>As</td>
<td>5</td>
<td>10</td>
<td>2.5</td>
<td>5.0</td>
</tr>
</tbody>
</table>

¹All calibration standards based on 500-ml final volume

7.2 To the calibration standards and a blank, also add the following chemicals: 25.0 mL HF; 3.75 mL HNO₃; 1.25 mL HCl; and 12.5 g granular Boric Acid. Make all standards and the blank to a final 500-mL volume with DDI water.

7.3 Use the TJA ICAP 61E spectrophotometer and analyze for the following elements: Fe, Mn, Al, Ca, Mg, Na, K, P, Si, Zr, Cu, Zn, As, Ti, Se, Cd, and Pb. No initial dilutions of samples are necessary prior to analysis. Use polypropylene spray chamber and HF resistant torch on ICP. Check instrument alignment and gas pressures to obtain optimum readings with maximum signal to noise ratio. The torch tip used for HF digestions should not be run dry or used with RF powers exceeding 1350. The HF torch tip should only be used with high flow torch.
7.4 Use the HF blank standard solution to dilute those samples with concentrations greater than the high standard. Rerun all elements and use only the data needed from the diluted analysis.

7.5 Run the detection limits using the blank standard solution. These values establish the lower detection limits for each element. Analyzed values lower than the detection limits are set equal to zero.

7.6 When ICP analyses are completed, transfer data from the hard drive storage to a 3.5 inch floppy disk as an ASCII file via the “Report Writer” in the TJA software Thermospec, Version 5.06. These data are imported into a LOTUS 123, Version 3.1 spreadsheet for data analysis. Refer to method 7C4a.

8. Calculations
   Refer to method 7C4a.

9. Report
   Refer to method 7C4a.

10. Precision
    No precision data are yet available for this procedure.

11. References
    Refer to digestion procedure.

Organic Carbon, Iron, Manganese, and Aluminum (6A, 6C, 6D, and 6G)
Sodium Pyrophosphate Extraction (6A4)
   CO₂ Evolution Gravimetric (6A4a)
   Sodium Pyrophosphate Extraction (6C8, 6D4, and 6G10)
   Atomic Absorption
   Perkin-Elmer 5000 AA (6C8a, 6D4a, and 6G10a)

1. Application
   Sodium pyrophosphate (0.1 \( M \) \( Na_4P_2O_7 \)) is used as a selective dissolution extractant for organically complexed Fe and Al (Wada, 1989). The \( Na_4P_2O_7 \) solution is a poor extractant for allophane, imogolite, amorphous aluminosilicates, and noncrystalline hydrous oxides of Fe and Al. The \( Na_4P_2O_7 \) solution does not extract opal, crystalline silicates, layer silicates, and crystalline hydrous oxides of Fe and Al (Wada, 1989). In *Soil Taxonomy*, sodium pyrophosphate extractable organic C, Fe, and Al are criteria for spodic placement (Soil Survey Staff, 1975).
2. Summary of Method

The soil sample is mixed with 0.1 M Na$_4$P$_2$O$_7$ and shaken overnight. Superfloc 16 is added, and the mixture is made to volume. The solution is allowed to settle and a clear extract is obtained. The Na$_4$P$_2$O$_7$ extracted solution is diluted with distilled deionized (DDI) water. The diluted extract is aspirated into an atomic absorption spectrophotometer (AA). The analyte is measured by absorption of the light from a hollow cathode lamp. An automatic sample changer is used to aspirate a series of samples. The AA converts absorption to analyte concentration. Percent sodium pyrophosphate extractable Fe, Mn, and Al are reported in methods 6C8a, 6D4a, and 6G10a, respectively. The organic C in the sodium pyrophosphate extract is wet oxidized and gravimetrically measured in method 6A4a.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the AA analyses of these elements. These interferences vary in importance, depending upon the particular analyte selected.

The concentration of Na$_4$P$_2$O$_7$ solution must be close to 0.1 M. Variable amounts of Fe, Al, Mn, and organic C may be extracted by varying the pyrophosphate concentration.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±0.0001 g
5.2 Shaker, Eberbach 6000 power unit, reciprocating speed of 60 to 260 epm, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
5.3 Nursing bottle, 240 mL (8 fl. oz.), graduated
5.4 Rubber stoppers, No. 2, to fit nursing bottles
5.5 Dispenser/diluters, Repipet, 0 to 10 mL, Labindustries, 1802 2nd St., Berkeley, CA
5.6 Atomic absorption spectrophotometer (AA), model 5000, Perkin-Elmer Corp., Norwalk, CT
5.7 Automatic burner control, model 5000, Perkin-Elmer Corp., Norwalk, CT
5.8 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.9 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.10 Heated regulator, single-stage, nitrous oxide, stock number 808 8039, Airco Welding Products, P.O. Box 486, Union, NJ
5.11 Diluter/dispenser, Microlab 500, Catalogue No. 69052, Hamilton Co., Bonaduz, GR, Switzerland
5.12 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.13 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.14 Absorption bulb, Nesbitt with stopper
5.15 Absorption bulb, Stetser-Norton
5.16 Flask, boiling, round bottom, short neck
5.17 Condenser, Allihn
5.18 Funnel, separatory, cylindrical, open top, with stopcock
5.19 Tube, drying, Schwartz
5.20 Containers, polypropylene
5.21 Dot matrix printer, P-132, Interdigital Data Systems, Inc.

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Hydrochloric acid (HCl), concentrated, 12 N
6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI H₂O.
6.4 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI H₂O.
6.5 Superfloc 16, 0.2% solution (w:v). Dissolve 2 g of Superfloc 16 in 1 liter of DDI H₂O. Do not shake the mixture as this breaks the polymer chains of the Superfloc. Gently swirl the mixture occasionally over the several days that the solution requires to completely dissolve the Superfloc. Suggested source is American Cyanamid Co., P.O. Box 32787, Charlotte, NC.
6.6 Sodium pyrophosphate solution, 0.1 M. Dissolve 800 g of Na₄P₂O₇·H₂O in 16 L of DDI H₂O. Dilute to 18 L with DDI H₂O.
6.7 Primary mixed standard, 4000 mg L\(^{-1}\) (4000 ppm) Fe; 2000 mg L\(^{-1}\) (3000 ppm) Al; and 600 mg L\(^{-1}\) (600 ppm) Mn. Dissolve 4.000 g of Fe wire, 3.000 g of Al wire, and 0.600 g of Mn metal powder in 1:1 HCl:DDI in a glass beaker. When dissolved, transfer to a 1-L volumetric flask and fill with 1% HCl solution. Store in a polypropylene bottle.

6.8 High calibration mixed standards solution (HCMSS), 80 mg/8 oz (80 ppm) Fe; 12 mg/8 oz (12 ppm) Mn; and 60 mg/8 oz (60 ppm) Al. Pipette 20 mL of primary mixed standard into an 8-oz bottle. Add 10.55 g of Na\(_4\)P\(_2\)O\(_7\)•H\(_2\)O and 3.5 mL of concentrated H\(_3\)PO\(_4\). Dilute to 8 oz with DDI H\(_2\)O. Store in a polypropylene bottle.

6.9 Low calibration mixed standards solution (LCMSS), 40 mg/8 oz (40 ppm) Fe; 6 mg/8 oz (6 ppm) Mn; and 30 mg/8 oz (30 ppm) Al. Pipette 10 mL of primary mixed standard into an 8-oz bottle. Add 10.55 g of Na\(_4\)P\(_2\)O\(_7\)•H\(_2\)O and 3.5 mL of concentrated H\(_3\)PO\(_4\). Dilute to 8 oz with DDI H\(_2\)O. Store in a polypropylene bottle.

6.10 Calibration reagent blank solution (CRBS). Add 10.55 g of Na\(_4\)P\(_2\)O\(_7\)•H\(_2\)O and 3.5 mL of concentrated H\(_3\)PO\(_4\). Dilute to 8 oz with DDI H\(_2\)O. Store in a polypropylene bottle.

6.11 Potassium dichromate (K\(_2\)Cr\(_2\)O\(_7\)), reagent

6.12 Potassium iodide solution. Dissolve 100 g of KI in 100 mL of DDI H\(_2\)O.

6.13 Silver sulfate, saturated aqueous solution

6.14 Digestion acid mixture. Mix 600 mL of concentrated H\(_2\)SO\(_4\) and 400 mL of 85% H\(_3\)PO\(_4\).

6.15 Indicarb or Mikohlbite

6.16 Soda lime

6.17 Zinc granules, 300 mesh

6.18 Anhydrole

6.19 Acetylene gas, purity 99.6%

6.20 Nitrous oxide gas, compressed

6.21 Compressed air with water and oil traps

7. Procedure

Extraction of Fe, Mn, and Al

7.1 Weigh 2.00 g of <2-mm, air-dry soil sample and place in an 8-oz nursing bottle.

7.2 Add 0.1 \(M\) Na\(_4\)P\(_2\)O\(_7\) solution to 7-oz level on bottle and securely stopper bottle.

7.3 Shake overnight (12 to 16 h) in a reciprocating shaker. After shaking, use a dispenser to add 4 mL of Superfloc 16 solution.
7.4 Fill bottle to 8-oz volume with \( \text{Na}_4\text{P}_2\text{O}_7 \) solution.

7.5 Stopper and shake vigorously for \( \approx 15 \) s.

7.6 Allow to settle for at least 3 days (4 to 6 days typical). The Fe, Mn, and Al are determined from a clear aliquot of solution.

**Dilution of Sample Extracts and Standards**

7.7 No ionization suppressant is required as the Na in the extractant is present in sufficient quantity. Set the digital diluter on Hamilton diluter to 66 for diluent and 35 for sodium pyrophosphate sample extracts, calibration reagent blanks, and calibration standards for a 1:20 dilution as follows:

7.8 Dilute 1 part sample sodium pyrophosphate sample extract with 19 parts of DDI \( \text{H}_2\text{O} \) (1:20 dilution).

7.9 Dilute 1 part CRBS with 19 parts of DDI \( \text{H}_2\text{O} \) (1:20 dilution).

7.10 Dilute 1 part LCMSS with 19 parts of DDI \( \text{H}_2\text{O} \) (1:20 dilution).

7.11 Dilute 1 part HCMSS with 19 parts of DDI \( \text{H}_2\text{O} \) (1:20 dilution).

7.12 Dispense the diluted solutions into test tubes which have been placed in the sample holder of the sample changer.

**AA Calibration**

7.13 Use the calibration reagent blank and high calibration standard to calibrate the AA. The AA requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

7.14 Use the low calibration standard as a check sample.

**AA Set-up and Operation**

7.15 Refer to the manufacturer’s manual for operation of the Perkin-Elmer 5000 AA. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>248.8</td>
</tr>
<tr>
<td>Mn</td>
<td>280.1</td>
</tr>
<tr>
<td>Al</td>
<td>309.3</td>
</tr>
<tr>
<td>Analyte</td>
<td>Burner Head</td>
</tr>
<tr>
<td>---------</td>
<td>--------------</td>
</tr>
<tr>
<td>Fe</td>
<td>5-cm parallel</td>
</tr>
<tr>
<td>Mn</td>
<td>5-cm parallel</td>
</tr>
<tr>
<td>Al</td>
<td>5-cm parallel</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>10 C₂H₂/25 Air</td>
</tr>
<tr>
<td>Mn</td>
<td>10 C₂H₂/25 Air</td>
</tr>
<tr>
<td>Al</td>
<td>30 C₂H₂/17 Air</td>
</tr>
</tbody>
</table>

Typical read delay is 6 s, and the integration by peak area is 8 s.

7.16 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.17 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep matrix the same after dilution by diluting with DDI H₂O (1:20 dilution).

**Organic C Determination**

7.18 Pipette 100 mL of the extract into a 100-ml flask.

7.19 Evaporate the extract to near dryness using a 50 °C water bath and a gentle stream of clean, filtered air.

7.20 Construct the wet combustion apparatus. Refer to figure 6A4-1 of the apparatus for gravimetric organic C determination.

7.21 Add 1 to 2 g of potassium dichromate.

7.22 Wash the neck of the flask with 3 mL of DDI H₂O and connect to condenser.

7.23 Attach a weighed Nesbitt bulb to the system and open the valve at the top.

7.24 Pour 25 mL of digestion-acid mixture into the funnel. Add the mixture to the flask and immediately close the stopcock. Use the digestion-acid mixture to lubricate the stopcock.

7.25 The tip of the air-delivery tube should be ≈0.5 cm below the digestion-acid mixture. Adjust the flow of the “carrier stream” to maintain 1 to 2 bubble s⁻¹ rate throughout the digestion. Apply suction on the outlet side of the Nesbitt bulb. Gentle air pressure and needle valve on the air pressure line aids flow adjustment.

7.26 With a gas flame or a variable power heating mantle, gently heat the flask until the mixture boils (≈3 to 4 min). Continue a gentle boiling for 10 min. Heating is too rapid if white fumes of SO₂ are visible above the second bulb of the reflux condenser.
Figure 6A4-1.—Apparatus for gravimetric organic carbon determinations of 0.1 M sodium pyrophosphate extracts.

7.27 Remove the heat and allow to aerate for 10 additional min at a rate of 6 to 8 bubbles s\(^{-1}\).

7.28 Close the stopcock on the Nesbitt bulb, disconnect the bulb from the system, and weigh to the nearest 0.0001 g.

8. Calculations

Analyte (%) = \(\frac{AA \times DR \times AD}{OD} \times 100\) / (Sample Weight (g) \times 1000)

where:
- Analyte = Fe, Mn, and Al
- AA = Analyte concentration AA reading
- DR = Dilution ratio of 1 if no additional dilution
- Sample = Sample weight (g)
- 100 = Factor to convert to percent
1000 = Conversion factor (mg g⁻¹)
AD/OD = Air-dry/oven-dry ratio
Organic C (%) = \[(Wt_F - Wt_I) \times 27.3 \times Volume \times AD/OD\]/(Sample Weight (g) \times 236.6)

where:
Wt_f = Nesbitt bulb weight after digestion (g)
Wt_i = Nesbitt bulb weight before digestion (g)
Volume = Volume of extract digested (mL)
AD/OD = Air-dry/oven-dry ratio (method 4B5)
27.3 = Conversion factor
236.6 = Total volume of extract (mL)

9. Report
   Report percent sodium pyrophosphate extractable Fe, Mn, and Al on oven-dry soil basis to the nearest whole number.

10. Precision
   Precision data are not available for this procedure.

11. References

Iron, Manganese, Aluminum, Silicon, and Phosphate (6C, 6D, 6G, 6V, 6S)
Ammonium Oxalate Extraction (6C9, 6D5, 6D6, 6G12, 6V2, 6S8)
Inductively Coupled Plasma Spectrometry
   Thermo Jarrell Ash, ICAP 61E (6C9b, 6D5b, 6G12b, 6V2b, 6S8a)
Optical Density (8J) (of ammonium oxalate extract)

1. Application
   Oxalic acid-ammonium oxalate (acid oxalate) is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, allophane, and amorphous aluminosilicates (Wada, 1989). Acid oxalate
is a poor extractant of imogolite and layer silicates and does not extract crystalline hydrous oxides of Fe and Al, opal, or crystalline silicate (Wada, 1989). A more reliable and accurate estimation of soil properties and a better understanding of soil exchange complex is provided when acid oxalate extraction is used in conjunction with other selective dissolution procedures, thermal techniques, and chemical tests. In Soil Taxonomy, acid oxalate extractable Fe and Al are criteria for andic soil properties (Soil Survey Staff, 1999).

2. Summary of Method

A soil sample is extracted with a mechanical vacuum extractor (Holmgren et al., 1977) in a 0.2 M acid oxalate solution buffered at pH 3.0 under darkness. The acid oxalate extract is weighed. The acid oxalate extract is diluted with 0.002 M DDBSA. The analytes are measured by an inductively coupled plasma emission spectrophotometer (ICP). Data are automatically recorded by a microcomputer and printer. The percent acid oxalate extractable Fe, Mn, Al, Si, and P are reported in methods 6C9b, 6D5b, 6G12b, 6V2b, and 6S8a respectively. In method 8J, the optical density of the extract is measured with a UV spectrophotometer at 430 nm.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the ICP analyses of these elements. These interferences vary in importance, depending upon the particular analyte chosen.

The acid oxalate buffer extraction is sensitive to light, especially UV light. The exclusion of light reduces the dissolution effect of crystalline oxides and clay minerals. If the sample contains large amounts of amorphous material (>2% Al), an alternate method should be used, i.e., shaking with 0.275 M acid oxalate, pH 3.25, 1:100 soil:extractant.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the UV spectrophotometer and ICP.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.3 Mechanical vacuum extractor, Mavco Sampletek, 5300 N. 57th St., Lincoln, NE
5.4 Syringes, polypropylene, disposable, 60 mL, for extractant reservoir, extraction vessel, and tared extraction syringe

5.5 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm, (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels

5.6 Pre-pulped tubes

5.7 Extraction vessel, 60-mL, 10u polypropylene, part no. 6986-6010, Whatman Inc., 9 Bridewell Place, Clifton, NJ

5.8 Disposable glass tubes

5.9 UV-visible spectrophotometer, Carey-50, Varian Instruments

5.10 Cuvettes, disposable, polystyrene, 1-cm light path

5.11 Inductively coupled plasma spectrometer, ICAP 61E, Thermo Jarrell Ash Corp., Franklin, MA

5.12 Nebulizers, High-Solids, 41 psgi, Thermo Jarrell Ash Corp., Franklin, MA

5.13 RF generator, floor mounted power unit, Model 7/90, Thermo Jarrell Ash Corp., Franklin, MA

5.14 Computer, AT&T 386 Starstation, Model CPU-G72, and printer, NEC Pinwriter, P2200XE, Dot Matrix

5.15 ThermoSpec software, Thermo Jarrell Ash Corp., Franklin, MA

5.16 Line conditioner, Unity/I, Model UT8K, Best Power Technology, Inc., Necedah, WI

5.17 Single-stage regulator, high-purity, high-flow, argon, product no. E11-X-N145DHF, Air Products and Chemicals, Inc., Box 538, Allentown, PA

5.18 Autosampler, Thermo Jarrell Ash Corp., Franklin, MA

5.19 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV

5.20 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gastight, Hamilton Co., P.O. Box 10030, Reno, NV

5.21 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX

5.22 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Acid oxalate buffer solution, 0.2 M, pH 3.0. Solution A (base): Dissolve 284 g of (NH₄)₂C₂O₄ • 2H₂O in 10 L of DDI water. Solution B (acid): Dissolve 252 g of H₂C₂O₄ • H₂O in 10 L of DDI water. Mix 4 parts solution A with 3 parts solution B. Adjust acid oxalate solution pH by adding either acid or base solution. Store in a polypropylene bottle.
6.3 pH buffers, pH 4.00 and 7.00, for electrode calibration


6.5 Primary Al standard, 1000 ppm. Certified Reference Solution, Fisher Chemical Scientific Co., Fairlawn, N.J.

6.6 Primary Si standard, 1000 ppm. Certified Reference Solution, Fisher Chemical Scientific Co., Fairlawn, N.J.


6.9 Dodecylbenzenesulfonic acid (DDBSA), Tech 97%, 0.1 M. Dissolve 32.2 g DDBSA in 1-L DDI water.

6.10 DDBSA solution. Dilution solution for acid oxalate extracts. Add 40.0 mL of 0.1 M DDBSA and make to 2-L volume with DDI water (0.002 M DDBSA solution).

6.11 High calibration standard. Mix 60 mL of each primary standard (Si, Fe, and Al) with 10 mL of primary Mn standard and 20 mL of primary P standard in 1 L volumetric flask. Add 50 mL of 0.4 M acid oxalate solution and 16.0 mL of 0.1 M DDBSA and make to 1-L volume with DDI water. The elements are added in the order (Si, Fe, Al, Mn, P) to avoid element precipitation. Resulting solution contains 60 ppm each of Si, Fe, and Al, 10 ppm Mn, and 20 ppm P. Store in a polypropylene bottle.

6.12 Medium calibration standard. Mix 30 mL of each primary standard (Si, Fe, and Al) with 5 mL of primary Mn standard and 10 mL of primary P standard in 1 L volumetric flask. Add 50 mL of 0.4 M acid oxalate solution and 16.0 mL of 0.1 M DDBSA and make to 1-L volume with DDI water. Resulting solution contains 30 ppm each of Si, Fe, and Al, 5 ppm Mn, and 10 ppm P. Store in a polypropylene bottle.

6.13 Low calibration standard. Mix 10 mL of each primary standard (Si, Fe, and Al) with 2 mL of primary Mn standard, and 3 mL primary P standard in 1 L volumetric flask. Add 30 mL of 0.4 M acid oxalate solution and 16.0 mL of 0.1 M DDBSA and make to 1-L volume with DDI water. Resulting solution contains 10 ppm each of Si, Fe, and Al, 2 ppm Mn, and 3 ppm P. Store in a polypropylene bottle.

6.14 Calibration reagent blank solution. Add 50 mL of 0.4 M acid oxalate solution and 16.0 mL of 0.1 M DDBSA and make to 1-L volume with DDI water.

6.15 Argon gas, purity 99.9%
7. Procedure

**Extraction of Fe, Mn, Al, Si, and P**

7.1 Prepare disposable sample tubes.
7.2 Weigh 0.500 g of <2-mm, air-dry soil and place in sample tube. Prepare two reagent blanks (no sample in tube) per set of 48 samples.
7.3 Place the sample tube on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.
7.4 Use a dispenser to add 15.00 mL of acid oxalate buffer to the sample tube. Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.
7.5 Set extractor for 30-min extraction rate and extract until the acid oxalate buffer solution is at a 0.5 to 1.0-cm height above sample. Turn off extractor.
7.6 Put reservoir tube on top of the sample tube.
7.7 Add 35 mL of acid oxalate buffer to the reservoir tube.
7.8 Cover the extractor with a black plastic bag to exclude light. Adjust the extraction rate for a 12-h extraction.
7.9 After the extraction, shut off the extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the extraction vessel.
7.10 Weigh each syringe containing acid oxalate extract to the nearest 0.01 g.
7.11 Mix extract in each syringe by manually shaking. Fill a disposable tube with extract solution. This solution is reserved for determinations of Fe, Mn, Al, Si, and P. If optical density is to be measured, fill a disposable cuvette with extract solution. Discard excess solution.

**Determination of Optical Density of Extract**

7.12 Place 4 mL of acid oxalate extract in disposable cuvette.
7.13 Place 4 mL of acid oxalate reagent blank in disposable cuvette.
7.14 On Varian spectrophotometer, select a 430-nm wavelength. Select normal slit width and height. Refer to the manufacturer’s manual for operation of the spectrophotometer.
7.15 Use the acid oxalate extract reagent blank to set spectrophotometer.
7.16 Record optical density of acid oxalate extract to nearest 0.000.

**Dilution of Sample Extracts and Standards**

7.17 Dilute acid oxalate extracts (1:10) with 0.002 M DDBSA solution. Add 1 part acid oxalate sample extract with 10 parts dilution solution.
7.18 Set the digital settings of the Hamilton diluter for a 1:10 dilution. Calibration reagent blanks and calibration standards are not diluted.

7.19 Dispense the diluted solutions into test tubes which have been placed in the sample holder of the sample changer.

**ICP Calibration**

7.20 Use a multipoint calibration for ICP analysis of acid oxalate extracts. The ICP calibrates the blank first, low standard, medium standard, followed by the high standard. Prepare a quality control (QC) standard with analyte concentration between the high and low calibration standards. The ICP reads the QC after the high standard. If the QC falls within the range set by operator, the instrument proceeds to analyze the unknowns. If the QC is outside the range, the instrument restandardizes. The QC is analyzed approximately every 12 samples.

**ICP Set-up and Operation**

7.21 Refer to the manufacturer’s manual for operation of the ICP. The following parameters are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gas Flow</strong></td>
<td></td>
</tr>
<tr>
<td>Torch Gas</td>
<td>High Flow</td>
</tr>
<tr>
<td>Auxiliary Gas Flow</td>
<td>Medium 1.0 LPM</td>
</tr>
<tr>
<td>Nebulizer Pressure</td>
<td>41 psi</td>
</tr>
<tr>
<td><strong>Power</strong></td>
<td></td>
</tr>
<tr>
<td>Approximate RF Power</td>
<td>1150</td>
</tr>
<tr>
<td><strong>Peristaltic Pump</strong></td>
<td></td>
</tr>
<tr>
<td>Analysis Pump Rate</td>
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</tr>
<tr>
<td>Flush Pump Rate</td>
<td>200 RPM</td>
</tr>
<tr>
<td>Relaxation Time</td>
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</tr>
<tr>
<td>Pumping Tube Type</td>
<td>Silicone-Orange-3 stop</td>
</tr>
<tr>
<td><strong>Argon Flow Rate</strong></td>
<td>2.0 LPM</td>
</tr>
<tr>
<td>Purged Optical Pathway Enclosure Purge</td>
<td>2.0 SLPM Air</td>
</tr>
</tbody>
</table>
Nebulizer pressure depends on the type of nebulizer that is being used, i.e., low flow nebulizer requires a higher nebulizer pressure whereas a higher flow nebulizer requires a lower nebulizer pressure. To check for correct nebulizer pressure, aspirate with 1000.0 ppm yttrium. Adjust pressure to correct yttrium bullet.

7.22 Analyte data are reported at the following wavelengths.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>259.940</td>
</tr>
<tr>
<td>Al</td>
<td>167.081</td>
</tr>
<tr>
<td>Si</td>
<td>251.611</td>
</tr>
<tr>
<td>Mn</td>
<td>257.610</td>
</tr>
<tr>
<td>P</td>
<td>178.28</td>
</tr>
</tbody>
</table>

7.23 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings. The instrument readings are usually programmed in ppm.

7.24 If sample exceeds calibration standard, dilute the sample (1:5) with 0.4 M acid oxalate solution (matrix) and then dilute (1:10) with the DDBSA solution.

8. Calculations

8a. Calculations (Fe, Al, Si,)

\[
\text{Analyte (\%)} = \frac{\text{ICP} \times (\text{Syr}_{\text{fin}} - \text{Syr}_{\text{init}}) \times \text{D.R.} \times \text{AD/OD}}{\text{Sample Weight (g)} \times 10,000 \times \text{Density}}
\]

where:
- ICP = ICP analyte concentration (ppm)
- Syr\(_{\text{fin}}\) = Weight of syringe + extract (g)
- Syr\(_{\text{init}}\) = Tare weight of syringe (g)
- D.R. = Dilution ratio of samples over calibration range
- Density = Density of acid oxalate solution (1.007)
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

8b. Calculations (Mn, P)

\[
\text{Analyte (ppm)} = \frac{\text{ICP} \times (\text{Syr}_{\text{fin}} - \text{Syr}_{\text{init}}) \times \text{D.R.} \times \text{AD/OD}}{\text{Sample Weight (g)} \times \text{Density}}
\]
where:
ICP = ICP analyte concentration (ppm)
Syr_{fin} = Weight of syringe + extract (g)
Syr_{init} = Tare weight of syringe (g)
D.R. = Dilution ratio of samples over calibration range
Density = Density of acid oxalate solution (1.007)
AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report
Report the percent acid oxalate extractable Fe, Al, and Si to the nearest 0.01%. Report the concentration of acid oxalate extractable Mn and P in ppm. Report the optical density of the acid oxalate extract to the nearest 0.001 unit.

10. Precision
Precision data are not available for this procedure.

11. References

Manganese and Aluminum (6D and 6G)
KCl, Automatic Extractor (6D3 and 6G9)
Inductively Coupled Plasma Spectrometry, Thermo Jarrell Ash, ICAP 61E (6D3b and 6G9c)

1. Application
The Al extracted by 1 N KCl approximates exchangeable Al and is a measure of the “active” acidity present in soils with a 1:1 water pH <5.5. Above pH 5.5, precipitation of Al occurs during analysis. This method does not measure the acidity component of hydronium ions (H_{3}O^{+}). If Al is present in measurable amounts, the hydronium is a minor component of the active acidity. Because the 1 N KCl extractant is an unbuffered salt and usually affects the soil pH one unit or less, the extraction is determined at or near the soil pH. The KCl extractable Al is related to the immediate lime requirement and existing CEC of the soil. The
"potential" acidity is better measured by the BaCl$_2$-TEA method (method 6H5a) (Thomas, 1982).

2. Summary of Method

A soil sample is leached with 1 N KCl using the mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed. The KCl extracted solution is diluted with distilled deionized water. The analytes are measured by inductively coupled plasma emission spectrophotometer (ICP). Data are automatically recorded by a microcomputer and printer. The Mn and Al are reported in mg kg$^{-1}$ (ppm) and meq 100 g$^{-1}$ oven-dry soil in methods 6D3b and 6G9c, respectively.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the ICP analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

The soil:extractant ratio must remain constant. A soil:extractant ratio of 1:10 (w:v) for batch procedures is most commonly used. Using a leaching technique, a 1:20 (w:v) ratio gives comparable results. If the sample size is changed, the amount of extractable Al is changed. No other significant interferences have been identified for this procedure.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the ICP.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.3 Mechanical vacuum extractor, Mavco Sampletek, 5300 N. 57th St., Lincoln, NE
5.4 Syringes, polypropylene, disposable, 60 mL, for extraction vessel, extractant reservoir and tared extraction syringe
5.5 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm, (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels
5.6 Analytical filter pulp, ash-free, Schleicher and Schuell, No. 289
5.7 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.8 Wash bottle, 20 mL, to dispense KCl
5.9 Polycons, Richards Mfg. Co.
5.10 Inductively coupled plasma spectrometer, ICAP 61E, Thermo Jarrell Ash Corp., Franklin, MA
5.11 RF generator, floor mounted power unit, Model 7/90, Thermo Jarrell Ash Corp., Franklin, MA
5.12 Computer, AT&T 386 Starstation, Model CPU-G72, and printer, NEC Pinwriter, P2200XE, Dot Matrix
5.13 ThermoSpec software, Thermo Jarrell Ash Corp., Franklin, MA
5.14 Line conditioner, Unity/I, Model UT8K, Best Power Technology, Inc., Necedah, WI
5.15 Single-stage regulator, high-purity, high-flow, argon, product no. E11-X-N145DHF, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.16 Autosampler, Thermo Jarrell Ash Corp., Franklin, MA
5.17 Nebulizers, Precision Glass, Type A, 2.3 mlpm, 35 psig, Precision Glass Co., 14775 Hinsdale Ave., Englewood, CO.
5.18 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.19 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gastight, Hamilton Co., P.O. Box 10030, Reno, NV
5.20 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.21 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Potassium chloride solution (KCl), 1.0 N. Dissolve 1342 g of KCl reagent in 16 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 18 L with DDI water. Use 1.0 N KCl for Al and Mn extraction.
6.3 Potassium chloride solution (KCl), 2.0 N. Dissolve 298.24 g of KCl reagent in 1.5 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 2 L with DDI water. Use 2.0 N KCl for standards.
6.4 Primary Al standard, 1000 ppm (111 meq L⁻¹). Certified Reference Solution, Fisher Chemical Scientific Co., Fairlawn, N.J.
6.5 Primary Mn standard, 1000 ppm (250 meq L⁻¹). Certified Reference Solution, Fisher Chemical Scientific Co., Fairlawn, N.J.
6.6 High calibration Al and Mn standard, 10 meq L⁻¹ and 8 ppm, respectively. Pipette 22.476 mL of primary Al standard into a 250-mL volumetric flask. Add 125.0 mL 2 N KCl solution to the flask and mix. Pipette 2.0 mL of primary Mn standard into flask and mix. Dilute to volume with DDI water.
6.7 Calibration reagent blank solution, 1.0 \( N \) KCl. Add 125 mL of 2.0 \( N \) KCl to a volumetric flask and make to 250-mL volume with DDI water. Store in polypropylene container.

6.8 Calibration Al and Mn check standard, 5 meq L\(^{-1}\) and 3.0 ppm, respectively. Pipette 11.24 mL of primary Al standard into a 250-mL volumetric flask. Add 125.0 mL 2 \( N \) KCl solution to the flask and mix. Pipette 0.75 mL of primary Mn standard into flask and mix. Dilute to volume with DDI water.

6.9 Dodecylbenzenesulfonic acid (DDBSA), tech 97\%, 0.1 \( M \). Dissolve 32.2 g DDBSA in 1-L DDI water.

6.10 DDBSA rinse solution. Dilute 40.0 mL 0.1 \( M \) DDBSA to 2-L volume with DDI water.

6.11 Argon gas, purity 99.9%.

7. Procedure

**Extraction of Al and Mn**

7.1 Prepare extraction vessel by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh exactly 2.50 g of <2-mm, air-dry soil and place in an extraction vessel. Prepare one quality control check sample per 48 samples.

7.3 Place the extraction vessel on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.

7.4 Use a squeeze bottle and fill extraction vessel to the 20-mL mark with 1.0 \( N \) KCl solution (≈10 mL). Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.

7.5 Put reservoir tube on top of the extraction vessel. Set extractor for fast extraction rate and extract until the KCl solution is at a 0.5- to 1.0-cm height above sample. Turn off extractor.

7.6 Add 45 mL KCl solution to reservoir tube. Set extractor for 45-min extraction.

7.7 After the extraction, shut off extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the extraction vessel.

7.8 Weigh each syringe containing KCl extract to the nearest 0.01 g.

7.9 Mix extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. This solution is reserved for extractable Al and Mn analyses.
Dilution of Extracts and Standards

7.10 Set the digital settings at a 1:5 dilution for the KCl sample extracts, calibration reagent blanks, calibration standards, and calibration check standards as follows:

7.11 Dilute 1 part KCl sample extract with 4 parts of DDI water (1:5 dilution).
7.12 Dilute 1 part calibration reagent blank with 4 parts of DDI water (1:5 dilution).
7.13 Dilute 1 part calibration standard with 4 parts of DDI water (1:5 dilution).
7.14 Dilute 1 part calibration check standard with 4 parts of DDI water (1:5 dilution).
7.15 Dispense the diluted solutions into test tubes and place in the sample holder of the sample changer.

ICP Calibration

7.16 Use the calibration reagent blank (1.0 N KCl), high standard (10 meq L⁻¹ Al and 8 ppm Mn), and the blank to calibrate the ICP.
7.17 Use the calibration check standard (5 meq L⁻¹ Al and 3 ppm Mn) as a check sample. Perform a calibration check every 12 samples.

ICP Set-up and Operation

7.18 Refer to the manufacturer’s manual for operation of the ICP. The following parameters are only very general guidelines for instrument conditions for the analytes.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas Flow</td>
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</tr>
<tr>
<td>Torch Gas</td>
<td>High Flow</td>
</tr>
<tr>
<td>Auxiliary Gas Flow</td>
<td>Medium 1.0 LPM</td>
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<tr>
<td>Nebulizer Pressure</td>
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</tr>
<tr>
<td>RF Power</td>
<td>1150</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Wavelength (nm)</th>
<th>High/Low</th>
<th>Peak (Offset)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>167.081</td>
<td>0/-21</td>
<td>-2</td>
</tr>
<tr>
<td>Mn</td>
<td>257.610</td>
<td>0/-1</td>
<td>0</td>
</tr>
</tbody>
</table>
7.19 Determine a set of 24 unknown samples for each successful calibration check.

7.20 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.21 If a sample exceeds the calibration standard, dilute the sample (1:5) as follows.

7.22 Analyze 4 quality control check sample for every 48 samples.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L\(^{-1}\) Al and ppm Mn) in undiluted extract. Use these values to calculate the analyte concentration on an oven-dry soil basis (meq 100 g\(^{-1}\)).

\[
\text{Analyte (meq 100 g}^{-1}\) = \frac{\text{ICP} \times (\text{Wt}_{syr+ext} - \text{Wt}_{syr}) \times \text{D.R.} \times 100 \times \text{AD/OD}}{\text{Sample Weight} \times 1.0412 \times 1000}
\]

where:
- ICP = ICP analyte reading
- Wt\(_{syr+ext}\) = Weight of extraction syringe & extract (g)
- Wt\(_{syr}\) = Weight of tared extraction syringe (g)
- D.R. = Dilution ratio of samples over calibration range
- 1.0412 = Density of 1 N KCl @ 20 °C
- 1000 = g L\(^{-1}\)
- 100 = Conversion factor (100-g basis)
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report KCl extractable Al and Mn in units of meq 100 g\(^{-1}\) of oven-dry soil to the nearest 0.01 meq 100 g\(^{-1}\).

10. Precision

Precision data are not available for this procedure.

11. References


Manganese and Aluminum (6G)
KCl, Automatic Extractor (6G9)
Atomic Absorption
  Thermo Jarrell Ash, Smith Hieftje 4000 (6D3c and 6G9d)

1. Application
   The Al extracted by 1 N KCl approximates exchangeable Al and is a measure of the "active" acidity present in soils with a 1:1 water pH <5.5. Above pH 5.5, precipitation of Al occurs during analysis. This method does not measure the acidity component of hydronium ions (H$_3$O$^+$). If Al is present in measurable amounts, the hydronium is a minor component of the active acidity. Because the 1 N KCl extractant is an unbuffered salt and usually affects the soil pH one unit or less, the extraction is determined at or near the soil pH. The KCl extractable Al is related to the immediate lime requirement and existing CEC of the soil. The “potential” acidity is better measured by the BaCl$_2$-TEA method (method 6H5a) (Thomas, 1982).

2. Summary of Method
   A soil sample is leached with 1 N KCl using the mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed. The KCl extract is diluted with distilled deionized (DDI) water. The analytes are measured by an atomic absorption spectrophotometer. The data are automatically recorded by a microcomputer and printer. The Al and Mn are reported in meq 100 g$^{-1}$ and mg kg$^{-1}$ (ppm) oven-dry soil in method 6D3c and 6G9d.

3. Interferences
   There are four types of interferences (matrix, spectral, chemical, and ionization) in the AA analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.
   The soil:extractant ratio must remain constant. A soil:extractant ratio of 1:10 (w:v) for batch procedures is most commonly used. Using a leaching technique, a 1:20 (w:v) ratio gives comparable results. If the sample is changed, the amount of extractable Al is changed. No other significant interferences have been identified for this procedure.

4. Safety
   Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets.
and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.3 Mechanical vacuum extractor, Mavco Sampletek, 5300 N. 57th St., Lincoln, NE
5.4 Syringes, polypropylene, disposable, 60 mL, for extraction vessel, extractant reservoir, and tared extraction syringe
5.5 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm, (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels
5.6 Analytical filter pulp, ash-free, Schleicher and Schuell, No. 289
5.7 Wash bottle, 20 mL, to dispense KCl
5.8 Polycons, Richards Mfg. Co.
5.9 Atomic absorption spectrophotometer (AA), Smith-Hieftje Model 4000, Thermo Jarrell Ash Corp., Franklin, MA
5.10 Autosampler, Model 150, Thermo Jarrell Ash Corp., Franklin, MA
5.11 ThermoSpec software, Version 3.01, Enable 4.0, DOS 5.0, Thermo Jarrell Ash Corp., Franklin, MA
5.12 Computer, CUi Advantage 486, Thermo Jarrell Ash Corp., Franklin, MA
5.13 Printer, NEC Pinwriter P3200
5.14 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.15 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.16 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.17 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.18 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Potassium chloride solution (KCl), 1.0 N. Dissolve 1342 g of KCl reagent in 16 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 18 L with DDI water. Use 1.0 N KCl solution for Al extraction.
6.3 Potassium chloride solution (KCl), 2.0 N. Dissolve 298.24 g of KCl reagent in 1.5 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 2 L with DDI water. Use 2.0 N KCl solution for standards.


6.5 Primary Mn standard, 1000 ppm. Certified Reference Solution, Fisher Chemical Scientific Co., Fairlawn, N.J.

6.6 High calibration Al (4 meq L\(^{-1}\)) and Mn (3 ppm) standard. Mix 9 mL of primary Al standard with 125 mL 2 N KCl. Add 0.75 mL of primary Mn standard and make to 250-mL volume with DDI water.

6.7 Low calibration Al (0.2 meq L\(^{-1}\)) and Mn (0.2 ppm) standard. Mix 1.8 mL of primary Al standard with 125 mL 2 N KCl. Add 0.05 mL of primary Mn standard and make to 250-mL volume with DDI water.

6.8 Calibration Al (2 meq L\(^{-1}\)) and Mn (1.5 ppm) check standard. Mix 4.5 mL of primary Al standard with 125 mL 2 N KCl. Add 0.375 mL of primary Mn standard and make to 250-mL volume with DDI water.

6.9 Calibration reagent blank solution, 1.0 N KCl. Add 125 mL of 2.0 N KCl to a volumetric flask and make to 250-mL volume with DDI water.

6.10 Nitrous oxide gas, compressed

6.11 Acetylene gas, compressed, purity 99.6%

6.12 Compressed air with water and oil traps

7. Procedure

**Extraction of Al**

7.1 Prepare extraction vessel by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh exactly 2.50 g of <2-mm, air-dry soil and place in an extraction vessel. Prepare one quality control check sample per 48 samples.

7.3 Place the extraction vessel on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.

7.4 Use a squeeze bottle and fill extraction vessel to the 20-mL mark with 1.0 N KCl solution (≈10 mL). Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.

7.5 Put reservoir tube on top of the extraction vessel. Set extractor for fast extraction rate and extract until the KCl solution is at a 0.5- to 1.0-cm height above sample. Turn off extractor.
7.6 Add 45 mL KCl solution to reservoir tube. Set extractor for 45-min extraction.

7.7 After the extraction, shut off extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the extraction vessel.

7.8 Weigh each syringe containing KCl extract to the nearest 0.01 g.

7.9 Mix extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. This solution is reserved for extractable Al and Mn analyses.

### Dilution of Sample Extracts and Standards

7.10 No ionization suppressant is required as the K in the extractant is present in sufficient quantity. Set the digital settings at a 1:2 dilution for the KCl sample extracts, calibration reagent blanks, calibration standards, and calibration check standards as follows:

7.11 Dilute 1 part KCl sample extract with 1 part of DDI water (1:2 dilution).

7.12 Dilute 1 part calibration reagent blank with 1 part of DDI water (1:2 dilution).

7.13 Dilute 1 part calibration standard with 1 part of DDI water (1:2 dilution).

7.14 Dilute 1 part calibration check standard with 1 part of DDI water (1:2 dilution).

7.15 Dispense the diluted solutions into test tubes and place in the sample holder of the sample changer.

### AA Calibration

7.16 Use the calibration reagent blank (1.0 N KCl), high standard (4 meq L\(^{-1}\) Al and 3 ppm Mn), and the low standard (2 meq L\(^{-1}\) Al and 1.5 ppm Mn) to calibrate the AA.

7.17 Use the calibration check standard (2 meq L\(^{-1}\) Al and 1.5 ppm Mn) as a check sample. Perform a calibration check every 12 samples.

### AA Set-up and Operation

7.18 Refer to the manufacturer’s manual for operation of the AA. The following parameters are only very general guidelines for instrument conditions for the analyte.

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength (nm)</th>
<th>Burner Head &amp; Angle</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>309.3</td>
<td>5-cm, parallel</td>
<td>20 C(_2)H(_2)/10 N(_2)O</td>
</tr>
<tr>
<td>Mn</td>
<td>280.1</td>
<td>5-cm, parallel</td>
<td>4 C(_2)H(_2)/16 Air</td>
</tr>
</tbody>
</table>
Typical read delay is 6 s, and integration by peak area is 8 s.

7.19 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.20 If a sample exceeds the calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with DDI water (1:2 dilution).

7.21 Analyze one quality control check sample for every 48 samples.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L\(^{-1}\) AI) in undiluted extract. Use these values to calculate the analyte concentration on an oven-dry soil basis (meq 100 g\(^{-1}\)).

\[
\text{Al (meq 100 g}^{-1}\text{)} = \left[\text{AAAl} \times (\text{Wt}_{\text{sy+ext}} - \text{Wt}_{\text{sy}}) \times \text{D.R.} \times 100 \times \text{AD/OD}\right] / \left[\text{Sample Weight (g)} \times 1.0412 \times 1000\right]
\]

where:
- AAAl = AA Al reading (meq L\(^{-1}\))
- Wt\(_{\text{sy+ext}}\) = Weight of extraction syringe and extract (g)
- Wt\(_{\text{sy}}\) = Weight of tared extraction syringe (g)
- D.R. = Dilution ratio for samples over calibration range
- 1.0412 = Density of 1 N KCl @ 20 °C
- 100 = Conversion factor (100-g basis)
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report KCl extractable Al in units of meq 100 g\(^{-1}\) of oven-dry soil to the nearest 0.1 meq 100 g\(^{-1}\).

10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples.

11. References


Chloride, Sulfate, Nitrate, Fluoride, and Nitrite (6K, 6L, 6M, 6U, and 6W)
Saturation Extract (6K1, 6L1, 6M1, 6U1, and 6W1)
Chromatograph, Anion Suppressor
   Dionex 2110i Ion Chromatograph (6K1d, 6L1d, 6M1d, 6U1b, and 6W1b)

1. Application
   The soluble anions that are commonly determined in saline and alkali soils are carbonate, bicarbonate, sulfate, chloride, nitrate, nitrite, fluoride, phosphate, silicate, and borate (Khym, 1974; U.S. Salinity Laboratory Staff, 1954). Carbonate and bicarbonate are determined by titration. Phosphate, silicate, and borate usually are not determined because they are found only occasionally in measurable amounts in soils. Chloride, sulfate, nitrate, fluoride, and nitrite are measured in solution by chromatography. In saline and alkali soils, carbonate, bicarbonate, sulfate, and chloride are the anions that are found in the greatest abundance. In general, soluble sulfate is usually more abundant than soluble chloride.

2. Summary of Method
   The saturation extract is diluted according to its electrical conductivity (EC<sub>s</sub>). The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to measure the anion. A chart recording is made of the chromatograph. Standard anions are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. A computer program automates these actions. The saturated extract anions, Cl<sup>−</sup>, SO<sub>4</sub><sup>2−</sup>, NO<sub>3</sub>−, F<sup>−</sup>, and NO<sub>2</sub>− are reported in meq L<sup>−1</sup> in methods 6K1d, 6L1d, 6M1d, 6U1b, and 6W1b, respectively.

3. Interferences
   Some saturation extracts contain suspended solids. Filtering after dilution removes the particles. Saturation extracts of acid soils that contain Fe and/or Al may precipitate and clog the separator column. Saturation extracts of very high pH may contain organic material which may clog or poison the column. Low molecular weight organic anions will co-elute with inorganic anions from the column.

4. Safety
   Wear protective clothing and safety glasses. When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.
5. Equipment

5.1 Ion chromatograph, Series 2110i, dual-channel system
5.2 Analytical column, AS4A 4mm P/N 37041, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.3 Guard column, AG4A 4mm P/N 37042, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.4 Analytical pumps, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.5 Automated sampler, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.6 Conductivity detectors, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.7 Anion self-regenerating suppressor (ASRS-1) with controller (SRC-1), Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.8 Computer interfaces, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.9 Computer software, A1-450 Chromatography Software Program Release 3.32, Microsoft Windows Operating Environment; Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.10 Computer, DFI
5.11 Printer, Epson, Fx-850
5.12 Poly-vials, 5 mL, P/N 038008, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.13 Poly-vials, filter caps, 5 mL, P/N 038009, Dionex Corp., 1228 Titan Way, Sunnyvale, CA
5.14 Digital diluter/dispenser, Microlab 500, Hamilton C., P.O. Box 10030, Reno, NV
5.15 Syringes, gas tight, Hamilton 1001 DX and 1010-TEF LL, Hamilton Co., P.O. Box 10030, Reno, NV
5.16 Disposable 0.2-µm pore size, 25-mm filter assembly, Gelman Sciences, Inc., 674 South Wagner Road, Ann Arbor, MI 48106. Use for saturation extracts and standards.
5.17 Disposable 0.2-µm pore size, Ultipor N₆₆ DFA3001NAEY, Pall Trinity Micro Corp., Cortland, NY. Use for filtering distilled deionized (DDI) water.

6. Reagents

6.1 Distilled deionized (DDI) filtered water
6.2 Sulfuric acid (H₂SO₄), concentrated, reagent
6.3 Toluene
6.4 Isopropanol to de-gas column
6.5 Stock NaHCO₃ solution, 0.480 M. Mix 40.34 g of dried NaHCO₃ with filtered DDI water and dilute to 1-L volume.
6.6 Stock Na₂CO₃ solution, 0.5040 M. Mix 53.42 g of dried Na₂CO₃ with filtered DDI water and dilute to 1-L volume.
6.7 Working eluent solution. Mix 100 mL of 0.5040 M NaHCO$_3$ and 100 mL of 0.4800 M Na$_2$CO$_3$ with filtered DDI water and dilute to 20-L volume. Add 8 drops of toluene to retard microbial growth.

6.8 Primary SO$_4^{2-}$ standard, 0.5 M (1.0 N). Mix 17.7560 g of Na$_2$SO$_4$ with filtered DDI water and dilute to 250-mL volume.

6.9 Primary Cl$^-$ standard, 1.0 M (1.0 N). Add 18.6392 g of KCl with filtered DDI water and dilute to 250-mL volume.

6.10 Primary F$^-$ standard, 0.125 M (0.125 N). Add 1.3122 g of NaF with filtered DDI water and dilute to 250-mL volume.

6.11 Primary NO$_3^-$ standard, 1.0 M (1.0 N). Add 25.2770 g of KNO$_3$ with filtered DDI water and dilute to 250-mL volume.

6.12 Primary mixed standard. Prepare 1 primary mixed standard by taking aliquots of each of the proceeding primary standards and diluting the combined aliquots to a 1-L volume with working eluent as follows:

<table>
<thead>
<tr>
<th>Primary standards</th>
<th>Aliquot</th>
<th>Final volume w/ eluent</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mL)</td>
<td>(mL)</td>
<td>(meq L$^{-1}$)</td>
</tr>
<tr>
<td>Na$_2$SO$_4$</td>
<td>50</td>
<td>1000</td>
<td>50</td>
</tr>
<tr>
<td>KCl</td>
<td>10</td>
<td>1000</td>
<td>10</td>
</tr>
<tr>
<td>NaF</td>
<td>100</td>
<td>1000</td>
<td>12.5</td>
</tr>
<tr>
<td>KNO$_3$</td>
<td>30</td>
<td>1000</td>
<td>30</td>
</tr>
</tbody>
</table>

Add eight drops of toluene to primary mixed standard to retard microbial growth and store in a glass container.

6.13 Mixed calibration standards. Prepare 4 mixed calibration standards (0.5, 1.0, 3.0, and 7.0 readings) by taking aliquots of primary mixed standard and diluting each aliquot to 100-mL volume with working eluent as follows:

<table>
<thead>
<tr>
<th>Primary mixed standards</th>
<th>Final volume w/ eluent</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(mL)</td>
<td>SO$_4^{2-}$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(meq L$^{-1}$)</td>
</tr>
<tr>
<td>0.5</td>
<td>100</td>
<td>0.25</td>
</tr>
<tr>
<td>1.0</td>
<td>100</td>
<td>0.50</td>
</tr>
<tr>
<td>3.0</td>
<td>100</td>
<td>1.5</td>
</tr>
<tr>
<td>7.0</td>
<td>100</td>
<td>3.5</td>
</tr>
</tbody>
</table>

6.14 NaNO$_2$, Baker reagent grade, 99.5% purity
6.15 Primary NO$_2^-$ standard, 1 N (1000 meq L$^{-1}$). Mix 69.3568 g of reagent grade NaNO$_2$ with filtered DDI water and dilute to 1-L volume. Take 5 mL aliquot of primary NO$_2^-$ standard and dilute with 500 mL of filtered DDI water (10 meq L$^{-1}$). Add eight drops of toluene to primary NO$_2^-$ standard to retard microbial growth and store in a glass container.

6.16 NO$_2^-$ calibration standards. Prepare 4 NO$_2^-$ calibration standards (0.5, 1.0, 3.0, and 7.0 readings) by taking aliquots of primary NO$_2^-$ standard (10 meq L$^{-1}$) and diluting each aliquot to 100-mL volume with working eluent as follows:

<table>
<thead>
<tr>
<th>Primary standard meq L$^{-1}$</th>
<th>Final volume w/ eluent (mL)</th>
<th>NO$_2^-$ Concentration (meq L$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>100</td>
<td>0.5</td>
</tr>
<tr>
<td>1.0</td>
<td>100</td>
<td>1.0</td>
</tr>
<tr>
<td>3.0</td>
<td>100</td>
<td>3.0</td>
</tr>
</tbody>
</table>

7. Procedure

**Dilution of extracts**

7.1 To estimate the total soluble anion concentration (meq L$^{-1}$), multiply the EC$_s$ (method 8A3a) by 10. Subtract the CO$_3^{2-}$ and HCO$_3^-$ concentrations (methods 6I1b and 6J1b) from the total anion concentration. The remainder is the ≈ concentration (meq L$^{-1}$) of anions to be separated by ion chromatography.

\[
\text{Anion concentration (meq L}^{-1}\text{)} = \text{EC}_s \times 10 - (\text{HCO}_3^- + \text{CO}_3^{2-})
\]

Refer to Table 1 for dilution of saturation extract with the working eluent.

7.3 Place the diluted samples in the Poly-vials and cap with filter caps.

7.4 Place the mixed calibration standards in the Poly-vials.

**Set-up and Operation of Ion Chromatograph (IC)**

7.5 Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex 2110i ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. Typical operation parameters are as follows:
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conductivity cell range</td>
<td>3 μS cm(^{-1}) full scale to 100 μS cm(^{-1})</td>
</tr>
<tr>
<td>Auto offset</td>
<td>“On”</td>
</tr>
<tr>
<td>Analytical pump flow rate</td>
<td>2.0 to 2.5 mL min(^{-1})</td>
</tr>
<tr>
<td>Low pressure limit</td>
<td>100 psi</td>
</tr>
<tr>
<td>High pressure limit</td>
<td>1200 psi</td>
</tr>
<tr>
<td>Regenerant flow rate</td>
<td>3 to 4 mL min(^{-1})</td>
</tr>
<tr>
<td>Injector loop</td>
<td>0.50 mL</td>
</tr>
<tr>
<td>Air pressure</td>
<td>3 to 8 psi</td>
</tr>
</tbody>
</table>

7.6 Load the sample holder cassettes with the capped samples, standards, and check samples.

7.7 Refer to the manufacturer’s manual for the operation of chromatograph.

8. Calculations

**Calibration Calculations**

8.1 Use the peak height of each anion standard to either construct a calibrated curve to plot anion concentration or use a least squares analysis to calculate anion concentration. The analytes are reported in meq L\(^{-1}\).

8.2 *Calibration Curve*: Plot the peak height against the meq L\(^{-1}\) of each anion standard on graph paper. Construct the calibration curve by finding the “best” line that fits the plotted standards.

8.3 *Linear Squares Analysis*: Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. An example for the anion Cl\(^-\) is as follows:

<table>
<thead>
<tr>
<th>Cl(^-) concentration (meq L(^{-1}))</th>
<th>Y = 0.1</th>
<th>1.5</th>
<th>4.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak height</td>
<td>X = 8.43</td>
<td>170.0</td>
<td>441.5</td>
</tr>
</tbody>
</table>

Number of standards = n = 3

\[ \Sigma Y = 5.6 \quad \Sigma X = 619.93 \]
\[ \Sigma Y/n = Y = 1.866 \quad \Sigma X/n = X = 206.6433 \]
\[ \Sigma X_i Y_i = 2021.843 \quad \Sigma X_i^2 = 223893.31 \]
\[ \Sigma X \Sigma Y = 3471.608 \]
\[
b = \frac{\sum X_i Y_i - \sum X_i \sum Y_i / n}{\sum X_i^2 - (\sum X_i)^2 / n} = \frac{2021.843 - 1157.2027}{223893.31 - 128104.4} = 0.0090265
\]

\(b\) = slope of the line, i.e., the amount that \(Y\) changes when \(X\) changes by 1 unit.

The equation is as follows:
\[
Y = Y + b (X - X)
\]
\[
Y = 1.866 + 0.0090265 (X) - 1.8653
\]

**Analyte Calculation**

8.4 *Calibration curve*: Read the analyte concentration (meq L\(^{-1}\)) directly from the calibration curve.

**Table 1.—Dilution factor for saturated paste soil extracts based on EC readings.**

<table>
<thead>
<tr>
<th>ECs ((\text{mmhos cm}^{-1}))</th>
<th>Dilution Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00 to 0.55</td>
<td>4</td>
</tr>
<tr>
<td>0.56 to 0.65</td>
<td>5</td>
</tr>
<tr>
<td>0.66 to 0.75</td>
<td>6</td>
</tr>
<tr>
<td>0.76 to 0.85</td>
<td>7</td>
</tr>
<tr>
<td>0.86 to 0.95</td>
<td>8</td>
</tr>
<tr>
<td>0.96 to 1.05</td>
<td>9</td>
</tr>
<tr>
<td>1.06 to 1.20</td>
<td>10</td>
</tr>
<tr>
<td>1.21 to 1.40</td>
<td>15</td>
</tr>
<tr>
<td>1.41 to 1.50</td>
<td>25</td>
</tr>
<tr>
<td>1.51 to 1.60</td>
<td>30</td>
</tr>
<tr>
<td>1.61 to 1.80</td>
<td>40</td>
</tr>
<tr>
<td>1.81 to 2.00</td>
<td>50</td>
</tr>
<tr>
<td>2.01 to 2.30</td>
<td>60</td>
</tr>
<tr>
<td>2.31 to 2.60</td>
<td>70</td>
</tr>
<tr>
<td>2.61 to 3.10</td>
<td>80</td>
</tr>
<tr>
<td>EC&lt;sub&gt;s&lt;/sub&gt; (mmhos cm&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>Dilution Factor</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>3.11 to 3.55</td>
<td>90</td>
</tr>
<tr>
<td>3.56 to 4.05</td>
<td>100</td>
</tr>
<tr>
<td>4.06 to 4.60</td>
<td>120</td>
</tr>
<tr>
<td>4.61 to 5.20</td>
<td>140</td>
</tr>
<tr>
<td>5.21 to 5.85</td>
<td>150</td>
</tr>
<tr>
<td>5.86 to 6.55</td>
<td>160</td>
</tr>
<tr>
<td>6.56 to 7.30</td>
<td>180</td>
</tr>
<tr>
<td>7.31 to 8.00</td>
<td>200</td>
</tr>
<tr>
<td>8.01 to 9.00</td>
<td>225</td>
</tr>
<tr>
<td>9.01 to 10.00</td>
<td>240</td>
</tr>
<tr>
<td>10.01 to 11.50</td>
<td>270</td>
</tr>
<tr>
<td>11.51 to 13.00</td>
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</tr>
<tr>
<td>13.01 to 14.50</td>
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<tr>
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<td>320</td>
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<td>400</td>
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<tr>
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</tr>
<tr>
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<td>720</td>
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<td>800</td>
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<tr>
<td>36.01 to 40.00</td>
<td>900</td>
</tr>
<tr>
<td>40.01 to 44.00</td>
<td>1000</td>
</tr>
</tbody>
</table>
8.5 Linear regression: Put the peak height in the preceding equation and solve for analyte concentration (meq L\(^{-1}\)). Thus, if sample extract has 204 peak height, the preceding equation is as follows:

\[ Y = 1.866 + 0.0090265 \times (204) - 1.8653 = 1.84 \text{ meq L}^{-1} \]

8.6 Repeat the calibration set and analyte calculation for each anion.

8.7 The chromatograph software automatically calculates the analyte concentrations and prints a report of the results.

9. Report

Report the saturation extract anions in units of meq L\(^{-1}\) to the nearest 0.1 meq L\(^{-1}\).

10. Precision

Precision data are not available for this procedure.

11. References


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Calcium, Magnesium, Sodium, and Potassium (6N, 6O, 6P, and 6Q)

Saturation Extraction (6N1, 6O1, 6P1, and 6Q1)

Atomic Absorption

Perkin-Elmer AA 5000 (6N1b, 6O1b, 6P1b, and 6Q1b)

1. Application

The commonly determined soluble cations are Ca\(^{2+}\), Mg\(^{2+}\), Na\(^{+}\), and K\(^{+}\). In soils with a low saturation pH, measurable amounts of Fe and Al may be present. Determination of soluble cations is used to obtain the relations between total cation concentration and other properties of saline solutions such as electrical conductivity and osmotic pressure (U.S. Salinity Laboratory Staff, 1954). The relative concentrations of the various cations in the soil-water extracts also provide information on the composition of the exchangeable cations in the soil. Complete analyses of the soluble ions provide a means to determine total salt content of the soils and salt content at field-moisture conditions.
2. Summary of Method

The saturation extract from method 8A3a is diluted with an ionization suppressant (LaCl₃). The analytes are measured by an atomic absorption spectrophotometer (AA). The data are automatically recorded by a microcomputer and printer. The saturation extracted cations, Ca²⁺, Mg²⁺, Na⁺, and K⁺, are reported in meq L⁻¹ in methods 6N1b, 6O1b, 6P1b, and 6Q1b, respectively.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the analysis of these cations. These interferences vary in importance, depending upon the particular analyte selected.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Filter paper, pre-pleated, 185-mm diameter, Schleicher and Schuell
5.3 Atomic absorption spectrophotometer (AA), model 5000, Perkin-Elmer Corp., Norwalk, CT
5.4 Automatic burner control, model 5000, Perkin-Elmer Corp., Norwalk, CT
5.5 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.6 Dot matrix printer, P-132, Interdigital Data Systems, Inc.
5.7 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.8 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.9 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.10 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Mathesen Scientific, Inc., Houston, TX
5.11 Containers, polypropylene
6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Hydrochloric acid (HCl), concentrated 12 N

6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.

6.4 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.

6.5 NH₄OH, reagent grade, sp gr 0.90

6.6 Glacial acetic acid, 99.5%

6.7 Primary stock mixed standards solution (PSMSS). Dissolve 0.8759 g of oven-dry reagent grade calcium carbonate (CaCO₃) in a minimum of volume of 1:1 HCl:DDI. Add 0.2127 g of clean Mg ribbon dissolved in 1:1 HCl. Add 1.0956 g of dry reagent grade sodium chloride (NaCl) and 0.1864 g of dry reagent grade KCl. Transfer to a 250-mL volumetric and bring to volume with 1% HCl solution. Resulting solution contains 70 meq L⁻¹ (1403 ppm) Ca; 70 meq L⁻¹ (851 ppm) Mg; 75 meq L⁻¹ (1724 ppm) Na; 10 meq L⁻¹ (391 ppm) K. Store in a polypropylene container.

6.8 NH₄OAc solution, 1.0 N, pH 7.0, reagent blank. Mix 57 mL of glacial acetic acid in 600 mL DDI water. While stirring, carefully add 68 mL concentrated of NH₄OH. Cool and adjust pH to 7.0 using NH₄OH or acetic acid. Dilute to 1 L with DDI water. The NH₄OAc solution is used for extraction of cations (method 5A8c).

6.9 Working stock mixed standards solution (WSMSS). Dilute 20 mL of the PSMSS with 80 mL DDI water (1:5). Resulting solution contains 14 meq L⁻¹ (281 ppm) Ca; 14 meq L⁻¹ (170 ppm) Mg; 15 meq L⁻¹ (345 ppm) Na; 2 meq L⁻¹ (78 ppm) K. Store in a polypropylene container.

6.10 Stock lanthanum ionization suppressant solution, 65,000 ppm. Wet 152.4 g of lanthanum oxide (La₂O₃) with 100 mL DDI water. Slowly and cautiously add 500 mL of 6 N HCl to dissolve the La₂O₃. Cooling the solution is necessary. Dilute to 2 L with DDI water. Filter solution. Store in polypropylene container.

6.11 Lanthanum ionization suppressant solution, 6500 ppm. Dilute 200 mL of stock lanthanum ionization suppressant solution with 1800 mL of DDI water (1:10). Store in polypropylene container.

6.12 Dilute calibration mixed standards solution (DCMSS). Dilute 1 part of the WSMSS with 39 parts of the lanthanum solution (1:40). Resulting solution contains 0.35 meq L⁻¹ (7 ppm) Ca; 0.35 meq L⁻¹ (4 ppm) Mg; 0.375 meq L⁻¹ (9 ppm) Na; 0.05 meq L⁻¹ (2 ppm) K. Store in polypropylene container.

6.13 Dilute calibration reagent blank solution (DCRBS). Dilute 1 part of DDI water with 39 parts of the lanthanum solution (1:40). Store in polypropylene container.
6.14 Compressed air with water and oil traps
6.15 Acetylene gas, purity 99.6%

7. Procedure

Dilution of Sample Extracts and Standards

7.1 The 10-mL syringe is for diluent (lanthanum ionization suppressant solution). The 1-mL syringe is for saturation sample extracts (method 8A3a), calibration reagent blanks, and calibration standards. Set the digital diluter at 1:40 dilution for saturation sample extracts, reagent blanks, and calibration standards as follows:

7.2 Dilute 1 part saturation sample extract with 39 parts of lanthanum ionization suppressant solution (1:40 dilution).

7.3 Dilute 1 part WSMSS with 39 parts of lanthanum ionization suppressant solution (1:40 dilution). This dilution is the DCMSS. Refer to reagents section.

7.4 Dilute 1 part DDI water with 39 parts of lanthanum ionization suppressant solution (1:40 dilution). This dilution is the DCRBS. Refer to reagents section.

7.5 Dispense the diluted solutions into test tubes which have been placed in the sample holders of the sample changer.

AA Calibration

7.6 Use the DCRBS and the DCMSS to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

AA Set-up and Operation

7.7 Refer to the manufacturer’s manual for operation of the AA. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Conc. meq/L</th>
<th>Burner &amp; Angle</th>
<th>Wavelength</th>
<th>Slit</th>
<th>Fuel/Oxidant C₂H₂/Air</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca</td>
<td>14.00</td>
<td>50 cm, @ 0°</td>
<td>422.7</td>
<td>0.7</td>
<td>10/25</td>
</tr>
<tr>
<td>Mg</td>
<td>14.00</td>
<td>50 cm, @ 30°</td>
<td>285.2</td>
<td>0.7</td>
<td>10/25</td>
</tr>
<tr>
<td>K</td>
<td>2.00</td>
<td>50 cm, @ 0°</td>
<td>766.5</td>
<td>1.4</td>
<td>10/25</td>
</tr>
<tr>
<td>Na</td>
<td>15.00</td>
<td>50 cm, @ 30°</td>
<td>589.0</td>
<td>0.4</td>
<td>10/25</td>
</tr>
</tbody>
</table>
7.8 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.9 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with the lanthanum ionization suppressant solution (1:40 dilution).

7.10 Analyze one quality control check sample for every 48 samples.

7.11 The instrument readings are usually programmed in meq L\(^{-1}\). Record analyte readings to 0.01 meq L\(^{-1}\).

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L\(^{-1}\) cation) in undiluted extract. Use these values and dilution ratio (if any) and calculate the analyte concentration in meq L\(^{-1}\) cation.

\[
\text{Analyte Concentration in Soil (meq L}^{-1}\) = \text{Analyte AA reading (meq L}^{-1}\) \times \text{Dilution ratio (if any)}
\]

9. Report

Report the saturation extraction cations of Ca\(^{2+}\), Mg\(^{2+}\), Na\(^+\), and K\(^+\) in units of meq L\(^{-1}\) to the nearest 0.1 meq L\(^{-1}\).

10. Precision

Precision data are not available for this procedure.

11. References


Calcium, Magnesium, Sodium, and Potassium (6N, 6O, 6P, and 6Q) Saturation Extraction (6N1, 6O1, 6P1, and 6Q1) Atomic Absorption

Thermo Jarrell Ash, Smith-Hieftje AA 4000 (6N1c, 6O1c, 6P1c, and 6Q1c)

1. Application

The commonly determined soluble cations are Ca\(^{2+}\), Mg\(^{2+}\), Na\(^+\), and K\(^+\). In soils with a low saturation pH, measurable amounts of Fe and Al may be present. Determination of soluble cations is used to obtain the relations between total cation concentration and other properties of saline solutions.
such as electrical conductivity and osmotic pressure (U.S. Salinity Laboratory Staff, 1954). The relative concentrations of the various cations in the soil-water extracts also provide information on the composition of the exchangeable cations in the soil. Complete analyses of the soluble ions provide a means to determine total salt content of the soils and salt content at field-moisture conditions.

2. Summary of Method

The saturation extract from method 8A3a is diluted with an ionization suppressant (LaCl$_3$). The analytes are measured by an atomic absorption spectrophotometer (AA). The data are automatically recorded by a microcomputer and printer. The saturation extracted cations, Ca$^{2+}$, Mg$^{2+}$, Na$^+$, and K$^+$, are reported in meq L$^{-1}$ in methods 6N1c, 6O1c, 6P1c, and 6Q1c, respectively.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the analysis of these cations. These interferences vary in importance, depending upon the particular analyte selected.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Filter paper, pre-pleated, 185-mm diameter, Schleicher and Schuell
5.3 Atomic absorption spectrophotometer (AA), Smith-Hieftje Model 4000, Thermo Jarrell Ash Corp., Franklin, MA
5.4 Autosampler, Model 150, Thermo Jarrell Ash Corp., Franklin, MA
5.5 ThermoSpec software, Version 3.01, Enable 4.0, DOS 5.0, Thermo Jarrell Ash Corp., Franklin, MA
5.6 Computer, CUi Advantage 486, Thermo Jarrell Ash Corp., Franklin, MA
5.7 Printer, NEC Pinwriter P3200
5.8 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
5.9 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.10 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.11 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX
5.12 Containers, polypropylene

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Hydrochloric acid (HCl), concentrated 12 N
6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.
6.4 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.
6.5 NH₄OH, reagent grade, sp gr 0.90
6.6 Glacial acetic acid, 99.5%
6.7 Primary stock mixed standards solution (PSMSS). Dissolve 0.8759 g of oven-dry reagent grade calcium carbonate (CaCO₃) in a minimum of volume of 1:1 HCl:DDI. Add 0.2127 g of clean Mg ribbon dissolved in 1:1 HCl. Add 1.0956 g of dry reagent grade sodium chloride (NaCl) and 0.1864 g of dry reagent grade KCl. Transfer to a 250-mL volumetric and bring to volume with 1% HCl solution. Resulting solution contains 70 meq L⁻¹ (1403 ppm) Ca; 70 meq L⁻¹ (851 ppm) Mg; 75 meq L⁻¹ (1724 ppm) Na; 10 meq L⁻¹ (391 ppm) K. Store in a polypropylene container.
6.8 NH₄OAc solution, 1.0 N, pH 7.0, reagent blank. Mix 57 mL of glacial acetic acid in 600 mL DDI water. While stirring, carefully add 68 mL concentrated of NH₄OH. Cool and adjust pH to 7.0 using NH₄OH or acetic acid. Dilute to 1 L with DDI water. The NH₄OAc solution is used for extraction of cations (method 5A8c).
6.9 Working stock mixed standards solution (WSMSS). Dilute 20 mL of the PSMSS with 80 mL DDI water (1:5). Resulting solution contains 14 meq L⁻¹ (281 ppm) Ca; 14 meq L⁻¹ (170 ppm) Mg; 15 meq L⁻¹ (345 ppm) Na; 2 meq L⁻¹ (78 ppm) K. Store in a polypropylene container.
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6.11 Lanthanum ionization suppressant solution, 6500 ppm. Dilute 200 mL of stock lanthanum ionization suppressant solution with 1800 mL of DDI water (1:10). Store in polypropylene container.

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7. Procedure

**Dilution of Sample Extracts and Standards**

7.1 The 10-mL syringe is for diluent (lanthanum ionization suppressant solution). The 1-mL syringe is for saturation sample extracts (method 8A3a), calibration reagent blanks, and calibration standards. Set the digital diluter at 1:40 dilution for saturation sample extracts, reagent blanks, and calibration standards as follows:

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7.5 Dispense the diluted solutions into test tubes which have been placed in the sample holders of the sample changer.

**AA Calibration**

7.6 Use the DCRBS and the DCMSS to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.
**AA Set-up and Operation**

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**7.8** Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

**7.9** If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with the lanthanum ionization suppressant solution (1:40 dilution).

**7.10** Analyze one quality control check sample for every 48 samples.

**7.11** The instrument readings are usually programmed in meq L⁻¹. Record analyte readings to 0.01 meq L⁻¹.

**8. Calculations**

**8.1** The instrument readings are the analyte concentration (meq L⁻¹ cation) in undiluted extract. Use these values and dilution ratio (if any) and calculate the analyte concentration in meq L⁻¹ cation.

$$\text{Analyte Concentration in Soil (meq L}^{-1}\text{)} = \text{Analyte AA reading (meq L}^{-1}\text{)} \times \text{Dilution ratio (if any)}$$

**9. Report**

Report the saturation extraction cations of Ca²⁺, Mg²⁺, Na⁺, and K⁺ in units of meq L⁻¹ to the nearest 0.1 meq L⁻¹.

**10. Precision**

Precision data are not available for this procedure.

**11. References**

Calcium, Magnesium, Sodium, and Potassium (6N, 6O, 6P, and 6Q) 
NH₄OAC Extraction (6N2, 6O2, 6P2, and 6Q2) Atomic Absorption
Perkin-Elmer AA 5000 (6N2e, 6O2d, 6P2b, and 6Q2b)

1. Application
The extractable bases (Ca²⁺, Mg²⁺, Na⁺, and K⁺) from the NH₄OAC extraction (method 5A8c) are generally assumed to be those exchangeable bases on the cation exchange sites of the soil. The abundance of these cations usually occurs in the sequence of Ca²⁺ > Mg²⁺ > K⁺ > Na⁺. Deviation from this usual order signals that some factor or factors, e.g., free CaCO₃ or gypsum, serpentine (high Mg²⁺), or natric material (high Na⁺), have altered the soil chemistry. The most doubtful cation extractions with this method are Ca²⁺ in the presence of free CaCO₃ or gypsum and K⁺ in soils that are dominated by mica or vermiculite (Thomas, 1982).

2. Summary of Method
The NH₄OAc extract from method 5A8c is diluted with an ionization suppressant (LaCl₃). The analytes are measured by an atomic absorption spectrophotometer (AA). The analyte is measured by absorption of the light from a hollow cathode lamp. An automatic sample changer is used to aspirate a series of samples. The AA converts absorption to analyte concentration. The data are automatically recorded by a microcomputer and printer. The NH₄OAc extracted cations, Ca²⁺, Mg²⁺, Na⁺, and K⁺, are reported in meq 100 g⁻¹ oven-dry soil in methods 6N2e, 6O2d, 6P2b, and 6Q2b, respectively.

3. Interferences
There are four types of interferences (matrix, spectral, chemical, and ionization) in the analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

4. Safety
Wear protective clothing and safety glasses. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.
Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.
5. Equipment

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6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.
6.4 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.
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6.6 Glacial acetic acid, 99.5%
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6.11 Lanthanum ionization suppressant solution, 6500 ppm. Dilute 200 mL of stock lanthanum ionization suppressant solution with 1800 mL of DDI water (1:10). Store in polypropylene container.

6.12 Dilute calibration mixed standards solution (DCMSS). Dilute 1 part of the WSMSS with 39 parts of the lanthanum solution (1:40). Resulting solution contains 0.35 meq L⁻¹ (7 ppm) Ca; 0.35 meq L⁻¹ (4 ppm) Mg; 0.375 meq L⁻¹ (9 ppm) Na; 0.05 meq L⁻¹ (2 ppm) K. Store in polypropylene container.

6.13 Dilute calibration reagent blank solution (DCRBS). Dilute 1 part of DDI water with 39 parts of the lanthanum solution (1:40). Store in polypropylene container.

6.14 Compressed air with water and oil traps

6.15 Acetylene gas, purity 99.6%

7. Procedure

**Dilution of Sample Extracts and Standards**

7.1 The 10-mL syringe is for diluent (lanthanum ionization suppressant solution). The 1-mL syringe is for NH₄OAc sample extracts (method 5A8c), calibration reagent blanks, and calibration standards. Set the digital diluter at a 1:40 dilution for the NH₄OAc sample extracts, reagent blanks, and calibration standards as follows:

7.2 Dilute 1 part NH₄OAc sample extract with 39 parts of lanthanum ionization suppressant solution (1:40 dilution).

7.3 Dilute 1 part WSMSS with 39 parts of lanthanum ionization suppressant solution (1:40 dilution). This is the DCMSS. Refer to reagents section.

7.4 Dilute 1 part DDI water with 39 parts of lanthanum ionization suppressant solution (1:40 dilution). This is the DCRBS. Refer to reagents section.

7.5 Dispense the diluted solutions into test tubes which have been placed in the sample holders of the sample changer.
AA Calibration

7.6 Use the DCRBS and the DCMSS to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

AA Set-up and Operation

7.7 Refer to the manufacturer’s manual for operation of the AA. The following are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Conc. meq/L</th>
<th>Burner &amp; Angle</th>
<th>Wavelength</th>
<th>Slit</th>
<th>Fuel/Oxidant C₂H₂/Air</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca</td>
<td>14.00</td>
<td>50 cm, @ 0°</td>
<td>422.7</td>
<td>0.7</td>
<td>10/25</td>
</tr>
<tr>
<td>Mg</td>
<td>14.00</td>
<td>50 cm, @ 30°</td>
<td>285.2</td>
<td>0.7</td>
<td>10/25</td>
</tr>
<tr>
<td>K</td>
<td>2.00</td>
<td>50 cm, @ 0°</td>
<td>766.5</td>
<td>1.4</td>
<td>10/25</td>
</tr>
<tr>
<td>Na</td>
<td>15.00</td>
<td>50 cm, @ 30°</td>
<td>589.0</td>
<td>0.4</td>
<td>10/25</td>
</tr>
</tbody>
</table>

7.8 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.9 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with the lanthanum ionization suppressant solution (1:40 dilution).

7.10 Analyze one quality control check sample for every 48 samples.

7.11 The instrument readings are usually programmed in meq L⁻¹. Record analyte readings to 0.01 meq L⁻¹.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L⁻¹ cation) in undiluted extract. Use these values and calculate the analyte concentration on an oven-dry soil basis (meq 100 g⁻¹).

Analyte Concentration in Soil (meq 100 g⁻¹) = \( \frac{A \times B \times C \times E}{10 \times D} \)

where:

- \( A \) = Analyte concentration in extract (meq L⁻¹)
- \( B \) = Extract volume (mL). Refer to method 5A8c.
- \( = Weight of extract in syringe (g)/Density of 1 N \text{NH}_4\text{OAc} (1.0124 \text{ g cm}^{-3}) \)
C = Dilution ratio, if needed
D = Soil sample weight (g)
E = AD/OD ratio (method 4B5)

9. Report
Report the extractable $\text{Ca}^{2+}$, $\text{Mg}^{2+}$, $\text{Na}^+$, and $\text{K}^+$ in units of meq 100 g$^{-1}$ of oven-dry soil to the nearest 0.1 meq 100 g$^{-1}$.

10. Precision
Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples. The number of observations, mean, standard deviation, and C.V. for the quality control check sample are as follows:

<table>
<thead>
<tr>
<th>Cation</th>
<th>n</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca</td>
<td>85</td>
<td>18.4</td>
<td>0.95</td>
<td>5.1%</td>
</tr>
<tr>
<td>Mg</td>
<td>84</td>
<td>7.5</td>
<td>0.23</td>
<td>3.1%</td>
</tr>
<tr>
<td>K</td>
<td>81</td>
<td>2.04</td>
<td>0.10</td>
<td></td>
</tr>
</tbody>
</table>

11. References

Calcium, Magnesium, Sodium, and Potassium (6N, 6O, 6P, and 6Q) NH$_4$OAC Extraction (6N2, 6O2, 6P2, and 6Q2)
Atomic Absorption
Thermo Jarrell Ash, Smith-Hieftje 4000 (6N2f, 6O2e, 6P2c, and 6Q2c)

1. Application
The extractable bases ($\text{Ca}^{2+}$, $\text{Mg}^{2+}$, $\text{Na}^+$, and $\text{K}^+$) from the NH$_4$OAC extraction (method 5A8c) are generally assumed to be those exchangeable bases on the cation exchange sites of the soil. The abundance of these cations usually occurs in the sequence of $\text{Ca}^{2+} > \text{Mg}^{2+} > \text{K}^+ > \text{Na}^+$. Deviation from this usual order signals that some factor or factors, e.g., free CaCO$_3$ or gypsum, serpentine (high Mg$^{2+}$), or natric material (high Na$^+$), have altered the soil chemistry. The most doubtful cation extractions with this method are $\text{Ca}^{2+}$ in the presence of free CaCO$_3$ or gypsum and $\text{K}^+$ in soils that are dominated by mica or vermiculite (Thomas, 1982).
2. Summary of Method

   The NH₄OAc extract from method 5A8c is diluted with an ionization suppressant (LaCl₃). The analytes are measured by an atomic absorption spectrophotometer (AA). The data are automatically recorded by a microcomputer and printer. The NH₄OAc extracted cations, Ca²⁺, Mg²⁺, Na⁺, and K⁺, are reported in meq 100 g⁻¹ oven-dry soil in methods 6N2f, 6O2e, 6P2c, and 6Q2c, respectively.

3. Interferences

   There are four types of interferences (matrix, spectral, chemical, and ionization) in the analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

4. Safety

   Wear protective clothing and safety glasses. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts.

   Follow standard laboratory procedures when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

   5.1 Electronic balance, ±1-mg sensitivity
   5.2 Filter paper, pre-pleated, 185-mm diameter, Schleicher and Schuell
   5.3 Atomic absorption spectrophotometer (AA), Smith-Hieftje Model 4000, Thermo Jarrell Ash Corp., Franklin, MA
   5.4 Autosampler, Model 150, Thermo Jarrell Ash Corp., Franklin, MA
   5.5 ThermoSpec software, Version 3.01, Enable 4.0, DOS 5.0, Thermo Jarrell Ash Corp., Franklin, MA
   5.6 Computer, CUi Advantage 486, Thermo Jarrell Ash Corp., Franklin, MA
   5.7 Printer, NEC Pinwriter P3200
   5.8 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA
   5.9 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
   5.10 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.11 Test tubes, 15-mL, 16 mm x 100, for sample dilution and sample changer, Curtin Matheson Scientific, Inc., Houston, TX

5.12 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Hydrochloric acid (HCl), concentrated 12 N

6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of concentrated HCl to 1 part DDI water.

6.4 HCl, 1% wt. Carefully dilute 25 mL of concentrated HCl to 1 L with DDI water.

6.5 NH₄OH, reagent-grade, sp gr 0.90

6.6 Glacial acetic acid, 99.5%

6.7 Primary stock mixed standards solution (PSMSS). Dissolve 0.8759 g of oven-dry reagent grade calcium carbonate (CaCO₃) in a minimum of volume of 1:1 concentrated HCl:DDI water. Add 0.2127 g of clean Mg ribbon dissolved in 1:1 HCl. Add 1.0956 g of dry reagent grade sodium chloride (NaCl) and 0.1864 g of dry reagent grade KCl. Transfer to a 250-mL volumetric and bring to volume with 1% HCl solution. Resulting solution contains 70 meq L⁻¹ (1403 ppm) Ca; 70 meq L⁻¹ (851 ppm) Mg; 75 meq L⁻¹ (1724 ppm) Na; 10 meq L⁻¹ (391 ppm) K. Store in a polypropylene container.

6.8 NH₄OAc solution, 1.0 N, pH 7.0, reagent blank. Mix 57 mL of glacial acetic acid in 600 mL of DDI water. While stirring, carefully add 68 mL of concentrated NH₄OH. Cool and adjust pH to 7.0 using NH₄OH or acetic acid. Dilute to 1 L with DDI water. The NH₄OAc solution is used for extraction of cations (method 5A8c).

6.9 Working stock mixed standards solution (WSMSS). Dilute 20 mL of the PSMSS with 80 mL DDI water (1:5). Resulting solution contains 14 meq L⁻¹ (281 ppm) Ca; 14 meq L⁻¹ (170 ppm) Mg; 15 meq L⁻¹ (345 ppm Na); 2 meq L⁻¹ (78 ppm) K. Store in a polypropylene container.

6.10 Stock lanthanum ionization suppressant solution, 65,000 ppm. Wet 152.4 g lanthanum oxide (La₂O₃) with 100 mL DDI water. Slowly and cautiously add 500 mL of 6 N HCl to dissolve the La₂O₃. Cooling the solution is necessary. Dilute to 2 L with DDI water. Filter solution. Store in polypropylene container.

6.11 Lanthanum ionization suppressant solution, 6500 ppm. Dilute 200 mL of stock lanthanum ionization suppressant solution with 1800 mL of DDI water (1:10). Store in polypropylene container.

6.12 Dilute calibration mixed standards solution (DCMSS). Dilute 1 part of the WSMSS with 39 parts of the lanthanum solution (1:40). Resulting solution contains 0.35 meq L⁻¹ (7 ppm) Ca; 0.35 meq L⁻¹ (4 ppm) Mg; 0.375 meq L⁻¹ (9 ppm) Na; 0.05 meq L⁻¹ (2 ppm) K. Store in polypropylene container.
6.13 Dilute calibration reagent blank solution (DCRBS). Dilute 1 part of DDI water with 39 parts of the lanthanum solution (1:40). Store in polypropylene container.

6.14 Compressed air with water and oil traps

6.15 Acetylene gas, purity 99.6%

7. Procedure

**Dilution of Sample Extracts and Standards**

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7.4 Dilute 1 part DDI water with 39 parts of lanthanum ionization suppressant solution (1:40 dilution). This is the DCRBS. Refer to reagents section.

7.5 Dispense the diluted solutions into test tubes which have been placed in the sample holders of the sample changer.

**AA Calibration**

7.6 Use the DCRBS and the DCMSS to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. Perform one calibration, i.e., blank plus standard, for every 12 samples.

**AA Set-up and Operation**

7.7 Refer to the manufacturer’s manual for operation of the AA. The following are only very general guidelines for instrument conditions for the various analytes.

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7.10 Analyze one quality control check sample for every 48 samples.

7.11 The instrument readings are usually programmed in meq L$^{-1}$. Record analyte readings to 0.01 meq L$^{-1}$.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L$^{-1}$ cation) in undiluted extract. Use these values and calculate the analyte concentration on an oven-dry soil basis (meq 100 g$^{-1}$).

\[
\text{Analyte Concentration in Soil (meq 100 g}^{-1} \text{)} = \frac{(A \times B \times C \times E)}{(10 \times D)}
\]

where:

\( A = \) Analyte concentration in extract (meq L$^{-1}$)
\( B = \) Extract volume (mL). Refer to method 5A8c.
\( C = \) Weight of extract in syringe (g)/Density of 1 N NH$_4$OAc (1.0124 g cm$^{-3}$)
\( D = \) Soil sample weight (g)
\( E = \) AD/OD ratio (method 4B5)

9. Report

Report the extractable Ca$^{2+}$, Mg$^{2+}$, Na$^+$, and K$^+$ in units of meq 100 g$^{-1}$ of oven-dry soil to the nearest 0.1 meq 100 g$^{-1}$.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples.

11. References

Total Sulfur (6R)
SO₂ Evolution, Infrared (6R3)
LECO SC-444 Sulfur Analyzer (6R3c)

1. Application

Organic and inorganic S forms are found in soils, with the organic S fraction accounting for >95% of the total S in most soils from humid and semi-humid (Tabatabai, 1982). Mineralization of organic S and its conversion to sulfate by chemical and biological activity may serve as a source of plant available S. Total S typically ranges from 0.01 to 0.05% in most mineral soils. In organic soils, total S may be >0.05%.

In well-drained, well-aerated soils, most of the inorganic S normally occurs as sulfate. In marine tidal flats, other anaerobic marine sediments, and mine spoils, there are usually large amounts of reduced S compounds which oxidize to sulfuric acid upon exposure to the air. In arid regions, significant amounts of inorganic S are found as sulfates such as gypsum and barite.

The typical use of total S is as an index of the total reserves of this element, which may be converted to plant available S. The SSL uses the combustion technique (LECO sulfur analyzer) for analysis of total S (method 6R3b). Extractable sulfate S (SO₄²⁻-S) is an index of readily plant-available S. Reagents that have been used for measuring SO₄²⁻-S include water, hot water, ammonium acetate, sodium carbonate and other carbonates, ammonium chloride and other chlorides, potassium phosphate and other phosphates, and ammonium fluoride (Bray-1). Extractable SO₄²⁻-S does not include the labile fraction of soil organic S that is mineralized during the growing season (Tabatabai, 1982). Extraction reagents for organic S include hydrogen peroxide, sodium bicarbonate, sodium hydroxide, sodium oxalate, sodium peroxide, and sodium pyrophosphate. There are other methods available for determination of soil S, especially for total S and SO₄²⁻-S. The investigator may refer to the review by Beaton et al. (1968).

2. Summary of Method

A fine-ground (<80-mesh) soil sample is oxidized at high temperature. The gases released are scrubbed, and the SO₂ in the combustion gases are measured using an infrared detector. Percent S is reported on an oven-dry soil basis.

3. Interferences

No significant interferences are known to affect the oxidizable S measurement.

4. Safety

Wear protective clothing and safety glasses. Magnesium perchlorate may form explosive mixtures. Magnesium perchlorate may contain traces of perchloric acid,
which remain from manufacturer’s operations. This acid is anhydrous because of the strong desiccating capability of the salt. Avoid prolonged contact with oxidizable material or material capable of forming unstable perchlorate esters or salts. Remove magnesium perchlorate by using an excess of water to thoroughly dilute the material.

The use of high temperatures in the oxidation of samples requires that extreme caution be used to prevent burns and fires. Follow standard laboratory procedures when handling compressed gases. Oxygen is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the sulfur analyzer.

5. Equipment

5.1 Sulfur analyzer, Leco Model SC-444, Sulfur and Carbon Analyzers, Leco Corp., St. Joseph, MI
5.2 Combustion boats, part no. 529-203, Leco Corp., St. Joseph, MI
5.3 Single-stage regulator, Oxygen Service, Part No. E11-W-N115BOX, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.4 Electronic balance, ±1-mg sensitivity

6. Reagents

6.1 Anhydrone, anhydrous magnesium perchlorate, granular
6.2 Glass wool
6.3 Compressed oxygen, >99.5% @ 30 psi
6.4 Calcium carbonate, CaCO₃, reagent grade
6.6 Soil Calibration Sample, part no. 502-062, Leco Corp., St. Joseph, MI

7. Procedure

7.1 Use a fine-ground 80-mesh, air-dry soil.
7.2 Prepare instrument as outlined in the operator’s instruction manual (Leco, 1994; Leco, 1993).
7.3 Methods are created with the method menu and stored in the instrument memory. System parameters are set as follows:
   Furnace operating temperature: 1450 °C
   Lance delay: 20 s
   Analysis time settings: 120 to 300 s
   Comparator level settings: 0.3%
7.3 Condition instrument by analyzing a few soil samples, until readings are stable.

7.4 Calibrate instrument by analyzing at least three replicates of each calibration standard. Use the soil calibration standard for samples with less than 0.01 percent TS and the sulfur standard for samples with more than 0.01 percent TS. Weigh standards in a range from 0.2 to 0.5 g.

7.5 Load samples on autoload rack, place in the analyzer, and press analyze key.

7.6 Weigh 0.2 to 0.5 g sample in a tared combustion boat. Add approximately 1 g of solid/powder combustion controller to sample.

7.7 Load samples on autoload rack, place in the analyzer, and press analyze key.

7.8 Repack the reagent (anhydrous magnesium perchlorate) tubes whenever the reagent becomes caked or moist or the warning alarm displays.

8. Calculations

$$S(\%) = S_i \times \frac{AD}{OD}$$

where:

- $S(\%)$ = S (%) on oven-dry basis
- $S_i$ = S (%) instrument
- $AD/OD$ = air-dry/oven-dry ratio (method 4B5)

9. Report

Report total S as a percentage of oven-dry weight to the nearest 0.1%.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run in every batch of 12 samples. A blank (crucible only) and a rerun of one of the 12 samples (unknowns) also are run in every batch. For 27 observations of the quality control check sample, the mean, standard deviation, and C.V. for total S are 0.57, 0.02, and 4.3%, respectively.

11. References


Bray P-1 Absorbed Phosphorus (6S)
Bausch and Lomb, Spectrophotometer 20 (6S3)

1. Application

The Bray P-1 procedure is widely used as an index of available P in the soil. The selectivity of the Bray extractant is designed to remove the easily acid-soluble P, largely calcium phosphates, and a portion of the phosphates of Al and Fe (Bray and Kurtz, 1945; Olsen and Sommers, 1982). In general, this method has been most successful on acid soils (Bray and Kurtz, 1945; Olsen and Sommers, 1982).

2. Summary of Method

A 1-g soil sample is shaken with 10 mL of extracting solution for 15 min at 100 oscillations per min\(^{-1}\). The solution is filtered. A 2-mL aliquot is transferred to a colorimetric tube to which 8-mL of ascorbic acid molybdate solution are added. The percent transmittance of the solution is read using a spectrophotometer. The Bray P-1 is reported in mg kg\(^{-1}\) (ppm) P.

3. Interferences

Many procedures may be used to determine P. Studies have shown that incomplete or excessive extraction of P to be the most significant contributor to inter-laboratory variation. The Bray P-1 procedure is sensitive to the soil/extractant ratio, shaking rate, and time. This extraction uses the ascorbic acid-potassium antimonyl-tartrate-molybdate method. The Fiske-Subbarrow method is less sensitive but has a wider range before dilution is required (North Central Regional Publication No. 221, 1988). For calcareous soils, the Olsen method is preferred. An alternative procedure for calcareous soils is to use the Bray P-1 extracting solution at a 1:50 soil:solution ratio. This procedure has been shown to be satisfactory for some calcareous soils (North Central Regional Publication No. 221, 1988; Smith et al., 1957).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the
use of concentrated H$_2$SO$_4$ and HCl to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Shaker, Eberbach 6000 power unit, reciprocating speed of 60 to 260 oscillations min$^{-1}$, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
5.3 Spectrophotometer 20, Bausch and Lomb
5.4 Pipettors, electronic digital, Rainin Instrument Co., Woburn, MA, 2500 µL and 10 mL
5.5 Cuvettes, glass, 10 mL, 1-cm light path
5.5 Funnel, 60° angle, long stem, 50-mm diameter
5.7 Filter paper, quantitative, Whatman grade 2, 9-cm diameter
5.8 Erlenmeyer flasks, 50 mL
5.9 Centrifuge, high-speed, International Equipment Co., IECB-22M

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Hydrochloric acid (HCl), concentrated, 12 N
6.3 HCl, 1 N. Carefully add 83.33 mL of concentrated HCl to DDI water and dilute to 1-L volume.
6.4 Sulfuric acid (H$_2$SO$_4$), concentrated, 36 N
6.5 Bray No. 1 Extracting solution, 0.025 N HCl and 0.03 N NH$_4$F. Dissolve 8.88 g of NH$_4$F in 4 L DDI H$_2$O. Add 200 mL of 1.0 N HCl and dilute to 8 L with DDI water. The solution pH should be 2.6 ±0.5. Store in a polyethylene bottle.
6.6 Stock standard P solution (SSPS), 100 ppm P. Add 0.2197 g of KH$_2$PO$_4$ in 25 mL of DDI water. Dilute to a final volume of 500 mL with extracting solution. Store in a refrigerator. Solution is stable to 1 yr.
6.7 Sulfuric-tartrate-molybdate solution (STMS). Dissolve 60 g of ammonium molybdate tetrahydrate [(NH$_4$)$_6$Mo$_7$O$_{24}$•4H$_2$O] in 200 mL of boiling DDI water. Allow to cool to room temperature. Dissolve 1.455 g of antimony potassium tartrate (potassium antimonyl tartrate hemihydrate K(SbO)C$_4$H$_7$O$_6$•½H$_2$O) in the ammonium molybdate solution. Slowly and carefully add 700 mL of concentrated H$_2$SO$_4$. Cool and dilute to 1 L with DDI water. Store in the dark in the refrigerator.
6.8 Ascorbic acid solution. Dissolve 33.0 g of ascorbic acid in DDI water and dilute to 250 mL with DDI water. Store in the dark in the refrigerator.
6.9 Working ascorbic acid molybdate solution (WAMS). Prepare fresh each day. Mix 25 mL of STMS solution with 800 mL of DDI water. Add 10 mL of ascorbic acid solution and dilute to 1 L with DDI water.

6.10 Standard P calibration solutions (SPCS), 0.2, 0.5, 1.0, 2.0, 3.0, 4.0, 5.0, 6.0, 7.0, and 8.0 ppm. Dilute the SSPS with the extracting solution as follows: 0.2 ppm = 0.5:250; 0.5 ppm = 0.5:100; 1.0 ppm = 1:100; 2.0 ppm = 2:100; 3.0 ppm = 3:100; 4.0 ppm = 4:100; 5.0 ppm = 5:100; 6.0 ppm = 6:100; 7.0 ppm = 7:100; 8.0 ppm = 8:100.

7. Procedure
7.1 Weigh 1.00 g of air-dry soil into a 50-mL Erlenmeyer flask.
7.2 Dispense 10.0 mL of extracting solution to flask.
7.3 Securely place the flask in the shaker. Shake for 15 min at 100 oscillations min$^{-1}$ at room temperature (20 °C).
7.4 Remove the sample from the shaker. Decant, filter, and collect extract.
7.5 Centrifuging or repeated filtering may be necessary to obtain clear extracts. Decant into 13-mL centrifuge tube and centrifuge at 10,000 RPM for 10 min.
7.6 Use the pipettor to transfer a 2-mL aliquot of the sample to a cuvette. Also transfer a 2-mL aliquot of each SPCS to a cuvette. Use a clean pipette tip for each sample and SPCS.
7.7 Dispense 8 mL of the WAMS to sample aliquot and to each SPCS (1:5 dilution).
7.8 The color reaction requires a minimum of 20 min before analyst records readings.
7.9 Set the spectrophotometer (red bulb) to read at 882 nm.
7.10 Set the 100% transmittance against the blank which has 8 mL of the WAMS solution and 2 mL of extracting solution.

8. Calculations
8.1 Transmittance of a solution is the fraction of incident radiation transmitted by the solution, i.e., $T = P/P_0$, and is often expressed as a percentage, i.e., $\%T = P/P_0 \times 100$. The absorbance of a solution is directly proportional to concentration and is defined by the equation, $A = -\log_{10} T$. These relationships are derived from Beer’s law.

Calibration Calculations

8.2 Use transmission of each SPCS to either construct a calibrated curve to plot P or use a least squares analysis to calculate P. The P is reported in ppm.
8.3 Calibration Curve: Plot the transmittance against the ppm P of each SPCS on semilog graph paper or convert to absorbances and plot on linear graph
paper. Construct the calibration curve by finding the “best” line that fits the plotted SPCS.

8.4 Linear Squares Analysis: Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. To facilitate data manipulation in a least squares analysis, the following standard curve is developed using the concentration of SPCS as a $f[\ln(\%T)]$. Final calculated analyte concentration with either $\log_{10}$ or $\ln$ base would be the same. Using the following example, calculate analyte concentration with $P$ (ppm) in extract = $Y$ variable and percent transmittance ($\%T$) = the $X$ variable. The $X$ variable is the natural logarithm of $T$.

<table>
<thead>
<tr>
<th>$P$ (ppm)</th>
<th>$T$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>77</td>
</tr>
<tr>
<td>4</td>
<td>59</td>
</tr>
<tr>
<td>6</td>
<td>45</td>
</tr>
<tr>
<td>8</td>
<td>33</td>
</tr>
<tr>
<td>0</td>
<td>24</td>
</tr>
</tbody>
</table>

Number of standards = $n=6$

$$\sum Y_i = 30 \quad \quad \sum X_i = 23.5077$$

$$\frac{\sum Y_i}{n} = Y = 5 \quad \quad \frac{\sum X_i}{n} = X = 3.9180$$

$$\sum X_i Y_i = 107.5902 \quad \quad \sum X_i^2 = 93.5185$$

$$\sum X_i \sum Y_i = 705.231$$

$$b = \frac{\sum X_i Y_i - \sum X_i \sum Y_i / n}{\sum X_i^2 - (\sum X_i)^2 / n} = \frac{107.5902 - 117.5385}{93.5185 - 92.102} = -7.023$$

$b =$ slope of the line, i.e., the amount that $Y$ changes when $X$ changes by 1 unit.

The equation is as follows:

$$Y = Y + b (X - X)$$

$$Y = 5 - 7.023 (\ln(X) - 3.9180)$$
Analyte Calculation

8.5 *Calibration Curve:* Read the P (ppm) directly from the calibration curve.

8.6 *Least Squares Analysis:* Put the ln(%)T in the preceding equation and solve for ppm P. Thus, if sample extract has 84% transmission, the preceding equation is as follows:

\[ Y = 5 - 7.023 \ln(84) + 27.516 = 1.40 \text{ ppm} \]

8.7 Convert the extract P (ppm) to soil P (ppm or lbs/A) as follows:

\[ \text{Soil P (ppm)} = \text{Extract P (ppm)} \times 10 \]

\[ \text{Soil P (lbs/A)} = \text{Extract P (ppm)} \times 20 \]

9. Report

Report the soil Bray P-1 mg kg\(^{-1}\) (ppm) to the nearest whole number.

10. Precision

Precision data are not available for this procedure.

11. References


North Central Regional Publication No. 221. 1988. Recommended chemical soil test procedures for the North Central region. Agric. Exp. Stn. of IL, IN, IA, KS, MI, MN, MS, NE, ND, OH, SD, WI, and USDA cooperating.


New Zealand P Retention (6S)

UV-Visible Spectrophotometer (6S4)

Beckmann DU-7 (6S4b)

1. Application

In *Soil Taxonomy*, the P retention of soil material is a criterion for andic soil properties (Soil Survey Staff, 1990). Andisols and other soils that contain large amounts of allophane and other amorphous minerals have capacities for binding
P (Gebhardt and Coleman, 1984). The factors that affect soil P retention are not well understood. However, allophane and imogolite have been considered as major materials that contribute to P retention in Andisols (Wada, 1985). Phosphate retention is also called P absorption, sorption, or fixation.

2. Summary of Method

A 5-g soil sample is shaken in a 1000-ppm P solution for 24 h. The mixture is centrifuged at 2000 rpm for 15 min. An aliquot of the supernatant is transferred to a colorimetric tube to which nitric vanadomolybdate acid reagent (NVAR) is added. The percent transmittance of the solution is read using a spectrophotometer. The New Zealand P retention is reported as percent P retained.

3. Interferences

No significant problems are known to affect the P retention measurement.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated HNO₃ to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Shaker, Eberbach 6000 power unit, reciprocating speed of 60 to 260 epm, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
5.3 Digital diluter/dispenser, Microlab 500, Hamilton Co., P.O. Box 10030, Reno, NV
5.4 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.5 Diluter/dispenser, 25 mL
5.6 UV-Visible Spectrophotometer, DU-7, Beckmann Instruments Inc.
5.7 Cuvettes, Labcraft Brand, disposable, polystyrene, square-bottom, 4.5 mL, 12.5 mm x 12.5 mm x 46 mm, Curtin Matheson Scientific, Inc., Houston, TX
5.9 Trunions, International no. 320, International Equip. Co., Boston, MA
5.10 Centrifuge tubes, 50 mL, Oak-Ridge, polyallomer, Nalgene 3119, Nalge Co., Box 20365, Rochester, NY
5.11 Plastic cups, 2 fl. oz.
5.12 Pipets, volumetric, class A, glass, various sizes of 1 to 20 mL

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Nitric acid (HNO₃), concentrated, 16 N
6.3 P retention solution, 1000 ppm P. Dissolve 35.2 g of KH₂PO₄ and 217.6 g of sodium acetate (Na₂C₂H₃O₂•3H₂O) in DDI water. Add 92 mL of glacial acetic acid. Dilute to 8 L with DDI water in a volumetric flask. The solution pH should range between 4.55 and 4.65.
6.4 Molybdate solution. Dissolve 16 g of ammonium molybdate (VI) [(NH₄)₆Mo₇O₂₄•4H₂O] in 50 °C DDI water. Allow the solution to cool to room temperature and dilute to 1 L with DDI water.
6.5 Nitric acid solution. Carefully and slowly dilute 100 mL of concentrated HNO₃ to 1 L of DDI water. Add the acid to the water.
6.6 Nitric vanadomolybdate acid reagent (NVAR), vanadate solution. Dissolve 0.8 g of NH₄VO₃ in 500 mL of boiling DDI water. Allow the solution to cool to room temperature. Carefully and slowly add 6 mL of concentrated HNO₃. Dilute to 1 L with DDI water. Mix the nitric acid solution with the vanadate solution and then add the molybdate solution. Mix well.
6.7 Stock P standard solution (SPSS), 4000 ppm P. Dissolve 17.6 g KH₂PO₄ in DDI water. Dilute to 1 L with DDI water.
6.8 Standard P calibration P solutions (SPCS), 100, 80, 60, 40, 20, and 0% P retained. Dilute the SPSS with a solution that contains 32.8 g of sodium acetate (CH₃COONa) and 23 mL of glacial acetic acid diluted to 2 L with DDI water as follows: 100% = DDI water (0 ppm); 80% = 1:20 (200 ppm); 60% = 1:10 (400 ppm); 40% = 3:20 (600 ppm); 20% = 1:5 (800 ppm); and 0% = 1:4 (1000 ppm). The percent amount refers to percent P retention.

7. Procedure
7.1 Weigh 5.00 g of air-dry soil into a 50-mL centrifuge tube.
7.2 Use the dispenser to add 25.0 mL of P-retention solution to centrifuge tube.
7.3 Cap centrifuge tube and place in shaker and shake for 24 h at room temperature (20 °C).
7.4 Add 2 to 3 drops of Superfloc, 0.02% w/v to each tube.
7.5 Centrifuge sample at 2000 rpm for 15 min. Filter using a Milipore filter, if necessary.
7.6 Pour sample supernatant into plastic cup.

7.7 Use the digital diluter to add the nitric vanadomolybdate acid reagent (NVAR) to each sample supernatant and to each SPCS. To fill a 4.5-mL cuvette, use a dilution of 1:20 sample dilution.

7.8 The color reaction requires a minimum of 30 min before the analyst records readings.

7.9 Set the spectrophotometer to read at 466 nm. Autozero using the DDI water (blank). A blank has all reagents contained in the sample extract except the soil.

7.10 Record the percent transmittance to the nearest 0.01 unit for the sample extract and each SPCS.

8. Calculations

8.1 Transmittance of a solution is the fraction of incident radiation transmitted by the solution, i.e., \( T = \frac{P}{P_0} \), and is often expressed as a percentage, i.e., \( T = \frac{P}{P_0} \times 100 \). The absorbance of a solution is directly proportional to concentration and is defined by the equation, \( A = -\log_{10} T \). These relationships are derived from Beer's law.

**Calibration Calculations**

8.2 Use the transmittance of each SPCS to either construct a calibrated curve to plot \( P \) or use a least squares analysis to calculate \( P \). The \( P \) is reported in percent retained.

8.3 Calibration Curve: Plot the transmittances against the ppm \( P \) of each SPCS on semilog graph paper or convert to absorbances and plot on linear graph paper. Construct the calibration curve by finding the “best” line that fits the plotted SPCS.

8.4 Least Squares Analysis: Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. To facilitate data manipulation in a least squares analysis, the following standard curve is developed using the concentration of SPCS as a \( f[\ln(\%T)] \). Final calculated analyte concentration with either \( \log_{10} \) or \( \ln \) base would be the same. Refer to method 6S3b for an example of least squares analysis.

**Analyte Calculation**

8.5 Calibration Curve: Read the percent \( P \) directly from the calibration curve.

8.6 Least Squares Analysis: Refer to method 6S3 for an example of least squares analysis.
9. Report
Report the percent New Zealand P retention to the nearest whole number.

10. Precision
Precision data are not available for this procedure.

11. References

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**Citric Acid Extractable Phosphorus (6S)**
**Beckmann DU-7, UV-Visible Spectrophotometer (6S5)**

1. Application
In *Soil Taxonomy*, citric acid soluble P$_2$O$_5$ is a criterion for distinguishing between mollic (<250 ppm P$_2$O$_5$) and anthropic epipedons (>250 ppm P$_2$O$_5$) (Soil Survey Staff, 1975). Additional data on anthropic epipedons from several parts of the world may permit improvements in this definition (Soil Survey Staff, 1994). The method 6S5 is used by N.A.A.S. (England and Wales) and is based on the method developed by Dyer (1894).

2. Summary of Method
A sample is checked for CaCO$_3$ equivalent. Sufficient citric acid is added to sample to neutralize the CaCO$_3$ plus bring the solution concentration of citric acid to 1%. A 1:10 soil:solution is maintained for all samples. The sample is shaken for 16 h and filtered. Ammonium molybdate and stannous chloride are added. The percent transmittance of the solution is read using a spectrophotometer. The 1% citric acid extractable P$_2$O$_5$ is reported in mg kg$^{-1}$ (ppm).

3. Interferences
Unreacted carbonates interfere with the extraction of P$_2$O$_5$. Sufficient citric acid is added to sample to neutralize the CaCO$_3$. However, a high citrate level in sample may interfere with the molybdate blue test. If this occurs, the method can
be modified by evaporating the extract and ashing in a muffle furnace to destroy the citric acid.

Positive interferences in the analytical determination of $P_2O_5$ are silica and arsenic, if the sample is heated. Negative interferences in the $P_2O_5$ determination are arsenate, fluoride, thorium, bismuth, sulfide, thiosulfate, thiocyanate, or excess molybdate. A concentration of Fe $>$1000 ppm interferes with $P_2O_5$ determination. Refer to Snell and Snell (1949) and Metson (1956) for additional information on interferences in the citric acid extraction of $P_2O_5$.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in fume hood. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Follow standard laboratory procedures.

5. Equipment

5.1 Electronic balance, ±0.01-g sensitivity
5.2 Shaker, Eberbach 6000 power unit, reciprocating speed of 60 to 260 oscillations min$^{-1}$, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
5.3 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. Box 10030, Reno, NV
5.4 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV
5.5 Centrifuge tubes, 50 mL, Oak-Ridge, polyallomer, Nalgene 3119, Nalge Co., Box 20365, Rochester, NY
5.6 Filter paper, quantitative, Whatman grade 2, 9-cm diameter
5.7 Funnel, 60° angle, long stem, 50-mm diameter
5.8 Erlenmeyer flasks, 50 ml
5.9 Bottles with gas release caps
5.10 Pipettors, electronic digital, Rainin Instrument Co., Woburn, MA, 2500 µL and 10 mL Digital Pipette, 10-ml
5.11 UV-visible spectrophotometer, DU-7, Beckmann Instruments, Inc.
5.12 Cuvettes, Labcraft Brand, disposable, polystyrene, square-bottom, 4.5 mL, 12.5 mm x 12.5 mm x 46 mm, Curtin Matheson Scientific, Inc., Houston, TX
6. Reagents

6.1 Distilled, deionized (DDI) water

6.2 Hydrochloric acid (HCl), concentrated, 12 N

6.3 Citric acid solution, 10%. Dissolve 100 g of anhydrous citric acid (C\textsubscript{6}H\textsubscript{8}O\textsubscript{7}) in 1-L volumetric flask.

6.4 Citric acid solution, 1%. Dilute 100.0 ml of 10% citric acid solution to 1-L with DDI water.

6.5 Ammonium molybdate solution, 1.5%. Dissolve 15.0 g of ammonium molybdate [(NH\textsubscript{4})\textsubscript{6}Mo\textsubscript{7}O\textsubscript{24}•4H\textsubscript{2}O] in 300 mL of distilled water. Transfer to a 1-L volumetric flask and carefully add 310 mL of concentrated HCl. Allow to cool. Make to 1-L volume with DDI water. Store in brown bottle in the dark. Solution is stable for ~3 months.

6.6 Stock stannous chloride solution (SSCS). Dissolve 10 g of stannous chloride (SnCl\textsubscript{2}•2H\textsubscript{2}O) in 100 mL of concentrated HCl.

6.7 Working stannous chloride solution (WSCS). Dilute 2 mL of SSCS with 100 mL of DDI water. Use immediately as solution is only stable for ~4 h.

6.8 Stock standard P\textsubscript{2}O\textsubscript{5} solution (SSPS), 250 ppm P. Dissolve 1.099 g of potassium dihydrogen orthophosphate (KH\textsubscript{2}PO\textsubscript{4}) with DDI water in 1-L volumetric flask. Add 5 ml of 2 N HCL. Make to 1-L volume with DDI water.

6.9 Working stock standard P\textsubscript{2}O\textsubscript{5} solution (WSSPS), 2.5 ppm P. Pipette 10.0 mL of SSPS and dilute to 1-L in a volumetric flask with DDI water.

6.10 Standard P\textsubscript{2}O\textsubscript{5} calibration solutions (SPCS). Pipette 0, 1, 2, 3, 4, and 5 mL of WSSPS into 50-mL oakridge tubes. Add 1 ml of 1% citric acid solution. Continue color development as for samples. Distilled water may be used as a blank.

7. Procedure

7.1 Weigh 3.00 g of <2-mm, air-dry soil into a bottle gas release tops. If the soil does not contain free carbonates, proceed to step 7.3.

7.2 If the soil contains free CaCO\textsubscript{3}, refer to Table 1 to determine the amount of 10% citric acid solution required to neutralize the CaCO\textsubscript{3}. Add required mLs of 10% citric acid into a graduated cylinder and bring to a volume of 30-ml with DDI water. Add this solution to the soil. Swirl the bottle over a period of 6 h at 100 oscillations min\textsuperscript{-1} dissolve and neutralize the CaCO\textsubscript{3}. Proceed to step 7.4.
Table 1.—Volume of 10% Citric Acid (mL) Required to Decompose CaCO₃ (%) and to Bring Solution Concentration to 1% in a Final Volume of 30 mL for 3-g Sample.

<table>
<thead>
<tr>
<th>%CC</th>
<th>mL CA²</th>
<th>% CC</th>
<th>mL CA</th>
<th>% CC</th>
<th>mL CA</th>
<th>%CC</th>
<th>mL CA</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3.0</td>
<td>16</td>
<td>9.7</td>
<td>32</td>
<td>16.4</td>
<td>48</td>
<td>23.2</td>
</tr>
<tr>
<td>1</td>
<td>3.4</td>
<td>17</td>
<td>10.2</td>
<td>33</td>
<td>49</td>
<td>23.6</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>3.8</td>
<td>18</td>
<td>10.6</td>
<td>34</td>
<td>50</td>
<td>24.0</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>4.3</td>
<td>19</td>
<td>11.0</td>
<td>35</td>
<td>17.7</td>
<td>51</td>
<td>24.4</td>
</tr>
<tr>
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<td>4.7</td>
<td>20</td>
<td>11.4</td>
<td>36</td>
<td>18.1</td>
<td>52</td>
<td>24.8</td>
</tr>
<tr>
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<td>5.1</td>
<td>21</td>
<td>11.8</td>
<td>37</td>
<td>18.6</td>
<td>53</td>
<td>25.3</td>
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<tr>
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<td>5.5</td>
<td>22</td>
<td>12.2</td>
<td>38</td>
<td>19.0</td>
<td>54</td>
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<td>12.7</td>
<td>39</td>
<td>19.4</td>
<td>55</td>
<td>26.1</td>
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<tr>
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<td>40</td>
<td>19.8</td>
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<tr>
<td>9</td>
<td>6.8</td>
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<td>13.5</td>
<td>41</td>
<td>20.2</td>
<td>57</td>
<td>27.0</td>
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<td>42</td>
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<td>58</td>
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<td>16.0</td>
<td>47</td>
<td>22.8</td>
<td>63</td>
<td>29.5</td>
</tr>
</tbody>
</table>

¹ %CC = percent calcium carbonate in a sample
² CA = ml of 10% citric acid needed to be diluted to 30-mL volume with RODI water and added to sample

7.3 If the soil contains no free CaCO₃, add 30 mL of 1% citric acid solution to the sample.
7.4 Cap the bottles, place in a shaker, and shake for 16 h at 100 oscillations min⁻¹.
7.5 Remove the sample from shaker and filter.
7.6 Pipette 1 mL of sample extract into a 50-mL oakridge tube. Add 4 mL of ammonium molybdate solution to all samples and standards. Bring up to 25 mL mark with DDI water. Add 2 mL stannous chloride. Shake to mix and allow to stand 20 min for color development.
7.7 Set the spectrophotometer to read at 660 nm. Set the zero against distilled water (blank). A blank has all reagents contained in the sample extract except the soil.

7.8 Record the percent transmittance to the nearest 0.01 unit for the sample extract and each SPCS.

8. Calculations

8.1 Transmittance of a solution is the fraction of incident radiation transmitted by the solution, i.e., \( T = \frac{P}{P_0} \), and is often expressed as a percentage, i.e., \( \%T = \frac{P}{P_0} \times 100 \). The absorbance of a solution is directly proportional to concentration and is defined by the equation, \( A = -\log_{10} T \). These relationships are derived from Beer’s law.

**Calibration Calculations**

8.2 Use transmission of each SPCS to either construct a calibrated curve to plot \( P_2O_5 \) or use a least squares analysis to calculate \( P_2O_5 \). The \( P_2O_5 \) is reported in ppm.

8.3 *Calibration Curve*: Plot the transmittances against the ppm \( P_2O_5 \) of each SPCS on semilog graph paper or convert to absorbances and plot on linear graph paper. Construct the calibration curve by finding the “best” line that fits the plotted SPCS.

8.4 *Linear Squares Analysis*: Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. To facilitate data manipulation in a least squares analysis, the following SPCS curve is developed using the concentration of SPCS as a f[ln(\%T)]. Final calculated analyte concentration with either \( \log_{10} \) or ln base would be the same. Refer to method 6S3 for an example of least squares analysis.

**Analyte Calculation**

8.5 *Calibration Curve*: Read the \( P_2O_5 \) (ppm) directly from the calibration curve.

8.6 *Least Squares Analysis*: Refer to method 6S3 for an example of least squares analysis.

8.7 Convert the extract \( P_2O_5 \) (ppm) to soil \( P_2O_5 \) (ppm or lbs/A) as follows:

\[
\text{Soil } P_2O_5 = \text{Extract } P_2O_5 \times DR \times 100 \times \text{AD}/\text{OD} / \text{Sample Weight (g)}
\]

where:

- Soil \( P_2O = P_2O_5 \) in soil (ppm)
- Extract \( P_2O_5 = P_2O_5 \) in extract (ppm)
- \( DR = \text{Dilution ratio, if necessary, otherwise 1} \)
100 = Conversion factor
AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report
Report the 1% citrate acid extractable P\textsubscript{2}O\textsubscript{5} in mg kg\textsuperscript{-1} (ppm) to nearest whole number.

10. Precision
Precision data are not available for this procedure.

11. References
Soil Survey Staff. 1975. Soil taxonomy: A basic system of soil classification
for making and interpreting soil surveys. USDA–SCS Agric. Handb. 436.
Washington, DC.
Print. Office, Washington DC.

MINERALOGY (7)

Instrumental Analyses (7A)
X-Ray Diffraction (7A2)
Phillips XRG-300 X-Ray Diffractometer
  Thin Film on Glass, Resin Pretreatment II (7A2i)
  (Mg Room Temp, Mg Glycerol Solvated, K 300°C, K 500°C)

1. Application
Clay fractions of soils are commonly composed of mixtures of one or more
phyllosilicate minerals together with primary minerals inherited directly from the
parent material (Whittig and Allardice, 1986). Positive identification of mineral
species and quantitative estimation of their proportions in these polycomponent
systems usually require the application of several complementary qualitative
and quantitative analyses (Whittig and Allardice, 1986). One of the most useful
methods to identify and to make semiquantitative estimates of the crystalline
mineral components of soil is X-ray diffraction analysis.
The operational strategy at the SSL and the preceding Lincoln Soil Survey Laboratory has been to adjust instrumental parameters to keep peak intensity of a soil reference constant from 1964 to present through the evolution of instrumentation. The intent is to keep the same quantitative interpretations consistent from sample to sample.

2. Summary of Method

Soils are dispersed and separated into fractions of interest. Sands and silts are mounted on glass slides as slurries or on double sticky tape for analysis. Clay suspensions are placed on glass slides to dry and to preferentially orient the clay minerals. The soil clay minerals of greatest interest are phyllosilicates, e.g., kaolinite, mica (illite), smectite, vermiculite, hydroxy-interlayered vermiculite, and chlorite.

Generally, no two minerals have exactly the same interatomic distances in three dimensions and the angle at which diffraction occurs is distinctive for a particular mineral (Whittig and Allardice, 1986). These interatomic distances within a mineral crystal result in a unique array of diffraction maxima, which help to identify that mineral. When several minerals are present in a sample, species identification is usually accomplished most easily and positively by determining the interatomic spacings that give rise to the various maxima and by comparing these with known spacings of minerals (Whittig and Allardice, 1986).

X-ray diffraction produces peaks on a chart that correspond to $2\theta$ angles on a goniometer. The angle of incidence of the goniometer is relative to the surface plane of the sample. Standard tables to convert $\theta$ or $2\theta$ angles to crystal "d" spacings are published in the U.S. Geological Survey Circular 29 (Switzer et al., 1948) and in other publications (Brown, 1980). At the SSL, conversions are made by the analysis program on the Philips diffractometer, d-spacings are recorded on an IBM-compatible 486 DOS-based computer system, and hard copies are printed for interpretation and filing. The crystal "d" spacings of minerals, i.e., the interval between repeating planes of atoms, can be calculated by Bragg's Law as follows:

$$n\lambda = 2d \sin \theta$$

where:
- $n =$ order of diffraction (integer)
- $\lambda =$ x-radiation wavelength (Angstroms, Å)
- $d =$ crystal “d” spacing (Å)
- $\theta =$ angle of incidence

When $n = 1$, diffraction is of the first order. The wavelength of radiation from an X-ray tube is constant and characteristic for the target metal in the tube. Copper radiation (CuK\textsubscript{α}) with a wavelength of 1.54 Å (0.154 nm) is used at the SSL. Because of similar structures of layer silicates commonly present in soil clays,
several treatments which characteristically affect the “d” spacings are necessary to identify components. At the SSL, four treatments are used, i.e., Mg\(^{2+}\) (room temperature); Mg\(^{2+}\)-glycerol (room temperature); K\(^{+}\) (300 °C); and K\(^{+}\) (500 °C).

3. Interferences

Intimate mixtures of similar phyllosilicate minerals on a fine scale cause problems in identification. The mixtures, differences in crystal size and purity, and background or matrix interferences affect quantification. No pretreatments other than dispersion with sodium hexametaphosphate are used for separation and isolation of the crystalline clay fraction. Impurities such as organic matter and iron oxides may act as matrix interferences causing peak attenuation during X-ray analysis or may interfere with clay dispersion and separation. The separation procedure to isolate the clay fraction from the other size fractions of the soil skews the <2-µm clay suspension toward the fine clay, but it minimizes the inclusion of fine silt in the fraction. Dried clay may peel from the XRD slide. One remedy is to rewet the peeled clay on the slide with 1 drop of glue-water mixture (1:7). Other remedies are as follows:

   a. Place double sticky tape on the slide prior to adding the dried clay.
   b. Dilute the suspension by half, if thick.
   c. Crush with ethanol and dry, and then add water to make a slurry slide.
   d. Roughen the slide surface with a fine grit sand paper.

Sufficient glycerol on the slides is required to solvate the clay, i.e., to expand smectites to 18 Å. X-ray analysis should be performed 1 to 2 days after glycerol addition. If excess glycerol is applied to the slide and free glycerol remains on the surface, XRD peaks are attenuated. Some suggestions to dry the slides and achieve optimum glycerol solvation are as follows:

   a. Use a desiccator to dry slide, usually when the clay is thin.
   b. If the center of slide is whitish and dry, usually with thick clay, brush slide with glycerol or add an additional drop of glycerol.

4. Safety

Operate the centrifuge with caution. Keep the centrifuge lid closed when in operation. Ensure that all hangers and tubes are seated firmly in proper location. Use tongs and appropriate thermal protection when operating the muffle furnace. The diffraction unit presents an electrical and radiation hazard. Analysts must receive radiation safety training before operating the equipment. Employees must wear a radiation film badge while in the room when the diffraction unit is in operation.

5. Equipment

5.1 Teaspoon (5 g)
5.2 Dispenser, 5 mL, for sodium hexametaphosphate solution
5.3 Centrifuge, International No. 2, with No. 240 head and carriers for centrifuge tubes, International Equip. Co., Boston, MA
5.4 Centrifuge tubes, plastic, 100 mL, on which 10-cm solution depth is marked
5.5 Rubber stoppers, No. 6, for centrifuge tubes
5.6 Mechanical shaker, reciprocal, 120 oscillations min\(^{-1}\)
5.7 Plastic cups, 60 mL (2 fl. oz.) with lids
5.8 Label machine
5.9 Hypodermic syringes, plastic, 12 mL, with tip caps
5.10 Screen, 80 mesh, copper
5.11 Dropper bottle, plastic, 30 mL (1 fl. oz.), for a 1:7 glycerol:water mixture
5.12 Muffle furnace
5.13 X-ray diffractometer, Philips XRG-300, with PW-1170 automated sample changer
5.14 PC-APD, Philips, software for Automatic Powder Diffraction (PW-1877), Version 3.5
5.15 Computer, IBM-compatible 486, Gateway 2000 4D X2-66V
5.16 Printer, Hewlett Packard LaserJet IV
5.17 Plotter, Hewlett Packard 7550 Plus
5.18 XRD slides, glass, 14 x 19 mm
5.19 XRD sample preparation board, wood, with 32 places for glass XRD slides
5.20 Slide holder. Accepts 14 x 19 mm XRD glass slides. Modified so slide surfaces rest flush with surface of holder.
5.21 Magazine for slide holder, 35 positions
5.22 Reference slides: quartz and clay from reference soil

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Sodium hexametaphosphate solution. Dissolve 35.7 g of sodium hexametaphosphate (NaPO\(_3\)\(_6\)) and 7.94 g of sodium carbonate (Na\(_2\)CO\(_3\)) in 1 L DDI water.
6.3 Potassium chloride (KCl), 1.0 \(N\). Dissolve 74.60 g KCl in 1 L DDI water or 671.40 g KCl in 9 L DDI water.
6.4 Magnesium chloride (MgCl\(_2\)), 1.0 \(N\). Dissolve 47.61 g MgCl\(_2\) in 1 L DDI water or 428.49 g MgCl\(_2\) in 9 L DDI water.
6.5 Glycerol:water mixture (1:7). Add 4 mL of glycerol to 28 mL DDI water plus 2 drops of toluene.
6.6 Exchange resin, Rexyn 101 (H), analytical grade. Pretreatment of resin as follows:
6.6.1 Divide equally Rexyn 101 (H), approximately 250-g portions, into two 600-mL beakers labeled K and Mg and add appropriate salt solution (1.0 N KCl or 1.0 N MgCl$_2$). Cover resin with salt solution.

6.6.2 Stir, let settle for 10 min, decant clear solution, and add salt solution. Repeat 3 times. Leave resin covered in salt solution for 8 to 12 h.

6.6.3 Repeat step 6.6.2 second day. Resin is ready for syringes. Saturated resin not used initially for syringes can be saved for future use.

6.7 White glue, diluted 1:7 with DDI water

7. Procedure

**Preparation (Recharge) of Resin-Loaded Syringes**

7.1 Place a small circle of 80-mesh screen in a 12-mL syringe and add 4 cm$^3$ of exchange resin from which salt solution has been drained. Our procedure requires each sample to have 2 Mg and 2 K slides prepared, so we produce our syringes in sets of two.

7.2 Saturate the resin in each of the four syringes with 4 mL of the appropriate 1.0 N salt solution (MgCl$_2$ or KCl) and expel. Repeat saturation of resin.

7.3 Fill syringe completely with the salt solution and allow to equilibrate for 4 to 20 h.

7.4 Rinse syringe twice with 4 mL of DDI water and rinse tip cap.

7.5 Completely fill syringe with DDI water and allow to equilibrate for 4 to 20 h.

7.6 Rinse syringe twice with DDI water.

7.7 Expel water, cap syringe, and store.

**Preparation of Clay Suspension**

7.8 Place $\approx$5 g (1 tsp) of air-dry $<$2-mm soil in a 100-mL plastic centrifuge tube. If the sample appears to be primarily sand, use 10 g (2 tsp) of $<$2-mm soil to obtain sufficient clay.

7.9 Add 5 mL of sodium hexametaphosphate solution. If the soil contains gypsum or is primarily calcium carbonate, use 10 mL of sodium hexametaphosphate dispersing agent.

7.10 Fill tube to 9.5-cm height with DDI water.

7.11 Place rubber stopper in tube and shake overnight in mechanical shaker.

7.12 Remove stopper from tube and rinse stopper and sides of tube with enough water to bring the volume to the 10-cm mark.
7.13 Balance the pairs of tubes and place in centrifuge. Centrifuge at 750 rpm for 3.0 min.

7.14 If the clay is dispersed, carefully decant 30 mL of suspension into a labeled, 60-mL, plastic cup. Place cap on cup.

7.15 If the clay did not disperse after being shaken overnight, remove the rubber stopper and carefully decant the clear supernatant liquid.

7.16 Add an additional 10 mL of sodium hexametaphosphate dispersing agent to sample and then add DDI water to 9.5-cm depth.

7.17 Stopper and shake overnight to disperse the clay. Rinse stopper and fill tube to 10-cm mark.

7.18 Centrifuge, decant, and store clay suspension.

7.19 Use the clay suspension for X-ray diffraction analysis and HF plus aqua regia dissolution analysis. Dry clay suspension for use in thermal analysis.

**Thin Film on Glass, Resin Pretreatment**

7.20 The SSL uses a sample board which holds 32 slides, i.e., 8 samples x 4 treatments. Prepare the sample board with glass XRD slides to receive the following 4 treatments per clay suspension sample.

- Mg$^{2+}$ — room temperature
- Mg$^{2+}$ — glycerol (room temperature)
- K$^+$ — 300 °C (heated for 2 h)
- K$^+$ — 500 °C (heated for 2 h)

7.21 Place one small drop of the glycerol:water mixture (1:7) on each Mg$^{2+}$-glycerol slide.

7.22 Draw 1 mL of <2-µm clay suspension into the resin-loaded syringe and invert back and forth to facilitate cation exchange.

7.23 Dispense 3 drops to clear the tip.

7.24 Dispense ≈0.1 mL (6 to 10 drops) to cover the appropriate XRD slide. Draw DDI water into the syringe and expel 3 times to remove all of the clay suspension. Recharge the syringe after 10 times of use.

7.25 When the clay suspension has dried, transfer the slides with the K$^+$-saturated clays to transite plates and heat for a minimum of 2 h in a muffle furnace.

7.26 Heat the following sample slides on the XRD sample board.

- K$^+$-300 °C — slides 3, 7, 11, 15, 19, 23, 27, and 31
- K$^+$-500 °C — slides 4, 8, 12, 16, 20, 24, 28, and 32

7.27 After heating, remove the transite plate from the furnace, cool to air temperature, and return slides to XRD sample board.
**X-Ray Diffraction Operation**

7.28 The X-ray analysis of the glycerol slide must be done within 1 to 2 days after the slide dries. If this is not possible, skip Step 7.21 when slide is prepared. Add one small drop of glycerol:water mixture (1:7) to dry slide 24 h prior to X-ray analysis.

7.29 Transfer the slides (1 to 32) from XRD sample board to slide holders (1 to 32) and place in slots (1 to 32) in a magazine for the automated sample changer.

7.30 Analyze one reference soil sample in each run. Place this sample in slot 33.

7.31 Analyze one quartz standard for 2θ and intensity calibrations in each run. Place this sample in slot 34. Intensity is measured at peak maximum at or near 26.66° 2θ for 10 s.

7.32 The 32 samples from one XRD board constitute one run on the diffraction unit. Prepare a run sheet for samples on each XRD sample board. Refer to example run instruction (7.33). Refer to the manufacturer's manual for operation of the X-ray diffractometer.

7.33 Place the magazine in the automated sample changer. Confirm that the XRD shutter is off when changing magazines. Set the XRD unit parameters as follows:

- CuKα radiation, λ: 1.54 Å (0.154 nm)
- Scan range: 2° to 34° 2θ
- Generator settings: 40 kv, 20 ma
- Divergence slit: 1°
- Receiving slit: 0.2 mm
- Monochrometer: Yes

Step size and scan speed vary depending on intensity of X-rays generated from tube. Adjust settings to maintain same long-term peak intensities on standard reference clay and quartz standard regardless of tube intensities.

7.34 Enter run instruction from the keyboard. Create a batch file for the automated run. File names specified are of the sample number. An example run instruction is as follows:

**Batch File Name:** Project number (e.g., CP95LA022)

**Raw Data File Name:** Run number

**First Sample:** 1

**Last Sample:** 33

(reference soil clay)
7.35 Activate program. The run stores raw data on the hard disk under the subdirectory designated by project type and year, e.g., CP95. Refer to example run instruction (7.34).

7.36 Print a hard copy of the “Detected Peaks File” for each sample and perform level 1 smoothing on diffraction patterns.

7.37 Prepare and print a 4-color graphics chart. The 4 colors are blue (Mg\(^{2+}\)); green (Mg\(^{2+}\)-glycerol); pink (K\(^+\) 300 °C); and red (K\(^+\) 500 °C). Stamp chart with label; enter run parameter information, and complete soil information, e.g., soil name, horizon designation, and depth. File hard copies of detected peaks and graphics chart in pasteboard binders by state, county, and chronology.

7.38 Record “d” spacing and intensity of quartz standard in the logbook. Record the peak intensities for designated peaks for the reference soil clay.

7.39 File the detected peaks printout and graph for the reference soil in the reference soil-clay folder.

Interpretation of X-Ray Diffraction Data

7.40 The angle in degrees two theta (2θ) measured in X-ray diffraction analyses is converted to angstroms (Å) using tables complied according to Bragg’s Law. Refer to summary of method. Angstroms convert to nanometers (nm) by a factor of 0.1, e.g., 14 Å = 1.4 nm.

7.41 Use the following X-ray diffraction criteria to identify some common crystalline minerals. The reported “d” values are for 00/ basal spacings. The Miller index (hkl) specifies a crystal face which has some orientation to the three crystallographic axes of a, b, and c. The Miller index (00l) indicates a crystal face that is parallel to the a and b axes, e.g., phyllosilicate minerals. The following X-ray diffraction criteria also has some questions (Q) that may aid the analyst in interpreting the diffraction patterns. These questions are a suggested procedural approach to help the analyst identify the relative locations of a few peaks and to confirm key criteria.

X-Ray Diffraction Criteria

1. Kaolinite and Halloysite
   a. Crystal structure missing at 500 °C.
   b. 7 Å (7.2 to 7.5 Å) with all other treatments.
   Q. Is there a 7 Å peak? Is it destroyed at 500 °C? Kaolinite or Halloysite.
   Q. Is the peak sharp and at ~7.1 Å? Kaolinite.
   Q. Is the peak broad and at 7.2 to 7.5 Å? Halloysite.
2. Mica (Illite)
   a. 10 Å with all treatments.
   b. 10 Å with Mg²⁺-saturation.
   Q. Is there a 10 Å peak with Mg²⁺-saturation? Mica (Illite).

3. Chlorite
   a. Crystal structure of Fe-chlorites destroyed at 650 to 700 °C.
   b. 14 Å with all other treatments.
   c. 14 Å at 500 °C.
   d. Generally also has strong 7 Å peak.
   Q. Is there a 14 Å peak when heated to 500 °C? Chlorite.

4. Vermiculite
   a. 14 Å with Mg²⁺-saturation.
   b. 14 Å with Mg²⁺-glycerol solvation.
   c. Nearly 10 Å with K⁺ saturation.
   d. 10 Å when K⁺-saturated and heated to 300 °C.
   Q. Is there an enhanced 10 Å peak with K⁺-saturation in comparison to Mg²⁺-saturation that cannot be attributed to smectite? Vermiculite.

5. Smectite
   a. 14 Å with Mg²⁺-saturation
   b. 12 to 12.5 Å with K⁺- or Na⁺-saturation.
   c. 17 to 18 Å with Mg²⁺-glycerol solvation.
   d. 10 Å with K⁺-saturation and heating to 300 °C.
   Q. Is there a 17 to 18 Å peak upon solvation? Smectite.

6. Gibbsite
   a. Peak at 4.83 Å with Mg²⁺ and Mg²⁺-glycerol but destroyed when heated to 300 °C.

7. Goethite
   a. Peak at 4.18 Å with Mg²⁺ and Mg²⁺-glycerol but destroyed when heated to 300 °C.

8. Hydroxy-interlayed Vermiculite or Smectite
   a. Incomplete collapse to 10 Å of smectite or vermiculite when K⁺-saturated and heated to 300 °C.

9. Quartz
   a. Peaks at 4.27 Å and 3.34 Å with all treatments (only 3.34 Å if small amounts).
10. Lepidocrocite
   a. Peak at 6.2 to 6.4 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol but destroyed when heated to 300 °C.

11. Potassium Feldspar
   a. Peak at 3.24 Å with all treatments.

12. Plagioclase Feldspar
   a. Twin peaks between 3.16 and 3.21 with all treatments.

13. Calcite
   a. Peak at 3.035 Å with all treatments.

14. Dolomite
   a. Peak at 2.88 to 2.89 Å with all treatments.

15. Gypsum
   a. Peak at 4.27 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol, but destroyed when heated to 300 °C.

16. Mixed Layer Vermiculite-Mica
   a. Peak at 11 to 13 Å with Mg$^{2+}$ that does not expand with Mg$^{2+}$-glycerol.
   b. Peak collapses to 10 Å with K$^+$-saturation and heating to 300 °C.

17. Mixed Layer Smectite-Mica
   a. Peak at 11 to 13 Å with Mg$^{2+}$ that expands to 14–16 Å with Mg-glycerol.
   b. Peak collapses to 10 Å with K$^+$-saturation and heating to 300 °C.

18. Mixed Layer Chlorite-Mica
   a. Peak at 14 Å with Mg$^{2+}$ and Mg$^{2+}$-glycerol.
   b. Peak collapses toward 10 Å with K$^+$-saturation and heating to 300 °C, and more completely with heating to 500 °C, but never to 10 Å.

19. Mixed Layer Chlorite-Smectite
   a. Peak at 11 to 13 Å with Mg$^{2+}$-saturation that expands to about 16 Å with Mg$^{2+}$-glycerol.
   b. Collapses to about 12 Å with K$^+$-saturation and heating to 300 °C and 500 °C.

7.42 Use the X-ray diffraction criteria, i.e., diagnostic basal 00l spacings (Å), in Table 1 for identification and ready reference of some common crystalline minerals as affected by differentiating sample treatments.
7.43 Preferential orientation of clay mineral samples enhances diffraction from the basal (00$l$) spacing and tends to minimize the number and intensity of peaks from diffraction by other $hkl$ planes. With preferential orientation, second, third, and fourth order peaks may be recorded in addition to the basal first order peaks. Groups of associated peaks that differ by order of diffraction are as follows:

**Smectite (Mg$^{2+}$-glycerol):**
- a. 17 to 18 Å.
- b. 8.5 to 9 Å (weak).

**Chlorite, vermiculite, and smectite:**
- a. 14, 7, 4.7, and 3.5 Å.
- b. 7, 4.7, and 3.5 Å weak for smectite.

**Mica:**
- a. 10, 5 (weak in biotites and moderate in muscovites), and 3.3 Å.

**Kaolinite:**
- a. 7 and 3.5 Å.

7.44 The differentiation of kaolinite and halloysite in a sample can be aided by the use of formamide (Churchman et al., 1984). The intercalation and expansion of halloysite to a d-spacing of ≈10.4 Å is relatively rapid (20 to 30 min), whereas kaolinite expansion requires ≈4 h upon treatment. The procedure is as follows:

a. Lightly spray formamide as an aerosol on the dried Mg$^{2+}$-saturated slide.

b. Wait 15 min but not more than 1 h and X-ray approximately 7.6 to 13.5° 2θ (d = 11.6 to 6.55 Å).

c. Halloysite will expand to ≈10.4 Å, whereas kaolinite will remain unchanged.

d. Heating the sample to 110 °C for 15 min will collapse the halloysite to ≈7 Å.

e. The total amount of kaolinite and halloysite can be determined by thermal analysis. The intensity ratio of the 10.4 to 7.2 Å peaks of the formamide-treated sample can be used to determine the relative percentage of halloysite and kaolinite.

8. Calculations

X-ray diffraction produces peaks on a chart that corresponds to 2θ angle on a goniometer. Standard tables to convert θ or 2θ to crystal “d” spacings are
Table 1.—X-ray diffraction parameters of common soil clay minerals.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Treatment</th>
<th>Na⁺</th>
<th>Mg²⁺</th>
<th>Mg²⁺ Gly</th>
<th>K⁺ 300 °C</th>
<th>K⁺ 500 °C</th>
<th>K⁺ 700 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kaolinite</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>LD¹</td>
<td>LD</td>
</tr>
<tr>
<td>Halloysite</td>
<td>7B²</td>
<td>7B</td>
<td>7B</td>
<td>7B</td>
<td>7B</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Mica (Illite)</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Chlorite</td>
<td>14⁺³</td>
<td>14⁺</td>
<td>14⁺</td>
<td>14⁺</td>
<td>14⁺</td>
<td>14⁺</td>
<td>T⁴⁻</td>
</tr>
<tr>
<td>Vermiculite</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Smectite</td>
<td>12.5</td>
<td>14</td>
<td>18</td>
<td>12.5</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Gibbsite</td>
<td>4.85</td>
<td>4.85</td>
<td>4.85</td>
<td>4.85</td>
<td>LD</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Goethite</td>
<td>4.18</td>
<td>4.18</td>
<td>4.18</td>
<td>4.18</td>
<td>LD</td>
<td>LD</td>
<td>LD</td>
</tr>
<tr>
<td>Interlayer</td>
<td>10-14</td>
<td>10-14</td>
<td>10-18</td>
<td>10-14</td>
<td>10-14</td>
<td>10-14</td>
<td>10-14</td>
</tr>
<tr>
<td>Quartz</td>
<td>3.14 and 4.27</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcite</td>
<td></td>
<td>3.035</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dolomite</td>
<td></td>
<td>2.88</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

¹ LD = Lattice destroyed
² B = Broad peak is common
³ * = Sometimes <14 Å
⁴ T = Temperature of decomposition varies with chemical composition, particle-size, and heating conditions.

9. Report

From the “Detected Peaks File” and graphics chart, identify the minerals present according to the registered “d” spacings. As a first approximation, use the following peak intensities, i.e., peak heights above background in counts s⁻¹, to assign each layer silicate mineral to one of the 5 semiquantitative classes.
Class Peak height above background (counts sec\(^{-1}\))

<table>
<thead>
<tr>
<th>Class</th>
<th>Peak height above background (counts sec(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 (Very Large)</td>
<td>&gt;1.88 \times 10^3</td>
</tr>
<tr>
<td>4 (Large)</td>
<td>1.12 to 1.88 \times 10^3</td>
</tr>
<tr>
<td>3 (Medium)</td>
<td>0.36 to 1.12 \times 10^3</td>
</tr>
<tr>
<td>2 (Small)</td>
<td>0.11 to 0.36 \times 10^3</td>
</tr>
<tr>
<td>1 (Very Small)</td>
<td>&lt;0.11 \times 10^3</td>
</tr>
</tbody>
</table>

Adjust class placement to reflect area under the curve if peak is broad relative to peak height or if thermal, elemental, clay activity data, or other evidence warrant class adjustment. If there are no peaks or no evidence of crystalline components, place the sample in NX class (noncrystalline).

**10. Precision**

Precision data are not available for this procedure. Method 7A2i (X-ray diffraction) is semiquantitative.

**11. References**


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**Total Analysis (7C)**

**HF Plus Aqua Regia (HF + HNO\(_3\) + HCl) Dissolution (7C4a)**

**1. Application**

Prior to the development of modern analytical techniques, e.g., X-ray diffraction and thermal analysis, identification of minerals was based on elemental analysis and optical properties (Washington, 1930; Bain and Smith, 1994). Chemical analysis is still essential to determine mineral structural formulas and to identify
and quantify specific mineral species through elemental allocation to minerals. Many clay mineral groups are subdivided based on composition.

Analysis of the entire fine-earth (<2-mm) fraction or specific particle-size separates provides information on parent material uniformity, pedon development, and mineral weathering within or between pedons. This interpretation is determined from differences between horizons or pedons in elemental concentrations, elemental ratios such as Si/Al, Si/Al+Fe, or Ti/Zr, or from differences in total elemental concentrations compared to concentrations determined by selective dissolution techniques.

The inherent fertility of a soil derived from its parent material can be examined by determination of the basic cations relative to the Si or Al content. Phosphorus fertility of a soil and potential water quality problems can be better understood by measurements of total P, especially when compared to other P measurements, such as water-soluble or Bray-extractable P.

Hydrofluoric acid (HF) is efficient in the digestion and dissolution of silicate minerals for elemental dissolution (Bernas, 1968; Sawhney and Stilwell, 1994). Aqua regia (HNO₃ and HCl) aids in digestion of soil components, especially the organic fraction. Method 7C4a is a digestion of 100 mg of dried clay suspension, the fine-earth (<2-mm) fraction, or other particle-size separate with HF and aqua regia. Closed digestion vessels (Parr Bombs) are heated in the oven at 110 °C for at least 6 hours. Elemental concentration of the digestate is determined by inductively coupled plasma-atomic emission spectrometry (ICP–AES).

2. Summary of Method

A clay suspension (method 7A2i) containing approximately 100 mg of clay material is pipeted into a Teflon digestion container and dried at 110 °C. An equal amount of suspension is pipeted into a tared aluminum-weighing dish and dried at 110 °C to obtain a dried sample weight. An oven-dry 100-mg soil sample (<80 mesh) or a specific particle-size separate may be substituted for the clay suspension. The P and Na content of the clay fraction is not measurable when the soil is dispersed in sodium hexametaphosphate (method 7A2i). Total P and Na are measurable on the fine-earth fraction or other particle-size separates not dispersed in Na- or P-containing reagents, and the analyses are included as a part of this procedure.

Following evaporation of the aqueous portion of the suspension, 0.75 mL HNO₃, 0.25 mL HCl, and 5 mL HF are added. The vessel is inserted into a stainless steel retainer vessel, heated, cooled, and 15 mL of 2.5 percent boric acid solution is added to neutralize the excess HF acid. The digestate is quantitatively transferred with boric acid solution, diluted to 100 mL, shaken, and allowed to stand overnight. Approximately 60 mL are saved for analysis. The concentration of Fe, Mn, Al, Ca, Mg, Na, K, P, Si, Zr, Cu, Zn, As, Ti, Se, Cd, and Pb are determined by ICP analysis in methods 6C7b, 6D6a, 6G11b, 6N5b, 6O5b, 6P3b, 6Q3b, 6S6a, 6V1b, 8K1a, 8L1a, 8M1a, 8N1a, 8O1a, 8P1a, 8Q1a, and 8R1a, respectively. Data are reported in method 7C4a.
3. Interferences

Insoluble fluorides of various metals may form. Formation of SiF₄ results in gaseous losses of Si, but additions of boric acid retards formation of this molecule as well as dissolves other metal fluorides.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated acids to the fume hood. Keep HF acid refrigerated and avoid skin contact with all acids. Wash hands thoroughly after handling reagents. Filling the Teflon cup of the acid digestion bomb to greater than 25 percent of the free volume or adding organic reagents or oxidizing agents to the cup may result in explosion of the digestion bomb.

5. Equipment

5.1 Pipette(s) capable of delivering 5, 0.75, and 0.25 mL
5.2 Volumetric flasks, Nalgene, 100 mL
5.3 Polypropylene bottles, 60 mL, with cap
5.4 Electronic balance, ±0.1 mg sensitivity
5.5 Acid digestion bombs: 25-mL Teflon containers with stainless steel retainer vessels
5.6 Oven, 110 °C
5.7 Desiccator with P₂O₅ drying agent
5.8 Disposable aluminum-weighing dishes

6. Reagents

6.1 Deionized distilled (DDI) water
6.2 Hydrofluoric acid (HF), 48%, low trace metal content
6.3 Concentrated hydrochloric acid (HCl), 12 N. Use instrumental grade reagents which contain low levels of impurities.
6.4 Concentrated nitric acid (HNO₃), 16 N. Use instrumental grade reagents which contain low levels of impurities.
6.5 Boric acid solution, 2.5 percent. Dissolve 25.0 g low trace metal, granular boric acid (H₃BO₃) in 1000 mL DDI water.

7. Procedure

**HF plus Aqua Regia Dissolution**

7.1 Prepare Na-saturated clay as in method 7A2i, Preparation of Clay Suspension, Steps 7.8 to 7.19. Clay dispersion by this method eliminates quantitative analysis of Na and P in the clay due to dispersion by sodium
hexametaphosphate. Digestion of the entire fine-earth (<2-mm) fraction or any fraction not derived by dispersion with sodium hexametaphosphate (or other Na- and P-containing dispersing agents) can be quantitatively analyzed for Na and P. Dispersion of clays and cleaning of test tubes and dishware should be with DDI water.

7.2 Pipette a known aliquot of clay suspension containing approximately 100 mg clay into a 25-mL Teflon container. The milliliters of suspension required depends on the clay concentration of the suspension but is generally from 2 to 6 mL. More dilute suspensions should be partially evaporated under a fume hood to concentrate the clay prior to transfer to the Teflon container. Fine-earth (<2-mm) or a specific particle-size separate ground to <80-mesh may be used instead of clay. Samples with greater than 3 percent organic C should be ashed in a muffle furnace at 400 °C for 2 h prior to analysis to destroy the organic matter. Oven-dry the sample (110 °C), cool over P₂O₅, and weigh to 100 ±0.1 mg. If a clay suspension is used, Steps 7.3 to 7.4 are performed. Proceed to Step 7.5 if using fine-earth or other oven-dried material.

7.3 Pipette a duplicate aliquot of suspension (as used in Step 7.2) into a tared Al weighing dish, dry at 110 °C, cool in a desiccator with P₂O₅, and weigh to the nearest 0.1 mg. Use this value as the sample weight in the calculations.

7.4 Dry the Teflon container and clay suspension in an oven for 4 h or until the aqueous portion of the suspension is completely evaporated. Remove from oven and cool on the bench top or in a fume hood. Cooling in a desiccator is not required.

7.5 Pipette 0.75 mL HNO₃ and 0.25 mL HCl into the sample and allow to completely wet and then pipette 5 mL HF into sample.

7.6 Place covered Teflon container in stainless steel retainer vessel. Place sample in oven at 110 °C for a minimum of 6 h. Samples can be left in the oven overnight at 110 °C.

7.7 Remove samples from oven and cool for at least 4 h.

7.8 Under a hood, remove Teflon container from steel retainer vessel, open the Teflon container, and add 15 mL 2.5 percent boric acid solution.

7.9 Quantitatively transfer contents of Teflon container to a 100 mL Nalgene volumetric flask and adjust to volume with 2.5 percent H₃BO₃.

7.10 Cap flask and mix well by inverting at least three times. Allow to stand overnight to dissolve any metal fluorides.

7.11 Invert the volumetric flask to mix and decant approximately 60 mL into a labeled polypropylene container.

7.12 Prepare working standards of a blank, a clay suspension from a SSL reference soil sample, and a National Institute of Standards and Technology
(NIST) standard reference material by the same digestion method. Run one of these standards with each set of 20 samples.

7.13 Solutions and standards are analyzed by ICP spectrometry. Refer to methods 6C7b, 6D6a, 6G11b, 6N5b, 6O5b, 6P3b, 6Q3b, 6S6a, 6V1b, 8K1a, 8L1a, 8M1a, 8N1a, 8O1a, 8P1a, 8Q1a, and 8R1a for analysis of Fe, Mn, Al, Ca, Mg, Na, K, P, Si, Zr, Cu, Zn, As, Ti, Se, Cd, and Pb, respectively.

8. Calculations

8.1 Data are transferred as an ASCII file from the ICP computer onto a 3.5-in floppy disk via “Report Writer” in the TJA software ThermoSpec, Version 5.06.

8.2 On a MS–DOS based PC computer, import the ASCII file of ICP data into the DOS editor and strip off unnecessary headers and data from standards. Save the file after editing, renaming using a format that can be imported into LOTUS, e.g., rename to .wk3 file for LOTUS 123, Version 3.1.

8.3 Import the file into an established total analysis spreadsheet in LOTUS 123. The spreadsheet has columns for sample number, soil fraction digested, soil weight, concentration of each element in ppm, and the calculated elemental percent. Each line of elemental data for a sample is imported as a single data string.

8.4 Parse the components of each data string into separate columns. Rearrange the data set in order to have all elemental values on a single line for a particular sample. Move the data into the correct columns of the spreadsheet.

8.5 Insert values for elements requiring dilution into the original line of sample data and replace all negative values with zero.

8.6 Input sample weights, or if possible, import sample weights (dried soil weights) from the ASCII file generated by computer attached to balance via RS-232.

8.7 Calculate the percent of an element in the soil from ppm in solution as shown in the Si example as follows:

\[
\text{Si (ppm) in solution} = 75.2 \text{ ppm (75.2 µg/mL)}
\]

Volume extract = 100 mL

Sample weight (110 °C) = 100.0 mg

Calculate as follows:

\[
\% \text{ Si} = \frac{75.2 \text{ µg mL}^{-1} \times 100 \text{ mL} \times (1 \text{ g/10}^6 \text{ µg}) \times (1/0.1 \text{ g soil}) \times 100}{100} = 7.52 \%
\]

8.8 The fraction digested needs to be identified with each sample. Use proper SSL database abbreviations.
8.9 Delete the Na and P data for clay samples dispersed in sodium hexametaphosphate.

8.10 Prepare the file to send to CMS. Save the file as an unformatted ASCII file using LOTUS.

8.11 Enter data for Si, Al, Fe, Mg, Mn, K, Ti, Ca, Zr, P, and Na into the SSL CMS database on a 110 °C weight basis as percent of the element in the fraction digested. Data are converted to the oxide form on the data sheet.

8.12 The factor for converting from an elemental form to an oxide form is based on the atomic weights of the element and oxygen. An example is as follows:

Atomic weight Si = 28.09

Atomic weight O = 16.0

Molecular weight SiO₂ = 60.09

Calculate percent Si in SiO₂ as follows:

\[
\text{Si (\%)} = \left( \frac{28.09}{60.09} \right) \times 100 = 46.7\%
\]

There is 46.7 percent Si in SiO₂. To convert from percent Si to percent Si oxide (SiO₂) in the soil, divide the percent Si by 0.467 or multiply by the inverse of this value. The following table lists the element, the oxide form, and the elemental percent in the oxide form.

<table>
<thead>
<tr>
<th>Element Form</th>
<th>Oxide</th>
<th>Elemental %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>SiO₂</td>
<td>46.7</td>
</tr>
<tr>
<td>Al</td>
<td>Al₂O₃</td>
<td>52.9</td>
</tr>
<tr>
<td>Fe</td>
<td>Fe₂O₃</td>
<td>69.9</td>
</tr>
<tr>
<td>Mg</td>
<td>MgO</td>
<td>60.3</td>
</tr>
<tr>
<td>Mn</td>
<td>MnO</td>
<td>77.4</td>
</tr>
<tr>
<td>K</td>
<td>K₂O</td>
<td>83.0</td>
</tr>
<tr>
<td>Ti</td>
<td>TiO₂</td>
<td>59.9</td>
</tr>
<tr>
<td>Ca</td>
<td>CaO</td>
<td>71.5</td>
</tr>
<tr>
<td>Zr</td>
<td>ZrO₂</td>
<td>74.0</td>
</tr>
<tr>
<td>P</td>
<td>P₂O₅</td>
<td>43.6</td>
</tr>
<tr>
<td>Na</td>
<td>Na₂O</td>
<td>74.2</td>
</tr>
</tbody>
</table>
9. Report

Data are reported as percent to the nearest tenth for Fe, Al, Mg, Na, K, and Si; to the nearest hundredth for Mn, Ca, P, and Ti; and to the nearest thousandth for Zr. The remaining trace elements (Cu, Zn, As, Se, Cd, and Pb) are reported in mg kg\(^{-1}\) (ppm).

10. Precision

The mean, standard deviation, and C.V. are calculated for each element for both the NIST standard and the SSL reference standard.

11. References


---

Surface Area (7D)
Ethylene Glycol Monoethyl Ether (EGME) Retention (7D2)

1. Application

Surface area determines many physical and chemical properties of materials. Water retention and movement, cation exchange capacity, pesticide adsorption, and many biological processes are closely related to specific surface (Carter et al., 1986). Soils vary widely in their reactive surface area because of differences in mineralogical and organic composition and in their particle-size distribution (Carter et al., 1965). Specific surface, defined as surface area per unit mass of soil, is usually expressed in units of m\(^2\) g\(^{-1}\) or cm\(^2\) g\(^{-1}\) soil. Specific surface has been measured for several clays, e.g., 810 m\(^2\) g\(^{-1}\) for smectite and 20 to 40 m\(^2\) g\(^{-1}\) for kaolinite and mica.

2. Summary of Method

Ethylene glycol monoethyl ether (EGME) retention is a surface-area determination. A soil sample is dried over phosphorus pentoxide (P\(_2\)O\(_5\)). The
sample is saturated with EGME. A monomolecular layer of EGME is established by desorbing the EGME by vacuum over EGME-saturated CaCl\textsubscript{2}. The solvate of CaCl\textsubscript{2} and EGME helps to maintain an EGME vapor pressure in the desiccator which results in the formation of a monomolecular layer of EGME on sample surfaces.

The weight of a monomolecular layer of EGME on the sample is determined by weighing the dried sample. EGME is determined by weighing the sample and sample plus EGME (Carter et al., 1965). The SSL determines EGME retention by method 7D2. The SSL reports EGME retention as mg EGME per g of soil to the nearest mg on a <2-mm base.

3. Interferences

The loss or contamination of sample and the variation in sample weight may cause erroneous results. Handle the weighing vessels with finger cots or tongs to prevent vessel contamination and the resulting weighing errors. High relative humidity in the laboratory may result in high moisture absorption by sample.

4. Safety

Wear protective clothing (e.g., coats, aprons, and gloves) and eye protection (e.g., face shields, goggles, or safety glasses) when handling reagents and working with vacuum desiccators. Follow standard laboratory safety procedures in handling reagents and vacuum devices. The P\textsubscript{2}O\textsubscript{5} is corrosive and reacts violently with water. Use caution in cleaning P\textsubscript{2}O\textsubscript{5} spills. The EGME is combustible and harmful if swallowed, inhaled, or absorbed through the skin. Keep samples and desiccators with EGME under fume hood at all times.

5. Equipment

5.1 Electronic balance, ±0.1-mg sensitivity, Mettler AE 160
5.2 Vacuum desiccator, 250 mm, Nalgene No. 5310, with desiccator plate, 230 mm
5.3 Laboratory vacuum or vacuum pump, 0.65 to 0.75 bars
5.4 EGME trap, anhydrous CaCl\textsubscript{2} in a large tube between desiccator and vacuum source
5.5 Syringe, polypropylene, 3 mL
5.6 Weighing bottle, cylindrical, low form, 50 x 30 mm

6. Reagents

6.1 Ethylene glycol monoethyl ether (EGME), reagent
6.2 Phosphorus pentoxide (P\textsubscript{2}O\textsubscript{5}), anhydrous
6.3 Calcium chloride (CaCl\textsubscript{2}), pellets, 40 mesh, reagent grade
7. Procedure

7.1 Dry 3 to 5 g of <2-mm, air-dry soil in a weighing bottle in a vacuum desiccator over P₂O₅ for 2 days.

7.2 Prepare solvated CaCl₂ by weighing 100 g oven-dried CaCl₂, without cooling, into a large beaker. Add 20 g EGME and mix by stirring. Transfer to a desiccator in which EGME-saturated samples equilibrate.

7.3 Weigh the P₂O₅-dried soil sample to the nearest 0.1 mg. When working outside the desiccator, cover the sample to avoid moisture adsorption from the atmosphere.

7.4 Use a 3-mL syringe to saturate the soil with EGME. Add 5 drops in excess of saturation.

7.5 Place the uncovered, EGME-soil mixture in a vacuum desiccator over solvated CaCl₂. Use a laboratory vacuum of 0.65 to 0.75 bar pressure.

7.6 Loosely cover the tops of weighing bottles with a piece of aluminum foil that is smaller than the inside diameter of desiccator.

7.7 Apply suction for 16 to 24 h.

7.8 Carefully release the suction. Remove weighing bottles and weigh the EGME-soil mixture.

7.9 If a 3-g sample is used, the difference between the EGME-soil mixture and P₂O₅-dry soil is ≈10 mg EGME/g P₂O₅-dry soil. When this difference is <10 mg, reduce the vacuum time to 1 h day⁻¹ and weigh twice daily.

7.10 Repeat the vacuum and weighing procedure until a constant weight is attained. Constant weight is defined as three successive daily weighings within 1 mg of EGME per gram P₂O₅-dry soil. When a constant weight is attained, make calculations.

8. Calculations

8.1 The EGME retention is calculated as follows:

Retention of EGME (mg g⁻¹) = (Wt₁ − Wt₂) x (1000/Wt₃)

where:

Wt₁ = Soil weight with monomolecular layer of EGME + Tare weight of bottle
Wt₂ = Soil weight after drying with P₂O₅ + Tare weight of bottle
Wt₃ = Soil weight after drying with P₂O₅ − Tare weight of bottle
1000 = Conversion factor (mg g⁻¹)

The surface area in units of mg EGME per g of soil is converted to m² g⁻¹, the convention commonly used in clay mineralogy. The conversion is as follows:
Surface area (m$^2$ g$^{-1}$) = (EGME retention (mg g$^{-1}$))/0.286

where:

0.286 = Conversion factor (mg EGME m$^{-2}$)

The constant, 0.286, is the amount of EGME (mg) that is required to cover a m$^2$ of clay surface with a monomolecular layer (Carter et al., 1986). This value is calculated from the measured value of 231.7 mg EGME per g of pure montmorillonite assumed to have 810 m$^2$ g$^{-1}$ on the basis of other measurements.

9. Report

Report EGME as mg EGME per g of soil to the nearest mg.

10. Precision

Precision data are not available for this procedure. Two quality control checks, a high and a low standard, are routinely analyzed in EGME. The mean (mg EGME per g soil), standard deviation, and C.V. for the quality control check sample are as follows:

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>n</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Std</td>
<td>109.0</td>
<td>10</td>
<td>7.4</td>
<td>7.3</td>
</tr>
<tr>
<td>Low Std.</td>
<td>37.5</td>
<td>10</td>
<td>0.64</td>
<td>4.8</td>
</tr>
</tbody>
</table>

11. References


MISCELLANEOUS (8)

Ratios and Estimates (8D)

To Noncarbonate Clay (8D2)

Divide the CEC-7 (method 5A8c), extractable Fe (method 6C2), or 15-bar water retention (method 4B2a or 4B2b) by the noncarbonate clay percentage. Noncarbonate clay is determined by subtracting the carbonate clay (method 3A1d or 3A2d) from total clay (method 3A1 or 3A2).
Ratios and Estimates (8D)
Ca to Mg (extractable) (8D3)
Divide extractable Ca\(^{2+}\) (method 6N2) by extractable Mg\(^{2+}\) (method 6O2).

Ratios and Estimates (8D)
Estimated Clay Percentage (8D4)
For most soils, clay percentage can be approximated as 2.5 \times 15-bar water percentage (method 4B2a or 4B2b). Use caution in applying this factor to any particular situation, especially if organic matter or other amorphous material is present in significant quantities.

Ratios and Estimates (8D)
Estimated Total Salt (8D5)
Use the charts and graphs available in U.S. Salinity Laboratory Staff (1954) to estimate total salt content from the electrical conductivity (EC\(_{s}\)) of the saturation extract (method 8A3a). The essential relations are summarized in the equations as follows:

\[
\text{Log total salt in soil (ppm)} = 0.81 + 1.08 \times \text{Log ECs (mmhos cm}^{-1}) + \text{Log SP}
\]

where:
- EC\(_{s}\) = Electrical conductivity of saturation extract
- SP = Saturation percentage of saturation extract

Total salt in soil (%) = Total salt (ppm) \times 10^{-4}

These equations are applicable to saturation extracts with an EC\(_{s}\) <20 mmhos cm\(^{-1}\). Deviations occur at higher salt concentrations.

Ratios and Estimates (8D)
Iron Plus Aluminum, Pyrophosphate Extractable to Dithionite-Citrate Extractable (8D6)
Divide the sum of the pyrophosphate-extractable Fe plus Al (methods 6C8a and 6G10a, respectively) by the sum of dithionite-citrate-extractable Fe plus Al (methods 6C2 and 6G7, respectively). Pyrophosphate and dithionite-citrate extractable Fe and Al are former criteria for spodic placement (Soil Survey Staff, 1975).
Ratios and Estimates (8D)
Index of Accumulation (8D7)

Subtract \( \frac{1}{2} \) the clay percentage (method 3A1 or 3A2) of a subhorizon from the CEC at pH 8.2 (method 5A3a) and multiply the remainder by the thickness of subhorizon (cm). The combined index of accumulation of amorphous material is a former criterion for spodic placement (Soil Survey Staff, 1975).

References


Use Table 1 with the SSL preparation methods 1B1, 1B2, 1B5, 1B6, and 1B7. Gravel codes are also defined in Table 1. In the “Code” column, “Char” refers to characterization sample. Laboratory preparation and >2-mm porosity are defined in footnotes on laboratory data sheet.

Table 1.—Laboratory Preparation Codes and Procedural Summaries.

<table>
<thead>
<tr>
<th>Code</th>
<th>Laboratory Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Char &gt;2 mm</td>
<td>Weigh sample at field-moisture content and record weight. Air-dry, weigh, and record weight. Sieve &gt;2-mm fractions, weigh, record weights, and discard. Report all analytical results on &lt;2-mm basis. Refer to method 1B1, Standard Air-dry.</td>
</tr>
<tr>
<td>S Blank</td>
<td>Lab preparation is same as S-blank. However, report clod parameters and Cm (correction factor for &gt;2-mm content moist soil) on an whole-soil basis. Refer to method 1B1, Standard Air-dry.</td>
</tr>
<tr>
<td>S P</td>
<td>Lab preparation is same as S-blank except do not record the weight of the &gt;2-mm fraction. All analytical results are reported on a &lt;2-mm basis. Refer to method 1B1, Standard Air-dry.</td>
</tr>
<tr>
<td>N Blank</td>
<td></td>
</tr>
</tbody>
</table>

824
<table>
<thead>
<tr>
<th>Code</th>
<th>Laboratory Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Char</strong></td>
<td>&gt;2 mm</td>
</tr>
<tr>
<td>M</td>
<td>Blank</td>
</tr>
<tr>
<td>S</td>
<td>K</td>
</tr>
<tr>
<td>S</td>
<td>R</td>
</tr>
<tr>
<td>G</td>
<td>P</td>
</tr>
<tr>
<td>W</td>
<td>P</td>
</tr>
<tr>
<td>H</td>
<td>Blank</td>
</tr>
</tbody>
</table>

Lab preparation is same as S-blank except:

- **Char >2 mm**: Lab preparation is same as S-blank except sieve <2-mm moist subsample for 15-bar moist analysis. Use <2-mm air-dry soil for all other analyses. Report all analytical results on <2-mm basis. Refer to method 1B2, Field-moist.

- **M Blank**: Lab preparation is same as S-blank except sieve <2-mm moist subsample for 15-bar moist analysis. Use <2-mm air-dry soil for all other analyses. Report all analytical results on <2-mm basis. Refer to method 1B2, Field-moist.

- **S K**: Lab preparation is same as S-blank except grind the 2- to 20-mm fraction to <2 mm and keep for CO₃ analyses, etc. Report the analytical results for the ground 2- to 20-mm fraction on a 2- to 20-mm basis and all other analytical results on a <2-mm basis. Refer to method 1B5, Coarse Fragments.

- **S R**: Lab preparation is same as S-blank except recombine the 2- to 20-mm fraction with the <2-mm fraction and grind the entire sample to <2 mm. Report all analytical results for ground sample on a <2-mm basis. Refer to method 1B5, Coarse Fragments.

- **G P**: Weigh sample at field-moisture content and record weight. Air-dry, weigh, and record weight. Grind entire sample to <2 mm. Report all analytical results for ground sample on a whole-soil basis. Refer to method 1B6, Whole-soil.

- **W P**: Weigh sample at field-moisture content and record weight. Air-dry, weigh, and record weight. Sieve >2-mm fractions, weigh, and record weights. Recombine the >2-mm fractions with the <2-mm fraction and grind entire sample to <2 mm. Report all analytical results on a whole-soil basis. This procedure is no longer performed at the SSL.

- **H Blank**: Obtain a moist whole-soil subsample for Histosol analysis. Obtain a <2-mm moist subsample for 15-bar moist analysis. Weigh remaining sample at field-moisture content and record weight. Air-dry, weigh, and record weight. Sieve >2-mm fractions, weigh, record weights, and discard. Pulverize subsample of <2-mm air-dry soil to a <80-mesh size and use for lab analyses. Use <80-mesh air-dry for all analyses except AD/OD, 15-, 1/10-, and 2-bar analyses. For the AD/OD, 15-, 1/10-, and 2-bar analyses, use <2-mm air-dry soil. Use <2-mm moist subsample for 15-bar moist. Report all analytical results except fabric on a <2-mm basis. Refer to method 1B7, Organic Material.
<table>
<thead>
<tr>
<th>Code</th>
<th>Laboratory Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Char &gt;2 mm</td>
<td>Lab preparation is same as N-blank except pulverize subsample of &lt;2-mm air-dry soil to a &lt;80-mesh size and use for lab analyses. Use &lt;80-mesh air-dry for all analyses except AD/OD and 15-bar analyses. For the AD/OD and 15-bar analyses, use &lt;2-mm air-dry soil. All analytical results are reported on a &lt;2-mm basis. Refer to method 1B1, Standard Air-dry.</td>
</tr>
<tr>
<td>A (L) Blank</td>
<td></td>
</tr>
</tbody>
</table>

**Gravel codes**

**P** = Porous >2-mm material that is considered soil is used for clod or core measurements.

**V** = Volume estimate is used to calculate the weight percentage of a >2-mm fraction. If that fraction is porous (P), code the samples with “P” rather than with “V”. 
ION EXCHANGE ANALYSES (5)

Cation Exchange Capacity (5A)
NH₄OAc, pH 7.0 (5A8)
Automatic Extractor (CEC-7)
   Steam Distillation (5A8b)

1. Application
   The CEC determined with 1 N NH₄OAc buffered at pH 7.0 is a commonly used method and has become a standard reference to which other methods are compared (Peech et al., 1947). The advantages of using this method are that the extractant is highly buffered so that the extraction is performed at a constant, known pH (7.0) and that the NH₄⁺ on the exchange complex is easily determined.

2. Summary of Method
   Displacement after washing is the basis for this procedure. The CEC is determined by saturating the exchange sites with an index cation (NH₄⁺); washing the soil free of excess saturated salt; displacing the index cation (NH₄⁺) adsorbed by the soil; and measuring the amount of the index cation (NH₄⁺). A sample is leached using 1 N NH₄OAc and a mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed and saved for analyses of the cations. The NH₄⁺ saturated soil is rinsed with ethanol to remove the NH₄⁺ that was not adsorbed. Steam distillation and titration are used to determine the NH₄⁺ adsorbed on the soil exchange complex. The CEC by NH₄OAc, pH 7 is reported in meq/100 g oven-dry soil in method 5A8b (Soil Conservation Service, 1984).

3. Interferences
   Incomplete saturation of the soil with NH₄⁺ and insufficient removal of NH₄⁺ are the greatest interferences to this method. Ethanol removes some adsorbed NH₄⁺ from the exchange sites of some soils. Isopropanol rinses have been used for some soils in which ethanol removes adsorbed NH₄⁺. Soils that contain large amounts of vermiculite can irreversibly “fix” NH₄⁺. Soils that contain large amounts of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract.
4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjeltec Auto 1030 Analyzer.

5. Equipment

5.1 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE

5.2 Syringes, polypropylene, disposable, 60 mL, for sample tube, extractant reservoir, and tared extraction syringe

5.3 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels

5.4 Polycorns, Richards Mfg. Co.

5.5 Kjeltec Auto 1030 Analyzer, Tecator, Fisher Scientific Inc.

5.6 Digestion tubes, straight neck, 250 mL

5.7 Analytical filter pulp, Schleicher and Schuell, no. 289

5.8 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.

5.9 Electronic balance, ±1-mg sensitivity

6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Ammonium acetate solution (NH₄OAc), 1 N, pH 7.0. Add 1026 mL of glacial acetic acid (CH₃COOH) to 15 L DDI water. Add 1224 mL of conc. ammonium hydroxide (NH₄OH). Mix and cool. Dilute with DDI water to 18 L and adjust to pH 7.0 with CH₃COOH or NH₄OH.

6.3 Ethanol (CH₃CH₂OH), 95%, U.S.P.

6.4 Nessler’s reagent. Add 4.56 g of potassium iodide (KI) to 30 mL DDI water. Add 5.68 g of mercuric iodide (Hgl₂). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL of DDI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with DDI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately.
6.5 Sodium chloride (NaCl), reagent, crystal
6.6 Antifoam agent, slipicone release spray, Dow Chemical Corp. Alternatively, mix equal parts of mineral oil and n-octyl alcohol.
6.7 Boric acid, 4% (w:v), with bromcresol green-methyl red indicator (0.075% bromcresol green and 0.05% methyl red), Ricca Chemical Co.
6.8 Hydrochloric acid (HCl), 0.1 N, standardized. Dilute 148 mL of conc. HCl in 16 L of DDI water.
6.9 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of DDI water. Dilute to 9 L with DDI water.

7. Procedure

Extraction of Bases

7.1 Prepare sample tube by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.
7.2 Weigh 2.50 g of <2-mm, air-dry soil and place in sample tube. Prepare one quality control check sample per 48 samples.
7.3 Place sample tube on upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1 in) length rubber tubing and insert the plunger in the slot of the stationary disk of the extractor.
7.4 Use a squeeze bottle to fill sample tube to the 20-mL mark with NH$_4$OAc solution (≈10 mL). Thoroughly wet the sample. Let stand for at least 20 min.
7.5 Put reservoir tube on top of the sample tube. Rapidly extract the NH$_4$OAc solution to a 0.5- to 1.0-cm height above sample. Turn off extractor. Add ≈45 mL of NH$_4$OAc solution to the reservoir tube. Set extractor for an overnight (12 to 16 h) extraction.
7.6 Next morning turn off the extractor. Pull the plunger of the syringe down. Do not pull plunger from the barrel of the syringe. Carefully remove the syringe containing the extract. Leave the rubber tubing on the sample tube. Weigh each syringe containing the NH$_4$OAc extract to the nearest 0.01 g.
7.7 Mix the extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. The solution in the polycon is reserved for analyses of extracted cations (methods 6N2e, 6O2d, 6P2b, and 6Q2b).

Removal of Excess Ammonium Acetate

7.8 Return the extractor to starting position. Attach syringe to the sample tube and rinse the sides of the sample tube with ethanol from a wash bottle. Fill the sample tube to the 20-mL mark with ethanol and let stand for 15 to 20 min.
7.9 Place reservoir tube on the sample tube. Rapidly extract the ethanol level to a 0.5- to 1.0-cm height above the sample. Turn off the extractor and add 55 to 60 mL of ethanol to the reservoir. Extract at a 45-min rate.

7.10 After the extractor has stopped, turn off the switch. Pull the plunger of the syringe down. Do not pull the plunger from the syringe barrel. Remove the syringe and discard the ethanol.

7.11 Repeat the ethanol wash.

7.12 After the second wash, collect a few drops of ethanol extract from the sample tube on a spot plate. Test for NH$_4^+$ by using Nessler’s reagent. A yellow, red to reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler’s reagent. Repeat until a negative test is obtained.

Steam Distillation: Samples and Reagent Blanks

7.13 Remove the sample tube and transfer the sample with filter pulp to a 250-mL digestion tube. Add 6 to 7 g of NaCl to the digestion tube. Use a gentle flow of compressed air to blow the filter pulp and sample out of the syringe. Wash the tube with DDI water and use a rubber policeman to complete transfer. The amount of distilled water that is added depends on the amount that is required to complete the transfer of tube contents.

7.14 Perform the same transfer and addition of reagents for blanks as for samples.

7.15 Spray silicone antifoam agent (or 2 drops of octyl alcohol) into the digestion tubes for each of the samples and reagent blanks.

7.16 When using new reagents, e.g., boric acid, reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.17 On bench worksheet, record the normality of standardized acid, i.e., $\approx 0.1 \, N$ HCl.

7.18 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

$$\text{CEC (meq/100 g)} = \frac{(\text{Titer} \times N \times 100 \times \text{AD/OD})}{(\text{Weight})}$$

where:

- Titer = Titer of sample (mL)
- $N$ = Normality of HCl titrant
9. Report

Report CEC-7 in units of meq/100 g of oven-dry soil to the nearest 0.1 meq/100 g.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples. With 113 observations of the quality control check sample, the mean, standard deviation, and C.V. for the CEC are 27.1, 0.57, and 2.1%, respectively.

11. References


NH₄Cl, pH 7.0 (5A9)
Steam Distillation (5A9b)

1. Application

The CEC determined with a neutral unbuffered salt, e.g., 1 N NH₄Cl, is an estimate of the “effective” CEC (ECEC) of the soil (Peech et al., 1947). For a soil with a pH of <7.0, the ECEC value should be < CEC measured with a buffered solution at pH 7.0. The NH₄Cl CEC is ≈ equal to the NH₄OAc extractable bases plus the KCl extractable Al for noncalcareous soils.

2. Summary of Method

Displacement after washing is the basis for this procedure. The CEC is determined by saturating the exchange sites with an index cation (NH₄⁺); washing the soil free of excess saturated salt; displacing the index cation (NH₄⁺) adsorbed by the soil; and measuring the amount of the index cation (NH₄⁺). A sample is leached using 1 N NH₄Cl and a mechanical vacuum extractor (Holmgren et al., 1977). The extract is weighed and saved for analyses of the cations. The NH₄⁺
saturated soil is rinsed with ethanol to remove the \( \text{NH}_4^+ \) that was not adsorbed. Steam distillation and titration are used to determine the \( \text{NH}_4^+ \) adsorbed on the soil exchange complex. The CEC by \( \text{NH}_4\text{Cl} \) is reported in meq/100 g oven-dry soil in method 5A9b (Soil Conservation Service, 1984).

3. Interferences

Incomplete saturation of the soil with \( \text{NH}_4^+ \) and insufficient removal of \( \text{NH}_4^+ \) are the greatest interferences to this method. Ethanol removes some adsorbed \( \text{NH}_4^+ \) from the exchange sites of some soils. Isopropanol rinses have been used for some soils in which ethanol removes adsorbed \( \text{NH}_4^+ \). Soils that contain large amounts of vermiculite can irreversibly “fix” \( \text{NH}_4^+ \). Soils that contain large amounts of soluble carbonates can change the extractant pH and/or can contribute to erroneously high cation levels in the extract.

4. Safety

Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses) when preparing reagents, especially concentrated acids and bases. Dispense concentrated acids and bases in a fume hood. Thoroughly wash hands after handling reagents. Use the safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

Ethanol is flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the vacuum extractor and the Kjeltec Auto 1030 Analyzer.

5. Equipment

5.1 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.2 Syringes, polypropylene, disposable, 60 mL, for sample tube, extractant reservoir, and tared extraction syringe
5.3 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in), for connecting syringe barrels
5.4 Polycons, Richards Mfg. Co.
5.5 Kjeltec Auto 1030 Analyzer, Tecator, Fisher Scientific Inc.
5.6 Digestion tubes, straight neck, 250 mL
5.7 Analytical filter pulp, Schleicher and Schuell, no. 289
5.8 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.9 Electronic balance, ±1-mg sensitivity
6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Ammonium chloride solution (NH₄Cl), 1 N. Dissolve 535 g of NH₄Cl reagent in DDI water and dilute to 10 L.

6.3 Ethanol (CH₃CH₂OH), 95%, U.S.P.

6.4 Nessler’s reagent. Add 4.56 g of potassium iodide (KI) to 30 mL DDI water. Add 5.68 g of mercuric iodide (HgI₂). Stir until dissolved. Dissolve 10 g of sodium hydroxide (NaOH) in 200 mL DDI water. Transfer NaOH solution to a 250-mL volumetric flask and slowly add K-Hg-I solution. Dilute to volume with DDI water and thoroughly mix. Solution should not contain a precipitate. Solution can be used immediately.

6.5 Sodium chloride (NaCl), reagent, crystal

6.6 Antifoam agent, slipicone release spray, Dow Chemical Corp. Alternatively, mix equal parts of mineral oil and n-octyl alcohol.

6.7 Boric acid, 4% (w:v), with bromcresol green-methyl red indicator (0.075% bromcresol green and 0.05% methyl red), Ricca Chemical Co.

6.8 Hydrochloric acid (HCl), 0.1 N, standardized. Dilute 148 mL of conc. HCl in 16 L of DDI water.

6.9 NaOH, 1 M. Add 500 mL of 50% NaOH solution to 8 L of DDI water. Dilute to 9 L with DDI water.

7. Procedure

Extraction of Bases

7.1 Prepare sample tube by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh 2.50 g of <2-mm, air-dry soil and place in sample tube. Prepare one quality control check sample per 48 samples.

7.3 Place sample tube on upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1 in) length rubber tubing and insert the plunger in the slot of the stationary disk of the extractor.

7.4 Use a squeeze bottle to fill sample tube to the 20-mL mark with NH₄Cl solution (~10 mL). Thoroughly wet the sample. Let stand for at least 20 min.

7.5 Put reservoir tube on top of the sample tube. Rapidly extract the NH₄Cl solution to a 0.5- to 1.0-cm height above sample. Turn off extractor. Add ≈45 mL of NH₄Cl solution to the reservoir tube. Set extractor for an overnight (12 to 16 h) extraction.
7.6 Next morning turn off the extractor. Pull the plunger of the syringe down. Do not pull plunger from the barrel of the syringe. Carefully remove the syringe containing the extract. Leave the rubber tubing on the sample tube. Weigh each syringe containing the NH₄Cl extract to the nearest 0.01 g.

7.7 Mix the extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. The solution in the polycon is reserved for analyses of extracted cations (methods 6N2e, 6O2d, 6P2b, and 6Q2b).

**Removal of Excess Ammonium Chloride**

7.8 Return the extractor to starting position. Attach syringe to the sample tube and rinse the sides of the sample tube with ethanol from a wash bottle. Fill the sample tube to the 20-mL mark with ethanol and let stand for 15 to 20 min.

7.9 Place reservoir tube on the sample tube. Rapidly extract the ethanol level to a 0.5- to 1.0-cm height above the sample. Turn off the extractor and add 55 to 60 mL of ethanol to the reservoir. Extract at a 45-min rate.

7.10 After the extractor has stopped, turn off the switch. Pull the plunger of the syringe down. Do not pull the plunger from the syringe barrel. Remove the syringe and discard the ethanol.

7.11 Repeat the ethanol wash.

7.12 After the second wash, collect a few drops of ethanol extract from the sample tube on a spot plate. Test for NH₄⁺ by using Nessler’s reagent. A yellow, red to reddish brown precipitate is a positive test. If the test is positive, repeat the ethanol wash and retest with Nessler’s reagent. Repeat until a negative test is obtained.

**Steam Distillation: Samples and Reagent Blanks**

7.13 Remove the sample tube and transfer the sample with filter pulp to a 250-mL digestion tube. Add 6 to 7 g of NaCl to the sample. Use a gentle flow of compressed air to blow the filter pulp and sample out of the syringe. Wash the tube with DDI water and use a rubber policeman to complete transfer. The amount of distilled water that is added depends on the amount that is required to complete the transfer of tube contents.

7.14 Perform the same transfer and addition of reagents for blanks as for samples.

7.15 Spray silicone antifoam agent (or 2 drops of octyl alcohol) into the digestion tubes for each of the samples and reagent blanks.
7.16 When using new reagents, e.g., boric acid, reagent blanks are distilled in 2 sets of 6, one set per Kjeltec machine. Each set of 6 is averaged and recorded on bench worksheet and manually set on each machine. During the steam distillation, the mean reagent blank titer is automatically subtracted from the sample titer.

7.17 On bench worksheet, record the normality of standardized acid, i.e., \( \approx 0.1 \text{ } N \text{ HCl} \).

7.18 Connect the tube to the distillation unit. Close the safety door. Distillation and titration are performed automatically. Record the titer in mL of titrant.

8. Calculations

\[
\text{CEC (meq/100 g)} = \frac{(\text{Titer} \times N \times 100 \times \text{AD/OD})}{\text{Weight}}
\]

where:
- Titer = Titer of sample (mL)
- \( N \) = Normality of HCl titrant
- Weight = Sample weight (g)
- 100 = Conversion factor to 100 g basis
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report neutral salts CEC in units of meq/100 g of oven-dry soil to the nearest 0.1 meq/100 g.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples. With 19 observations of the quality control check sample, the mean, standard deviation, and C.V. for the CEC are 26.0, 0.37, and 1.4\%, respectively.

11. References


CHEMICAL ANALYSES (6)

Total Carbon (6A)
Dry Combustion (6A2)
LECO CR-12 Carbon Analyzer (6A2d)

1. Application

Total C in soils is the sum of organic and inorganic C. Most of the organic C is associated with the organic matter fraction, and the inorganic C is generally found with carbonate minerals. The organic C in mineral soils generally ranges from 0 to 12%.

Total C is quantified by two basic methods, i.e., wet or dry combustion. The SSL uses dry combustion. In total C determinations, all forms of C in a soil are converted to CO$_2$ followed by a quantification of the evolved CO$_2$. Total C can be used to estimate the organic C content of a soil. The difference between total and inorganic C is an estimate of the organic C. Organic C also can be determined directly (method 6A1c). The inorganic C should be equivalent to carbonate values measured by CO$_2$ evolution with strong acid (Nelson and Sommers, 1982).

Organic C defines mineral and organic soils. In Soil Taxonomy, organic C is also used at lower taxonomic levels, e.g., ustollic and fluventic subgroups (Soil Survey Staff, 1975).

2. Summary of Method

An 80-mesh soil sample is oxidized at high temperatures. The released gases are scrubbed, and the CO$_2$ in the combustion gases is measured by using an infrared detector. Percent total C is reported on an oven-dry soil basis.

3. Interferences

This procedure simultaneously measures inorganic and organic C.

4. Safety

Wear protective clothing and safety glasses. Magnesium perchlorate may form explosive mixtures. Magnesium perchlorate may contain traces of perchloric acid, which remain from manufacturer’s operations. This acid is anhydrous because of the strong desiccating capability of the salt. Avoid prolonged contact with oxidizable material or material capable of forming unstable perchlorate esters or salts. Remove magnesium perchlorate by using an excess of water to thoroughly dilute the material.

The use of high temperatures in the oxidation of samples requires that extreme caution be used to prevent burns and fires. Follow standard laboratory procedures when handling compressed gases. Oxygen is highly flammable.
Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the carbon analyzer.

5. Equipment
5.2 Data transmit card, part no. 772-573, Leco Corp., St. Joseph, MI
5.3 Combustion boats, part no. 529-203, Leco Corp., St. Joseph, MI
5.4 Single-stage regulator, oxygen service, part no. E11-W-N115Box, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.4 Electronic balance, ±1-mg sensitivity

6. Reagents
6.1 Anhydrone, anhydrous magnesium perchlorate, granular
6.2 Glass wool
6.3 Compressed oxygen, >99.5% @ 30 psi
6.4 Calcium carbonate, CaCO₃, reagent grade

7. Procedure
7.1 Use a fine-ground 80-mesh, air-dry soil
7.2 Weigh sample in a tared combustion boat. The sample size is dependent upon the C content. The product of sample weight (g) multiplied by C percentage should not be >10%. In most cases, the sample size is 1.00 g, unless the C content is >10%.
7.3 Refer to the manufacturer’s manual for operation of carbon analyzer.
7.4 Combust sample in an O₂ atmosphere in which the C is oxidized to CO₂. Moisture and dust are removed by the instrument, and the CO₂ gas is then measured by a solid state infrared detector. The microprocessor formulates the analytical results (Cᵢ) by combining the outputs of the infrared detector and the system ambient sensors with pre-programmed calibration, linearization and weight compensation factors. Analytical results are displayed and printed on the control console.

8. Calculations

\[ C(\%) = Cᵢ \times \frac{AD}{OD} \]

where:
\[ C(\%) = C(\%), \text{ oven-dry basis} \]
C_i=C (%) instrument
AD/OD=air−dry/oven-dry ratio (method 4B5)

9. Report
   Report total C percentage on an oven-dry basis to the nearest 0.1%.

10. Precision
   Precision data are not available for this procedure. A quality control check sample is included in every batch of ten samples. For 41 observations of the quality control check sample, the mean, standard deviation, and C.V. for total carbon are 11.38, 0.062, and 5.5%, respectively.

11. References

Nitrogen (6B)
Kjeldahl Digestion II (6B3)
Ammonia Steam Distillation, Automatic Titrator (6B3a)

1. Application
   The total N content of the soil may range from <0.02% in subsoils, 2.5% in peats, and 0.06 to 0.5% in surface layers of many cultivated soils (Bremmer and Mulvaney, 1982). The total N data may be used to determine the soil C:N ratio, the soil potential to supply N for plant growth, and the N distribution in the soil profile. The C:N ratio generally ranges between 10 to 12. Variations in the C:N ratio may serve as an indicator of the amount of soil inorganic N. Uncultivated soils usually have higher C:N ratios than do cultivated soils.
   Soils with large amounts of illites or vermiculites can “fix” significant amounts of N compared to those soils dominated by smectites or kaolinites (Young and Aldag, 1982; Nommik and Vahtras, 1982). Since the organic C of many soils diminishes with depth while the level of “fixed” N remains constant or increases, the C:N ratio narrows (Young and Aldag, 1982). The potential to “fix” N has important fertility implications as the “fixed” N is slowly available for plant growth.
2. Summary of Method

A soil sample is digested using the Kjeldahl technique. The digest is made alkaline, the steam is distilled to release NH$_4^{+}$-N, and the NH$_4^{+}$-N is complexed with boric acid. The complexed NH$_4^{+}$-N is titrated with HCl, and the total N is calculated against a reagent blank (Soil Conservation Service, 1984).

3. Interferences

The total N that is measured by the Kjeldahl method does not distinguish among the types of N that are present in the soil. Practically all of the N is measured, but some forms of N are not recovered. Generally, soils have small amounts of N in the nonrecoverable forms, i.e., NO$_3^-$ and NO$_2^-$. Soils with significant amounts of NO$_3^-$ or NO$_2^-$ are usually saline. The anion analysis of the saturated paste extracts measures NO$_3^-$ and NO$_2^-$ (methods 6M1c and 6W1a, respectively).

The most significant error in the Kjeldahl method is the heating of the digestion mixture over 400 °C. Loss of N occurs when the temperature of the digestion is >400 °C (Bremmer and Mulvaney, 1982).

4. Safety

Wear protective clothing (coats, aprons, sleeve guards and gloves) and eye protection (face shields, goggles, or safety glasses) when handling acids and bases. Use heat resistant gloves when handling hot digestion tubes during digestion and steam distillation. Use the provided safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids. Digestion blocks are used at high temperatures, i.e., 250 and 400 °C. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Use the fume hood and fume aspiration devices to control and dispose of the acid fumes when digesting samples.

Boric acid is toxic and must not be ingested. Hengar granules contain Se which is toxic. Concentrated H$_2$SO$_4$ reacts violently with water and must be handled with caution. The 50% NaOH solution is very corrosive. Follow prudent laboratory safety precautions when handling these chemicals. Follow the manufacturer’s safety precautions when using the Kjeltec Auto 1030 Analyzer.

5. Equipment

5.1 Electronic balance, ±0.001-g sensitivity
5.2 Digestion tubes, 250 mL, with constricted neck, Ace Glass Co., Inc.
5.3 Digestion blocks, 250 and 400 °C
5.4 Dispenser, Zippette, 30 mL or equivalent, for conc. sulfuric acid (H$_2$SO$_4$), Brinkmann Instruments Inc.
5.5 Kjeltec Auto 1030 Analyzer, Tecator, Fisher Scientific Inc.
6. Reagents

6.1 Distilled water
6.2 Distilled deionized (DDI) water
6.3 Hydrochloric acid (HCl), conc., 12 N
6.4 Sodium hydroxide (NaOH), 50% (w:v), reagent
6.5 Hengar granules (selenized)
6.6 Digestion salt mixture. Mix 1000 g of potassium sulfate powder, 55 g of ferrous sulfate powder (anhydrous), and 32 g of copper II sulfate powder (anhydrous) in a tumbling mill for at least 30 min.
6.7 Antifoam, silicone spray bottle, Slipicone release spray, Dow Chemical Corp.
6.8 Boric acid, 4% (w:v), with bromcresol green-methyl red (0.075% bromcresol green and 0.05% methyl red) indicator, Ricca Chemical Co.
6.9 HCl, 0.1 N, standardized. Dilute 148 mL of conc. HCl in 16 L of DDI water.
6.10 Sucrose

7. Procedure

**Kjeldahl Digestion of Sample**

7.1 Weigh 3.000 g of <2-mm, air-dry soil into a 250-mL digestion tube. Refer to Table 1 for sample size.
7.2 Prepare 3 to 5 reagent blanks in every batch of 20 analyses. Reagent blanks contain 0.5 g of sucrose plus all reagents used in sample analysis, i.e., 12 mL of H₂SO₄, 4.5 g of digestion salt mixture, and 1 or 2 Hengar granules. Samples do not receive the 0.5 g of sucrose. Reagent blanks are run as samples and are not automatically subtracted during distillation procedure.
7.3 Use a dispenser to add 5 mL of distilled water to sample tube. Shake the tube to wet the sample.
7.4 Use a dispenser to add 12 mL of conc. H₂SO₄ to sample.
7.5 Allow sample to stand overnight.
7.6 Use a calibrated scoop to add 4.5 g of digestion salt mixture to sample.
7.7 Add 1 or 2 Hengar granules to sample.
7.8 Preheat one digestion heating block to 250 °C and the other to 400 °C.
7.9 Place the tube in the 250 °C block, attach a fume aspirator, and digest for at least 30 min.
7.10 Remove the tube, place in the 400 °C block, and digest sample for 1 h.
7.11 Remove the tube, place on a cooling board, and allow sample to cool for at least 15 min.

7.12 Remove the aspirator. Add 50 mL of distilled water.

Table 1.—Sample Size for Total N Based on Volume of Titrant (FeSO₄) Used in Organic C Analysis (method 6A1c).

<table>
<thead>
<tr>
<th>Fe₂SO₄ (mL)</th>
<th>Sample Size (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;6.00</td>
<td>3.0</td>
</tr>
<tr>
<td>4.00 to 5.00</td>
<td>2.0</td>
</tr>
<tr>
<td>3.00 to 4.00</td>
<td>1.5</td>
</tr>
<tr>
<td>&lt;2.00</td>
<td>1.0</td>
</tr>
</tbody>
</table>

If >10.00 mL of K₂Cr₂O₇ (method 6A1c) is used and/or sample size is <1.00 g, then divide the volume of FeSO₄ by 2 and then use Table 1. To obtain a representative sample, do not use a sample size <0.5 g for total N analysis.

**Ammonia Steam Distillation, Automatic Titrator**

7.13 Spray silicone antifoam solution (or 2 drops of octyl alcohol) into the digestion tube and connect to the distillation unit.

7.14 Close the safety door.

7.15 Distillation and titration are performed automatically.

7.16 On bench worksheet, record the mL that are titrated for samples and reagent blanks. On bench worksheet, also record the normality of standardized HCl.

**8. Calculations**

\[
N (%) = \frac{[(\text{Titer}_{\text{sample}} - \text{Titer}_{\text{blank}}) \times N \times 1.4 \times \text{AD/OD}]}{\text{Sample Weight}}
\]

where:

- \( \text{Titer}_{\text{sample}} \) = Titer of sample (mL)
- \( \text{Titer}_{\text{blank}} \) = Average titer of reagent blank (mL)
- \( N \) = Normality of HCl titrant solution
- 1.4 = Conversion factor
- \( \text{AD/OD} \) = Air-dry/oven-dry ratio (method 4B5)
9. Report

Report total N as a dimensionless value to the nearest 0.001 unit on an oven-dry basis.

10. Precision

Precision data are not available for this procedure. For 105 observations of the quality control check sample, the mean, standard deviation, and C.V. for total N are 0.143, 0.004, and 2.7%, respectively.

11. References


Iron, Aluminum, and Potassium (6C, 6G, and 6Q)
HF Dissolution (6C7, 6G11, and 6Q3)
Atomic Absorption (6C7a, 6G11a and 6Q3a)

1. Application

Historically, elemental analysis was developed for the analysis of rocks and minerals (Washington, 1930). The elemental analysis of soils, sediments, and rocks necessitates their decomposition into soluble forms. Hydrofluoric acid (HF) is efficient in the digestion and dissolution of silicate minerals for elemental decomposition. Elemental concentrations of Fe, Al, and K are determined by atomic absorption using 100 mg of clay suspension contained in a closed vessel with boric acid (H₃BO₃) to neutralize excess acid (Berdanier, Lynn, and Threlkeld, 1978; Soil Conservation Service, 1984).

2. Summary of Method

To 100 mg of clay suspension (method 7A2i), 5 mL of HF acid are added. The solution is heated, cooled, and 2 to 3 g of H₃BO₃ are added to neutralize excess acid. The solution is diluted to 100 mL, allowed to stand overnight, and 20 mL are decanted (method 7C3). The concentrations of Fe, Al, and K are determined by
atomic absorption (AA) in methods 6C7a, 6G11a, and 6Q3a, respectively. Data are reported in method 7C3.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in AA analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

The stable matrix system (HBF₄-H₃BO₃-ionic constituents of silicates) provides a suitable salt-free single matrix that greatly diminishes the chemical ionization, matrix, and instrumental interferences for AA determinations. One of the principal advantages of this technique is that all elements may be determined from a single sample solution (Lim and Jackson, 1982).

4. Safety

There are no significant hazards to analyst by this procedure. Wear protective clothing, e.g., coats and aprons. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Atomic Absorption spectrophotometer (AA), Perkin-Elmer Corp., Norwalk, CT

5.2 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT

5.3 Microcomputer, 7500 Professional Computer, Perkin-Elmer Corp., Norwalk, CT

5.4 Dot matrix printer, P-132, Interdigital Data Systems, Inc.

5.5 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. box 10030, Reno, NV, 89510

5.6 Syringes, 10,000 and 1000 µL, 1001 DX an 10110-TEL LL gastight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510

5.7 Centrifuge tubes, polystyrene, 15 mL, conical bottom, graduated, part no. 2087, for sample dilution and sample changer, Becton Dickinson Labware, Becton Dickinson and Co., 2 Bridgewater Lane, Lincoln, Park, NJ 07035

5.8 Containers, polypropylene or teflon

6. Reagents

6.1 Distilled Deionized (DDI) water

6.2 Sodium chloride (NaCl) solution, 1143 ppm Na. Dissolve 5.81 g of NaCl in 2 L of DDI water.
6.3 Boric acid, $\text{H}_2\text{BO}_3$

6.4 Hydrofluoric acid (HF) solution, 2.47 $N$. Fill a polyethylene volumetric flask ⅓ full with DDI water. In hood, slowly and carefully add 49.36 g of HF. Slowly and carefully add 20 g of $\text{H}_2\text{BO}_3$. Hot reaction. May not completely dissolve. Make to 1-L volume with DDI water. Store HF solution in refrigerator. Use HF solution as reagent blank.

6.5 Fe stock solution, 1000 ppm. Commercial. Weigh 1.0000 g of Fe wire, dissolve in HCl, and make to 1-L volume with DDI water. Store in polypropylene container.

6.6 Al stock solution, 1000 ppm. Commercial. Weigh 1.0000 g of Al wire, dissolve in HCl, and make to 1-L volume with DDI water. Store in polypropylene container.

6.7 K stock solution, 50 meq L$^{-1}$. Dissolve 3.7279 g of KCl in 1 L of DDI. Store in polypropylene container.

6.8 Fe standard, 200 ppm. To 50 ml of Fe stock solution, add 12.34 ml of HF solution and 5 g of $\text{H}_2\text{BO}_3$. Make to 250-ml volume with DDI water. Store in polypropylene container.

6.9 Al standard, 200 ppm. To 50 ml of Al stock solution add 12.34 ml of HF solution and 5 g of $\text{H}_2\text{BO}_3$. Store in polypropylene container.

6.10 K standard, 1 meq L$^{-1}$. Add 12.34 ml of HF solution and 5 g of $\text{H}_2\text{BO}_3$ to 10 ml of K stock solution. Store in polypropylene container.

6.11 NaCl solution (1143 ppm Na). Dissolve 2.54 g of NaCl in DDI and make to 1-L volume.

7. Procedure

**Dilution of Sample Extracts and Standards**

7.1 Set the digital settings at 60 for the diluent (NaCl solution) and 99 for the HF sample, calibration reagent blanks, and calibration standards.

7.2 Dilute 1 part HF sample with 7 parts of NaCl solution (1:7 dilution).

7.3 Dilute 1 part calibration reagent blank (HF solution) with 7 parts NaCl solution (1:7 dilution).

7.4 Dilute 1 part of each calibration standard (200 ppm Fe, 200 ppm Al, and 1 meq$^{-1}$ K) with 7 parts of NaCl solution (1:7 dilution).

7.5 Dispense the diluted solutions into 15-mL conical polystyrene centrifuge tubes. Place tubes in carousels of the sample changer.

**AA Calibration**

7.6 Use calibration reagent blank (HF solution) and calibration standards to calibrate the AA. The AA program requires a blank and a standard, in that
order, to establish a single point calibration curve for element determination. During AA determinations, perform one calibration, i.e., blank plus standard, for every 8 samples.

**AA Operation**

7.7 The following parameters are only very general guidelines for instrument conditions for the analyte.

<table>
<thead>
<tr>
<th>Element</th>
<th>Wavelength</th>
<th>Angle</th>
<th>Fuel/Oxidant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>302.1</td>
<td>Parallel</td>
<td>C₂H₂/Air, 20/25</td>
</tr>
<tr>
<td>Al</td>
<td>308.2</td>
<td>Parallel</td>
<td>C₂H₂/N₂O, 30/17</td>
</tr>
<tr>
<td>K</td>
<td>766.5</td>
<td>30°</td>
<td>C₂H₂/Air, 20/25</td>
</tr>
</tbody>
</table>

7.8 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.9 If a sample exceeds the calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record the dilution. Remember to keep the matrix the same after dilution.

8. Calculations

Calculations are reported in method 7C3.

9. Report

Report concentrations of Fe, Al, and K by atomic absorption. Elemental concentrations are converted to percent oxides. Data are reported in method 7C3.

10. Precision

Precision data are not available for this procedure.

11. References


Iron, Aluminum, and Silicon (6C, 6G, and 6V)
Ammonium Oxalate Extraction (6C6, 6G12, and 6V2)
Inductively Coupled Plasma Spectrometry (6C9a, 6G12a, 6V2a)

Optical Density (8J) (of Ammonium Oxalate Extract)

1. Application

Oxalic acid-ammonium oxalate (acid oxalate) is used as a selective dissolution extractant for organically complexed Fe and Al, noncrystalline hydrous oxides of Fe and Al, allophane, and amorphous aluminosilicates (Wada, 1989). Acid oxalate is a poor extractant of imogolite and layer silicates and does not extract crystalline hydrous oxides of Fe and Al, opal, or crystalline silicate (Wada, 1989). A more reliable and accurate estimation of soil properties and a better understanding of soil exchange complex is provided when acid oxalate extraction is used in conjunction with other selective dissolution procedures, thermal techniques, and chemical tests. In “Soil Taxonomy,” acid oxalate extractable Fe and Al are criteria for andic soil properties (Soil Survey Staff, 1990).

2. Summary of Method

A soil sample is extracted with a mechanical vacuum extractor (Holmgren et al., 1977) in a 0.2 M acid oxalate solution buffered at pH 3.0 under darkness. The acid oxalate extract is weighed. The acid oxalate extract is diluted with 0.1 N HCl. The diluted extract is vaporized and atomized by an inductively coupled plasma emission spectrophotometer (ICP). The atoms or ions of the analyte are energized in high temperatures, resulting in the movement of valence electrons to higher orbits from the nucleus. As the electrons fall back to a lower orbit, electromagnetic energy at a specific wavelength for a given atom is emitted in measurable amounts (Soltanpour et al., 1982). Data are automatically recorded by a microcomputer and printer. The percent acid oxalate extractable Fe, Al, and Si are reported in methods 6C9a, 6G12a, and 6V2a, respectively (Soil Conservation Service, 1984). On a less routine basis, Mn is also measured. To date, however, a National Soil Survey Laboratory (NSSL) method code has not been assigned to the Mn determination by acid oxalate extraction. In method 8J, the optical density of the extract is measured with a UV spectrophotometer at 430 nm.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the ICP analyses of these elements. These interferences vary in importance, depending upon the particular analyte chosen.

The acid oxalate buffer extraction is sensitive to light, especially UV light. The exclusion of light reduces the dissolution effect of crystalline oxides and clay.
minerals. If the sample contains large amounts of amorphous material (>2% Al), an alternate method should be used, i.e., shaking with 0.275 \( M \) acid oxalate, pH 3.25, 1:100 soil:extractant.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the UV spectrophotometer and ICP.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.3 Syringes, polypropylene, disposable, 60 mL, for sample tube, extractant reservoir, and tared extraction syringe
5.4 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels
5.5 Polycons, Richards Mfg. Co.
5.6 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.7 UV-visible spectrophotometer, DU-7, Beckmann Instruments, Inc.
5.8 Cuvettes, disposable, polystyrene, 1-cm light path
5.9 Inductively coupled plasma spectrophotometer (ICP), Perkin-Elmer model 6000
5.10 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.11 Microcomputer, 7500 Professional Computer, Perkin-Elmer Corp., Norwalk, CT
5.12 Dot matrix printer, P-132, Interdigital Data Systems, Inc.
5.13 Single-stage regulator, high-purity, high-flow, argon, product no. E11-X-N145DHF, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.14 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.15 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gastight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.16 Centrifuge tubes, polystyrene, 15 mL, conical, graduated, part no. 2087, for sample dilution and sample changer, Becton Dickinson Labware, Becton Dickinson and Co., 2 Bridgewater Lane, Lincoln Park, NJ 07035

5.17 Containers, polypropylene

6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Hydrochloric acid (HCl), conc. 12 N

6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of conc. HCl to 1 part DDI water.

6.4 HCl, 1% wt. Carefully dilute 25 mL of conc. HCl to 1 L with DDI water.

6.5 HCl, 0.1 N. Add 8.33 mL of conc. HCl to DDI water and make to 1-L volume.

6.6 Acid oxalate buffer solution, 0.2 M, pH 3.0. Solution A (base): Dissolve 284 g of (NH₄)₂C₂O₄•H₂O in 10 L of DDI water. Solution B (acid): Dissolve 252 g of H₂C₂O₄•H₂O in 10 L of DDI water. Mix 4 parts solution A with 3 parts solution B. Adjust acid oxalate solution pH by adding either acid or base solution. Store in a polypropylene bottle.

6.7 pH buffers, pH 4.00 and 7.00, for electrode calibration

6.8 Primary Fe standard, 1000 ppm. Dissolve 1.000 g of Fe wire in a minimum volume of 1:1 HCl:DDI. Dilute to 1-L volume in a volumetric flask using 1% HCl. Store in a polypropylene bottle.

6.9 Primary Al standard, 1000 ppm. Dissolve 1.000 g of Al wire in a minimum volume of 1:1 HCl:DDI. Dilute to 1-L volume in a volumetric flask using 1% HCl water. Store in a polypropylene bottle.

6.10 Primary Si standard, 1000 ppm. Fuse 0.2139 g of SiO₂ with 2 g of Na₂CO₃ in a platinum crucible. Dissolve the melt with DDI water and transfer to a 100-mL volumetric flask. Dilute to 1-L volume with DDI water. Store in a polypropylene bottle.

6.11 Primary Mn standard, 1000 ppm. Dissolve 1.000 g of Mn wire in a minimum volume of 1:1 HCl:DDI. Dilute to 1-L volume in a volumetric flask using 1:1 HCl:DDI. Store in a polypropylene bottle.

6.12 High calibration standard. Mix 30 mL of each primary standard (Al, Fe, and Si) with 5 mL of primary Mn standard. Add 50 mL of 0.4 M acid oxalate solution, 20 mL of conc. HCl, and make to 1-L volume with DDI water. Resulting solution contains 5 ppm Mn and 30 ppm each of Al, Fe, and Si. Store in a polypropylene bottle.

6.13 Low calibration standard. Mix 10 mL of each primary standard (Al, Fe, and Si) with 2 mL of primary Mn standard. Add 30 mL of 0.4 M acid oxalate solution, 20 mL of conc. HCl, and make to 1-L volume with DDI water.
Resulting solution contains 2 ppm Mn and 10 ppm each of Al, Fe, and Si. Store in a polypropylene bottle.

6.14 Calibration reagent blank solution. Add 30 mL of 0.4 M acid oxalate solution, 20 mL of conc. HCl, and make to 1-L volume with DDI water.

6.15 Argon gas, purity 99.9

7. Procedure

**Extraction of Fe, Al, Si, and Mn**

7.1 Prepare sample tube by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh 0.500 g of <2-mm, air-dry soil and place in sample tube. Prepare two reagent blanks (no sample in tube) per set of 48 samples.

7.3 Place the sample tube on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.

7.4 Use a dispenser to add 15.00 mL of acid oxalate buffer to the sample tube. Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.

7.5 Set extractor for 30-min extraction rate and extract until the acid oxalate buffer solution is at a 0.5 to 1.0-cm height above sample. Turn off extractor.

7.6 Put reservoir tube on top of the sample tube.

7.7 Add 35 mL of acid oxalate buffer to the reservoir tube.

7.8 Cover the extractor with a black plastic bag to exclude light. Adjust the extraction rate for a 12-h extraction.

7.9 After the extraction, shut off the extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the sample tube.

7.10 Weigh each syringe containing acid oxalate extract to the nearest 0.01 g.

7.11 Mix extract in each syringe by manually shaking. Fill a polycon with extract solution. This solution is reserved for determinations of Fe, Mn, Al, and Si. If optical density is to be measured, fill a disposable cuvette with extract solution. Discard excess solution.

**Determination of Optical Density of Extract**

7.12 Place 4 mL of acid oxalate extract in disposable cuvette.

7.13 Place 4 mL of acid oxalate reagent blank in disposable cuvette.

7.14 On DU-7 spectrophotometer, select a 430-nm wavelength. Select normal slit width and height.
7.15 Use the acid oxalate extract reagent blank to set spectrophotometer.
7.16 Record optical density of acid oxalate extract to nearest 0.000.

**Dilution of Sample Extracts and Standards**

7.17 For better nebulization, add one drop of DDBSA solution to each tube (sample extracts, calibration standards, and reagent blanks) to reduce surface tensions. Add DDBSA to tube before the addition of diluted solution.

7.18 Set the digital settings of the Hamilton diluter at 63 for the diluent (0.1 N HCl) and 70 for the acid oxalate extracts for a 1:10 dilution. Calibration reagent blanks and calibration standards are not diluted.

7.19 Dilute 1 part acid oxalate sample extract with 10 parts of 0.1 N HCl (1:10 dilution).

7.20 Dispense the diluted solutions into 15-mL conical polystyrene centrifuge tubes which have been placed in carousels of the sample changer.

**ICP Calibration**

7.21 Use high calibration standard and calibration reagent blank to calibrate ICP. The ICP requires a standard and a blank, in that order, for calibration. During ICP determinations, perform one calibration, i.e., standard plus blank, for every 6 samples.

7.22 Use the low calibration standard as a check sample.

**ICP Set-up and Operation**

7.23 The following parameters are only very general guidelines for instrument conditions for the various analytes.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICP power</td>
<td>1250 W</td>
</tr>
<tr>
<td>Plasma gas flow</td>
<td>Ar 12 L min⁻¹</td>
</tr>
<tr>
<td>Nebulizer gas flow</td>
<td>Ar 0.5 L min⁻¹</td>
</tr>
<tr>
<td>Auxiliary gas flow</td>
<td>Ar 0.05 to 1 L min⁻¹</td>
</tr>
</tbody>
</table>

Use a high solids nebulizer instead of the cross flow nebulizer.

7.24 Analyte data for some elements are reported at 2 wavelengths which serve as data checks.
7.25 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings. The instrument readings are usually programmed in ppm.

7.26 If sample exceeds calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with 0.1 N HCl at the 1:1 ratio.

8. Calculations

Analyte (%) = \([\text{ICP} \times (\text{Syr}_{\text{fin}} - \text{Syr}_{\text{init}}) \times \text{DR} \times \text{AD/OD}] / [\text{Sample} \times 10,000 \times \text{Density}]\]

where:
- ICP = ICP analyte concentration (ppm)
- Syr\(_{\text{fin}}\) = Weight of syringe + extract (g)
- Syr\(_{\text{init}}\) = Tare weight of syringe (g)
- DR = Dilution ratio of samples over calibration range
- Sample = Weight of sample (g)
- Density = Density of acid oxalate solution (1.007)
- AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report the percent acid oxalate extractable Fe, Al, and Si to the nearest 0.01%. Percent acid oxalate extractable is also reported. To date, however, no method code has been assigned to Mn determination by acid oxalate extraction. Report the optical density of the acid oxalate extract to the nearest 0.001 unit.

10. Precision

Precision data are not available for this procedure. The mean, standard deviation, and CV for Fe, Al, Si, and optical density for both the low and high standards are as follows:

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Wave-length</th>
<th>Low Standards</th>
<th>High Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(nm)</td>
<td>(ppm)</td>
<td>(ppm)</td>
</tr>
<tr>
<td>Fe</td>
<td>238.204 / 239.562</td>
<td>30.00</td>
<td>10.00</td>
</tr>
<tr>
<td>Al</td>
<td>394.400 / 396.150</td>
<td>30.00</td>
<td>10.00</td>
</tr>
<tr>
<td>Si</td>
<td>212.412 / 251.611</td>
<td>30.00</td>
<td>10.00</td>
</tr>
<tr>
<td>Mn</td>
<td>257.610</td>
<td>5.00</td>
<td>2.00</td>
</tr>
</tbody>
</table>
### High Standard

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optical density</td>
<td>18</td>
<td>0.18</td>
<td>0.03</td>
<td>14.2</td>
</tr>
<tr>
<td>Fe</td>
<td>17</td>
<td>0.94</td>
<td>0.17</td>
<td>18.2</td>
</tr>
<tr>
<td>Al</td>
<td>17</td>
<td>2.6</td>
<td>0.19</td>
<td>7.6</td>
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<tr>
<td>Si</td>
<td>17</td>
<td>1.2</td>
<td>0.09</td>
<td>7.3</td>
</tr>
</tbody>
</table>

### Low Standard

<table>
<thead>
<tr>
<th></th>
<th>N</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optical density</td>
<td>25</td>
<td>0.06</td>
<td>0.00</td>
<td>7.5</td>
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<tr>
<td>Fe</td>
<td>24</td>
<td>0.26</td>
<td>0.03</td>
<td>9.7</td>
</tr>
<tr>
<td>Al</td>
<td>25</td>
<td>0.17</td>
<td>0.01</td>
<td>8.4</td>
</tr>
<tr>
<td>Si</td>
<td>26</td>
<td>0.02</td>
<td>0.01</td>
<td>53.2</td>
</tr>
</tbody>
</table>

### 11. References


### Manganese and Aluminum (6D and 6G)

1. **1 N KCl Extractable, Automatic Extractor (6D3 and 6G9)**

   Inductively Coupled Plasma Spectrometry (6D3 and 6G9b)

#### 1. Application

The Al extracted by 1 N KCl approximates exchangeable Al and is a measure of the “active” acidity present in soils with a 1:1 water pH <5.5. Above pH 5.5, precipitation of Al occurs during analysis. This method does not measure the acidity component of hydronium ions (H₃O⁺). If Al is present in measurable amounts, the hydronium is a minor component of the active acidity. Because the
1 N KCl extractant is an unbuffered salt and usually affects the soil pH one unit or less, the extraction is determined at or near the soil pH. The KCl extractable Al is related to the immediate lime requirement and existing CEC of the soil. The “potential” acidity is better measured by the BaCl₂-TEA method (method 6H5a) (Thomas, 1982).

2. Summary of Method

A soil sample is leached with 1 N KCl using the mechanical vacuum extractor (Holmgren et al., 1977). The leachate is weighed. The KCl extracted solution is diluted with 0.5 N HCl. The diluted extract is vaporized and atomized by an inductively coupled plasma emission spectrophotometer (ICP). The atoms or ions of the analyte are energized in high temperatures, resulting in the movement of valence electrons to higher orbits from the nucleus. As the electrons fall back to a lower orbit, electromagnetic energy at a specific wavelength for a given atom is emitted in measurable amounts (Soltanpour et al., 1982). Data are automatically recorded by a microcomputer and printer. The Mn and Al are reported in meq/100 g oven-dry soil in methods 6D3 and 6G9b (Soil Conservation Service, 1984).

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the ICP analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

The soil:extractant ratio must remain constant. A soil:extractant ratio of 1:10 (w:v) for batch procedures is most commonly used. Using a leaching technique, a 1:20 (w:v) ratio gives comparable results. If the sample size is changed, the amount of extractable Al is changed. No other significant interferences have been identified for this procedure.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Follow the manufacturer’s safety precautions when using the ICP.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Analytical filter pulp, Schleicher and Schuell, no. 289
5.3 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.4 Syringes, polypropylene, disposable, 60 mL, for sample tube, extractant reservoir, and tared extraction syringe
5.5 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in), for connecting syringe barrels
5.6 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.7 Wash bottle, 20 mL, to dispense KCl
5.8 Polycons, Richards Mfg. Co.
5.9 Inductively coupled plasma (ICP) atomic emission spectrophotometer, Perkin-Elmer model 6000
5.10 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.11 Microcomputer, 7500 Professional Computer, Perkin-Elmer Corp., Norwalk, CT
5.12 Dot matrix printer, P-132, Interdigital Data Systems, Inc.
5.13 High-purity, high-flow, single-stage regulator, argon, product no. E11-X-N145DHF, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.14 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.15 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510

6. Reagents
6.1 Distilled deionized (DDI) water
6.2 Hydrochloric acid (HCl), conc., 12 N
6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of conc. HCl to 1 part DDI water.
6.4 HCl, 1% wt. Carefully dilute 25 mL of conc. HCl to 1 L with DDI water.
6.5 HCl, 0.5 N. Add 1 part of conc. HCl to 24 parts DDI water (1:25 dilution).
6.6 Potassium chloride solution (KCl), 1.0 N. Dissolve 1342 g of KCl reagent in 16 L DD water. Allow solution to equilibrate to room temperature. Dilute to 18 L with DDI water. Use 1.0 N KCl for Al and Mn extraction.
6.7 Potassium chloride solution (KCl), 2.0 N. Dissolve 298.24 g of KCl reagent in 1.5 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 2 L with DDI water. Use 2.0 N KCl for standards.
6.8 Primary Al standard, 2248.5 ppm (250 meq L⁻¹). Dissolve 2.2485 g of Al wire in a minimum volume of 1:1 HCl:DDI. This is a very slow reaction. Dilute to 1 L in a volumetric flask using 1% HCl solution. Store in polypropylene container.
6.9 Primary Mn standard, 1000 ppm (36 meq L⁻¹). Commercial. Dissolve 1.000 g of Mn metal in a minimum volume of 1:1 HCl:DDI. When dissolved, dilute to 1 L in a volumetric flask using 1% HCl solution. Store in a polypropylene container.

6.10 Mn standard, 250 ppm (9 meq L⁻¹). Mix 25 mL of primary Mn standard (1000 ppm) with 10 mL of 1:1 HCl:DDI and dilute to 100-mL volume with DDI water. Store in a polypropylene bottle.

6.11 Calibration Al and Mn standard, 10 meq L⁻¹ Al and 5 ppm Mn. Mix 10 mL of primary Al standard (250 meq L⁻¹) with 125 mL 2.0 N KCl solution. Add 5 mL of Mn standard (250 ppm). Make to 250-mL volume with DDI water. Store in a polypropylene container.

6.12 Calibration Al and Mn check standard, 5 meq L⁻¹ Al and 2 ppm Mn. Mix 5 mL of primary Al standard (250 meq L⁻¹) with 125 mL 2.0 N KCl solution. Add 2 mL of Mn standard (250 ppm). Store in a polypropylene container.

6.13 Calibration reagent blank solution, 1.0 N KCl. Add 125 mL of 2.0 N KCl to a volumetric flask and make to 250-mL volume with DDI water. Store in a polypropylene container.

6.14 Dodecylbenzenesulfonic acid (DDBSA), tech 97%. Working stock is 0.1 M. Dilute 25 mL of 0.1 M DDBSA to 1-L volume with DDI water.

6.15 Argon gas, purity 99.9%

7. Procedure

**Extraction of Al and Mn**

7.1 Prepare sample tube by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh exactly 2.50 g of <2-mm, air-dry soil and place in sample tube. Prepare one quality control check sample per 48 samples.

7.3 Place the sample tube on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.

7.4 Use a squeeze bottle and fill sample tube to the 20-mL mark with 1.0 N KCl solution (≈10 mL). Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.

7.5 Put reservoir tube on top of the sample tube. Set extractor for fast extraction rate and extract until the KCl solution is at a 0.5- to 1.0-cm height above sample. Turn off extractor.
7.6 Add 45 mL KCl solution to reservoir tube. Set extractor for 45-min extraction.

7.7 After the extraction, shut off extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the sample tube.

7.8 Weigh each syringe containing KCl extract to the nearest 0.01 g.

7.9 Mix extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. This solution is reserved for extractable Al and Mn analyses.

**Dilution of Extracts and Standards**

7.10 For better nebulization, add one drop of DDBSA solution to KCl sample extracts, calibration reagent blanks and calibration standards to reduce surface tensions. Add DDBSA to tube before adding diluted solution.

7.11 Set the digital settings at 40 for the diluent (0.5 N HCl) and 99 for the KCl sample extracts, calibration reagent blanks, calibration standards, and calibration check standards for a 1:5 dilution as follows:

7.12 Dilute 1 part KCl sample extract with 4 parts of 0.5 N HCl (1:5 dilution).

7.13 Dilute 1 part calibration reagent blank with 4 parts of 0.5 N HCl (1:5 dilution).

7.14 Dilute 1 part calibration standard (10 meq L⁻¹ Al and 5 ppm Mn) with 4 parts of 0.5 N HCl (1:5 dilution).

7.15 Dilute 1 part calibration check standard (5 meq L⁻¹ Al and 2 ppm Mn) with 4 parts of 0.5 N HCl (1:5 dilution).

7.16 Dispense the diluted solutions into 15-mL conical polystyrene centrifuge tubes which have been placed in carousels of the sample changer.

**ICP Calibration**

7.17 Use calibration standard (10.00 meq L⁻¹ Al and 5.00 ppm Mn,) and calibration reagent blank (1.0 N KCl) to calibrate ICP. The ICP requires a standard and a blank, in that order, for calibration. During ICP determinations, perform one calibration, i.e., standard plus blank, for every 6 samples.

7.18 Use the calibration check standard (5.00 meq L⁻¹ Al and 2.00 ppm Mn) as a check sample.

7.19 The following parameters are only very general guidelines for instrument conditions for the analytes.
ICP Set-up and Operation

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Al</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma flow (ml Ar min⁻¹)</td>
<td>12.0</td>
<td>12.0</td>
</tr>
<tr>
<td>Nebulizer flow (mL Ar min⁻¹)</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Auxiliary flow (mL Ar min⁻¹)</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Viewing Height (nm)</td>
<td>15.0</td>
<td>15.0</td>
</tr>
</tbody>
</table>

**Wavelength 1**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavelength (nm)</td>
<td>394.400</td>
</tr>
<tr>
<td>Scan speed (s)</td>
<td>2.0</td>
</tr>
<tr>
<td>Bkg. Correction (nm)</td>
<td>−0.069, +0.055</td>
</tr>
</tbody>
</table>

**Wavelength 2**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavelength (nm)</td>
<td>396.152</td>
</tr>
<tr>
<td>Scan speed (s)</td>
<td>2.0</td>
</tr>
<tr>
<td>Bkg. Correction (nm)</td>
<td>−0.069, +0.055, −0.046, +0.049</td>
</tr>
</tbody>
</table>

High solids nebulizer has a 20 s read delay.

7.20 Load sample tubes in the carousel so that the calibration standard, reagent blank, calibration check standard, and 6 unknown samples are determined in order. Determine a set of 24 unknown samples with each carousel.

7.21 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.

7.22 If a sample exceeds the calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with the 0.5 N HCl solution (1:5 dilution).

7.23 Analyze one quality control check sample for every 48 samples.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L⁻¹ Al and ppm Mn) in undiluted extract. Use these values to calculate the analyte concentration on an oven-dry soil basis (meq/100 g).

\[
\text{Analyte (meq/100 g)} = \frac{[\text{ICP} \times (\text{Wt}_{\text{syr+ext}} - \text{Wt}_{\text{syr}})\times \text{DR} \times 100 \times \text{AD/OD}] / [\text{Smp. Wt.} \times 1.0412 \times 1000]}{
\]}

where:

- ICP=ICP analyte reading
- Wt_{syr+ext} = Weight of extraction syringe & extract (g)
Wt<sub>sy</sub> = Weight of tared extraction syringe (g)
DR. = Dilution ratio of samples over calibration range
Smp. Wt = Sample weight (g)
1.0412 = Density of 1 N KCl @ 20 °C
1000 = g L<sup>−1</sup>
100 = Conversion factor (100-g basis)
AD/OD = Air-dry/oven-dry ratio (method 4B5)

9. Report
Report KCl extractable Al and Mn in units of meq/100 g of oven-dry soil to the nearest 0.01 meq/100 g.

10. Precision
Precision data are not available for this procedure.

11. References

Aluminum (6G)
KCl, Automatic Extractor (6G9)
Atomic Absorption (6G9a)

1. Application
The Al extracted by 1 N KCl approximates exchangeable Al and is a measure of the “active” acidity present in soils with a 1:1 water pH < 5.5. Above pH 5.5, precipitation of Al occurs during analysis. This method does not measure the acidity component of hydronium ions (H<sub>3</sub>O<sup>+</sup>). If Al is present in measurable amounts, the hydronium is a minor component of the active acidity. Because the 1 N KCl extractant is an unbuffered salt and usually affects the soil pH one unit
or less, the extraction is determined at or near the soil pH. The KCl extractable Al is related to the immediate lime requirement and existing CEC of the soil. The “potential” acidity is better measured by the BaCl₂-TEA method (method 6H5a) (Thomas, 1982).

2. Summary of Method

A soil sample is leached with 1 N KCl using the mechanical vacuum extractor (Holmgren et al., 1977). The leachate is weighed. The KCl extract is diluted with distilled deionized (DDI) water. The diluted extract is aspirated into an atomic absorption spectrophotometer (AA). The analyte is measured by absorption of the light from a hollow cathode lamp. An automatic sample changer is used to aspirate a series of samples. The AA converts absorption to analyte concentration. The data are automatically recorded by a microcomputer and printer. The Al is reported in meq/100 g oven-dry soil in method 6G9a.

3. Interferences

There are four types of interferences (matrix, spectral, chemical, and ionization) in the AA analyses of these cations. These interferences vary in importance, depending upon the particular analyte selected.

The soil:extractant ratio must remain constant. A soil:extractant ratio of 1:10 (w:v) for batch procedures is most commonly used. Using a leaching technique, a 1:20 (w:v) ratio gives comparable results. If the sample is changed, the amount of extractable Al is changed. No other significant interferences have been identified for this procedure.

4. Safety

Wear protective clothing and eye protection. When preparing reagents, exercise special care. Restrict the use of concentrated HCl to a fume hood. Many metal salts are extremely toxic and may be fatal if ingested. Follow standard laboratory practices when handling compressed gases. Gas cylinders should be chained or bolted in an upright position. Acetylene gas is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the AA.

5. Equipment

5.1 Electronic balance, ±1-mg sensitivity
5.2 Mechanical vacuum extractor, 24 place, Centurion International, Inc., Lincoln, NE
5.3 Syringes, polypropylene, disposable, 60 mL, for sample tube, extractant reservoir, and tared extraction syringe
5.4 Rubber tubing, 3.2 ID x 6.4 OD x 25.4 mm (⅛ ID x ¼ OD x 1 in) for connecting syringe barrels
5.5 Wash bottle, 20 mL, to dispense KCl
5.6 Plunger, modified. Remove rubber and cut plastic protrusion from plunger end.
5.7 Polycons, Richards Mfg. Co.
5.8 Atomic absorption spectrophotometer (AA), Perkin-Elmer model 5000
5.9 Autosampler, AS-50, Perkin-Elmer Corp., Norwalk, CT
5.10 Microcomputer, 7500 Professional Computer, Perkin-Elmer Corp., Norwalk, CT
5.11 Dot matrix printer, P-132, Interdigital Data Systems, Inc.
5.12 Single-stage regulator, acetylene service, part number E11-0-N511A, Air Products and Chemicals, Inc., Box 538, Allentown, PA 18105
5.13 Heated regulator, single-stage, nitrous oxide, stock number 808 8039, Airco Welding Products, P.O. Box 486, Union, NJ 07083
5.14 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.15 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
5.16 Centrifuge tubes, polystyrene, 15 mL, conical bottom, graduated, part no. 2087, for sample dilution and sample changer, Becton Dickinson Labware, Becton Dickinson and Co., 2 Bridgewater Lane, Lincoln Park, NJ 07035
5.17 Containers, polypropylene or teflon

6. Reagents

6.1 Distilled deionized (DDI) water
6.2 Hydrochloric acid (HCl), conc., 12 N
6.3 HCl, 1:1 HCl:DDI, 6 N. Carefully mix 1 part of conc. HCl to 1 part DDI water.
6.4 HCl, 1% wt. Carefully dilute 25 mL of conc. HCl to 1 L with DDI water.
6.5 Potassium chloride solution (KCl), 1.0 N. Dissolve 1342 g of KCl reagent in 16 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 18 L with DDI water. Use 1.0 N KCl solution for Al extraction.
6.6 Potassium chloride solution (KCl), 2.0 N. Dissolve 298.24 g of KCl reagent in 1.5 L DDI water. Allow solution to equilibrate to room temperature. Dilute to 2 L with DDI water. Use 2.0 N KCl solution for standards.
6.7 Primary Al standard, 2248.5 ppm (250 meq L⁻¹). Dissolve 2.2485 g of Al wire in a minimum volume of 1:1 HCl:DDI. This is a very slow reaction.
Dilute to 1 L in a volumetric flask using 1% HCl solution. Store in polypropylene bottle.

6.8 Calibration Al standard, 10 meq L\(^{-1}\). Mix 10 mL of primary Al standard (250 meq L\(^{-1}\)) with 125 mL of 2.0 \(N\) KCl solution. Make to 250-mL volume with DDI water. Store in polypropylene bottle.

6.9 Calibration Al check standard, 5 meq L\(^{-1}\). Mix 5 mL of primary Al standard (250 meq L\(^{-1}\)) with 125 mL of 2.0 \(N\) KCl solution. Make to 250-mL volume with DDI water. Store in polypropylene bottle.

6.10 Calibration reagent blank solution, 1.0 \(N\) KCl. Add 125 mL of 2.0 \(N\) KCl to a volumetric flask and make to 50-mL volume with DDI water. Store in polypropylene bottle.

6.11 Nitrous oxide gas, compressed
6.12 Acetylene gas, compressed, purity 99.6%
6.13 Compressed air with water and oil traps

7. Procedure

**Extraction of Al**

7.1 Prepare sample tube by tightly compressing a 1-g ball of filter pulp into the bottom of a syringe barrel with a modified plunger.

7.2 Weigh exactly 2.50 g of <2-mm, air-dry soil and place in sample tube. Prepare one quality control check sample per 48 samples.

7.3 Place the sample tube on the upper disk of the extractor and connect a tared extraction syringe. Use 25.4-mm (1-in) length rubber tubing to insert the handle of the plunger in the slot of the stationary extractor disk.

7.4 Use a squeeze bottle and fill sample tube to the 20-mL mark with 1.0 \(N\) KCl solution (~10 mL). Make sure that the sample is thoroughly wetted. During the addition, wash sides of the tube and wet the sample. Shaking, swirling, or stirring may be required to wet organic samples. Allow sample to stand for at least 30 min.

7.5 Put reservoir tube on top of the sample tube. Set extractor for fast extraction rate and extract until the KCl solution is at a 0.5- to 1.0-cm height above sample. Turn off extractor.

7.6 Add 45 mL KCl solution to reservoir tube. Set extractor for 45-min extraction.

7.7 After the extraction, shut off extractor and pull plunger of syringe down. Do not remove the plunger from syringe barrel. Carefully remove the syringe with extract leaving the rubber tubing on the sample tube.

7.8 Weigh each syringe containing KCl extract to the nearest 0.01 g.
7.9 Mix extract in each syringe by manually shaking. Fill a polycon with extract solution and discard the excess. This solution is reserved for extractable Al analysis.

**Dilution of Sample Extracts and Standards**

7.10 No ionization suppressant is required as the K in the extractant is present in sufficient quantity. Set the digital settings at 40 for the diluent (DDI water) and 99 for the KCl sample extracts, calibration reagent blanks, calibration standards, and calibration check standards for a 1:5 dilution as follows:

7.11 Dilute 1 part KCl sample extract with 4 parts of DDI water (1:5 dilution).
7.12 Dilute 1 part calibration reagent blank with 4 parts of DDI water (1:5 dilution).
7.13 Dilute 1 part calibration standard (10 meq L\(^{-1}\) Al) with 4 parts of DDI water (1:5 dilution).
7.14 Dilute 1 part calibration check standard (5 meq L\(^{-1}\) Al) with 4 parts of DDI water (1:5 dilution).
7.15 Dispense the diluted solutions into 15-mL conical polystyrene centrifuge tubes which are placed in carousels of the sample changer.

**AA Calibration**

7.16 Use calibration reagent blank (1.0 N KCl) and calibration standard (10 meq L\(^{-1}\) Al) to calibrate the AA. The AA program requires a blank and a standard, in that order, to establish a single point calibration curve for element determination. During AA determinations, perform one calibration, i.e., blank plus standard, for every 12 samples.
7.17 Use the calibration check standard (5 meq L\(^{-1}\) Al) as a check sample.

**AA Set-up and Operation**

7.18 The following parameters are only very general guidelines for instrument conditions for the analyte.
- Element Head = Al
- Wavelength (nm) = 309.3
- Burner head & angle = 5 cm Parallel
- Fuel/Oxidant (C\(_2\)H\(_2\)/N\(_2\)O) = 30/17
- Typical read delay is 6 s, and integration by peak area is 8 s.
7.19 Use the microcomputer and printer to set instrument parameters and to collect and record instrument readings.
7.20 If a sample exceeds the calibration standard, dilute the sample (dilution ratio in calculation) with appropriate matrix and record dilution. Remember to keep the matrix the same after dilution by diluting with DDI water (1:5 dilution).

7.21 Analyze one quality control check sample for every 48 samples.

8. Calculations

8.1 The instrument readings are the analyte concentration (meq L\(^{-1}\) Al) in undiluted extract. Use these values to calculate the analyte concentration on an oven-dry soil basis (meq/100 g).

\[
\text{Al (meq/100 g)} = \frac{[\text{AA Al} \times (W_{\text{syr+ext}} - W_{\text{syr}}) \times DR \times 100 \times AD/OD]}{\text{Smp. Wt.} \times 1.0412 \times 1000}
\]

where:

- \(\text{AA Al}\) = AA Al reading (meq L\(^{-1}\))
- \(W_{\text{syr+ext}}\) = Weight of extraction syringe and extract (g)
- \(W_{\text{syr}}\) = Weight of tared extraction syringe (g)
- \(DR\) = Dilution ratio for samples over calibration range
- \(\text{Smp. Wt.}\) = Sample weight (g)
- 1.0412 = Density of 1 N KCl @ 20 °C
- 1000 = g L\(^{-1}\)
- 100 = Conversion factor (100-g basis)
- \(AD/OD\) = Air-dry/oven-dry ratio (method 4B5)

9. Report

Report KCl extractable Al in units of meq/100 g of oven-dry soil to the nearest 0.1 meq/100 g.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run with every batch of 48 samples. For 21 observations of the quality control sample, the mean, standard deviation, and C.V. for extractable Al are 3.1, 0.18, and 5.7 %, respectively.

11. References


Chloride, Sulfate, Nitrate, Fluoride, and Nitrite (6K, 6L, 6M, 6U, and 6W)
Saturation Extract (6K1, 6L1, 6M1, 6U1, and 6W1)
Chromatograph (6K1c, 6L1c, 6M1c, 6U1a, and 6W1a)

1. Application
The soluble anions that are commonly determined in saline and alkali soils are carbonate, bicarbonate, sulfate, chloride, nitrate, nitrite, fluoride, phosphate, silicate, and borate (Khym, 1974; U.S. Salinity Laboratory Staff, 1954). Carbonate and bicarbonate are determined by titration in methods 6I1b and 6J1b, respectively (Soil Conservation Service, 1984). Phosphate, silicate, and borate are not determined because they are found only occasionally in measurable amounts in soils. Chloride, sulfate, nitrate, fluoride, and nitrite are measured in solution in methods 6K1c, 6L1c, 6M1c, 6U1a, and 6W1a, respectively (Soil Conservation Service, 1984). In saline and alkali soils, carbonate, bicarbonate, sulfate, and chloride are the anions that are found in the greatest abundance. In general, soluble sulfate is usually more abundant than soluble chloride.

2. Summary of Method
The saturation extract is diluted according to its electrical conductivity (ECs). The diluted sample is injected into the ion chromatograph, and the anions are separated. A conductivity detector is used to determine the anion. A chart recording is made of the chromatograph. Standard anions are used to calibrate the system. A calibration curve is determined, and the anion concentrations are calculated. The saturated extract anions, Cl\(^-\), SO\(_4\)\(^{2-}\), NO\(_3\)\(^-\), F\(^-\), and NO\(_2\)\(^-\), are reported in meq L\(^-1\) in methods 6k1c, 6L1c, 6M1c, 6U1a, and 6W1a, respectively (Soil Conservation Service, 1984).

3. Interferences
Some saturation extracts contain suspended solids. Filtering after dilution removes the particles. Saturation extracts of acid soils that contain Fe and/or Al may precipitate and clog the separator column. Saturation extracts of very high pH may contain organic material which may clog or poison the column. Low molecular weight organic anions will co-elute with inorganic anions from the column.

4. Safety
Wear protective clothing and safety glasses. When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if
ingested. Thoroughly wash hands after handling these metal salts. Follow the manufacturer’s safety precautions when using the chromatograph.

5. Equipment

5.1 Ion chromatograph, Series 2110i, with conductivity detector, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.2 HPIC AS3 analytical column, P/N 030985, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.3 HPIC AG3 analytical guard column, P/N 030986, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.4 Anion micro membrane suppressor, P/N 037072, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.5 Automated sampler, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.6 Poly-vials, 5 mL, P/N 038008, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.7 Poly-vials, filter caps, 5 mL, P/N 038009, Dionex Corp., 1228 Titan Way, Sunnyvale, CA 94086

5.8 Chart recorder, Honeywell Corp., chart speed 0.5 cm min⁻¹, span 1000 mV F.S.

5.9 Digital diluter/dispenser, product number 100004, with hand probe and actuator, product number 230700, Hamilton Co., P.O. Box 10030, Reno, NV 89510

5.10 Syringes, gas tight, Hamilton 1001 DX and 1010-TEF LL, Hamilton Co., P.O. Box 10030, Reno, NV 89510

5.11 Syringes, disposable, polypropylene, 12 mL

5.12 Disposable 0.2-µm pore size, 25-mm filter assembly, Gelman Sciences, Inc., 674 South Wagner Road, Ann Arbor, MI 48106. Use for saturation extracts and standards.

5.13 Disposable 0.2-µm pore size, Ultipor N₆₀ DFA3001NAEY, Pall Trinity Micro Corp., Cortland, NY 13045. Use for filtering distilled deionized (DDI) water.

6. Reagents

6.1 Distilled deionized (DDI) filtered water

6.2 Sulfuric acid (H₂SO₄), conc., reagent

6.3 Toluene

6.4 Isopropanol to de-gas column
6.5 Regenerant solution for membrane suppressor columns, 0.025 \( N \) \( \text{H}_2\text{SO}_4 \). Carefully mix 22.92 g of conc. \( \text{H}_2\text{SO}_4 \) with filtered DDI water and dilute to 18-L volume. Store in a clean glass carboy with a solid stopper. Cover the carboy top with aluminum foil to protect the contents from dust.

6.6 Stock \( \text{NaHCO}_3 \) solution, 0.480 \( M \). Mix 40.34 g of dried \( \text{NaHCO}_3 \) with filtered DDI water and dilute to 1-L volume.

6.7 Stock \( \text{Na}_2\text{CO}_3 \) solution, 0.3838 \( M \). Mix 40.68 g of dried \( \text{Na}_2\text{CO}_3 \) with filtered DDI water and dilute to 1-L volume.

6.8 Working eluent solution. Mix 112.5 mL of 0.480 \( M \) \( \text{NaHCO}_3 \) and 112.5 mL of 0.3838 \( M \) \( \text{Na}_2\text{CO}_3 \) with filtered DDI water and dilute to 18-L volume. Add 3 drops of toluene to retard microbial growth.

6.9 Primary \( \text{SO}_4^{2-} \) standard, 0.5 \( M \) (1.0 \( N \)). Mix 17.7560 g of \( \text{Na}_2\text{SO}_4 \) with filtered DDI water and dilute to 250-mL volume.

6.10 Primary \( \text{Cl}^- \) standard, 1.0 \( M \) (1.0 \( N \)). Add 18.6392 g of KCl with filtered DDI water and dilute to 250-mL volume.

6.11 Primary \( \text{F}^- \) standard, 0.125 \( M \) (0.125 \( N \)). Add 1.3122 g of NaF with filtered DDI water and dilute to 250-mL volume.

6.12 Primary \( \text{NO}_3^- \) standard, 1.0 \( M \) (1.0 \( N \)). Add 25.2770 g of \( \text{KNO}_3 \) with filtered DDI water and dilute to 250-mL volume.

6.13 Primary mixed standard. Prepare 1 primary mixed standard by taking aliquots of each of the proceeding primary standards and diluting the combined aliquots to a 1-L volume with working eluent as follows:

<table>
<thead>
<tr>
<th>Primary Stds.</th>
<th>Aliquot</th>
<th>Final</th>
<th>Conc. Vol. w/Eluent</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Na}_2\text{SO}_4 )</td>
<td>50</td>
<td>1000</td>
<td>50</td>
</tr>
<tr>
<td>KCl</td>
<td>10</td>
<td>1000</td>
<td>10</td>
</tr>
<tr>
<td>NaF</td>
<td>100</td>
<td>1000</td>
<td>12.5</td>
</tr>
<tr>
<td>KNO(_3)</td>
<td>30</td>
<td>1000</td>
<td>30</td>
</tr>
</tbody>
</table>

Add eight drops of toluene to primary mixed standard to retard microbial growth and store in a glass container.

6.14 Mixed calibration standards. Prepare 4 mixed calibration standards (0.5, 1.0, 3.0, and 7.0 readings) by taking aliquots of primary mixed standard and diluting each aliquot to 100-mL volume with working eluent as follows:
### Table 7.1. Anion Standards

<table>
<thead>
<tr>
<th>Primary Mixed Stds.</th>
<th>Final Vol. w/Eluent</th>
<th>SO$_4^{2-}$ (meq L$^{-1}$)</th>
<th>Cl$^-$ (meq L$^{-1}$)</th>
<th>F$^-$ (meq L$^{-1}$)</th>
<th>NO$_3^-$ (meq L$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(mL)</td>
<td>(mL)</td>
<td>(meq L$^{-1}$)</td>
<td>(meq L$^{-1}$)</td>
<td>(meq L$^{-1}$)</td>
<td>(meq L$^{-1}$)</td>
</tr>
<tr>
<td>0.5</td>
<td>100</td>
<td>0.25</td>
<td>0.05</td>
<td>0.0625</td>
<td>0.15</td>
</tr>
<tr>
<td>1.0</td>
<td>100</td>
<td>0.50</td>
<td>0.10</td>
<td>0.125</td>
<td>0.30</td>
</tr>
<tr>
<td>3.0</td>
<td>100</td>
<td>1.5</td>
<td>0.30</td>
<td>0.375</td>
<td>0.90</td>
</tr>
<tr>
<td>7.0</td>
<td>100</td>
<td>3.5</td>
<td>0.70</td>
<td>0.875</td>
<td>2.1</td>
</tr>
</tbody>
</table>

#### 6.15 NaNO$_2$, Baker reagent grade, 99.5% purity

#### 6.16 Primary NO$_2^-$ standard, 1 N (1000 meq L$^{-1}$). Mix 69.3568 g of reagent grade NaNO$_2$ with filtered DDI water and dilute to 1-L volume. Take 5 mL aliquot of primary NO$_2^-$ standard and dilute with 500 mL of filtered DDI water (10 meq L$^{-1}$). Add eight drops of toluene to primary NO$_2^-$ standard to retard microbial growth and store in a glass container.

#### 6.17 NO$_2^-$ calibration standards. Prepare 4 NO$_2^-$ calibration standards (0.5, 1.0, 3.0, and 7.0 readings) by taking aliquots of primary NO$_2^-$ standard (10 meq L$^{-1}$) and diluting each aliquot to 100-mL volume with working eluent as follows:

### Table 7.1. NO$_2^-$ Standards

<table>
<thead>
<tr>
<th>Primary Mixed Stds.</th>
<th>Final Vol. w/Eluent</th>
<th>NO$_2^-$ (meq L$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(mL)</td>
<td>(mL)</td>
<td>(meq L$^{-1}$)</td>
</tr>
<tr>
<td>0.5</td>
<td>100</td>
<td>0.5</td>
</tr>
<tr>
<td>1.0</td>
<td>100</td>
<td>1.0</td>
</tr>
<tr>
<td>3.0</td>
<td>100</td>
<td>3.0</td>
</tr>
<tr>
<td>7.0</td>
<td>100</td>
<td>7.0</td>
</tr>
</tbody>
</table>

#### 7. Procedure

**Dilution of extracts**

#### 7.1 To estimate the total soluble anion concentration (meq L$^{-1}$), multiply the EC$_s$ (method 8A3a) by 10. Subtract the CO$_3^{2-}$ and HCO$_3^-$ concentrations (methods 6I1b and 6J1b) from the total anion concentration. The remainder is the ≈ concentration (meq L$^{-1}$) of anions to be separated by ion chromatography.

Anion concentration (meq L$^{-1}$) = EC$_s$ x 10 – (HCO$_3^-$ + CO$_3^{2-}$)

#### 7.2 Dilute the saturation extract with the working eluent. Some typical dilutions are as follows:
### EC<sub>s</sub> Dilution Factor

<table>
<thead>
<tr>
<th>(mmhos cm&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>Dilution Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0 to 0.4</td>
<td>1:3</td>
</tr>
<tr>
<td>0.4 to 0.7</td>
<td>1:5</td>
</tr>
<tr>
<td>0.8 to 1.2</td>
<td>1:9</td>
</tr>
<tr>
<td>1.2 to 1.8</td>
<td>1:17</td>
</tr>
<tr>
<td>1.8 to 2.9</td>
<td>1:39</td>
</tr>
<tr>
<td>3.0 to 5.5</td>
<td>1:80</td>
</tr>
<tr>
<td>5.5 to 7.5</td>
<td>1:150</td>
</tr>
<tr>
<td>7.5 to 9.7</td>
<td>1:200</td>
</tr>
<tr>
<td>9.7 to 13.5</td>
<td>1:290</td>
</tr>
<tr>
<td>13.5 to 15.5</td>
<td>1:350</td>
</tr>
<tr>
<td>15.5 to 25.0</td>
<td>1:660</td>
</tr>
<tr>
<td>25.0 to 40.0</td>
<td>1:1100</td>
</tr>
<tr>
<td>40.0 to 55.0</td>
<td>1:2100</td>
</tr>
<tr>
<td>55.0 to 75.0</td>
<td>1:4800</td>
</tr>
<tr>
<td>+75.0</td>
<td>1:15,500</td>
</tr>
</tbody>
</table>

### 7.3
Place the diluted samples in 12-mL syringes. Cap syringes to prevent evaporation or contamination.

### 7.4
Place the mixed calibration standards in 12-mL syringes.

### Set-up and Operation of Ion Chromatograph (IC)

### 7.5
Because any number of factors may cause a change in IC operating conditions, only a general set-up of the Dionex 2110i ion chromatograph is presented. Individual analysts may modify some or all of the operating conditions to achieve satisfactory results. The μS cm<sup>-1</sup> units are equivalent to mmhos cm<sup>-1</sup>. Typical operation parameters are as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conductivity cell range</td>
<td>3 μS cm&lt;sup&gt;-1&lt;/sup&gt; full scale to 100 μS cm&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
<tr>
<td>Auto offset</td>
<td>&quot;On&quot;</td>
</tr>
<tr>
<td>Analytical pump flow rate</td>
<td>2.0 to 2.5 mL min&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
<tr>
<td>Low pressure limit</td>
<td>200</td>
</tr>
<tr>
<td>Parameter</td>
<td>Range</td>
</tr>
<tr>
<td>--------------------------</td>
<td>----------------------------</td>
</tr>
<tr>
<td>High pressure limit</td>
<td>1000</td>
</tr>
<tr>
<td>Regenerant flow</td>
<td>3 to 4 mL min⁻¹</td>
</tr>
<tr>
<td>Injector loom</td>
<td>0.50 mL</td>
</tr>
<tr>
<td>Air pressure</td>
<td>3 to 6 psi</td>
</tr>
<tr>
<td>Chart recorder speed</td>
<td>0.5 cm min⁻¹</td>
</tr>
<tr>
<td>Chart recorder span</td>
<td>1000 mV full scale</td>
</tr>
</tbody>
</table>

7.6 Initial IC operation should be long enough to establish a stable baseline.
7.7 Inject the most concentrated standard. The IC adjustment may be necessary to obtain adequate stability, resolution, and reproducibility.
7.8 Inject standards in random order to detect if memory effects are evident.
7.9 Analyze blanks at frequent intervals.
7.10 The injection loop requires complete flushing, i.e., 3 to 5x the loop volume.
7.11 Inject samples, standards, and blanks in the IC after achievement of stability. The analyst may change the detector range to suit the sample.
7.12 The analyst records the detector range and peak height for each detected anion. The anion identity may be determined by comparison to standards. Peak height is determined from the baseline to the peak.

8. Calculations

**Calibration Calculations**

8.1 Use the peak height of each anion standard to either construct a calibrated curve to plot anion concentration or use a least squares analysis to calculate anion concentration. The analytes are reported in meq L⁻¹.

8.2 *Calibration Curve:* Plot the peak height against the meq L⁻¹ of each anion standard on graph paper. Construct the calibration curve by finding the “best” line that fits the plotted standards.

8.3 *Linear Squares Analysis:* Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. An example for the anion Cl⁻ is as follows:

\[ \text{Cl}^- \text{ (meq L}^{-1}\text{)} = Y = 0.1 \times 1.5 = 4.0 \]

Peak height = \(X = 8.43 \times 170.0 = 441.5\)

Number of standards = \(n = 3\)


\[ \sum Y_i = 5.6 \quad \sum X_i = 619.93 \]

\[ \frac{\sum Y_i}{n} = Y = 1.866 \quad \frac{\sum X_i}{n} = X = 206.6433 \]

\[ \sum X_i Y_i = 2021.843 \quad \sum X_i^2 = 223893.31 \]

\[ \sum X_i \sum Y_i = 3471.608 \]

\[
\begin{align*}
b &= \frac{\sum X_i Y_i - \sum X_i \sum Y_i/n}{\sum X_i^2 - (\sum X_i)^2/n} \\
&= \frac{2021.843 - 1157.2027}{223893.31 - 128104.4} = 0.0090265
\end{align*}
\]

\( b = \text{slope of the line, i.e., the amount that } Y \text{ changes when } X \text{ changes by 1 unit.} \)

The equation is as follows:

\[ Y = \bar{Y} + b (X - \bar{X}) \]

\[ Y = 1.866 + 0.0090265 (X) - 1.8653 \]

**Analyte Calculation**

**8.4 Calibration curve:** Read the analyte concentration (meq L\(^{-1}\)) directly from the calibration curve.

**8.5 Linear regression:** Put the peak height in the preceding equation and solve for analyte concentration (meq L\(^{-1}\)). Thus, if sample extract has 204 peak height, the preceding equation is as follows:

\[ Y = 1.866 + 0.0090265 (204) - 1.8653 = 1.84 \text{ meq L}^{-1} \]

**8.6 Repeat the calibration set and analyte calculation for each anion.**

**9. Report**

Report the saturation extract anions in units of meq L\(^{-1}\) to the nearest 0.1 meq L\(^{-1}\).

**10. Precision**

Precision data are not available for this procedure.

**11. References**


Total Sulfur (6R)
SO$_2$ Evolution, Infrared (6R3)
LECO SC-132 Sulfur Analyzer (6R3b)

1. Application
Organic and inorganic S forms are found in soils, with the organic S fraction accounting for >95% of the total S in most soils from humid and semi-humid (Tabatabai, 1982). Mineralization of organic S and its conversion to sulfate by chemical and biological activity may serve as a source of plant available S. Total S typically ranges from 0.01 to 0.05% in most mineral soils. In organic soils, total S may be >0.05%.

In well-drained, well-aerated soils, most of the inorganic S normally occurs as sulfate. In marine tidal flats, other anaerobic marine sediments, and mine spoils, there are usually large amounts of reduced S compounds which oxidize to sulfuric acid upon exposure to the air. In arid regions, significant amounts of inorganic S are found as sulfates such as gypsum and barite.

The typical use of total S is as an index of the total reserves of this element, which may be converted to plant available S. The SSL uses the combustion technique (LECO sulfur analyzer) for analysis of total S (method 6R3b).

Extractable sulfate S (SO$_4^{2-}$-S) is an index of readily plant-available S. Reagents that have been used for measuring SO$_4^{2-}$-S include water, hot water, ammonium acetate, sodium carbonate and other carbonates, ammonium chloride and other chlorides, potassium phosphate and other phosphates, and ammonium fluoride (Bray-1). Extractable SO$_4^{2-}$-S does not include the labile fraction of soil organic S that is mineralized during the growing season (Tabatabai, 1982). Extraction reagents for organic S include hydrogen peroxide, sodium bicarbonate, sodium hydroxide, sodium oxalate, sodium peroxide, and sodium pyrophosphate. There are other methods available for determination of soil S, especially for total S and SO$_4^{2-}$-S. The investigator may refer to the review by Beaton et al. (1968).

2. Summary of Method
A fine-ground (<80-mesh) soil sample is oxidized at high temperature. The gases released are scrubbed, and the SO$_2$ in the combustion gases are measured using an infrared detector. Percent S is reported on an oven-dry soil basis.

3. Interferences
No significant interferences are known to affect the oxidizable S measurement.
4. Safety
Wear protective clothing and safety glasses. Magnesium perchlorate may form explosive mixtures. Magnesium perchlorate may contain traces of perchloric acid, which remain from manufacturer’s operations. This acid is anhydrous because of the strong desiccating capability of the salt. Avoid prolonged contact with oxidizable material or material capable of forming unstable perchlorate esters or salts. Remove magnesium perchlorate by using an excess of water to thoroughly dilute the material.

The use of high temperatures in the oxidation of samples requires that extreme caution be used to prevent burns and fires. Follow standard laboratory procedures when handling compressed gases. Oxygen is highly flammable. Avoid open flames and sparks. Standard laboratory equipment includes fire blankets and extinguishers for use when necessary. Follow the manufacturer’s safety precautions when using the sulfur analyzer.

5. Equipment
5.2 Data transmit card, part no. 772-573, Leco Corp., St. Joseph, MI
5.3 Combustion boats, part no. 529-203, Leco Corp., St. Joseph, MI
5.5 Electronic balance, ±1-mg sensitivity

6. Reagents
6.1 Anhydrone, anhydrous magnesium perchlorate, granular
6.2 Glass wool
6.3 Compressed oxygen, >99.5% @ 30 psi

7. Procedure
7.1 Weigh an air-dry, fine-ground (<80-mesh) soil sample in a tared combustion boat. Sample size depends upon S content. The product of sample weight in g multiplied by the S percent must not be >2. In most cases, the sample size is 1.00 g, unless the S content is >2%

7.2 Refer to the manufacturer’s manual for operation of sulfur analyzer. An overview of the sulfur analyzer is as follows:
   a. Samples are combusted in an O₂ atmosphere in which the S is oxidized to SO₂.
   b. Moisture and dust are removed, and the SO₂ gas is then measured by a solid state infrared detector.
c. The microprocessor formulates the analysis results. The control console displays and prints results by combining the outputs of the infrared detector and system ambient sensors with pre-programmed calibration, linearization, and weight compensation factors.

8. Calculations

\[ S(\%) = S_i \times \frac{AD}{OD} \]

where:

- \( S(\%) \) = \( S(\%) \) on oven-dry basis
- \( S_i \) = \( S(\%) \) instrument
- \( \frac{AD}{OD} \) = air-dry/oven-dry ratio (method 4B5)

9. Report

Report total S as a percentage of oven-dry weight to the nearest 0.1%.

10. Precision

Precision data are not available for this procedure. A quality control check sample is run in every batch of 12 samples. A blank (crucible only) and a rerun of one of the 12 samples (unknowns) also are run in every batch. For 27 observations of the quality control check sample, the mean, standard deviation, and C.V. for total S are 0.57, 0.02, and 4.3%, respectively.

11. References


Phosphorus (6S)
New Zealand P Retention (6S4)

1. Application

In Soil Taxonomy, the P retention of soil material is a criterion for andic soil properties (Soil Survey Staff, 1990). Andisols and other soils that contain large amounts of allophane and other amorphous minerals have capacities for binding P (Gebhardt and Coleman, 1984). The factors that affect soil P retention are not well understood. However, allophane and imogolite have been considered...
as major materials that contribute to P retention in Andisols (Wada, 1985). Phosphate retention is also called P absorption, sorption, or fixation.

2. Summary of Method
   A 5-g soil sample is shaken in a 1000-ppm P solution for 24 h. The mixture is centrifuged at 2000 rpm for 15 min. An aliquot of the supernatant is transferred to a colorimetric tube to which nitric vanadomolybdate acid reagent (NVAR) is added. The percent transmittance of the solution is read using a colorimeter. The New Zealand P retention is reported as percent P retained.

3. Interferences
   No significant problems are known to affect the P retention measurement.

4. Safety
   Wear protective clothing (coats, aprons, sleeve guards, and gloves) and eye protection (face shields, goggles, or safety glasses). When preparing reagents, exercise special care. Many metal salts are extremely toxic and may be fatal if ingested. Thoroughly wash hands after handling these metal salts. Restrict the use of concentrated HNO₃ to a fume hood. Use safety showers and eyewash stations to dilute spilled acids and bases. Use sodium bicarbonate and water to neutralize and dilute spilled acids.

5. Equipment

   5.1 Electronic balance, ±0.01-g sensitivity
   5.2 Shaker, Eberbach 6000 power unit, reciprocating speed of 60 to 260 epm, with 6040 utility box carrier and 6110 floor stand, Eberbach Corp., Ann Arbor, MI
   5.3 Digital diluter/dispenser, product no. 100004, with hand probe and actuator, product no. 230700, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
   5.4 Syringes, 10,000 and 1000 µL, 1001 DX and 1010-TEL LL gas tight, Hamilton Co., P.O. Box 10030, Reno, NV, 89510
   5.5 Diluter/dispenser, 25 mL
   5.6 Calorimeter, Bausch and Lomb
   5.7 Calorimeter tubes, glass, 10 mL, 1-cm light path, Bausch and Lomb
   5.8 Centrifuge, International no. 2, Model V, with no. 250 A head, International Equip. Co., Boston, MA
   5.9 Trunions, International no. 320, International Equip. Co., Boston, MA
   5.10 Centrifuge tubes, 50 mL, Oak-Ridge, polyallomer, Nalgene 3119, Nalge Co., Box 20365, Rochester, NY 14602.
   5.11 Plastic cups, 2 fl. oz.
   5.12 Pipets, volumetric, class A, glass, various sizes of 1 to 20 mL
6. Reagents

6.1 Distilled deionized (DDI) water

6.2 Nitric acid (HNO₃), conc.

6.3 P retention solution, 1000 ppm P. Dissolve 8.80 g of KH₂PO₄ and 32.8 g of sodium acetate (CH₃COONa) in DDI water. Add 23 mL of glacial acetic acid. Dilute to 2 L with DDI water in a volumetric flask. The solution pH should range between 4.55 and 4.65.

6.4 Molybdate solution. Dissolve 16 g of ammonium molybdate (VI) [(NH₄)₆MO₇O₂₄•4H₂O] in 50 °C DDI water. Allow the solution to cool to room temperature and dilute to 1 L with DDI water.

6.5 Nitric acid solution. Carefully and slowly dilute 100 mL of conc. HNO₃ to 1 L of DDI water. Add the acid to the water.

6.6 Nitric vanadomolybdate acid reagent (NVAR), vanadate solution. Dissolve 0.8 g of NH₄VO₃ in 500 mL of boiling DDI water. Allow the solution to cool to room temperature. Carefully and slowly add 6 mL of conc. HNO₃. Dilute to 1 L with DDI water. Mix the nitric acid solution with the vanadate solution and then add the molybdate solution. Mix well.

6.7 Stock P standard solution (SPSS), 4000 ppm P. Dissolve 17.6 g K₂HPO₄ in DDI water. Dilute to 1 L with DDI water.

6.8 Standard P calibration P solutions (SPCS), 100, 80, 60, 40, 20, and 0% P retained. Dilute the SPSS with a solution that contains 32.8 g of sodium acetate (CH₃COONa) and 23 mL of glacial acetic acid diluted to 2 L with DDI water as follows: 100% = DDI water (0 ppm); 80% = 1:20 (200 ppm); 60% = 1:10 (400 ppm); 40% = 3:20 (600 ppm); 20% = 1:5 (800 ppm); and 0% = 1:4 (1000 ppm). The percent amount refers to percent P retention.

7. Procedure

7.1 Weigh 5.00 g of air-dry soil into a 50-mL centrifuge tube.

7.2 Use the dispenser to add 25.0 mL of P-retention solution to centrifuge tube.

7.3 Cap centrifuge tube and place in shaker and shake for 24 h at room temperature (20 °C).

7.4 Add 2 to 3 drops of Superfloc, 0.02% w/v to each tube.

7.5 Centrifuge sample at 2000 rpm for 15 min.

7.6 Pour sample supernatant into plastic cup.

7.7 Use the digital diluter to add the nitric vanadomolybdate acid reagent (NVAR) to each sample supernatant and to each SPCS. To fill a 10-mL Calorimeter tube, the diluter setting is 66 for diluent (NVAR) and 35 for sample (1:20 dilution).
7.8 The color reaction requires a minimum of 30 min before the analyst records readings.

7.9 Set the Calorimeter (blue bulb) to read at 466 nm. Set the zero against DDI water (blank). A blank has all reagents contained in the sample extract except the soil.

7.10 Record the percent transmittance to the nearest 0.01 unit for the sample extract and each SPCS.

8. Calculations

8.1 Transmittance of a solution is the fraction of incident radiation transmitted by the solution, i.e., \( T = \frac{P}{P_o} \), and is often expressed as a percentage, i.e., \( T = \frac{P}{P_o} \times 100 \). The absorbance of a solution is directly proportional to concentration and is defined by the equation, \( A = -\log_{10} T \). These relationships are derived from Beer's law.

**Calibration Calculations**

8.2 Use the transmittance of each SPCS to either construct a calibrated curve to plot \( P \) or use a least squares analysis to calculate \( P \). The \( P \) is reported in percent retained.

8.3 *Calibration Curve*: Plot the transmittances against the ppm \( P \) of each SPCS on semilog graph paper or convert to absorbances and plot on linear graph paper. Construct the calibration curve by finding the “best” line that fits the plotted SPCS.

8.4 *Least Squares Analysis*: Use a least squares criterion, i.e. best moving average. Refer to a statistical analysis book for additional information on least squares analysis. To facilitate data manipulation in a least squares analysis, the following standard curve is developed using the concentration of SPCS as a \( f[\ln(\%T)] \). Final calculated analyte concentration with either \( \log_{10} \) or \( \ln \) base would be the same. Refer to method 6S3b for an example of least squares analysis.

**Analyte Calculation**

8.5 *Calibration Curve*: Read the percent \( P \) directly from the calibration curve.

8.6 *Least Squares Analysis*: Refer to method 6S3 for an example of least squares analysis.

9. Report

Report the percent New Zealand P retention to the nearest whole number.

10. Precision

Precision data are not available for this procedure.
11. References


MINERALOGY (7)

Instrumental Analyses (7A)
Thermal Gravimetric Analysis (7A4)
Perkin-Elmer 7 Series (7A4b)

1. Application

Thermal analysis defines a group of analyses that determine some physical parameter, e.g., energy, weight, or evolved substances, as a dynamic function of temperature (Tan et al., 1986). Thermogravimetric analysis (TGA) is a technique for determining weight loss of a sample as it is being heated at a controlled rate. The weight changes are recorded as a function of temperature, i.e., a thermogravimetric curve, and provide quantitative information about substances under investigation, e.g., gibbsite (Al(OH)₃), kaolinite (Al₂Si₂O₅(OH)₄), and 2:1 expandable minerals (smectite and vermiculite).

2. Summary of Method

A 5- to 10-mg sample of soil clay is weighed into a platinum sample pan and placed in the TGA balance. The instrument records the initial sample weight. The analyst zeros the balance. The sample is then heated from a temperature of 30 to 900 °C at a rate of 20 °C min⁻¹ in a flowing N₂ atmosphere. The computer collects weight changes as a function of temperature and records a thermogravimetric curve. Gibbsite and kaolinite are quantified by noting the weight loss between 250 to 350 °C and 450 to 550 °C, respectively, and then relating these data to the theoretical weight loss of pure gibbsite or kaolinite (Soil Conservation Service, 1984). The weight loss is due to dehydroxylation, i.e., loss of crystal lattice water. Though not presently performed by the National Soil Survey Laboratory (NSSL), quantification of the 2:1 expandable minerals (smectite + vermiculite) is related to weight loss at <250 °C, i.e., loss of adsorbed water (Karathanasis and Hajek, 1982; Tan et al., 1986). At this low temperature, adsorbed water is proportional...
to the specific area of the sample (Jackson, 1956; Karathanasis and Hajek, 1982; Mackenzie, 1970; Tan and Hajek, 1977).

3. Interferences

Organic matter is objectionable because it has a weight loss by dehydrogenation and by oxidation to CO$_2$ between 300 to 900 °C (Tan, et al., 1986). Analysis in an inert N$_2$ atmosphere alleviates this problem. Mineral salts that contain water of crystallization also may be interferences. Samples should be washed free of any soluble salts.

A representative soil sample is important as sample size is small (<10 mg). Avoid large aggregates in sample, the presence of which may cause thermal interferences, i.e., differential kinetics of gas diffusion through the sample and physical movement of sample in a reaction.

In general, the same reactions that interfere with DSC/DTA also interfere with TGA determinations of kaolinite, gibbsite, and 2:1 expandable minerals. However, TGA is more sensitive to small water losses at slow rates, whereas DSC/DTA is more sensitive to large water losses at rapid rates (Tan, et al., 1986). This sensitivity difference may help to explain why kaolinite and gibbsite quantifications in TGA vs. DSC/DTA often are not equivalent, i.e., TGA estimates tend to be greater than the corresponding DSC/DTA estimates. In TGA, there is a greater probability of measuring water losses in specific temperature regimes that are not specifically associated with dehydroxylation reactions of interest. This problem is particularly apparent with illitic samples, which characteristically contain more "structural" water than ideal structural formulae would indicate (Rouston, et al., 1972; Weaver and Pollard, 1973).

Even though it is well established that various minerals lose the major portion of their crystal lattice water at different temperature ranges (Tan et al., 1986), there are overlaps in these weight loss regions (WLR) of minerals which interfere in the identification and measurement of the minerals of interest. The goethite WLR (250 to 400 °C) overlaps the gibbsite WLR (250 to 350 °C) (Mackenzie and Berggen, 1970). The illite WLR (550 to 600 °C) overlaps the high end of the kaolinite WLR (450 to 550 °C) (Mackenzie and Caillere, 1975). The WLR of hydroxy-Al interlayers in hydroxy-Al interlayered vermiculite (HIV) (400 to 450 °C) overlaps the low end of the kaolinite WLR (450 to 550 °C), especially in the poorly crystalline kaolinites (Mackenzie and Caillere, 1975). Similarly, the dehydroxylation of nontronites, Fe-rich dioctahedral smectites, (450 to 500 °C) may interfere with kaolinite identification and measurement (Mackenzie and Caillere, 1975).

4. Safety

Secure high pressure N$_2$ tanks and handle with care. When changing the tanks, protect valves with covers. Do not program the analyzer for >950 °C.
because it may present a safety hazard during sample analysis and cleaning cycles. Do not heat aluminum sample pans >600 °C. Aluminum melts at 660 °C, and the pans alloy with and destroy the sample holders. Always use high quality purge gases with the TGA. Minimum purity of 99.9% is recommended.

5. Equipment
5.1 Thermal analysis system, Perkin-Elmer 7 series, 7500 computer, TAC7 instrument controllers
5.2 Thermogravimetric analyzer module, TGA7, Hewlett-Packard digital plotter
5.3 Pressure tanks (2), N₂, purity 99.99%
5.4 Two-stage gas regulators (2), 50 psi outlet pressure
5.5 One-stage gas regulator for compressed air
5.6 Electronic balance, ±0.1-mg sensitivity, Mettler AE160
5.7 Forceps, flat-tipped
5.8 Weighing spatula
5.9 Desiccator, glass
5.10 Mortar and pestle
5.11 Sieves, 100 mesh or 80 mesh
5.13 Gibbsite, standard, Surinam Gibbsite, National Soil Survey Laboratory (NSSL), 67L022.

6. Reagents
6.1 Magnesium nitrate saturated solution [Mg(NO₃)₂•6H₂O]
6.2 Ethanol

7. Procedure

Derive <2µm Clay Fractions

7.1 Prepare Na-saturated clay as in method 7A2i, preparation of clay suspension, 7.8 to 7.19.
7.2 Dry the clay suspension and transfer to mortar. Moisten sample with ethanol and grind with pestle to make a homogeneous slurry.
7.3 Air-dry sample using flowing air in hood. Lightly grind sample with pestle to make a homogeneous powder.
7.4 Sieve sample with 80-mesh screen. Equilibrate sample overnight over a saturated magnesium nitrate solution (55% rh) in a glass desiccator.
TGA Operation

7.5 Set-up the instrument and calibrate.
7.6 Turn on the N₂ purge gases and set to 6 and 3.5 psi for balance and sample purge, respectively. The balance purge pressure should always be greater than the sample purge pressure.
7.7 Turn on compressed air and set to 25 psi.
7.8 Place the platinum sample pan in the balance stirrup. Use the computer to raise the furnace tube and to zero the balance. Lower the furnace tube.
7.9 Remove the sample pan from the stirrup. Weigh ≈5 mg of sample, i.e., <100-mesh whole-soil or derived <2-µm clay fraction, into tared sample pan. Refer to section on derived <2-µm clay fractions, 7.1 to 7.4.
7.10 Use flat-tipped forceps to remove the sample pan from the analytical balance. Tap the sample pan against a hard surface several times to uniformly distribute the sample.
7.11 Carefully place sample pan in the stirrup of the TGA microbalance.
7.12 The standard sample run heating program has a heating rate of 20 °C min⁻¹, a starting temperature of 30 °C, and an ending temperature of 900 °C.
7.13 Raise the furnace tube and allow it to seat. Press “Read Weight” key (usually twice) until a relative weight percentage of 100.0% is displayed. The computer then reads the weight.
7.14 Immediately start the “Run” program.
7.15 At the end of the sample run (≈45 min), remove the sample pan from the microbalance stirrup. The furnace tube is lowered automatically at the end of run.
7.16 To store data, enter the appropriate file name on the computer for the completed run. If data are not stored by appropriate file name, data are stored under a default file name of “gsav”. Only four of these files can be saved at any one time, after which files are overwritten. Once a file is named, it cannot be changed.

8. Calculations

8.1 The thermogravimetric curve is displayed on the computer monitor. The ordinate (Y) is expressed in a relative weight percentage, i.e., the initial sample weight is 100.0%. Use the computer to calculate the total change in sample weight (ΔY), within the predetermined temperature range, as a sample weight percent.

\[% \text{ Kaolinite} = \left( \frac{\Delta \text{sample weight }%}{450-550°C} \right) / 14 \times 100\]
where:
Δ sample weight = total Δ in sample weight expressed as relative percent
14 = percent weight of hydroxyl water lost from pure kaolinite

% Gibbsite = \([\Delta \text{ sample weight } \%_{250-350{\degree}C} / 34.6] \times 100\)

where:
Δ sample weight = total Δ in sample weight expressed as relative percent
34.6 = percent weight of hydroxyl water lost from pure gibbsite

The percent weights of hydroxyl water lost from kaolinite and gibbsite are derived from the following assumed dehydroxylation reactions.

\[ \text{Si}_2\text{Al}_2\text{O}_5(\text{OH})_4 \rightarrow 2\text{SiO}_2 + \text{Al}_2\text{O}_3 + 2\text{H}_2\text{O} \]  
(kaolinite)

\[ 2\text{Al(OH)}_3 \rightarrow \text{Al}_2\text{O}_3 + 3\text{H}_2\text{O} \]  
(gibbsite)

Using kaolinite as an example, percent weight of hydroxyl water lost is calculated from the following formula weights.

\[ \text{Si}_2\text{Al}_2\text{O}_5(\text{OH})_4 = 258 \text{ g mol}^{-1} \]
\[ 2\text{H}_2\text{O} = 36 \text{ g mol}^{-1} \]

Percent weight of hydroxyl water lost = \((36/258) \times 100 = 34.6\%\)

9. Report
Report percent gibbsite and/or kaolinite to nearest whole number.

10. Precision
Precision data are not available for this procedure.

11. References
Jackson, M.L. 1956. Soil chemical analysis. Advan. course. M. L. Jackson, Madison, WI.

**Instrumental Analyses (7A)**

**Differential Scanning Calorimetry (7A6)**

**1. Application**

Calorimetry measures specific heat or thermal capacity of a substance. Differential scanning calorimetry (DSC) is a calorimetric technique in which the rate of heat flow between a sample and a reference material is measured as materials are held isothermal to one another. The DSC directly measures the magnitude of an energy change (H, enthalpy or heat content) in a material undergoing an exothermic or endothermic reaction. DSC is commonly used to quantify gibbsite \((\text{Al(OH)}_3)\) and kaolinite \((\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4)\) in soils and clays by measuring the magnitude of their dehydroxylation endotherms which are between 250 to 350 °C and 450 to 550 °C, respectively (Karathanasis and Hajek, 1982; Jackson, 1956; Mackenzie and Berggen, 1970; Mackenzie, 1970).

**2. Summary of Method**

An 8 mg sample of soil clay is weighed into an aluminum sample pan and placed in the DSC sample holder. The sample and reference are heated under flowing \(\text{N}_2\) atmosphere from a temperature of 30 to 600 °C at a rate of 10 °C min\(^{-1}\). Data are collected by the computer and a thermogram is plotted. Gibbsite and kaolinite are quantified by measuring the peak area of any endothermic reactions.
between 250 to 350 °C and 450 to 550 °C, respectively, and by calculating the H of the reaction. These values are related to the values for the respective known quantities of the two minerals (gibbsite and kaolinite).

3. Interferences

Organic matter is objectionable because it produces irregular exothermic peaks in air or O₂, commonly between 300 to 500 °C, which may obscure important reactions from the inorganic components of interest (Schnitzer and Kodama, 1977). Analysis in an inert N₂ atmosphere alleviates this problem. Mineral salts that contain water of crystallization also may be interferences. Samples should be washed free of any soluble salts.

Use a representative soil sample as sample size is small (<10 mg). Avoid large aggregates in sample, the presence of which may cause thermal interferences because of differential kinetics of gas diffusion through the sample and physical movement of sample in a reaction.

The dehydroxylation of goethite is between 250 to 400 °C and may interfere with the identification and integration of the gibbsite endotherm (250 to 350 °C) (Mackenzie and Berggen, 1970). The dehydroxylation of illite is between 550 to 600 °C and partially overlaps the high end of the kaolinite endotherm (450 to 550 °C), resulting in possible peak integrations (Mackenzie and Caillere, 1975). The dehydroxylation of hydroxy-Al interlayers in hydroxy-Al interlayered vermiculite (HIV) is between 400 to 450 °C and may interfere with the low end of the kaolinite endotherm (450 to 550 °C), especially in the poorly crystalline kaolinites (Mackenzie and Caillere, 1975). Similarly, the dehydroxylation of nontronites, Fe-rich dioctahedral smectites is between 450 to 500 °C and may interfere with kaolinite identification and measurement (Mackenzie and Caillere, 1975).

4. Safety

Secure high pressure N₂ tanks and handle with care. When changing the tanks, valves should be protected with covers. Do not program the analyzer for >950 °C because it may present a safety hazard during sample analysis and cleaning cycles. Do not heat aluminum sample pans >600 °C. Aluminum melts at 660 °C, and the sample pans alloy with and destroy the sample holders. Always use high quality purge gases with the DSC. Minimum purity of 99.9% is recommended.

5. Equipment

5.1 Thermal analysis system, Perkin-Elmer 7 series, 7500 computer, TAC7 instrument controllers

5.2 Differential scanning calorimeter module, DSC7, Hewlett-Packard digital plotter

5.3 Pressure tanks (2), N₂, purity 99.99%
5.4 Two-stage gas regulators (2), 50 psi outlet pressure
5.5 Electronic balance, ±0.1-mg sensitivity, Mettler AE160
5.6 Forceps, flat-tipped
5.7 Weighing spatula
5.8 Desiccator, glass
5.9 Mortar and pestle
5.10 Sieves, 100 mesh or 80 mesh
5.11 Kaolinite, standard, poorly crystalline, Georgia Kaolinite, Clay Minerals Society, Source Clay Minerals Project, sample KGa-2
5.12 Gibbsite, standard, Surinam Gibbsite, National Soil Survey Laboratory (NSSL), 67L022

6. Reagents
6.1 Magnesium nitrate saturated solution \([\text{Mg(NO}_3\text{)}_2\cdot6\text{H}_2\text{O}]\)
6.2 Ethanol

7. Procedure

**Derive <2µm Clay Fractions**

7.1 Prepare Na-saturated clay as in method 7A2i, preparation of clay suspension, 7.8 to 7.19.
7.2 Dry the clay suspension and transfer to mortar. Moisten sample with ethanol and grind with pestle to make a homogeneous slurry.
7.3 Air-dry sample using flowing air in hood. Lightly grind sample with pestle to make a homogeneous powder. Transfer to original container for storage until use.
7.4 Prior to TGA analysis, sieve sample with 80-mesh screen. Equilibrate sample overnight over a saturated magnesium nitrate solution (55% rh) in a glass desiccator.

**DSC Operation**

7.5 Set-up the instrument and calibrate.
7.6 Weigh ≈8 mg of sample, i.e., <100-mesh whole-soil or derived <2-µm clay fraction, into tared aluminum sample pan. Refer to section on derived <2-µm clay fractions, 7.1 to 7.4.
7.7 Use flat-tipped forceps to remove aluminum sample pan from balance. Drop sample from a 4- to 5-mm height to uniformly distribute sample in pan. Return the sample pan with sample to the balance and record weight to nearest ±0.1 mg. This weight is entered into computer in appropriate menu.
7.8 Carefully place aluminum sample pan in the center of DSC platinum sample side (left side) of sample holder. Place platinum two-hole lid on holder that covers the sample pan. Align lid holes with purge gas exit hole in DSC head.

7.9 Place empty aluminum sample pan in reference side (right side) of sample holder. Place remaining platinum two-hole lid on holder that covers the sample pan. Align lid holes as in previous step.

7.10 Close DSC head cover and lock.

7.11 The standard sample run heating program has a heating rate of 10 °C min⁻¹, 5.3 min data delay, 5.0 min N₂ purge.

7.12 Start the “Run” program.

7.13 Observe the milliwatts (mW) readout on the computer display terminal and when reading stabilizes (≈5 to 10 s), remove the sample pan and sample from the sample side of sample holder. Do not disturb the reference side.

7.14 To store data, enter the appropriate file name on the computer for the completed run. If data are not stored by appropriate file name, data are stored under a default file name of “gsav”. Only four of these files can be saved at any one time, after which files are overwritten. Once a file is named, it cannot be changed.

8. Calculations

The thermogram is displayed on the computer monitor. The area under the DSC curve is proportional to the enthalpy (H). Use the computer to calculate the H or enthalpy of reaction per g of kaolinite and/or gibbsite (joules g⁻¹) as appropriate.

8.1 % Kaolinite weight=H/12.62

where:

12.62=factor obtained from standard curve of kaolinite mixtures using China clay of undetermined purity

8.2 % Gibbsite weight=H/15.03

where:

15.03=factor obtained from standard curve of gibbsite values using deferrated Surinam gibbsite of undetermined purity

9. Report

Report percent kaolinite and/or gibbsite to the nearest whole number.

10. Precision

Precision data are not available for this procedure.
11. References

Jackson, M.L. 1956. Soil chemical analysis. Advan. course. M. L. Jackson, Madison, WI.


Total Analysis (7C)
HF Dissolution (7C3)

1. Application

Historically, elemental analysis was developed for the analysis of rocks and minerals (Washington, 1930). The elemental analysis of soils, sediments, and rocks necessitates their decomposition into soluble forms. Hydrofluoric acid (HF) is efficient in the digestion and dissolution of silicate minerals for elemental decomposition. Method 7C3 is an HF acid digestion. Elemental concentration is determined by atomic absorption using 100 mg of clay suspension contained in a
closed vessel with boric acid \((H_3BO_3)\) to neutralize excess acid (Berdanier, Lynn, and Threlkeld, 1978; Soil Conservation Service, 1984).

2. Summary of Method

To 100 mg of clay suspension (method 7A2i), 5 mL of HF acid are added. The solution is heated, cooled, and 2 to 3 g of \(H_3BO_3\) are added to neutralize excess acid. The solution is diluted to 100 mL, allowed to stand overnight, and 20 mL are decanted. The concentrations of Fe, Al, and K are determined by atomic absorption in methods 6C7a, 6G11a, and 6Q3a, respectively. Data are reported in method 7C3.

3. Interferences

Organic material may remain as a residue with this method.

4. Safety

Perform procedure in hood. Keep HF acid refrigerated and avoid contact with skin.

5. Equipment

5.1 Pipette, 5 mL,
5.2 Volumetric flask, Nalgene, 100 mL
5.3 Polyethylene container, 25 mL, with cover
5.4 Electronic balance, ±0.1-mg sensitivity

6. Reagents

6.1 Distilled water
6.2 Hydrofluoric acid (HF), 48%,
6.3 Boric acid, \((H_3BO_3)\), granular

7. Procedure

HF Dissolution

7.1 Prepare Na-saturated clay as in method 7A2i, preparation of clay suspension, 7.8 to 7.19.
7.2 Pipette 2 mL of clay suspension into a 25-mL Teflon cup and add 5 mL of HF. A 100 mg of 100-mesh whole-soil sample may be substituted for the clay suspension.
7.3 Pipette a duplicate sample into a weighing dish, dry at 105 °C, and weigh. Use this sample for calculations.
7.4 Place covered Teflon cup in stainless steel retainer and tighten Teflon cap. Place sample in oven at 105 °C for ≈4 h.
7.5 Turn off oven, open door, and let stand overnight to cool.
7.6 Remove sample from oven.
7.7 Under a hood, remove Teflon cup from steel retainer vessel and add 2 to 3 g of \( H_3BO_3 \) acid.
7.8 Rinse contents of Teflon cup into a 100-mL Nalgene volumetric flask and adjust to volume with distilled water. Allow to stand overnight.
7.9 Decant ≈20 mL into a 25-mL polyethylene container for elemental analysis by atomic absorption. Refer to methods 7C7a, 6G11a, and 6Q3a.

8. Calculations

Use the MR 2.0 to perform calculations. Inputs are as follows: project number; sample number; tare value; tare + sample value; Al and Fe readings (mg L\(^{-1}\)); and K readings (meq L\(^{-1}\)). Review data for internal consistency. Request a rerun, if necessary, at this time. Store data on a data disk.

The following example illustrates the conversion calculations of atomic absorption readings or element concentrations of Fe, Al, and K to appropriate oxide forms. The concentrations of Fe and Al (mg L\(^{-1}\)) and K (meq L\(^{-1}\)) are converted to percent Fe\(_2O_3\), Al\(_2O_3\), and K\(_2O\), respectively. Refer to method 4A5 for air-dry/oven-dry ratio (AD/OD).

\[
\begin{align*}
\text{Sample weight} & = S = 0.1071 \text{ g} \\
\text{Fe reading} & = [\text{Fe}] = 34.2 \text{ mg L}^{-1} \\
\text{Fe}_2O_3 \text{ molecular weight} & = \text{Fe}_2O_3 = 159.70 \\
\text{Fe atomic weight} & = \text{Fe} = 55.85 \\
\text{Al reading} & = [\text{Al}] = 72.8 \text{ mg L}^{-1} \\
\text{Al}_2O_3 \text{ molecular weight} & = \text{Al}_2O_3 = 101.94 \\
\text{Al atomic weight} & = \text{Al} = 26.98 \\
\text{K reading weight} & = [\text{K}] = 0.61 \text{ meq L}^{-1} \\
\text{K}_2O \text{ molecular weight} & = \text{K}_2O = 94.19 \\
\text{K atomic weight} & = \text{K} = 39.10 \\
\text{K equivalent weight} & = 39 = 39 \\
\text{AD/OD} & = 1.024 = \text{AD/OD} \\
100 \text{ mL}/1000 \text{ mL} = \text{dil. factor} & = 100/1000 \\
1/1000 \text{ mg g}^{-1} = \text{conv. factor} & = 1/1000 \\
100/1 \text{ P/100 pts.} = \text{conv. factor} & = 100/1 \\
\% \text{ Fe}_2O_3 & = \\
[\text{Fe}]x1/Sx100/1000x1/1000x100/1x\text{AD/ODxFe}_2O_3/\text{Fe} \\
& = 34.2x1/0.1071x0.1x0.001x100x1.024x159.7/111.70 \\
& = 4.68
\end{align*}
\]
% $\text{Al}_2\text{O}_3$ =

$$[\text{Al}] \times \frac{1}{S} \times \frac{100}{1000} \times \frac{1}{100} \times \frac{\text{AD/OD}}{\text{Al}} \times \frac{\text{Al}_2\text{O}_3}{\text{Al}} =$$

$$72.8 \times \frac{1}{0.1071} \times 0.1 \times 0.001 \times 100 \times 1.024 \times 101.94/53.96$$

% $\text{Al}_2\text{O}_3$ = 13.15%

% $\text{K}_2\text{O}$ =

$$[\text{K}] \times \frac{1}{S} \times \frac{100}{1000} \times \frac{1}{100} \times \frac{\text{AD/OD}}{\text{K}_2\text{O}} \times \frac{\text{K}_2\text{O}}{\text{K}} \times 39 =$$

$$0.61 \times \frac{1}{0.1071} \times 0.1 \times 0.001 \times 100 \times 1.024 \times 94.19/78.2 \times 39 =$$

% $\text{K}_2\text{O}$ = 2.74

9. Report

Report data to nearest whole percent.

10. Precision

Precision data are not available for this procedure. A quality control check sample is routinely run in HF analyses. For 38 observations of the quality control check sample, the mean, standard deviation, and C.V. for percent Fe, $\text{Al}_2\text{O}_3$, and $\text{K}_2\text{O}$ are as follows:

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Mean</th>
<th>Std. Dev.</th>
<th>C.V.</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Fe</td>
<td>2.8</td>
<td>0.40</td>
<td>14</td>
</tr>
<tr>
<td>% $\text{Al}_2\text{O}_3$</td>
<td>12.1</td>
<td>1.04</td>
<td>9</td>
</tr>
<tr>
<td>% $\text{K}_2\text{O}$</td>
<td>2.4</td>
<td>0.38</td>
<td>16</td>
</tr>
</tbody>
</table>

11. References


LABORATORY PREPARATION OF SOIL SAMPLES (1B)
Carbonate-Containing Material (1B3)

Procedure

Prepare dialysis membrane sacks from 5½-inch cellulose casing (Visking Company), using large rubber bands to tie the bottoms. Place the sample (as much as 6 kg if very gravelly and highly calcareous) in a dialysis membrane and add about 1 L pH 5, NaOAc buffer. Tie the top of the dialysis membrane around a glass breather tube 4-in long and hang the assembly in a 60-L reservoir of buffer held in a 20-gal plastic garbage can. If carbonate is dissolving, knead the membrane to release bubbles of CO₂. When bubbles of CO₂ no longer form on kneading, open the dialysis membrane and use strong acid to check the coarser material for carbonate coatings (carbonate remains longer in the coarser material). When sample is free of carbonate, desalt it by dialysis against tap water flowing continuously through a large plastic garbage can. Check the ionic concentration inside the membrane by measuring conductivity of a small volume of the supernatant liquid poured out through the breather tube. Continue dialysis until the salt concentration is less than 10 meq/L.

The procedure used to dry the sample depends on whether the particles larger than 2 mm have been removed before buffer treatment. If they have been removed, withdraw excess water from the sample in the membrane with filter candles. Knead the membrane to mix the sample and place it in contact with ethanol to desiccate further. Remove the sample from the membrane and air-dry.

If the buffer-treated sample contains particles larger than 2 mm, wet sieve the sample through a 2-mm sieve. Then dry sieve the material remaining on the sieve (>2 mm) and add the <2-mm fraction from this sieving to the <2-mm fraction separated by the wet sieving. Remove most of the water from the <2-mm fraction with filter candles. Use ethanol to transfer the samples to shallow pans and dry. Ethanol prevents aggregation of clay into durable flakes during drying.
Discussion

The time required for carbonate removal varies greatly, depending on particle size, percentage and type of carbonate, and sample size. Samples from horizons strongly cemented by carbonate have required as long as 2 months. The concentration of alkaline-earth ions in the buffer greatly affects the rate of carbonate removal. Changing the buffer in the reservoir well before the buffer capacity has been exhausted, thereby keeping the alkaline-earth ion concentration low, increases the rate markedly. Desalting usually takes about 4 days.

For carbonate-cemented horizons, the whole sample, not just the <2-mm material, must be buffer treated. Furthermore, for horizons without carbonate cementation, buffer treatment of the whole sample has the advantage of washing the >2-mm skeletal material free of adhering fines and organic material. This problem is considered further in 1B4.

For very gravelly horizons, large samples (several kilograms) are necessary for buffer treatment because of the small amount of <2-mm material. Using large samples also increases precision of the >2-mm percentage.

References


Carbonate-Indurated Material Containing Coarse Fragments (1B4)

Break the field sample to get several representative subsamples. Remove the carbonate from one subsample by acid treatment and separate the coarse fragments from the fine earth (1B3). Weigh the two fractions. Use the noncarbonate fine earth for the standard characterization and mineralogical measurements (sections 6-7).

Grind another subsample of the whole field sample to pass 80-mesh sieve. Determine the carbonate content (weight) of this whole ground subsample (6E).

These weights can be used to calculate the CaCO₃ percentage of the fine earth. Any analytical value based on the noncarbonate fine earth can be converted to the whole-soil basis as well as to the basis of the carbonate-containing fine earth.

PARTICLE-SIZE ANALYSIS (3)

Particles <2 mm (Pipette Method) (3A)

An automated balance system, consisting of a Radio Shack Model II microcomputer interfaced to a Mettler PL2000 electronic balance (for sand) and a Mettler AE160 electronic balance (for silt and clay), is used for determining, storing, and processing sample weights.
Air-Dry Samples (3A1)

Apparatus

- Fleaker, 300 ml (tare to 1 mg)
- Pasteur-Chamberlain filter candles, fineness “F”
- Shaker, horizontal, 120 oscillations per minute
- Cylinders, 1000 ml
- Stirrer, motor-driven
- Stirrer, hand. Fasten a circular piece of perforated plastic to one end of a brass rod.
- Shaw pipette rack
- Pipets, 25 ml automatic (Lowy with overflow bulb)
- Polyurethane foam, pipe-insulating cover
- Shaker with ½-in vertical and lateral movements and 500 oscillations per minute. Accommodates a nest of sieves.
- Wide-mouth glass pill bottles with screw caps, 90 ml (tare to 1 mg)
- Electronic balance (0.1-mg sensitivity)
- Set of sieves. Square-mesh woven phosphor bronze wire cloth. U.S. Series and Tyler Screen Scale equivalent designations as follows:

<table>
<thead>
<tr>
<th>Sand Size</th>
<th>Opening (mm)</th>
<th>U.S. No.</th>
<th>Tyler Mesh Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>VCS</td>
<td>1.0</td>
<td>18</td>
<td>16</td>
</tr>
<tr>
<td>CS</td>
<td>0.5</td>
<td>35</td>
<td>32</td>
</tr>
<tr>
<td>MS</td>
<td>0.25</td>
<td>60</td>
<td>60</td>
</tr>
<tr>
<td>FS</td>
<td>0.105</td>
<td>140</td>
<td>150</td>
</tr>
<tr>
<td>VFS</td>
<td>0.047</td>
<td>300</td>
<td>300</td>
</tr>
</tbody>
</table>

Reagents

- Hydrogen peroxide ($\text{H}_2\text{O}_2$), 30 to 35 percent
- Sodium hexametaphosphate ($\text{NaPO}_3\text{O}_6$). Dissolve 35.7 grams of ($\text{NaPO}_3\text{O}_6$) and 7.94 grams of $\text{Na}_2\text{CO}_3$ per liter of water.
- Demineralized water

Procedure

  Removing organic matter.—Place about 10 air-dry soil containing no particles larger than 2 mm in a tared fleaker. Add about 50-ml of demineralized water (referred to subsequently as water) and then add 5 ml of $\text{H}_2\text{O}_2$. Cover the
fleaker with a watchglass. If a violent reaction occurs, repeat the cold H₂O₂
treatment periodically until no more frothing occurs. Heat the fleaker to about 90
°C on an electric hot plate. Add H₂O₂ in 5-ml quantities at 45-min intervals until
the organic matter is destroyed, as determined visually. Continue heating for
about 30 min to remove any excess H₂O₂.

Removing cementing agents (optional).—Treat the sample with about 200
ml of 1 N sodium acetate buffered at pH 5 to remove carbonates. When CO₂
bubbles are no longer evident, wash free of salts with a filter candle system.
Highly calcareous samples may need a second treatment.

Remove siliceous cementing agents by soaking the sample overnight in
0.1 N NaOH. Iron oxide cementing agents are removed by shaking overnight
in sodium dithionite (6C2). Wash free of salts with filter candle system before
proceeding.

Removing dissolved mineral and organic components.—After the H₂O₂
treatment, place the fleaker in a rack and add about 150 ml of water in a jet
strong enough a short Pasteur-Chamberlain filter of “F” fineness. Five such
washings and filterings are usually enough except for soils containing much
coarse gypsum. Remove soil adhering to the filter by gentle back pressure;
use finger as policeman. Dry the sample overnight in an oven at 105 °C, cool
in a desiccator, and weigh to the nearest milligram. Use the weight of the oven-
dry, H₂O₂-treated sample as the base weight for calculating percentages of the
various fractions.

Dispersing the sample.—Add 10 ml of sodium hexametaphosphate
dispersing agent to the Fleaker containing oven-dry treated sample. Make the
volume to approximately 200 ml. Stopper and shake overnight on a horizontal
reciprocating shaker at 120 oscillations per minute.

Separating sands from silt and clay.—Wash the dispersed sample with water
on a 300-mesh sieve. Silt and clay pass through the sieve into a 1-L cylinder.
Use a clamp and stand to hold the sieve above the cylinder. Avoid using jets
of water in washing the sample. Gently tap the sieve clamp with the side of the
hand to facilitate sieving. Continue washing until the suspension volume in the
cylinder is about 800 ml. Sand and some coarse silt remain on the sieve. It
is important to wash all particles of less than 20 µ diameter through the sieve.
Remove the sieve from the holder, wash the sands into an evaporating dish
with water, and dry at 105 to 110 °C. Bring the silt and clay suspension in the
cylinder to 1 L with water and cover with a watchglass.

Pipeting.—First pipette the <20 µ fraction at a 10-cm depth. Vary
sedimentation times according to temperature. Next, pipette the <2µ fraction
after a predetermined setting time (usually 4½ to 6½ hr). Vary depth according
to time and temperature. Use a Lowy 25-ml automatic pipette and regulate
filling time to about 12 s. Before each pipeting, stir material in the sedimentation
cylinder, and stir the suspension for 30 s with a hand stirrer, using an up-and-
down motion. Note the time at completion of stirring. About 1 min before
sedimentation is complete, lower the tip of the pipette slowly into the suspension to the proper depth with a Shaw pipette rack. At the appropriate time, fill the pipette and empty into a 90-ml, wide-mouth bottle. Rinse the pipette into the bottle once. Dry in an oven overnight at 105 °C. Cool in a desiccator containing phosphorus pentoxide (P₂O₅). Weigh.

**Sieving and weighing the sand fractions.**—Transfer the dried sands to a nest of sieves. Shake for 3 min on a shaker that has ½-in vertical and lateral movements and oscillates at 500 strokes per minute. Record the weights of the individual sand fractions.

**Calculations**

Pipetted fractions:

Percentage of pipetted fractions = \((A - B)KD\)

where:

- \(A\) = Weight (g) of pipeted fraction
- \(B\) = Weight correction for dispersing agent (g)
- \(K\) = \(1000 \div \text{(ml in pipette)}\)
- \(D\) = \(100 \div \text{(g of H₂O₂-treated oven-dry total sample)}\)

The <20-µ fraction minus the <2-µ fraction equals fine silt.

Sand fractions:

Percentage of sieved fractions = weight (g) of fraction on sieve times D.

Coarse silt fraction:

Obtain by difference. Subtract the sum of the percentages of sand plus the <20-µ fraction from 100.

**References**


**Carbonate and Noncarbonate Clay I (3A1a)**

**Apparatus**

- Warburg manometer
- ¼-oz (5-ml) gelatin capsules
- 30-ml plastic cups

**Reagents**

- Hydrochloric acid (HCl), 6 \(N\)
Procedure

If carbonate is present, use the glass bottle containing clay residue from regular pipette analysis and determine carbonate as in 6E1b. Use Warburg manometer.

Calculations

\[ Cc = \left( \frac{\left( A - B - C \right) \times \text{Factor}}{D} \right) \times 100 \]

where:
- \( Cc \) = Carbonate clay (pct <2 mm)
- \( A \) = Upper reading
- \( B \) = Lower reading
- \( C \) = Blank
- \( \text{Factor} \) = Factor derived from standard curve and includes pipette volume factor
- \( D \) = Total sample weight (3A1)

\[ Nc = \text{Total clay} - Cc \]

where:
- \( Nc \) = Noncarbonate clay (pct <2 mm)
- \( Cc \) = Carbonate clay (pct <2 mm)

References

Shields and Meyer (1964).

Moist Samples (3A2)

If drying affects dispersion of treated sample, oven-drying may be avoided by removal of a pipette sample to estimate the total weight of the sample. Pipette 50 ml at a depth of 20 cm at time zero while the suspension is still turbulent. Use the oven-dry weight of the aliquot to calculate the total weight of the <0.05-mm fraction. Add this weight to the total weight of the sands to obtain the total weight of the sample.

An optional procedure is to carefully weigh out two identical samples and pretreat to remove organic matter and dissolved mineral matter. The first sample is continued through the standard procedure, excluding oven-drying. The second sample is oven-dried, weighed, and discarded. The oven-dry weight of the second sample is substituted in the calculations for the first sample.

Carbonate and Noncarbonate Clay (3A2a)

Proceed as in 3A1a except use field-moist sample.
FABRIC-RELATED ANALYSES (4)

Bulk Density (4A)

Density is defined as mass per unit volume. Soil density as commonly used differs from most density measurements in that the volume of interparticle space is included but the mass of the liquid phase is excluded. Therefore, soil density has been called bulk density, Dw, to distinguish it from the more usual density that is based on intraparticle volume only. Since the volume of a shrinking-swelling soil changes with a change in its water content, subscripts are added to designate the moisture condition when the measurement was made. Thus, Dw, is the bulk density of a moist sample, Dw, is the bulk density of a clod sample equilibrated at 1/3-bar tension, and Dw, is the bulk density of a dry sample.

Saran-Coated Clods (4A1)

Reagents

- Methyl ethyl ketone
- Dow Saran F310.—The saran resin dissolves readily in acetone or methyl ethyl ketone. In this method, methyl ethyl ketone is used as a solvent because it is less soluble in water than is acetone and there is less penetration of the Saran-solvent solution into a moist clod. However, acetone is adequate for a first (field) coat and is more readily available. Saran-solvent ratios of 1:4 to 1:7 are used, depending on the porosity of the soil to be coated.
- Coating solution.—To prepare the solution, fill a weighted container with a solvent to about three-fourths its volume. From the weight of the solvent, calculate the weight of resin required to obtain a predetermined resin-solvent ratio and add to the solvent. Since the solvent is flammable and its vapors form explosive mixtures with air, mix the components with an air-powered or nonsparking electric stirrer under an exhaust hood. Information on the safe handling and use of methyl ethyl ketone is available in Chemical Safety Data Sheet SD-83, Manufacturing Chemists’ Association, Inc., 1825 Connecticut Avenue NW, Washington, D.C. The threshold limits of methyl ethyl ketone are 200 ppm as given in OSHA standards, Part 2, Section 1910.93, table G1.
- If a high-speed stirrer is used, the resin dissolves in about 1 hr. In the field, mix with a wooden stick. Metal cans (1 gal) are satisfactory containers for mixing and storing the plastic. Keep the containers tightly closed to prevent evaporation of the solvent.
Procedure

Collect natural clods (three per horizon) of about 100 to 200 cm³ in volume (fist-sized). Remove a piece of soil larger than the clod from the face of a sampling pit with a spade. From this piece, prepare a clod by directly cutting or breaking off protruding peaks and material sheared by the spade. If roots are present, they can be cut conveniently with scissors or side cutters. In some soils, clods can be removed directly from the face of a pit with a knife or spatula. No procedure for taking clod samples fits all soils; the procedure must be adjusted to meet the conditions in the field at the time of sampling.

The clods are tied with the fine copper wire or placed in hairnets and suspended from a rope or string, hung out like a clothesline. Moisten dry clods with a fine mist spray. The suspended clods are dipped by raising a container of the dipping mixture upward around each clod, so it is immersed momentarily. The saran-coated clods should be allowed to dry for 30 min or longer. Clods coated in this way can be transported to the laboratory and examined microscopically in an undisturbed state. For convenience, either of two concentrations of plastic solution is usually used—a 1:7 solution for most soil samples or a 1:4 solution for clods that have larger pores. If bulk density at field-moisture content is desired, store the clods in waterproof plastic bags as soon as the coating dries since the coating is permeable to water vapor. Although the coating keeps the clods intact, they may be crushed in transport unless they are packed in rigid containers.

In the laboratory, additional coatings of plastic are applied to make the clod waterproof and to prevent its disruption during wetting. Then, weigh the clod, either in its natural moisture condition or in an adjusted moisture condition (e.g., ½-bar tension) in air and in water to obtain its volume by Archimedes’ principle. Subsequent changes in moisture condition and volume of the soil sample can be followed by reweighing the coated clod in air and in water. Finally, weigh the oven-dry clod in air and in water.

Be careful not to lose any soil material because the weight of material lost is calculated as soil moisture, and calculated bulk densities depend on the final oven-dry weight of the clod.

Bulk-density values determined by this method are reported on the basis of fine-earth fabric. Weight and volume measurements are made on clod samples that may contain particles >2 mm; however, after the measurements are made, the weight and volume of the coarse fraction are subtracted. The remainder consists of the weight of <2-mm material and the volume of these fine-earth particles and the pore space associated with them.

Sometimes it is necessary to correct bulk density for weight and volume of the plastic coating. The coating has a density of about 1.3 g/cm³ and it loses 10 to 20 percent of its air-dry weight on oven-drying at 105 °C. Thus, the amount of correction becomes smaller as bulk density of the soil approaches the density of the coating and as moisture content of the soil approaches the weight loss of the coating.
Calculations

\[
Db_{\frac{1}{3}} = \frac{wt\text{clod}_{od} - wt>2\text{ mm} - tcoat_{od}}{vol\text{clod}_{\frac{1}{3}} - vol>2\text{ mm} - vol\text{ coat}}
\]

\[
Db_{od} = \frac{wt\text{clod}_{od} - wt>2\text{ mm} - wt\text{coat}_{od}}{vol\text{clod}_{od} - vol>2\text{ mm} - vol\text{ coat}}
\]

\[
W_{\frac{1}{3}} = \frac{wt\text{clod}_{\frac{1}{3}} - wt\text{clod}_{od} - (wt\text{clod}_{ad} - wt\text{coat}_{od})}{wt\text{clod}_{od} - wt>2\text{ mm} - wt\text{coat}_{od}} \times 100
\]

where:
\(Db_{\frac{1}{3}}\) = bulk density of <2-mm fabric at \(\frac{1}{3}\)-bar tension in grams per cubic centimeter
\(Db_{od}\) = bulk density of <2-mm fabric at oven-dryness in grams per cubic centimeter
\(W_{\frac{1}{3}}\) = the weight percentage of water retained at \(\frac{1}{3}\)-bar tension
\(wt\text{ clod}_{od}\) = weight of oven-dry coated clod
\(wt\text{ clod}_{\frac{1}{3}}\) = weight of coated clod equilibrated at \(\frac{1}{3}\)-bar tension
\(vol\text{ clod}_{od}\) = volume of oven-dry coated clod
\(vol\text{ clod}_{\frac{1}{3}}\) = volume of coated clod equilibrated at \(\frac{1}{3}\)-bar tension
\(vol>2\text{ mm}\) = volume of material >2 mm separated from clod after oven-drying
\(wt>2\text{ mm}\) = weight of material >2 mm separated from after oven-drying
\(wt\text{ coat}_{ad}\) = weight of Saran coating before oven-drying
\(wt\text{ coat}_{od}\) = weight of Saran coating after oven-drying
\(vol\text{ coat}\) = volume of Saran coating (estimated)

The field coat (initial coat) of plastic penetrates the clod to some extent. Weight of the field coat, estimated to 1.5 times the weight of each additional coat, is computed by:

\[
W_{\text{coat}_{\text{init}}} = \frac{(Wt\text{clod}_{B} - Wt\text{clod}_{A})}{3} \times 1.5
\]

where:
\(Wt\text{coat}_{\text{init}}\) = Weight of field (initial) coat
\(Wt\text{clod}_{A}\) = Weight of clod with one coat of plastic
\(Wt\text{clod}_{B}\) = Weight of clod with three additional coats of plastic
References
Brasher et al. (1966).

**Air-Dry (Db) (4A1b)**
After measuring field-state volume, place clods in a drying room kept at 90 °F. Weigh a few clods each day until they reach a constant weight. Assume then that all the clods are air-dry. Coat them again with Saran and measure “air-dry” volume as described in 4A1a. Determine oven-dry weight and calculate bulk density as described in 4A1.

**30-cm Absorption (Db) (4A1c)**
After measuring air-dry volume, remove a patch of the Saran coating from one side of each clod. Next, place the clods on a sand tension table with the exposed side in contact with very fine sand that has been equilibrated to 30-cm water tension. Again weigh a few clods each day until they reach constant weight and assume that all the clods are at 30-cm water tension. Most clods reach equilibrium in 7 to 10 days. Remove the clods from the tension table and coat with Saran until waterproof. Measure volume of the clods and calculate bulk density as described in 4A1.

**⅓-Bar Desorption II (Db/3) (4A1e)**
Cut a flat surface on the coated field-moist clods with a sharp knife or diamond saw. Seat the clods on saturated ceramic plates with the flat surface in contact with the plates. Place the plates in pans and add water to just cover the surface of the plates. After the clods become wet by capillary movement, place the plates in a pressure cooker and equilibrate at ⅓ bar. After equilibration, carefully remove the clods from the plates and dip in Saran until waterproof. Measure volume of the clods and calculate bulk density as described in 4A1.

**⅓-Bar Desorption III (Db/3) (4A1f)**
Proceed as in 4A1e except prewet the clods at 10-cm tension on porous bricks (cheesecloth layer between clod and brick) instead of saturating them on ceramic plates.

**⅓-10-Bar Desorption (Db/40) (4A1g)**
Proceed as in 4A1d, e, or f except make final desorption at ⅓-10 bar.

**Paraffin-Coated Clods (4A2)**
**Oven-dry (Db) (4A2a)**
Oven-dry the clods, coat with paraffin, and weigh in water and in air. Calculate bulk density as follows:
\[ \text{Db}_d (\text{g/cc}) = \frac{\text{Wt}_{\text{air}} - \text{Wt}>2 \text{ mm}}{\text{Wt}_{\text{air}} - \text{Wt}_{\text{H}_2\text{O}} - (\text{Wt}>2 \text{ mm}/2.65)} \]

where:
- \( \text{Wt}_{\text{air}} \): Weight in air
- \( \text{Wt}>2 \text{ mm} \): Weight of >2-mm fraction in clod
- \( \text{Wt}_{\text{H}_2\text{O}} \): Weight in water

**Nonpolar-Liquid Saturated Clods (4A4)**

**Procedure**

Place a natural clod in a nonpolar liquid of low viscosity, e.g., high-purity kerosene. Evacuate under vacuum until bubbles cease to appear and weigh the clod suspended in the nonpolar liquid. Remove the clod, place it on a sand table under 3-cm tension against the nonpolar liquid to drain off excess nonpolar liquid, and weigh it in air. The difference in weight of the clod in air and suspended in the nonpolar liquid divided by the density of the nonpolar liquid is the clod volume. Determine the oven-dry weight and calculate bulk density as in 4A1. The difference between the clod’s initial weight before immersion in the nonpolar liquid and its oven-dry weight is the moisture content.

**References**

Rennie (1957).

**Water Retention (4B)**

**Pressure-Plate Extraction (4B1)**

After measuring the \( \frac{1}{3} \)-bar volume (4A1d), the Saran coating is removed from the flat surface of the clods. The clods are allowed to air-dry (4 to 6 days) and then placed in the drying room 2 or 3 days. They are then placed on a tension table of very fine sand and equilibrated to 5-cm tension as in 4A1d. After about 2 weeks, some of the highly organic clods that have not rewetted are placed in a pan of free water overnight to make certain that wetting is complete. The clods are again desorbed to \( \frac{1}{3} \)-bar as in 4A1d and volume measurements of the clod are made and bulk density is calculated as described in 4A1.

**Soil Pieces (4B1b)**

**Procedure**

Make desorption measurements of soil pieces concurrently with the sieved-sample measurements. Cover the sieved samples in retainer rings with small
squares of industrial tissues (Kimwipes). Place the soil pieces (about 2.5 cm in diameter) on the tissues before adding water to the plate. Proceed as in 4B1a. If the soil pieces contain >2-mm material, wet sieve and weigh the oven-dry >2-mm material. Report moisture content as percentage of oven-dry weight of <2 mm material.

References
Young (1962).

Sand-Table Absorption (4B3)
Saran-coated clods that have been equilibrated on a sand table to determine bulk density (4A) can also be used to determine water content at these tensions.

Micromorphology (4E)
Thin Sections (4E1)
Moved-Clay Percentage (4E1c)

Apparatus
• Diamond tile saw
• Thin-section equipment
• Point-counting eyepiece

Reagents
• Aroclor 5460 (Monsanto)
• Polyester resin
• Styrene

Procedure
Impregnate an undisturbed field sample with Aroclor (4E1b). With a diamond saw, cut the clods into pieces about 3 by 1 by 1 cm. Mount about 10 pieces side by side with polyester resin (use a little styrene) to form a block. Cut this assembly to form slices 3 by 1 by 1 cm. Slice all the field sample, composite, and withdraw subsamples of 10 to 15 slices. Stack these slices, tape them together, and mount in plastic (polyester resin plus styrene). Cut a section through the stack parallel to the direction of stacking and along the longer of the two remaining axes. Mount one such section from each stack on a glass slide and prepare a thin section.

To estimate the moved-clay volume insert a point-counting eyepiece into the microscope and run a transect along each strip. Keep the transect length and the number of fields in a transect constant. Count the number of points that fall on
moved clay. Divide this number by the total number of points to get an estimate of the proportion of moved clay. To convert these volume estimates to weight estimates, multiply by the ratio of the bulk density of the moved clay to the bulk density of the appropriate dry fabric. Assume that the moved clay has a bulk density of 2.00 g per cubic centimeter.

References

Scanning Electron Microscopy (4E2)
Electronically reproduced images of fabric surfaces can be obtained at magnifications ranging from 50 to 30,000 diameters. Depth of focus by this technique is large compared to that by light microscope. Stereoscopic pictures can be taken to give three-dimensional viewing.

Procedure
Take a sample of fabric up to 10 mm in diameter and 2 to 3 mm thick. Coat with a thin metallic layer and insert in the instrument. The image is displayed on a cathode ray tube.

ION EXCHANGE ANALYSES (5)

Cation Exchange Capacity (CEC) (5A)
NH₄OAc, pH 7.0 (Buchner funnel) (5A1)

Reagents
- Ammonium acetate (NH₄OAc), 1 N, pH 7.0. Mix 68 ml ammonium hydroxide (NH₄OH), specific gravity 0.90, and 57 ml 99.5-percent acetic acid (CH₃COOH) per liter of solution desired. Cool, dilute to volume with water, and adjust to pH 7.0 with CH₃COOH or NH₄OH. Optionally prepare from NH₄OAc reagent salt and adjust pH.
- Ethanol (CH₃CH₂OH), 95-percent, U.S.P.
- Nessler’s reagent (optional). Prepare according to Yuen and Pollard.

Procedure
Weigh 25 g air-dry <2-mm soil (some early work was done with 50-g samples) into a 250-ml Erlenmeyer flask and add 35 to 50 ml NH₄OAc solution. Stopper, shake the flask for several minutes, and allow to stand overnight. Transfer contents of the flask to a Buchner funnel (Coors No. 1) fitted with moist Whatman No. 42 filter paper. Filter, using gentle suction if needed. Leach with 200 ml
NH₄OAc, adding small amounts at a time so that leaching requires no less than 1 hour. Transfer leachate from suction flask to volumetric flask and retain for analysis of NH₄OAc-extractable cations (methods 6N2, 6O2, 6P2, 6Q2).
Add 95-percent ethanol in small amounts to the ammonium-saturated soil remaining on the Buchner funnel until the leachate gives a negative test for ammonia with Nessler’s reagent or leach with 100 ml ethanol.

References
Peech et al. (1947) and Yuen and Pollard (1952).

**Direct Distillation of Adsorbed Ammonia, Kjeldahl (5A1a)**

**Reagents**
- Sodium chloride (NaCl)
- Antifoam mixture. Mix equal parts of mineral oil and n-octyl alcohol.
- Sodium hydroxide (NaOH), 1 N
- Hydrochloric acid (HCl), 0.2 N, standardized
- Boric acid (H₃BO₃), 4-percent
- Mixed indicator. Mix 1.250 g methyl red and 0.825 g methylene blue in 1 liter 95-percent ethanol
- Brom cresol green, 0.1-percent, aqueous solution

**Procedure**
Transfer the soil plus filter paper from method 5A1 to a Kjeldahl flask. Add 400 ml water and about 10 g NaCl, 5 drops antifoam mixture, a gram or two of granular zinc, and 40 ml 1 N NaOH. Connect the flask with the condenser and distill 200 ml into 50 ml 4-percent H₃BO₃ solution. Titrate the distillate to the first tinge of purple with 0.2 N HCl, using 10 drops mixed indicator and 2 drops brom cresol green.

**Calculations**

\[
\text{CEC (meq/100 g)} = (\frac{A}{B}) \times N \times 100
\]

where:
- \(A\) = Volume HCl (mL)
- \(B\) = Sample weight (g)
- \(N\) = Normality of acid

Report on oven-dry basis.

**References**
Peech et al. (1947).
Displacement of Adsorbed Ammonia, Semimicro Kjeldahl (5A1b)

Reagents

- Sodium chloride (NaCl), acidified, 10-percent. Dissolve 100 g NaCl, reagent-grade, ammonia-free, in 750 ml warm water; add 25 ml 2 N hydrochloric acid (HCl) and bring to 1000-ml volume.
- Sodium hydroxide (NaOH), 1 N
- Boric acid (H₃BO₃), 2-percent
- Sulfuric acid (H₂SO₄), 0.01 N, standardized
- Ethanol, 95-percent
- Mixed indicator. Dissolve 0.1 g methyl red and 0.1 g brom cresol green in 250 ml ethanol.

Procedure

Leach soil from method 5A1 with 240 ml 10-percent acidified NaCl solution, using small increments. Drain completely between each increment. Transfer the leachate to a 250-ml volumetric flask and adjust volume to mark. Pipette a suitable aliquot of the leachate into a micro-Kjeldahl distillation flask and attach to steam-distillation apparatus. Start steam distillation and slowly add 10 ml 1 N NaOH. Catch distillate in a 250-ml Erlenmeyer flask containing 10 ml H₃BO₃ and 10 drops of mixed indicator. Distill for 5 minutes after H₃BO₃ turns green, lower receiving flask, and rinse condenser and outlet hose into receiving flask. Titrate the ammonia with 0.01 N H₂SO₄ to a red end point, using a blank for comparison.

Calculations

\[
\text{CEC (meq/100 g)} = \frac{A}{B} \times N \times \frac{C}{D} \times 100
\]

where:
- \(A\) = Volume H₂SO₄ (mL)
- \(B\) = Sample weight (g)
- \(N\) = Normality of acid
- \(C\) = Volume leachate (mL)
- \(D\) = Volume aliquot (mL)

Report on oven-dry basis.

NaOAc, pH 8.2 (5A2)
Carbonifuge Method (5A2a)

Reagents

- Sodium acetate (NaOAc), 1 N, pH 8.2
• Ethanol, 95-percent
• Ammonium acetate (NH₄OAc), 1 N, pH 7.0. Add 57 ml concentrated acetic acid and 68 ml concentrated NH₄OH, specific gravity 0.90, to about 800 ml water. Cool and dilute to 1 liter and adjust to pH 7.0 by adding more NH₄OH or acetic acid.

**Procedure**

Weigh 5-g samples to an accuracy of 1 percent and place in centrifuge tubes. Add 33 ml NaOAc, stopper the tubes, and shake for 5 minutes. Remove stopper and centrifuge until the supernatant liquid is clear (usually 5 min). Decant the supernatant liquid as completely as possible and discard. Repeat four times, discarding the supernatant liquid each time. After the last saturation, wash the rubber stoppers and use absorbent paper to remove any acetate crystals remaining on lip of centrifuge tube. Add about 30 ml ethanol to each tube, stopper, shake for 5 minutes, remove stopper, and centrifuge until the supernatant liquid is clear. Decant and discard the supernatant liquid. Continue washing until the electrical conductivity of the supernatant liquid from the last washing is between 55 and 40 µmho per centimeter. Optionally, decrease volume by about 5 ml each washing. Replace the absorbed sodium from the sample by extracting with three 30-ml portions of NH₄OAc solution. Dilute to 100 ml and determine the sodium concentration as described in 6P2a.

**Calculations**

CEC (meq/100 g) = (A/B) x dilution x 10

where:
A = Na from curve (meq/L)
B = Sample weight (g)

Report on oven-dry basis.

**References**

Richards (1954).

---

**KOAc, pH 7.0 (5A4)**

**Procedure**

Proceed as in 5A1 except substitute 1 N KOAc, pH 7.0, for NH₄OAc. Determine potassium with flame photometer.
BaCl₂, pH 8.2 (5A5)

Apparatus

- Leaching tubes
- Flame photometer

Reagents

- Buffer solution. Barium chloride (BaCl₂), 0.5 N, and triethanolamine (TEA), 0.2 N. Adjust to pH 8.2 with HCl. Protect from CO₂ of the air by attaching a drying tube containing soda lime (sodium calcium hydrate) to the air intake.
- Replacement solution. Barium chloride (BaCl₂), 0.5 N. Add 0.4 ml buffer solution per liter and mix. Protect from CO₂ with soda-lime tube.
- Magnesium nitrate (Mg(NO₃)₂), 1 N

Procedure

Transfer a 5-g sample to a leaching tube. For field-moist samples use a sample large enough to give an oven-dry weight of about 5 g. Leach with 50 ml BaCl₂-TEA solution, controlling the leaching rate to give at least 4 hours of soil-solution contact time. Follow with 100 ml BaCl₂ replacement solution, controlling the leaching rate so that the soil and BaCl₂ solutions are in contact for a total of 20 to 24 hours. Rinse walls of leaching tube with 15 to 20 ml H₂O, collecting this washing with leachates from BaCl₂ solutions. Extractable acidity can be determined by using this solution (6H1a). Place leaching tube on a clean flask and wash with methanol until free of chloride ion. For many samples 100 ml methanol is enough, but more methanol may be needed for some soils, particularly those of heavy texture and containing large amounts of hydrous oxides. Leach with 100 ml 0.001 N BaCl₂ to remove methanol.

Disconnect leaching tube and flask, rinse underside of leaching tube, place over a 250-ml volumetric flask, and leach with 100 ml 1 N Mg(NO₃)₂ solution. Control leaching rate to give a soil-solution contact time of 16 hours or more. Rinse walls of leaching tube with 15 to 20 ml H₂O; collect rinse in the Mg(NO₃)₂ leachate. Make to volume.

Barium by Flame Photometry (5A5a)

Make standards in 1 N Mg(NO₃)₂. Determine barium by flame photometry at 489 mū.

Calculations

CEC (meq/100 g) = (A/B) x dilution x 25
where:
A = Ba from curve (meq/L)
B = Sample weight (g)

NH₄OAc, pH 7.0, Leaching Tube (5A6)

Apparatus
- Allihn leaching tubes or 50-ml plastic syringe barrels

Reagents
- Same as in 5A1

Procedure
Prepare the Allihn tubes by placing either filter paper (Reeve Angel No. 934 AH, 3-cm fiber glass) or filter paper pulp on the fritted glass plate. If the syringe barrel is used as a leaching tube, compress the filter paper pulp in the barrel bottom with the syringe plunger. Place a Gooch perforated plate over the filter paper to permit stirring the soil without damage to the filter. (This plate is not necessary if an adequate pulp pad is used.) Place 5 or 10 g soil and a teaspoon of Celite into the tubes. (Optionally place a layer of Celite under the soil.) Add 25 ml N NH₄OAc; stir and leach. Add an additional 25 ml N NH₄OAc and let stand overnight. Stop the leaching with a pinch clamp or by stoppering the leaching tube. Add the NH₄OAc directly to the leaching tube or use a constant level device (fig. 6N3-1). A volumetric flask can be substituted for the 250-ml Erlenmeyer flask and tubing.

Make the leachate to volume if a volumetric flask is used or, if tared suction flasks are used, make to the appropriate calibrated weight for 100 ml NH₄OAc. Set aside the leachate for further analysis. Add about 10 ml ethanol to the soil pad, stir, and leach. Leach with 100 ml ethanol and check for NH₄⁺ in leachate. If NH₄⁺ is present, leach with an additional 100 ml ethanol. Some soils, particularly those containing amorphous material, require as much as 400 ml ethanol to clear the ammonia from the leachate.

Direct Distillation (5A6a)
Transfer soil cake to Kjeldahl flask and determine ammonia as described in 5A1a.

NH₄Cl, pH 7.0, Mechanical Extraction (5A7)
Direct Distillation (5A7a)
Determine ammonia by Kjeldahl distillation as described in 5A1a.
**NH₄OAc, pH 7.0, Automatic Extractor (5A8)**

**Direct Distillation (5A8a)**

**Reagents**
- Sodium chloride (NaCl)
- Antifoam mixture. Mix equal parts of mineral oil and n-octyl alcohol.
- Sodium hydroxide (NaOH), 1 N
- Hydrochloric acid (HCl), 0.2 N, standardized
- Boric acid (H₃BO₃), 4-percent

**Procedure**
Transfer the soil plus filter pulp from methods 5A8 or 5A9 to a Kjeldahl flask. Add 400 ml water and about 10 g NaCl, 5 drops antifoam mixture, a gram or two of granular zinc, and 40 ml of 1 N NaOH. Connect the flask with the condenser and distill 140 ml into 50 ml of 4-percent H₃BO₃ solution in 250-ml titrator beaker. Titrate with automatic titrator to end point pH setting of 4.60.

**Calculations**

\[
\text{CEC (meq/100 g)} = \frac{(A/B) \times N \times 100}{100}
\]

where:
\[
A = \text{Volume HCl (mL)}
\]
\[
B = \text{Sample weight (g)}
\]
\[
N = \text{Normality of acid}
\]

Report on oven-dry basis.

**References**
Peech et al. (1947).

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**NH₄Cl, pH 7.0, Automatic Extractor (5A9)**

**Direct Distillation (5A9a)**

Determine ammonia by Kjeldahl distillation as described in 5A8a.

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**Extractable Bases (5B)**

**NH₄OAc, pH 7.0, Buchner Funnel (5B1)**

**Procedure**
Analyze the NH₄OAc leachate from method 5A1a for calcium, magnesium, sodium, and potassium (methods 6N2, 6O2, 6P2, 6Q2).
Uncorrected (Extractable) (5B1a)

If a soil does not contain soluble salts, the extractable bases are presumed to equal the exchangeable bases. They are, however, reported as extractable bases.

Corrected (Exchangeable) (5B1b)

If a soil contains soluble salts, estimate their amount from the saturation extract as follows. Multiply cation concentration in the saturation extract (meq/L) by the saturation percentage (divided by 1000) to convert to milliequivalents per 100 g. Subtract this quantity from the concentration of the extracted cation. This procedure is not valid for calcium and magnesium in the presence of carbonates or for calcium in the presence of gypsum because these salts are soluble in NH$_4$OAc.

References

Peech et al. (1947).

KCl-Triethanolamine Extraction, pH 8.2 (5B2)

Reagents

- Buffer solution. Potassium chloride (KCl), 1.0 $N$, and triethanolamine (TEA), 0.2 $N$, pH 8.2.

Procedure

Proceed as in 5B1 except leach with 1 $N$ KCl buffered at pH 8.2 with triethanolamine. Determine calcium by method 6N4, magnesium by 6O4.

References

North-Central Regional Research Committee (1955).

KCl-Triethanolamine Extraction, pH 8.2 (Revised) (5B3)

Reagents

- Buffer solution. Potassium chloride (KCl), 1.0 $N$, and triethanolamine (TEA), 0.2 $N$, pH 8.2.

Procedure

Weigh 10-g samples and transfer to 100-ml beakers. Add 40 ml buffer solution. Stir thoroughly at least three times over a period of not less than 1 hour. Filter the suspension and collect the leachate in a 100-ml volumetric flask.
Analyze the leachate for Ca and Mg by an appropriate method (6N4, 6O4).

**Uncorrected (Extractable) (5B3a)**
If a soil does not contain soluble salts, the extractable bases are presumed to equal the exchangeable bases. They are, however, reported as extractable bases.

**Corrected (Exchangeable) (5B3b)**
If a soil contains soluble salts, estimate their amounts from the saturation extract and correct as in 5B1b.

### NH₄OAc, pH 7.0, Leaching Tube (5B4)
Analyze the NH₄OAc leachate from method 5A6 for Ca, Mg, Na, and K (methods 6N2, 6O2, 6P2, 6Q2).

**Uncorrected (Extractable) (5B4a)**
If a soil does not contain soluble salts, the extractable bases are presumed to equal the exchangeable bases. They are, however, reported as extractable bases.

**Corrected (Exchangeable) (5B4b)**
If a soil contains soluble salts, estimate their amounts from the saturation extract and correct as in 5B1b.

### NH₄OAc, pH 7.0, Automatic Extractor (5B5)
**Corrected (Exchangeable) (5B5b)**
If a soil contains soluble salts, estimate their amount from the saturation extract as follows. Multiply cation concentration in the saturation extract (meq/L) by the saturation percentage (divided by 1000) to convert to milliequivalents per 100 g. Subtract this quantity from the concentration of the extracted cation. This procedure is not valid for calcium and magnesium in the presence of carbonates that contain those elements, or for calcium in the presence of gypsum, because these compounds are soluble in NH₄OAc.

### Base Saturation (5C)
**NaOAc, pH 8.2 (5C2)**
Divide sum of NH₄OAc-extracted bases by the exchange capacity determined by method 5A2a.
**Exchangeable Sodium Percentage (ESP) (5D)**
NaOAc, pH 8.2 (5D1)

Divide exchangeable sodium (meq/100 g) by the exchange capacity determined by method 5A2a.

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**Calcium Saturation (Exchangeable-Calcium Percentage) (5F)**
NH₄OAc, pH 7.0 (5F1)

Divide the NH₄OAc-extracted calcium by the exchange capacity determined by method 5A1 or 5A6.

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**Organic Carbon (6A)**

Determine carbon for each horizon that may contain organic matter. Report as carbon percentage by weight of <2-mm material.

To calculate total carbon per unit area, convert these weight percentages to volume percentages. Multiply each value by the bulk density Dbm, where m is usually ⅓ bar or 30 cm, and by the thickness (inches) of that horizon. If coarse fragments are present, further multiply by Cm (4A). Sum the organic-matter percentages and multiply by 0.254 to convert to kilograms of carbon per square meter.

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**Acid-Dichromate Digestion (6A1)**

FeSO₄ Titration (6A1a)

**Reagents**

- Potassium dichromate (K₂Cr₂O₇), 1.00 N (49.04 g per liter)
- Ferrous sulfate, 1.0 N. Dissolve 280 g reagent-grade FeSO₄·7H₂O in water, add 80 ml concentrated H₂SO₄, cool, and dilute to 1 liter. Standardize this reagent each day by titrating against 10 ml N K₂Cr₂O₇ as directed.
- Barium diphenylaminesulfonate indicator, 0.16 percent aqueous solution
- Orthophenanthroline-ferrous complex (optional), 0.025 M solution of one of the phenanthroline-ferrous complex indicators
- H₂SO₄, at least 96-percent
- Phosphoric acid (H₃PO₄), 86-percent

**Procedure**

Transfer 1 g (0.5 g or less if high in organic matter) soil ground to pass an 80-mesh sieve to a 500-ml Erlenmeyer flask. Add 10 ml N K₂Cr₂O₇. Add 20 ml concentrated H₂SO₄ rapidly, directing the stream into the solution. Immediately
swirl vigorously or place in wrist-action shaker for 1 minute. Let the flask stand on a sheet of asbestos for about 30 minutes. Add 200 ml water and 10 ml H₃PO₄. Add 0.5 ml barium diphenylaminesulfonate just before titrating. Titrate by adding FeSO₄ drop by drop to a light green end point. If more than 8 ml of the available 10 ml K₂Cr₂O₇ are reduced, repeat the determination, using less soil. If orthophenanthroline-ferrous complex is the indicator, it is not necessary to add H₃PO₄.

Calculations

\[
\text{Organic carbon (pct.)} = \left( \frac{A-B}{C} \right) \times N \times \left( \frac{0.30}{0.77} \right)
\]

where:
- \( A \) = Volume FeSO₄ blank (mL)
- \( B \) = Volume FeSO₄ sample (mL)
- \( C \) = Sample weight (g)
- \( N \) = Normality of FeSO₄
- 0.77 = Recovery factor proposed by Walkley (1935)

Report on oven-dry basis.

References

Peech et al. (1947) and Walkley (1935).

\textbf{CO₂ Evolution, Gravimetric (6A1b)}

Apparatus

- See figure 6A1b-1.

Reagents

- Digestion-acid mixture. Mix 600 ml concentrated H₂SO₄ and 400 ml 85-percent H₃PO₄.
- Potassium dichromate (K₂Cr₂O₇), reagent grade
- Potassium iodide (KI). Dissolve 100 g KI in 100 ml water.
- Silver sulfate (Ag₂SO₄), saturated aqueous solution
- Concentrated sulfuric acid (H₂SO₄)
- Other reagents: Indicarb or Mikohbite, soda lime, 30-mesh zinc, and anhydrole (anhydrous magnesium perchlorate)

Procedure

Place a soil sample containing 20 to 40 mg carbon (usually 0.5 to 3 g oven-dry soil) in digestion flask and add 1 to 2 g K₂Cr₂O₇. Wash the neck of the flask with 3 ml water and connect the flask to reflux condenser. Attach the weighed Nesbitt
Figure 6A1b-1.—Apparatus for gravimetric organic carbon determination by wet combustion with potassium dichromate (6A1b).

Pour 25 ml digestion-acid mixture into funnel, let it enter the flask, and close the stopcock immediately to prevent loss of CO$_2$. Use digestion-acid mixture to lubricate the funnel stopcock. The tip of the air-delivery tube should extend about 0.5 cm below the surface of the acid during digestion. Adjust the “carrier stream” to a flow rate of one or two bubbles per second and maintain this rate during digestion. Heat with a gas flame of sufficient intensity to bring the sample to boiling in 3 to 4 min. Continue gentle boiling for a total heating period of 10 min (avoid excessive frothing). Heating is too rapid if white fumes of SO$_3$ are visible above the second bulb of the reflux condenser during boiling. At the end of the digestion period, remove the flame and aerate for 10 min at the rate of six to eight bubbles per second. Then close the stopcock on the Nesbitt bulb, disconnect the bulb from the system, and weigh.

Calculations

\[
\text{Organic carbon (pct.)} = ((A - B)/C) \times 27.3
\]
where:
A = Final bulb weight (g)
B = Initial bulb weight (g)
C = Sample weight (g)

Report on oven-dry basis.

References
Allison (1960).

Dry Combustion (6A2)
CO₂ Evolution, Gravimetric I (6A2a)

Apparatus
• See figure 6A2a-1.

Reagents
• Powdered manganese oxide (MnO₂)

Procedure
Place 0.5 to 1.5 g soil that has been ground to 80 mesh in an Alundum boat containing 0.25 g powdered MnO₂. Insert the boat into the quartz tube of the

Figure 6A2a-1.—Apparatus for organic carbon determination by dry combustion, carbon dioxide evolution I (6A2a).
multiple-unit combustion furnace shown. Before inserting the soil, preheat the long part of the quartz tube to 900 °C or more (1000 or 1100 °C) and clear of CO₂ by passing CO₂-free oxygen through the combustion train until the weighing bottle shows a constant weight. While oxygen is passing slowly through the apparatus, heat to a temperature of 900 °C or higher (15 to 30 min). Continue heating in a streaming oxygen atmosphere for 30 minutes more or until the Nesbitt absorption bulb has reached a constant weight.

Calculations
Report on oven-dry basis as in 6A1b.

References
Robinson (1930).

CO₂ Evolution, Gravimetric II (6A2b)

Apparatus
- See figure 6A2b-1.

Procedure
Heat tube to approximately 950 °C. Sweep with oxygen until weight of Nesbitt bulb is constant. Remove rubber stopper in the oxygen inlet end of the tube and insert the boat containing 0.5 to 1.5 g soil. Reinsert the stopper and use the push rod to move the boat into the hot zone. Heat for 10 minutes, remove bulb, and record weight gain. Remove boat and repeat process with fresh sample, using the same Nesbitt bulb.

Calculations
Report on oven-dry basis as in 6A1b.

References
Robinson (1930) and Post. (Post, G.J. A study of three methods for determination of organic carbon in Ohio soils of several great soil groups and the profile distribution of carbon-nitrogen ratios. M.Sc. thesis. The Ohio State University, 34 pp. 1956.)

CO₂ Evolution III (6A2c)

Apparatus
- LECO 70-second carbon analyzer, model 750-100
- LECO induction furnace, model 521-000
Figure 6A2b-1.—Apparatus used for organic carbon determination by dry combustion, carbon dioxide evolution II (6A2b).

Reagents
- Manganese dioxide
- Antimony
- 1-g standard sample rings containing 0.870 percent carbon
- 1-g standard sample rings containing 0.073 percent carbon
- Metal accelerator
- Iron chip accelerator
- Anhydrone

Procedure
For noncalcareous soils, weigh approximately ½ g of <2-mm soil into crucibles in duplicate. Add to the soil in the crucibles one scoop of copper accelerator and one scoop of iron chip accelerator. Mix by stirring. Add an additional scoop of iron chips to the stirred mixture. Four standard soils containing 0.8, 2.1, 3.5, and 6.5 percent organic carbon are run with each group of soils. Follow LECO instruction
 manuals for instrument operation. Record readings from digital voltmeter as percent carbon.

References
Tabatabai and Bremner (1970).

Peroxide Digestion (6A3)
Gravimetric Weight Loss (6A3a)

Reagents
• Hydrogen peroxide (H₂O₂), 6-percent

Procedure
Digest soil for several hours in a covered beaker with 6-percent H₂O₂. Remove soluble material by washing three to five times with a Pasteur-Chamberlain clay filter, "F" fineness. Dry the beaker and soil, and weigh.

Calculations
Organic matter (pct.)=((A+B)/C) x 100

where:
A=Weight loss on heating (g)
B=Weight of dry matter in solution (g)
C=Sample weight (g)

Note that organic matter differs from organic carbon (see 6A1a).

References
North-Central Regional Research Committee on Soils (1955).

 Nitrogen (6B)
Kjeldahl Digestion I (6B1)

Reagents
• Concentrated sulfuric acid (H₂SO₄)
• Salt mixture:
  • Potassium sulfate (K₂SO₄), 1000 g
  • Ferrous sulfate (anhydrous) FeSO₄, 55 g
  • Copper sulfate (anhydrous) CuSO₄, 32 g
  • Hengar granules (selenized)
Procedure
  Weigh 5 g soil into 800-ml Kjeldahl flask, add 20 ml distilled water, and let
stand overnight. Add 10 g salt mixture, 2 or 3 Hengar granules, and 30 ml H₂SO₄.
Digest on Kjeldahl digestion heaters, rotating flasks frequently. Continue digestion
1 hr after mixture is clear.

References
  Association of Official Agricultural Chemists (1945).

Ammonia Distillation (6B1a)

Reagents
  • Mixed indicator. Methyl red, 0.125-percent, and methylene blue,
    0.0825-percent, in 95-percent ethanol.
  • Methyl red (optional), 0.25-percent
  • Brom cresol green, 0.1-percent aqueous solution
  • Boric acid (H₃BO₃), 4-percent
  • HCl, standardized, 0.1 N or 0.05 N

Procedure
  Cool digestion flask (6B1) and dilute contents with about 400 ml water. Add 2 to
3 g mossy zinc, 5 drops antifoam mixture, and 70 ml concentrated NaOH solution.
Connect flask to condenser and distill ammonia into 25 or 50 ml H₃BO₃ solution.
Titrate with standard HCl to purple end point, using 10 drops mixed indicator and 2
drops brom cresol green or 3 drops brom cresol green and 1 drop methyl red.

Calculations
  \[ N (\text{pct}) = \frac{(A - B)}{C} \times N \times 1.4 \]
  where:
  \[ A = \text{Volume HCl sample (mL)} \]
  \[ B = \text{Volume HCl blank (mL)} \]
  \[ C = \text{Sample weight (g)} \]
  \[ N = \text{Normality of HCl} \]
  Report on oven-dry basis.

Ammonia Distillation, Automatic Titrator (6B1b)

Reagents
  • Boric acid (H₃BO₃), 4-percent
  • HCl, standardized, 0.1 N or 0.05 N
• Concentrated sodium hydroxide (NaOH) solution, 50-percent
• Antifoam mixture: Equal parts n-octyl alcohol and mineral oil
• Mossy zinc

**Procedure**

Cool digestion flask (6B1) and dilute contents with about 400 ml water. Add 2 to 3 g mossy zinc, 5 drops antifoam mixture, and 70 ml concentrated NaOH solution. Connect flask to condenser and distill ammonia into 250-ml titrator beaker containing 50 ml H₃BO₃ solution. Titrate with standard HCl to end point pH setting of 4.60 on automatic titrator.

**Calculations**

\[ N \text{ (pct.)} = \frac{(A - B)}{C} \times N \times 1.4 \]

where:
- A = Volume HCl sample (mL)
- B = Volume HCl blank (mL)
- C = Sample weight (g)
- N = Normality of acid

Report on oven-dry basis.

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**Semimicro Kjeldahl (6B2)**

**Apparatus**

- Aminco-Koegel semimicro rotary digestion rack and steam-distillation apparatus

**Reagents**

- Concentrated sulfuric acid (H₂SO₄)
- H₂SO₄, 0.01 N, standardized
- Sodium hydroxide (NaOH), 50-percent
- Boric acid (H₃BO₃), 2-percent
- Mixed indicator. Mix 0.1 g methyl red and 0.1 g brom cresol green and dissolve in 250 ml ethanol.
- Salt mixture. Mix 790 g potassium sulfate (K₂SO₄), 100 g ferrous sulfate (FeSO₄), 100 g copper sulfate (CuSO₄), and 10 g selenium metal.

**Procedure**

Using an analytical balance, weigh on a cigarette paper either 0.500 or 1.000 g oven-dry soil that has been ground to about 0.2 mm. Roll soil in cigarette paper
and drop into a 100-ml digestion-distillation flask. Add 2 g salt mixture 1 ml water, and 5 ml concentrated H$_2$SO$_4$. Swirl vigorously and digest, rotating the flask frequently until fumes are emitted. Continue digestion for at least 1 hour after mixture becomes white. Cool to room temperature and add 15 ml water. Shake until the contents of the flask are thoroughly mixed.

**Ammonia Distillation (6B2a)**

**Procedure**

Measure 10 ml 2-percent H$_3$BOa with an automatic pipette into a 125-ml flask and add 0.5 ml mixed indicator. Place this flask under delivery tube. Connect digestion-distillation flask containing soil digested according to method 6B2 to the distillation unit by the ground-glass connection. Start steam passing through the system and slowly add 15 ml 50-percent NaOH. Distill for 12 minutes, add 0.5 ml more mixed indicator, and titrate the absorbed ammonia with 0.01 N H$_2$SO$_4$.

**Calculations**

\[
N \text{ (pct)} = \frac{A}{B} \times N \times 1.4
\]

where:

- A = Volume H$_2$SO$_4$ (mL)
- B = Sample weight (g)
- N = Normality of H$_2$SO$_4$

Report on oven-dry basis.

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**Iron (6C)**

**Dithionite Extraction (6C1)**

**Reagents**

- Sodium dithionite powder (Na$_2$S$_2$O$_4$)
- Hydrochloric acid (HCl), 10-percent

**Apparatus**

- 8-oz Pyrex nursing bottles or 250-ml flat-bottomed centrifuge bottles

**Procedure**

Place 4 g soil, ground to 80 mesh, in a nursing or centrifuge bottle. Add 4 g Na$_2$S$_2$O$_4$ and 75 ml water. Stopper and shake overnight or for 16 hours. Then adjust the pH to 3.5 to 4.0, if necessary, with 10-percent HCl. Let stand for no less than 1 hour, stirring four or five times. Transfer the suspension to a graduated
cylinder, dilute to 200 ml with water, and mix. Centrifuge or filter a part of the suspension and transfer 50 ml of the clear solution to a 250-ml beaker.

References
Kilmer (1960).

**Dichromate Titration (6C1a)**

**Reagents**
- Hydrogen peroxide ($\text{H}_2\text{O}_2$), 35-percent
- Ammonium hydroxide ($\text{NH}_4\text{OH}$), 1:1
- Hydrochloric acid ($\text{HCl}$), 1:1
- Stannous chloride ($\text{SnCl}_2$). Dissolve 1 g $\text{SnCl}_2$ in 2 to 4 ml concentrated HCl and dilute to 50 ml with water; prepare fresh each time.
- Mercuric chloride ($\text{HgCl}_2$), saturated aqueous solution
- Phosphoric acid ($\text{H}_3\text{PO}_4$), 85-percent
- Potassium dichromate ($\text{K}_2\text{Cr}_2\text{O}_7$), 0.100 $N$, standard
- Barium diphenylaminesulfonate, 0.16-percent aqueous solution

**Procedure**
Add 10 to 15 ml $\text{H}_2\text{O}_2$ (6C1) to the solution to destroy any excess reducing agent. Cover the beaker with a watchglass and warm on a hot plate until the reaction starts. Set the solution aside until the reaction subsides and then boil for 10 to 15 minutes. Add a slight excess of 1:1 $\text{NH}_4\text{OH}$ and boil the solution for 15 to 20 minutes to ensure complete removal of $\text{H}_2\text{O}_2$. Dissolve $\text{Fe(OH)}_3$ by adding 1:1 HCl through the lip of the beaker. Usually 10 to 15 ml are enough. Heat the solution to 90 °C and reduce by adding $\text{SnCl}_2$ by drops, stirring until the yellow color just disappears. Add three to four drops more. Cool the solution to room temperature and add 15 ml $\text{HgCl}_2$ solution all at once. A light silky precipitate of $\text{Hg}_2\text{Cl}_2$ forms if the proper amount of $\text{SnCl}_2$ has been added. Dilute the solution to 100 to 150 ml and add 5 ml $\text{H}_3\text{PO}_4$. Add 10 drops of barium diphenylaminesulfonate and titrate the solution with standard $\text{K}_2\text{Cr}_2\text{O}_7$ to a violet-blue end point.

**Calculations**

$$\text{Fe (pct.)} = \frac{(A/B) \times N \times (C/D) \times 5.58}{\text{where:}}$$

- $A$ = Volume $\text{K}_2\text{Cr}_2\text{O}_7$ (mL)
- $B$ = Sample weight (g)
- $N$ = Normality of $\text{K}_2\text{Cr}_2\text{O}_7$
- $C$ = Volume extract (mL)
- $D$ = Volume aliquot (mL)
\[ \text{Fe}_2\text{O}_3 \text{ (pct.)} = \text{Fe (pct.)} \times 1.43 \]

Report on oven-dry basis.

**EDTA Titration (6C1b)**

**Reagents**
- Hydrogen peroxide (H\(_2\)O\(_2\)), 35-percent.
- Ammonium persulfate ((NH\(_4\))\(_2\)S\(_2\)O\(_8\)).
- Salicylic acid, 1-percent in 95-percent ethanol
- EDTA, standardized as g iron per ml EDTA. Prepare EDTA as described in 6N1a.
- Iron standard, 0.500 g iron per liter

**Procedure**

Pipette a 5- to 25-ml aliquot from the centrifuge tube of method 6C1 into a 250-ml beaker. Add 50 ml water to the beaker. Then add by drops 5 ml H\(_2\)O\(_2\) and digest over low heat until bubbling from the decomposing H\(_2\)O\(_2\) ceases. Remove immediately to avoid precipitation of Fe\(_2\)O\(_3\) in samples high in iron. Caution: Add H\(_2\)O\(_2\) slowly to prevent liberation of elemental sulfur from any remaining Na\(_2\)S\(_2\)O\(_4\). Keep the volume in the beaker to about 50 ml during the digestion by adding water if necessary. Remove from heat and cool. Adjust the pH between 2.0 and 3.0 with a pH meter, using either concentrated acetic acid or a 20-percent NaOAc solution. Add a few milligrams (NH\(_4\))\(_2\)S\(_2\)O\(_8\) to the solution to ensure total oxidation of iron. Then add 1 ml indicator (1-percent salicylic acid) and titrate with 0.02 N EDTA to a pale yellow or colorless end point.

**Calculations**

\[ \text{Fe (pct.)} = \frac{A \times V \times (C/D)}{B} \times 100 \]

where:
- \(A\) = Volume EDTA (mL)
- \(B\) = Sample weight (g)
- \(V\) = Titer of EDTA in g Fe/ml EDTA
- \(C\) = Volume extract (mL)
- \(D\) = Volume aliquot (mL)

\[ \text{Fe}_2\text{O}_3 \text{ (pct.)} = \text{Fe (pct.)} \times 1.43 \]

Report on oven-dry basis.

**References**

Cheng, Bray, and Kurtz (1953).
Dithionite-Citrate Extraction (6C2)

Reagents
- Sodium dithionite ($Na_2S_2O_4$)
- Sodium citrate
- Superfloc flocculating agent, 0.2 percent in water

Procedure
Weigh 1 to 4 g of soil (approximately 0.2 g maximum extractable iron) into an 8-oz nursing bottle. Add 2 g sodium dithionite and 20 to 25 g sodium citrate. Make up to 4 oz with water, and shake overnight in a reciprocating shaker. Add 2 ml Superfloc solution to the suspension, make up to 8 oz with water, shake vigorously for 15 s, and allow to settle for at least 1 hr. This extract is used for analysis of iron (6C2b), aluminum (6G7a), and manganese (6D2a).

References
Holmgren (1967).

Orthophenanthroline Colorimetry (6C2a)

Apparatus
- Seligson pipette, 0.1-ml

Reagents
- Orthophenanthroline, 0.25-percent
- Iron solution, 1000 mg per liter, standard
- Sodium dithionite powder ($Na_2S_2O_4$)
- Sodium citrate crystals
- Superfloc flocculating agent, 0.2-percent, in water

Procedure
Add 5 drops Superfloc solution to the dithionite-treated soil suspension (6C2) and make to 8 oz. Shake vigorously for about 15 seconds and allow to settle. Pipette a 0.1-ml aliquot with a Seligson pipette into a 25-ml volumetric flask. Add water to about 10 ml. Using a small scoop, tap a pinch of dithionite and a pinch of sodium citrate into the flask. Add 0.5 ml 0.25-percent orthophenanthroline and make to volume. Shake and read in a colorimeter at 508 Mµ after 1 hour. To prepare the standards, pipette 5-, 10-, 25-, 50-, 100-, 150-, and 200-ml aliquots of standard iron solution (1000 mg/L) into 8-oz shaking bottles and make to 8 oz after adding reagents as in 6C2. Transfer 0.1-ml aliquots to 25-ml volumetrics and develop color by the above procedure.
Plot the standard curve as milligrams iron per 8-oz bottle against percentage transmission.

**Calculations**

\[
\text{Fe (pct.)} = \frac{A}{B} \times 10^{-1}
\]

- \(A\) = Fe in bottle (mg)
- \(B\) = Sample weight (g)

\[
\text{Fe}_2\text{O}_3 \text{ (pct.)} = \text{Fe (pct.)} \times 1.43
\]

Report on oven-dry basis.

**References**

Holmgren (1967).

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**Dithionite-Citrate-Bicarbonate Extraction (6C3)**

**Reagents**

- Sodium bicarbonate (NaHCO\(_3\)), 1 \(M\)
- Sodium citrate, 0.3 \(M\)
- Sodium chloride (NaCl), saturated solution
- Acetone

**Procedure**

Weigh 4 g soil (1 g clay) into a 100-ml centrifuge tube. Add 40 ml 0.3 \(M\) Na-citrate and 5 ml 1 \(M\) NaHCO\(_3\). Bring temperature to 80 °C in water bath. Add 1 g solid Na\(_2\)S\(_2\)O\(_4\), stir constantly for 1 minute and occasionally for 15 minutes. Add 10 ml NaCl solution and 10 ml acetone to promote flocculation. Mix, warm in water bath, and centrifuge 5 minutes at 1,600 to 2,200 rpm. Decant clear supernatant into 500-ml volumetric flask and make to volume.

**References**

Mehra and Jackson (1960).

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**Potassium Thiocyanate Colorimetry (6C3a)**

**Apparatus**

- Colorimeter

**Reagents**

- Hydrochloric acid (HCl), 6 \(N\)
• Potassium thiocyanate (KSCN), 20-percent
• Hydrogen peroxide (H₂O₂), 30-percent

Procedure
Transfer suitable aliquot (0.5 to 3 ppm iron in final solution) to 50 ml-volumetric flask. Add water to 35 ml, 1 drop H₂O₂, 5 ml HCl, and 5 ml KSCN solution. Make to volume and read at 490 mµ in colorimeter.

Calculations
Fe (pct.) = (A/B) x (C/D) x 0.005
where:
A = Fe from curve (mg)
B = Sample weight (g)
C = Volume extract (mL)
D = Volume aliquot (mL)

Fe₂O₃ (pct.) = Fe (pct.) x 1.43

Report on oven-dry basis.

References
Jackson (1956).

Pyrophosphate-Dithionite Extraction (6C4)

Reagents
• Pyrophosphate solution. Dissolve 89.2 g Na₄P₂O₇•10H₂O in 800 to 900 ml water. Adjust the pH of this solution to 8.0 by adding hydrogen saturated exchange resin. Decant or filter, wash the resin, and dilute the solution to 1000 ml to make 0.2 M Na₄P₂O₇.
• Sodium dithionite (Na₂S₂O₄)
• Digestion acid. 10 parts concentrated HNO₃, 4 parts concentrated H₂SO₄, and 4 parts concentrated HClO₄.

Procedure
Mix 80 ml pyrophosphate solution and 2.0 g solid sodium dithionite in a beaker and add this solution to 4 g soil in a centrifuge tube (pH 8.0 pyrophosphate solution and dithionite combined in this ratio result in a solution having a pH of about 7.3). Continue the extraction for 30 minutes at 50 °C, shaking the suspension in the tube every 5 minutes. Centrifuge the suspension 5 to 10 minutes at 2000 rpm. Dilute the extract to 100 ml (solution A).
Immediately transfer 5 ml solution A to a beaker. Add 1 to 2 ml digestion acid and heat on a hot plate until almost dry to destroy the organic and hydrolyze pyrophosphate to orthophosphate. Allow to cool, dissolve the salts in HCl, and dilute to 100 ml (solution B). Determine iron and aluminum in solution B by appropriate methods, such as 6C3a and 6G1a.

References
Franzmeier, Hajek, and Simonson (1965).

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**Sodium Pyrophosphate Extraction (6C5)**

**Reagents**
- Sodium pyrophosphate \( (\text{Na}_4\text{P}_2\text{O}_7) \), 0.1 \( M \)
- Superfloc solution, 0.4 percent

**Procedure**
Place 2 g soil into 250-ml centrifuge bottle (polypropylene). Add 200 ml 0.1 \( M \) \( \text{Na}_4\text{P}_2\text{O}_7 \), cap, and shake overnight. Add 5 to 10 drops 0.4-percent Superfloc, shake, and centrifuge at 2000 rpm (Int. No. II centrifuge). Transfer the supernatant liquid to a plastic or glass container and reserve for Fe and Al analyses.

The supernatant liquid must be clear in reflected light. If fine colloids are visible, repeat the procedures. If fine colloids are still present, spin the suspension in a super centrifuge until the supernatant liquid is clear. Foam rubber can be used in the centrifuge cups as a cushion for the 250-ml flat-bottom plastic bottles.

References
Bascomb (1968).

**Atomic Absorption (6C5a)**

**Apparatus**
- Atomic absorption spectrophotometer

**Reagents**
- Standard Fe solution, 0 to 50 ppm

**Procedure**
Establish standard curve and match readings from extract to curve readings. Dilute where necessary.
Calculations

\[ \text{Fe (pct.)} = A \times \frac{B}{C} \times \frac{1}{10,000} \times \text{dilution} \]

where:
\[ A = \text{Fe (ppm)} \]
\[ B = \text{Volume extract (mL)} \]
\[ C = \text{Sample weight (g)} \]

Report on oven-dry basis.

Ammonium Oxalate Extraction (6C6)

Reagents
- Ammonium oxalate \((\text{NH}_4)_2\text{C}_2\text{O}_4\) 0.2 \(M\), pH 3.0
- Adjust the pH of 0.2 \(M\) \((\text{NH}_4)_2\text{C}_2\text{O}_4\) to 3.0 with 0.2 \(M\) oxalic acid \((\text{H}_2\text{C}_2\text{O}_4)\).
- Superfloc solution, 0.4 percent

Procedure
Place 2 g soil into 250-ml centrifuge bottle (polypropylene). Add 200 ml 0.2 \(M\) \((\text{NH}_4)_2\text{C}_2\text{O}_4\), cap, and shake immediately in the dark for 4 hours. Add 5 to 10 drops 0.4-percent Superfloc, shake, and centrifuge at 2000 rpm (Int. No. II centrifuge). Transfer the supernatant liquid to a plastic or glass container. Store in the dark and reserve for Fe and Al analyses.

The supernatant liquid must be clear in reflected light. If fine colloids are visible, repeat the procedure. If fine colloids are still present, spin the suspension in a supercentrifuge until the supernatant liquid is clear.

References
McKeague and Day (1966).

Atomic Absorption (6C6a)

Proceed as in 6C5a except use extract from 6C6.

Manganese (6D)

Dithionite Extraction (6D1)

Extract 4.00 g soil as described in 6C1.
Permanganate Colorimetry (6D1a)

Reagents
- Concentrated nitric acid (HNO₃)
- Hydrogen peroxide (H₂O₂), 30-percent
- Phosphoric acid (H₃PO₄), 85-percent
- Sodium para periodate (Na₃H₂IO₆) or sodium meta periodate (NaIO₄). 
- Purified water diluent. Add 100 ml 80-percent H₃PO₄ and 1 g Na₃H₂IO₆ to 1 liter water (Mn-free); heat to boiling and digest for 1 hour; stopper with foil-covered stopper. About 85 ml of this diluent is needed for each sample.
- KMnO₄, standard

Procedure
Take a 10- to 25-ml aliquot from the dithionite extract and place in a 150-ml beaker. Add 5 ml 30-percent H₂O₂, digest on hot plate, and evaporate until dry. Cool beaker and contents and add 3 ml concentrated HNO₃ and 2 ml 30-percent H₂O₂. Digest on hot plate for 30 minutes, using a close fitting cover glass, then raise cover glass, and evaporate until dry. Take up residue with 10 ml 85-percent H₃PO₄, heat to boiling, remove, and cool to about 50 °C. Dilute with 10 ml water and add 0.2 g Na₃H₂IO₆. Cover beaker and heat to boiling. Cool to 50 °C and add 62 ml purified water diluent and 0.1 g Na₃H₂IO₆. Digest at 90 °C for 40 minutes or until no further color develops. Transfer the hot solution to a 100-ml volumetric flask, using purified water diluent to rinse the beaker. Cool, make up to volume with the diluent, stopper, and shake. Determine percentage transmittance with a photoelectric colorimeter at 540 mµ. Interpolate concentration from a standard absorbance concentration.

Calculations

\[ \text{Mn (pct.)} = \frac{A}{B} \times \frac{C}{D} \times 54.9 \]

where:

- \( A = \text{MnO}_4^- \) (meq/L)
- \( B = \text{Sample weight (g)} \)
- \( C = \text{Volume extract (mL)} \)
- \( D = \text{Volume aliquot (mL)} \)

\[ \text{MnO (pct.)} = \text{Mn (pct.)} \times 1.291 \]

Report on oven-dry basis.

References

Jackson (1956).
Calcium Carbonate (6E)
HCl Treatment (6E1)
   Gas Volumetric (semiquantitative) (6E1a)

This procedure uses a simple leveling device to measure the volume of gas released when the soil is treated with HCl. It has an inherent error caused by the solubility of CO₂ in the HCl solution. Data on file at the laboratory at Lincoln, Nebraska, indicate that the results are about 10 percent (8 to 12 percent) low for CaCO₃ equivalents ranging from 40 percent to 6 percent (1-g basis). For 1-percent equivalents the values are about 20 percent low and for less than 1 percent, the values have doubtful significance.

References
   Association of Official Agricultural Chemists (1945).

Manometric (6E1b)

Apparatus
   • Wide-mouth prescription bottles, 3-oz, with bakelite cap; drill 7/16-in hole in cap for serum bottle stopper. Rubber gasket, 1⅜ in OD x 15/16 in ID.
   • Serum bottle stopper
   • Mercury manometer and a 26-gauge hypodermic needle attached to manometer tube
   • Gelatin capsule, ¼ oz

Reagents
   • Hydrochloric acid (HCl), 6 N
   • Glycerin

Procedure
   Place 2 g of soil in prescription bottle and add 5 ml water. Moisten lip of bottle with a drop of glycerin to ensure a good seal with rubber gasket. Fill gelatin capsule with HCl, put cap in place, and invert to seal cap on capsule. Place capsule in bottle and immediately cap the bottle. In a minute or two the HCl will dissolve the capsule. After 1 hr insert hypodermic needle through serum stopper and read manometer. Compare reading with those for standards prepared by treating aliquots of standard Na₂CO₃ solution in same manner as samples.

   Vary sample weight according to CaCO₃ content as follows: For <25 percent CaCO₃, use 2 g soil; for 25 to 50 percent CaCO₃, 1 g soil; and for >50 percent CaCO₃, 0.5 g soil. For trace amounts, add a few drops 6 N HCl to soil and observe under binocular microscope. Evolution of gas bubbles indicates the presence of CaCO₃.
References
Williams (1948).

Gravimetric (weight loss) (6E1c)

Apparatus
- See figure 6E1c-1.

Reagents
- Hydrochloric acid (HCl), 6 N
- Anhydrone (Mg(ClO₄)₂)

Procedure
Assemble apparatus as shown in figure 6E1c-1. Place a sample of soil containing less than 1 g CaCO₃ equivalent in a 125-ml Erlenmeyer flask. Wash down the sides of the flask with 10 ml water. Place 7 ml 6 N HCl into vial C and then place the vial upright in the flask without spilling any acid. Moisten stopper G with glycerin, sprinkle with a small amount of 180-mesh abrasive to overcome slipperiness, and place the apparatus with stopcocks D and E, tubes I and J, attached firmly in position in the flask. Close stopcocks D and E. Place the apparatus beside the balance. Wait 30 minutes before weighing to allow temperature of the apparatus to equilibrate with temperature of air in the balance. Do all weighing with stopcock D open since a change in temperature of the flask with the stopcock closed results in a change in weight of the apparatus. Use tongs to place apparatus on the weighing pan, open stopcock D, weigh to 0.1 mg, and then immediately close stopcock D. Check the weight 10 minutes later to be certain that the weight of the flask has stabilized. Open stopcock D and then shake apparatus to upset the vial, allowing the acid to react with the carbonates. After 10 minutes, attach the rubber tube from the air-drying vessel to stopcock E. Open stopcock E and apply suction at stopcock D to give 5 to 10 bubbles per second at the base of tube J to sweep out CO₂. Shake the flask after 10 minutes and again after 20 minutes. After 30-minutes sweeping time, stop the suction and close stopcocks D and E. Return apparatus to the balance. Delay weighing for 1 hour to allow the heat generated by absorption of water by the anhydrone to be dissipated. Weigh apparatus with stopcock D open. Check the weight after 10 minutes.

Calculations
Carbonate as CaCO₃ (pct.) = ((A − B)/C) x 228
where:
A = Initial weight of flask (g)
B = Final weight of flask (g)
C = Sample weight (g)

Report on oven-dry basis.
Figure 6E1c-1.—Apparatus used for calcium carbonate determination by weight loss (6E1c).

References
Erickson et al. (1947).
Gravimetric (weight gain) (6E1d)

Proceed as in 6E1c except add additional trap containing CO$_2$-absorbing Ascarite to end of gas train. Weigh Ascarite bulb before and after CO$_2$ evolution. Weight gain equals the CO$_2$ evolved from the sample. Better results are obtained if the Ascarite is size-graded so that CO$_2$ passes through the coarser material first. Indicarb can be used in place of Ascarite.

Titrimetric (6E1e)

Reagents
- Hydrochloric acid (HCl), 0.5 N, standardized
- Sodium hydroxide (NaOH), 0.25 N, standardized
- Phenolphthalein, 1 percent in 60-percent ethanol

Procedure
Place 5 to 25 g soil in a 150-ml beaker, add exactly 50 ml HCl, cover with a watchglass, and boil gently for 5 minutes. Cool, filter, and wash all the acid from the soil with water. Determine the amount of unused acid by adding 2 drops of phenolphthalein and back-titrating with NaOH.

Calculations

\[ \text{Carbonate as CaCO}_3 (\%) = \frac{(50 \times A - B \times C)}{D} \times 5 \]

where:
- A = Normality of HCl
- B = Volume NaOH (mL)
- C = Normality of NaOH
- D = Sample weight (g)

Report on oven-dry basis.

References
Richards (1954).

Warburg Method (6E1f)

Apparatus
- Warburg manometer, mercury filled
- Warburg reaction vessel, 15-ml capacity, with vented stopper for sidearm
- Constant temperature bath

Reagents
- HCl, 1:1. Na$_2$CO$_3$ solution for standard curve. Dissolve 1.06 g Na$_2$CO$_3$ in
water and make to 1 liter. Solution contains 1.06 mg Na$_2$CO$_3$ per ml or the equivalent of 1 mg CaCO$_3$ per ml. Obtain standard curve by measuring CO$_2$ pressure from 1, 2, 4, 6, 8, and 10 ml Na$_2$CO$_3$ solution.

**Procedure**

Weigh 100 mg sample of finely ground soil and transfer to Warburg reaction vessel. Be careful not to get any sample in center well. Pipette 1 ml water into vessel and mix well with sample. Pipette 1 ml 1:1 HCl into sidearm, insert greased stopper, and leave in vent-open position. Attach reaction vessel to manometer and fasten with rubber bands or spring supports. Place reaction vessel in constant temperature bath at 25 °C for 5 to 10 minutes to bring flask contents to temperature of water bath. Remove flask from bath, close stopper vent, and fasten with rubber bands or springs. Tilt flask to allow acid to flow from sidearm into reaction vessel, mix contents, and return vessel to water bath. Let stand for at least 30 minutes before reading manometer. Use the standard curve to convert the difference between the two manometer arm readings (mm) to milligrams CaCO$_3$. Gently tap the manometer holder occasionally to prevent low readings caused by mercury adhering to manometer walls.

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**Sensitive Qualitative Method (6E2)**

**Visual, Gas Bubbles (6E2a)**

Add few drops 6 $N$ H$_2$SO$_4$ to soil and observe under binocular microscope. Evolution of gas bubbles indicates the presence of CaCO$_3$.

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**H$_2$SO$_4$ Treatment (6E3)**

**Gravimetric (weight gain) (6E3a)**

**Apparatus**

- See figure 6A1b-1.

**Reagents**

- Sulfuric acid (H$_2$SO$_4$). Dissolve 57 ml concentrated H$_2$SO$_4$ and 92 g of FeSO$_4$•7H$_2$O in 600 ml water, cool, and dilute to 1000 ml. This solution is approximately 2 $N$ in acidity and contains 5-percent FeSO$_4$ as antioxidant. Keep well stoppered.

**Procedure**

Place a 1- to 5-g sample of oven-dry soil in the digestion flask E and connect condenser D. Weigh the Nesbitt bulb, attach to the system, and adjust the carrier stream to a flow rate of 1 or 2 bubbles per second. Pour 25 ml of the acid
solution into the funnel and let it enter the digestion flask E. Close the stopcock immediately. Apply heat slowly and bring contents of flask to a boil in about 4 minutes. Continue gentle boiling for exactly 3 minutes more for a total heating period of 7 minutes. Remove the flame, adjust the carrier stream to 6 or 8 bubbles per second, and continue aerating for 10 minutes. Disconnect the Nesbitt bulb and weigh.

**Calculations**

Carbonate as CaCO$_3$ (%) = ((A - B) / C) x 227

where:
- A = Final weight of bulb (g)
- B = Initial weight of bulb (g)
- C = Sample weight (g)

Report on oven-dry basis.

**References**

Allison (1960).

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**Gypsum (6F)**

**Water Extract (6F1)**

**Indirect Estimate (6F1b)**

Add a weighed quantity of soil to enough water to dissolve all the gypsum by overnight shaking. The concentration of sulfate in this dilute soil:water extract should be <10 meq/L. Gypsum can be estimated by method 6F2. If crystals are observed or estimated gypsum content is >5 percent, the <2-mm sample should be ground to approximately 80 mesh. Determine total sulfate in this extract by any appropriate procedure. Also determine Ca and SO$_4$ in a saturation extract by any appropriate procedure.

**Calculations**

Gypsum = (SO$_4$)$_{DE}$ - (SO$_4$)$_{non-gypsum \ SE}$

but SO$_4$$_{non-gypsum \ SE}$ = (SO$_4$)$_{SE}$ - (SO$_4$)$_{gypsum \ SE}$

∴ gypsum = (SO$_4$)$_{DE}$ + (SO$_4$)$_{gypsum \ SE}$ - (SO$_4$)$_{SE}$

(SO$_4$)$_{DE}$ = SO$_4$ in dilute water extract

(SO$_4$)$_{SE}$ = SO$_4$ in saturation extract
(SO₄)ₓ gypsum SE = 30 meq/L if SO₄ and Ca are ≥ 30 meq/L

= (SO₄)ₓ SE if (Ca)ₓ SE > (SO₄)ₓ SE

= (Ca)ₓ SE if (Ca)ₓ SE < (SO₄)ₓ SE

All quantities are reported in meq/100 g.

Gypsum (%) = \text{Gypsum (meq/100 g)} \times 0.0861 \text{ (g/meq)}

References
Lagerwerff, Akin, and Moses (1965).

Ion Chromatograph (6F1c)

Apparatus
- DIONEX Model 2110i ion chromatograph
- Recorder (1 volt input)
- Voltage stabilizer

Reagents
- \(M\) \(Na_2CO_3\)
- \(0.003 M\) \(NaHCO_3\)
- \(0.0024 M\) \(Na_2CO_3\)
- \(N\) \(H_2SO_4\)
- Mixed standard solutions:
  - Fluoride 0.0125 to 5.0 meq/L
  - Chloride 0.01 to 4.0 meq/L
  - Nitrate 0.025 to 10.0 meq/L
  - Sulfate 0.05 to 20.0 meq/L

All solutions are filtered through a polycarbonate membrane having 0.4-µm pore size. Soil extracts are filtered with a disposable filter unit (Millix™) having 0.22-µm pore size.

Procedure
The soil extract is obtained as described in 6F1a. Fill a plastic syringe (3 to 10 cc) with a solution having a concentration within the range of the sulfate standard. Baseline is established using a full-scale µmhos setting of 3 before each determination. This setting is adjusted as needed, keeping in the range used for making the determinations on the mixed standard. Peak height readings are made on the mixed standard using eight concentrations. A curve fitting linear regression equation \([y(\text{meq/L}) = a_1 \text{(PKH)} + a_0]\) is established for the sulfate standards. Sulfate concentration in the soil extracts is determined by this equation.
Calculations
See 6F1b.

Weight Loss (6F2)

Apparatus
- Vacuum desiccator
- Aluminum dish
- Balance, 0.001-g sensitivity

Reagents
- Phosphorus pentoxide ($P_2O_5$)

Procedure
Place about 10 g of soil in a tared (Wt A) aluminum dish. Saturate sample with water and let stand overnight to air-dry. Place in a vacuum desiccator with $P_2O_5$ desiccant. Evacuate desiccator and allow to stand 48 hr. Remove dish from desiccator and weigh (Wt B), then place in oven at 105 °C for 24 hr. Allow dish to cool in desiccator and weigh (Wt C).

Calculations

\[
\text{Gypsum} \text{ (\%)} = \frac{(\text{WtB} - \text{WtC}) \times 100}{(\text{WtB} - \text{WtA}) \times 0.1942}
\]

The theoretical crystal water content of gypsum is 20.91 percent. However, Nelson et al. have determined that, in practice, this content averages 19.42 percent.

References

Gypsum Requirement (6F5)
The amount of gypsum needed to replace all of the sodium on the exchange complex with calcium is the gypsum requirement.

Reagents
- Saturated gypsum solution. Place about 25 g gypsum (CaSO$_4$•2H$_2$O) in 5 L water in a large flask, stopper, and shake by hand periodically for 1 hr or more. Let settle and decant through a filter into storage bottle. Determine calcium concentration by titration of an aliquot with standard EDTA solution using Eriochrome black T as indicator.
EDTA solution. Dissolve 1.25 g di-sodium ethylenediamine tetraacetate in water and dilute to 1 L. Standardize against solutions containing known concentrations of Ca and Mg.

Buffer solution. Dissolve 6.75 g ammonium chloride in about 400 ml water. Add 570 ml concentrated ammonium hydroxide and dilute to 1 L with distilled water.

Eriochrome black T indicator. Dissolve 1 g Eriochrome black T in 100 ml triethanolamine.

Procedure

Weigh 5 g soil into flask, add 100 ml saturated gypsum solution, stopper, and shake for 5 min in mechanical shaker. Filter through folded filter paper, discarding the first few milliliters of filtrate, which may be cloudy. Pipette a 5-ml aliquot of filtrate into a 125-ml Erlenmeyer flask and dilute to 25 or 30 ml with distilled water. Add 10 drops of buffer solution, 2 drops Eriochrome black T indicator, and titrate with standard EDTA solution to blue end point.

Calculations

Gypsum requirement (meq/100 g) = (A − B) x 2

where:
A = Ca concentration of gypsum solution (meq/L)
B = Ca + Mg concentration of filtrate (meq/L)

References
Richards (1954).

Aluminum (6G)
KCl Extraction I (30 min) (6G1)

Reagents

Potassium chloride (KCl), 1 N

Procedure

Weigh 10-g soil samples into 125-ml Erlenmeyer flasks. Add 50 ml 1 N KCl to each flask, mix several times, and let stand for 30 minutes. Filter through 5.5-cm Whatman No. 42 filter paper in Buchner funnel, using suction as necessary. Leach each sample as rapidly as possible with about five 9-ml portions of KCl, using the first to help transfer the remaining soil in the Erlenmeyer flasks to the Buchner funnels. Transfer the extract to 100-ml volumetric flasks and dilute to
volume with the extracting solution. Or use Allihn leaching tubes and bring to standard weight in tared suction flasks.

References
Lin and Coleman (1960) and Pratt and Bair (1961).

**Aluminon Colorimetry I, Hot Color Development (6G1a)**

**Reagents**
- Thioglycolic acid (HSCH\(_2\)COOH). Dilute 1 ml purified acid to 100 ml with water.
- Aluminon reagent. Dissolve in separate containers 0.75 g Aluminon (ammonium aurine tricarboxylate), 15 g gum acacia, and 200 g NH\(_4\)OAc crystals. To the NH\(_4\)OAc solution add 189 ml concentrated HCl, then the gum acacia, and finally the Aluminon. Mix, filter, and dilute to 1500 ml with water. To get the gum acacia in suspension, add slowly to boiling water while stirring constantly.
- Aluminum standard. Add 2.24 g AlCl\(_3\)•6H\(_2\)O per liter of water. This solution should be nearly 250 ppm aluminum. Check concentration of an aliquot containing 10 ppm aluminum by analyzing for chloride.

**Procedure**
If samples contain less than 5 meq per 100 g aluminum, pipette a 1-ml aliquot of each extract into numbered and calibrated test tubes. If more aluminum is present, dilute before the aliquot is taken. Dilute to approximately 20 ml with distilled water. Add 2 ml dilute thioglycolic acid to each tube, stopper, and shake all the tubes. Pipette 10 ml Aluminon into each tube and dilute to exactly 50 ml. The pH should be between 3.7 and 4.0. Stopper and shake all tubes. Place tubes in a rack and heat in a boiling-water bath for 4 minutes. Cool in running water to room temperature. Transfer samples to reading tubes and measure light transmittance at 535 m\(\mu\) and compare with a standard curve.

**Calculations**

\[
\text{Al (meg/100 g)} = \frac{A}{B} \times \frac{C}{D} \times \frac{9}{5}
\]

where:
- A = Al from curve (mg/L)
- B = Sample weight (g)
- C = Volume extract (mL)
- D = Volume aliquot (mL)

Report on oven-dry basis.
References
Chenery (1948) and Yoe and Hill (1927).

**Aluminon Colorimetry II, HCl Predigestion (6G1b)**

**Procedure**
Proceed as in 6G1a but first add 3 ml $N$ HCl to the aliquot and heat for 30 minutes at 80 to 90 °C.

References
Hsu (1963).

**Aluminon Colorimetry III, Overnight Color Development (6G1c)**
Proceed as in 6G1a except eliminate boiling-water bath, adjust pH to 4.0, and allow color to develop overnight before reading.

**Fluoride Titration (6G1d)**

**Reagents**
- Potassium fluoride (KF), 1 $N$. Titrate with NaOH to a phenolphthalein end point. This eliminates the need for a blank correction in the Al titration.
- Sodium hydroxide (NaOH), 0.1 $N$, standardized
- Sulfuric acid ($\text{H}_2\text{SO}_4$), 0.1 $N$, standardized
- Phenolphthalein, 0.1 percent

**Procedure**
Add 6 to 8 drops phenolphthalein to the leachate in the suction flask (6G1). Titrate with standard NaOH to a pink color that persists for 30 seconds or more. Correct for a KCl blank to obtain KCl extractable acidity. Then add 10 ml KF, and titrate with standard $\text{H}_2\text{SO}_4$ until the pink color disappears. Set aside while other samples are titrated and then complete to a lasting colorless end point. If there is a considerable amount of Al, add a few more drops of phenolphthalein.

**Calculations**
\[
\text{Acidity (meq/100 g)} = \frac{A}{B} \times N \times 100
\]
where:
- $A =$ Volume NaOH (mL)
- $B =$ Sample weight (g)
- $N =$ Normality of NaOH

\[
\text{Al (meq/100)} = \frac{A}{B} \times N \times 100
\]
where:
A = Volume H₂SO₄ (mL)
B = Sample weight (g)
N = Normality of H₂SO₄

References
Yuan (1959).

Atomic Absorption (6G1e)

Apparatus
- Perkin-Elmer Model 290 atomic absorption spectrophotometer with nitrous oxide burner attachment

Reagents
- Standard Al solution, 0 to 5 meq per liter

Procedure
Dilute sample to within range of standard curve. Compare absorbance with standard curve.

Calculations

\[
\text{Al (meq/100 g)} = \frac{A}{B} \times \text{dilution} \times \frac{C}{10}
\]

where:
A = Al from curve (meq/L)
B = Sample weight (g)
C = Volume extract (mL)

KCl Extraction II, Overnight (6G2)
Weigh 10 g soil into 125-ml Erlenmeyer flask. Add 50 ml 1 N KCl and let stand overnight. In the morning transfer to filter funnels and leach with an additional 50 ml KCl.

Aluminon Colorimetry I (6G2a)
Follow procedure for aluminum analysis described in 6G1a.

NH₄OAc Extraction (6G3)
Prepare soil as described in 5A1.
Aluminon Colorimetry III (6G3a)
Follow method of 6G1c.

NaOAc Extraction (6G4)
Prepare soil as described in 5A2.

Aluminon Colorimetry III (6G4a)
Follow method of 6G1c.

Sodium Pyrophosphate Extraction (6G5)
Prepare extract as described in 6C5.

Atomic Absorption (6G5a)

Apparatus
- Atomic absorption spectrophotometer

Reagents
- Standard Al solution, 0 to 50 ppm or 0 to 160 ppm

Procedure
Establish standard curve and match readings from extract to curve readings. Dilute where necessary.

Calculations
\[ \text{Al (pct.)} = A \times (B/C) \times (1/10,000) \times \text{dilution} \]
where:
- \( A = \text{Al (ppm)} \)
- \( B = \text{Volume extract (mL)} \)
- \( C = \text{Sample weight (g)} \)

Report on oven-dry basis.

Ammonium Oxalate Extraction (6G6)
Prepare extract as described in 6C6.

Atomic Absorption (6G6a)
Analyze extract as described in 6G5a.
**NH₄Cl, Automatic Extractor (6G8)**
Prepare extract as described in 5A9.

**Atomic Absorption (6G8a)**

**Apparatus**
- Atomic absorption spectrophotometer

**Reagents**
- Standard Al solutions, 0 to 6 meq/L

**Procedure**
Compare absorbance of samples from 5A9 with that of standards at 309.3 nm, diluting if necessary.

**Calculations**

\[ \text{Al (meq/100 g)} = \frac{A}{B} \times \text{dilution} \times \frac{C}{10} \]

where:
- \( A \) = Al (meq/L)
- \( B \) = Sample weight (g)
- \( C \) = Volume extract (mL)

Report on oven-dry basis.

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**Extractable Acidity (6H)**
**BaCl₂-Triethanolamine I (6H1)**
Extractable acidity data are reported on some data sheets as exchange acidity and on others as extractable H⁺.

**Reagents**
- Buffer solution. Barium chloride, 0.5 \( N \), and triethanolamine, 0.2 \( N \). Add about 90 ml 1 \( N \) HCl per liter to adjust pH to 8.2. Protect the buffer solution from \( \text{CO}_2 \) of the air by attaching a drying tube containing soda lime (sodium calcium hydrate) to the air opening at the top of the solution bottle.
- Replacement solution. Barium chloride, 0.5 \( N \). Add 5 ml buffer solution per liter. Protect the replacement solution from \( \text{CO}_2 \) of the air by attaching a drying tube similar to that used for the buffer solution.
Procedure

Weigh 5 g soil into a 125-ml Erlenmeyer flask. Add 15 ml buffer solution and let stand for 30 minutes, swirling occasionally to mix. Use 35 ml buffer solution to transfer all the soil solution to a No. 4 Gooch crucible containing a moist Whatman No. 540 filter paper and filter into a 500-ml suction flask. The rate of filtration should be such that at least 30 minutes is needed to complete the filtering and leaching. Then leach the soil with 100 ml of the replacement solution, adding small amounts at a time. It may be necessary to use a larger amount of buffer solution to leach allophanic soils high in organic matter with extractable acidity of more than 35 meq per 100 g.

Back-Titration with HCl (6H1a)

Reagents

- Hydrochloric acid (HCl), 0.2 N, standardized
- Brom cresol green, 0.1-percent aqueous solution
- Mixed indicator. Dissolve 1.250 g methyl red indicator and 0.825 g methylene blue in 1 liter 90-percent ethanol.

Procedure

Run a blank by adding 100 ml replacement solution, 2 drops brom cresol green, and 10 drops mixed indicator to 50 ml buffer solution. Titrate with HCl to a chosen end point in the range from green to purple. Add 2 drops brom cresol green and 10 drops mixed indicator to the leachate and titrate to the same end point chosen for the blank. Calculate exchange acidity (EA) as follows.

Calculations

\[
EA \text{ (meq/100 g)} = \frac{(A - B)}{C} \times N \times 100
\]

where:
- \(A\) = Volume HCl blank (mL)
- \(B\) = Volume HCl sample (mL)
- \(C\) = Sample weight (g)
- \(N\) = Normality of HCl

Report on oven-dry basis.

References

Peech et al. (1947).
BaCl₂-Triethanolamine II (6H2)

**Apparatus**
- Sulfur absorption tubes
- Whatman No. 41 filter paper or glass-fiber filter paper cut to fit sulfur absorption tubes

**Reagents**
- Buffer solution. BaCl₂, 0.5 N, and triethanolamine, 0.2 N as in 6H1.
- Mixed indicator. Dissolve 1.250 g methyl red and 0.825 g methylene blue in 1 liter 90 percent ethanol.
- Celite

**Procedure**
Stopper bottom of sulfur absorption tubes with medicine-dropper bulbs and fit to a 300-ml suction flask with a rubber stopper. Place Whatman No. 41 filter paper in bottom of absorption tube, cover with ¼ inch of acid-washed sand, and add exactly 25 ml buffer solution. Weigh 10 g soil and mix with teaspoonful of Celite. Add to the absorption tube by means of a funnel. After 30 minutes remove the medicine-dropper bulbs, wash bulbs out with a little water, and add washings to absorption tubes. Leach with 25 ml more buffer solution and then leach with 100 ml replacement solution in small increments. If necessary, use suction to facilitate leaching.

**Back-Titration with HCl (6H2a)**

**Reagents**
- Same as in 6H1a.

**Procedure**
Titrated with standard HCl, using either 2 drops brom cresol green and 10 drops methyl red or 10 drops mixed indicator. Use same end point as that chosen for a blank run by leaching sand and Celite with 50 ml buffer solution and 100 ml replacement solution.

**Calculations**
Use same calculation as in 6H1a.
KCl-Triethanolamine (6H3)
Back-titration with NaOH (6H3a)

Procedure
Leach 10 g soil with 50 ml KCl-triethanolamine solution and follow by washing with 50 ml unbuffered 1 N KCL. Add a known volume of standard acid to leachate and washings and back-titrate with standard alkali (NaOH). Titrate an equal volume of acid to the same end point for a blank.

Calculations
\[
EA \text{ (meq/100 g)} = \frac{(A - B)}{C} \times N \times 100
\]
where:
- \(EA\) = Extractable acidity
- \(A\) = Volume NaOH sample (mL)
- \(B\) = Volume NaOH blank (mL)
- \(C\) = Sample weight (g)
- \(N\) = Normality of NaOH

References
North-Central Regional Research Committee (1955).

BaCl₂-Triethanolamine III (6H4)

Apparatus
- 60-ml plastic syringe barrels.
- Buffer solution. Barium chloride, 0.5 \(N\), and triethanolamine, 0.2 \(N\). Add 1 \(N\) HCl (about 90 ml/L) to adjust pH to 8.2. Protect the buffer solution from CO₂ of the air by attaching a drying tube containing soda lime (sodium calcium hydrate) to the air opening at the top of the solution bottle.
- Replacement solution. Barium chloride, 0.5 \(N\). Add 5 ml of above buffer solution per liter. Protect the replacement solution from CO₂ of the air by attaching a drying tube similar to that used for the buffer solution.
- “Celite” filter pulp.

Procedure
Prepare syringe barrels as leaching tubes by forcing a 1-g ball of filter pulp into bottom of barrel with syringe plunger. Measure 1.5 g celite and 5 g soil sample into tube. Attach pinch clamp to delivery tube of syringe barrel and
add approximately 25 ml buffer solution to sample. Let stand 30 min, stirring occasionally. Remove pinch clamp and filter with low suction into titrator beaker using a total of 50 ml buffer solution followed by 100 ml replacement solution.

References
Peech et al. (1947).

**Back-Titration with HCl, Automatic Titrator (6H4a)**

**Reagents**
- Hydrochloric acid (HCl), 0.33 \( N \), standardized

**Procedure**
Titrate the leachate contained in the 250-ml beaker to an end-point pH setting of 4.60 with automatic titrator. Carry reagent blank through procedure.

**Calculations**

\[
EA (\text{meq/100 g}) = ((A-B)/C) \times N \times 100
\]

where:
- \( EA \) = Extractable acidity
- \( A \) = Volume HCl blank (mL)
- \( B \) = Volume HCl sample (mL)
- \( C \) = Sample weight (g)
- \( N \) = Normality of HCl

Report on oven-dry basis.

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**Carbonate (6l)**

**Saturation Extract (6l1)**

**Acid titration (6l1a)**

**Reagents**
- Sulfuric acid (\( H_2SO_4 \)), 0.05 \( N \), standardized
- Phenolphthalein

**Procedure**
Pipette an appropriate aliquot of saturation extract into a 250-ml Erlenmeyer flask or a porcelain crucible. The electrical conductivity (EC \( \times 10^3 \)) of the saturation extract (8A1a) can be used to determine the aliquot to be used for carbonate,
bicarbonate, and chloride determinations. Where EC $\times 10^3$ is 1.0 or less, use a 10-
ml aliquot; if 1.0 to 10.0, use a 5-ml aliquot; if more than 10.0, use a 2-ml aliquot.

Make volume to 50 ml (10 ml for porcelain crucible) with water. To the 50 ml
in the Erlenmeyer flask, add a drop or two of phenolphthalein. If a pink color is
produced, titrate with 0.05 $N \text{H}_2\text{SO}_4$, adding a drop every 2 or 3 seconds until the
pink color disappears. Use this solution to determine bicarbonate (6J1a).

Calculations

\[
\text{Carbonate (meq/L)} = \frac{(A/B) \times N \times 2000}{1000}
\]

where:

- $A$ = Volume $\text{H}_2\text{SO}_4$ (mL)
- $B$ = Volume aliquot (mL)
- $N$ = Normality of $\text{H}_2\text{SO}_4$

References

Association of Official Agricultural Chemists (1945) and Richards (1954).

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Bicarbonate (6J)
Saturation Extract (6J1)
Acid Titration (6J1a)

Reagents

- Sulfuric acid ($\text{H}_2\text{SO}_4$), 0.05 $N$, standardized
- Methyl orange, 0.01-percent aqueous solution

Procedure

Use solution remaining from carbonate titration (6I1a). To the colorless
solution from this titration or to the original solution if no color is produced with
phenolphthalein, add 4 drops methyl orange and continue titration to the methyl
orange end point without refilling the burette. Retain this solution for the chloride
determination (6K1a). Make a blank correction for the methyl orange titration.

Calculations

\[
\text{Bicarbonate (meq/L)} = \frac{((A - (2 \times B)) / C) \times N \times 1000}{1000}
\]

where:

- $A$ = Total volume $\text{H}_2\text{SO}_4$ (mL)
- $B$ = Volume $\text{H}_2\text{SO}_4$ from 6I1a (mL)
- $C$ = Volume aliquot (mL)
- $N$ = Normality of $\text{H}_2\text{SO}_4$
Chloride (6K)
Saturation Extract (6K1)
Mohr Titration (6K1a)

Reagents
- Potassium chromate (K₂CrO₄) indicator. Dissolve 5 g K₂CrO₄ in water and add a saturated solution of AgNO₃ until a permanent slight red precipitate is produced, filter, and dilute to 100 ml.
- Silver nitrate (AgNO₃), 0.05 N, standardized
- Sodium bicarbonate (NaHCO₃), saturated solution (optional)
- Nitric acid (HNO₃), 0.1 N (optional)

Procedure
To the solution from the bicarbonate titration (6J1a) add 6 drops K₂CrO₄ indicator and titrate with AgNO₃ to a reddish-orange end point. Make a correction with a blank of 50 ml water containing the indicators of both titrations. The laboratory at Riverside, California, modifies this procedure by adding saturated NaHCO₃ solution to a pink end point and neutralizing to a colorless end point with HNO₃ before adding the indicator.

Calculations

\[
\text{Chloride (meq/L) = } \frac{(A - B)}{C} \times N \times 1000
\]

where:
- A = Volume AgNO₃ sample (mL)
- B = Volume AgNO₃ blank (mL)
- C = Volume aliquot (mL)
- N = Normality of AgNO₃

References
Association of Official Agricultural Chemists (1945).
Reagents

- Standard silver nitrate (AgNO₃), 0.025 N
- Buffer solutions. Either potassium acid phthalate or trisodium citrate and citric acid. To prepare phthalate buffer, weigh 37.5 g potassium acid phthalate and bring to a volume of 500 ml with water; 4 ml of this buffer added to a 46-ml solution brings the pH to about 4. To prepare trisodium citrate buffer, weigh 43.8 g trisodium citrate and 43.3 g citric acid into 500-ml volumetric flask and bring to volume with water. Add a small amount of toluene to the solution for storage; 10 ml of this buffer added to a 40-ml solution brings pH to about 4.

Procedure

Standardize the pH meter by adjusting the needle to a convenient setting (about 0.8) on the expanded scale when the electrode is immersed in buffer solution (4 or 10 ml made to 50 ml) without chloride. To titrate the sample, pipette an aliquot containing as much as 2.0 meq chloride into a beaker and add 4 ml buffer. Make to 50 ml. Immerse the electrode and burette tip into the beaker and titrate with AgNO₃ to the end point previously established for the buffer without chloride.

Calculations

Chloride (meq/L) = \((A/B) \times N \times 1000\)

where:

- \(A\) = Volume AgNO₃ (mL)
- \(B\) = Volume aliquot (mL)
- \(N\) = Normality of AgNO₃

Sulfate (6L)

Saturation Extract (6L1)

Gravimetric, BaSO₄ Precipitation (6L1a)

Reagents

- Concentrated hydrochloric acid (HCl)
- Barium chloride (BaCl₂), 10-percent
- Methyl orange, 0.01-percent

Procedure

Pipette an aliquot of saturation extract into a 250-ml beaker. Dilute to approximately 100 ml with water. Add 2 drops methyl orange and 0.5 ml concentrated HCl to the beaker. Heat to boiling and add BaCl₂ solution by drops,
stirring constantly until precipitation is complete. Let stand on hot plate for several hours. Remove from heat and let samples stand overnight. Filter through Gooch crucibles, which have been ignited and weighed. Dry in 105 °C oven and ignite in muffle furnace at 1200 °F (650 °C) for 30 minutes. Cool in desiccator and weigh.

Calculations

\[ \text{SO}_4 \text{(meq/L)} = \frac{A}{B} \times 8.568 \]

where:

\[ A = \text{BaSO}_4 \text{(mg)} \]
\[ B = \text{Volume aliquot (mL)} \]

References

Richards (1954).

**EDTA Titration (6L1b)**

Apparatus

- Repipet, automatic dilutor, pipette range 0.1 to 1.0 ml
- Titration assembly including a 10-ml burette with magnetic stirrer

Reagents

- Thymol blue indicator, 0.04-percent
- Nitric acid (HNO\(_3\)), 0.4 \(N\)
- Calcium nitrate (Ca(NO\(_3\))\(_2\)), 0.05 \(N\). Dissolve 5.90 g Ca(NO\(_3\))\(_2\)•4H\(_2\)O in 1 liter CO\(_2\)-free water. EC is 5.15 ±0.15 mmhos per cm at 25 °C.
- Acetone, reagent grade, boiling range 55.5 to 57.5 °C
- Ethanol, 95-percent, reagent grade
- Hydrochloric acid (HCl), 0.01 \(N\)
- EDTA solution, 0.02 \(N\). Standardize against CaCl\(_2\).

Procedure

Pipette an aliquot containing 0.01 to 0.05 meq SO\(_4\) from soil-water extracts and transfer to a 100-ml beaker. Bring volume to 7.5 ±0.5 ml with water. Add 2 drops 0.04-percent thymol blue and 0.4 \(N\) HNO\(_3\) drop by drop until color changes from yellow to distinct red. Add 2 ml 0.05 \(N\) Ca(NO\(_3\))\(_2\), 20 ml acetone, and stir. Allow 30 minutes for the precipitate to flocculate. Place a 9.0-cm Whatman No. 42 filter paper in a 5.0-cm fluted funnel and fit snugly with water. Wash the sides of filter paper with 5 ml 95-percent ethanol from a wash bottle. Transfer the precipitate and supernatant to the filter paper with alcohol. Rinse the beaker twice and wash filter paper three times, using 3 to 5 ml ethanol per rinse. Allow the alcohol in the filter paper to evaporate. Wash the funnel stem thoroughly with water. Place the
beaker that contained the CaSO₄ precipitate under the funnel and wash the filter paper with 3 to 5 ml portions of 0.01 N HCl until approximately 25 ml is leached. Proceed as in 6N1a, except eliminate carbamate and add an extra drop 4 N NaOH to neutralize the 25 ml 0.01 N HCl.

The amount of sulfate is determined from the Ca²⁺ content in the CaSO₄ precipitate.

**Calculations**

\[
\text{SO}_4 \text{ (meq/L)} = \frac{(A/B) \times N \times 1000}{V}
\]

where:

- \( A \) = Volume EDTA (mL)
- \( B \) = Volume aliquot (mL)
- \( N \) = Normality of EDTA

**References**

Bower and Wilcox (1965); Lagerwerff, Akin, and Moses (1965); and Nelson (1970).

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**NH₄OAc Extraction (6L2)**

Obtain extract by method 5B1.

**Gravimetric, BaSO₄ Precipitation (6L2a)**

Proceed as in 6L1a. A greater quantity of acid will be needed to lower the pH. Otherwise the procedures are the same.

---

**Nitrate (6M)**

**Saturation Extract (6M1)**

**Phenoldisulfonic Acid Colorimetry (6M1a)**

**Reagents**

- Phenoldisulfonic acid. Dissolve 25 g phenol in 150 ml concentrated \( \text{H}_2\text{SO}_4 \), add 75 ml fuming \( \text{H}_2\text{SO}_4 \) (13 to 15 percent \( \text{SO}_3 \)), and heat at 100 °C for 2 hours.
- Standard potassium nitrate (KNO₃), 0.010 \( N \)
- Silver sulfate (Ag₂SO₄), 0.020 \( N \)
- Ammonium hydroxide solution (NH₄OH), 1:1, approximately 7 \( N \)
- Calcium oxide (CaO)
Procedure

First determine the chloride concentration in an aliquot of saturation extract as directed in 6K1a. Pipette another aliquot containing 0.004 to 0.04 meq of nitrate into a 25-ml volumetric flask. Add an amount of Ag$_2$SO$_4$ equivalent to the amount of chloride present, dilute to volume, and mix. Separate the precipitate by centrifuging the suspension in a 50-ml centrifuge tube. Transfer the solution to another centrifuge tube, flocculate any suspended organic matter by adding about 0.1 g CaO, and clear by centrifuging again. Pipette a 10-ml aliquot into an 8-cm evaporating dish. Evaporate the aliquot to dryness, cool, and dissolve the residue in 2 ml phenoldisulfonic acid. After 10 minutes, add 10 ml water and transfer to a 100-ml volumetric flask. Make alkaline by adding NH$_4$OH, dilute to volume, and mix. Measure light transmission through a 460 mµ filter of solution in an optical cell against that of water in a similar cell.

Prepare a calibration curve by pipetting 0-, 0.2-, 0.4-, 0.8-, 1.2-, and 1.6-ml aliquots of standard KNO$_3$ into evaporating dishes and treating as for sample except for additions of Ag$_2$SO$_4$ and CaO and the clarifying procedure.

Calculations

\[ \text{NO}_3 \text{ (meq/L)} = \frac{A}{B} \times 1000 \]

where:

A = NO$_3$ from curve (meq/L)
B = Volume aliquot (mL)

References

Richards (1954).

**Diphenylamine (Qualitative) (6M1b)**

Use this procedure to test for nitrates if there is a significant excess of cations over anions in the extract. A quantitative measurement can be made if there is a positive indication of NO$_3$ (6M1a).

Reagents

- Diphenylamine in H$_2$SO$_4$. Dissolve 0.05 g diphenylamine in 25 ml concentrated sulfuric acid. Store in polyethylene dropper bottle.

Procedure

Place a drop of extract in a spot plate and add 3 or 4 drops diphenylamine reagent. Nitrate is present if a blue color develops.

References

Treadwell and Hall (1943).
Calcium (6N)
Saturation Extract (6N1)
   EDTA Titration (6N1a)

Reagents
   - Sodium hydroxide (NaOH), approximately 4 N
   - Calcium chloride (CaCl₂), 0.02 N. Dissolve calcite crystals in HCl and make to volume.
   - Murexide. Thoroughly mix 0.5 g ammonium purpurate with 100 g powdered potassium sulfate (K₂SO₄).
   - EDTA solution, 0.02 N. Standardize against CaCl₂.
   - Sodium diethyldithiocarbamate, 1-percent

Procedure
   Pipette an aliquot containing 0.02 to 0.20 meq of calcium into a beaker. Add 5 drops carbamate, 1 drop NaOH for each 5-ml aliquot, and a suitable amount (15 to 20 mg for a 10-ml aliquot) of murexide, mixing after each addition. A magnetic stirrer is helpful. Titrate with EDTA to a lavender end point. A blank containing NaOH, murexide, carbamate, and a drop or two of EDTA helps to distinguish the end point. If the sample is overtitrated with EDTA, it can be back-titrated with standard CaCl₂. Retain solution for magnesium determination (6O1a).

Calculations
   \[ \text{Ca (meq/L)} = \frac{A}{B} \times N \times 1000 \]
   where:
   \[ A = \text{Volume EDTA (mL)} \]
   \[ B = \text{Volume aliquot (mL)} \]
   \[ N = \text{Normality of EDTA} \]

References
   Cheng and Bray (1951).

NH₄OAc Extraction (6N2)
   Prepare NH₄OAc extract as described in 5A1. EDTA-alcohol extraction.

EDTA-Alcohol Separation (6N2a)

Reagents
   - Standard calcium chloride (CaCl₂), 5 mg per ml. Dissolve calcite crystals in HCl and make to volume.
• Ethanol, 95-percent
• Standard EDTA. Dissolve 1.25 g disodium ethylenediaminetetraacetate in water and dilute to a volume of 1 liter. Standardize against solutions containing known amounts of calcium and magnesium. Run the standards through the separation procedure before titrating.
• Sodium hydroxide (NaOH), 10-percent aqueous solution
• Calcon. Dissolve 1 g Calcon (Eriochrome Blue Black R) in 100 ml triethanolamine.

Procedure
Pipette 25-ml aliquots from the pH 7, \( \text{NH}_4 \text{OAc} \) extracts obtained in the total exchange-capacity method (5A1) into 100-ml beakers and evaporate to dryness at moderate heat. Cool and add 3 ml \( \text{N} \text{HNO}_3 \) to dissolve the residue. Transfer the solution quantitatively to 50-ml conical centrifuge tubes with ethanol, using a wash bottle with a fine delivery tip. Add 1 ml 6 \( \text{N} \text{H}_2\text{SO}_4 \). While mixing the contents of the tube by swirling, add approximately 34 ml 95-percent ethanol. Cover the tubes and let stand overnight. The next morning remove the covers and centrifuge the tubes at about 2000 rpm (Int. No. II centrifuge) for 15 minutes. Decant the alcohol solution into 250-ml Erlenmeyer flasks and retain for the magnesium determination. Use the \( \text{CaSO}_4 \) precipitate for calcium determination.

Break up the \( \text{CaSO}_4 \) precipitate with a small steam of water from a wash bottle and transfer the precipitate and solution to 250-ml Erlenmeyer flasks. Dilute the solution to a total volume of about 100 ml. Place the sample on a magnetic stirrer, add 5 ml 10-percent NaOH, 2 drops Calcon indicator solution, and titrate with the standard EDTA solution to the blue color of a blank carried through the procedure. The pH of the solution should be about 12.5. The color change is from red to clear blue. Titrate until the color in the sample and in the blank are the same.

Calculations

\[
\text{Ca (meq/100 g)} = \frac{A}{B} \times N \times \frac{C}{D} \times 100
\]

where:

\[
A = \text{Volume EDTA (mL)}
\]

\[
B = \text{Sample weight (g)}
\]

\[
N = \text{Normality of EDTA}
\]

\[
C = \text{Volume extract (mL)}
\]

\[
D = \text{Volume aliquot (mL)}
\]

References
Barrows and Simpson (1962).
**Oxalate Precipitation I, KMnO₄ Titration (6N2b)**

**Reagents**

- Oxalic acid (C₂H₂O₄), 5-percent aqueous solution
- Brom cresol green, 0.04-percent aqueous solution
- Ammonium hydroxide (NH₄OH), 1 N
- Sulfuric acid (H₂SO₄), 1 N
- Standard potassium permanganate (KMnO₄), 0.05 N
- Wash solution, saturated calcium oxalate (CaC₂O₄)
- Asbestos. Digest asbestos in 1 N HNO₃ solution containing just enough KMnO₄ to give a deep purple color. Add more permanganate if the color disappears; digest for 24 hours or until the permanganate color is permanent. Destroy the excess permanganate with oxalic acid and wash thoroughly on a Buchner funnel.

**Procedure**

Transfer an aliquot of the filtrate (5A1) to a 400-ml Pyrex beaker and evaporate to complete dryness. Cool, cover the beaker with a watchglass, and slowly add through the lip 10 ml concentrated HNO₃ and 2 ml concentrated HCl. Warm until the reaction has subsided and no more brown fumes are given off. Rinse the watchglass into the beaker. Evaporate to dryness at low heat to prevent spattering and continue to heat for about 10 minutes to dehydrate the salts. Then place the beaker in an electric muffle furnace at about 150 °C, heat to 390° ±10°, and hold at this temperature for about 20 minutes. Remove the beaker from the muffle furnace and cool. Treat the residue with 3 ml 6 N HCl, evaporate to dryness at low heat, and continue heating for about 30 minutes longer to dehydrate silica. Cool and dissolve the residue in 0.1 N HNO₃, using a rubber policeman to loosen the residue.

Add 5 ml oxalic acid, heat the contents of the beaker almost to boiling, and add 1 ml brom cresol green. Adjust the pH of the hot solution to approximately 4.6 by slowly adding 1 N NH₄OH, stirring constantly. Let digest at about 80 °C for 1 hour or until the supernatant liquid is clear. Collect the CaC₂O₄ precipitate on a compact asbestos pad in a Gooch crucible or in a Whatman No. 42 filter paper in filter funnel. Rinse the beaker four times with water or water saturated with CaC₂O₄ and pour the washings into the crucible. Wash the precipitate five more times with water saturated with CaC₂O₄.

Remove the Gooch crucible from its holder, rinse the outside, and replace crucible in the beaker. If filter paper is used, pierce the paper and wash most of the precipitate into the beaker with 3.6 N H₂SO₄. Wash off excess H₂SO₄ with water and place filter paper on watchglass. Add 100 ml water and 7 ml concentrated H₂SO₄. Heat to 90 °C and stir until CaC₂O₄ is dissolved. Titrate with standard KMnO₄ solution to a pink color. Add filter paper to solution and titrate to a permanent pink color.
Calculations

\[ \text{Ca (meq/100 g)} = \frac{(A/B)}{N} \times \frac{(C/D)}{100} \]

where:
\( A = \text{KMnO}_4 \text{ (mL)} \)
\( B = \text{Sample weight (g)} \)
\( N = \text{Normality of KMnO}_4 \)
\( C = \text{Volume extract (mL)} \)
\( D = \text{Volume aliquot (mL)} \)

Report on oven-dry basis.

References

Peech et al. (1947).

**Oxalate Precipitation II, KMnO₄ Titration (Fe, Al, and Mn removed) (6N2c)**

Proceed as in 6N2b but after muffle treatment and before oxalate precipitation, remove iron, aluminum, and manganese by the following procedure.

**Reagents**
- Hydrochloric acid (HCl), 6 \(N\)
- Ammonium hydroxide (NH₄OH), 2 \(N\)
- Bromine water, saturated
- Ammonium chloride (NH₄Cl), 6 \(N\)
- Concentrated nitric acid (HNO₃)

**Procedure**

Dissolve salts and oxides by adding 5 ml 6 \(N\) HCl and heating on a hot plate until all salts and oxides are in solution. Add 75 to 100 ml water and heat the solution until it is nearly boiling. Immerse the pH electrodes into the hot solution and precipitate the hydroxides of iron, aluminum, and titanium by slowly adding 2 \(N\) NH₄OH until the meter indicates a pH of 6.2 to 6.4. Add 2 more drops of NH₄OH to neutralize the acidifying effect of the 15 ml saturated bromine water, which is slowly added next to precipitate manganese hydroxide. Since bromine water lowers the pH of the solution, readjust it to 6.2 to 6.4 with 2 \(N\) NH₄OH. Heat the solution with precipitate until it just begins to boil (1 or 2 min on a Bunsen burner) and remove from the heat.

Place on a hot plate at a temperature of 80 to 90 °C for 1 hour. Filter when the breaker has cooled enough to handle easily. Use an 11-cm Whatman No. 42 filter paper or its equivalent. Collect the filtrate in a beaker of the same size as those used for precipitating calcium. Wash and police the beaker containing the precipitate with hot 2-percent NH₄Cl. Wash the precipitate on the filter with the same solution. Five washings are usually enough. To the filtrate add 10
ml concentrated HNO₃ and evaporate to dryness; add 5.0 ml 6 N HCl, take to dryness, and use high heat to dehydrate silica. Proceed with the calcium precipitation (6N2b).

References
Washington (1930) and Fieldes et al. (1951).

Oxalate Precipitation, Cerate Titration (6N2d)
Proceed as in 6N2b except substitute the following for the permanganate titration.

Reagents
- Ammonium hexanitrate cerate \((\text{NH}_4)_2 \text{Ce(NO}_3)_3\) in molar perchloric acid \((\text{HClO}_4)_4\), 0.1 N. Add 85 ml 70- to 72-percent perchloric acid to 500 ml water. Dissolve 56 g ammonium hexanitrate cerate in the acid solution and dilute to 1 liter.
- Ammonium hexanitrate cerate in molar perchloric acid, 0.05 N. Follow the directions for the preparation of the 0.1 N solution but use only 28 g cerate.
- Perchloric acid \((\text{HClO}_4)_4\), 2 N. Add 170 ml 70 to 72-percent perchloric acid to 500 ml water and dilute to 1 liter.
- Nitro-ferroin indicator solution. Dilute a solution of nitro-orthophenanthroline ferrous sulfate with water to a convenient working strength. Two to four drops of the solution should give a sharp color change at the end point.
- Standardize the cerate solutions against accurately weighed quantities of primary standard grade sodium oxalate. Convenient weights of sodium oxalate are 0.10 to 0.11 g for the 0.05 N solution and 0.10 to 0.18 g for the 0.1 N cerate solution. Dissolve the sodium oxalate in 100 to 150 ml 2 N perchloric acid and titrate as directed in the following procedure.

Procedure
Dissolve the filtered and washed (use water) calcium oxalate in 100 to 200 ml 2 N perchloric acid. If a paper filter has been used, macerate it before titration. Add 2 to 4 drops of nitro-ferroin indicator solution and titrate with 0.05 N or 0.1 N cerate solution, depending upon the amount of oxalate present. The solution changes from red to colorless at the end point.

Calculations
\[ \text{Ca (meq/100 g)} = (A/B) \times N \times (C/D) \times 100 \]
where:
- \(A\) = Volume cerate (mL)
- \(B\) = Sample weight (g)
- \(N\) = Normality of cerate
- \(C\) = Volume extract (mL)
$D =$ Volume aliquot (mL)

Report on oven-dry basis.

**NH$_4$Cl-Ethanol Extraction (Calcareous Soils) (6N3)**

**Apparatus**
- See figure 6N3-1.

**Reagents**
- Ammonium chloride (NH$_4$Cl), 1 N, in 60-percent ethanol.
  To make 9 liters of extraction solution, dissolve 482 g NH$_4$Cl in 2,835 ml water and add 5,985 ml 95-percent ethanol. Adjust pH to 8.5 with 140 to 145 ml NH$_4$OH.
- Celite

**Procedure**
Fill extraction tube with water, set tube upright in holder, and let most of the water drain out. Close screw clamp and place filter paper on plate with a stirring rod. Let remainder of the water drain out of tube. The filter paper provides enough tension to keep the bottom part of the tube filled with water. Place tube on the rack and add about 1½ teaspoons washed sand. Place an extra perforated plate (inverted) on top of the sand and cover the plate with more sand. Place heaping teaspoon of Celite on the sand and pour about 20 ml extraction solution into the tube. Pour remainder of 400 ml extraction solution into a 500-ml Erlenmeyer flask. Add soil sample slowly and then stir with a rod to mix soil and Celite. Allow sample to settle and then place filter paper on top of the soil column. Put upper tube in place, stopper, and let stand overnight.

![Figure 6N3-1.—Apparatus for ammonium chloride-ethanol extraction for calcium (6N3).](image-url)
In the morning, place a 500-ml volumetric flask under the delivery tip and open screw clamp on lower extraction tube slowly. When level of liquid is a few milliliters above the soil, invert the 500-ml Erlenmeyer flask containing remainder of extraction solution (delivery tube in place), place glass tip in the upper tube, and open the pinch clamp. Use the screw clamp on lower tube to adjust flow rate through soil column. When all the extraction solution has passed through the soil column, remove volumetric flask, make to volume with water, and mix.

**EDTA Titration (6N3a)**

Pipette a 50-ml aliquot for determination of Ca and Mg into a 100-ml beaker and evaporate to dryness. Add 10-ml concentrated HNO₃ and 1 or 2 ml concentrated HCl. Cover with watchglass, place on hot plate, and heat until no more brown fumes are evolved. Remove cover glass, rinse into beaker, and evaporate solution to dryness. Take up residue with 3 ml \( \text{N}_\text{HNO}_3 \). Quantitatively transfer solution with ethanol to a 50-ml conical centrifuge tube and proceed with determination of Ca according to 6N2a.

References

Tucker (1954).

**KCl-Triethanolamine Extraction (6N4)**

Prepare extract as in method 5B2.

**Oxalate-Permanganate Titration (6N4a)**

Proceed as in 6N2b.

**EDTA Titration (6N4b)**

Reagents

- Sodium hydroxide (NaOH), 4 \( N \)
- EDTA 0.02 \( N \). Dissolve 3.723 g disodium dihydrogen ethylenediamine tetraacetate in water and dilute to 1 liter. Standardize the solution against standard CaCl₂ prepared in the TEA buffer solution.
- Ammonium purpurate (murexide) indicator. Thoroughly mix 0.5 g ammonium purpurate with 100 g powdered potassium sulfate.
- Eriochrome Black T (Erio T) indicator. Dissolve 0.5 g Erio T in 100 ml of triethanolamine.

Procedure

Pipette a 5-ml aliquot of extract from method 5B3 into a 100-ml beaker. Add 20 ml water, 5 drops 4 \( N \) NaOH, and 50 mg murexide. Titrate with standard EDTA.
using a 10-ml microburet. Approach the end point slowly (orange-red to lavender or purple). Save the solution for the Mg\textsuperscript{2+} determination.

Calculations
\[
\text{Ca (meq/100 g)} = \frac{(A/B) \times N \times (C/D) \times 100}{B}
\]
where:
- \(A\) = Volume EDTA (mL)
- \(B\) = Sample weight (g)
- \(N\) = Normality of EDTA
- \(C\) = Volume extract (mL)
- \(D\) = Volume aliquot (mL)

References
Bower and Wilcox (1965).

Atomic Absorption (6N4c)
Proceed as in 6N1b except use sample from KCl-TEA extraction.

HF Dissolution (6N5)
Obtain extract as in 7C3.

Atomic Absorption (6N5a)

Apparatus
- Diluter
- Atomic absorption spectrophotometer

Reagents
- Standard Ca solutions, 0 to 30 meq/L

Procedure
Dilute HF extracts from 7C3 and Ca standards fivefold to twentyfold with water. Compare absorbance of samples with that of standards at 442.7 nm.

Calculations
\[
\text{Ca (pct.)} = \frac{(A \times 10 \times \text{dilution} \times 20.04 \text{ mg/meq})}{B}
\]
where:
- \(A\) = Ca (meq/L)
- \(B\) = Sample weight (mg)
CaO (pct.) = Ca (pct.) x 1.40

Report on oven-dry basis.

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**Magnesium (6O)**

**Saturation Extract (6O1)**

**EDTA Titration (6O1a)**

### Reagents

- **Buffer solution.** Mix 33.75 g NH₄Cl with 285 ml concentrated (15 N) NH₄OH, add 5 g disodium Mg-versenate, and dilute to 500 ml.
- **Eriochrome Black T indicator.** Dissolve 0.5 g Eriochrome Black T (F241) and 4.5 g hydroxylamine hydrochloride (NH₂OH·HCl) in 100 ml 95-percent ethanol or dissolve 1.0 g Eriochrome Black T in 100 ml triethanolamine.
- **EDTA 0.02 N.** Standardize with magnesium solution.

### Procedure

To the sample just titrated for calcium (6N1a), add 3 or 4 drops concentrated HCl, stir until the murexide is destroyed, add 1 ml NH₄Cl•NH₄OH buffer solution, 1 or 2 drops Eriochrome Black T indicator, and complete the titration for magnesium, using EDTA. The end point should be a clear blue with no tinge of red.

### Calculations

\[ \text{Mg (meq/100 g)} = \frac{(A/B) \times N \times (C/D) \times 100}{\text{Report on oven-dry basis.}} \]

where:

- \( A \) = Volume EDTA (mL)
- \( B \) = Sample weight (g)
- \( N \) = Normality of EDTA
- \( C \) = Volume extract (mL)
- \( D \) = Volume aliquot (mL)

### References

Cheng and Bray (1951).

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**NH₄OAc Extraction (6O2)**

Prepare NH₄OAc extraction as described.
EDTA Titration, Alcohol Separation (6O2a)

Reagents

- Buffer solution. Dissolve 67.5 g NH₄Cl in about 400 ml water. Add 570 ml concentrated NH₄OH and dilute to 1 liter with water.
- Hydroxylamine hydrochloride (NH₂OH·HCl), 5-percent aqueous solution. Prepare fresh solution every 10 days.
- Potassium ferrocyanide (K₄Fe(CN)₆•3H₂O), 4-percent aqueous solution
- Triethanolamine, U.S.P.
- Eriochrome Black T. Dissolve 1 g Eriochrome Black T (Superchrome Black TS) in 100 ml triethanolamine.
- Standard magnesium solution, 5.0 mg per milliliter. Transfer 2.500 g unoxidized reagent-grade magnesium metal to a 500-ml volumetric flask. Add 150 ml water and 20 ml concentrated HCl. When in solution, make to volume with water and mix. Dilute an aliquot of this solution to get a solution containing 0.5 mg magnesium per milliliter.
- EDTA, 0.02 N. Standardize with magnesium standard solution.

Procedure

Place the Erlenmeyer flasks containing the alcohol solution retained from the CaSO₄ separation (6N2a) on a hot plate and evaporate the alcohol at moderate heat. Do not evaporate to complete dryness. Cool and dilute to 100 ml with water and add 5 ml buffer solution and 10 drops each of hydroxylamine hydrochloride, potassium ferrocyanide, and triethanolamine. Stir and let stand 5 to 10 minutes. Place the sample on the stirrer, add 2 drops Eriochrome Black T, and titrate with standard EDTA to the ice-blue end point. The color change is from red through wine to ice blue. A blank carried through this procedure usually requires 0.3 to 0.8 ml EDTA to get the proper ice-blue color. Correct for a blank carried through this procedure and use the corrected titration to calculate the magnesium in the sample.

Calculations

\[ \text{Mg (meq/100 g) = } \frac{A}{B} \times N \times \frac{C}{D} \times 100 \]

where:

- A = Volume EDTA (mL)
- B = Sample weight (g)
- N = Normality of EDTA
- C = Volume extract (mL)
- D = Volume aliquot (mL)

Report on oven-dry basis.
References
Barrows and Simpson (1962).

Phosphate Titration (6O2b)

Reagents
- Sodium hydroxide (NaOH), 0.1 N, standardized. Protect from CO\textsubscript{2} of the air with a soda lime trap.
- Sulfuric acid (H\textsubscript{2}SO\textsubscript{4}), 0.1 N
- Ammonium hydroxide (NH\textsubscript{4}OH), concentrated
- Diammonium hydrogen phosphate ((NH\textsubscript{4})\textsubscript{2} HPO\textsubscript{4}), 10-percent solution
- Brom cresol green, 0.1-percent aqueous solution
- Hydrochloric acid (HCl), 1:1
- Carbon-dioxide-free water. Boil water in a 5-liter round-bottom boiling flask for about 15 minutes. Cool and protect from CO\textsubscript{2} of the air with a soda lime trap.

Procedure
Transfer the filtrate from the calcium determination (6N2b, 6N2c, or 6N2d) to a 400-ml beaker, add 10 ml concentrated HN\textsubscript{3}, cover with a 3.5-inch Speedyvap watchglass and evaporate to dryness. Dissolve the residue in 5 ml 1:1 HCl and transfer to a 250-ml Erlenmeyer flask, policing twice and rinsing the beaker twice after final policing. The volume of solution should be about 75 ml or more. Using 3 to 4 drops brom cresol green indicator, neutralize the solution with concentrated NH\textsubscript{4}OH added by drops. Add 5 ml 10-percent (NH\textsubscript{4})\textsubscript{2} HPO\textsubscript{4} and 10 ml concentrated NH\textsubscript{4}OH. Heat the solution just to boiling, cool, stopper, and let stand overnight.

Filter through a 9-cm Whatman No. 40 filter paper, pouring the solution down a stirring rod. Rinse the flask five times with 1 N NH\textsubscript{4}OH and pour the rinsings onto the filter. Wash the precipitate on the filter five more times with 1 N NH\textsubscript{4}OH. Place the wet filter paper with precipitate on a watchglass and let dry at no more than 40 °C until free of ammonia. Place the dry filter in the original flask, add 5 drops brom cresol green and 10 ml 0.1 N H\textsubscript{2}SO\textsubscript{4} or more if necessary to dissolve the precipitate. The solution should be yellow. After most of the precipitate has dissolved, add 50 ml CO\textsubscript{2}-free water, stopper the flask, and shake vigorously until the filter paper is macerated. Remove the stopper and rinse it and the flask walls with CO\textsubscript{2}-free water. Back-titrante with standard 0.1 N NaOH to pH 4.5.

To determine the correct end point, prepare a color standard by pipetting 5 ml potassium dihydrogen phosphate (2-percent solution) into a 250-ml Erlenmeyer flask, adding 65 ml water, 5 drops brom cresol green, and a macerated filter paper.

Calculations
\[
Mg \text{ (meq/100 g)} = ((A−B)/C) \times N \times (D/E) \times 100
\]
where:
A = Volume NaOH blank (mL)
B = Volume NaOH sample (mL)
C = Sample weight (g)
N = Normality of NaOH
D = Volume extract (mL)
E = Volume aliquot (mL)

Report on oven-dry basis.

References
Peech et al. (1947).

Gravimetric, Magnesium Pyrophosphate (6O2c)

Reagents
- Diammonium hydrogen phosphate ((NH₄)₂HPO₄), 10-percent solution
- Nitric acid (HNO₃), concentrated
- Ammonium hydroxide (NH₄OH), concentrated
- Ammonium hydroxide (NH₄OH), 1:1
- Hydrochloric acid (HCl), 6 N

Procedure
Continue analysis on filtrate from oxalate precipitation (6N2b). This filtrate will probably fill a 150-ml beaker. Place cover glass on filtrate and heat at a low temperature. When volume has been reduced, add 20 ml concentrated HNO₃. Evaporate to complete dryness and wash cover glass and sides of beaker with water. Dissolve residue in 5 ml 6 N HCl and then dilute to about 75 ml. Add 2 or 3 drops brom cresol green and bring pH to 4.6 with 1:1 NH₄OH. Add 5 ml 10-percent diammonium hydrogen phosphate (make up fresh each time). Add 10 ml concentrated NH₄OH, stir solution vigorously until a precipitate forms, and let stand overnight.

On the next day filter on a 11.0-cm Whatman No. 42 filter paper, rinse beaker five times with 1 N NH₄OH, and pour washings into the filter. Wash the precipitate in the filter five more times with 1 N NH₄OH. Place filter in oven to dry (2 to 3 hours) and evolve NH₄OH to prevent any explosion in the muffle furnace. Place crucibles (Coors 000) with filters containing magnesium precipitate in muffle furnace. Raise temperature gradually to 1000 °C and hold at 1000 °C for 1 hour. Allow muffle furnace to cool down and remove crucibles. Place in desiccator and dry over phosphorus pentoxide (P₂O₅). Weigh Mg₂P₂O₇ and record.
Calculations

\[ \text{Mg (meq/100 g)} = \frac{A}{B} \times \frac{C}{D} \times 1.797 \]

where:
\[ A = \text{Mg}_2\text{P}_2\text{O}_7 \text{ (mg)} \]
\[ B = \text{Sample weight (g)} \]
\[ C = \text{Volume extract (mL)} \]
\[ D = \text{Volume aliquot (mL)} \]

---

**NH\textsubscript{4}Cl-Ethanol Extraction (Calcareous Soils) (6O3)**

Proceed as in 6N3.

**EDTA Titration (6O3a)**

Proceed as in 6N3a except determine magnesium in alcohol extract by method 6O2a.

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**KCl-Triethanolamine Extraction (6O4)**

Prepare extract as described in 5B2 or 5B3.

**Phosphate Titration (6O4a)**

Proceed as in 6O2b except use extract from 5B2 or 5B3.

**EDTA Titration (6O4b)**

**Reagents**

- Concentrated hydrochloric acid (HCl)
- Concentrated ammonium hydroxide (NH\textsubscript{4}OH)
- EDTA 0.02 \( N \). Dissolve 3.723 g disodium dihydrogen ethylenediaminetetraacetate in water and dilute to a volume of 1000 ml. Standardize the solution against standard MgCl\textsubscript{2}.
- Eriochrome Black T (Erio T) indicator. Dissolve 0.5 g Erio T in 100 ml triethanolamine.

**Procedure**

Add 4 or 5 drops concentrated HCl to the solution used for the calcium determination (6N4b). Set aside until the murexide turns colorless. Add 15 to 20 drops concentrated NH\textsubscript{4}OH. This should bring the pH between 10.0 and 10.3. Add 1 drop Erio T and titrate with EDTA to a clear blue end point.
Calculations

\[ \text{Mg (meq/100 g)} = \frac{(A \times B \times N \times C \times D)}{100} \]

where:
- \( A \) = Volume EDTA (mL)
- \( B \) = Sample weight (g)
- \( N \) = Normality of EDTA
- \( C \) = Volume extract (mL)
- \( D \) = Volume aliquot (mL)

Report on oven-dry basis.

**Atomic Absorption (6O4c)**

Proceed as in 6O1b except use samples from the KCl-TEA extract.

**HF Dissolution (6O5)**

Obtain extract as in 7C3.

**Atomic Absorption (6O5a)**

**Apparatus**
- Diluter
- Atomic absorption spectrophotometer

**Reagents**
- Standard Mg solutions, 0 to 10 meq/L

**Procedure**
Dilate HF extracts from 7C3 and Mg standards fivefold to twentyfold with water. Compare absorbance of samples with that of standards at 285.2 nm.

**Calculations**

\[ \text{Mg (pct.)} = \frac{(A \times 10 \times \text{dilution} \times 12.16 \text{ mg/meq})}{B} \]

where:
- \( A \) = Mg (meq/L)
- \( B \) = Sample weight (mg)

\[ \text{MgO (pct.)} = \text{Mg (pct.)} \times 1.66 \]

Report on oven-dry basis.
Sodium (6P)
Saturation Extract (6P1)
   Flame Photometry (6P1a)

Apparatus
Beckman Model DU spectrophotometer with flame attachment.

Reagents
- Standard sodium solutions, 0.0 to 2.0 meq per liter
- Concentrated hydrochloric acid (HCl)
- Hydrochloric acid (HCl), 6 N
- Hydrochloric acid (HCl), 0.4 N
- Concentrated nitric acid (HNO₃)

Procedure
Pipette an aliquot of appropriate size (5 to 25 ml) of the saturation extract into a 100-ml beaker and evaporate to dryness on a hot plate. Treat the residue with 1 ml concentrated HCl and 3 ml concentrated HNO₃ and again evaporate to dryness on the hot plate. Repeat the acid treatment on the residue. Add 5 ml 6 N HCl to the residue and bring to dryness. Then raise the temperature to high for 20 minutes to render the silica insoluble. Wash and filter the residue into 50-ml volumetric flasks, using 0.4 N HCl. Determine flame luminosity of samples appropriately diluted and compare with luminosity of standard solutions made up with 0.4 N HCl. The evaporation and dehydration steps are used only where there is enough silica to clog the burner. If they are not used, merely dilute the sample.

Calculations
Na (meq/L) = A x dilution
where:
A = Na from curve (meq/L)

NH₄OAc Extraction (6P2)
Flame Photometry (6P2a)
Proceed as in 6P1a except make standard solutions in NH₄OAc. The evaporation and dehydration steps can be eliminated.

Calculations
Na (meq/100 g) = (A/B) x dilution x (C/10)
where:
A = Na from curve (meq/L)
B = Sample weight (g)
C = Volume extract (mL)

Report on oven-dry basis.

References
Fieldes et al. (1951).

HF Dissolution (6P3)
Obtain extract as in 7C3.

Atomic Absorption (6P3a)

Apparatus
• Diluter
• Atomic absorption spectrophotometer

Reagents
• Standard Na solutions, 0 to 20 meq/L in HF and boric acid

Procedure
Dilute HF extracts from 7C3 and Na standards fivefold to twentyfold with water. Compare absorbance of samples with that of standards at 589 nm.

Calculations
Na (pct.) = (A x dilution x 23.00 mg/meq) / B
where:
A = Na (meq/L)
B = Sample weight (mg)

Na₂O (pct.) = Na (pct.) x 1.35

Report on oven-dry basis.

Potassium (6Q)
Saturation Extract (6Q1)
Flame photometry (6Q1a)

Apparatus
• Beckman Model DU spectrophotometer with flame attachment
Reagents

- Standard potassium chloride (KCl) solutions ranging from 0.0 to 1.0 meq per liter

Procedure

Proceed as in 6P1a. Determine flame luminosity of potassium at 768 mµ and compare with that of the standard solutions.

Calculations

\[ K \text{ (meq/L)} = A \times \text{dilution} \]

where:

\[ A = K \text{ from curve (meq/L)} \]

References

Fieldes et al. (1951).

NH₄OAc Extraction (6Q2)
Flame Photometry (6Q2a)

Proceed as in 6Q1a except make up standards in NH₄OAc solution.

Calculations

\[ K \text{ (meq/100 g)} = \frac{A}{B} \times \text{dilution} \times \frac{C}{10} \]

where:

\[ A = K \text{ from curve (meq/L)} \]
\[ B = \text{Sample weight (g)} \]
\[ C = \text{Volume extract (mL)} \]

Report on oven-dry basis.

Sulfur (6R)
NaHCO₃ Extract, pH 8.5 (6R1)
Methylene Blue Colorimetry (6R1a)

References

Kilmer and Nearpass (1960).
HCl Release (Sulfide) (6R2)

Apparatus
- Nitrogen tank (water pumped)
- Scrubber. 250-ml Erlenmeyer flask equipped with three-hole rubber stopper to accommodate entry and exit for sweep gas, and a 4-foot glass tube to serve as a manometer.
- Reaction flask. 250-ml Erlenmeyer equipped with three-hole rubber stopper to accommodate entry and exit for sweep gas, and a burette for adding acid. Reaction flask sits on a magnetic stirrer.
- Collection bottles. Two 500-ml bottles (No. 8 rubber stopper), each fitted with a two-hole rubber stopper to accommodate entry and exit tubes for sweep gas. The entry tubes should be detachable below the rubber stopper. Attach a pinch clamp to the exit tube to help control the flow rate of gases.

Procedure
Place 10 ml zinc acetate solution in collection flask. Add water to 150 ml volume and place flask in train. Add moist sample (collected as in 1A2b) in tared reaction flask (250-ml Erlenmeyer), introduce N$_2$ gas (unless sample is run immediately), stopper, and weigh. Determine moisture content on a separate sample. Place flask in collection train. Sweep with N$_2$ gas for about 5 minutes. Reduce flow until pressure drops enough so that 50 to 60 ml of 6 N HCl can be added to reaction flask. Adjust flow of N$_2$ to about 4 bubbles per second in collection flask and turn on stirrer. Collect sample for 45 to 60 minutes. Second collection bottle should be a blank. Cut flow, disconnect entry tube but leave in collection bottle, remove collection bottles, and stopper until ready to titrate.

Iodine Titration (6R2a)

Apparatus
- Iodine applicator, approximately a 50-ml reservoir with stopcock delivery in a two-hole rubber stopper (No. 8). Fit a glass tube for air exit through the stopper.
- Burette for thiosulfate
- Magnetic stirrer

Reagents
- Iodine 0.1 N, standardized
- Sodium thiosulfate 0.1 N, standardized
- Starch indicator
- Hydrochloric acid (HCl), 6 N
Procedure

Mix an aliquot of standardized iodine solution and 5 ml of 6 \( N \) HCl in iodine applicator. Place applicator on bottle and add acidified iodine. Wash contents of applicator, quantitatively, into bottle. Remove applicator, stopper bottle, and swirl so that iodine enters the top of the entry tube from collection train. Any white precipitate of ZnS should dissolve off entry tube. Remove stopper and titrate with standardized thiosulfate until iodine color becomes faint. Add 1 or 2 ml starch indicator and titrate until blue color changes to clear. The end point is abrupt.

Stopper bottle and again swirl so that solution passes through the entry tube. Blue color should reappear. Again titrate to the end point. Magnetic stirrer can be used to mix the sample.

Calculations

\[
S \text{ (meq/100 g)} = \frac{(A - B)}{C} \times N \times 100
\]

where:

- \( A \) = Volume thio for blank (mL)
- \( B \) = Volume thio for sample (mL)
- \( C \) = Sample weight (g)
- \( N \) = Normality of thio

References

Pierce and Haenisch (1955); Johnson and Ulrich (1959); and Chapman and Pratt (1961).

SO\(_2\) Evolution (6R3)

KIO\(_3\) Titration (6R3a)

Apparatus

- LECO induction furnace model 521
- LECO automatic sulfur titrator model 532
- LECO crucibles and lids
- Oxygen tank and regulator
- LECO starch dispenser and 0.2-ml scoop

Reagents

- Potassium iodate (KIO\(_3\))
- Potassium iodide (KI)
- Arrowroot starch
- Hydrochloric acid (HCl) 7.7 \( N \)
- Hydrochloric acid (HCl) 0.18 \( N \)
• Magnesium oxide, (MgO)
• Iron-chip accelerator
• Copper metal accelerator

Procedure
Into a tared crucible, weigh approximately ½ g of 60-mesh soil, recording gross weight. Where high sulfur content might be present, either ¼ or ¹/₁₀ g sample should be run. Add 2 scoops of MgO and a scoop of iron chips. Mix thoroughly. Add a half scoop of copper accelerator and a scoop of iron chips. Magnesium oxide scoops are heaping; all others are level. A cover is placed on the crucible, which is placed on the pedestal and raised into the combustion tube for ignition. The LECO instruction manual is followed in setting up the furnace and titrator. The timer is set to 8 min and grid tap switch to midposition. These settings should be adjusted as needed to get complete fusion of the mixture in the crucible; however, plate current should not exceed 350 mA. When the burette reading does not change for 2 min and plate current has achieved 300 to 350 mA, the titration is complete and the titer is recorded. A blank is run using all ingredients except soil. Sulfate removal before analysis may be desirable in some instances. Sample is leached with 50 ml of 7.7 \( \text{N} \) HCl followed by 500 ml of distilled water.

Calculations
The KIO\(_3\) burette is direct reading in percent for a 1-g sample containing up to 0.2 percent sulfur, provided the KIO\(_3\) concentration is 0.444 g/L. With 1.110 g KIO\(_3\)/L, multiply burette readings by 5 (½-g sample, 0.005 to 1.00-percent sulfur range).

References

Phosphorus (6S)
Perchloric Acid Digestion (6S1)
Perchloric acid is extremely hazardous and subject to explosion if improperly handled. Do not attempt this procedure unless the hazards are well understood and the laboratory is specially equipped to handle perchloric acid digestion.

Reagents
• Perchloric acid (HClO\(_4\)), 60-percent
• Concentrated nitric acid (HNO\(_3\))
• Concentrated hydrochloric acid (HCl)

Procedure
Weigh 2.000 g oven-dry soil, ground to approximately 100 mesh, into a 300-ml Erlenmeyer flask, add 30 ml 60-percent HClO\(_4\), and boil until the soil is white.
Continue boiling 20 minutes longer to ensure complete extraction. Soils high in organic matter should be pretreated with HNO₃ and HCl to destroy the readily oxidize organic matter.

**Molybdovanadophosphoric Acid Colorimetry (6S1a)**

**Apparatus**
- Spectrophotometer

**Reagents**
- Solution I. Dissolve 20 g ammonium molybdate ((NH₄)₆Mo₇O₂₄•4H₂O) in 250 ml water.
- Solution II. Dissolve 1.25 g ammonium metavanadate (NH₄VO₃) in 300 ml boiling water, cool, and add 425 ml 60-percent HClO₄. Mix solution I and II and dilute to 1 liter in a volumetric flask. Store in a brown bottle.
- Standard phosphorus solution. Weigh out 0.2194 g oven-dry KH₂PO₄ and dilute to 1 liter. This solution contains 50 ppm phosphorus.
- Concentrated nitric acid (HNO₃) for samples high in organic matter only.
- Concentrated hydrochloric acid (HCl) for samples high in organic matter only.

**Procedure**
Transfer the extract into 250-ml volumetric flasks, bring to volume, and let residue settle out. Pipette a 25-ml aliquot into a 50-ml volumetric flask, add 10 ml molybdovanadate reagent, bring to volume, and mix.

After 10 minutes, the color is fully developed on most samples and can be read at 460 mµ. Prepare a standard curve covering the range 0 to 5 ppm phosphorus in 50 ml solution. Plot on semilog paper.

**Calculations**

Total P (pct.)=(A/400)×(250/B)

where:
- A = P from curve (ppm)
- B = Volume aliquot (mL)

**Comments**
- The color developed is molybdovanadophosphoric acid and is very stable, lasting 2 weeks or more.
- To destroy organic matter in samples high in organic matter, add 15 ml HNO₃ and 5 ml HCl. When brown fumes stop coming off, add HClO₄, and follow the usual procedure.
Sediment disturbance during aliquot removal makes it impossible to take more than one aliquot a day. If more aliquots are necessary, remove the sediment by filtering the suspension into a 250-ml volumetric flask, using Whatman No. 50 filter paper.

Comparison of results by Na$_2$CO$_3$ fusion and by perchloric acid on lava samples indicates that extraction may not be complete for some silicate minerals. Extraction by HClO$_4$ should be complete on common phosphate minerals.

The volume of molybdo-vanadate reagent added is not critical but must be constant. The presence of chlorides slows down color development but does not interfere otherwise.

References
Sherman (1942); Kitson and Mellons (1944); and Jackson (1956).

Adsorption Coefficient (6S2)

Apparatus
- Automatic extractor, 24 place
- Syringes, 60 cc polypropylene. Use one sample tube and one extraction syringe per sample.

Reagents
- Extractant. Dissolve 4.5 g ammonium fluoride (NH$_4$F) and 85.6 g ammonium chloride (NH$_4$Cl) in about 4 L of distilled water, add 92 ml glacial acetic acid and 10 ml concentrated HCl, make to 8 L and mix.
- Sulfuric-molybdate-tartrate solution. Dissolve 100 g ammonium molybdate [(NH$_4$)$_6$Mo$_7$O$_{24}$•4H$_2$O] and 2.425 g antimony potassium tartrate [K(SbO)(C$_4$H$_4$O$_6$)•½H$_2$O] in 500 ml distilled water, heating if necessary but not to exceed 60 °C. Slowly add 1,400 ml concentrated H$_2$SO$_4$ and mix well. Cool, dilute to 2 L with water, and store in refrigerator in polyethylene or Pyrex bottle.
- Ascorbic acid solution. Dissolve 88.0 g ascorbic acid in distilled water, dilute to 1 L, mix, and store in glass bottle in refrigerator.
- Phosphorus stock standard, 100 ppm. Weigh 0.4394 g dried monobasic potassium phosphate (KH$_2$PO$_4$) into a 1-L volumetric flask, dissolve, and make to volume with extractant solution.
- Phosphorus working standards, 2 to 10 ppm. Pipette 2, 4, 6, 8, and 10-ml aliquots of phosphorus stock standard into a series of 100-ml volumetric flasks and make to volume with extractant solution. The standards contain 2, 4, 6, 8, and 10 ppm P.
• Saturate stock solution. Dissolve 4.394 g dried monobasic potassium phosphate (KH$_2$PO$_4$) in distilled water and make to 1 L.
• Saturate working solution. Pipette 20- and 80-ml aliquots of Saturate stock solution into two 1-L volumetric flasks. The resulting solutions contain 20 and 80 ppm P.
• Color solution. Measure 40 ml ascorbic solution and 80 ml sulfuric-molybdate-tartrate solution into 2 L of distilled water. Bring to 4 L, mix, and store in refrigerator.

**Apparatus**
- Colorimeter
- Automatic extractor
- Shaker

**Procedure**

**A. Saturation**
Weigh three 2-g subsamples of oven-dried soil into 50-ml Erlenmeyer flasks. To the first add 2 ml distilled water. To the second add 2 ml 20 ppm P solution. To the third add 2 ml 80 ppm P solution. Let stand for 1 hr then place in oven at 60 °C and dry overnight.

**B. Extraction**
To each of the dried samples in the 50-ml Erlenmeyer flasks, add 20 ml extractant reagent, and shake for 20 min (Burrell shaker). Extract samples using the automatic extractor.

**C. Developing the color**
**Standard curve.** Using 50-ml Erlenmeyer flasks, pipette aliquots from the phosphorus working standards as follows:
- Flask 1—2 ml extractant
- Flask 2—2 ml 2 ppm P
- Flask 3—2 ml 4 ppm P
- Flask 4—2 ml 6 ppm P
- Flask 5—2 ml 8 ppm P
- Flask 6—2 ml 10 ppm P

**Samples.** For each sample extracted in part B, pipette 2 ml of extract into clean 50-ml Erlenmeyer flasks corresponding with sample numbers. To all flasks, standards, and samples, add 25 ml of color solution, swirl to mix, and let stand for 15 min to allow color to develop. After color has developed fully, transfer to colorimeter tubes.
D. Reading the color

Using a wavelength setting of 880 µm, set colorimeter to 100 percent transmittance (T) with No. 1 standard containing 2 ml extractant. Read percent transmittance of remaining standards and samples. Generally, the standard curve is around the following values:

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<tr>
<th>ppm</th>
<th>%T</th>
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<tr>
<td>0</td>
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<td>2</td>
<td>77</td>
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<tr>
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<tr>
<td>8</td>
<td>33</td>
</tr>
<tr>
<td>10</td>
<td>24</td>
</tr>
</tbody>
</table>

E. Calculations

1. Develop standard curve by the least squares analysis using concentration of standards as a f(ln%t). This results in the equation:
   \[ \text{Concentration} = m(\ln%t) + b \]

2. Use this equation to determine solution concentrations of unknowns (leachate). Concentration of leachate x 10 is desorbed P in ppm of dry soil.

3. P retained of that added = P added (desorbed P at that conc. minus desorbed P at zero P addition).

4. Pa (adsorption coefficient) is the slope of the least square regression of P retained as a function of phosphorus added, f(P added).

Example

<table>
<thead>
<tr>
<th>P</th>
<th>T</th>
<th>Added P</th>
<th>T</th>
</tr>
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<tbody>
<tr>
<td>ppm</td>
<td>(%)</td>
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</tr>
<tr>
<td>10</td>
<td>24</td>
<td>80</td>
<td>45</td>
</tr>
</tbody>
</table>
1. Concentration
   \[ \text{Conc.}= -7.023(\ln\%t) + 32.5158 \]
   \[ \text{Concentration} = -7.023 (\ln84) + 32.5158 = 1.40 \]
   \[ = -7.023 (\ln73) + 32.5158 = 2.38 \]
   \[ = -7.023 (\ln45) + 32.5158 = 5.78 \]

   Desorbed P (ppm) of dry soil
   \[ = 1.40 \times 10 = 14.0 \]
   \[ = 2.38 \times 10 = 23.8 \]
   \[ = 5.78 \times 10 = 57.8 \]

2. P (ppm) retained of that added
   \[ = 0 - (14.0 - 14.0) = 0 \]
   \[ = 20 - (23.8 - 14.0) = 10.2 \]
   \[ = 80 - (57.8 - 14.0) = 36.2 \]

3. \( y = 0.4478(P \text{ added}) + 0.5083 \)
   \( P_\alpha = 0.4478 \)

References
Mehlich (1978).

Boron (6T)
Saturation Extract (6T1)
Carmine Colorimetry (6T1a)
Refer to USDA Handbook 60, method 17 (p. 100) and method 73b (p. 142).

Silicon (6V)
HF Dissolution (6V1)
Obtain extract as in 7C3.

Atomic Absorption (6V1a)

Apparatus
- Diluter
- Atomic absorption spectrophotometer

Reagents
- Standard Si solutions, 400 and 800 mg/L in HF
**Procedure**

Dilute HF extracts from 7C3 and Si standards fivefold to twentyfold with water. Compare absorbance of samples with that of standards at 252 nm.

**Calculations**

\[
\text{Si (pct.)} = \frac{(A \times 10 \times \text{dilution})}{B}
\]

where:

\[A = \text{Si (mg/L)}\]

\[B = \text{Sample weight (g)}\]

\[
\text{SiO}_2 \text{ (pct)} = (\text{pct}) \times 2.14
\]

Report on oven-dry basis.

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**MINERALOGY (7)**

**Instrumental Analyses (7A)**

**Preparation (7A1)**

The treatment to be used in preparing samples depends on analysis objective and sample composition.

**Carbonate Removal (7A1a)**

**Reagents**

- Sodium acetate (NaOAc), N, pH 5.0. Dissolve 82 g NaOAc, 27 mL glacial acetic acid L\(^{-1}\), adjust to pH 5.0.

**Procedure**

Place 5 g soil in a 90-ml centrifuge tube, add 40 ml N NaOAc, pH 5.0, and heat at 95 °C for 30 minutes, stirring occasionally. Centrifuge at 1600 rpm for 5 minutes and decant supernatant liquid. Repeat if necessary until carbonates are removed. Wash twice with N NaOAc, pH 5.0. Save decantates for calcium and magnesium analysis. One washing is enough to prepare neutral or basic noncalcareous soils for optimum hydrogen peroxide treatment.

**References**

Jackson (1956).
Organic-Matter Removal (7A1b)

Reagents

- Hydrogen peroxide (H$_2$O$_2$), 30- to 35-percent

Procedure

Transfer sample to a 250-ml tall breaker, using a minimum of water, and add 5 ml H$_2$O$_2$. When frothing subsides, heat at 90 °C. Continue to watch for frothing. Add 5- to 10-ml aliquots of H$_2$O$_2$ each hour until 25 to 30 ml H$_2$O$_2$ has been used. Wash five times, removing water by filter candles. Transfer to a 90-ml centrifuge tube.

References

Kilmer and Alexander (1949).

Iron Removal (7A1c)

Reagents

- Sodium bicarbonate (NaHCO$_3$), N
- Sodium citrate (Na$_3$C$_6$H$_5$O$_7$), 0.3 M
- Sodium dithionite (Na$_2$S$_2$O$_4$) powder

Procedure

If iron reduction is intended, add 5 ml N NaHCO$_3$ and as much as 40 ml 0.3 M sodium citrate. Heat to 70 °C (not above 80 °C) in a water bath and add 1 to 2 g Na$_2$S$_2$O$_4$, stir for 1 minute, and then stir intermittently for 15 minutes. Decant supernatant liquid and save for iron analysis. Repeat treatment as needed if soil is high in iron. Wash twice with 0.3 M sodium citrate.

References

Aguilera and Jackson (1953) and Mehra and Jackson (1960).

Disaggregation and Particle-Size Fractionation (7A1d)

Reagents

- Sodium bicarbonate (NaHCO$_3$), pH 9.5 to 10.0
- Sodium metaphosphate (NaPO$_3$). Prepare as in 3A1.

Procedure

Use NaHCO$_3$ solution or sodium hexametaphosphate-sodium bicarbonate as a dispersing agent. Use hexametaphosphate carefully with amorphous materials since phosphates may be precipitated. A dilute HCl treatment may be useful for
some highly allophanic soils. Do not use mechanical blenders for disaggregation if silt and sand are to be studied because they fracture large quartz grains. Separate sand from silt with a 300-mesh sieve and further separate the sands, using a nest of sieves (3A1). Separate clay (<2 μ) from silt by centrifuging at 750 rpm for 3 minutes (International No. 2 centrifuge with No. 240 head and solution depth of 10 cm). If silt and sand are to be studied, save these fractions in small vials after drying and weighing. If interested in separating fine clay (<0.2 μ) from the coarse clay (0.2 μ to 2 μ), centrifuge at 2400 rpm for 30 minutes with a solution depth of 10 cm. Adjust time according to temperature. Add 50-ml aliquots of clay suspension after each centrifugation until the required amount of clay is obtained. Make each suspension of coarse and fine clay to known volume and determine its concentration and the concentration of the whole clay.

References
Kilmer and Alexander (1949) and Jackson (1956).

**Particle-Size Distribution Analysis (PSDA) Pretreatment (7A1e)**
Mineralogical analysis can be performed on samples from the particle-size distribution analysis (3A1). These samples have undergone peroxyde digestion and sodium metaphosphate dispersion.

**X-Ray Diffraction (7A2)**
The minerals in soil clays of greatest interest are mostly flaky or platy, e.g., kaolinite, illite (mica), vermiculite, chlorite, and montmorillonite. They are most readily identified and distinguished from one another by observing the effect of different treatments on the interplanar spacings along the axis perpendicular to the platy surfaces. X-ray diffraction produces peaks on a chart corresponding to the various angles (2R) of goniometer from which the crystallographic spacing of the mineral or minerals can be calculated by Bragg’s law. Tables of spacings corresponding to angles have been published in U.S. Geological Survey Circular 29 (Switzer et al. 1948).

The pretreatment used to distinguish montmorillonite from vermiculite and chlorite and to identify illite is saturation of the exchange complex of the clay with magnesium and treatment with ethylene glycol or glycerol. With this treatment, montmorillonite has a distinctive interplanar spacing of 17 Angstroms (17 Å) to 19 Å. Chlorite and vermiculite keep a 14 Å spacing and mica a spacing of 10 Å. To distinguish vermiculite from chlorite and to identify kaolinite, which has a 7 Å spacing, the pretreatment consists of saturating the clay with potassium and heating on a glass slide at 500 °C. Intermediate heat treatments 110 and 250 °C, can be used to study interlayering in the collapsing minerals or other special problems. After the 500 °C treatment, vermiculite and montmorillonite collapse.
completely to 10 Å, kaolinite becomes amorphous, and chlorite still shows 14 Å and sometimes 7 Å peaks. Interstratified forms of these minerals are indicated by spacings intermediate between those of the individual components.

Clay suspensions are dried as thin films so that the plates are parallel to one another (preferred orientation). This results in greater X-ray diffraction peak intensities. For identification and semiquantitative estimation of nonplaty minerals such as quartz, feldspars, and crystalline iron and aluminum oxides, randomly oriented dry-powder samples can be used. This dry-powder method was used for nearly all analysis, including clay fractions, before 1951.

Various techniques are used to prepare the ion-saturated clays and to improve their parallel orientation. Details can be obtained from the soil survey laboratories.

References
Brindley (1951), Brown (1961), Brunton (1955), Grim (1953), Jackson (1956), and Switzer et al. (1948).

**Thin Film on Glass. Solution Pretreatment (7A2a)**

**Reagents**
- Potassium chloride (KCl), \( N \)
- Magnesium acetate (Mg(OAC)\(_2\)), \( N \)
- Magnesium chloride (MgCl\(_2\)), \( T N \)
- Glycerol, 10-percent in ethanol by volume

**Procedure**
Place an aliquot containing 50 mg clay in a 50-ml centrifuge tube. All a few ml 1 \( N \) KCl, centrifuge, and discard the clear supernatant. Combine sediments if necessary to get 50 mg in the tube. Wash four times by suspending and centrifuging in 20-ml portions \( N \) KCl. After the last washing with \( N \) KCl, wash with water until some of the clay remains suspended after centrifuging. Add a few drops of acetone or centrifuge at higher speed, or both, to flocculate the clay. Discard the supernatant. Clays are now free of chloride. Suspend the sediment in water and adjust the volume of the suspension to yield the desired weight of clay per slide. For most clays 50 mg per slide (27 by 46 mm) gives maximum intensity of reflection with minimum peeling of clay films. For amorphous clays, 25 mg per slide is adequate if glass slides are dried in a low-humidity atmosphere.

For magnesium saturation and glycerol solvation place an aliquot containing 100 mg clay in a 50-ml centrifuge tube. Wash twice with \( N \) Mg(OAC)\(_2\) acetate and then three times with \( N \) MgCl\(_2\). Wash the suspension free of chloride or until clay disperses. Place 2.5 ml clay suspension containing 50 mg clay on a glass slide (25 mg clay if the clay is amorphous). Solvate the remaining clay in the test tube with glycerol (about \( \frac{1}{2} \) ml of 10-percent glycerol in ethanol per 50 mg clay).
Mix well and pipette 50 mg clay onto the glass slide. The slide should be moist but not wet. Or prepare the glycerol slide by adding 10-percent glycerol, a drop at a time, to the slides until the clay film is moist.

**Thin Film on Glass, Resin Pretreatment (7A2b)**

**Reagents**
- Potassium-charged resin (Dowex 50W-X8)
- Magnesium-charged resin (Dowex 50W-X8)
- Glycerol, 10-percent in ethanol by volume

**Procedure**
Add ¼ teaspoon K-charged resin to 50 mg clay in a 1 ml volume in a 50-ml centrifuge tube. Mix and transfer a 1-ml aliquot to a glass slide (27 by 46 mm). Take the aliquot from the top of the suspension to avoid removing the resin.

Magnesium-clay and Mg-glycerol-clay slides can be prepared using a Mg-charged cation exchange resin. Add ½ teaspoon Mg-charged resin to the clay suspension (100 mg clay in a 4-ml volume) in a 50-ml centrifuge tube. Mix with the suspension, remove 1-ml aliquot, and place it on a glass slide. Add approximately ½ ml 10-percent glycerol in ethanol to the tube. Mix and transfer a 1-ml aliquot to a glass slide or use a Mg-clay slide for both Mg and Mg-glycerol solvated slides. Record a diffraction pattern for the Mg-saturated clay film. After solvating the clay film with 10-percent glycerol solution, record a second X-ray pattern.

**References**
Rex (1967).

**Thin Film on Glass, Sodium Metaphosphate Pretreatment (7A2c)**
Shake soil overnight in sodium metaphosphate solution (3A2). Centrifuge to separate the clay or siphon off the clay. Pipette about 50 mg clay to a glass slide (47 by 26 mm). Concentrate the clay suspension if necessary. Scan the clay film at room temperature, again after heating to 500 °C. The clay film is Na⁺ saturated. The sodium metaphosphate peaks do not interfere with peaks of the more common clay minerals in this quick check method.

**Thin Film on Tile, Solution Pretreatment (7A2d)**

**Apparatus**
- Ceramic tile (porous precipitate drying plate, sawed into 27- by 46- by 7-mm blocks)
Procedure

Prepare clay suspensions as in 7A2a except dry the suspensions on ceramic tile blocks. Clay suspensions dry in a few seconds on tile, preventing particle-size segregation. Partly immerse the Mg-saturated clay films in a 10-percent glycerol solution. The porous tile rapidly transfers the glycerol to the clay film. Blot off excess glycerol before recording the X-ray pattern.

**Thin Film on Tile, Resin Pretreatment (7A2e)**

Prepare clay suspensions as in 7A2b. Dry on ceramic tile blocks as in 7A2d. Solvate with glycerol as in 7A2d.

**Thin Film on Tile, Sodium Metaphosphate Pretreatment (7A2f)**

Prepare the sample as in 7A2c. Pipette the clay onto ceramic tile blocks as in 7A2d. Follow method 7A2c for the other treatments. Or solvate with glycerol as in 7A2d.

**Powder Mount, Diffractometer Recording (7A2g)**

Distinguishing dioctahedral and trioctahedral minerals requires random orientation of the sample. There is no completely satisfactory method for preparing a random mount, but several techniques are used.

Pack the sample in a box mount against a glass slide. When the box is full, tape the back of the box. Invert the box and remove the slide to expose the sample to X-rays. For more random packing, sprinkle the dry sample (ground to <100 mesh) on double stick tape fixed on a glass slide or on a thin film of Vaseline on a glass slide. Scan the sample by X-ray and measure the reflections with a Geiger, proportional, or other counter.

Quick checks for whole samples, particularly for nonlayered minerals, can be made with a modified powder mount. Form the sample into a thick slurry, apply to a glass slide, and let dry. This is for convenience rather than random orientation.

**Powder Mount, Camera Recording (7A2h)**

Photographic plates are still the best means of identifying minerals. Mount the sample in the center of a circular X-ray camera. Record the X-ray reflections on photographic film placed in a cylindrical film holder inside the camera. All diffraction peaks are recorded simultaneously.

**Thin Film on Glass, NaPO₃ Pretreatment II (7A2j)**

**Apparatus**

- Hypodermic syringe (1.0 cc)
- Glass slides 24 x 46 mm or 14 x 19 mm
- International No. 2 centrifuge with a No. 240 head
- 100-ml centrifuge tubes (plastic)
Reagents
- Glycerol-water mixture (1:8 glycerol-water)
- Sodium hexametaphosphate solutions

Procedure
Shake approximately 5 g oven-dried soil (<2 mm) overnight with 5 ml sodium hexametaphosphate solution (3A1) and 35 ml of water in a 100-ml centrifuge tube, centrifuge at 750 rpm for 3 min for a 10-cm suspension depth, and decant clays. Draw about 0.5 cc of clay suspension into the syringe. Expel approximately 0.2 cc of the clay suspension onto an area approximately 20 x 27 mm in a band across the middle of a 46- x 27-mm slide or expel approximately 0.1 cc of clay suspension, containing approximately 6 mg of clay, onto and covering the 14- x 19-mm slide. Prior to the deposition of the clay suspension, one small drop of glycerol-water mixture is placed on the slide which is to be solvated. Prepare four slides for X-ray diffraction: 1) Na⁺—room temperature, 2) Na⁺—solvated, 3) Na⁺—heated 2 hr at 300 °C, and 4) Na⁺—heated for 2 hr at 500 °C.

Powder Mounts (7A2k)
Two procedures are used for random orientation of mineral separates. In the first procedure, double-stick tape is affixed to a glass slide, a surplus of the sample is sprinkled onto the tape, the excess material is removed, and the slide is scanned by X-ray analysis. In the second procedure, a <2-mm soil sample is ground finer than 100 mesh prior to slide preparation. A thin film of Vaseline is applied to a glass slide, the 100-mesh sample is added, the excess removed, and the slide is scanned by X-ray analysis.

For quick check of a <2-mm sample, particularly for nonlayered minerals, a small portion is ground to less than 100 mesh and placed on a glass slide. Water is applied a little at a time until a thick slurry is formed. The slurry is allowed to dry and the slide is scanned by X-ray analysis. This method is also applicable for specific mineral separates, very fine sands or silts.

References
Brown (1961) and Jackson (1956).

Differential Thermal Analysis (7A3)
Differential thermal analysis (DTA) is a measurement of the difference in heat absorbed by or evolved from a sample of soil material and a thermally inert material as the two are heated simultaneously at a constant rate. Thermocouples are in contact with two platinum pans; one pan contains an unknown and
the other pan contains an inert material of similar composition. If a reaction
occurs, a difference in temperature is registered on a strip chart recorder or
photographically. The magnitude of the difference depends on the nature of the
reaction and amount of reacting substance in the unknown. The temperature at
which the reaction occurs identifies the substance if enough is known about the
sample to predict the possibilities.

**Apparatus**
- Columbia scientific instrument (CSI) system 200
- Mortar and pestle
- Analytical balance
- Desiccator

**Reagents**
- Reference sample, calcined kaolinite, 2 to 20 µ
- Ethyl alcohol, 95 percent
- Magnesium nitrate (Mg(NO₃)₂•6H₂O

**Procedure**

The decanted clay from 7A2i or 7A2j is air−dried, ground in alcohol to
approximately 100 mesh, and stored in a desiccator with Mg(NO₃)₂•6H₂O. A
3− to 7−mg sample is placed on a small platinum pan in the sample holder. The
temperature of the kaolinite reference sample and clay sample is increased at a
rate of 20 °C per minute to a maximum of 900 °C. The sample can be heated in
air or nitrogen.

The common endothermic reactions studied or recorded are loss of structural
water in gibbsite, goethite, and kaolin and loss of carbon dioxide in carbonates.
Change of state or rearrangement of crystal lattices can be either exothermic or
endothermic. Oxidation reactions such as burning of carbon and oxidation of
ferrous iron are exothermic.

Loss of structural hydroxyls can be measured quantitatively by calibrating
areas of peaks of known mixtures of standard minerals, as is done commonly to
determine the percentage of kaolin and gibbsite in soils. The standard curves
are prepared by running the known mixtures under the same conditions as the
unknowns. Kaolin has an endotherm at 500 to 600 °C and gibbsite, at 310 °C.
Each worker should prepare a set of standard curves.

Endotherms at about 120 °C indicate surface−adsorbed water. Montmoril-
onite produces a double peak at a low temperature if saturated with a divalent
cation. The proportion of this mineral can be estimated if samples are kept in
an atmosphere with a high (70 to 80 percent) relative humidity for 24 hr or more
before analysis. Allophane has a broad endotherm at about 160 °C.
Samples can be any well-powdered material, whole-soil, or separated fractions. Organic matter is objectionable because it produces irregular exothermic reactions that obscure the important peaks. If a clay separate is used, it must be washed free of hygroscopic salts or salts containing water of crystallization.

References
Grim (1968), McKenzie (1957), and Tan and Hajek (1977).

Thermal Gravimetric Analysis (7A4)
CSI Stone Model 10002B (7A4a)
Thermal gravimetric analysis is the detection and measurement of weight changes in a sample of soil material as the sample is being heated or cooled over a specific temperature range.

Apparatus
- CSI Stone Model 10002B used in conjunction with an RC−202 recorder−controller. Furnace is water cooled, with a rapid cooling Kanthal element. Furnace is capable of operation at temperatures of up to 1,200 °C.

Procedure
Prepare sample as described in 7A3 and place in balance pan suspended above thermocouple assembly. Heat sample at rate of 20 °C/min to desired temperature. If a weight loss occurs, it is registered on a strip chart recorder. The magnitude of the weight loss depends on the reaction and the amount of reacting substance in the unknown. The temperature at which the reaction occurs usually identifies the substance.

Infrared Analyses (7A5)
Soil or clay samples (7A2j) are incorporated into a potassium bromide (KBr) pellet for infrared analyses. Sample concentration in the pellet ranges from 0.1 to 1 percent.

Reagents
- Potassium bromide, spectroscopic grade

Apparatus
- Infrared spectrometer. Perkin Elmer Model 283.
- Pellet die
• Hydraulic press
• Analytical balance

**Procedure**

Mix 0.30 g KBr and 1 mg of sample in mortar and pestle. Transfer the mixture to the pellet die, and place die in hydraulic press. Apply 8 tons of pressure for 1 min. Place pellet in instrument holder and scan for 12 min. Peaks produced on chart recorder are used to identify the substance.

**References**


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**Optical Analyses (7B)**

**Grain Studies (7B1)**

**Grain Mounts, Canada Balsam (7B1b)**

For Canada balsam, heat slide plus balsam for 15 min at 125 °C. Add mineral grains, stir, heat for an additional 5 min, place cover glass in position and press firmly, remove slide from hot plate, and cool.

The refractive index of Canadian balsam is close to that of quartz, which helps to distinguish quartz from other colorless minerals, particularly the feldspars. Other available commercial media cover the refractive index range of 1.53 to 1.55. Piperine with a refractive index of 1.68, which is close to that of many of the common heavy minerals, is best for mounting them.

**Electron Microscopy (7B2)**

Electron microscopy gives information on particle size and morphology of clay-size particles. Evidence of clay formation or weathering can also be seen. Positive identification of halloysite often depends on observation of rolled structures under the electron microscope.

**Procedure**

Place a drop of dilute clay suspension on a 200-mesh copper grid. After drying, insert this grid in the microscope.

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**Total Analysis (7C)**

**Chemical (7C1)**

The procedures follow the standard procedures for rock analysis set forth by Hillebrand and Lundell (1929) and modified by Robinson (1920) and by Shapiro and Brannock (1956).
X-Ray Emission Spectrography (7C2)

X-ray emission spectrography is elemental analysis by measuring the X-ray fluorescence produced by bombarding a sample with high-energy X-rays. Each element yields fluorescent radiation of unique wavelengths, one of which is selected for measurement by using an analyzing crystal that diffracts according to Bragg’s law. The intensity of the fluorescent radiation is generally proportional to the amount of the element present, but this is affected by sample homogeneity, particle size, and the absorption and enhancement of radiation by any other elements present in the sample (matrix effects). These effects can be overcome or compensated for by (1) comparing the intensities with those of standards of similar composition prepared in a similar manner, (2) fusing both samples and standards in borax or lithium borate to eliminate particle-size effects and to reduce matrix effects, and (3) making matrix corrections by calculation the absorption-enhancement coefficient of the sample for the particular radiation being measured.

References

Vanden Heuvel (1965).

Surface Area (7D)
Glycerol Retention (7D1)

Apparatus

- Weighing cans

Reagents

- Glycerol, 2-percent

Procedure

Oven-dry a clay sample (about 0.2 g) at 110 °C for 2 hours. Cool and weigh. Add 5 ml 2-percent glycerol solution and mix. Heat in oven containing free glycerol at 110 °C to constant weight. Record weight.

Calculations

To calculate the percent of glycerol retained, subtract weight of oven-dry sample from weight of glycerol and oven-dry sample, divide by weight of oven-dry sample, and multiply by 100. For the surface area of noncollapsible minerals (m²/g), multiply glycerol retained by 19.1.

References

Kinter and Diamond (1958).
MISCELLANEOUS (8)

Saturation Extract, Mixed (8A)
Saturation Extract (8A1)

Apparatus
• Richards or Buchner funnels
• Filter rack or flask
• Filter paper
• Vacuum pump
• Extract containers such as test tubes or 1-oz bottles

Procedure
Transfer the saturated soil paste to a filter funnel with a filter paper in place and apply vacuum until air begins to pass through the filter. Collect the extract in a bottle or test tube. If carbonate and bicarbonate are to be determined on the extract, add 1 drop of 1000 ppm sodium hexametaphosphate solution for each 25 ml of extract to prevent precipitation of calcium carbonate on standing.

References
Richards (1954).

Conductivity of Saturation Extract (8A1a)

Apparatus
• Conductivity bridge
• Conductivity cell

Procedure
Determine temperature of the saturation extract obtained by methods 8A1 or 8B1. Draw the extract into the cell and read the meter. Correct for temperature and cell constant using Table 1 (Table 15, Richards 1954) and report as electrical conductivity, mmhos per centimeter at 25 °C. If the instrument fails to balance, dilute the extract 1:9 with distilled water and redetermine. The conductivity of the diluted extract is approximately one-tenth the conductivity of the saturation extract.

References
Richards (1954).
Conductivity of Saturation Extract (Quick Test) (8A1b)

Apparatus
- Extractor, miniature Richards-type (fig. 8a1b-1)
- Conductivity cell, micropipet
- Filter paper, glass fiber, 3.0 cm
- Vacuum pump

Procedure
Add 1 tablespoon soil to a 100-ml beaker or container. Make a saturated paste as in 8A. Place filter paper in recess of extractor and moisten with water. Insert the tip of the pipette conductivity cell through the other end of the extractor and into the paste. Apply suction to the cell until full. Proceed as in 8A1a.

Figure 8a1b-1.—Miniature Richards-type extractor made of polymethyl methacrylate (Lucite).

Bureau of Soils Cup, Resistance (8A2)

Apparatus
- Wheatstone bridge
- Bureau of Soils electrode cup
Procedure
Rinse the soil cup with water, dry, and fill with soil paste (8A). Jar cup to remove air bubbles, strike off excess paste so the cup is level full, and connect cup to the bridge. Record resistance (ohms) and temperature of soil paste (°F).

Calculations
Convert resistance of the soil paste in ohms to percentage of soluble salt by using the tables and formulas on pages 346–349, Soil Survey Manual.

References
Richards (1954) and Soil Survey Staff (1951).

Saturated, Capillary Rise (8B)

Apparatus
- Sand table. Mariotte bottle.
- Filter paper
- Polyethylene dish with lid

Procedure
Weigh 250 g air-dry soil into cups made from Whatman No. 52 (15-cm) filter paper and place them on a sand table wetted at 5-cm tension with water. The sand table used consists of two nested plastic dishpans. The outer pan holds distilled water, which is kept at a constant level by a Mariotte bottle. The inner pan, containing medium to fine (35 to 80 mesh) pure quartz sand, rests on rubber stoppers and is suspended in the distilled water. Its perforated bottom is covered with a fine cloth-mesh screen that permits water to move upward by capillarity through the sand to the table surface. The sand on the table surface is then smoothed and covered with an absorbent paper towel. Lightweight porous firebricks can be used in place of the sand table.

Keep the samples on the sand table 16 to 18 hours, remove them, and weigh. Water adsorption drops rapidly after an initial wetting of 2 hours and the rate becomes very slow after 6 to 9 hours. Moisture moves toward the top and center of the sample, which is wetted last, ensuring retention of soluble salts in the soil. Calculate moisture at saturation from the wet- and dry-soil weights, correcting for the wet and dry filter paper weights. Add air-dry moisture percentage to moisture at saturation and report on oven-dry basis. After the wet weighing, transfer the sample to a pint polyethylene refrigerator dish, mix briefly with a spatula, and determine the pH. Keep a lid on the dish whenever possible to reduce evaporation.
References
Longenecker and Lyerly (1964).

Saturation Extract (8B1)
Proceed as in 8A1, using the saturated paste obtained by method 8B.

Conductivity of Saturation Extract (8B1a)
Proceed as in method 8A1a except use saturation extract obtained by method 8B1.

Reaction pH (8C)
Soil Suspensions (8C1)
Water dilution (8C1a)

Procedure
For 1:1 dilution add an equal weight of water to 20 or 30 g soil in a 50-ml beaker or paper cup. Stir at regular intervals for about an hour. Measure pH of the soil suspension with a glass electrode, stirring well just before immersing the electrodes in the suspension. For other dilutions vary the amount of soil, keeping the volume of water constant.

KCl (8C1c)

Procedure
Proceed as in method 8C1a except use N KCl instead of water.

CaCl₂ (8C1e)
Proceed as in 8C1a except use 0.01 M CaCl₂. This procedure can be combined with 8C1a by adding an equal volume of 0.02 M CaCl₂ to the soil suspension prepared for the water pH. Stir twice at 15-minute intervals before reading. The soil-solution ratio will be 1:2, but the pH difference between 1:1 and 1:2 suspensions is negligible.

References
Schofield and Taylor (1955) and Peech (1965).
LITERATURE CITED


APPENDIX

Figure 1.—Laboratory data sheet for Hayter silt loam, McCreary County, KY.

Figure 2. —Profile description of Hayter silt loam, McCreary County, KY.
Figure 1.—Laboratory data sheet for Hayter silt loam, McCreary County, KY.

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<th>CEC</th>
<th>6G1d</th>
<th>Base saturation</th>
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<td>6O2b Mg</td>
<td>6P2a Na</td>
<td>6Q2a K</td>
<td>Ext. acidity</td>
<td>S3a cations</td>
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**Figure 2.—Profile description of Hayter silt loam, McCreary County, KY.**

<table>
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<tr>
<th>Horizon and Beltsville Lab. No.</th>
<th>Description</th>
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<td>01</td>
<td>1-½ to 0 inches; hardwood leaf litter.</td>
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<td>Ap1</td>
<td>0 to ½ inch; very dark grayish brown (10YR 3/2) silt loam; moderate fine granular structure; very friable; 12 percent sandstone fragments (&gt;3 in. diameter); many roots; pH 7.0.</td>
</tr>
<tr>
<td>Ap2 63776</td>
<td>½ to 5 inches; brown (10YR 4/3) silt loam; weak medium granular structure; very friable; 12 percent sandstone fragments; many roots; pH 5.0.</td>
</tr>
<tr>
<td>B1 63777</td>
<td>5 to 10 inches; brown (7.5YR 4/4) silt loam; weak to moderate fine subangular blocky structure; friable; 25 percent sandstone fragments; many roots; pH 5.0.</td>
</tr>
<tr>
<td>B21 63778</td>
<td>10 to 19 inches; brown (7.5YR 4/4) silt clay loam/silt loam; moderate medium blocky structure; friable; 25 percent sandstone fragments; common roots; pH 5.0.</td>
</tr>
<tr>
<td>B22t 63779</td>
<td>19 to 34 inches; brown to dark brown (7.5YR 4/4, 3/2) silty clay loam; moderate medium blocky structure; friable; common clay films; 20 percent sandstone fragments; few roots; pH 5.0.</td>
</tr>
<tr>
<td>Layer (ID)</td>
<td>Depth Range</td>
</tr>
<tr>
<td>------------</td>
<td>-------------</td>
</tr>
<tr>
<td>B23t 63780</td>
<td>34 to 48 inches; brown (7.5YR 4/4) silty clay loam; moderate medium subangular blocky structure; friable to firm; 30 percent sandstone fragments; common clay films; few roots; pH 5.0.</td>
</tr>
<tr>
<td>B3t 63781</td>
<td>48 to 60 inches; brown (7.5YR 5/4) silty clay loam; weak to moderate medium subangular blocky structure; friable to firm; common clay films; 35 percent sandstone fragments; few roots; pH 5.0.</td>
</tr>
</tbody>
</table>

Notes: Colors are given for moist soil. The B21 and B23t layers were sampled for the Bureau of Public Roads. Reaction was determined by Soiltex.
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