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Standard Guide for Preparation of Working Reference Materials for Use in the Analysis of Nuclear Fuel Cycle Materials¹

This standard is issued under the fixed designation C 1128; 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—Figure 1 was corrected editorially in February 1997.

1. Scope

1.1 This guide covers the preparation and characterization of working reference materials (WRM) that are produced by a laboratory for its own use in the analysis of nuclear materials. Guidance is provided for establishing traceability of WRMs to certified reference materials by a defined characterization process. The guidance provided is generic; it is not specific for a given material.

1.2 The information provided by this guide is found in the following sections:

	Section
Planning	6
Preparation	7
Packaging and Storage	8
Characterization	9
Statistical Analysis	10
Documentation	11

1.3 The values stated in SI units are to be regarded as the standard.

1.4 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:

- C 1009 Guide for Establishing a Quality Assurance Program for Analytical Chemistry Laboratories Within the Nuclear Industry²
- C 1068 Guide for Qualification of Measurement Methods by a Laboratory Within the Nuclear Industry²
- C 1215 Guide for Preparing and Interpreting Precision and Bias Statements in Test Method Standards in the Nuclear Industry²
- 2.2 ISO Standards:

ISO Guide 25 General Requirements for the Competence of

Calibration and Testing Laboratories³

ISO Guide 30 Terms and Definitions Used in Connection with Reference Materials³

3. Terminology

3.1 Definitions of Terms Specific to This Standard:

3.1.1 certified reference material (CRM)—a reference material with one or more property values that are certified by a technically valid procedure, accompanied by or traceable to a certificate or other documentation that is issued by a certifying body (as defined by ISO Guide 30). A certifying body is a technically competent body (organization or firm, public or private) that issues a reference material certificate (as defined by ISO Guide 30). Such an organization could be the National Institute of Standards and Technology (NIST) or the New Brunswick Laboratory. A reference material certificate is a document certifying one or more property values for a certified reference material, stating that the necessary procedures have been carried out to establish their validity (as defined by ISO Guide 30).

3.1.2 reference material $(RM)^4$ —a material or substance one or more properties of which are sufficiently well established to be used for the calibration of an apparatus, the assessment of a measurement method, or assigning values to materials (as defined by ISO Guide 30). A reference material may be referred to in this guide also as a standard, such as calibration standard or control standard.

3.1.3 working reference material $(WRM)^4$ —a RM usually prepared by a single laboratory for its own use as a calibration standard, as a control standard, or for the qualification of a measurement method (see Guide C 1068) as indicated in Fig. 1.

4. Summary of Guide

4.1 This guide covers the preparation of WRMs from nuclear fuel cycle materials. These materials are compounds and metal of uranium and plutonium, absorber materials such

¹ This guide is under the jurisdiction of ASTM Committee C-26 on Nuclear Fuel Cycle and is the direct responsibility of Subcommittee C26.8 on Quality Assurance Applications.

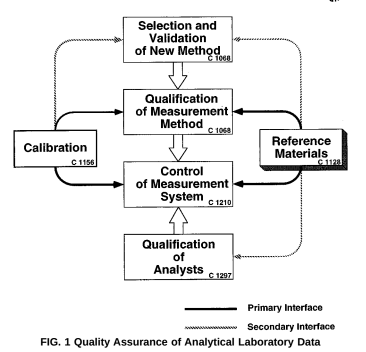
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² Annual Book of ASTM Standards, Vol 12.01.

³ Available from American National Standards Institute, 11 W. 42nd St., 13th Floor, New York, NY 10036.

⁴ It is important that some well defined indication of the uncertainty in the stated values be given in the certificate.

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as boron carbide, and cladding materials such as zirconium and stainless steel. The criteria governing the preparation of reliable WRMs are identified and discussed. Because this guide is generic, requirements and detailed information for specific nuclear materials are not given. A flow diagram to illustrate an approach to producing WRMs is given in Fig. 2.

5. Significance and Use

5.1 Certified reference materials (CRMs) prepared from nuclear materials are generally of high purity, possessing chemical stability or reproducible stoichiometry. Usually they are certified using the most unbiased and precise measurement methods available, often with more than one laboratory being involved in making certification measurements. CRMs are generally used on a national or international level, and they are at the top of the metrological hierarchy of reference materials. A graphic description of the United States nuclear measurement system is shown in Fig. 3.

5.2 Working reference materials (WRMs) need to have quality characteristics that are similar to CRMs, although the rigor used to achieve those characteristics is not usually as stringent as for CRMs. Where possible, CRMs are often used to calibrate the methods used for establishing the concentration values (reference values) assigned to WRMs, thus providing traceability to CRMs. A WRM is normally prepared for a specific application.

5.3 Because of the importance of having highly reliable measurement data from nuclear materials, particularly for control and accountability purposes, CRMs are sometimes used for calibration when available. However, CRMs prepared from nuclear materials are not always available for specific applications. Thus, there may be a need for a laboratory to prepare WRMs from nuclear materials. Also, CRMs are often too expensive, or their supply is too limited for use in the quantities needed for long-term, routine use. When properly prepared, WRMs will serve equally well as CRMs for most

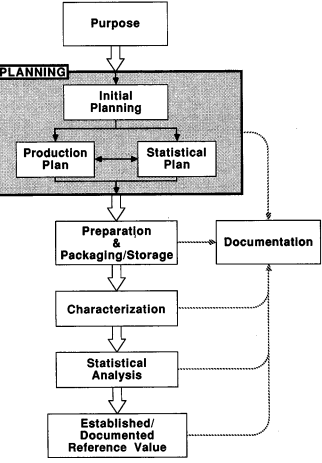


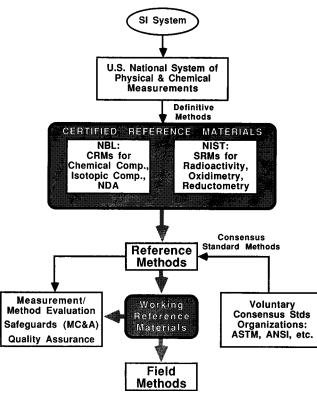
FIG. 2 Producing a Working Reference Material

applications, and using WRMs will preserve supplies of CRMs.

5.4 Difficulties may be encountered in the preparation of RMs from nuclear materials because of the chemical and physical properties of the materials. Chemical instabilities, problems in ensuring stoichiometry, and radioactivity are factors involved, with all three factors being involved with some materials. Those preparing WRMs from nuclear materials must be aware of how these factors affect preparation, as well as being aware of the other criteria governing the preparation of reliable WRMs.

6. Planning

6.1 Producing a WRM requires forethought before the work starts to ensure that the credibility of the completed WRM will be established. Planning is also important to ensure that the necessary resources are available. Time, funding, and materials can be wasted easily without thorough planning, which should include developing an outline or general scheme for preparing the WRM. The intended use of the WRM, the sources available for obtaining needed materials, and the equipment required are some areas of planning that must be considered. These considerations and others are discussed in this section in terms of initial planning, a production plan, and a statistical plan (see Fig. 2). Initial planning generally starts with the application or need for a WRM and the quantity needed. As planning





progresses into the actual preparation, a production plan and a statistical analysis plan will be developed.

6.2 Initial Planning:

6.2.1 Application of WRM-A WRM can be prepared for a single method of analysis or for several methods. For example, one might be prepared for the determination of uranium in uranium dioxide. If a standard is also required for the isotopic analysis of uranium, it might be possible to prepare and characterize that WRM for isotopic analysis as well. In situations involving the determination of impurities, it may or may not be desirable to prepare WRMs. Often, determinations of impurities do not require highly reliable results, and the preparation of a WRM might not be cost effective. A reference material lower in the metrological hierarchy could be adequate. On the other hand, during the preparation of a WRM for the determination of a major constituent, it might be possible to add desired impurities and to establish values for those impurities. This would give a multi-purpose WRM. Careful consideration should be given to the preparation of multipurpose WRMs, however, because they tend to be difficult to prepare and characterize.

6.2.2 *Quantity*—The quantity of WRM prepared will depend on such factors as the length of time required for its use, the frequency of use, the amount of material available, and its anticipated shelf life. Consideration should be given to the amount of WRM that will be needed for characterization and perhaps for archival purposes. Needs may develop during the use of a WRM such as the exchange of materials with another laboratory for an interlaboratory testing program. For this and other possible contingencies, the preparation of a quantity over the anticipated amount should be planned.

6.3 *Production Plan*—An outline should be prepared during planning that specifies how the WRM will be produced. The subjects discussed in 6.2 and in this section must be considered and addressed if appropriate. A preparation procedure should be written and included as a part of the production plan (see 7.4). The production plan must be integrated with the statistical plan (see 6.4).

6.3.1 *Materials*—The selection of materials is an important part of planning because proper selection is critical to achieving credible WRMs. Selection depends on availability (source), cost, chemical and physical properties, and stability or reproducible stoichiometry. The material selected for a WRM must be as similar as possible to the sample material in chemical and physical properties, particularly in those that will affect the method of analysis. One way to achieve similarity in composition is to prepare the WRM material by the same or similar process used to prepare the sample material. Probably the most important criterion for selection is stability. The WRM composition must be sufficiently stable to make the preparation of the WRM cost effective, and the stability must be known well enough to establish a shelf life with a high degree of confidence. Somewhat unstable materials whose stoichiometries can be reproduced easily can be used for WRMs.

6.3.2 *Equipment*—Generally, standard laboratory equipment will be involved in preparing a WRM. Analytical setups and instrumentation will be required, possibly to analyze starting materials for impurities and other constituents and certainly to analyze the prepared material during final characterization of the WRM. Depending on packaging requirements, equipment may be required for such things as sealing glass ampoules or packaging a WRM in a special atmosphere.

6.3.3 *Use*—The degree of attention given to some steps in producing a WRM may vary depending on its planned use. Usually, WRMs are used for calibration and measurement control. A common approach to producing a control standard is to take material from a batch of production material, treat it as necessary to ensure homogeneity, and establish initial measurement control limits by using the same method and conditions used for sample analysis. To produce a calibration standard, more care in preparation and rigor in characterization are required.

6.3.4 *Characterization of Materials*—Planning must provide for the characterization of materials used for a WRM. Characterization may include the analysis of starting materials for impurities and major constituents. It must include a scheme for establishing the concentration value to be assigned (reference value) to each constituent of interest. In planning for characterization, consideration must be given to the degree of reliability required for a reference value. This will involve planning for the statistical collection and analysis of characterization data (see 6.4).

6.3.5 *Packaging*—Packaging of the WRM must be planned. Decisions need to be made about dividing the WRM into portions, selecting containers, sealing containers, and using additional means to protect the integrity of the WRM. For some WRMs, it may be necessary to package soon after preparation to preserve integrity, and packaging materials and equipment must be ready then. Inadequate packaging may lead to loss of the WRM's integrity through such consequences as contamination, evaporation, and absorption.

6.4 Statistical Plan-A statistical plan for characterization must be developed during planning (see Appendix X1). Such a plan is necessary to allow an uncertainty to be determined for each reference value. The statistical plan establishes how characterization will be done regarding the sampling of the WRM, the frequency and number of measurements to be made of the WRM and any reference material to be measured with the WRM, and the order of those measurements (see 9.3 and 9.4). The validation or calibration of the measurement method to be used for characterization must be addressed in the plan also (see 9.2.3). It is essential to have a gualified statistician involved in developing the plan, and the statistician must be brought into the planning process early (see Fig. 2). Developing a statistical plan is an iterative process that will go on throughout planning, and it must be integrated with the production plan (see 6.3).

7. WRM Preparation

7.1 The objective of preparation is to make physical and chemical manipulations so as to produce a homogeneous and stable material in the form required for a WRM. For a given WRM, the physical and chemical manipulations that will be used depend on the starting material(s), the WRM form required, and the physical and chemical properties of the materials involved. Various aspects of preparation are discussed in this section.

7.2 Starting Materials—The starting materials for the preparation of WRMs may be the WRM forms desired or may be other materials that are processed into those forms. In the former case, the starting material is process material. For example, a batch of uranium dioxide pellets, boron carbide powder, or plutonium nitrate solution might be taken directly from a process run, treated as necessary, characterized, and packaged as a WRM. In the latter case, various approaches are used to produce the form desired. For example, high-purity uranium metal might be dissolved, given amounts of impurities added to the solution, and the solution converted into uranium dioxide after thorough mixing to produce a WRM with specified impurity levels.

7.3 *WRM Form*—The form of the WRM can be any stable state of the element of interest or a somewhat unstable state whose stoichiometry is easily reproducible. The forms most commonly used for nuclear materials have been oxides as powder or pellets, metal, and nitrate solutions.

7.4 *Procedure*—Write a preparation procedure using a scheme for preparing the WRM developed during the planning stage (see 6.3). The procedure must include the necessary steps for making the required chemical and physical manipulations, and it should include requirements for recording data generated during preparation. If it is planned that the reference value will be calculated based on process or make-up parameters (weights, volumes, etc.), write the procedure accordingly to minimize the possibilities of losing any material during processing (see 9.1). Procedures to illustrate the preparation of two WRM solutions are given in Appendix X2.

8. Packaging and Storage

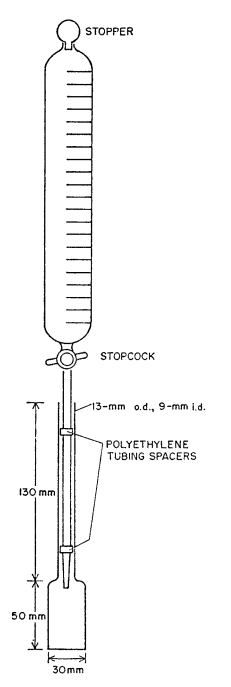
8.1 *Packaging*—Once preparation is complete, the WRM must be packaged for use. A frequent practice is to divide the WRM material into essentially equal portions or units, each of which represents enough material for a one-time use. If a WRM is sufficiently stable, it could be divided into larger portions for multiple use. There is a risk here, however, because each time a container is opened there is a potential for loss of WRM integrity. Key to packaging is to contain the WRM portions in a manner to preserve their integrity for the life of the WRM (see 6.3.5). A technique sometimes used for solutions is to evaporate each weighed portion to near-dryness in its packaging container, giving a weighed amount of the element of interest for a one-time use. Various aspects of packaging are discussed in this section. A procedure to illustrate packaging a WRM solution is given in Appendix X2.

8.1.1 *Container*—Containers often used for WRMs are screw-cap glass vials or bottles and glass ampoules. It is important that the container material is compatible chemically with the WRM matrix and that the material will not contribute to the contamination of the WRM. To avoid contamination, containers are often specially cleaned before packaging. When radioactive material such as plutonium is involved, the primary container is often packaged in a secondary or outer container to protect against radioactive contamination.

8.1.2 Addition to Container—The manner of adding WRM material to containers depends on the nature of the material, the type of container, and whether the weight of each WRM portion is required. It is exceedingly important that the WRM material be delivered into each container without any part of the material adhering to the neck or top of the container (or outside of the container), particularly when solution is added to glass ampoules that will be heat sealed. Special apparatus is sometimes used for delivery to glass ampoules (see Fig. 4 as an example). When a WRM is to be apportioned by weight, WRM material is usually added to tared containers, which are reweighed after addition. When radioactive material is involved, special care is required to keep the outsides of the containers free of contamination. Each container should be surveyed after addition, and those contaminated should be discarded.

8.1.3 *Cover Gas*—With some materials, stability is enhanced by packaging the WRM in an inert gas or dry air. A common way to do this is to package in a glove box containing the atmosphere desired. The materials most often packaged in an inert and dry atmosphere or simply in dry air are the oxides, particularly powders. This is done to give more ensurance of stability and integrity, even when an oxide is basically stable. When a special atmosphere is used, care must be taken to ensure that containers will not lose the atmosphere over the shelf life of the WRM.

8.1.4 Sealing Containers—If a special atmosphere is used as discussed in 8.1.3, the method of sealing the containers is important. For screw cap containers, sealing the caps with a sealant over the cap is one way. Using glass ampoules that are heat sealed is another approach (a procedure for sealing glass ampoules is given in Appendix X2). Glass ampoules are commonly used for solutions to avoid loss of integrity through



NOTE 1—The capacity of the buret is dependent on the number of vials to be filled.

FIG. 4 Apparatus for Filling Ampoules

evaporation. When simply closing a vial or bottle with a screw cap is satisfactory, a cap liner that provides a reasonably air-tight seal should be used.

8.1.5 *Labeling*—Each WRM container must be labeled for identification. Individual identification of each container or unit is not usually required unless each unit is uniquely identifiable by a characteristic that affects the use of the WRM, such as the net weight of the WRM in the container. As a minimum, information on a label must provide traceability to the WRM. It would be desirable to have the date of preparation and any limitations on shelf life indicated on the label. It is

essential that labels be firmly attached to the containers and that their markings be nonsmearing and nonfading. If capability exists to use a bar-code labeling technique, that approach may be desirable since more information can be added in a smaller space.

8.2 *Storage*—Although a major purpose of packaging is to preserve the integrity of WRMs, attention should also be given to how and where the packaged WRMs are stored. Exposure over time to large fluctuations in temperature, or to above-ambient temperatures, could adversely affect the container seals and the WRM materials themselves. Exposure to conditions that would damage or destroy labels, or even damage containers, must be avoided.

9. Characterization

9.1 Characterization, as discussed in this section, applies to the analysis or measurement of a prepared WRM to establish a reference value for the WRM. Characterization normally begins after the prepared WRM has been packaged. The required number of WRM units is selected, and the specified number of measurements (analyses) is made using the designated measurement method or methods. If a WRM is to be used for calibration purposes, the decision might be made to use two methods if two comparable and applicable methods are available. In some instances, the reference value is based on a make-up value in which the starting material is weighed and processed quantitatively through the preparation procedure with a final weight or volume determination. Even then, the make-up value is often confirmed or verified by measurement. The selection and use of the measurement method is briefly discussed below. An outline to illustrate the characterization of a WRM is given in Appendix X1.

9.2 Measurement Method:

9.2.1 *Type*—Often the measurement method selected is the method used for the analysis of the samples for which the WRM is prepared. If another method is used, it must be equal to or better than the sample method in terms of precision and bias.

9.2.2 Conditions of Measurement—A decision, which is based on the intended use of the WRM, must be made regarding how much care will be taken when measuring the WRM. If the WRM is to be used as a control standard, the measurements might be made under the routine conditions used for sample analysis. The alternative is to make the measurements under more rigidly controlled conditions. For example, the method might be qualified first using the criteria given in Guide C 1068, and then only highly qualified analysts might be permitted to make the measurements.

9.2.3 Validation of Method—Before the measurement method is used for characterization, the method must be validated in the sense that it is calibrated by using a selected calibration standard and by following a prescribed calibration procedure. The ideal standard would be a CRM that has the same matrix as the WRM. A second choice could be a CRM with a different matrix but still certified for the element of interest. If possible, the calibration standard should have a higher standing in the metrological hierarchy of standards than the WRM will have. A calibration procedure should be prepared and integrated with the statistical plan (see 6.4).

9.3 *Sampling*—After preparation and packaging, there will be a finite number of WRM units that will be sampled for characterization. Sampling must be based on a random selection of the required number of units. Consideration should be given to minimizing the number of units taken for characterization to maximize the number available for the planned use of the WRM. Sampling is addressed in the statistical plan.

9.4 *Measurement Scheme*—There are various factors that could be considered when devising a measurement scheme. In addition to the possibility of using more than one measurement method, more than one analyst might be used. Instead of two different methods, there might be duplicate setups for one method. The degree of replication of each step in the analysis and the time period for the analysis would be considerations. These and other factors will affect the measurement scheme and the amount of work required. A balance must be decided upon between the cost of characterization and the degree of reliability desired. The measurement scheme is addressed in the statistical plan.

10. Statistical Analysis

10.1 A statistical analysis of the characterization data is made to derive the reference value and to determine an appropriate uncertainty for that value. The statistical analysis is based on the statistical plan, and it should be done by a statistician if possible. The meaning of the uncertainty value assigned to the reference value should be defined (see Guide C 1215).

11. Documentation

11.1 Records generated during the preparation and characterization of a WRM provide the documentary evidence and support for the technical interpretation, judgments, and decisions regarding the quality of the WRM. Records provide the historical evidence needed for future reviews and evaluations should the credibility of the WRM ever be questioned. They provide linkage (traceability) between the WRM's assigned value(s) and a nationally recognized measurement base as represented by CRMs or other recognized standards (see 9.2.3). Thus, consideration must be given to the records that will be generated and retained from planning through characterization. The types of records that might be generated and the record controls that should be established are discussed in this section.

11.2 *Types*:

11.2.1 *Preparation*—The more obvious types of preparation records are the preparation procedure and the data generated during preparation such as weights of materials, volumes of solutions, blending or mixing times, and temperature and other process conditions used (see 7.4). Information about starting materials such as source, treatment history, composition, and physical characteristics are important record items. Other types are data generated from preliminary or test preparation work, process control data, literature references supporting the preparation techniques used, and names of those doing the work and dates the work was done.

11.2.2 *Characterization*—Characterization data are important record material. The method(s) used for characterization must be documented, and the statistical plan used to obtain and evaluate the characterization data must be included in the records. Those doing the characterization work and the statistical evaluation of data should be identified in the records, as well as the dates the work was done.

11.2.3 Other Records—There may be information generated during the planning stage that should become records. An example might be memos or letters that initiated planning for the WRM and that contain documentation of the need for the WRM. External documents associated with the production of the WRM could be useful records. Examples are CRM certifications and NIST certifications for standard weights. There will be records related to packaging and storage that must be included. These are the records identifying individual units of the packaged WRM and information related to storage and shelf life requirements. Basically, any piece of information or document that would help support the credibility of the WRM should be considered for inclusion in the records.

11.3 *Record Control*—Records generated for a WRM should be incorporated into the laboratory's records management system (see Guide C 1009). It is important to establish a retention time for each type of record to preserve traceability and documentary evidence over the expected life of the WRM, and perhaps for some time beyond that point (see 11.1). The record system must provide for easy retrievability of the records and adequate storage facilities to protect the records from damage. If adequate records are not available when needed, loss of credibility is very possible.

12. Keywords

12.1 certified reference material (CRM); characterize; documentation; package; reference material (RM)

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APPENDIXES

(Nonmandatory Information)

X1. CHARACTERIZATION OF A WRM

X1.1 The purpose of this appendix is to illustrate, through an example, the characterization of a prepared WRM as a calibration standard and as a control standard. It is assumed that the WRM was prepared by following this guide for its planning, preparation, and packaging. That includes the preparation of a statistical plan, in which a statistician was closely involved. The measurement schemes presented are for illustration and would normally be a part of the statistical plan.

X1.2 Prepared WRM—For the purpose of this illustration, the WRM was prepared from uranium dioxide taken from a fuel fabrication process in which fuel pellets are produced from uranium dioxide. The WRM was packaged in 1-g units. It was decided to prepare 500 units. The element of interest is uranium, for which the reference value will be determined by characterization. It is planned that the WRM will be used during the routine analysis of uranium dioxide.

X1.3 *Method of Sample Analysis*—The method of analysis is the Davies-Gray method, which requires putting solid samples into solution for the uranium measurement (titration). The following conditions are assumed for this illustration. Samples are dissolved one day and titrated the next day. On the average, 20 titrations can be made per day by a single analyst. The measurement is based on the titration of uranium with NIST potassium dichromate, which is a CRM. Under routine conditions, the method is capable of a relative standard deviation (RSD) of 0.15 % for 100-mg samples, with no significant bias. A high-precision version of the method is capable of a RSD of 0.05 %.

X1.4 *Characterization of WRM*—Units of the WRM are sampled for characterization, which will establish the reference value and its associated uncertainty. The measurement method for characterization is the method used for sample analysis (Davies-Gray).

X1.4.1 *Characterization for Calibration Standard*—For this example, the following measurement scheme is assumed for characterization of the WRM as a calibration standard:

X1.4.1.1 One of the most experienced analysts will make the measurements using the high-precision version of the method.

X1.4.1.2 A solution of a uranium metal CRM (U-CRM) will be prepared and a makeup value calculated. This solution will be titrated to compare one CRM with another CRM (NIST dichromate CRM) as a validation step for the method.

X1.4.1.3 The WRMs will be dissolved one day and titrated the next. For each titration of the WRM solutions and the U-CRM, 100 mg of uranium will be taken.

X1.4.1.4 Initially, the analyst will titrate ten aliquots of the U-CRM. If there is not a significant difference at the 0.05 level of significance between the mean of the results and the makeup value, characterization will continue.

X1.4.1.5 The analyst will select randomly for analysis *n* WRMs from the 500 units using a table of random numbers. Determining the value of *n* will depend on the expected variability of the analytical method (RSD), how close (*e*) the user wants the determined reference value, \bar{x} , of the WRM to be to the true value, μ , and the degree of confidence [(1 – α) 100 %] wanted in the established reference value.

X1.4.1.6 The value of *n* will be calculated based on the following: RSD = 0.05 %, e = 0.03 % relative, and $\alpha = 0.05$. The value of *n* is 11 using the following equation:

$$n \ge \left(\frac{\zeta_{0.025} \cdot \sigma}{e}\right)^2 = \left(\frac{1.96 \cdot 0.0005 \,\mu}{0.0003 \,\mu}\right)^2 = 10.67$$
 (X1.1)

where it is assumed that the results are at least approximately normally distributed (see Note X1.1).

NOTE X1.1— $\zeta_{0.025} = 1.96$ is that value such that (for a normal random variable, X having mean μ and standard deviation σ) P ((1X – μ l) > 1.96 σ) = 2 (0.025) = 0.05.

X1.4.1.7 The analyst will titrate the 11 WRM solutions and 6 U-CRMs in one day. The order of titration will be as follows: U W W U W W U W W U W W U W W U W W U. The U-CRM solution will be titrated with the WRM solutions to serve as a control during their titration, giving additional confidence in characterization.

X1.4.1.8 The resulting data from the analyst will be evaluated statistically, and a reference value for the WRM will be calculated and an uncertainty will be established.

NOTE X1.2—A statistician should be involved with X1.4.1.5-X1.4.1.8. There are a number of variables that must be considered in planning the specific details for each situation. For example, the value of n and the number of titrations made of the WRM solutions and U-CRM may depend on limitations in cost and time. Such limitations could affect the degree of confidence that must be accepted in the reference value.

X1.4.2 Characterization for Control Standard—The following measurement and data treatment scheme will be assumed for the characterization of the WRM as a measurement control standard. In reality, if an established measurement control program exists, the scheme would be dictated by that program.

X1.4.2.1 The routine version of the method will be used to analyze the WRM for preparation of the initial control chart.

X1.4.2.2 Twenty WRMs will be randomly selected from the 500 units using a table of random numbers.

X1.4.2.3 Two WRMs will be analyzed each work day for 10 consecutive work days.

X1.4.2.4 The analysts doing the routine work will make the analyses. The analysts will be assigned as normally assigned to run samples.

X1.4.2.5 Using an analysis of variance technique, the resulting data can be analyzed to confirm that the process mean is constant during this period. The standard deviation of the daily measurements can be compared to the method standard deviation to check for consistency.

X2. PREPARATION AND PACKAGING WRM

X2.1 The following examples of procedures illustrate the preparation and packaging of two WRMs, one starting with plant material and the other starting with metal. The plant material is plutonium nitrate solution and the metal is high purity uranium metal.

X2.2 Preparation from Plant Material⁵—A plant plutonium nitrate solution selected as the starting material for a WRM shall have measured and representative impurity levels and isotopic abundance values. Also, the selected material must be single phase and have no heterogeneously distributed organic matter. This procedure provides treatment designed to destroy plutonium polymers. In this procedure and subsequent treatments to produce the WRM, dilution relative to the plutonium concentration of the plant material occurs. The plant material, therefore, should be selected or should be concentrated by low-temperature evaporation to contain 1.5 times the normal plutonium concentration of the plant stream.

X2.2.1 Reagents:

X2.2.1.1 HF, 29m, and

X2.2.1.2 HNO₃, 15.7M, 8M, and 2M.

X2.2.2 Apparatus:

X2.2.2.1 *Beaker*, appropriate size for the volume of WRM to be prepared, with unribbed watch glass as a cover.

X2.2.2.2 Hot plate.

X2.2.3 Procedure:

X2.2.3.1 Transfer a desired volume of the selected plant plutonium nitrate solution to a beaker having a capacity at least four times the volume attained in X2.2.3.2.

X2.2.3.2 Adjust the HNO₃ concentration to 8*M* by adding an amount of 15.7*M* HNO₃ or 2*M* HNO₃ calculated based on the HNO₃ concentration of the plant plutonium nitrate solution. Do not use water as a diluent because localized concentrations of low acidity in solution may cause hydrolysis of the plutonium and formation of an insoluble hydrous oxide.

X2.2.3.3 Add a calculated volume of 29*M* HF to produce a solution that is 0.05*M* in HF.

X2.2.3.4 Heat the solution at 80 to 90°C for at least 2 h at a moderate rate so that no solution is sprayed onto the beaker walls.

X2.2.3.5 Allow the solution to age for about two weeks. Cover the beaker with a cover glass to decrease evaporation. See Note X2.1.

X2.2.3.6 Filter the aged solution, collecting the filtrate in a tared volumetric flask. Rinse the filter with 8M HNO₃. See Note X2.2.

X2.2.3.7 Dilute to volume to give a final acid concentration of about 8*M*, using 15.7M HNO₃ to increase acidity or not less than 2*M* to decrease acidity.

X2.2.3.8 Stopper the flask and weigh.

X2.2.3.9 Mix the solution by inverting the flask at least 10

times or mix using a TFE-fluorocarbon coated magnetic stir bar for at least 5 min.

X2.2.3.10 Package the WRM. See X2.3.

NOTE X2.1—All plutonium nitrate WRMs prepared for plutonium concentration measurements, except those prepared by dissolving plutonium metal in HCl, are aged to provide time for radiolytically generated hydrogen peroxide to reduce any Pu(VI) to Pu(IV). Two weeks ensure that >99 % of the plutonium is Pu(IV). Because the reduction of Pu(VI) by hydrogen peroxide produces gas, the solution is open to the atmosphere during the aging period. The aging period also provides a reasonable time for precipitate formation, an undesirable effect often occurring in plutonium nitrate solutions.

NOTE X2.2—Every plutonium nitrate WRM intended for use in plutonium concentration measurements is filtered, even when no precipitate is apparent. The recommended filter is a plastic membrane-type, resistant to 8M HNO₃, with a 0.45-µm pore size. The filter, with residue, is analyzed for its plutonium content, and the assay value of the WRM either is corrected for the insoluble plutonium or the WRM is characterized for its plutonium concentration using two different assay methods.

X2.3 Packaging Solution in Glass Ampoules⁴—The containers are flame-sealed, thick-walled glass ampoules designed for 10 mL or less of solution. Normally, the packaging is for WRMs used for plutonium concentration. If the packaging is to be for total plutonium content per ampoule, the ampoules must be weighed (tared) before transfer of the WRM and weighed after transfer (see X2.3.1.3).

X2.3.1 Filling Ampoules:

X2.3.1.1 Clean, dry, and label the ampoules.

X2.3.1.2 Insert the buret (see Fig. 4) into an ampoule, being careful not to let the tip of the buret touch the inside wall of the ampoule.

X2.3.1.3 Transfer the desired amount of WRM solution to the ampoule, then temporarily seal the ampoule with a rubber stopper.

X2.3.1.4 Repeat X2.3.1.2 and X2.3.1.3 until all ampoules are filled.

X2.3.2 Sealing Ampoules:

X2.3.2.1 Within 1 h of filling, flame seal each ampoule's neck about 70 mm above the body, using an oxygen-gas flame. Complete the seal in less than 1 min to avoid warming the solution.

X2.3.2.2 Anneal in an air-gas flame and allow to cool. See Note X2.3.

X2.3.2.3 Test the integrity of the seals by inverting each ampoule, smearing the seal with a filter paper, and alpha counting the paper. Reject any leaky ampoule.

NOTE X2.3—The sealing and annealing should be done by an experienced glassblower.

X2.4 *Preparation from Uranium Metal*⁶—Clean high purity uranium metal of any surface oxide and weigh. Dissolve

⁵ Based on NUREG-0118 (also designated LA-NUREG-6348), Preparation of Working Calibration and Test Materials: Plutonium Nitrate Solution, Nuclear Regulatory Commission. Available from the Superintendent of Documents, U.S. Government Printing Office, Washington, DC 20402.

⁶ Based on NBL-CAL-U(E)-2, Preparation of Uranium Standard Solution from Uranium Metal, New Brunswick Laboratory, United States Department of Energy. Available from the Superintendent of Documents, U.S. Government Printing Office, Washington, DC 20402.

the metal and dilute the resulting solution to volume to give the desired uranium concentration. Measure the mass of the solution and calculate the uranium concentration (g U/g soln). Dispense weighed aliquants of the solution into individual containers for use as measurement standards.

X2.4.1 Reagents:

X2.4.1.1 *HNO*₃, 8*M*, and

X2.4.1.2 Uranium Metal, 99.9 %.

X2.4.2 Apparatus:

X2.4.2.1 *Glass Bottles*, 500 mL, equipped with polycone seal caps.

X2.4.2.2 Glass Flask, 5 L.

X2.4.2.3 Polyethylene Dispensing Bottles, 60 mL and 125 mL.

X2.4.2.4 Titration Beakers.

X2.4.2.5 Petri Dish, for each beaker.

X2.4.3 Preparation and Storage of Master Solution:

X2.4.3.1 Clean about 65 g of metal in 8M HNO₃. Rinse thoroughly with distilled water and then with acetone. Dry with a lint-free towel and weigh to the nearest 0.01 mg. Calculate the actual mass of uranium by applying the proper bouyancy and purity corrections. See Note X2.4.

X2.4.3.2 Transfer the metal carefully to a preweighed 5-L glass flask. Add 150 mL of 8M HNO₃ and warm the flask over low heat. Replenish the acid as needed until dissolution is completed. Record the total amount of acid added.

X2.4.3.3 Cool the solution to ambient temperature and bring to a total volume of 4 L by adding distilled water in 300 to 500-mL portions and swirling after each addition. See Note X2.5N.

X2.4.3.4 Transfer the solution to 500-mL glass bottles,

filling only to the shoulders of the bottles. Wipe the mouth of each bottle before adding and tightening the cap. Weigh each bottle to the nearest 0.01 g and seal with plastic tape. The master solution is stored in these bottles.

X2.4.3.5 Check the weight whenever a bottle is opened for the first time. Do not use the solution in a bottle if a discrepancy in the weights of greater than 0.1 g is observed. See Note X2.6.

Note X2.4—The uranium metal buoyancy correction (assuming a uranium metal density of 18.7) for balance weights with an apparent density of 8.0 g per cm is -0.086 mg per g (at sea level) or -0.082 (at 740 mm Hg, 25°C, and 30 % relative humidity).

Note X2.5—The final uranium concentration should be about 15 mg U per g solution.

NOTE X2.6—Data accumulated from several master solutions have shown that, with proper care in handling, these solutions can be stored for periods in excess of one year.

X2.4.4 Preparation of Measurement Standards:

X2.4.4.1 Transfer about 125 g of master solution to a 125 mL polyethylene dispensing bottle or 60 g to a 60 mL bottle. Weigh the bottle to the nearest 0.01 mg.

X2.4.4.2 Dispense a selected amount of the master solution into an appropriately identified titration beaker. Reweigh the bottle and record the weight.

X2.4.4.3 Continue X2.4.4.2 until the dispensing bottle is empty.

X2.4.4.4 Rinse the walls of each beaker with distilled water and then evaporate the solution in each beaker to dryness on a steam bath. Cool and cover each beaker with a petri dish.

X2.4.4.5 Store the beakers until a measurement standard is needed. Then, dissolve the dried contents of a beaker.

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