

Designation: D5744 – $13^{\varepsilon 1}$

Standard Test Method for Laboratory Weathering of Solid Materials Using a Humidity Cell¹

This standard is issued under the fixed designation D5744; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ε) indicates an editorial change since the last revision or reapproval.

 ϵ^1 NOTE—Editorially updated Research Report footnote information in Section 13 in December 2013.

1. Scope

1.1 This kinetic test method covers a laboratory weathering procedure that (1) enhances reaction-product transport in the aqueous leach of a solid material sample of specified mass, and (2) measures rates of weathering-product mass release. Soluble weathering products are mobilized by a fixed-volume aqueous leach that is performed and collected weekly. Leachate samples are analyzed for pH, alkalinity/acidity, specific conductance, sulfate, and other selected analytes.

1.1.1 This test method is intended for use to meet kinetic testing regulatory requirements for mining waste rock and ores sized to pass a 6.3-mm (0.25-in.) Tyler screen.

1.1.2 Interlaboratory testing of this method has been confined to mine waste rock. Application of this test method to metallurgical-processing waste (for example, mill tailings) is outside the scope of the test method.

1.2 This test method is a modification of a laboratory weathering procedure developed originally for mining wastes (1-3).² However, it may have useful application wherever gaseous oxidation coupled with aqueous leaching are important mechanisms for contaminant mobility.

1.3 This test method calls for the weekly leaching of a well-characterized solid material sample (weighing at least 1000-g), with water of specified purity, and the collection and chemical characterization of the resulting leachate. Test duration is determined by the user's objectives of the test.

1.4 As described, this test method may not be suitable for some materials containing plastics, polymers, or refined metals. These materials may be resistant to traditional particle size reduction methods. 1.5 Additionally, this test method has not been tested for applicability to organic substances and volatile matter.

1.6 This test method is not intended to provide leachates that are identical to the actual leachate produced from a solid material in the field or to produce leachates to be used as the sole basis of engineering design.

1.7 This test method is not intended to simulate site-specific leaching conditions. It has not been demonstrated to simulate actual disposal site leaching conditions. Furthermore, the test is not designed to produce effluents that are in chemical equilibrium with the solid phase sample.

1.8 This test method is intended to describe the procedure for performing the laboratory weathering of solid materials. It does not describe all types of sampling and analytical requirements that may be associated with its application.

1.9 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.9.1 *Exception*—The values given in parentheses are for information only.

1.10 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:³
D75 Practice for Sampling Aggregates
D276 Test Methods for Identification of Fibers in Textiles
D420 Guide to Site Characterization for Engineering Design and Construction Purposes (Withdrawn 2011)⁴

¹ This test method is under the jurisdiction of ASTM Committee D34 on Waste Management and is the direct responsibility of Subcommittee D34.01.04 on Waste Leaching Techniques.

Current edition approved Sept. 1, 2013. Published September 2013. Originally approved in 1996. Last previous edition approved in 2012 as D5744-12. DOI: 10.1520/D5744-13E01.

 $^{^{2}}$ The boldface numbers in parentheses refer to the list of references at the end of this standard.

³ For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards*volume information, refer to the standard's Document Summary page on the ASTM website.

⁴ The last approved version of this historical standard is referenced on www.astm.org.

- D653 Terminology Relating to Soil, Rock, and Contained Fluids
- D737 Test Method for Air Permeability of Textile Fabrics
- D1067 Test Methods for Acidity or Alkalinity of Water
- D1125 Test Methods for Electrical Conductivity and Resistivity of Water
- D1193 Specification for Reagent Water
- D1293 Test Methods for pH of Water
- D1498 Test Method for Oxidation-Reduction Potential of Water
- D2234/D2234M Practice for Collection of a Gross Sample of Coal
- D3370 Practices for Sampling Water from Closed Conduits
- E276 Test Method for Particle Size or Screen Analysis at No. 4 (4.75-mm) Sieve and Finer for Metal-Bearing Ores and Related Materials
- E877 Practice for Sampling and Sample Preparation of Iron Ores and Related Materials for Determination of Chemical Composition and Physical Properties
- E1915 Test Methods for Analysis of Metal Bearing Ores and Related Materials for Carbon, Sulfur, and Acid-Base Characteristics
- E2242 Test Method for Column Percolation Extraction of Mine Rock by the Meteoric Water Mobility Procedure
- D5744 Test Method for Laboratory Weathering of Solid Materials Using a Humidity Cell
- E691 Practice for Conducting an Interlaboratory Study to Determine the Precision of a Test Method

3. Terminology

3.1 Definitions:

3.1.1 *acid-producing potential, AP, n*—maximum potential for a solid material sample to produce acidic effluent can be determined based on the total sulfur present in the sample.

3.1.1.1 Discussion—It is assumed that this sulfur is present as iron sulfides (for example, pyrite) (4). This assumption leads to overestimation of the acid-producing potential of samples containing non-ferrous sulfide minerals such as galena (PbS) or non-acid-producing sulfur-bearing minerals such as gypsum (CaSO₄). The AP is commonly converted to the amount of calcium carbonate required to neutralize the resulting amount of the acidic effluent produced by the oxidation of contained iron sulfide minterals; it is expressed as the equivalent tonnes of calcium carbonate per 1000 tonnes of solid material (3). The AP is, therefore, calculated by multiplying the percent of sulfur contained in the material by a stoichiometric factor of 31.2 (5).

3.1.2 *interstitial water*, *n*—residual water remaining in the sample pore spaces at the completion of the fixed-volume weekly leach.

3.1.3 *leach*, *n*—weekly addition of water to solid material that is performed either dropwise or by flooding for a specified time period.

3.1.4 *loading*, *n*—mass of a chemical species, which is the product of the species concentration and the mass of the weekly leachate collected.

3.1.5 *mill tailings*, *n*—finely ground ore processing waste (commonly passing a $150-\mu m$ [100-mesh screen]) resulting from the mill processing of ore.

3.1.6 *neutralizing potential, NP, n*—capacity of a solid material sample to neutralize an acidic effluent while maintaining a drainage pH of at least 6.0. NP is expressed in terms of tonnes of calcium carbonate equivalent per 1000 tonnes of solid material (3).

3.1.6.1 *Discussion*—NP can be estimated using several techniques, including the following (1) determining the amount of calcium and magnesium carbonate in the sample; (2) digesting the solid material with an excess of standardized acid and back titrating with a standardized base to measure and convert the residual acid to calcium carbonate equivalents (2, 6); and (3) determining the carbonate carbon content in the sample (for example Test Method E1915 acid base classification).

3.1.6.2 *Discussion*—The AP and NP are specifically applicable to the determination of AP from mining wastes comprised of iron-sulfide and carbonate minerals. These terms may be applicable to any solid material containing iron-sulfide and carbonate minerals.

3.1.6.3 *Discussion*—Calcium plus magnesium carbonate determination generally provides a reasonably accurate NP quantification for samples in which carbonate minerals are present. Digestion and back-titration techniques generally overestimate the capacity of mine waste samples to neutralize acid while maintaining drainage pH \geq 6.0. These techniques can yield negative values if there is excess soluble acidity on the sample. Carbonate-carbon determinations will overestimate the capacity of mine-waste samples to neutralize acid if they contain metal carbonate minerals that are not net neutralizing (for example, iron carbonates such as siderite [FeCO₃] (7).

3.1.6.4 *Discussion*—AP and NP comprise most acid-base classifications and these two components have historically been determined by several different analytical methods (7). However, only one acid-base classification is currently an ASTM standard, Test Method E1915. Test Method E1915 uses either pyrolysis or chemical treatment of the mine-waste sample to speciate and quantify sulfide-sulfur and carbonate-carbon concentrations, which are expressed as acid-generating potential (AGP) and acid neutralizing potential (ANP), respectively. Through this speciation, it provides a better estimate of acid generation than historic AP determinations in which non-ferrous and non-acid-generating sulfur minerals are present (for example, galena [PbSO₄] and gypsum [CaSO₄], respectively.

3.1.7 *run-of-mine, adj*—usage in this test method refers to ore and waste rock produced by excavation (with attendant variable particle sizes) from open pit or underground mining operations.

3.1.8 *waste rock, n*—rock produced by excavation from open pit or underground mining operations that has an economic mineral content less than a specified economic cutoff value for metallurgical processing.

4. Summary of Test Method

4.1 This laboratory-weathering procedure is designed to enhance the mass release of acidity/alkalinity, metals, and other pertinent analytes from a sample of solid material weighing at least 1000 g. This is done by providing conditions conducive to sample oxidation and then leaching the sample with a fixedvolume aqueous leach. Ratio of leach volume to sample mass is 0.5 : 1 or 1 : 1 depending upon the efficiency of sample wetting and amount of effluent required for chemical analyses. The weekly effluent produced is characterized for dissolved weathering products. This test method is performed on each sample in a cylindrical cell. Multiple cells can be arranged in parallel. This configuration permits the simultaneous testing of multiple splits of the same solid material sample, or of solid material samples each characterized by different compositions.

4.2 Two protocol options (Options A and B) comprise the test procedure, and these options differ only in the way that the oxygen is supplied to samples in the individual humidity cells. Option A protocol calls for weekly cycles composed of three days of dry air (less than 10 % relative humidity) and three days of water-saturated air (approximately 95 % relative humidity) pumped up through the sample, followed by a leach with water on Day 7. Option B protocol differs from Option A in that each cell is stored for six days under conditions of controlled and relatively constant temperature and humidity, and oxygen is supplied to the sample by diffusion (and possibly advection) of ambient air rather than by pumping. Although a test duration as short as 20 weeks may be suitable for some samples, more recent research indicates that a test duration well beyond 20 weeks may be required depending upon the objectives of the test (8, 9).

5. Significance and Use

5.1 The laboratory weathering procedure will generate data that can be used to: (1) determine whether a solid material will produce an acidic, alkaline, or neutral effluent, (2) identify solutes in the effluent that represent dissolved weathering products formed during a specified period of time, (3) determine the mass of solute release, and (4) determine the rate at which solutes are released (from the solids into the effluent) under the closely controlled conditions of the test.

5.2 Data generated by the laboratory weathering procedure can be used to address the following objectives: (1) determine the variation of drainage quality as a function of compositional variations (for example, iron sulfide and calcium+magnesium carbonate contents) within individual mine-rock lithologies, (2) determine the amount of acid that can be neutralized by the sample while maintaining drainage pH \geq 6.0 under the conditions of the test, (3) estimate mine-rock weathering rates to aid in predicting the environmental behavior of mine rock, and (4) determine mine-rock weathering rates to aid in experimental design of site-specific kinetic tests.

5.3 The laboratory-weathering procedure provides conditions conducive to oxidation of solid material constituents and enhances the transport of weathering reaction products contained in the resulting weekly effluent. This is accomplished by controlling the exposure of the solid material sample to such environmental parameters as reaction environment temperature and application rate of water and oxygen.

5.4 Because efficient removal of reaction products is vital to track mineral dissolution rates during the procedure, laboratory leach volumes are large per unit mass of rock to promote the rinsing of weathering-reaction products from the mine-rock sample. A comparison of laboratory kinetic tests with field tests has shown that more reaction products from mineral dissolution are consistently released per unit weight and unit time in laboratory weathering tests (9). For example, sulfate release rates observed in laboratory tests of metal-mine rock have been reported to be 3 to 8 times those for small-scale field test piles of Duluth Complex rock (10), and from 2 to 20 times those for small-scale field test piles of Archean greenstone rock (11). A greater increase is anticipated when laboratory rates are compared with field rates measured from operational waste-rock piles.

5.5 Fundamental assumptions governing Options A and B of the procedure:

5.5.1 Option A-An excess amount of air pumped up through the sample during the dry- and wet-air portions of the weekly cycle reduces the potential for oxidation reaction rates being limited by low-oxygen concentrations. Weekly leaches with low ionic strength water promote the removal of leachable mineral dissolution products produced from the previous week's weathering cycle. The purpose of the three-day dry-air portion of the weekly cycle is to evaporate some of the water that remains in the pores of the sample after the weekly leach without totally drying out the sample. Consequently, sample saturation is reduced and air flow is enhanced. During the dry-air portion of the cycle, the oxygen diffusion rate through the sample may increase several orders of magnitude as compared to its diffusion rate under more saturated conditions of the leach. This increase in the diffusion rate under neardryness conditions helps promote the oxidation of such constituents as iron sulfide. Additionally, evaporation from the three days of dry air increases pore water cation/anion concentrations and may also cause increased acidity (for example, by increasing the concentration of hydrogen ion generated from previously oxidized iron sulfide). Increased acid generation will enhance the dissolution of additional sample constituents. As evaporation continues, the remaining water may become over-saturated with respect to some mineral phases, consequently causing them to precipitate. Some precipitated minerals are potential sources of acidity when re-dissolved (for example, melanterite, $FeSO_4 \cdot 7H_2O;$ and jarosite, $K_2Fe_6(OH)_{12}(SO_4)_4$). Compared to the three days of dry air where the pore-water mass decreases over time, the wet (saturated)-air portion of the weekly cycle helps maintain a relatively constant mass of pore water in the sample (12). This may help promote some diffusion of weathering products (for example, re-dissolved precipitation products) in the remaining pore water without totally saturating the sample and adversely affecting oxygen diffusion.

Note 1—Under idealized conditions (that is, infinite dilution in air and water), published oxygen diffusion rates in air are five orders of magnitude greater than in water ($0.178 \text{ cm}^2 \text{ s}^{-1}$ versus $2.5 \times 10^{-5} \text{ cm}^2 \cdot \text{s}^{-1}$ at 0 and 25° C, respectively) (13).

5.5.2 *Option B*—In contrast to Option A, Option B protocol does not include dry air or wet air introduction to the humidity cells during the weekly cycle. Instead, Option B requires that temperature and relative humidity be maintained within a constant range by storing the cells in an environmentally

controlled enclosure during the 6 days following the weekly 500- or 1000-mL leach. Consequently, oxygen is delivered to the cells by diffusion (and possibly advection) of ambient air, rather than by pumping. Because it lacks a dry-air cycle, more interstitial water is retained in the Option B sample than in the Option A sample during the weekly cycle. Furthermore, the interstitial water content for Option B is more constant than that in Option A during the weekly dry-air cycle. In addition, the interstitial water content for Option B is less variable over the course of testing than that in Option A (14).

5.6 This test method has been conducted on metal-mine wastes to classify their tendencies to produce acidic, alkaline, or neutral effluent, and to measure the concentrations of selected inorganic components leached from the waste (2-3, 14-16).

Note 2—Interlaboratory testing of this method to date has been confined to mine waste rock. The method has not been tested for applicability to metallurgical-processing waste. Although the method has been applied by some practitioners to finely ground metallurgicalprocessing wastes such as mill tailings, those materials were not included in the interlaboratory testing of the method. Consequently, modifications of this method might be necessary to deal with problems associated with finely ground materials, which would make this method as written, inappropriate for kinetic testing of finely ground materials. For kinetic testing of finely ground materials, please refer to the biological acid production potential method in the appendix of Test Methods E1915 or other kinetic methods accepted by the regulatory jurisdiction.

5.7 The following are examples of parameters for which the scheduled weekly, semi-monthly, or monthly collected effluent may be analyzed (see 11.5.2 for suggested effluent collection frequency):

5.7.1 pH, Eh (oxidation/reduction potential), and conductivity (see Test Methods D1293, Practice D1498, and Test Methods D1125, respectively, for guidance);

5.7.2 Alkalinity/acidity values (see Test Methods D1067 for guidance); dards televised acceleration of the second standards st

5.7.3 Cation and anion concentrations;

5.7.4 Metals and trace metals concentrations.

5.8 An assumption used in this test method is that the pH of each of the leachates reflects the progressive interaction of the interstitial water with the acid-generating or acid-neutralizing capacity, or both, of the solid material under specified laboratory conditions.

5.9 This test method produces leachates that are amenable to the determination of both major and minor constituents. It is important that precautions be taken in sample collection, filtration, preservation, storage, and handling to prevent possible contamination of the samples or alteration of the concentrations of constituents through sorption or precipitation.

5.10 The leaching technique, rate of leach water addition, liquid-to-solid ratio, and apparatus size may not be suitable for all types of solid material.

5.11 Notable differences have been observed between Option A and Option B protocols:

5.11.1 Water retention in the solid-material sample between weekly leaches is more variable for Option A than in Option B; for Option A, standard deviations from the mean water retention can range from 20 to 60% of the mean value; comparable values for Option B have been reported at less than 9% (14).

5.11.2 Greater water retention in Option B cells may favor dissolution of, and consequent acid neutralization by, magnesium-bearing minerals; increased retention may facilitate transport of acidic reaction products from iron-sulfide minerals to magnesium-bearing minerals (14).

5.11.3 Comparisons of sulfate mass release from the same sample subjected to Option A and Option B protocols indicate no significant difference in sulfate concentration as a result of water-retention variation between protocols (14). This suggests the increased water retention of Option B does not limit oxygen diffusion to the extent that sulfide mineral oxidation rates are reduced (14). However, samples containing greater than 7 % sulfur have not as yet been subjected to comparable Option A and Option B protocol studies.

Note 3—Examples of products from the test include the following: (1) effluent pH, acidity/alkalinity, and specific conductance; (2) cumulative mass release of individual solutes; and (3) release rates for individual solutes (for example, the average release of μ g sulfate/g of solid material sample/week). The dissolution time required for depletion of estimated NP and the subsequent duration of acid generation can be estimated using the values generated in items (2) and (3) above (15).

6. Apparatus

OPTIONS A AND B

6.1 *Humidity Cell*—A modified column constructed of materials suitable to the nature of the analyses to be performed (see Practices D3370 for guidance). Multiple humidity cells can be arranged in an array to accommodate the simultaneous laboratory weathering of different solid material types (Fig. 1). Two different sets of humidity cell dimensions are used to accommodate particle size differences present in the solid material:

6.1.1 Cells having suggested dimensions of 10.2-cm (4.0in.) inside diameter (ID) by 20.3-cm (8.0-in.) height can be used to accommodate coarse solid material samples that have been either screened or crushed to 100 % passing 6.3 mm (0.25 in.).

6.1.2 Cells with suggested dimensions of 20.3-cm (8.0-in.) ID by 10.2-cm (4.0-in.) height can be used to accommodate solid material samples that pass a 150-µm (100-mesh) screen.

Note 4—Some coarse solid material samples may break down into finer-grained weathering products that could inhibit airflow and result in material being ejected from the cell during Option A's dry-air cycle. Consequently, use of the 20.3-cm ID cell rather than the 10.2-cm ID cell may be more appropriate (9). It should be noted that there are no published ruggedness testing results for this cell.

Note 5—For Option A, if samples are to be tested in the 20.3-cm ID cell, the air-entry port to the 20.3 cm ID cell needs to be moved from beneath the sample to just slightly above the sample so that air flow is directed across the sample surface rather than attempting to infiltrate the sample up through its bottom surface. The air-exit port is centered in the lid.

6.1.3 For cell wall thicknesses, 0.635-cm (0.25-in.) and 0.318-cm (0.125-in) cm thickness have been used for Options A and B, respectively.

6.1.4 A perforated disk (constructed of materials suitable to the nature of analyses to be performed), approximately





FIG. 1 Side View of 16-Cell Array (Option A)

0.315-cm (0.125-in.) thick, with an outside diameter (OD) suitable to the suggested vessel ID (6.1.1 and 6.1.2) is elevated approximately 1.25 cm (0.5 in.) above the cell bottom to support the solid material sample (see Fig. 1).

6.1.5 For Option A, the cell lid and base are 1.27 cm (0.5 in.) thick and machined so they each include a lip and plug; the plug portion fits into the ID of the humidity-cell top/bottom, and the lip fits over the rim of the cell opening. A hole is drilled in the center of the lid and base and tapped to accommodate a barbed NPT fitting for attachment to flexible tubing. The tubing from the lid leads to the air-exit port bubbler described in 6.19 and 6.20. The tubing from the base drains into a collection vessel.

Note 6—Lids for Option A can have an "O"-ring seal installed (machined into the plug surface) if air leakage makes it difficult to maintain constant airflow among individual cells. Both the "O"-ring seal and the air-exit port bubbler (described in 6.20) have been helpful in maintaining airflow through individual cells of a multiple cell array during the dry- and wet-air portions of the weekly cycle. However, flow rates may still differ somewhat from cell to cell because of porosity differences between samples of differing particle-size distribution.

6.1.6 Lids for Option B do not require a barbed NPT fitting. The centered hole in the Option B lid is left open to allow for exchange of ambient air during the six-day portion of the weekly cycle. A hole is drilled in the center of the base and tapped to accommodate a barbed NPT fitting. Leachate from the cell drains directly through this fitting into a collection vessel.

NOTE 7—The cell and particle size dimensions described above are those used commonly for assessing the potential of waste-rock samples associated with metal-mining operations to produce acidic effluent. A "shoe box"-shaped cell design with similar dimensions is preferred by some researchers (6).

6.2 *Separatory-Funnel Rack*, capable of holding 500-mL or 1-L separatory funnels above the humidity cells.

6.3 *Filter Media*, such as a 12-oz/yd² polypropylene felt characterized by 22-µm (0.009-in.) diameter filaments. The media should be able to transmit dry air at a rate of 20 to 30 cfm (see Test Methods D276 and D737 for guidance).

Note 8—Caution must be used in the selection of filter media materials since they may affect the effluent pH and chemistry adversely. Both pyrex wool and quartz wool retain as much as 10 to 15 g of water per g of wool (retained water tends to re-humidify the dry-air cycle to as much as 85 % relative humidity). Additionally, pyrex wool causes the neutral effluent pH to be raised by as much as 2 pH units due to leaching of the wool (11). In addition, pyrex (borosilicate) can contribute boron if this is a constituent of interest.

6.4 *Two Riffle Splitters*, with 0.63-cm (0.25-in.) and 2.5-cm (1.0-in.) wide riffles, respectively; the riffle splitter is a commonly used device for obtaining representative splits of dry, free-flowing granular materials.

6.5 Laboratory Balance, capable of weighing to 0.1 g.

6.6 Analytical Balance, capable of weighing to 1.0 mg.

6.7 Screen, 6.3 mm (0.25 in.).

6.8 Screen, 150 mm (100 mesh).

6.9 Drying Oven—Any thermostatically controlled drying oven capable of maintaining a steady temperature of $40 \pm 2^{\circ}$ C.

6.10 *pH Meter*—Any pH meter with readability of 0.01 units and an accuracy of ± 0.05 units at 25°C; two-channel operation (that is, pH and Eh) is desirable.

6.11 *Conductivity Meter*, capable of reading in micromhos (microseimens); calibrate at 25°C.

6.12 Separatory Funnel, 500 mL or 1 L, one per each humidity cell.

6.13 Collection Vessel (vessel such as an Erlenmeyer flask or Nalgene bottle), 500 mL or 1 L, one per each humidity cell.

6.14 Volumetric Flask, 500 mL or 1 L.

OPTION A

6.15 *Digital Hygrometer/Thermometer*, with a relative humidity range of 5 to 95 %, and temperature range of -40 to 104° C (-40 to 220° F).

6.16 *Cylindrical Humidifier*, with suggested dimensions of 12.1-cm (4.75-in.) ID by 134.6-cm (53.0-in.) length. The following associated equipment are needed to provide saturated air for the three-day wet-air portion of the weekly cycle:

6.16.1 A thermostatically controlled heating element to maintain the water temperature at 25°C during the wet-air cycle.

6.16.2 An aeration stone (similar to aquarium-aeration equipment) or commercially available gas dispersion fritted cylinders or disks to bubble air into the humidifier water.

6.17 *Flow meter*, capable of delivering air to each humidity cell at a rate of approximately 1 to 10 L/min/cell.

6.18 *Oil/Water Trap*, 0.01-µm, for inclusion in the feed-air line.

6.19 *Air-Exit Port Bubbler*—A 50-mL Erlenmeyer flask with a rubber stopper containing a vent and air-inlet tube (Fig. 1). The bubbler is connected to the air exit port in the humidity cell lid with flexible tubing. This helps maintain similar positive air pressure throughout all of the humidity cells.

6.20 *Flexible-Tubing Quick Disconnect*—A fitted, two-piece connection placed in the middle of the air-exit port flexible tubing so that the bubbler can be disconnected from the humidity cell to facilitate the measurement of air flow and relative humidity.

6.21 *Desiccant Column*, 5.1-cm (2-in.) ID by 50.8-cm (20-in.) length, plastic or glass cylinder capped on both ends (one cap should be removable for desiccant replacement), with an air inlet port on the bottom and an air exit port on the top.

6.22 *Dry Air Manifold*—A cylindrical manifold constructed from 2.25-in. ID schedule 40 acrylic plastic tubing, 28 in. long and fitted with 16 NPT barbed fittings. The airline exiting the desiccant column is routed directly to the cylinder, which then supplies dry air to each cell through an airline attached to its corresponding NPT barbed fitting. The cylindrical manifold fits atop the separatory-funnel rack.

OPTION B

6.23 Environmentally-Controlled Enclosure—Any enclosure suitably sized to accommodate the number of samples being tested and associated equipment, and capable of maintaining consistent humidity (± 10 %) and temperature ($\pm 2^{\circ}$ C).⁵ 6.23.1 Temperature Control—Any commercially-available heater capable of maintaining consistent temperature within the enclosure.

6.23.2 *Humidity Control*—Any commercially-available humidifier and dehumidifier capable of maintaining consistent humidity within the enclosure.

6.23.3 Instruments to Measure Temperature and Humidity— Any commercially-available manual or digital hygrometer/ thermometer (see 6.15). Temperature should be readable to at least 1°C and relative humidity to 1 %.

6.23.4 *Fan*—Any commercially-available fan to provide air circulation within the enclosure.

7. Reagents

7.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society, where such specifications are available.⁶

7.2 *Purity of Water*—Unless otherwise indicated, references to water shall be understood to mean reagent water as defined by Type III at 18 to 27°C conforming to Specification D1193.

⁵ The tolerance ranges for humidity and temperature are the range of differences of maximum and minimum values from the mean of the respective data.

⁶ Reagent Chemicals, American Chemical Society Specifications, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, seeAnalar Standards for Laboratory Chemicals, BDH Ltd., Poole, Dorset, U.K., and the United States Pharmacopeia and National Formulary, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD

The method by which the water is prepared, that is, distillation, ion exchange, reverse osmosis, electrodialysis, or a combination thereof, should remain constant throughout testing.

7.3 *Purity of Air*—The feed air line shall contain a 0.01-µm oil/water trap in advance of the flow meter.

8. Sampling

8.1 Collect the samples to be tested using available sample methods developed for the specific industry (see Practices D75 and E877, Guide D420, Terminology D653, and Test Methods D2234/D2234M).

8.2 The sampling methodology for materials of similar physical form shall be used where no specific methods are available.

8.3 The amount of material recommended to be sent to the laboratory should be sufficient to provide 8 to 10 kg of bulk sample for splitting, analysis, and testing (see 9.3).

Note 9—Additional information on theory and methods for obtaining representative samples is contained in Pitard (16).

8.4 To prevent sample contamination or constituent loss prior to testing, store the samples in closed containers that are appropriate to the sample type and desired analyses (see Guide D420 for guidance).

8.5 The time elapsed between sample collection and subsequent humidity cell testing should be minimized to reduce the amount of sample pre-oxidation (see Practices D3370 for guidance). Report the length of time between sample collection and testing.

9. Sample Preparation

9.1 Air dry as-received bulk samples of solid material to prevent the additional oxidation of reactive minerals or compounds. If air-drying is not practicable, oven dry the solid material at a maximum temperature of 40°C for 24 h, or until a constant weight is reached.

Note 10—Oven drying at temperatures above 40° C may introduce chemical and physical changes in certain mineral species comprising the sample (9). These potential changes should be evaluated and accounted for in the analysis of the test data.

9.2 After reserving any coarse material needed for Method E2242-02 (Meteoric Water Mobility Procedure) or other possible testing and analyses, screen the air-dried bulk samples through a 6.3-mm (0.25-in.) screen in accordance with Test Method E276. Crush any oversize material so that 100 % passes the screen. For particles finer than 15.2 cm (6 in.), the generation of excessive fines can be limited by stage crushing the oversize material in three steps: (1) large jaw crusher set at 1.92 cm, (2) small jaw crusher set at 0.95 cm, and (3) roll crusher set at 0.64 cm. After each of the first two crushing steps, the -0.64 cm fraction is collected and the oversize is passed to the next crushing phase.

9.2.1 **Caution**: Crushing a bulk sample so it passes a 6.3-mm (0.25-in.) screen may change the character of the sample by artificially increasing liberation and consequent surface areas of acid-producing and acid-consuming minerals contained in the + 6.3-mm (0.25-in.) material. A suggestion for

avoiding this problem is to segregate the -6.3-mm (0.25-in.) fraction by screening rather than crushing, and to test that fraction according to the protocol and equipment described in this test method. The + 6.3-mm (0.25-in.) material can either be stage-crushed (as in 9.2), or tested separately. For example, column testing could be conducted, although no standard protocol has been established for this testing. Samples from the drill core and cuttings also present material sizing problems, which must be considered when interpreting drill core and cuttings laboratory weathering data. The drill core must be crushed to - 6.3-mm (0.25-in.) to fit the cell described in this test method. The resulting size distribution from crushing will differ from that of run-of-mine due to differences in fracture patterns inherent to blasting practices that produce run-of-mine material. By contrast, drill cuttings size fractions are commonly less than 6.3-mm (0.25-in.) due to the rotary-percussive nature of obtaining the sample. The effects of particle size distribution changes resulting from the more finely crushed sample or from rotary-drill cuttings should be considered in the interpretation of data. In particular, particle size reduction will increase specific surface area of acid-generating and acidneutralizing minerals and is likely to increase liberation of these minerals. Both of these effects will tend to increase the surface area of these minerals available for reaction. If this increase is biased towards either acid-generating or acidneutralizing minerals, the balance of acid-generating and acid-neutralizing reactions will shift.

9.3 Mix and divide the bulk sample to obtain a representative test unit with a weight in the range of 8 to 10 kg, using a riffle splitter with 1-in. (2.54-cm) chutes (perform in accordance with Test Method E877, Sampling and Preparation Procedure-Riffling). Divide the test unit into eight nominal 1-kg test specimens. Store each test specimen in a resealable plastic bag (to prevent continued atmospheric oxidation of sulfide mineral-bearing samples, samples could be vacuum sealed or refrigerated).

Note 11—The dried sample should be mixed through the riffle splitter at least once before making any splits; recombine the splits resulting from the sample mixing exercise by pouring individual splits either over each other or through the splitter again. Once the actual split is made, it is wise to re-mix it (according to the above procedure) prior to making the next split. Mixing the sample through the riffle splitter may still result in segregation of the sample. If segregation persists, use of a rotary sample divider is advised.

9.4 *Head-Sample Analysis*—Select one 1-kg test specimen at random, and crush the dried test specimen so that at least 95 % passes a 1.7-mm (10-mesh) screen, in accordance with Test Method E276.

9.4.1 Divide the crushed test specimen in half twice, using a riffle splitter with 6.35-mm (0.25-in.) chutes (in accordance with Test Method E877, Sampling and Preparation Procedure-Riffling), and select a 250-g subsample at random for head sample analysis.

9.4.2 Transfer the selected 250-g subsample to a ring and puck grinding mill and grind to a nominal 95 % passing a 150- μ m (100-mesh) screen, in accordance with Test Method E276. Use the pulverized subsample to perform the following determinations: (*1*) total sulfur, sulfide, and carbonate analyses in accordance with Test Method E1915, (2) whole rock and

trace-element chemistry analyses, and (3) mineral characterization to identify and quantify the different mineral species comprising the sample.

Note 12—Because accurate estimation of a sample's capacity to neutralize acid (NP) requires identification of carbonate mineral speciation (that is, calcite, dolomite, ferroan dolomite, siderite, and so forth), and quantification of calcium and magnesium content in iron-bearing minerals (that is, ankerite, ferroan dolomite, and siderite, and so forth), these determinations are strongly recommended. It is also recommended that sample whole-rock chemistry and mineralogy be compared to ensure that chemistry is consistent with mineralogy and vice versa. Additionally, leach extraction testing of the pre-test sample, compared with leach extraction testing of post-test sample, may be beneficial in determining the extent of solutes released by mineral dissolution and subsequently sequestered in secondary solid phases during testing (**17**).

9.5 *Screen-Fraction Analysis*—Select one 1-kg test specimen at random, and determine the particle size distribution in accordance with Test Method E276. Sieve openings of 6, 10, 28, 35, 48, 100, 200, and 270 Tyler mesh are suggested.

9.6 The following analyses are recommended:

9.6.1 Determine the total sulfur, sulfide, and carbonate contents of individual size fractions in accordance with Test Method E1915. Whole rock or trace element chemical analyses, or both, may also be performed on these fractions.

9.6.2 Determine the extent of acid-generating (for example, iron sulfide, iron sulfate) and acid-neutralizing (for example, calcium carbonate, magnesium carbonate) mineral liberation of the individual size fractions.

NOTE 13—Paragraphs 9.6.1 and 9.6.2 are recommended as best practice to quantify the amount of sulfide and carbonate minerals present in each fraction and their degree of liberation within these fractions. The objective of the analyses described in 9.6.1 and 9.6.2 is to aid the user in correlating drainage quality with solid-phase composition by providing more detailed description of the exposed surface areas of acid-producing and acidneutralizing minerals. These surface areas strongly affect the rates of acid production and neutralization. With regard to the applications of testing presented in 5.1 and 5.2, the analyses described will help: (1) identify whether the material being tested will produce acidic, alkaline, or neutral effluent; (2) determine the variation of drainage quality as a function of solid-phase composition; (3) measure the amount of estimated NP accessible in the solid; and (4) in general, aid in predicting the environmental behavior of the solid (see Refs 8, 18, and 19). To perform all of the determinations described in 9.6.1 and 9.6.2 a larger mass of sample material may be required than the stated 1 Kg (18).

9.7 Select one 1-kg test specimen at random for use in the laboratory weathering test method. Divide the test specimen into four nominal 250-g subsamples using the riffle splitter with 25.4-mm (1-in.) chutes, and label and store in resealable plastic bags until it is time to load the humidity cells.

Note 14—If the leach volume from the 1 kg sample mass does not provide sufficient effluent to meet analytical needs, a 2-kg test specimen could be used as long as the 0.5 : 1 or 1 : 1 leach-volume to sample-mass ratio is maintained. Split the sample between two cells. Record and then combine recovered volumes of the weekly effluent from both cells. Also record the combined volume.

9.8 Reserve the remaining test specimens for replicated testing or to resolve disputed results (recommend sample preservation by vacuum seal or refrigerated storage).

10. Apparatus Assembly

10.1 Option A—The humidity cells are table-mounted at a height sufficient to accommodate the placement of both the

humidifier and one collection flask for effluent collection from the bottom of each cell (Fig. 1). During the water-saturated and dry-air portions of each weekly cycle, feed air is metered to the bottom of each cell (or midway up the side in cells designed for minus 100 mesh material) at the selected rate (1 to 10 L/min). Feed air for the three-day dry-air portion is routed first through a desiccant column and then to each of the cells through a dry-air manifold (Fig. 2, Fig. 3). Feed air for the watersaturated air portion is routed through a water-filled humidifier by means of aeration stones or gas dispersion fritted cylinders/ disks, and then to each humidity cell (Fig. 2). If necessary, a water-bubbling vessel can be attached to the air-exit port of each humidity cell lid to maintain constant airflow among the individual cells (Fig. 1).

10.2 Option B—The humidity cells are mounted on a rack of sufficient height to accommodate placement of vessels for collection of effluent from the bottom of each cell. The upper portion of the rack doubles as a separatory-funnel rack, and is of sufficient height to accommodate placement of the funnel spigot above the humidity-cell lid. A simple rack of wood construction is shown in Fig. 4. Note that holes are drilled in the humidity-cell shelf to accommodate the barbed fitting (drain) that is centered at the bottom of each cell. Unlike the Option A apparatus, no air plumbing is required. Unlike Option A, Option B cells are stored in an enclosure in which temperature and humidity are controlled during the 6 days following the leach. Shelves for cell storage and space for temperature- and humidity-control equipment are required in the enclosure.

10.3 Option A and B—A separatory funnel rack is mounted on the table that holds the cells if the weekly water leach is applied dropwise (drip leach). Multiple separatory funnels (one for each cell) are held in the rack during the drip leach that is performed on the seventh day of each weekly cycle (Fig. 2, Fig. 4). The separatory funnel can be used to meter the required water volume slowly down the sides of the cell wall until the sample is flooded if the weekly leach is to be a flooded leach.

11. Procedure

OPTIONS A AND B

11.1 Cell Loading:

11.1.1 If more than one humidity cell is used at one time, label each with a sequential number, and use the same number for the matching collection vessel.

11.1.2 Weigh each humidity cell (without its lid) and each collection vessel; record the tare weights of each to the nearest 0.1 g.

11.1.3 Cut the filter media (such as 12-oz/yd^2 polypropylene described in 6.3) to the humidity cell's inside diameter dimensions so that it fits snugly yet lies flat on the perforated support.

11.1.4 Re-weigh the humidity cell, and record the resulting tare to the nearest 0.1 g; the original cell tare (11.1.2) minus the new cell tare is the weight of the filter media.

11.1.5 Transfer the contents from each of the four bags containing the 250-g samples (9.7) into the humidity cell (Fig. 5). Prior to the transfer, mix the contents of each bag by gentle



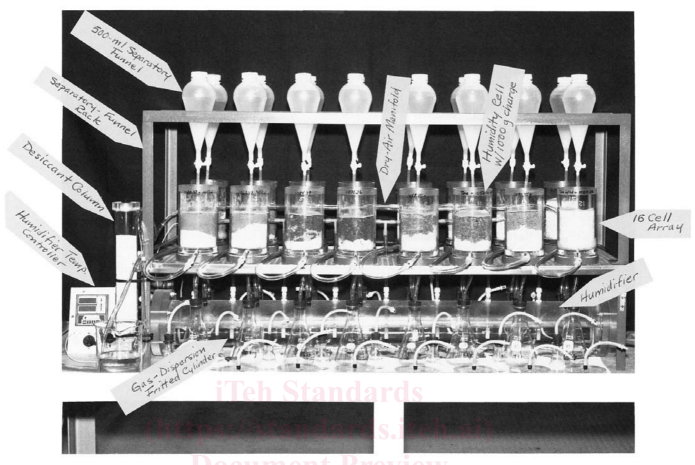


FIG. 2 Front View of 16-Cell Array (Option A) with Separatory Funnel Rack

rolling to eliminate possible stratification that may have occurred during sample storage.

11.1.6 Re-weigh the loaded cell, and record the weight to the nearest 0.1 g; the loaded cell weight minus the combined cell and filter-media tare weight is the weight of the sample charge.

11.2 First Water Leach:

11.2.1 The leach application method for the first water leach (and all subsequent weekly leaches) is the flood leach.

Note 15—Both drip and flood-leach alternatives were provided in the original D5744-96 method. However, a subsequent interlaboratory study (20) that compared drip with flood-leach alternatives determined that there was little difference in results from the drip and flood-leach applications. Results from the two laboratories participating in the study showed that the drainage pH values for the end of rate periods were within 0.2 units of the mean, and sulfate release rates were consistently within 10% of the mean. Consequently, because there was little difference between results from the two leach alternatives, and to simplify the method protocol, the flood-leach alternative is designated as the preferred water-leach application if the flood-leach application becomes impracticable.

11.2.2 The first flooded leach, designated as the Week 0 leach, initiates the humidity cell test and establishes the starting or initial characteristics of the leachate. Because the laboratory-weathering test is made up of seven-day cycles, the sample is always leached on the same day of the week (for example, Monday). On the scheduled leaching day for Week 0,

if removal of pre-leaching oxidation products is identified as a necessary component of the test objective, perform three consecutive leaches (using either a 500-mL or 1-L volume per leach) (12, 14). Report the rationale for using the three consecutive leaches. Otherwise, use a single 500 mL or 1-L volume leach. Initial selection of either the 500-mL or 1-L volume of water is dependent upon the pore volume desired, quantity of solution required for analytical purposes, and the detection limits desired or available for analyses. However, after the Week 0 leach (that is, initial 500-mL or 1-L leaches), the selected weekly volume (500 mL or 1 L) may need to be changed because the volume originally selected is either determined to be insufficient to flush the reaction products from the sample, or rates of release may be too slow to produce detectable solutes of interest. Consequently, a weekly review of the drainage quality data from the selected weekly leach volume is recommended to ensure that the selected volume is adequate for the objectives of the test. Report any changes in the weekly leach volume and document the rationale for the change.

Note 16—Prior studies of humidity-cell tests on mining wastes showed that cation and anion loadings were commonly high in the leachate recovered from the Week 0 500-mL leach. These high loadings were due to the dissolution of pre-existing soluble oxidation salts present in the sample prior to sample collection. The first 3 to 5 weeks of weekly 500-mL leaches were required to flush these pre-existing salts from the

∰ D5744 – 13^{ε1}



FIG. 3 Detail of Desiccant Column and Flow Meter

samples being studied (15). The objective of performing three consecutive leaches during the Week 0 leach is to reduce the influence of pre-existing salts on succeeding weekly drainage quality, if this influence is identified as a critical component of the test objective. It is possible that some pre-existing salts may continue to remain in the sample despite the three consecutive Week-0 leaches. Therefore, caution should be used in interpreting the rate of mass release observed during the first 3 to 5 weeks of the test.

Note 17—If objectives of the test require more effluent volume for chemical analyses than is provided by either the 500 or 1000 mL leach, run two cells of the same material as described in Note 14 while maintaining the leach-volume to sample-mass ratio as 0.5:1 or 1:1.

Note 18—Assuming a sample density of 2.6 to 3.0 g/cm³ and a porosity of 40 to 45 %, a 500-mL leach represents approximately 1.5 to 2 pore volumes and a 1000-g sample would fill a 4-in. ID humidity cell to a height of approximately 2.5 to 3.5 in. above its perforated base.

11.2.3 Fill a separatory funnel for each cell with de-ionized water using a volumetric flask. Set each separatory funnel above its corresponding cell, place the collection vessel beneath its corresponding cell, and remove the cell lid. Plug the humidity cell air-entry and effluent-drainage ports with removable waterproof caps to prevent premature drainage of the

flooded water leach (see Fig. 4). Using the separatory funnel, discharge the selected water volume slowly down the sides of the cell wall until the sample is flooded. This application method reduces hydraulic agitation of the sample surface commonly caused by pouring liquid from an open-mouthed vessel. Alternatively, flooding may be accomplished by any application apparatus (for example, a peristaltic pump) that supplies the selected volume of leachant at a reasonable rate without causing agitation and suspension of the finer fractions contained in the sample charge.

11.2.4 Allow the flooded cell to sit for a period of at least 1 h before draining the leachate into the collection vessel by removing the drainage-port plug. The leach time commences after all of the leachant has been placed in the cell. The solid material sample should be saturated and covered with leachant to a depth sufficient to maintain sample saturation. In testing mining wastes, the observed depth of leachant cover from a 500-mL flooded leach performed in 10.2-cm (4.0-in.) ID cells is approximately 2.5 cm (1.0 in.).