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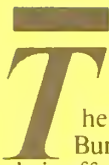
U.S. DEPARTMENT OF COMMERCE/National Bureau of Standards

Standard Reference Materials:

**Summary of the
Environmental Research,
Analysis, and Control
Standards Issued by the
National Bureau of
Standards**

R. Mavrodineanu and S. D. Rasberry

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Provides the national system of physical and chemical measurement; coordinates the system with measurement systems of other nations and furnishes essential services leading to accurate and uniform physical and chemical measurement throughout the Nation's scientific community, industry, and commerce; provides advisory and research services to other Government agencies; conducts physical and chemical research; develops, produces, and distributes Standard Reference Materials; and provides calibration services. The Laboratory consists of the following centers:

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Standard Reference Materials:

Summary of the Environmental Research, Analysis, and Control Standards Issued by the National Bureau of Standards

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Preface

Standard Reference Materials (SRM's) as defined by the National Bureau of Standards (NBS) are well-characterized materials, produced in quantity and certified for one or more physical or chemical properties. They are used to assure the accuracy and compatibility of measurements throughout the Nation. SRM's are widely used as primary standards in many diverse fields in science, industry, and technology, both within the United States and throughout the world. They are also used extensively in the fields of environmental and clinical analysis. In many applications, traceability of quality control and measurement processes to the national measurement system is carried out through the mechanism and use of SRM's. For many of the Nation's scientists and technologists it is therefore of more than passing interest to know the details of the measurements made at NBS in arriving at the certified values of the SRM's produced. An NBS series of papers, of which this publication is a member, called the NBS Special Publication - 260 Series, is reserved for this purpose.

The 260 Series is dedicated to the dissemination of information on different phases of the preparation, measurement, certification and use of NBS SRM's. In general, much more detail will be found in these papers than is generally allowed, or desirable, in scientific journal articles. This enables the user to assess the validity and accuracy of the measurement processes employed, to judge the statistical analysis, and to learn details of techniques and methods utilized for work entailing the greatest care and accuracy. These papers also should provide sufficient additional information not found on the certificate so that new applications in diverse fields not foreseen at the time the SRM was originally issued will be sought and found.

Inquiries concerning the technical content of this paper should be directed to the author(s). Other questions concerned with the availability, delivery, price, and so forth, will receive prompt attention from:

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Gaithersburg, MD 20899

Stanley D. Rasberry, Chief
Office of Standard Reference Materials

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** May be ordered from: National Technical Information Services (NTIS). Springfield Virginia 22161.

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Abstract

This publication is a summary of the environmental research, analysis, and control standards issued by NBS as Standard Reference Materials (SRM's). The material, composition, certification, use, and remarks concerning each of the SRM's described are presented in tabular form. Copies of the certificates of these SRM's are contained in the appendix for more detailed information.

Key Words: chemical composition; environmental standards;
Standard Reference Materials.

Introduction

Since its inauguration in 1901, the National Bureau of Standards (NBS) has issued nearly 2000 different Standard Reference Materials (SRM's). Many of these have been renewed several times, many have been replaced or discontinued as technology changed. Today, over 900 SRM's are available, together with a large number of scientific publications related to the fundamental and applied characteristics of these materials. Each material is certified for chemical composition, chemical properties, or its physical or mechanical characteristics. Each SRM is provided with a Certificate or Certificate of Analysis that contains the essential data concerning its properties or characteristics. The SRM's currently available cover a wide range of chemical, physical, and mechanical properties, and a corresponding wide range of measurement interests in practically all aspects of fundamental and applied science. These SRM's constitute a unique and invaluable means of transferring to the user accurate data obtained at NBS, and provide essential tools that can be used to improve accuracy in practically all areas where measurements are performed.

In addition to SRM's, the National Bureau of Standards issues a variety of Research Materials (RM's) having various properties described in individual "Reports of Investigation." They are intended primarily to further the scientific or technical research on that particular material. Other materials, called Special Reference Materials (GM's), are also available from NBS. These are materials produced and certified by other Government agencies, standard organizations, or other nonprofit organizations, that are considered useful to the public and for which no alternate method of national distribution exists.

The various categories of materials available from NBS are given in table 1. This table lists these materials according to their chemical composition, physical properties, or engineering characteristics. A more detailed alphabetic enumeration of these materials is given in appendix I. Table 1 and appendix I were taken from NBS Special Publication 260, NBS Standard Reference Materials Catalog, 1984-85 Edition¹. This publication lists every material available from the NBS Office of Standard Reference Materials.

Further information on the reference materials available from NBS may be obtained from the Office of Standard Reference Materials, National Bureau of Standards, Gaithersburg, MD 20899. Information on other NBS services may be obtained from the Technical Information and Publications Division, National Bureau of Standards, Gaithersburg, MD 20899.

In addition to reference materials, NBS provides many additional services. These include: Measurement Assurance Programs, Calibration and Related Measurement Services, Proficiency Sample Programs, a National Voluntary Laboratory Accreditation Program, Standards Information Services, Standard Reference Data, and Technical Information and Publications.

¹For sale by the Superintendent of Documents, U.S. Government Printing Office, Washington, DC 20402, under Stock No. 003-003-02558-5 (Price \$5.50, add 25 percent for foreign orders.)

Table 1. Categories of Standard Reference Materials available from the National Bureau of Standards.

CERTIFIED CHEMICAL COMPOSITION STANDARDS

| | |
|--|--|
| Steels (chip form) | Gases in Metals |
| Plain carbon | High-Purity Metals |
| Low alloy | Electron Probe Microanalytical Standards |
| High alloy | Primary, Working, and Secondary |
| Stainless | Standard Chemicals |
| Tool | Microchemical Standards |
| Steels (granular form) | Clinical Laboratory Standards |
| Steels (solid form) | Biological Standards |
| Ingot iron and low alloy | Environmental Standards |
| Special ingot irons and low alloy | Analyzed gases |
| Stainless | Analyzed liquids and solids |
| Specialty | Permeation tubes |
| High-temperature alloys | Industrial Hygiene Standards |
| Tool | Spectrochemical Standards |
| Steelmaking Alloys | Hydrocarbon Blends |
| Cast Irons (chip form) | Metallo-Organic Compounds |
| Cast Steels, White Cast Irons, Ductile | Fertilizers |
| Irons, and Blast Furnace Irons | Ores |
| (solid form) | Minerals, Refractories, Glasses, and |
| Nonferrous Alloys (chip form) | Carbides |
| Aluminum "Benchmarks" | Cement |
| Cobalt | Trace Element Standards |
| Copper | Nuclear Materials |
| Copper "Benchmarks" | Special nuclear materials |
| Lead | Plutonium assay |
| Magnesium | Plutonium isotopic |
| Nickel | Uranium assay |
| Nickel Superalloy, Trace Elements | Uranium isotopic |
| Nickel oxide | Neutron density standards |
| Selenium | Fission track glass standards |
| Tin | Isotopic Reference Standards |
| Titanium | |
| Zinc | |
| Zirconium | |
| Nonferrous Alloys (solid form) | |
| Aluminum "Benchmarks" | |
| Copper | |
| Copper "Benchmarks" | |
| Lead | |
| Nickel | |
| Titanium | |
| Zinc | |
| Zirconium | |

CERTIFIED PHYSICAL PROPERTY STANDARDS

Ion Activity Standards

pH standards
pD standards
Ion selective electrodes

Optical Standards

Spectrophotometric
Thermal emittance
Refractive index

Mechanical and Metrology Standards

Magnification
Coating thickness
Glass
Elasticity
Density
Polymer
Rheology

Radioactivity Standards

Alpha-particle standards
Beta-particle and gamma-ray gas standards
Alpha-particle, beta-particle, gamma-ray, and electron-capture solution standards
Contemporary standard for carbon-14 dating laboratories
Environmental standards
Low energy photon sources
Gamma-ray "point-source" standards
Radium gamma-ray solution standards
Radium solution standards for random analysis
Radioactivity standard reference materials currently not in stock

Heat Standards

Superconductive thermometric fixed point devices
Freezing Points

Defining fixed points
Determined reference points

Melting points
Calorimetric

Combustion
Solution
Heat source
Enthalpy and heat capacity

Vapor pressure
Thermal expansion
Thermocouple materials
Thermal resistance

Metallurgical

Mossbauer

X-ray Diffraction

Gas Transmission

Permittivity

Reference Fuels

Resistivity

Magnetic Standards

Magnetic susceptibility
Magnetic moment
Paramagnetic resonance

ENGINEERING TYPE STANDARDS

Standard Rubber and Rubber-Compounding Materials

Reference Magnetic Tapes

Lubricant Standards

Sizing Standards

Glass spheres for particle size
Turbidimetric and fineness (cement)

X-ray and Photographic Standards

Surface Flammability Standards

Semiconductor Production Standards

Water Vapor Permeance

Tape Adhesion Testing Standards

Color Standards

RESEARCH MATERIALS

SPECIAL REFERENCE MATERIALS

Environmental Research, Analysis, and Control Standards

This work is the fourth in a series of NBS Special Publications dedicated to the description of the Standard Reference Materials (SRM's) issued by the National Bureau of Standards (NBS).

The first volume, NBS SP 260-71, "Summary of the Clinical Laboratory Standards Issued by the National Bureau of Standards" (176 pp., November 1981), describes in a tabular form 41 SRM's available in that field and includes copies of the corresponding Certificates of Analysis for further information.

The second volume, NBS SP 260-97, "Summary of the Coal, Ore, Mineral, Rock, and Refractory Standards Issued by the National Bureau of Standards" (134 pp., September 1985), describes in the same manner 39 SRM's available in that field.

The third volume, NBS SP 260-104, "Summary of the Biological and Botanical Standards Issued by the National Bureau of Standards" (68 pp., October 1985), presents in a similar manner the essential data concerning 9 SRM's issued in that field.

The present volume, NBS SP 260-105, "Summary of the Environmental Research, Analysis, and Control Standards Issued by the National Bureau of Standards" (97 pp., March 1986), describes in a tabular form the characteristic properties of 22 SRM's issued in the field of environmental analytical instrumentation and methodology. Copies of the Certificates of Analysis are reproduced in this work also, for further information.

Tables 2 and 3 contain the essential information concerning the material composition, the certification parameters, and use. Under "Remarks," additional data such as storage conditions and stability is provided.¹ All the data and information contained in these tables were extracted from the Certificates or Certificates of Analysis issued for the SRM's included in the table. An examination of these tables gives the reader a general view of these SRM's. For more detailed information, the individual Certificates reproduced in appendix II should be consulted as well as the references cited in each Certificate. The SRM's listed in the tables include all of the environmental standards that were issued or were in preparation by the end of 1984. These SRM's are the result of the concerted efforts of a number of scientists from the NBS National Measurement Laboratory. Each Certificate lists the individuals who contributed to development of the SRM.

Appendix III provides a guide to the reader to assist in requesting NBS to develop new SRM's. A final appendix, appendix IV, is a guide to ordering SRM's.

In addition to the SRM's and their Certificates, NBS issues a series of Special Publications, called the "260 Series," that relate directly to Standard Reference Materials as stated in the preface. The list of available publications in the "260 Series" is given at the beginning of this publication.²

¹NOTE: The use of proprietary designations in table 2 is for information only, and should not be construed as an endorsement of the product by either the Department of Commerce or the National Bureau of Standards.

²For complete bibliographic reference and ordering information, see "Other NBS Publications in This Series," pp. iv.

**TABLE 2. SUMMARY OF THE ENVIRONMENTAL
RESEARCH, ANALYSIS, AND
CONTROL STANDARDS.**

| | | |
|---|---|--|
| 1579 Powdered Lead Based Paint | Collected by the Philadelphia Dept. Public Health, and sieved to a powder passing through a 325 mesh sieve. | Pb: 11.87 ± 0.04 % by wt., based on samples >0.1 g of as-received material. |
| 1580 Organics in Shale Oil | The shale oil came from the Laramie Energy Technology Center, Laramie, Wyoming, and was collected from the Mahogany Zone of the Colorado Green River Formation. | Fluoranthene: 54; pyrene: 104; benzo[a]pyrene: 21; benzo[e]pyrene: 18; perylene: 3.4; phenol: 407; o-cresol: 385; 2,6-dimethylphenol: 175; benzol[f]quinoline: 16, $\mu\text{g/g}$. |
| 1620a Sulfur in Residual Fuel Oil | The material is a commercial "No. 5 Heavy" residual fuel oil as defined by the American Society for Testing and Materials. | S: 4.504 ± 0.010 wt. %. |
| 1621b Sulfur in Residual Fuel Oil | The material is a commercial "No. 6" residual fuel oil as defined by the American Society for Testing and Materials. | S: 0.950 ± 0.005 wt. %. |
| 1622b Sulfur in Residual Fuel Oil | Same as for SRM 1621b. | S: 1.982 ± 0.018 wt. %. |
| 1623a Sulfur in Residual Fuel Oil | Same as for SRM 1620a. | S: 0.240 ± 0.003 wt. %. |
| 1624a Sulfur in Distillate (Diesel) Fuel Oil | The material is a commercial "No. 2-D" distillate fuel oil as defined by the American Society for Testing and Materials. | S: 0.141 ± 0.002 wt. %. |
| 1630 Trace Mercury in Coal | Commercial coal crushed to 210-500 micrometer particle size. | Hg: $0.13 \mu\text{g/g} \pm 1$ in the last significant figure. |

| Certification | Use | Remarks |
|---|--|---|
| By x-ray fluorescence, atomic absorption spectrometry, polarography. | In the calibration of apparatus and methods for determining Pb in paint removed from the interior surfaces of old housing. | |
| By gas chromatography, gas chromatography/mass spectrometry, and high performance liquid chromatography. | For evaluating the reliability of analytical methods used for trace organic compounds in oil materials. | Seven additional organic compounds are included for information only and are not certified. |
| By gravimetry, ion chromatography, and x-ray fluorescence. | As analytical standard in the determination of total S in fuel oils and similar materials. | Four additional physical properties are indicated but are not certified; similarly, 16 trace elements are mentioned semi-quantitatively only. |
| As for SRM 1620a. Is valid for 3 years from date of purchase. | As for SRM 1620a. | |
| As for SRM 1621b. | As for SRM 1621b. | |
| As for SRM 1620a. | As for SRM 1620a. | |
| By gravimetry and ion chromatography. Is valid for 3 years from date of purchase. | As for SRM 1620a. | |
| By neutron activation and flameless atomic absorption spectrometry (0.14 µg/g). The neutron activation procedure is described in the certificate. | As analytical standard for the determination of trace mercury in coal. | Selenium is also given for information, but is not certified (2.1 µg/g). |

Table 2. Continued.

| SRM | Material | Composition |
|---|--|---|
| 1632b Trace Elements in Coal (Bituminous) | Obtained from an underground mine that recovers coal from the Pittsburgh seam, crushed and sieved through a -60 mesh at the Humphrey No. 7 mine and coal preparation plant of the Consolidation Coal Co., Christopher Coal Co. Div., Osage, W. Va. | Material should be vacuum dried at ambient temperature for 24 hours prior to use. Values based on a minimum sample size of 250 mg. C (total) 78.11; H 5.07; N 1.56; S 1.89; Volatile matter 35.4; Al 0.855; Ca .204; Fe .759; Mg .0383; K .0748; Na .0515; Ti .0454; values % by wt. As 3.72; Ba 67.5; Cd 0.0573; Co 2.29; Cu 6.28; Pb 3.67; Mn 12.4; Ni 6.10; Rb 5.05; Se 1.29; Th 1.342; U 0.436; Zn 11.89; values $\mu\text{g/g}$. Not certified: 17 additional constituents (see certificate). |
| 1633a Trace Elements in Coal Fly Ash | Obtained from a coal fired power plant using Pennsylvania and West Virginia Coal. The ash was sieved through a No. 170 sieve. | Determined on 250 mg or more sample dried to constant weight. Al 14.3; Ca 1.11; Fe 9.40; K 1.88; Mg 0.455; Na 0.17; Si 22.8; wt. %. Sb 6.8; As 145; Cd 1.0; Cr 196; Cu 118; Hg 0.16; Ni 127; Pb 72.4; Rb 131; Se 10.3; Sr 830; Th 24.7; Tl 5.7; U 10.2; V 2.97; Zn 220; $\mu\text{g/g}$. Additional 15 elements determined but not certified. |
| 1634a Trace Elements in Fuel Oil | Commercial No. 6 residual fuel oil as defined by ASTM. | Determined on at least 1 g sample. Pb 2.80; Mn 0.19; Ni 29; Se 0.15; Na 87; V 56; Zn 2.7; $\mu\text{g/g}$. S 2.85 wt. %. Values for additional 11 elements and 4 physical properties are given but not certified. |
| 1635 Trace Elements in Coal (Subbituminous) | Subbituminous coal from Eagle Mine of the Imperial Coal Co. of Erie, Colorado; sieved through a No. 65 sieve. | Determined on at least 250 mg of dried sample. As 0.42; Cd 0.03; Cr 2.5; Cu 3.6; Pb 1.9; Mn 21.4; Ni 1.74; Se 0.9; Th 0.62; U 0.24; V 5.2; Zn 4.7; $\mu\text{g/g}$; and Fe 0.239; S 0.33 wt %. Additional 10 elements determined but not certified. |
| 1636a, 1637a, 1638a Lead in Reference Fuel | Supplied by Phillips Petroleum Co., Bartlesville, Okla. Lead was added as tetraethyl lead motor mix. | Determined on at least 1 g sample. Vial I-11.2; Vial II-18.8; Vial III-25.1; Vial IV-764 $\mu\text{g/g}$. |

| Certification | Use | Remarks |
|--|--|--|
| Analyses performed in the NBS Center for Analytical Chemistry. Estimated uncertainty 0.006 depending on constituent. | For the calibration of apparatus and the evaluation of techniques employed in the analysis of coal or similar materials. | Should be kept in its original bottle. Should not be exposed to intense sources of radiation, including ultraviolet lamps or sunlight. |
| Using two to four different analytical procedures for each element as shown in the Certificate, on the dried sample. | For calibration of apparatus and methods used in the analysis of coal ash and similar materials. | The materials should be dried as indicated in the Certificate. Stability (>3 years) not yet established. |
| Using two to three independent analytical procedures for each element. | For calibration of apparatus and methods used in the analysis of fuel oils and similar materials for trace elements. | Store in tightly sealed bottle. Certificate valid for 3 years from date of purchase. |
| Using two to three independent analytical procedures for each element. For sample drying, see the Certificate. | For calibration of apparatus and techniques used in trace element analysis of coal and similar materials. | Store in tightly sealed bottle in a cool, dark place. Long term (>1 year) stability not yet established. |
| By isotope dilution mass spectrometry. SRM 1636a is made from Vials I, II, III, and IV. SRM 1637a is made from Vials I, II, and III. SRM 1638a is made from Vial IV. | For calibration of instruments and techniques used for the analysis of Pb in gasoline. | The vials should be stored at 10-30 °C in darkness, and not reused after first opening. |

Table 2. - Continued.

| SRM | Material | Composition |
|---|---|--|
| 1641b Mercury in Water - $\mu\text{g/mL}$ | This SRM was prepared at NBS and is delivered in 6 ampoules of 20 mL each. | Hg: $1.52 \pm 0.04 \mu\text{mL}$. |
| 1642b Mercury in Water - ng/mL | Same as SRM 1641b. | Hg: $1.49 \pm 0.06 \text{ng/mL}$. |
| 1643b Trace Elements in Water | Prepared at the U.S. Geological Survey, National Water Quality Laboratory, Arvada, Colo., by using high-purity reagents and sterilization. | Ba 44; Be 19; Cd 20; Cr 18.6; Co 26; Cu 21.9; Fe 99; Pb 23.7; Mn 28; Mo 85; Ni 49; Se 9.7; Ag 9.8; Sr 227; Tl 8; V 45.2; Zn 66 ng/g . Additional 3 elements determined but not certified. |
| 1644 Generator Columns for Polynuclear Aromatic Hydrocarbons | Three 50 cm x 0.6 cm coiled stainless steel tubes each packed with fine quintus quartz coated with 0.5 wt. % of polynuclear aromatic hydrocarbon (PAH). | At 20 °C: anthracene 30.7; benzo(a)anthracene 6.45; benzo(a)pyrene 1.13, all in $\mu\text{g/Kg}$. |
| 1645 River Sediment | Deposit dredged from the bottom of Indiana Harbor Canal, Gary, Ind., freeze-dried, sieved (No. 80 screen) and radiation-sterilized. | Al 2.26; Cr 2.96; Fe 11.3; K 1.26; Mg 0.74; Na 0.54; Zn 0.172, wt %. Cd 10.2; Cu 109; Co 10.1; Pb 714; Mn 785; Hg 1.1; Ni 45.8; Tl 1.44; Th 1.62; U 1.11; V 23.5; $\mu\text{g/g}$. Additional 13 values determined but not certified. |
| 1646 Estuarine Sediment | Dredged from the Chesapeake Bay, freeze-dried, radiation-sterilized and sieved (No. 100 sieve). | Al 6.25; Ca 0.83; Fe 3.35; Mg 1.09; P 0.054, wt. %. As 11.6; Cd 0.36; Cr 76; Co 10.5; Cu 18; Pb 28.2; Mn 375; Hg 0.063; Ni 32; V 94; Zn 138, $\mu\text{g/g}$. |
| 1648 Urban Particulate Matter | Urban particulate matter was collected in St. Louis, Mo., over a period of 12 months. | Al 3.42; Fe 3.91; K 1.05; Pb 0.655; Na 0.425; Zn 0.476, wt. %. As 115; Cd 75; Cr 403; Cu 609; Ni 82; Se 27; U 5.5; V 140 $\mu\text{g/g}$. Non-certified values are given for 26 elements. |

| | | |
|---|---|---|
| By atomic absorption spectrometry and neutron activation. Use of blank samples is necessary. | For primary calibration of instruments and techniques and as a spike sample in method-of-addition procedures for the determination of Hg in natural waters. | Stability limited to 1 year from date of purchase. |
| Same as SRM 1641b. | Same as SRM 1641b. | Should be used without dilution. For precautions in use see Certificate. |
| At least two from the nine analytical procedures employed were used for the determination of each element. | For evaluating the accuracy of trace element determination in fresh water and for instrument calibration. | The certification is valid for two years from the shipping date. For precautions in use see Certificate. |
| Performed by two independent analytical procedures. Concentrations for the three PAH at other temperatures (10-30 °C) are given in the Certificate. | This SRM is intended to provide accurate concentrations of the three PAH. | For further properties, use, and stability, see Certificate. |
| Six independent analytical procedures and 100 mg to 1 g of sample were used for certification. | For calibration of apparatus and methods used in the analysis of river sediments or similar materials. | For details on stability, use, and homogeneity, see Certificate. Data valid for 5 years from date of purchase. |
| By 6 independent analytical procedures on 500 mg or more dried sample, using two to four different procedures for each element. | For calibration of instrumentation and evaluation of analytical methods in sediments and similar matrices. | Certified data valid for 5 years after date of shipping. |
| On 100 mg or more of dried sample using 9 independent analytical procedures. | Calibration of apparatus and evaluation of methods used in the analysis of atmospheric particulate matter and similar matrices. | Certification valid for 5 years from date of purchase, for samples kept in the original bottle at 10-30 °C in a desiccator and in dark. |

Table 3. Comparative List of Chemical Elements for the SRM's Described in Table 2.

Analyzed Liquids and Solids

MULTI-ELEMENT

Concentrations:

Values expressed as microgram per gram, except:

Weight Percent -- %

Nanogram per gram -- *italics*

Parenthesis indicates elements not certified and given for information only.

| SRM | Type | Unit Size | Al | Sb | As | Ba |
|-------|---|-----------|---------|--------|--------|---------|
| 1632b | Trace Elements in Coal (Bituminous) | 75 g | 0.855% | (0.24) | 3.72 | 67.5 |
| 1633a | Trace Elements in Coal Fly Ash | 75 g | 14.3% | 6.8 | 145 | (0.15%) |
| 1634a | Trace Elements in Fuel Oil | 100 mL | -- | -- | (0.12) | -- |
| 1635 | Trace Elements in Coal (Subbituminous) | 75 g | (0.32%) | (0.14) | 0.42 | -- |
| 1643b | Trace Elements in Water (ng/g) | 950 mL | -- | -- | (49) | 44 |
| 1645 | River Sediment | 70 g | 2.26% | (51) | (66) | -- |
| 1646 | Estuarine Sediment | 75 g | 6.25% | (0.4) | 11.6 | -- |
| 1648 | Urban Particulate | 2 g | 3.42% | (45) | 115 | (737) |

| SRM | Be | Bi | B | Br | Cd | Ca | C | Ce |
|-------|-----------|-------------|-------------|--------|-----------|--------|--------|-------|
| 1632b | -- | -- | -- | (0.17) | 0.0573 | 0.204% | 78.11% | (9) |
| 1633a | (12) | -- | -- | -- | 1.00 | 1.11% | -- | (180) |
| 1634a | (0.006) | -- | -- | (<1) | (0.002) | (16) | -- | -- |
| 1635 | -- | -- | -- | -- | 0.03 | -- | -- | (3.6) |
| 1643b | <i>19</i> | <i>(11)</i> | <i>(94)</i> | -- | <i>20</i> | -- | -- | -- |
| 1645 | -- | -- | -- | -- | 10.2 | (2.9%) | -- | -- |
| 1646 | (1.5) | -- | -- | -- | 0.36 | 0.83% | -- | (80) |
| 1648 | -- | -- | -- | (500) | 75 | -- | -- | (55) |

| SRM | Cs | Cl | Cr | Co | Cu | Eu | F |
|-------|--------|---------|-------------|-----------|-------------|--------|---------|
| 1632b | (0.44) | (1260) | (11) | 2.29 | 6.28 | (0.17) | -- |
| 1633a | (11) | -- | 196 | (46) | 118 | (4) | -- |
| 1634a | -- | (31) | (0.7) | (0.3) | -- | -- | -- |
| 1635 | -- | -- | 2.5 | (0.65) | 3.6 | (0.06) | -- |
| 1643b | -- | -- | <i>18.6</i> | <i>26</i> | <i>21.9</i> | -- | -- |
| 1645 | -- | -- | 2.96% | 10.1 | 109 | -- | (0.09%) |
| 1646 | (3.7) | -- | 76 | 10.5 | 18 | (1.5) | -- |
| 1648 | (3) | (0.45%) | 403 | (18) | 609 | (0.8) | -- |

| SRM | Ga | Ge | Hf | H | In | I | Fe | La |
|-------|--------|-------|--------|-------|-------|------|-----------|-------|
| 1632b | -- | -- | (0.43) | 5.07% | -- | -- | 0.759% | (5.1) |
| 1633a | (58) | -- | (8) | -- | -- | -- | 9.4% | -- |
| 1634a | -- | -- | -- | -- | -- | -- | (31) | -- |
| 1635 | (1.05) | -- | (0.29) | -- | -- | -- | 0.239% | -- |
| 1643b | -- | -- | -- | -- | -- | -- | <i>99</i> | -- |
| 1645 | -- | -- | -- | -- | -- | -- | 11.3% | (9) |
| 1646 | -- | (1.4) | -- | -- | -- | -- | 3.35% | -- |
| 1648 | -- | -- | (4.4) | -- | (1.0) | (20) | 3.91% | (42) |

Table 3. Continued.

| SRM | Pb | Li | Mg | Mn | Hg | Mo | Ni | N |
|-------|--------|------|---------|-------|----------|--------|------|------|
| 1632b | 3.67 | (10) | 0.0383% | 12.4 | -- | (0.9) | 6.10 | 1.56 |
| 1633a | 72.4 | -- | 0.455% | 179 | 0.16 | (29) | 127 | -- |
| 1634a | 2.80 | -- | -- | 0.19 | (<0.002) | (0.12) | 29 | -- |
| 1635 | 1.9 | -- | -- | 21.4 | -- | -- | 1.74 | -- |
| 1643b | 23.7 | -- | -- | 28 | -- | 85 | 49 | -- |
| 1645 | 714 | -- | 0.74% | 785 | 1.1 | -- | 45.8 | -- |
| 1646 | 28.2 | (49) | 1.09% | 375 | 0.063 | (2.0) | 32 | -- |
| 1648 | 0.655% | -- | (0.8%) | (860) | -- | -- | 82 | -- |

| SRM | P | K | Rb | Sm | Sc | Se | Si | Ag |
|-------|--------|---------|------|--------|--------|-------|--------|-----|
| 1632b | -- | 0.0748% | 5.05 | (0.87) | (1.9) | 1.29 | (1.4%) | -- |
| 1633a | -- | 1.88% | 131 | -- | (40) | 10.3 | 22.8% | -- |
| 1634a | -- | -- | -- | -- | -- | 0.15 | -- | -- |
| 1635 | -- | -- | -- | -- | (0.63) | 0.9 | -- | -- |
| 1643b | -- | -- | -- | -- | -- | 9.7 | -- | -- |
| 1645 | -- | 1.26% | -- | -- | (2) | (1.5) | -- | -- |
| 1646 | 0.054% | (1.4%) | (87) | -- | (10.8) | (0.6) | (31%) | -- |
| 1648 | -- | 1.05% | (52) | (4.4) | (7) | 27 | -- | (6) |

| SRM | Na | Sr | S | Te | Tl | Th | Ti | W |
|-------|---------|-------|---------|-------|-------|-------|---------|--------|
| 1632b | 0.0515% | (102) | 1.89% | -- | -- | 1.342 | 0.0454% | (0.48) |
| 1633a | 0.17% | 830 | (0.18%) | -- | 5.7 | 24.7 | (0.8%) | -- |
| 1634a | 87 | -- | 2.85% | -- | -- | -- | -- | -- |
| 1635 | (0.24%) | -- | 0.33% | -- | -- | 0.62 | (0.02%) | -- |
| 1643b | -- | 227 | -- | -- | 8.0 | -- | -- | -- |
| 1645 | 0.54% | -- | (1.1%) | -- | 1.44 | 1.62 | -- | -- |
| 1646 | (2.0%) | -- | (0.96%) | (0.5) | (0.5) | (10) | (0.51%) | -- |
| 1648 | 0.425% | -- | (5.0%) | -- | -- | (7.4) | (0.40%) | (4.8) |

| SRM | U | V | Zn |
|-------|-------|------|--------|
| 1632b | 0.436 | (14) | 11.89 |
| 1633a | 10.2 | 297 | 220 |
| 1634a | -- | 56 | 2.7 |
| 1635 | 0.24 | 5.2 | 4.7 |
| 1643b | -- | 45.2 | 66 |
| 1645 | 1.11 | 23.5 | 0.172% |
| 1646 | -- | 94 | 138 |
| 1648 | 5.5 | 140 | 0.476% |

Analyzed Liquids and Solids

SINGLE ELEMENT

| SRM | Type | Unit Size | Lead | Sulfur | Mercury |
|-------|-----------------------------|-----------|--------|--------|------------|
| 1579 | Powdered Lead Base Paint | 35 g | 11.87% | -- | -- |
| 1620a | Sulfur in Residual Fuel Oil | 100 mL | -- | 4.504% | -- |
| 1621b | Sulfur in Residual Fuel Oil | 100 mL | -- | 0.950% | -- |
| 1622b | Sulfur in Residual Fuel Oil | 100 mL | -- | 1.982% | -- |
| 1623a | Sulfur in Residual Fuel Oil | 100 mL | -- | 0.240% | -- |
| 1624a | Sulfur in Residual Fuel Oil | 100 mL | -- | 0.141% | -- |
| 1630 | Trace Mercury in Coal | 50 mg | -- | -- | 0.13 µg/g |
| 1641b | Mercury in Water - µg/mL | 120 mL | -- | -- | 1.52 µg/mL |
| 1642b | Mercury in Water - ng/mL | 950 mL | -- | -- | 1.49 ng/mL |

| SRM | Type | Element Certified | Nominal Concentration | No. Units |
|-------|------------------------|-------------------|---------------------------------|--------------|
| 1636a | Lead in Reference Fuel | Pb | 0.03, 0.05, 0.07, and 2.0 g/gal | 3 vials each |
| 1637a | Lead in Reference Fuel | Pb | 0.03, 0.05, 0.07 g/gal | 4 vials each |
| 1638a | Lead in Reference Fuel | Pb | 2.0 g/gal | 12 vials |

Organic Constituents

| SRM | Type | Unit of Issue |
|------|--|------------------------|
| 1580 | Shale Oil | Set of 5-2 mL ampoules |
| 1644 | Polyaromatic Hydrocarbon Generator Columns | Set of 3 columns |

| Constituents | SRM 1580 (µg/g) | SRM 1644 (µg/kg) |
|--------------------|-----------------|------------------|
| Anthracene | -- | 30.7 |
| Benzo[a]anthracene | -- | 6.45 |
| Benzo[a]pyrene | 21 | 1.13 |
| Benzo[e]pyrene | 18 | -- |
| Fluoranthene | 54 | -- |
| o-Cresol | 385 | -- |
| Phenol | 407 | -- |
| Perylene | 3.4 | -- |
| Pyrene | 104 | -- |
| 2,6-Dimethylphenol | 175 | -- |
| Benzo[f]quinoline | 16 | -- |

Appendix I.

Alphabetical Index by Standard Reference Material Name

| Name | SRM | Name | SRM |
|---|-------|---|-------|
| Acetanilide | 141c | Aluminum, Freezing Point Standard | 44f |
| Acid Open-Hearth Steel, 0.2% Carbon | 19G | Aluminum, Magnetic Gram Susceptibility | 763 |
| Acid Potassium Phthalate | 84j | Aluminum Oxide, Melting Point | 742 |
| AISI 1045 Steel | 20g | Aluminum Rod Ultra Purity | RM 1R |
| AISI 4340 Steel | 36l | Aluminum-26 Radioactivity Standard | 4229 |
| AISI 4340 Steel | 1261a | Americium-241 Alpha-Particle Standard | 4904F |
| AISI 94B17 Steel (Modified) | 362 | Americium-241 Gamma-ray Standard | 4213 |
| AISI 94B17 Steel (Modified) | 1262a | Ammonium Dihydrogen Phosphate | 194 |
| Albacore Tuna | RM 50 | Angiotensin I (Human) | 998 |
| Alkali Lead Silicate Glass | 712 | Anisic Acid | 142 |
| Alpha Quartz | 1878 | Anticonvulsant Drug Level Assay Standard | 1599 |
| Alumina (Reduction Grade) | 699 | Antiepilepsy Drug Level Assay Standard | 900 |
| Alumina Silicate Glass | 714 | Antimony-125-Tellurium-125m, Europium-154, Europium-155 Mixed- Radionuclide Point-Source Standard | 4275B |
| Aluminosilicate Glass | 715 | Antimony-125-Tellurium-125m, Europium-154, Europium-155 Mixed- Radionuclide Solution Standard | 4276B |
| Aluminum Alloy | 85B | A.O.H., 0.4C Spectrographic Steel Standard | 413 |
| Aluminum Alloy 6011 (Modified) | 858 | Argillaceous Limestone | 1C |
| Aluminum Alloy 6011 (Modified) | 1258 | Arsenic Trioxide Reductometric Standard | 83d |
| Aluminum Alloy 7075 | 859 | Assay-Isotopic Standard for Potassium | 985 |
| Aluminum Alloy 7075 | 1259 | Assay-Isotopic Standard for Rhenium | 989 |
| Aluminum Block, Eddy Current Conductivity | 1860 | Assay-Isotopic Standard for Silicon | 990 |
| Aluminum Block, Eddy Current Conductivity | 1861 | Assay-Isotopic Standard for Strontium | 987 |
| Aluminum Block, Eddy Current Conductivity | 1862 | 2% Austenite in Ferrite | 488 |
| Aluminum Block, Eddy Current Conductivity | 1863 | 5% Austenite in Ferrite | 485a |
| Aluminum Brass Standard for Optical Emission and X-ray Spectroscopic Analysis | 1118 | 15% Austenite in Ferrite | 486 |
| Aluminum Brass Standard for Optical Emission and X-ray Spectroscopic Analysis | C1118 | 30% Austenite in Ferrite | 487 |
| Aluminum Brass Standard for Optical Emission and X-ray Spectroscopic Analysis | 1119 | | |
| Aluminum Brass Standard for Optical Emission and X-ray Spectroscopic Analysis | C1119 | | |
| Aluminum Casting Alloy 356 | 855 | | |
| Aluminum Casting Alloy 380 | 856 | | |
| Aluminum Cube Ultra Purity | RM 1C | | |
| Aluminum 2-Ethylhexanoate | 1075a | | |

| Name | SRM | Name | SRM |
|---|-------|--|-------|
| Austenitic Stainless Steel, Thermal Conductivity and Electrical Resistivity | 1460 | Beryllium on Filter Media | 2675 |
| Austenitic Stainless Steel, Thermal Conductivity and Electrical Resistivity | 1461 | Bessemer Steel (Simulated) 0.1% Carbon | 8j |
| Austenitic Stainless Steel, Thermal Conductivity and Electrical Resistivity | 1462 | Bilirubin | 916 |
| Barium Crown Glass | 713 | Bis(1-phenyl-1, 3-butanediono) copper (II) | 1080a |
| Barium Cyclohexanebutyrate | 1051b | Bis(1-phenyl-1, 3-butanediono) oxovanadium (IV) | 1052b |
| Barrium-133 Radioactivity Point-Source Standard | 4241B | Black Porcelain Enamel for Directional Hemispherical Reflectance | 2021 |
| Barium-133 Radioactivity Standard | 4251B | Black Porcelain Enamel for Directional Hemispherical Reflectance | 2022 |
| Basalt Rock | 688 | Blast Furnace Iron Standard (Chill Cast White) | 1143a |
| Base Oil | 1083 | Blast Furnace Iron Standard (Chill Cast White) | 1144a |
| Basic Electric Spectrographic Steel Standard | 404a | B.O.H., 0.4C Spectrographic Steel Standard | 417a |
| Basic Open-Hearth Steel, 0.1% Carbon | 15g | Boric Acid | 951 |
| Basic Open-Hearth Steel, 0.1% Carbon | 335 | Boron-Doped Silicon Slices for Resistivity Measurements | 1521 |
| Basic Open-Hearth Steel, 0.1% Carbon | 1228 | Borosilicate Glass | 93a |
| Basic Open-Hearth Steel, 0.2% Carbon | 11h | Borosilicate Glass | 623 |
| Basic Open-Hearth Steel, 0.4% Carbon | 12H | Borosilicate Glass | 717 |
| Basic Open-Hearth Steel, 0.5% Carbon | 152A | Borosilicate Glass | 1825 |
| Basic Open-Hearth Steel, 0.8% Carbon | 14f | Borosilicate Glass, Thermal Expansion | 731 |
| Basic Open-Hearth Steel, 1% Carbon (Disk) | 1227 | Bovine Liver | 1577a |
| Basic Open-Hearth Steel, 1.1% Carbon | 16f | Bovine Serum Albumin | 926 |
| Basic Open-Hearth Steel, 1.1% Carbon | 337 | Bovine Serum Albumin (7% Solution) | 927 |
| 0.4C Basic Oxygen Furnace Steel | 178 | Branched Polyethylene | 1476 |
| Bauxite (Arkansas) | 69b | Brewers Yeast | 1569 |
| Bauxite (Dominican) | 697 | Bright Copper Microhardness Standard | 1894 |
| Bauxite (Jamaican) | 698 | Bright Nickel Microhardness Standard | 1895 |
| Bauxite (Surinam) | 696 | Bromobenzoic Acid | 2142 |
| Benzene in Nitrogen | 1805 | Burnt Refractory | 76a |
| Benzene in Nitrogen | 1806 | Burnt Refractory | 77a |
| Benzene Permeation Device | 1911 | Burnt Refractory | 78a |
| Benzoic Acid | 140b | Cadmium Cyclohexanebutyrate | 1053a |
| Benzoic Acid | 350a | Cadmium, Vapor Pressure | 746 |
| Benzoic Acid Calorimetric Standard | 39i | Calcium Carbonate | 915 |
| Benzothiazyl Disulfide Rubber Compound | 373f | Calcium 2-Ethylhexanoate | 1074a |
| Beryllium-Copper Standard | 1122 | Calcium in Low-Alloy (Silicon) Steel | 1254 |
| Beryllium-Copper Standard | C1122 | Calcium Molybdate | 71 |
| Beryllium-Copper Standard | C1123 | Calibrated Glass Beads | 1004 |
| | | Calibrated Glass Beads | 1017a |
| | | Calibrated Glass Beads | 1018a |
| | | Calibrated Glass Spheres | 1003a |
| | | Carbon Dioxide in Air | 1670 |
| | | Carbon Dioxide in Air | 1671 |
| | | Carbon Dioxide in Air | 1672 |
| | | Carbon Dioxide in Nitrogen | 1674b |
| | | Carbon Dioxide in Nitrogen | 1675b |
| | | Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2619a |
| | | Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2620a |

| Name | SRM | Name | SRM |
|--|-------|--|---------|
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2621a | Catalyst Package for Lubricant Oxidation | 1817 |
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2622a | Centerline Drawings for Optical Character Recognition, B | 1901 |
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2623a | Characters | |
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2624a | Centroid Color Chart | 2106 |
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2625a | Centroid Color Kit | 2107 |
| Carbon Dioxide in Nitrogen (Combustion Efficiency Gas Standard) | 2626a | Cesium-137, Barium-137m Point-Source Radioactivity Standard | 4200B |
| Carbon Dioxide in Nitrogen (Mobile Source Emission Gas Standard) | 2632 | Cesium-137, Barium-137m Point-Source Radioactivity Standard | 4207 |
| Carbon Dioxide in Nitrogen (Mobile Source Emission Gas Standard) | 2633 | Cesium-137 Burn-Up Standard | 4233B |
| Carbon Monoxide in Air (Ambient Air Quality Gas Standard) | 2612a | Cesium-134 Radioactivity Standard | 4250B |
| Carbon Monoxide in Air (Ambient Air Quality Gas Standard) | 2613a | Channel Black Rubber Compound | 375g |
| Carbon Monoxide in Air (Ambient Air Quality Gas Standard) | 2614a | Chlorine-36 Beta-ray Standard | 4943 |
| Carbon Monoxide in Nitrogen | 1677c | Chlorine-36 Radioactivity Standard | 4422L |
| Carbon Monoxide in Nitrogen | 1678c | Chlorobenzoic Acid | 2144 |
| Carbon Monoxide in Nitrogen | 1679c | Chrome Refractory | 103a |
| Carbon Monoxide in Nitrogen | 1680b | Chromium-Molybdenum-Aluminum Steel | 106B |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2635 | Chromium-Molybdenum Steel | 36b |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2636 | Chromium-Molybdenum Steel | 133B |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2637 | Chromium-Nickel-Molybdenum Steel | 139b |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2638 | Chromium-Nickel-Molybdenum Steel | 1222 |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2639 | 17Chromium-9 Nickel-0.2 Selenium Steel | 339 |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2640 | Chromium-Nickel Spectrographic Steel Standard | 408a |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2641 | 15Chromium-7 Nickel Steel | 344 |
| Carbon Monoxide in Nitrogen (Mobile Source Emission Gas Standard) | 2642 | 16Chromium-4 Nickel Steel | 345 |
| Carbon-14 Radioactivity Standard | 4245 | Chromium-51 Radioactivity Standard | 4400L-F |
| Carbon-14 Radioactivity Standard | 4246 | Chromium Steel | 163 |
| Carbon Steel | 1224 | Chromium-Tungsten Steel | 155 |
| Carbon Steel, 0.6% | 13g | Chromium-Vanadium Spectrographic Steel Standard | 407a |
| Cast Iron | 4k | Cholesterol | 911a |
| Cast Iron | 5L | Chrysotile Asbestos Fibers | 1876 |
| Cast Iron | 6g | Citrus Leaves | 1572 |
| Cast Iron | 7G | Clinical Laboratory Thermometer | 934 |
| Cast Iron Car Wheel | 122h | Cobalt Cyclohexanebutyrate | 1055b |
| Cast Steel 3 | C1173 | Cobalt-Molybdenum-Tungsten Steel | 153A |
| Cast Steel Standard | 1138a | Cobalt-57 Radioactivity Standard | 4408L-C |
| Cast Steel Standard | 1139a | Cobalt-60 Radioactivity Standard | 4915D |
| | | Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | 1115 |
| | | Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | C1115 |
| | | Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | 1116 |

| Name | SRM | Name | SRM |
|--|-------|---|--------|
| Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | C1116 | Cupro-Nickel, 10% (CDA 706) High Purity | 874 |
| Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | 1117 | Cystine | 143c |
| Commercial Bronze Standard for Optical Emission and X-ray Spectroscopic Analysis | C1117 | Dextrose | 41b |
| Common Lead Isotopic Standard | 981 | D-Glucose | 917 |
| Copper Concentrate | 332 | Dibutyltin Bis(2-ethylhexanoate) | 1057b |
| Copper Heat Capacity Test Specimen | RM5 | Didymium Glass Filter for Checking the Wavelength Scale of Spectrophotometers | 2009 |
| Copper-Nickel-Chromium Cast Iron | 115A | Didymium Glass Filter for Checking the Wavelength Scale of Spectrophotometers | 2010 |
| Copper Ore, Mill Heads | 330 | Disodium Hydrogen Phosphate | 186IIc |
| Copper Ore, Mill Tails | 331 | Disodium Hydrogen Phosphate | 2186II |
| Copper-Thermal Expansion | 736a | D-Mannitol | 920 |
| Copper, Secondary Freezing Point Standard | 45d | Dolomitic Limestone | 88a |
| Cortisol (Hydrocortisone) | 921 | Doped Platinum | 681L1 |
| Creatinine | 914 | Doped Platinum | 681L2 |
| Cr-Mo Low Alloy Steel | 1270 | Ductile Cast Iron | 341 |
| Cr-Mo Steel (ASTM A-213) | 291 | Electrical Residual Resistivity Ratio Standard | 769 |
| Cr-Mo (SAE 4140) Spectrographic Steel Standard | 414 | Electrolytic Iron | 365 |
| Cr-Mo (SAE 4150) Spectrographic Steel Standard | 427 | Electrolytic Iron | 1265a |
| Cr-Mo (SAE X4130) Spectrographic Steel Standard | 418a | Electrolytic Iron, Thermal Conductivity and Electrical Resistivity | 1463 |
| Cr-Ni-Mo Steel (AISI 8620) | 293 | Electrolytic Iron, Thermal Conductivity and Electrical Resistivity | 1464 |
| 18Cr-10Ni Steel (AISI 304L) | 101f | Electronic and Magnetic Alloy Standard | 1159 |
| Cr-V Steel (Modified) | 363 | Electronic and Magnetic Alloy Standard | 1160 |
| Cr-V Steel (Modified) | 1263a | Enriched Boric Acid | 952 |
| Cr-V Steel (SAE 6150) | 30f | Equal-Atom Lead Isotopic Standard | 982 |
| Crystalline Potassium Dichromate | 935 | Estuarine Sediment | 1646 |
| Crystalline Potassium Iodide, Heterochromatic Stray Radiant Energy Standard | 2032 | Europium-152 Point-Source Standard | 4218E |
| Crystalline (Ruby) Electron Paramagnetic Resonance Absorption Intensity Standard | 2601 | Europium-152 Radioactivity Standard | 4370B |
| Cupro-Nickel (CDA 706) | 1275 | Extra Dense Lead Glass | 709 |
| Cupro-Nickel (CDA 715) | 1276 | Fe-Cr-Ni Alloy Microprobe Standard | 479a |
| Cupro-Nickel, 10% (CDA 706) Doped | 875 | Fe-3Si Alloy Microprobe Standard | 483 |
| | | Feldspar | 70a |
| | | Feldspar | 99a |
| | | Ferrochromium (Low Carbon) | 196 |
| | | Ferrochromium Silicon | 689 |
| | | Ferroniobium | 340 |
| | | Ferrophosphorus | 90 |
| | | Ferrosilicon | 58a |
| | | Ferrosilicon | 59a |
| | | Ferrosilicon (75% Si) | 195 |
| | | First Surface Aluminum Mirror for Specular Reflectance | 2003a |
| | | First Surface Mirror, Gold on Glass | 2008a |

| Name | SRM | Name | SRM |
|-------------------------------------|---------|--------------------------------------|---------|
| Fission Track Glass Standard | 961 | Gold-198 Radioactivity Standard | 4405L-B |
| Fission Track Glass Standard | 962a | Gold-Silver Wires for Microprobe | 481 |
| Fission Track Glass Standard | 963a | Analysis | |
| Fission Track Glass Standard | 964 | Gold, Vapor Pressure | 745 |
| Flint Clay | 97a | Gray Cast Iron | 334 |
| Fluorobenzoic Acid | 2143 | Halocarbons (in methanol) for Water | 1639 |
| Fluorspar | 79a | Analysis | |
| Free-Cutting Brass | 1103 | High-Alloy Steel (A-743) | C1288 |
| Free-Cutting Brass | C1104 | High-Alloy Steel (AISI 310 Mod.) | C1287 |
| Freeze-Dried Urine | 2670 | High-Alloy Steel, (AISI 414 Mod.) | C1289 |
| Freeze-Dried Urine Certified | 2671a | High-Alloy White Cast | 892 |
| for Fluoride | | High-Alloy White Cast Iron | 890 |
| Freeze-Dried Urine Certified | 2672a | High-Alloy White Cast Iron | 891 |
| for Mercury | | High-Carbon Ferrochromium | 64c |
| Fused-Silica Thermal Expansion | 739 | High-Carbon Ferromanganese | 68c |
| Gadolinium-148 Alpha-Particle | 4907 | High-Carbon Steel (Modified) | 364 |
| Standard | | High-Carbon Steel (Modified) | 1264a |
| Gallium Melting-Point Standard | 1968 | High-Grade Fluorspar | 180 |
| Gallium-67 Radioactivity Standard | 4416L-D | High-Nickel Steel | 126c |
| Gas Furnace Black Rubber Compound | 382a | High-Nickel Steel | 1158 |
| Gasometric Set (1095-1099) | 1089 | High-Purity Gold | 685 |
| Gasometric Standard for Unalloyed | 357 | High-Purity Platinum | 680L1A |
| Zirconium | | High-Purity Platinum | 680L2A |
| Gasometric Standard for Unalloyed | 358 | High-Purity Platinum Thermoelement | 1967 |
| Zirconium | | High-Purity Zinc | 682 |
| Generator Columns for Polynuclear | 1644 | High-Silicon Steel | 179 |
| Aromatic Hydrocarbons | | High-Silicon Steel | 1134 |
| Gilding Metal | 1112 | High-Silicon Steel | 1135 |
| Gilding Metal | C1112 | High-Silicon Steel (Calcium Bearing) | 125b |
| Gilding Metal | 1113 | High-Sulfur Steel | 105 |
| Gilding Metal | C1113 | High-Sulfur Steel | 129c |
| Gilding Metal | 1114 | High-Sulfur Steel | 1136 |
| Gilding Metal | C1114 | High Temperature Alloy A286 | 348 |
| Glasses for Microchemical Analysis | 1871 | High Temperature Alloy M308 | 1197 |
| Glasses for Microchemical Analysis | 1872 | High Temperature Alloy L605 and | S1199 |
| Glasses for Microchemical Analysis | 1873 | S816 | |
| Glasses for Microchemical Analysis | 1874 | High-Temperature Alloy | 1206-2 |
| Glasses for Microchemical Analysis | 1875 | High-Temperature Alloy | 1207-1 |
| Glass Fibers for Microanalysis | RM 31 | High-Temperature Alloy | 1207-2 |
| Glass Filter for Transmittance | 2030 | High-Temperature Alloy | 1208-1 |
| Measurement | | High-Temperature Alloy | 1208-2 |
| Glass Filters for Spectrophotometry | 930D | Homogeneous River Sediment for | RM 45B |
| Glass Fluorescence Source | 477 | Radioactivity Measurements | |
| Glass Sand | 81a | Human Liver, Environmental | 4352 |
| Glass Sand | 165a | Radioactivity | |
| Glass Spheres | 1019a | Human Lung, Environmental | 4351 |
| Gold Coating on Glass Sealing Alloy | 1398a | Radioactivity | |
| Gold Coating on Nickel | 1379 | Human Serum | 909 |
| Gold Coating on Nickel | 1380 | | |
| Gold Coating on Nickel | 1399b | | |
| Gold-Copper Wires for Microprobe | 482 | | |
| Analysis | | | |
| Gold-195 Radioactivity Standard | 4421L | | |

| Name | SRM | Name | SRM |
|---|---------|--|---------|
| Hydrogen in Unalloyed Titanium | 352b | Iron Ore (Sibley) | 27f |
| Hydrogen in Unalloyed Titanium | 1086 | Iron Ore Concentrate (Canada) | 690 |
| Hydrogen in Unalloyed Titanium | 1087 | Iron-59 Radioactivity Standard | 4411L-B |
| Hydrogen in Unalloyed Titanium | 1088 | Isobutylene-Isoprene (Butyl) Rubber | 1495 |
| Hydrogen-3 Radioactivity Standard | 4361 | Isobutylene-Isoprene (Butyl) Rubber | 388L |
| Hydrogen-3 Radioactivity Standard | 4926C | Isotopic Standard for Bromine | 977 |
| Hydrogen-3 Toluene Radioactivity Standard | 4947 | Isotopic Standard for Chlorine | 975 |
| 4-Hydroxy-3 methoxy-DL-mandelic Acid (VMA) | 925 | Isotopic Standard for Chromium | 979 |
| ICTA High Temperature Set Differential Thermal Analysis | GM 760 | Isotopic Standard for Copper | 976 |
| ICTA Low Temperature Set Differential Thermal Analysis | GM 757 | Isotopic Standard for Magnesium | 980 |
| ICTA Mod Temperature Set Differential Thermal Analysis | GM 759 | Isotopic Standard for Silver | 978 |
| ICTA Mid Temperature Set Differential Thermal Analysis | GM 758 | Krypton-85 Gaseous Radioactivity Standard | 4308C |
| ICTA Polystyrene Differential Thermal Analysis | GM 754 | Krypton-85 Radioactivity Standard | 4235 |
| ICTA Thermogravimetry Set | GM 761 | Krypton-85 Radioactivity Standard | 4935C |
| Incoloy, 901 and Hastelloy X | S1198 | Lead-Barium Glass | 89 |
| Inconels, Alloy 600 (Chips) | 864 | Lead-Base Bearing Metal | 53e |
| Inconels, Alloy 600 (Solid) | 1244 | Lead-Base Bearing Metal | 1132 |
| Inconels, Alloy 625 (Chips) | 865 | Lead Cyclohexanebutyrate | 1059c |
| Inconels, Alloy 625 (Solid) | 1245 | Lead in Reference Fuel | 1636a |
| Incoloy, Alloy 800 (Chips) | 866 | Lead in Reference Fuel | 1637a |
| Incoloy, Alloy 800 (Solid) | 1246 | Lead in Reference Fuel | 1638a |
| Incoloy, Alloy 825 (Chips) | 867 | Lead Nitrate | 928 |
| Incoloy, Alloy 825 (Solid) | 1247 | Lead on Filter Media | 2674 |
| Indium-111 Radioactivity Standard | 4417L-C | Lead-203 Radioactivity Standard | 4420L |
| Ingot Iron Spectrographic Steel Standard | 420a | Lead, Secondary Freezing Point Standard | 49e |
| Intermediate Purity Selenium | 726 | Lead-Silica Glass | 1827 |
| Intermediate-Purity Zinc | 728 | Lead-Silica Glass (Viscosity) | 711 |
| Iodine-123 Radioactivity Standard | 4414L-C | Lead-Silica Glass for dc Volume Resistivity | 624 |
| Iodine-125 Radioactivity Standard | 4407L-H | Lead-Silica Glass for Dielectric Constant | 774 |
| Iodine-129 Radioactivity Standard | 4949B | Lead 206 Spike Assay and Isotopic Solution Standard | 991 |
| Iodine-131 Radioactivity Standard | 4401L-I | Leaded-Tin Bronze Alloy | 1035 |
| Iron Foil Mössbauer Standard | 1541 | Light-Sensitive Paper | 700d |
| Iron-55 Low-Energy Photon Standard | 4260C | Light-Sensitive Paper | 701d |
| Iron Metal (Clinical Standard) | 937 | Light-Sensitive Plastic Chip | 703 |
| Iron Ore (Labrador) | 692 | Linear Polyethylene | 1475 |
| Iron Ore (Nimba) | 693 | Linear Polyethylene | 1482 |
| | | Linear Polyethylene | 1483 |
| | | Linear Polyethylene | 1484 |
| | | Linerboard, Standard for Tape Adhesion Testing | 1810 |
| | | Liquid Absorbance Standard for Ultraviolet and Visible Spectrophotometry | 931c |
| | | Lithium Carbonate | 924 |
| | | Lithium Ore | 181 |
| | | Lithium Ore | 182 |
| | | Lithium Ore | 183 |
| | | Low-Alloy Steel, (AISI 4130) | 1225 |
| | | Low Alloy Steel | 1226 |
| | | Low Alloy Steel (A242 Mod.) | C1285 |
| | | Low-Alloy Steel, AISI 4130 | 72g |
| | | Low Alloy Steel (AISI 1526, Modified) | 1269 |
| | | Low-Alloy Steel (Hy 80) | 1286 |

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| Low-Alloy Steel Set (661-665) | S668 | Naval Brass Standards for Optical Emission and Spectroscopic Analysis | 1108 |
| Low-Carbon Silicon Steel | 131c | | |
| Low-Carbon Silicon Steel | 1036 | Naval Brass Standards for Optical Emission and Spectroscopic Analysis | C1108 |
| Low-Carbon Stainless Steel (AISI 316L) | 166c | | |
| Magnesium-base Alloy | 171 | Neutral Glass | 716 |
| Magnesium Cyclohexanebutyrate | 1061c | Neutron Density Monitor Wire | 953 |
| Magnesium Gluconate Dihydrate | 929 | Nickel-Chromium Cast Iron | 82b |
| Magnetic Coating on Magnetic Substrate (Nickel on Steel) | 1365a | Nickel-Chromium-Molybdenum Cast Iron | 107C |
| Magnetic Coating on Magnetic Substrate (Nickel on Steel) | 1366a | Nickel-Chromium Steel | 32E |
| Magnetic Coating on Non-Magnetic Substrate (Nickel and Chromium on brass) | 1367a | Nickel-Copper Alloy | 882 |
| Magnetic Tape, High Density | 6250 | Nickel Cyclohexanebutyrate | 1065b |
| Manganese Fluoride, Magnetic Gram Susceptibility | 766 | Nickel Oxide, No. 1 | 671 |
| Manganese Ore | 25d | Nickel Oxide, No. 2 | 672 |
| Manganese-54 Point-Source Radioactivity Standard | 4997E | Nickel Oxide, No. 3 | 673 |
| Manganese-54 Radioactivity Standard | 4257 | Nickel-63 Radioactivity Standard | 4226 |
| Manganese Steel | 100B | Nickel Silver (CDA 762) | 879 |
| Manganous Cyclohexanebutyrate | 1062b | Nickel Silver (CDA 770) | 880 |
| Maraging Steel | 1156 | Nickel Spectrographic Steel Standard | 409b |
| Metal on Quartz Filters for Spectrophotometry | 2031 | Nickel Sphere, Magnetic Moment | 772 |
| Metals on Filter Media | 2676b | Nickel Steel | 33d |
| Methane in Air | 1658a | Ni-Cr-Mo-V Steel | 1173 |
| Methane in Air | 1659a | Nicotinic Acid | 148 |
| Methane in Air | 1660a | Niobium-94 Gamma-ray Standard | 4201B |
| Medium Manganese Spectrographic Steel Standard | 405a | Nitric Oxide in Nitrogen | 1683b |
| Mercaptobenzothiazole | 383a | Nitric Oxide in Nitrogen | 1684b |
| Mercury, Freezing Point | 743 | Nitric Oxide in Nitrogen | 1685b |
| Mercury-203 Radioactivity Standard | 4418L | Nitric Oxide in Nitrogen | 1686b |
| Mercury in Water, µg/mL | 1641b | Nitric Oxide in Nitrogen | 1687b |
| Mercury in Water, ng/mL | 1642b | Nitric Oxide in Nitrogen (Mobile Source Emission Gas Standard) | 2627 |
| Microcopy Resolution Test Chart | 1010a | Nitric Oxide in Nitrogen (Mobile Source Emission Gas Standard) | 2628 |
| Microprobe Standard - Cartridge Brass | 478 | Nitric Oxide in Nitrogen (Mobile Source Emission Gas Standard) | 2629 |
| Mineral Glasses for Microanalysis | 470 | Nitric Oxide in Nitrogen (Mobile Source Emission Gas Standard) | 2630 |
| Molybdenum Concentrate | 333 | Nitric Oxide in Nitrogen (Mobile Source Emission Gas Standard) | 2631 |
| Molybdenum, Heat Capacity | 781 | Nitrogen Dioxide in Air (Stationary Source Emission Gas Standard) | 2653 |
| Molybdenum-99 Radioactivity Standard | 4412L-H | Nitrogen Dioxide in Air (Stationary Source Emission Gas Standard) | 2654 |
| Molybdenum-Tungsten-Chromium-Vanadium Steel | 134A | Nitrogen Dioxide in Air (Stationary Source Emission Gas Standard) | 2655 |
| Naval Brass Standards for Optical Emission and Spectroscopic Analysis | 1106 | Nitrogen Dioxide in Air (Stationary Source Emission Gas Standard) | 2656 |
| Naval Brass Standards for Optical Emission and Spectroscopic Analysis | C1106 | Nitrogen Dioxide Permeation Device | 1629a |
| Naval Brass Standards for Optical Emission and Spectroscopic Analysis | 1107 | 4-Nitrophenol | 938 |
| Naval Brass Standards for Optical Emission and Spectroscopic Analysis | C1107 | | |

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| Nodular Cast Iron | 342a | Organics in Shale Oil | 1580 |
| Nominal One Micrometer Polystyrene Spheres | 1690 | Oxalic Acid | 4990C |
| Non-Fat Powdered Milk | 1549 | Oxygen in Ferrous Materials Ingot Iron | 1090 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1359 | Oxygen in Ferrous Materials (Stainless Steel AISI 431) | 1091 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1360 | Oxygen in Ferrous Materials Vacuum Melted Steel | 1092 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1361b | Oxygen in Maraging Steel | 1094 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1362a | Oxygen in Nitrogen (Gas Standard) | 2657 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1363a | Oxygen in Nitrogen (Gas Standard) | 2658 |
| Nonmagnetic Coating on Magnetic Substrate (Copper and Chromium on Steel) | 1364a | Oxygen in Nitrogen (Gas Standard) | 2659 |
| NPL GM Alpha Alumina | 8005 | Oxygen in Titanium-Base Materials | 355 |
| NPL GM Alpha Alumina | 8006 | Oxygen in Valve Steel | 1093 |
| NPL GM Alpha Alumina | 8007 | Oyster Tissue | 1566 |
| NPL GM Alpha Alumina | 8008 | Palladium, Magnetic Gram Susceptibility | 765 |
| NPL GM Graphitized Carbon Black | 8001 | Penetrant Test Block | 1850 |
| NPL GM Graphitized Carbon Black | 8002 | Peruvian Soil, Environmental Radioactivity | 4355 |
| NPL GM Melting Point Set | 8000 | Petroleum Crude Oil | 1582 |
| NPL GM Non-porous Silica | 8003 | Phosphate Rock (Florida) | 120b |
| NPL GM Non-porous Silica | 8004 | Phosphor Bronze (CDA 521) | 871 |
| N-tertiary-Butyl-2-benzothiazolesulfenamide Rubber Compound | 384d | Phosphor Bronze (CDA 544) | 872 |
| Obsidian Rock | 278 | Phosphorized Copper, Cu VIII | C1251 |
| Octaphenylcyclotetrasiloxane | 1066a | Phosphorized Copper, Cu IX | C1252 |
| Oil Furnace Black Rubber Compound | 378b | Phosphorized Copper, Cu X | C1253 |
| Opal Glass Powder | 91 | Phosphorus-32 Radioactivity Standard | 4406L-G |
| Optical Emission and X-ray Spectroscopic Analysis | 1102 | Photographic Step Tablet | 1008 |
| Optical Microscope Linewidth Measurement Standard | 474 | Pine Needles | 1575 |
| Optical Microscope Linewidth Measurement Standard | 475 | Plastic Clay | 98a |
| Optical Microscope Linewidth Measurement Standard | 476 | Platinum, Magnetic Gram Susceptibility | 764 |
| | | Plutonium-238 Alpha-Particle Standard | 4906B |
| | | Plutonium-240 Alpha-Particle Emission-Rate Solution Standard | 4338 |
| | | Plutonium-239 Alpha-Particle Solution Standard | 4331 |
| | | Plutonium-242 Alpha-Particle Solution Standard | 4334B |
| | | Plutonium Isotopic Standard | 946 |
| | | Plutonium Isotopic Standard | 947 |
| | | Plutonium Isotopic Standard | 948 |
| | | Plutonium Metal | 949f |
| | | Plutonium Metal (Standard Matrix Material) | 945 |
| | | Plutonium-244 Spike Assay and Isotopic Standard | 996 |
| | | Polychlorinated Biphenyls in Oil | 1581 |
| | | Polycrystalline Alumina Elasticity Standard | 718 |
| | | Polyester Plastic Film for Oxygen Gas Transmission | 1470 |
| | | Polyisobutylene Solution in Cetane | 1490 |
| | | Polystyrene | 1478 |
| | | Polystyrene | 1479 |
| | | Polystyrene (Broad Molecular Weight) | 706 |
| | | Polystyrene (Narrow Molecular Weight) | 705 |
| | | Polystyrene Spheres | 1691 |
| | | Portland Cement (Black) | 1880 |

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| Portland Cement (Blue) | 635 | Quartz on Filter Media | 2679a |
| Portland Cement (Clear) | 639 | Quinine Sulfate Dihydrate | 936 |
| Portland Cement (Gold) | 634 | Radiogenic Lead Isotopic Standard | 983 |
| Portland Cement (Green) | 638 | Radium-226 Gamma-ray Standard | 4956 |
| Portland Cement (Pink) | 637 | Radium-226 Gamma-ray Standard | 4957 |
| Portland Cement (Red) | 633 | Radium-226 Gamma-ray Standard | 4958 |
| Portland Cement (White) | 1881 | Radium-226 Gamma-ray Standard | 4959 |
| Portland Cement (Yellow) | 636 | Radium-226 Gamma-ray Standard | 4960 |
| Portland Cement Fineness Standard | 114n | Radium-226 Gamma-ray Standard | 4961 |
| Potassium Chloride | 2202 | Radium-226 Gamma-ray Standard | 4962 |
| Potassium Chloride (Clinical Standard) | 918 | Radium-226 Gamma-ray Standard | 4963 |
| Potassium Chloride (Primary Chemical) | 999 | Radium-226 Gamma-ray Standard | 4964B |
| Potassium Chloride for Solution Calorimetry | 1655 | Radium Standard (Blank Solution) | 4952B |
| Potassium Dichromate | 136d | Radon-226 for Radon Analysis | 4953C |
| Potassium Dihydrogen Phosphate | 200 | Red Brass | 1109 |
| Potassium Dihydrogen Phosphate | 1861c | Red Brass | C1109 |
| Potassium Dihydrogen Phosphate | 21861 | Red Brass | 1110 |
| Potassium Erucate | 1076 | Red Brass | C1110 |
| Potassium Feldspar | 607 | Red Brass | 1111 |
| Potassium Fluoride | 2203 | Red Brass | C1111 |
| Potassium Hydrogen Phthalate | 185e | Reduced Iron Oxide | 691 |
| Potassium Hydrogen Tartrate | 188 | Reference Fuel Isooctane | 1816a |
| Potassium Iodide with Attenuator | 2033 | Reference Fuel n-Heptane | 1815a |
| Potassium Nitrate | 193 | Reflection Step Tablet | 2061 |
| Potassium Tetroxalate | 189 | Refractive Index Glass | 1820 |
| Powdered Lead Based Paint | 1579 | Refractive Index Silicone Liquids | 1823 |
| Priority Pollutant Polynuclear Aromatic Hydrocarbons (in Acetonitrile) | 1647 | Refractive Index, Soda-Lime Glass | 1822 |
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| Propane in Air | 1666b | Resulfurized-Rephosphorized Steel | C1221 |
| Propane in Air | 1667b | Rice Flour | 1568 |
| Propane in Air | 1668b | River Sediment | 1645 |
| Propane in Air | 1669b | River Sediment, Environmental Radioactivity | 4350B |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2643 | Rocky Flats Soil Number 1, Environmental Radioactivity | 4353 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2644 | Rubidium Melting Point | 1969 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2645 | Rutile Ore | 670 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2646 | Scanning Electron Microscope Magnification Standard | 484c |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2647 | Scanning Electron Microscope Performance Standard | 2069 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2648 | Secondary Standard Flexible Disk Cartridge (Computer Amplitude Reference) | 3210 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2649 | Secondary Standard Magnetic Tape | 3200 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2650 | Secondary Standard Magnetic Tape Cassette | 1600 |
| Propane in Nitrogen (Mobile Source Emission Gas Standard) | 2651 | Secondary Standard Magnetic Tape Cartridge (Computer Amplitude Reference) | 3216 |
| Propane in Nitrogen and Oxygen (Mobile Source Emission Gas Standard) | 2652 | Second Surface Aluminum Mirror for Specular Reflectance | 2023 |
| Quartz Cuvette for Spectrophotometry | 932 | | |
| Quartz for Hydrofluoric Acid Solution Calorimetry | 1654 | | |

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| Second Surface Aluminum Mirror for Specular Reflectance | 2024 | Soda-Lime Sheet Glass | 1831 |
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| Selenium-Bearing Steel | 1170b | Soda-Lime Silica Glass | 710 |
| Selenium-75 Radioactivity Standard | 4409L-D | Soda-Lime Silica Glass for Liquidus Temperature | 773 |
| Sheet Brass | 37E | Sodium Bicarbonate | 191a |
| Silica Brick | 198 | Sodium Bicarbonate | 2191 |
| Silica Brick | 199 | Sodium Carbonate | 192a |
| Silicon-Aluminum Alloy | 87a | Sodium Carbonate | 2192 |
| Silicon Bronze | 158A | Sodium Chloride | 2201 |
| Silicon Density Standard | 1840 | Sodium Chloride (Clinical Standard) | 919 |
| Silicon Density Standard | 1841 | Sodium Cyclohexanebutyrate | 1069b |
| Silicon Metal | 57a | Sodium Oxalate Reductometric Standard | 40h |
| Silicon Powder, Spacing Standard for X-ray Diffraction | 640a | Sodium Pyruvate | 910 |
| Silicon Power Device Level Resistivity Standard | 1522 | Sodium Tetraborate Decahydrate (Borax) | 187b |
| Silicon Resistivity Standard for Eddy Current Testers | 1523 | Solder | 127b |
| Silver 2-Ethylhexanoate | 1077a | Solder | 1131 |
| Silver-Gold Thermocouple Wire | 733 | Special Nuclear Container DOT 6M, 15 gal. | 9940 |
| Silver, Vapor Pressure | 748 | Special Nuclear Container, 55 gal. | 9941 |
| Sintered and Arc-Cast Tungsten, Thermal Conductivity and Electrical Resistivity | 1465 | Special Nuclear Container Type A, 10 gal. | 9942 |
| Sintered and Arc-Cast Tungsten, Thermal Conductivity and Electrical Resistivity | 1466 | Special Nuclear Container, Type A, 55 gal. | 9943 |
| Sintered and Arc-Cast Tungsten, Thermal Conductivity and Electrical Resistivity | 1467 | Special Nuclear Material Package | 9910 |
| Sintered and Arc-Cast Tungsten, Thermal Conductivity and Electrical Resistivity | 1468 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 461 |
| Smoke Density Chamber Standard (Flaming Exposure Condition) | 1007a | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 462 |
| Smoke Density Chamber Standard (Non-flaming Exposure Condition) | 1006b | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 463 |
| Soda-Lime Container Glass | 621 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 464 |
| Soda-Lime Flat Glass | 620 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 465 |
| Soda-Lime Float Glass | 1830 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 466 |
| Soda-Lime Glass | 1826 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 467 |
| Soda-Lime Glass Powder | 92 | Spectrographic Ingot Iron and Low-Alloy Steel Standard (Rod) | 468 |
| | | Spectrographic Ingot Iron and Low-Alloy Steel Standard | 1166 |
| | | Spectrographic Stainless Steel Standard | 442 |
| | | Spectrographic Stainless Steel Standard | 443 |
| | | Spectrographic Stainless Steel Standard | 444 |
| | | Spectrographic Stainless Steel Standard (Disc) | D849 |
| | | Spectrographic Stainless Steel Standard (Disc) | D850 |
| | | Spectrographic Stainless Steel Standard (Group II) | 445 |

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| Spectrographic Stainless Steel Standard (Group II) | 446 | Spectroscopic Titanium-Base Standard | 644 |
| Spectrographic Stainless Steel Standard (Group II) | 447 | Spectroscopic Titanium-Base Standard | 645 |
| Spectrographic Stainless Steel Standard (Group II) | 448 | Spectroscopic Titanium-Base Standard | 646 |
| Spectrographic Stainless Steel Standard (Group II) | 449 | Spheroidized Iron Carbide in Ferrite | 493 |
| Spectrographic Stainless Steel Standard (Group II) | 450 | Spreading Resistance Calibration (100) n-Type Silicon | 2529 |
| Spectrographic Stainless Steel Standard (Group II) | 849 | Spreading Resistance Calibration (100) p-Type Silicon | 2528 |
| Spectrographic Stainless Steel Standard (Rod) | 850 | Spreading Resistance Calibration (111) n-Type Silicon | 2527 |
| Spectrographic Steel Standard (Disc) | D803a | Spreading Resistance Calibration (111) p-Type Silicon | 2526 |
| Spectrographic Steel Standard (Disc) | D807a | Stabilized Wine | 1590 |
| Spectrographic Steel Standard (Rod) | 803a | Stainless Steel | 121d |
| Spectrographic Steel Standard (Rod) | 804a | Stainless Steel | 123c |
| Spectrographic Steel Standard (Rod) | 805a | Stainless Steel | 160b |
| Spectrographic Steel Standard (Rod) | 807a | Stainless Steel (AISI 446) | 367 |
| Spectrographic Steel Standard (Rod) | 808a | Stainless Steel (AISI 446) | 1267 |
| Spectrographic Steel Standard (Rod) | 809a | Stainless Steel, 13% Chromium | 73c |
| Spectrographic Steel Standard (Rod) | 817b | Stainless Steel, Cr-Ni | C1151 |
| Spectrographic Steel Standard (Rod) | 820a | Stainless Steel, Cr-Ni | 1151a |
| Spectrographic Steel Standard (Rod) | 821 | Stainless Steel, Cr-Ni | C1152 |
| Spectrographic Steel Standard (Rod) | 827 | Stainless Steel, Cr-Ni | 1152a |
| Spectrographic Tool Steel Standard | 436 | Stainless Steel, Cr-Ni | C1153 |
| Spectrographic Tool Steel Standard | 437 | Stainless Steel, Cr-Ni | 1153a |
| Spectrographic Tool Steel Standard | 438 | Stainless Steel, Cr-Ni | C1154 |
| Spectrographic Tool Steel Standard | 439 | Stainless Steel, Cr-Ni | 1154a |
| Spectrographic Tool Steel Standard | 440 | Stainless Steel, Cr-Ni-Mo | 1155 |
| Spectrographic Tool Steel Standard | 441 | Stainless Steel, Cr-Ni-Nb | 1172 |
| Spectrographic Tool Steel Standard | 837 | Stainless Steel, Cr-Ni-Ti | 1171 |
| Spectrographic Tool Steel Standard (Disc) | D837 | Stainless Steel for Pitting or Crevice Corrosion | 1890 |
| Spectrographic Tool Steel Standard (Disc) | D840 | Stainless Steel Thermal Expansion | 738 |
| Spectrographic Tool Steel Standard (Disc) | D841 | Stearic Acid Rubber Compound | 372h |
| Spectrographic Zinc-Base Die-Casting Alloy A | 625 | Steel (AISI 1211) | 368 |
| Spectrographic Zinc-Base Die-Casting Alloy B | 626 | Steel (Lead-Bearing) | 1169b |
| Spectrographic Zinc-Base Die-Casting Alloy C | 627 | Strontium Cyclohexanebutyrate | 1070a |
| Spectrographic Zinc-Base Die-Casting Alloy D | 628 | Strontium-85 Radioactivity Standard | 4403L-B |
| Spectrographic Zinc-Base Die-Casting Alloy E | 629 | Strontium-89 Radioactivity Standard | 4945D |
| Spectrographic Zinc-Base Die-Casting Alloy F | 630 | Styrene-butadiene Rubber (Type 1500) | 386h |
| Spectrographic Zinc Spelter Standard | 631 | Succinonitrile Freezing Point | 1970 |
| Spectroscopic Titanium-Base Standard | 641 | Sucrose | 17c |
| Spectroscopic Titanium-Base Standard | 642 | Sulfate and Nitrate on Filter Media | 2673 |
| Spectroscopic Titanium-Base Standard | 643 | Sulfur Dioxide in Nitrogen | 1661a |
| | | Sulfur Dioxide in Nitrogen | 1662a |
| | | Sulfur Dioxide in Nitrogen | 1663a |
| | | Sulfur Dioxide in Nitrogen | 1664a |
| | | Sulfur Dioxide in Nitrogen | 1693 |
| | | Sulfur Dioxide in Nitrogen | 1694 |

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| Sulfur Dioxide in Nitrogen | 1696 | Titanium-Base Alloy (Unalloyed) | 650 |
| Sulfur Dioxide Permeation Tube (2 cm tube) | 1627 | Titanium-Base Alloy (Unalloyed) | 651 |
| Sulfur Dioxide Permeation Tube (5 cm tube) | 1626 | Titanium-Base Alloy (Unalloyed) | 652 |
| Sulfur Dioxide Permeation Tube (10 cm tube) | 1625 | Titanium Dioxide | 154b |
| Sulfur in Coal | 2682 | Toluene | 211c |
| Sulfur in Coal | 2683 | Tomato Leaves | 1573 |
| Sulfur in Coal | 2684 | Tool Steel (AISI M2) | 132b |
| Sulfur in Coal | 2685 | Tool Steel (AISI M2) | 1157 |
| Sulfur in Residual Fuel Oil | 1619 | Tool Steel Abrasive Wear Standard | 1857 |
| Sulfur in Residual Fuel Oil | 1620a | Tracealloy (Nickel-Base High-Temperature Alloy) | 897 |
| Sulfur in Residual Fuel Oil | 1621b | Tracealloy (Nickel-Base High-Temperature Alloy) | 898 |
| Sulfur in Residual Fuel Oil | 1622b | Tracealloy (Nickel-Base High-Temperature Alloy) | 899 |
| Sulfur in Residual Fuel Oil | 1623a | Trace Elements in a Glass Matrix | 610 |
| Sulfur in Residual Fuel Oil | 1624a | Trace Elements in a Glass Matrix | 611 |
| Sulfur Rubber Compound | 371g | Trace Elements in a Glass Matrix | 612 |
| Superconductive Thermometric Fixed Point Device | 767a | Trace Elements in a Glass Matrix | 613 |
| Superconductive Thermometric Fixed Point Device | 768 | Trace Elements in a Glass Matrix | 614 |
| Surface Flammability Standard | 1002c | Trace Elements in a Glass Matrix | 615 |
| Synthetic Sapphire | 720 | Trace Elements in a Glass Matrix | 616 |
| Technetium-99 Radioactivity Standard | 4288 | Trace Elements in a Glass Matrix | 617 |
| Technetium-99m Radioactivity Standard | 4410H-I | Trace Elements in Coal (Bituminous) | 1632a |
| Tetrachloroethylene in Nitrogen | 1808 | Trace Elements in Coal (Sub- bituminous) | 1635 |
| Thallium-201 Radioactivity Standard | 4404L-F | Trace Elements in Coal Fly Ash | 1633a |
| Thermal Resistance, Fibrous Glass Batt | 1451 | Trace Elements in Fuel Oil | 1634a |
| Thermal Resistance, Fibrous Glass Board | 1450b | Trace Elements in Water | 1643a |
| Thorium-228, Thallium-208 Gamma-ray Point-Source Standard | 4206C | Trace Mercury in Coal | 1630 |
| Tin-Base Bearing Metal | 54D | 2,2,4-Trimethylpentane | 217c |
| Tin, Freezing Point | 741 | Tripalmitin | 1595 |
| Tin-113-Indium-113m Radioactivity Standard | 4402L-C | Tris, Basimetric | 723a |
| Tin-121m Point-Source Gamma-ray Emission-Rate Standard | 4264B | Tris, for Solution Calorimetry | 724a |
| Tin, Secondary Freezing Point Standard | 42g | Tris(hydroxymethyl)aminomethane | 922 |
| Titanium Alloy | 654a | Tris(hydroxymethyl)aminomethane hydrochloride | 923 |
| Titanium-Base Alloy | 173b | Tris(1-phenyl-1, 3-butanediono) Chromium (III) | 1078b |
| Titanium-Base Alloy | 176 | Tris(1-phenyl-1, 3-butanediono) Iron (III) | 1079b |
| | | Triphenyl Phosphate | 1071b |
| | | Tungsten Carbide | 276a |
| | | Tungsten-Chromium-Vanadium Steel | 50c |
| | | Tungsten Concentrate | 277 |
| | | Tungsten, Heat Capacity | 782 |
| | | Tungsten-20% Molybdenum Alloy Electron Microprobe Standard | 480 |
| | | Tungsten Thermal Expansion | 737 |
| | | Unalloyed Copper | 1034 |
| | | Unalloyed Copper, Cu "O" | 393 |
| | | Unalloyed Copper, Cu IV | 457 |
| | | Unalloyed Copper, Cu XI | 454 |
| | | Unalloyed Copper, Cu I (Chip) | 394 |
| | | Unalloyed Copper, Cu II (Chip) | 395 |
| | | Unalloyed Copper, Cu III (Chip) | 396 |
| | | Unalloyed Copper, Cu V (Chip) | 398 |
| | | Unalloyed Copper, Cu VI (Chip) | 399 |
| | | Unalloyed Copper, Cu VII (Chip) | 400 |
| | | Unalloyed Copper, Cu I (Rod) | 494 |

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| Unalloyed Copper, Cu II (Rod) | 495 | Wear-Metals in Lubricating Oil (300 ppm) | 1085 |
| Unalloyed Copper, Cu III (Rod) | 496 | Wheat Flour | 1567 |
| Unalloyed Copper, Cu V (Rod) | 498 | White Cast Iron | 338 |
| Unalloyed Copper, Cu VI (Rod) | 499 | White Cast Iron (Disc) | 1145 |
| Unalloyed Copper, Cu VII (Rod) | 500 | White Cast Iron (Disc) | 1146 |
| Unalloyed Titanium | 354 | White Cast Iron (Disc) | 1150 |
| Uranium Isotopic Standard (Nominally depleted to 0.02%) | U-0002 | White Ceramic Tile for Directional Hemispherical Reflectance | 2019b |
| Uranium Isotopic Standard | U-005a | White Ceramic Tile for Directional Hemispherical Reflectance | 2020 |
| Uranium Isotopic Standard (Nominally 1% Enriched) | U-010 | White Iron | 3d |
| Uranium Isotopic Standard (Nominally 1.5% Enriched) | U-015 | White Opan Glass Diffuse Spectral Reflectance Standard for the Visible Spectrum | 2015 |
| Uranium Isotopic Standard | U-020 | Xenon-127 Gaseous Radioactivity Standard | 4309G |
| Uranium Isotopic Standard | U-030a | Xenon-133 Gaseous Radioactivity Standard | 4307I |
| Uranium Isotopic Standard (Nominally 5% Enriched) | U-050 | Xenon-133 Gaseous Radioactivity Standard | 4415L-I |
| Uranium Isotopic Standard (Nominally 10% Enriched) | U-100 | Xenon-133, Xenon-137, Krypton-85 Mixed Gaseous Radioactivity Standard | 4310B |
| Uranium Isotopic Standard (Nominally 15% Enriched) | U-150 | X-ray Film Step Tablet | 1001 |
| Uranium Isotopic Standard (Nominally 20% Enriched) | U-200 | X-ray Powder Diffraction Intensity Standard | 674 |
| Uranium Isotopic Standard (Nominally 35% Enriched) | U-350 | X-ray Powder Diffraction (Mica) Low 2 Theta | 675 |
| Uranium Isotopic Standard (Nominally 50% Enriched) | U-500 | Ytterbium-169 Radioactivity Standard | 4419L-B |
| Uranium Isotopic Standard (Nominally 75% Enriched) | U-750 | Zinc-Base Alloy (Die Casting) | 94c |
| Uranium Isotopic Standard (Nominally 80% Enriched) | U-800 | Zinc Concentrates | 113a |
| Uranium Isotopic Standard (Nominally 85% Enriched) | U-850 | Zinc Concentrates | 329 |
| Uranium Isotopic Standard (Nominally 90% Enriched) | U-900 | Zinc Cyclohexanebutyrate | 1073b |
| Uranium Isotopic Standard (Nominally 93% Enriched) | U-930 | Zinc, Freezing Point | 740 |
| Uranium Isotopic Standard (Nominally 97% Enriched) | U-970 | Zinc, Freezing Point Standard | 43h |
| Uranium Metal | 960 | Zinc Metal | 683 |
| Uranium Oxide | 950b | Zinc Oxide Rubber Compound | 370e |
| Uranium Oxide | 969 | Zircaloy-2 | 360a |
| Uranium-233 Spike Assay and Isotopic Solution Standard | 995 | Zircaloy-4 Metal | 1237 |
| Uranium-235 Spike Assay and Isotopic Solution Standard | 993 | Zircaloy-4 Metal | 1238 |
| Urban Dust/Organics | 1649 | Zircaloy-4 Metal | 1239 |
| Urban Particulate Matter | 1648 | Zirconium-Barium Chromate Formulation for Heat-Source Powder Calorimetry | 1651 |
| Urea | 912a | Zirconium-Barium Chromate Formulation for Heat-Source Powder Calorimetry | 1652 |
| Urea | 2141 | Zirconium Metal | 1234 |
| Urea | 2152 | Zirconium Metal | 1235 |
| Uric Acid | 913 | Zirconium Metal | 1236 |
| Vanadium and Nickel in Residual Fuel Oil | 1618 | | |
| Vanadium in Curde Oil | 8505 | | |
| Vanadium-49 Low-Energy Photon Standard | 4266 | | |
| Waspaloy | 349 | | |
| Wear-Metals in Lubricating Oil (100 ppm) | 1084 | | |

U. S. Department of Commerce
Frederick B. Dent
Secretary
National Bureau of Standards
Richard W. Roberts, Director

Appendix II. Certificates for the Environmental Research,
Analysis, and Control Standards (listed in
numerical order).

National Bureau of Standards Certificate of Analysis Standard Reference Material 1579 Powdered Lead Based Paint

This Standard Reference Material is intended for use in the calibration of apparatus and methods used in the determination of lead in paint removed from the interior surfaces of old housing. The certified value is based on at least a 100 milligram sample of the as-received, total material.

Lead Content 11.87 ± 0.04 Weight Percent

The certified value of 11.87 percent lead is the weighted average value determined by a statistical analysis of the results of 32 determinations by atomic absorption spectrometry (average 11.84 percent lead, $s = 0.13$ percent lead), and 16 determinations by polarography (average 11.93 percent lead, $s = 0.13$ percent lead). The standard error of the weighted average is 0.02 percent lead, and the half-width of the 95 percent confidence interval is taken to include ± 0.04 percent lead by weight.

X-ray fluorescence spectrometry showed the bottle-to-bottle inhomogeneity of the material with respect to lead content to be no greater than 0.02 percent lead; no within-bottle inhomogeneity was detected.

Analyses for lead and determinations of homogeneity were carried out in the NBS Analytical Chemistry Division by the following persons:

X-ray Fluorescence: S. D. Rasberry
Atomic Absorption Spectrometry: T. C. Rains and T. A. Rush
Polarography: E. J. Maienthal

Statistical calculations were carried out by J. Mandel of the NBS Institute for Materials Research.

The overall direction and coordination of the technical measurements leading to this certificate were performed under the chairmanship of B. Greifer.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. W. Mears.

Washington, D. C. 20234
January 23, 1973

J. Paul Cali, Chief
Office of Standard Reference Materials

(over)

Preparation, Testing, and Analysis

Collection

The paint for this Standard Reference Material was collected by the staff of the Philadelphia Department of Public Health from the interior surfaces of dwellings undergoing renovation. The paint was softened with a hand torch, scraped from the plaster and wood substrates, and collected in plastic bags as a heterogeneous mixture of many different kinds of paints. In the laboratory, non-paint matter such as bits of metal, plastic, glass, and wood were removed and the paint mixture was ground in a disk mill to produce a material suitable for feeding into a jet mill. The paint was comminuted in a jet mill operating at 100 psig air pressure, then sieved through a 100-mesh vibrating screen to remove the coarse, non-grindable fraction. Two additional passes through the jet mill at 97 to 107 psig gave a fine powder with 99.31 weight percent passing through a 325 mesh sieve.

Homogeneity

Sample homogeneity was ascertained by x-ray fluorescence analysis for lead content on 17 samples chosen at random from the total lot. A statistical analysis of the data from 136 observations showed the bottle-to-bottle variability among the samples to be no greater than 0.02 percent lead. No within-bottle variation with respect to lead was detected.

Dissolution

A procedure used to dissolve the sample is summarized briefly: dry ash the weighed paint for 2 hours at 450 ° C, digest with 2:5 HCl - HNO₃ containing HF, evaporate to dryness; treat with HNO₃, evaporate to dryness; treat twice with HCl and evaporate to dryness each time. Extract the solids twice with portions of acetic acid - ammonium acetate solution, heating for several hours just below boiling. Combine the extracts and heat the mixture (including solids) for one hour, just below boiling. Cool the mixture and determine lead in solution. (The solids need not be removed for polarographic analysis.)

An alternate procedure for sample dissolution is: dry ash the weighed paint for 6 hours at 500 ° C, cool, then digest for 2 hours in 1:1 HCl - HNO₃. Separate the insoluble solids from the solution by centrifuging, and wash 3 times with 1:10 HNO₃ combining the rinsings with the principal solution. Determine lead in solution.

Details of the dissolution procedures, the analytical procedures, and results will be published in the 260 series of NBS Special Publications.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1580

Organics in Shale Oil

This SRM is intended primarily for evaluating the reliability of analytical methods for the determination of trace level organic compounds in an oil matrix, i.e., shale oil, petroleum crude oil, or coal-derived liquids.

Certified Values of Constituent Organic Compounds: The certified values for selected organic constituents are shown in Table 1. These values are based on results obtained by two independent, analytical methods (see Table 2). Non-certified values, which are given for information only, are listed in Table 3.

NOTICE AND WARNINGS TO USER

Expiration of Certification: This certification is valid, within the limits certified, for 3 years from the date of purchase. In the event that the certification should become invalid before then, purchasers will be notified by NBS.

Storage: Sealed ampoules, as received, should be stored in the dark at temperatures between 10-30 °C.

Use: Samples for analysis should be withdrawn from ampoules immediately after opening and processed without delay for any certified value in Table 1 to be valid within the stated uncertainty. Certified values are not applicable to ampoules stored after opening, even if resealed.

PREPARATION AND ANALYSIS

The shale oil for this SRM came from a 150-ton retort for *in-situ* simulated combustion of oil shale, operated by the Laramie Energy Technology Center, Laramie, Wyoming. The shale was from the Mahogany Zone of the Colorado Green River Formation. The shale oil had been supplied in November 1975 to the Oak Ridge National Laboratory (ORNL) where it underwent centrifugation to separate the oil from water and sludge. The shale oil was provided to NBS by Bruce R. Clark, ORNL, Oak Ridge, Tennessee.

At NBS, the centrifuged sample was filtered through fine filter paper and mixed in a 20-liter, Teflon-stoppered, glass bottle by rolling for 40 hours. Samples were aliquoted into 2-mL amber glass ampoules. Although not intended to be representative of all shale oils, SRM 1580 provides a typical specimen of this matrix for use in developing analytical methods.

Randomly selected ampoules were analyzed. Each analyst examined at least six ampoules, sometimes measuring replicates from one ampoule. No trend was found in measured values with the ampouling sequence.

Two independent techniques were employed for the determination of the certified values for the organic constituents. Three different methods of sample preparation were used prior to analysis: simple dilution of the shale oil with methylene chloride (or other suitable solvent); acid/base extraction to isolate acidic, basic, and neutral components; and a high performance liquid chromatographic fractionation. The following techniques were employed for the final quantitative analysis: gas chromatography (GC), gas chromatography/mass-spectrometry (GC/MS) with single ion monitoring for selective detection, and high performance liquid chromatography (HPLC) with selective fluorescence detection. All GC/MS analyses used the standard addition method for quantitation. The GC and HPLC analyses employed either internal standard, external standard, or standard addition methods. The analytical methods and the corresponding values are summarized in Table 2.

Consultation on the statistical design of the experimental work was provided by K. R. Eberhardt of the Statistical Engineering Division.

The coordination of the technical measurements leading to certification were performed under the direction of H. S. Hertz, S. N. Chesler, L. R. Hilpert, W. E. May, and S. A. Wise.

The technical and support aspects involved in preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Washington, D.C. 20234
November 24, 1980
(Revision of Certificate
dated 3-10-80)

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

The following members of the staff of the Center for Analytical Chemistry, Organic Analytical Research Division, performed the analytical determinations.

1. J. M. Brown-Thomas
2. S. N. Chesler
3. F. R. Guenther
4. L. R. Hilpert
5. P. L. Konash
6. W. E. May
7. R. M. Parris
8. K. L. Richie

TABLE 1. Certified Values of Organic Constituents

| <u>Compound</u> | <u>Concentration ($\mu\text{g/g}^a$)</u> |
|--|---|
| Fluoranthene | 54 \pm 10 |
| Pyrene | 104 \pm 18 |
| Benzo[<i>a</i>]pyrene | 21 \pm 6 |
| Benzo[<i>e</i>]pyrene | 18 \pm 8 |
| Perylene | 3.4 \pm 2.2 |
| Phenol | 407 \pm 50 |
| <i>o</i> -Cresol | 385 \pm 50 |
| 2,6-Dimethylphenol | 175 \pm 30 |
| Benzo[<i>f</i>]quinoline (5,6-Benzoquinoline) | 16 \pm 4 |

^aThe estimated uncertainty listed for a constituent is based on judgment and represents an evaluation of the combined effects of method imprecision, possible systematic errors among methods, and material inhomogeneity. The estimated uncertainty is intended to correspond to approximately 95% confidence limits.

TABLE 2. Summary of Results by the Analytical Methods Used in Certification

| <u>Compound</u> | <u>Concentration ($\mu\text{g/g}^a$)</u> | <u>Number of Ampoules Analyzed</u> | <u>Sample Preparation Technique</u> | <u>Analytical Technique</u> |
|--|---|------------------------------------|-------------------------------------|-----------------------------|
| Fluoranthene | 55 \pm 5 | 6 | Direct Injection | GC/MS |
| | 53 \pm 2 | 9 | HPLC | HPLC |
| Pyrene | 101 \pm 5 | 6 | Direct Injection | GC/MS |
| | 107 \pm 8 | 10 | HPLC | HPLC |
| Benzo[<i>a</i>]pyrene | 20 \pm 1 | 6 | Direct Injection | GC/MS |
| | 23 \pm 1 | 8 | HPLC | HPLC |
| Benzo[<i>e</i>]pyrene | 17 \pm 1 | 6 | Direct Injection | GC/MS |
| | 20 \pm 3 | 8 | HPLC | HPLC |
| Perylene | 2.8 \pm 0.6 | 5 | Direct Injection | GC/MS |
| | 3.9 \pm 0.6 | 11 | HPLC | HPLC |
| Phenol | 412 \pm 35 | 8 | HPLC | GC/MS |
| | 402 \pm 4 | 8 | Acid/Base Extraction | GC |
| <i>o</i> -Cresol | 386 \pm 42 | 8 | HPLC | GC/MS |
| | 384 \pm 9 | 8 | Acid/Base Extraction | GC |
| 2,6-Dimethylphenol | 183 \pm 23 | 9 | HPLC | GC/MS |
| | 168 \pm 8 | 8 | Acid/Base Extraction | GC |
| Benzo[<i>f</i>]quinoline (5,6-Benzoquinoline) | 16 \pm 1 | 7 | HPLC | HPLC |
| | 15 \pm 1 | 8 | Acid/Base Extraction | Multi-dimensional GC |

^aUncertainty is the standard deviation of a single measurement.

TABLE 3. Non-Certified Values of Organic Compounds in Shale Oil

NOTE: The values shown in this table are not certified because they are not based on the results of two independent methods. These values are included for information only.

| <u>Compound</u> | <u>Concentration ($\mu\text{g/g}$)</u> |
|-----------------------|---|
| <i>p</i> -Cresol | (270) ^a |
| <i>m</i> -Cresol | (330) ^a |
| 2,5-Dimethylphenol | (320) ^a |
| 2,4-Dimethylphenol | (380) ^a |
| 2,5,6-Trimethylphenol | (360) ^a |
| 2,4,6-Trimethylphenol | (120) ^a |
| Phenanthridine | (45) ^b |

^aAcid/base extraction - GC analysis
^bHPLC extraction - HPLC analysis

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1620a

Sulfur in Residual Fuel Oil

Sulfur Concentration 4.504 ± 0.010 weight percent

This Standard Reference Material is intended for use as an analytical standard in the determination of total sulfur in fuel oils or materials of similar matrices. SRM 1620a is a commercial "No. 5 Heavy" residual fuel oil as defined by American Society for Testing and Materials, ASTM.

Sulfur was certified using three independent methods of analysis: gravimetry, ion chromatography, and x-ray fluorescence.

The standard error of the certified value includes observed variability within and between measurement methods and any observed material heterogeneity.

NOTICE AND RECOMMENDED USE: Due to the high sulfur content of SRM 1620a, it is recommended that the bottle be shaken vigorously before sampling. Homogeneity and stability testing at NBS indicates that the best results are achieved when the material is shaken before use.

Analyses for certification were performed by W. F. Koch and E. R. Deardorff of the Inorganic Analytical Research Division and P. A. Pella of the Gas and Particulate Science Division.

The statistical analysis of the certification data was performed by R. C. Paule of the National Measurement Laboratory.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
December 22, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

SRM 1620a was also tested and found to exhibit the physical properties shown in Table 1. In addition, semi-quantitative values obtained by emission spectrometry are given in Table 2. These values are not certified, but supplied for information only.

Table 1
Physical Properties for SRM 1620a

| Flash Point ^a °C | Kinematic Viscosity ^b 50 °C (cSt) | Pour Point ^c °C | Density @ 20 °C ^d g/cm ³ |
|--------------------------------|---|-------------------------------|---|
| 70 | 47.75 | 2 | 1.096 |

These measurements were performed by S. Weeks, Materials Chemistry Division, Center for Material Science.

Methods Used for Physical Tests

- a. ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- b. ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- c. ASTM D97-66 (1978) Pour Point of Petroleum Oils
- d. ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

Table 2
Semi-Quantitative Emission Spectrometry
Analysis for SRM 1620a

| Element | µg/mL | Element | µg/mL |
|---------|-------|---------|-------|
| Al | 20 | Mo | <1 |
| B | <1 | Na | 31 |
| Ca | 9 | Ni | <1 |
| Cr | <1 | Si | 13 |
| Cu | <1 | Sn | <1 |
| Fe | <5 | Ti | <1 |
| Mg | <1 | V | <1 |
| Mn | <1 | Zn | 23 |

Note: SRM 1620a was analyzed using the rotating disc method. This method is based on absolute amounts of sample since no internal standard is used to correct for the amount of sample actually analyzed. Differences in actual values may range from factors of 1-3.

These measurements were performed by J. A. Norris, Inorganic Analytical Research Division, Center for Analytical Chemistry.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1621b

Sulfur in Residual Fuel Oil

Sulfur Concentration 0.950 ± 0.005 weight percent

This Standard Reference Material is intended for use as an analytical standard in the determination of total sulfur in fuel oils or materials of similar matrices. SRM 1621b is a commercial "No. 6" residual fuel oil as defined by American Society for Testing and Materials, ASTM.

Sulfur was certified using three independent methods of analysis: gravimetry, ion chromatography, and x-ray fluorescence.

The standard error of the certified value includes observed variability within and between measurement methods and any observed material heterogeneity.

NOTICE: The certification of SRM 1621b is valid for 3 years from date of purchase.

Analyses for certification were performed by W. F. Koch and E. R. Deardorff of the Inorganic Analytical Research Division and P. A. Pella of the Gas and Particulate Science Division.

The statistical analysis of the certification data was performed by R. C. Paule of the National Measurement Laboratory.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
December 22, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

SRM 1621b was also tested and found to exhibit the physical properties shown in Table 1. In addition, semi-quantitative values obtained by emission spectrometry are given in Table 2. These values are not certified, but supplied for information only.

Table 1
Physical Properties for SRM 1621b

| Flash Point ^a °C | Kinematic Viscosity ^b 50 °C (cSt) | Pour Point ^c °C | Density @ 20 °C ^d g/cm ³ |
|--------------------------------|---|-------------------------------|---|
| 111 | 89.2 | 11 | 0.929 |

These measurements were performed by S. Weeks, Materials Chemistry Division, Center for Material Science.

Methods Used for Physical Tests

- a. ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- b. ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- c. ASTM D97-66 (1978) Pour Point of Petroleum Oils
- d. ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

Table 2
Semi-Quantitative Emission Spectrometry
Analysis for SRM 1621b

| Element | µg/mL | Element | µg/mL |
|---------|-------|---------|-------|
| Al | 6 | Mo | <1 |
| B | <1 | Na | 8 |
| Ca | 9 | Ni | 6 |
| Cr | 3 | Si | 6 |
| Cu | <1 | Sn | <1 |
| Fe | <5 | Ti | <1 |
| Mg | <1 | V | 15 |
| Mn | 1 | Zn | 15 |

Note: SRM 1621b was analyzed using the rotating disc method. This method is based on absolute amounts of sample since no internal standard is used to correct for the amount of sample actually analyzed. Differences in actual values may range from factors of 1-3.

These measurements were performed by J. A. Norris, Inorganic Analytical Research Division, Center for Analytical Chemistry.



National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1622b

Sulfur in Residual Fuel Oil

Sulfur Concentration 1.982 ± 0.018 weight percent

This Standard Reference Material is intended for use as an analytical standard in the determination of total sulfur in fuel oils or materials of similar matrices. SRM 1622b is a commercial "No. 6" residual fuel oil as defined by American Society for Testing and Materials, ASTM.

Sulfur was certified using three independent methods of analysis: gravimetry, ion chromatography, and x-ray fluorescence.

The standard error of the certified value includes observed variability within and between measurement methods and any observed material heterogeneity.

NOTICE: The certification of SRM 1622b is valid for 3 years from date of purchase.

Analyses for certification were performed by W. F. Koch and E. R. Deardorff of the Inorganic Analytical Research Division and P. A. Pella of the Gas and Particulate Science Division.

The statistical analysis of the certification data was performed by R. C. Paule of the National Measurement Laboratory.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
December 22, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

SRM 1622b was also tested and found to exhibit the physical properties shown in Table 1. In addition, semi-quantitative values obtained by emission spectrometry are given in Table 2. These values are not certified, but supplied for information only.

Table 1
Physical Properties for SRM 1622b

| Flash Point ^a °C | Kinematic Viscosity ^b 50 °C (cSt) | Pour Point ^c °C | Density @ 20 °C ^d g/cm ³ |
|--------------------------------|---|-------------------------------|---|
| 65 | 377.34 | -7 | 0.984 |

These measurements were performed by S. Weeks, Materials Chemistry Division, Center for Material Science.

Methods Used for Physical Tests

- a. ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- b. ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- c. ASTM D97-66 (1978) Pour Point of Petroleum Oils
- d. ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

Table 2
Semi-Quantitative Emission Spectrometry
Analysis for SRM 1622b

| Element | µg/mL | Element | µg/mL |
|---------|-------|---------|-------|
| Al | 8 | Mo | <1 |
| B | <1 | Na | 25 |
| Ca | 24 | Ni | 15 |
| Cr | 1 | Si | 13 |
| Cu | <1 | Sn | <1 |
| Fe | <5 | Ti | <1 |
| Mg | 2 | V | 50 |
| Mn | 1 | Zn | 11 |

Note: SRM 1622b was analyzed using the rotating disc method. This method is based on absolute amounts of sample since no internal standard is used to correct for the amount of sample actually analyzed. Differences in actual values may range from factors of 1-3.

These measurements were performed J. A. Norris, Inorganic Analytical Research Division, Center for Analytical Chemistry

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1623a

Sulfur in Residual Fuel Oil

Sulfur Concentration.0.240 ± 0.003 weight percent

This Standard Reference Material is intended for use as an analytical standard in the determination of total sulfur in fuel oils or materials of similar matrices. SRM 1623a is a commercial "No. 5 Heavy" residual fuel oil as defined by American Society for Testing and Materials, ASTM.

Sulfur was certified using three independent methods of analysis: gravimetry, ion chromatography, and x-ray fluorescence.

The standard error of the certified value includes observed variability within and between measurement methods and any observed material heterogeneity.

NOTICE: The certification of SRM 1623a is valid for 3 years from date of purchase.

Analyses for certification were performed by W. F. Koch and E. R. Deardorff of the Inorganic Analytical Research Division and P. A. Pella of the Gas and Particulate Science Division.

The statistical analysis of the certification data was performed by R. C. Paule of the National Measurement Laboratory.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
December 22, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

SRM 1623a was also tested and found to exhibit the physical properties shown in Table 1. In addition, semi-quantitative values obtained by emission spectrometry are given in Table 2. These values are not certified, but supplied for information only.

Table 1
Physical Properties for SRM 1623a

| Flash Point ^d °C | Kinematic Viscosity ^b 50 °C (cSt) | Pour Point ^c °C | Density @ 20 °C ^d g/cm ³ |
|--------------------------------|---|-------------------------------|---|
| 140 | 53.82 | 17 | 0.918 |

These measurements were performed by S. Weeks, Materials Chemistry Division, Center for Material Science.

Methods Used for Physical Tests

- a. ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- b. ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- c. ASTM D97-66 (1978) Pour Point of Petroleum Oils
- d. ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

Table 2
Semi-Quantitative Emission Spectrometry
Analysis for SRM 1623a

| Element | µg/mL | Element | µg/mL |
|---------|-------|---------|-------|
| Al | 5 | Mo | <1 |
| B | <1 | Na | 9 |
| Ca | 9 | Ni | 1 |
| Cr | 1 | Si | <1 |
| Cu | <1 | Sn | <1 |
| Fe | <5 | Ti | <1 |
| Mg | <1 | V | 3 |
| Mn | <1 | Zn | 15 |

Note: SRM 1623a was analyzed using the rotating disc method. This method is based on absolute amounts of sample since no internal standard is used to correct for the amount of sample actually analyzed. Differences in actual values may range from factors of 1-3.

These measurements were performed by J. A. Norris, Inorganic Analytical Research Division, Center for Analytical Chemistry.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1624a

Sulfur in Distillate (Diesel) Fuel Oil

Sulfur Concentration 0.141 ± 0.002 weight percent

This Standard Reference Material is intended for use as an analytical standard in the determination of total sulfur in fuel oils or materials of similar matrices. SRM 1624a is a commercial "No. 2-D" distillate fuel oil as defined by American Society for Testing and Materials, ASTM.

Sulfur was certified using two independent methods of analysis: gravimetry and ion chromatography.

The standard error of the certified value includes observed variability within and between measurement methods and any observed material heterogeneity.

NOTICE: The certification of SRM 1624a is valid for 3 years from date of purchase.

Analyses for certification were performed by W. F. Koch and E. R. Deardorff of the Inorganic Analytical Research Division and P. A. Pella of the Gas and Particulate Science Division.

The statistical analysis of the certification data was performed by R. C. Paule of the National Measurement Laboratory.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
December 22, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

SRM 1624a was also tested and found to exhibit the physical properties shown in Table 1. In addition, semi-quantitative values obtained by emission spectrometry are given in Table 2. These values are not certified, but supplied for information only.

Table 1
Physical Properties for SRM 1624a

| Flash Point ^a °C | Kinematic Viscosity ^b 40 °C (cSt) | Cloud Point ^c °C | Density @ 20 °C ^d g/cm ³ |
|--------------------------------|---|--------------------------------|---|
| 53 | 2.57 | -14 | 0.848 |

These measurements were performed by S. Weeks, Materials Chemistry Division, Center for Material Science.

Methods Used for Physical Tests

- a. ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- b. ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- c. ASTM D2500-66 (1976) Cloud Point of Petroleum Oils
- d. ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

Table 2
Semi-Quantitative Emission Spectrometry
Analysis for SRM 1624a

| Element | µg/mL | Element | µg/mL |
|---------|-------|---------|-------|
| Al | 1 | Mo | <1 |
| B | <1 | Na | <1 |
| Ca | 7 | Ni | <1 |
| Cr | <1 | Si | <1 |
| Cu | <1 | Sn | <1 |
| Fe | <5 | Ti | <1 |
| Mg | <1 | V | <1 |
| Mn | <1 | Zn | <1 |

Note: SRM 1624a was analyzed using the rotating disc method. This method is based on absolute amounts of sample since no internal standard is used to correct for the amount of sample actually analyzed. Differences in actual values may range from factors of 1-3.

These measurements were performed by J. A. Norris, Inorganic Analytical Research Division, Center for Analytical Chemistry.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1630

Trace Mercury in Coal

This Standard Reference Material is intended as an analytical standard for the determination of trace mercury in coal. The material is a commercially available coal that was crushed to a size of 210 to 500 micrometers with a roll crusher. From a total of 500 packaged bottles, 30 were randomly selected for analysis. Duplicate determinations were made on 0.5 g portions of 25 of these bottles, and single determinations were made on the other five. The mercury content of this material was obtained by destructive neutron activation analysis.

The recommended value is the average of these 55 determinations on 30 bottles, which was found to be:

$$\text{Mercury content} = 0.13 \mu\text{g/g}$$

The recommended value is not expected to change by more than ± 1 in the last significant figure.

A study of homogeneity showed no variability among bottles that could not be accounted for by analytical error. Duplicate samples from the same bottle indicated a homogeneity for mercury of $\pm 5\%$ (relative).

The mercury content was also determined by flameless atomic absorption spectrometry, yielding an average value of $0.14 \mu\text{g/g}$.

Selenium was also determined using destructive neutron activation analysis. The value obtained, which is not certified but included for information only, was found to be $2.1 \mu\text{g/g}$.

The homogeneity testing and analyses for certification were performed in the NBS Analytical Chemistry Division by T. E. Gills and H. Rook under the direction of P. D. LaFleur.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by C. L. Stanley.

Washington, D.C. 20234
August 1, 1979
(Revision of Certificate dated 11-2-71
Editorial Revision only.)

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

ANALYTICAL PROCEDURE

The bottles containing the samples were allowed to remain open at room temperature (about 25 °C) for twenty-four hours.

The coal samples, along with solution standards of mercury and NBS Standard Reference Material 1571 (Orchard Leaves) used as a control, were encapsulated in cleaned quartz vials. The geometry of both the samples and the standards were optimized so that flux monitors were not needed. The samples were irradiated for four hours at a thermal flux of $6 \times 10^{13} \text{ n}\cdot\text{cm}^{-2} \text{ sec}^{-1}$. The samples were allowed to decay for three days to minimize the personnel dose rate. The samples were postweighed into porcelain boats and burned in a combustion tube. The volatile mercury compounds and other volatile products liberated during burning were trapped in a liquid nitrogen cold trap. The cold trap was allowed to warm to room temperature. The mercury compounds were then transferred to polyethylene bottles by washing the cold trap with concentrated nitric acid and water. For this analysis, ^{197}Hg produced by $^{196}\text{Hg}(n,\gamma) ^{197}\text{Hg}$ was used as the measuring activity.

Bromine-82, an interfering isotope, was separated from the sample by using the classical silver bromide precipitation.

The samples were counted on a 22 cm^3 Ge(Li) detector connected to a 2048-multichannel analyzer. The accumulated data was processed by computer for peak identification and integration. The concentrations were determined by using a Standard Comparator Method.

NOTE TO USER

It is suggested that persons using SRM 1630 to check their analytical technique should adopt the following criteria. If the average, \bar{X} , of N replicate measurements on this SRM is found to lie in the interval—

$$0.127 - \frac{0.013}{\sqrt{N}} < \bar{X} < 0.127 + \frac{0.013}{\sqrt{N}}$$

then the analytical technique used gives a result compatible with that found at NBS. However, if the value \bar{X} lies outside this interval, then the technique should be examined for possible bias or miscalibration.

NOTE: The above expression is not rigorously correct. It does not include a possible component for between laboratory variability nor sources of systematic error.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1632b

Trace Elements in Coal

(Bituminous)

This Standard Reference Material (SRM) is intended for use in the calibration of apparatus and the evaluation of techniques employed in the analysis of coal or similar materials. SRM 1632b is a bituminous coal with a nominal sulfur content of 1.9%. It is in the form of a fine powder (-60 mesh).

Certified Values of Constituent Elements: The certified values for the constituent elements are given in Table 1. The certified values are based on measurements using proven techniques and methods. Noncertified values are given in Table 2 and are provided for information only. These values are based on measurements made using a single technique or method. While no reason exists to suspect systematic bias in the information values, no attempt was made to determine if such a bias exists that is attributable to the technique and/or method used. A list of analytical techniques and methods used for the different analyses is given in Table 3. As part of its update certification program, NBS will periodically update many of these values to certification status.

Expiration of Certification: The certification of SRM 1632b will be valid up to 5 years from the purchase date. Should any of the certified constituents become invalid prior to that date, purchasers will be notified by NBS.

Use: This material should be vacuum dried at ambient temperature for 24 hours prior to use. The certified concentrations are reported on a "dry-weight" basis, thus the concentration determined on undried samples should be adjusted for the moisture content of the sample. Typical moisture loss using the drying procedure stated above is 1.3%.

A minimum sample size of 250 mg of the dried material is required for the certified values to be valid.

This SRM should be kept in its original bottle. It should not be exposed to intense source of radiation, including ultraviolet lamps or sunlight.

The statistical analysis of the certification data was performed by R.C. Paule of the National Measurement Laboratory.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T.E. Gills.

Gaithersburg, MD 20899
June 20, 1985

Stanley D. Rasberry, Chief
Office of Standard Reference Materials

(over)

Source and Preparation of Material: The coal for this SRM was obtained from the Humphrey No. 7 mine and coal preparation plant of the Consolidation Coal Company, Christopher Coal Company Division, Osage, West Virginia. This mine produces bituminous coal with a sulfur content of 1.8- 1.9 percent (dry basis). This coal was obtained from an underground mine that recovers coal from the Pittsburgh seam, which is considered the single most valuable and extensive coal seam in the United States.

Approximately 900 kg of the coal for SRM 1632b was oven dried prior to processing, in accordance with procedures outlined in ASTM D2013. The coal was reduced in size to -60 mesh and sieved prior to blending. The coal was then blended in a stainless steel cone blender (approximate capacity 0.85 cubic meter). After blending the coal was packaged in polyethylene-lined aluminum cans and was subsequently repackaged in fifty gram units.

Analysis

Major, Minor, and Trace Constituents: In general, the major, minor, and trace constituents were certified using two or more independent methods of analysis or two or more different laboratories. For those constituents that were determined using a single method, technique, or laboratory, the values are given for information only. (See Table 3).

Calorific Value: The calorific value was determined using measurements made in an isoperibol calorimeter, an isothermal calorimeter, and an adiabatic calorimeter at two different laboratories.

Moisture, Ash, and Volatile Matter: The moisture, ash, and volatile matter values were determined on measurements made using the standard ASTM methods, D3173, D3174, and D3175, respectively. In addition, commercial instruments commonly used for the determination of the parameters provided additional values.

Table 1. Certified Values of Constituent Elements

| <u>Major Constituents</u> | | <u>Minor Constituents</u> | |
|---------------------------|--------------------------------------|---------------------------|--------------------------------------|
| <u>Elements</u> | <u>Content</u> <u>Wt. Percent</u> | <u>Elements</u> | <u>Content</u> <u>Wt. Percent</u> |
| Carbon (Total) | 78.11 ± 0.37 ^a | Aluminum | 0.855 ± 0.019 |
| Hydrogen | 5.07 ± 0.06 | Calcium | 0.204 ± 0.006 |
| Nitrogen | 1.56 ± 0.07 | Iron | 0.759 ± 0.045 |
| Sulfur | 1.89 ± 0.06 | Magnesium | 0.0383 ± 0.0008 |
| Volatile matter | 35.4 ± 1.1 | Potassium | 0.0748 ± 0.0028 |
| | | Sodium | 0.0515 ± 0.0011 |
| | | Titanium | 0.0454 ± 0.0017 |

Trace Constituents

| <u>Element</u> | <u>Content</u> <u>µg/g</u> | <u>Element</u> | <u>Content</u> <u>µg/g</u> |
|----------------|-------------------------------|----------------|-------------------------------|
| Arsenic | 3.72 ± 0.09 | Manganese | 12.4 ± 1.0 |
| Barium | 67.5 ± 2.1 | Nickel | 6.10 ± 0.27 |
| Cadmium | 0.0573 ± 0.0027 | Rubidium | 5.05 ± 0.11 |
| Cobalt | 2.29 ± 0.17 | Selenium | 1.29 ± 0.11 |
| Copper | 6.28 ± 0.30 | Thorium | 1.342 ± 0.036 |
| Lead | 3.67 ± 0.26 | Uranium | 0.436 ± 0.012 |
| | | Zinc | 11.89 ± 0.78 |

Calorific Value^{b,c}

14005 ± 35 Btu/lb (32.57 ± 0.08 MJ kg⁻¹)

Ash, wt. %

6.79 ± 0.16

^aThe listed ± uncertainties for carbon, hydrogen, volatile matter, and calorific value are two standard deviations of the certified value. The listed ± uncertainties for all other constituents are two standard deviations for the certified values and include an allowance for minor sample heterogeneity. The observed sample variability was generally less than two percent of the constituent value.

^bThe calorific value (MJ kg⁻¹) may decrease upon aging or normal oxidation of the coals. NBS will continue to monitor this value and report any substantive change in the certified calorific value to the purchaser. The reference date for the calorific value is May 1985.

^cThe calorific value is determined as HHV2 (Higher Heating Value-Moisture Free).

Table 2. Noncertified Values for Constituent Elements

| <u>Trace Constituents</u> | | | |
|---------------------------|-------------------------------|----------------|-------------------------------|
| <u>Element</u> | <u>Content</u> <u>µg/g</u> | <u>Element</u> | <u>Content</u> <u>µg/g</u> |
| Antimony | (0.24) | Lithium | (10) |
| Bromine | (17) | Molybdenum | (0.9) |
| Cerium | (9) | Samarium | (0.87) |
| Cesium | (0.44) | Scandium | (1.9) |
| Chlorine | (1260) | Silicon, wt % | (1.4) |
| Chromium | (11) | Strontium | (102) |
| Europium | (0.17) | Tungsten | (0.48) |
| Hafnium | (0.43) | Vanadium | (14) |
| Lanthanum | (5.1) | | |

Table 3. Analytical Techniques and Methods Used for the Characterization of SRM 1632b

| Method/ Element | A | B | C | D | E | F | G | H | I | J | K | L | M |
|--------------------|---|---|---|---|---|---|-----|---|---|---|---|---|---|
| Al | | | • | • | | | | | | | | • | |
| As | | | • | | • | | | | | | | | |
| Ash Content | | | | | | | • 2 | | • | | | | |
| Ba | | | • | | | | | | | | | | |
| Br | | | • | | | | | | | | | | |
| C (Total) | | | | | | | • 5 | • | • | | • | | |
| Ca | | • | • | • | | | | | | | | • | |
| Cal Val | | | | | | | | | • | • | | | |
| Cd | • | • | | | | | | | | | | | |
| Ce | | | • | | | | | | | | | | |
| Cl | | | • | | | | | | | | | | |
| Co | | | • | • | | | | | | | | | |
| Cr | | | • | | | | | | | | | | • |
| Cs | | | • | | | | | | | | | | |
| Cu | | | | | • | | | | | | | | • |
| Eu | | | • | | | | | | | | | | |
| Fe | • | | • | | | | | | | | | • | |
| H | | | | | | | • 5 | • | • | | | | |
| Hf | | | • | | | | | | | | | | |
| K | | • | • | • | | | | | | | | • | |
| La | | | • | | | | | | | | | | |
| Li | | | | • | | | | | | | | | |
| Mg | • | • | • | | | | | | | | | | |
| Mn | | | • | • | | | | | | | | | |
| Mo | | | • | | | | | | | | | | |
| N | | | | | | | • 6 | | | | | | |
| Na | | | • | • | | | | | | | | | |
| Ni | • | | | | | | | | | | | | • |
| Pb | • | • | | | | | | | | | | | |
| Rb | | • | • | • | | | | | | | | | |
| S | | | | | | • | • 4 | | • | | | • | |
| Sb | | | • | | | | | | | | | | |
| Sc | | | • | | | | | | | | | | |
| Se | | | • | | • | | | | | | | | |
| Si | | | • | | | | | | | | | • | |
| Sm | | | • | | | | | | | | | | |
| Sr | | | • | | | | | | | | | | |
| Th | | • | • | | | | | | | | | | |
| Ti | | | • | • | | | | | | | | • | • |
| U | | • | • | | | | | | | | | | |
| V | | | • | | | | | | | | | | • |
| Volatile Matter | | | | | | | • 3 | | • | | | | |
| W | | | • | | | | | | | | | | |
| Zn | • | • | • | | | | | | | | | | |

Analytical Methods

- A. Atomic absorption spectrometry
- B. Isotope dilution mass spectrometry
- C. Instrumental neutron activation analysis
- D. Flame emission spectrometry
- E. Flameless atomic absorption spectrometry
- F. Ion chromatography
- G. ASTM Methods: (1)D3173, (2)D3174, (3)D3175, (4)D3177, (5)D3178, (6)D3179
- H. Combustion coulometry
- I. Commercial coal analyzers: moisture, ash, sulfur, Btu, volatile matter, carbon, hydrogen, nitrogen
- J. Commercial calorimeter
- K. Gas chromatography
- L. X-ray fluorescence
- M. Inductively coupled plasma emission spectrometry

Analysts

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National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1633a

Trace Elements in Coal Fly Ash

This Standard Reference Material (SRM) is intended for use in the evaluation of analytical methods for the determination of constituent elements in coal fly ash or materials with a similar matrix.

SRM 1633a is a fly ash that was sieved through a No. 170 sieve with a nominal sieve opening of 90 μm .

Certified Values of Constituent Elements: The certified values for the constituent elements are shown in Table 1. The analytical techniques used and the analysts are given in Table 3. The certified values are based on results obtained by reference methods of known accuracy or from two or more independent, reliable analytical methods. Noncertified values are given for information only in Table 2.

Notice and Warnings to Users: This certification is invalid 5 years from date of purchase of the SRM. The constituents certified or analyzed are reviewed periodically and may be updated to reflect improved measurement. Updated certificates will be made available upon request.

Use: This material should be dried to a constant weight before using. Recommended procedures for drying are: (1) Vacuum drying for 24 hours at ambient temperature using a cold trap at or below -50°C and a pressure not greater than 30 Pa (0.2 mm Hg); (2) drying for 2 hours in an oven at 105°C ; (3) drying in a desiccator over P_2O_5 or Mg_2ClO_4 . Samples of the dried material weighing at least 250-mg should be used for analysis. When not in use the material should be kept in a tightly sealed bottle.

Source and Preparation of Material: The fly ash material was supplied by a coal fired power plant and is a product of Pennsylvania and West Virginia coals. It was selected as a typical fly ash and is not intended as a fly ash from a specific coal or combustion process. The material was sieved and blended for 2 hours in a Vee blender. The material was then removed and placed in a series of bulk containers from which specific samples were taken for homogeneity testing and certification analysis. Twelve bottles were selected for the homogeneity test. Samples from each bottle were analyzed for cobalt, chromium, europium, iron, scandium, and thorium using nondestructive neutron activation analysis. The observed standard deviations for both 50 and 250 mg sample sizes were consistent with counting statistics, indicating that the fly ash is homogeneous within $\pm 5\%$ (relative) based on these elements. The homogeneity testing and certification analyses were performed in the NBS Center for Analytical Chemistry.

The overall direction and coordination of the analytical measurements leading to the initial certification were performed in the Center for Analytical Chemistry under the chairmanship of L.A. Machlan.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by W.P. Reed and T.E. Gills.

Gaithersburg, MD 20899
January 5, 1985
(Revision of certificate
dated April 18, 1979)

Stanley D. Rasberry, Chief
Office of Standard Reference Materials

(over)

Table 1. Certified Values of Constituent Elements

| <u>Major Constituents</u> | <u>Content Wt. Percent</u> | <u>Minor Constituents</u> | <u>Content Wt. Percent</u> |
|-------------------------------|--------------------------------|-------------------------------|--------------------------------|
| Aluminum | 14.3 ± 1.0 ^a | Magnesium | 0.455 ± 0.010 |
| Iron | 9.4 ± 0.1 | Sodium | 0.17 ± 0.01 |
| Potassium | 1.88 ± 0.06 | | |
| Silicon | 22.8 ± 0.8 | | |
| Calcium | 1.11 ± 0.01 | | |

Trace Constituents

| <u>Element</u> | <u>Content µg/g</u> | <u>Element</u> | <u>Content µg/g</u> |
|----------------|---------------------|----------------|---------------------|
| Antimony | 6.8 ± 0.4 | Rubidium | 131 ± 2 |
| Arsenic | 145 ± 15 | Selenium | 10.3 ± 0.6 |
| Cadmium | 1.00 ± 0.15 | Strontium | 830 ± 30 |
| Chromium | 196 ± 6 | Thorium | 24.7 ± 0.3 |
| Copper | 118 ± 3 | Thallium | 5.7 ± 0.2 |
| Manganese | 179 ± 8 | Uranium | 10.2 ± 0.1 |
| Mercury | 0.16 ± 0.01 | Vanadium | 297 ± 6 |
| Nickel | 127 ± 4 | Zinc | 220 ± 10 |
| Lead | 72.4 ± 0.4 | | |

^aThe uncertainties of the certified values are based on judgment and represent an evaluation of the combined effects of method imprecision, possible systematic errors among methods, and material variability for samples of 250-mg or more. (No attempt was made to derive exact statistical measures of imprecision because several methods were involved in the determination of most constituents).

Supplemental Information

Note: The following values are not certified because they are not based on the results of either a reference method or of two or more independent methods. These values are included for information only.

Table 2. Noncertified Values for Constituent Elements

| <u>Element</u> | <u>Content Wt. Percent</u> | <u>Element</u> | <u>Content µg/g</u> |
|----------------|--------------------------------|----------------|-------------------------|
| Barium | 0.15 | Beryllium | 12 |
| Titanium | 0.8 | Cerium | 180 |
| Sulfur | 0.18 | Cobalt | 46 |
| | | Cesium | 11 |
| | | Europium | 4 |
| | | Gallium | 58 |
| | | Hafnium | 8 |
| | | Molybdenum | 29 |
| | | Scandium | 40 |

Table 3. Analytical Methods Used for Certified Constituent Elements

| Method/ Element | A | B | C | D | E | F | G | H | I |
|--------------------|---|---|---|---|---|---|---|---|---|
| Aluminum | • | | • | | | | | | • |
| Antimony | | | • | | | | • | | |
| Arsenic | • | | • | | | | | | |
| Cadmium | | • | • | • | | | • | | |
| Calcium | • | • | | | • | | | | |
| Chromium | • | • | • | | | | | | |
| Copper | • | • | • | | | | | | |
| Iron | • | • | • | | | | | | |
| Lead | | • | | • | • | | | | |
| Magnesium | • | • | | | | | | | |
| Manganese | • | | • | | | | | | • |
| Mercury | • | | • | | | | | | |
| Nickel | • | • | | • | • | | | | |
| Potassium | • | • | | | • | | | | |
| Rubidium | • | • | • | | • | | | | |
| Selenium | • | | • | | | | • | | |
| Silicon | | | | | • | | | • | |
| Sodium | • | | • | | | | | | |
| Strontium | • | | | | • | • | | | |
| Thallium | | • | | | | | • | | |
| Thorium | | • | • | | | | | | |
| Uranium | | • | | | | | | | |
| Vanadium | • | • | • | | | | | | |
| Zinc | • | • | | • | • | • | | | |

Analytical Methods

- A. Atomic Absorption Spectrometry or Flame Emission Spectrometry
- B. Isotope Dilution Mass Spectrometry
- C. Neutron Activation Analysis
- D. Polarography
- E. X-ray Fluorescence Spectrometry
- F. Inductively-Coupled Plasma Emission Spectrometry
- G. Isotope Dilution Spark Source Mass Spectrometry
- H. Gravimetry
- I. Direct Coupled Plasma Emission Spectrometry

Analysts

NBS Center for Analytical Chemistry

- | | |
|----------------------|------------------------|
| 1. J. B. Baldwin | 16. W. R. Kelly |
| 2. T. J. Brady | 17. H. M. Kingston |
| 3. E. R. Deardorff | 18. E. C. Kuehner |
| 4. M. G. Dias | 19. R. M. Lindstrom |
| 5. L. J. Powell | 20. L. A. Machlan |
| 6. M. S. Epstein | 21. E. J. Maienthal |
| 7. R. F. Fleming | 22. J. S. Maples |
| 8. E. L. Garner | 23. J. D. Messman |
| 9. T. E. Gills | 24. L. J. Moore |
| 10. C. A. Grabnegger | 25. P. J. Paulsen |
| 11. J. W. Gramlich | 26. P. A. Pella |
| 12. R. R. Greenberg | 27. T. C. Rains |
| 13. S. Hanamura | 28. K. J. R. Rosman |
| 14. S. H. Harrison | 29. T. A. Rush |
| 15. E. G. Heald | 30. P. A. Sleeth |
| | 31. R. L. Watters, Jr. |

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1634a

Trace Elements in Fuel Oil

This Standard Reference Material is intended for use in the evaluation of methods and the calibration of apparatus used in the analysis of fuel oils and other materials with similar matrices for trace elements. SRM 1634a is a commercial "No. 6" residual fuel oil as defined by the American Society for Testing and Materials (ASTM). This SRM was certified using two or more independent methods of analysis and a single method that has been carefully evaluated with respect to its accuracy and precision. Methods were selected to include those that are commonly used in the field and in laboratories.

The certified values are given in table 1 and are based on at least a 1.0 g sample of the material which is the minimum amount that should be used for analysis.

Table 1

| Element ¹ | Content ² $\mu\text{g/g}$ | Element ¹ | Content ² , Wt % |
|---------------------------|--------------------------------------|-------------------------|-----------------------------|
| Lead ^d | 2.80 ± 0.08 | Sulfur ^{f,g,h} | 2.85 ± 0.05 |
| Manganese ^{b,c} | 0.19 ± 0.02 | | |
| Nickel ^{d,c} | 29 ± 1 | | |
| Selenium ^{b,c} | 0.15 ± 0.02 | | |
| Sodium ^{b,c} | 87 ± 4 | | |
| Vanadium ^{a,b,d} | 56 ± 2 | | |
| Zinc ^{b,d} | 2.7 ± 0.2 | | |

1. Method of Analysis

- | | |
|---------------------------------------|--|
| a. Isotope Dilution Mass Spectrometry | e. Inductive Coupled Plasma Spectrometry |
| b. Neutron Activation Analysis | f. Gravimetry |
| c. Atomic Absorption Spectrometry | g. Ion Chromatography |
| d. Spark Source Mass Spectrometry | h. X-ray Fluorescence |

2. The uncertainties shown are expressed at the 95% confidence level and include any observed material heterogeneity, possible method differences, and errors of measurement.

NOTICE: The certification of SRM 1634a is valid for 3 years from date of purchase.

The statistical analysis of the certification data was performed by K.R. Eberhardt of the Statistical Engineering Division.

The overall direction and coordination of the analytical measurements leading to certification were performed in the Inorganic Analytical Research Division, E.L. Garner, Chief.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T.E. Gills.

Washington, D.C. 20234
 February 19, 1982

George A. Uriano, Chief
 Office of Standard Reference Materials

(over)

PREPARATION, TESTING, AND ANALYSIS

A random scheme for sample selection was used in assessing the homogeneity of this material. The elements calcium and vanadium were measured by x-ray fluorescence as indicators of homogeneity. Based on these elements, the material variability for this lot of 1634a is within $\pm 2\%$ relative.

Long-term stability of this SRM has not been rigorously established. When not in use, the material should be stored in the tightly sealed bottle. NBS will continue to monitor this material and any substantive change in its certification will be reported to the purchasers.

Analyses for the various elements were performed in the Center for Analytical Chemistry, Inorganic Analytical Research Division, by I. L. Barnes, T.A. Butler, E.R. Deardorff, J.W. Gramlich, S. Hanamura, H.M. Kingston, W.F. Koch, G.M. Lambert, R.M. Lindstrom, L.A. Machlan, J.R. Moody, P.J. Paulson, T.C. Rains, T.A. Rush, and R. Zeisler.

The homogeneity studies were performed in the Gas and Particulate Science Division by P.A. Pella and M. Watson.

The physical properties were measured by S. Weeks, Materials Chemistry Division, Center for Materials Science.

The values and physical properties data in table 2 are *not certified* because they are based on a non-reference method or were not determined by two or more independent methods. The values are included for information only.

Table 2

Supplemental Information

| Element | Content, $\mu\text{g/g}$ | Physical Properties | |
|------------|--------------------------|----------------------------------|-------------------------|
| Arsenic | (0.12) | | |
| Beryllium | (0.006) | Flash Point ^a | 64 °C |
| Bromine | (<1) | | |
| Cadmium | (0.002) | Kinematic Viscosity ^b | 321.66 |
| Calcium | (16) | at 50 °C | |
| Chlorine | (31) | Pour Point ^c | -10 °C |
| Chromium | (0.7) | | |
| Cobalt | (0.3) | Density at 20 °C ^d | 0.995 g/cm ³ |
| Iron | (31) | | |
| Mercury | (<0.002) | | |
| Molybdenum | (0.12) | | |

Methods Used for Physical Tests

- ASTM D-93-80 Flash Point by Pensky-Martens Closed Tester
- ASTM D445-79 Kinematic Viscosity of Transparent and Opaque Liquids
- ASTM D97-66 (1978) Pour Point of Petroleum Oils
- ASTM D4052-81 Density and Relative Density of Liquids by Digital Density Meter (modified)

National Bureau of Standards Certificate of Analysis

Standard Reference Material 1635

Trace Elements in Coal (Subbituminous)

This Standard Reference Material is intended for use in the calibration of apparatus and the evaluation of techniques employed in the trace element analysis of coal and similar materials. The material should be dried without heat to constant weight before use.

The recommended procedures for drying are either vacuum drying at ambient temperature for 24 hours, or freeze drying in which the drying chamber is kept at room temperature. The moisture content of this material is approximately 20%. Because of this moisture level, it is recommended that small individual samples be dried immediately before use. Drying of large samples may result in a violent discharge of water vapor and resultant loss of sample. When not in use, the material should be kept in a tightly sealed bottle and stored in a cool, dark place. Long-term (>1 year) stability of this SRM has not been rigorously established. NBS will continue to monitor this material and any substantive change will be reported to purchasers.

The certified values given below are based on at least a 250-mg sample of the dried material, the minimum amount that should be used for analysis.

| Element ¹ | Content, $\mu\text{g/g}^2$ | Element ¹ | Content, $\mu\text{g/g}^2$ |
|--------------------------|----------------------------|-------------------------|----------------------------|
| Arsenic ^{a,b} | 0.42 ± 0.15 | Thorium ^{c,e} | 0.62 ± 0.04 |
| Cadmium ^{c,d,e} | 0.03 ± 0.01 | Uranium ^c | 0.24 ± 0.02 |
| Chromium ^{c,e} | 2.5 ± 0.3 | Vanadium ^{e,g} | 5.2 ± 0.5 |
| Copper ^{a,c,e} | 3.6 ± 0.3 | Zinc ^{c,d} | 4.7 ± 0.5 |
| Lead ^{c,d} | 1.9 ± 0.2 | | |
| Manganese ^{a,e} | 21.4 ± 1.5 | Element ¹ | Wt. % ² |
| Nickel ^{e,d} | 1.74 ± 0.10 | Iron ^{c,d,e,f} | 0.239 ± 0.005 |
| Selenium ^{a,e} | 0.9 ± 0.3 | Sulfur ^{f,h} | 0.33 ± 0.03 |

1. Methods of Analysis:

- | | |
|---------------------------------------|--------------------------------|
| a. Atomic Absorption Spectrometry | e. Neutron Activation |
| b. Photon Activation | f. Spectrophotometry |
| c. Isotope Dilution Mass Spectrometry | g. Flame Emission Spectrometry |
| d. Polarography | h. Gravimetry |

2. The estimated uncertainty is based on judgment and represents an evaluation of the combined effects of method imprecision, possible systematic errors among methods, and material variability for samples of 250-mg or more. (No attempt was made to derive exact statistical measures of imprecision because several methods were involved in the determination of most constituents.)

The overall direction and coordination of the analytical measurements leading to this certificate were performed in the Analytical Chemistry Division under the chairmanship of L. J. Moore.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by W. P. Reed.

Washington, D.C. 20234
 August 22, 1979
 (Revision of Certificate
 dated 1-23-78)

George A. Uriano
 Office of Standard Reference Materials

(over)

PREPARATION, TESTING, and ANALYSIS

This material was prepared from one lot of subbituminous coal from the Eagle Mine of The Imperial Coal Company, Erie, Colorado. The material was ground and sieved thru a No. 65 (230 μm) sieve by the Colorado School of Mines Research Institute. The material was then blended in a V-type blender.

Samples for homogeneity testing were taken from the top, middle, and bottom of three bulk containers of coal, and analyzed by neutron activation analysis for sodium, scandium, chromium, iron, cobalt, lanthanum, cerium, and thorium. Replicate analyses of 250-mg samples indicated a homogeneity for these elements of $\pm 2.5\%$ (relative) except for chromium, which was homogeneous within counting statistics of $\pm 6\%$. The homogeneity measurements were performed in the NBS Analytical Chemistry Division by R. R. Greenberg. Certification analyses for the various elements were made in the NBS Analytical Chemistry Division by T. J. Brady, B. I. Diamondstone, L. P. Dunstan, M. S. Epstein, M. Gallorini, E. L. Garner, T. E. Gills, J. W. Gramlich, R. R. Greenberg, S. H. Harrison, G. M. Hyde, G. J. Lutz, L. A. Machlan, E. J. Maienthal, J. D. Messman, T. J. Murphy, and T. C. Rains.

The following values are *not certified* because they were based on a non-reference method, or were not determined by two or more independent methods. They are included for information only.

| <u>Element</u> | <u>Content</u> <u>($\mu\text{g/g}$)</u> |
|----------------|---|
| Antimony | (0.14) |
| Cerium | (3.6) |
| Cobalt | (0.65) |
| Europium | (0.06) |
| Gallium | (1.05) |
| Hafnium | (0.29) |
| Scandium | (0.63) |
| | <u>(wt. %)</u> |
| Aluminum | (0.32) |
| Sodium | (0.24) |
| Titanium | (0.02) |

National Bureau of Standards

Certificate of Analysis

Standard Reference Materials 1636a, 1637a, 1638a

Lead in Reference Fuel

This Standard Reference Material is intended for use in the calibration of instruments and the evaluation of techniques used for the analysis of lead in gasoline. Samples of the reference fuel are supplied at four lead concentrations, nominally 0.03, 0.05, 0.07, and 2.0 g/gal. These Standard Reference Materials are made up of various combinations of the four concentrations, see Table 1 on the back of this certificate.

Certified Values: The certified values for the lead content, expressed in units of $\mu\text{g/g}$, are shown below. These certified values are based on results obtained by isotope dilution mass spectrometry, a definitive method of known accuracy.

| <u>Vial Identification</u> | <u>Nominal Lead Concentration g/gal</u> | <u>Certified Lead Concentration $\mu\text{g/g}$</u> |
|--------------------------------|---|--|
| I | 0.03 | 11.2 ± 0.2^a |
| II | 0.05 | 18.8 ± 0.1 |
| III | 0.07 | 25.1 ± 0.2 |
| IV | 2.0 | 764 ± 4 |

^aThe uncertainties shown are the 95 percent confidence intervals for a single determination plus allowance for known sources of possible error.

Use: The certification of these materials is based on a minimum sample size of 1.0 gram and only samples equal to or greater than 1 gram should be used for any analytical determination to be related to the certified values of this certificate.

Stability: The ampoules should be stored at temperatures between 10-30 °C. They should not be exposed to intense sources of radiation, including ultraviolet lamps or sunlight. The ampoules should be opened only at time of use. No attempt should be made to keep the material in opened ampoules for future use.

Source and Preparation of Material: The reference fuel containing lead at the four concentration levels were supplied by Phillips Petroleum Company of Bartlesville, Oklahoma. The 91-octane number (Research Octane Number) reference fuel is a mixture of 91 percent by volume (0.899 mole-fraction) 2,2,4-trimethylpentane and 9 percent by volume (0.101 mole-fraction) n-heptane. Lead was added in the form of tetraethyl lead motor mix.

Analyses leading to certification were performed in the Inorganic Analytical Research Division by T. J. Murphy and I. L. Barnes.

The overall direction and coordination of the technical measurements leading to this certificate were performed by E. L. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of these Standard Reference Materials were coordinated through the Office of Standard Reference Materials by T. E. Gills.

Washington, D.C. 20234
February 5, 1980

George A. Uriano, Chief
Office of Standard Reference Material

(over)

Table 1
Composition of SRM's 1636a, 1637a, 1638a

| <u>SRM</u> | <u>Nominal Concentration</u> | <u>No. Units.</u> |
|------------|------------------------------|-------------------|
| 1636a | 0.03,0.05,0.07, 2.0 g/gal | 3 vials each |
| 1637a | 0.03,0.05,0.07 g/gal | 4 vials each |
| 1638a | 2.0 g/gal | 12 vials each |

Additional Information: Because the volume of the reference material varies with temperature, the various concentrations of lead are certified by weight, i.e., micrograms of lead per gram of fuel. For convenience to the user, information is given for the concentration in the customary units, grams per gallon and grams per liter, at 23 °C. These data are shown in Table 2.

Table 2

| <u>Vial Identification</u> | <u>Nominal Concentration</u> <u>g/gal</u> | <u>Density^a</u> <u>at 23 °C</u> <u>g/mL</u> | <u>Lead Concentration^b</u> <u>at 23 °C</u> | |
|----------------------------|--|--|--|------------|
| | | | <u>g/gal</u> | <u>g/L</u> |
| I | 0.03 | 0.6888 | 0.0292 | 0.0077 |
| II | 0.05 | 0.6888 | 0.0490 | 0.0129 |
| III | 0.07 | 0.6888 | 0.0654 | 0.0173 |
| IV | 2.0 | 0.6895 | 1.994 | 0.527 |

^aThe density (ρ) of each concentration was measured at 23 °C using a modification of ASTM Method D1217. The temperature coefficient of these materials is $0.0008 \text{ g}(\text{mL})^{-1}(\text{°C})^{-1}$.

^bThe conversion of the certified values ($\mu\text{g/g}$) to C(g/gal) and C(g/L) was done using equations 1 and 2 respectively.

$$\text{Eq. 1} \quad C_{\text{g/gal}} = \frac{3.785 \rho C_{\mu\text{g/g}}}{10^3}$$

$$\text{Eq. 2} \quad C_{\text{g/L}} = \frac{\rho C_{\mu\text{g/g}}}{10^3}$$



National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1641b

Mercury in Water – $\mu\text{g/mL}$

This Standard Reference Material is intended for use in the primary calibration of instruments and techniques used for the determination of mercury in natural waters. It is designed for the preparation of calibration solutions and for use as a “spike” sample in a “method-of-additions” type analytical procedure.

Mercury concentration $1.52 \pm 0.04 \mu\text{g/mL}$

The estimated uncertainty, $0.04 \mu\text{g/mL}$, includes the effects of the observed random variability and an upper bound estimate of possible systematic errors. The random variability expressed as two standard deviations of the certified value is $\pm 0.02 \mu\text{g/mL}$ and reflects both internal and between-method variabilities for the NBS atomic absorption and neutron activation measurements. The upper bound estimate of possible systematic errors is $\pm 0.02 \mu\text{g/mL}$.

Stability: The long-term stability of trace mercury solutions has been a constant problem. At or below the $\mu\text{g/mL}$ level, mineral acid stabilization is not sufficient. However, the addition of trace gold to a nitric acid solution of mercury was found to stabilize the concentration of mercury in the two previous issues of this Mercury in Water SRM. Although the mercury concentration of SRM 1641b has not changed significantly in eight months, the stability will continue to be monitored. However, SRM 1641b should *not* be used after ONE YEAR FROM date of purchase.

Precautions: Traces of mercury vapor are present in most laboratory situations. Therefore, contamination of reagents, equipment, and common laboratory materials may cause a severe problem. Apparatus for analysis at this level must be scrupulously cleaned immediately before use, and only the purest-grade reagents, with respect to mercury, should be used.

SRM 1641b was prepared by J.R. Moody. Atomic absorption analyses were performed by T.C. Rains and T.A. Butler; and neutron activation analyses by R. Zeisler, Inorganic Analytical Research Division.

The overall direction and coordination of technical measurements leading to certification were performed under the chairmanship of E.L. Garner, Inorganic Analytical Research Division. The statistical evaluation was done by R.C. Paule.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Analytical: Two independent techniques were used in the certification of this Standard Reference Material: atomic absorption spectrometry and instrumental neutron activation analysis.

Use: This SRM consists of six ampoules, each containing approximately 20 mL of solution. Dilutions may be made by the addition of accurately measured aliquots, withdrawn from an ampoule, to known volumes of pure or natural water (spiking mode) using conventional techniques. Blank determinations should be made of the water and other reagents used.

The reliability of the dilution process will depend on the care exercised and the reliability of the calibration of the volumetric apparatus, which should have an uncertainty no greater than one percent. The volumetric apparatus should be scrupulously cleaned. Diluted solutions should be used without delay, as their stability cannot be guaranteed. SRM 1642b, which is certified for mercury at the ng/mL level, should be used to validate methodology for these concentrations. The long-term retention of unused portions of this Standard Reference Material in opened ampoules is not recommended.

Washington, D.C. 20234
April 13, 1983

George A. Uriano, Chief
Office of Standard Reference Materials

National Bureau of Standards

Certificate

Standard Reference Material 1642b

Mercury in Water – ng/mL

This Standard Reference Material is intended for use in the primary standardization of instruments and techniques used for the determination of mercury in water. It is intended for use as received, without dilution or other alteration. The concentration of mercury in this Standard Reference Material is at, or near, the detection limit of most commercial instruments used for the determination of mercury in water. It is to be used for the primary standardization of these instruments near these detection limits where many analytical problems occur.

Mercury Concentration 1.49 ± 0.06 ng/mL

The estimated uncertainty, 0.06, includes the effects of the observed random variability and an upper bound estimate of possible systematic errors. The random variability expressed as two standard deviations of the certified value is ± 0.04 and reflects both internal and between-method variabilities for the NBS atomic absorption and neutron activation measurements. The upper bound estimate of possible systematic errors is ± 0.02 .

Stability: Trace mercury solutions have been a constant problem when long-term storage is required. Below the μg . mL level, mineral acid stabilization is not sufficient. A stabilizing technique has been applied to this Standard Reference Material that allows for prolonged storage. Gold, as the tetrachloride, has been added in a concentration 10 times that of the mercury. The gold ion, in conjunction with the normal mineral acid, has proven to be an effective stabilizer. It is recommended that this Standard Reference Material not be used after ONE YEAR FROM DATE OF PURCHASE.

Precautions: Traces of mercury vapor are present in most laboratory situations. Therefore, contamination of reagents, equipment, and common laboratory materials is a severe problem. Apparatus for analyses at this level must be scrupulously cleaned immediately before use, and only the purest-grade reagents should be employed. After use, the bottle should be capped tightly and placed inside the aluminized bag, which should be folded and sealed with a sealing tape. This safeguard will assist in maintaining the integrity of the sample.

Analytical: Two independent techniques were used in the certification of this Standard Reference Material: atomic absorption spectroscopy and neutron activation analysis.

Use: This Standard Reference Material should be used, as received, without dilution. It may be carried through the chemical manipulations required for the analytical procedure normally used for the analysis of natural waters.

This Standard Reference Material was prepared by J.R. Moody. Atomic absorption analyses were performed by I.C. Rains and T.A. Butler and neutron-activation analyses were performed by R. Zeisler.

The overall direction and coordination of the technical measurements leading to the certification were performed under the chairmanship of E.L. Garner. The statistical evaluation was done by R.C. Paule.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Washington, D.C. 20234
July 15, 1982

George A. Uriano, Chief
Office of Standard Reference Materials

National Bureau of Standards

Certificate

Standard Reference Material 1643b

Trace Elements in Water

This Standard Reference Material (SRM) is intended primarily for use in evaluating the accuracy of trace element determinations in filtered and acidified fresh water and for calibrating instrumentation used in these determinations. SRM 1643b consists of approximately 950 mL of water in a polyethylene bottle, which is sealed in an aluminized bag to maintain stability. SRM 1643b simulates the elemental composition of fresh water. Nitric acid is present at a concentration of 0.5 moles per liter to stabilize the trace elements.

Concentrations of Constituent Elements: The concentrations of the trace elements that were determined are shown in Table 1. The certified values are based on results obtained either by reference methods of known accuracy or by two or more independent, reliable analytical methods. Noncertified values, which are given for information only, appear in parentheses.

Notice and Warnings to Users:

Expiration of Certification: This certification is invalid two years after the shipping date.

Precautions: The bottle should be shaken before use because of possible water vapor condensation. To prevent possible contamination of the SRM, do not insert pipets into the bottle. After use, the bottle should be capped tightly and placed inside the aluminized bag, which should be folded and sealed with sealing tape. This safeguard will protect the SRM from possible environmental contamination and long-term loss of water.

Elemental determinations of ng/g levels are limited by contamination. Apparatus should be scrupulously cleaned and only the purest grade reagents employed. Sampling and manipulations, such as evaporations, should be done in a clean environment, for example, a Class 100 clean hood.

The overall direction and coordination of the technical measurements leading to this certification were performed under the direction of E. Garner, Chief of the Inorganic Analytical Research Division.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Washington, DC 20234
May 18, 1984

Stanley D. Rasberry, Chief
Office of Standard Reference Materials

(over)

(Table 1)
Concentrations of Constituent Elements

| Element | Concentration,* ng/g | Element | Concentration,* ng/g |
|---------------------------|-------------------------|-----------------------------|-------------------------|
| Arsenic ^{1,5} | (49)** | Lead ^{3,4b} | 23.7 ± 0.7 |
| Barium ^{2a,2b,5} | 44 ± 2 | Manganese ^{1,2a,3} | 28 ± 2 |
| Beryllium ^{1,2a} | 19 ± 2 | Molybdenum ^{2a,5} | 85 ± 3 |
| Bismuth ¹ | (11) | Nickel ^{2a,3} | 49 ± 3 |
| Boron ^{2a} | (94) | Selenium ^{1,5} | 9.7 ± 0.5 |
| Cadmium ^{2b,3,5} | 20 ± 1 | Silver ^{1,5} | 9.8 ± 0.8 |
| Chromium ^{4b} | 18.6 ± 0.4 | Strontium ^{2a,5} | 227 ± 6 |
| Cobalt ^{1,5} | 26 ± 1 | Thallium ^{4b} | 8.0 ± 0.2 |
| Copper ^{3,4b} | 21.9 ± 0.4 | Vanadium ^{4b} | 45.2 ± 0.4 |
| Iron ^{2a,4a,5} | 99 ± 8 | Zinc ^{2a,5} | 66 ± 2 |

* The estimated uncertainty is based on judgment and represents an evaluation of the combined effects of method imprecision and possible systematic errors among methods. To convert to nanograms per milliliter, multiply by the density of the SRM. The density at 23 °C is 1.017 grams per milliliter.

** Values in parentheses are not certified.

- | | |
|---|---|
| 1. Atomic absorption spectrometry, electrothermal | 4. Isotopic dilution mass spectrometry, |
| 2. Atomic emission spectrometry, | a. resonance ionization |
| a. dc plasma | b. thermal ionization |
| b. flame | 5. Neutron activation, |
| 3. Laser enhanced ionization flame spectrometry | instrumental |

Source and Preparation of Material: SRM 1643b was prepared at the U.S. Geological Survey, National Water Quality Laboratory, Arvada, Colorado, under the direction of V.J. Janzer of that laboratory and J.R. Moody of the NBS Center for Analytical Chemistry. Only high-purity reagents were used and the containers were acid-cleaned and sterilized before use. In the preparation, a polyethylene cylindrical tank was filled with distilled water and sufficient nitric acid to make the solution approximately 0.5 moles HNO₃ per liter. Solutions containing known amounts of calcium, sodium, magnesium, potassium, and the elements to be determined were added to the acidified water solution with constant stirring. After thoroughly mixing, the solution was filtered, sterilized, and then transferred to one-liter polyethylene bottles. The approximate concentrations, in µg/mL, of Ca, Na, Mg, and K are respectively 35, 8, 15, and 3.

Analysts:

Center for Analytical Chemistry, National Bureau of Standards

- | | |
|--------------------|------------------------|
| 1. K. A. Brletic | 10. J. R. Moody |
| 2. T. A. Butler | 11. L. J. Powell |
| 3. E. C. Deal | 12. T. C. Rains |
| 4. M. S. Epstein | 13. T. A. Rush |
| 5. J. D. Fassett | 14. S. F. Stone |
| 6. K. Fitzpatrick | 15. G. C. Turk |
| 7. H. M. Kingston | 16. R. L. Watters, Jr. |
| 8. R. M. Lindstrom | 17. R. Zeisler |
| 9. L. A. Machlan | |

National Bureau of Standards Certificate

Standard Reference Material 1644

Generator Columns for Polynuclear Aromatic Hydrocarbons

SRM 1644 is intended to provide accurate concentrations of anthracene, benzo(a) anthracene (1,2-benzanthracene), and benzo(a)pyrene (3,4-benzpyrene) in water. The SRM consists of three 50 cm x 0.6 cm (coiled) stainless steel tubes, each packed with fine quintus quartz (sea sand) coated with approximately 0.5 percent by weight of the polynuclear aromatic hydrocarbon (PAH) of interest.

Principle of Operation: A saturated aqueous solution of the PAH of interest is generated by flowing high-purity water slowly through the column. Because the aqueous solubility of a compound is a well-defined thermodynamic quantity, a saturated solution has a fixed concentration (1, 2, 3).

Equilibration and Use of Generator Columns: To equilibrate a new column before initial use, purge with high-purity water, such as commercial HPLC grade water. The volume required for equilibration of each column is: 500 mL for anthracene, 1000 mL for benzo(a)anthracene, and 500 mL for benzo(a)pyrene. After equilibration, pump the high-purity water at a constant temperature (± 0.1 °C) through the column at a flow rate between 0.1 and 5 mL/min to produce a saturated solution. Record the temperature. The solution should be used immediately after generation to avoid sorption losses.

If either the temperature is changed by as much as 1 °C or the flow is interrupted for less than one hour, pump 25 mL water through the column under the new conditions to restore equilibrium prior to sample collection. However, if the flow is interrupted for more than one hour, pump 50 mL water prior to sample collection. During periods of frequent use, a column can be kept equilibrated by maintaining a steady, but low, flow rate of approximately 0.1 mL/min through the column. The flow rate can be increased to collect a large volume of sample and then decreased again. Columns should be purged with 10 liters of oil-free nitrogen prior to storage periods of more than one month.

Certified Concentrations: When used as directed, these columns generate saturated solutions. The concentrations of the compounds in these solutions at temperatures between 10 and 30 °C were determined by two independent analytical methods. The data obtained were combined by fitting an empirical expression of the form $\ln[\text{Conc}] = A + B(1/T) + C(1/T^2)$ by least squares. In this equation, $\ln[\text{Conc}]$ is the natural logarithm of the concentration, T is the absolute temperature, and A, B, and C are constants for each compound. The derived equations for anthracene, benzo(a)anthracene, and benzo(a)pyrene were used to calculate the certified concentrations at one degree intervals between 10 and 30 °C. These certified concentrations are given in Tables I-III.

Service Life of Columns: Generator columns for anthracene, benzo(a)anthracene, and benzo(a)pyrene are certified for either two years or for total aqueous purge volumes of 7.5×10^2 , 3×10^3 , and 1.5×10^4 liters, respectively, whichever comes first.

Consultation on the statistical design of the experimental work and statistical analysis of the data was provided by K. R. Eberhardt of the Statistical Engineering Division. Coordination of the technical measurements leading to certification was performed by W. E. May, R. A. Velapoldi, and H. S. Hertz. Technical measurements leading to the development and certification of SRM 1644 were performed by the following members of the Center for Analytical Chemistry: W. E. May, J. M. Brown-Thomas, W. J. Sonnefeld, R. A. Velapoldi, and P. A. White.

The technical and support aspects concerning the certification and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Washington, D.C. 20234
April 27, 1981

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

The anthracene generator column, after equilibration, produces aqueous solutions that contain very small amounts of phenanthrene. Because of its relatively high solubility (1000 $\mu\text{g}/\text{kg}$), residual amounts of phenanthrene are depleted from the column at a rapid rate.

The benzo(a)anthracene generator column, after equilibration, produces aqueous solutions that contain as much as 0.3 $\mu\text{g}/\text{kg}$ anthracene in addition to the certified concentrations of benzo(a)anthracene. The concentration of anthracene in the generator column effluent varies with time (volume) and, therefore, is not certified.

The benzo(a)pyrene generator column, after equilibration, produces aqueous solutions that contain as much as 0.1 $\mu\text{g}/\text{kg}$ benzo(a)anthracene and 1 $\mu\text{g}/\text{kg}$ chrysene in addition to the certified concentrations of benzo(a)pyrene. The concentration of these non-analyte components vary with time and, therefore, are not certified.

Analyses of the Saturated Aqueous Solutions: The concentrations of the saturated aqueous solutions of anthracene, benzo(a)anthracene, and benzo(a)pyrene in the effluent from the respective generator columns were determined by two independent analytical techniques. The first technique was high-performance liquid chromatography (HPLC). It involved quantitative extraction of the PAH of interest from the aqueous effluent by an "extractor column" packed with an octadecylsilane (C_{18}) bonded phase; use of an acetonitrile-water eluant to transfer components from the extractor column to an analytical C_{18} column for separation of the analyte from non-analyte components; and detection of the analyte by measuring its absorbance at 254 nm. The second technique involved the use of a "standard addition" spectrofluorimetric technique for "on stream" analysis. The aqueous effluent from the generator column was mixed with PAH standards dissolved in acetonitrile and the PAH concentration of the resultant mixture was determined by fluorescence at the following excitation (λ_{ex}) and emission (λ_{em}) wavelengths: anthracene, $\lambda_{\text{ex}} = 254$ nm, $\lambda_{\text{em}} = 384$ and 404 nm; benzo(a)anthracene, $\lambda_{\text{ex}} = 290$ nm, $\lambda_{\text{em}} = 395$ nm; benzo(a)pyrene, $\lambda_{\text{ex}} = 296$ nm, $\lambda_{\text{em}} = 414$ nm. The PAH effluent concentration was determined mathematically. Where necessary, corrections for inner filter effects or the emission-absorbance contributions by impurities were made to obtain the values for determining the certified PAH concentrations listed in Tables I-III.

References:

1. May, W. E., The Solubility Behavior of Polycyclic Aromatic Hydrocarbons in Aqueous Systems, American Chemical Society Advances in Chemistry Series 185 (7), 143-192 (1980).
2. May, W. E., Wasik, S. P., and Freeman, D. H., Determination of the Aqueous Solubility of Polynuclear Aromatic Hydrocarbons by a Coupled-Column Liquid Chromatographic Technique, Anal. Chem. 50, 1 (1978).
3. Schwarz, F. P. and Miller, J. M., Determination of the Aqueous Solubilities of Organic Liquids at 10 °C, 20 °C, and 30 °C by Elution Chromatography, Anal. Chem. 52, 2162-2164 (1980).

Table I. Certified Aqueous Concentrations of Anthracene and Their Uncertainties, in Micrograms/Kilogram and Nanomoles/Liter, as a Function of Temperature

| Temperature °C | Concentration and Its Uncertainty ¹ , | |
|-------------------|--|------------|
| | µg/kg | nmol/L |
| 10 | 16.6 ± 0.7 | 93.1 ± 4.0 |
| 11 | 17.6 ± 0.6 | 98.7 ± 3.6 |
| 12 | 18.7 ± 0.6 | 105 ± 3.3 |
| 13 | 19.8 ± 0.5 | 111 ± 3.1 |
| 14 | 21.1 ± 0.5 | 118 ± 2.9 |
| 15 | 22.4 ± 0.5 | 126 ± 2.9 |
| 16 | 23.8 ± 0.5 | 134 ± 2.9 |
| 17 | 25.4 ± 0.5 | 142 ± 2.9 |
| 18 | 27.0 ± 0.5 | 151 ± 3.0 |
| 19 | 28.8 ± 0.5 | 161 ± 3.1 |
| 20 | 30.7 ± 0.6 | 172 ± 3.2 |
| 21 | 32.8 ± 0.6 | 184 ± 3.2 |
| 22 | 35.0 ± 0.6 | 196 ± 3.2 |
| 23 | 37.4 ± 0.6 | 210 ± 3.2 |
| 24 | 39.9 ± 0.6 | 224 ± 3.3 |
| 25 | 42.7 ± 0.6 | 239 ± 3.7 |
| 26 | 45.7 ± 0.8 | 256 ± 4.4 |
| 27 | 48.9 ± 1.0 | 273 ± 5.5 |
| 28 | 52.4 ± 1.3 | 293 ± 7.2 |
| 29 | 56.1 ± 1.7 | 313 ± 9.5 |
| 30 | 60.1 ± 2.2 | 336 ± 12 |

¹The uncertainties are 99 percent (Working-Hotelling) confidence bands for the entire regression curve. The difference between the true and certified concentrations should be less than the stated uncertainty at the 99 percent confidence level.

Table II. Certified Aqueous Concentrations of Benzo(a)anthracene and Their Uncertainties, in Micrograms/Kilogram and Nanomoles/Liter, as a Function of Temperature

| Temperature °C | Concentration and Its Uncertainty ¹ , | |
|-------------------|--|------------|
| | µg/kg | nmol/L |
| 10 | 3.38 ± 1.2 | 14.8 ± 5.3 |
| 11 | 3.60 ± 1.1 | 15.8 ± 4.6 |
| 12 | 3.83 ± 0.91 | 16.8 ± 4.0 |
| 13 | 4.09 ± 0.79 | 17.9 ± 3.4 |
| 14 | 4.36 ± 0.68 | 19.1 ± 3.0 |
| 15 | 4.65 ± 0.59 | 20.4 ± 2.6 |
| 16 | 4.96 ± 0.54 | 21.7 ± 2.4 |
| 17 | 5.29 ± 0.55 | 23.2 ± 2.4 |
| 18 | 5.65 ± 0.60 | 24.8 ± 2.6 |
| 19 | 6.04 ± 0.68 | 26.4 ± 3.0 |
| 20 | 6.45 ± 0.77 | 28.2 ± 3.4 |
| 21 | 6.90 ± 0.87 | 30.2 ± 3.8 |
| 22 | 7.38 ± 0.94 | 32.3 ± 4.1 |
| 23 | 7.90 ± 1.0 | 34.5 ± 4.4 |
| 24 | 8.45 ± 1.0 | 36.9 ± 4.5 |
| 25 | 9.05 ± 1.0 | 39.5 ± 4.6 |
| 26 | 9.69 ± 1.0 | 42.3 ± 4.6 |
| 27 | 10.4 ± 1.1 | 45.4 ± 4.7 |
| 28 | 11.1 ± 1.2 | 48.4 ± 5.0 |
| 29 | 11.9 ± 1.3 | 51.9 ± 5.9 |
| 30 | 12.8 ± 1.7 | 55.8 ± 7.3 |

¹The uncertainties are 99 percent (Working-Hotelling) confidence bands for the entire regression curve. The difference between the true and certified concentrations should be less than the stated uncertainty at the 99 percent confidence level.

Table III. Certified Aqueous Concentrations of Benzo(a)pyrene and Their Uncertainties, in Micrograms/Kilogram and Nanomoles/Liter, as a Function of Temperature

| Temperature °C | Concentration and Its Uncertainty ¹ , | |
|-------------------|--|-------------|
| | µg/kg | nmol/L |
| 10 | 0.59 ± 0.06 | 2.34 ± 0.23 |
| 11 | 0.63 ± 0.05 | 2.50 ± 0.20 |
| 12 | 0.67 ± 0.04 | 2.65 ± 0.17 |
| 13 | 0.71 ± 0.04 | 2.81 ± 0.15 |
| 14 | 0.76 ± 0.03 | 3.01 ± 0.12 |
| 15 | 0.81 ± 0.03 | 3.21 ± 0.11 |
| 16 | 0.87 ± 0.03 | 3.44 ± 0.10 |
| 17 | 0.93 ± 0.03 | 3.68 ± 0.10 |
| 18 | 0.99 ± 0.03 | 3.92 ± 0.11 |
| 19 | 1.06 ± 0.03 | 4.19 ± 0.12 |
| 20 | 1.13 ± 0.03 | 4.47 ± 0.13 |
| 21 | 1.21 ± 0.04 | 4.79 ± 0.15 |
| 22 | 1.30 ± 0.04 | 5.14 ± 0.16 |
| 23 | 1.39 ± 0.04 | 5.50 ± 0.16 |
| 24 | 1.49 ± 0.04 | 5.89 ± 0.17 |
| 25 | 1.59 ± 0.04 | 6.28 ± 0.17 |
| 26 | 1.71 ± 0.04 | 6.76 ± 0.18 |
| 27 | 1.83 ± 0.05 | 7.23 ± 0.19 |
| 28 | 1.96 ± 0.05 | 7.74 ± 0.22 |
| 29 | 2.11 ± 0.07 | 8.33 ± 0.27 |
| 30 | 2.26 ± 0.09 | 8.92 ± 0.35 |

¹The uncertainties are 99 percent (Working-Hotelling) confidence bands for the entire regression curve. The difference between the true and certified concentrations should be less than the stated uncertainty at the 99 percent confidence level.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1645

River Sediment

This Standard Reference Material (SRM) is intended for use for the calibration of apparatus and the verification of methods used in the analysis of river sediments and material with a similar matrix.

Certified Values of Constituent Elements: The certified values for the constituent elements are shown in Table 1. The analytical techniques used and the names and affiliations of the analysts are shown in Table 3. Certified values are based on results obtained by reference methods of known accuracy and analyses performed by two or more analysts; or alternatively, from results obtained by two or more independent, reliable analytical methods. Noncertified values are given for information only in Table 2. All values are based on measurements made on a dried sample of at least 100 mg for all constituents except iron and chromium for which a 1-g sample was used.

Notice and Warnings to Users:

Expiration of Certification: This certification is invalid 5 years from the date of purchase.

Stability: This material has been freeze-dried and is essentially free of moisture. However, its stability has not been rigorously assessed. NBS will continue to monitor this material and if substantive changes in certification occur the purchasers will be notified. The material should be kept in its original bottle and stored at temperatures between 10-30 °C. The material should be dried without heat to a constant weight before using. Recommended procedures for drying are: (1) drying for 24 hours using a cold trap at or below -50 °C and a pressure not greater than 30 Pa (0.2 mm Hg); (2) drying for 24 hours in a desiccator over P₂O₅ or Mg (ClO₄)₂.

Use: Material of this kind is intrinsically heterogeneous. Consequently, the analyst should endeavor to minimize any segregation by thoroughly mixing the contents of the bottle by shaking and/or rolling before each use. In addition, when taking a portion for analysis, the analyst should strive to remove as representative a sample as possible.

Source and Preparation of Material: The material for this SRM was prepared from material dredged from the bottom of the Indiana Harbor Canal near Gary, Indiana. The material was screened to remove foreign objects, freeze-dried, and sieved to pass a No. 80 (180 μm) screen. The material was thoroughly mixed in a V-blender and bottled. The bulk material was radiation-sterilized to minimize alteration due to biological activity.

The collection, freeze-drying and homogenization of this SRM were performed under the supervision and direction of H.L. Rook, Gas and Particulate Science Division.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of J.K. Taylor, Center for Analytical Chemistry.

The technical and support aspects involved in the preparation, current and previous certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T.E. Gills and W.P. Reed.

Washington, D.C. 20234
May 5, 1982
(Revision of Certificate
dated 11-16-78)

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

Homogeneity Assessment and Certification: The homogeneity of this material was established using a minimum sample size of 100 milligrams for all constituents except iron and chromium for which the sample size was 1.0 gram.

Randomly selected bottles were used for the analytical measurements. Each analyst examined at least 6 different bottles, some of them measuring replicate samples from each bottle. Accordingly, it is believed that all bottles of this SRM have substantially the same composition. Measurements and calibrations were made to reduce random and systematic errors to no more than one percent, relative.

Table 1. Certified Values of Constituent Elements

| <u>Major Constituents</u> | | <u>Minor Constituents</u> | |
|---------------------------|--|---------------------------|--|
| <u>Element</u> | <u>Content</u> <u>wt. percent^a</u> | <u>Element</u> | <u>Content</u> <u>wt. percent^a</u> |
| Aluminum ^b | 2.26 ± 0.04 | Magnesium ^b | 0.74 ± 0.02 |
| Chromium | 2.96 ± 0.28 | Sodium ^b | 0.54 ± 0.01 |
| Iron | 11.3 ± 1.2 | Zinc | 0.172 ± 0.017 |
| Potassium ^b | 1.26 ± 0.05 | | |

| <u>Trace Constituents</u> | | | |
|---------------------------|---|----------------|---|
| <u>Element</u> | <u>Content</u> <u>µg/g^a</u> | <u>Element</u> | <u>Content</u> <u>µg/g^a</u> |
| Cadmium | 10.2 ± 1.5 | Nickel | 45.8 ± 2.9 |
| Copper | 109 ± 19 | Thallium | 1.44 ± 0.07 |
| Cobalt ^b | 10.1 ± 0.6 | Thorium | 1.62 ± 0.22 |
| Lead | 714 ± 28 | Uranium | 1.11 ± 0.05 |
| Manganese | 785 ± 97 | Vanadium | 23.5 ± 6.9 |
| Mercury | 1.1 ± 0.5 | | |

^aThe uncertainties of the certified values for the elements, except those noted by superscript "b," include those errors associated with both measurement and material variability. They represent the 95 percent tolerance limits for an individual sub-sample, i.e., 95 percent of the sub-samples from a unit of this SRM would be expected to have a composition within the indicated range of values 95 percent of the time.

^bThese elements are certified as a part of the NBS update certification program. For each element a "best value" is given based on all methods of measurement that were used as well as a *standard error* of this value. Both are based on considerations of variability both within and between analytical methods.

Supplemental Information

Note: The following values are not certified because they are not based on the results of either a reference method or of two or more independent methods. These values are included for information only.

Table 2. Noncertified Values for Constituent Elements

| <u>Element</u> | <u>Content wt. Percent</u> | <u>Element</u> | <u>Content µg/g</u> |
|----------------|--------------------------------|----------------|-------------------------|
| Calcium | (2.9) | Antimony | (51) |
| Fluorine | (0.09) | Arsenic | (66) |
| Sulfur | (1.1) | Lanthanum | (9) |
| | | Scandium | (2) |
| | | Selenium | (1.5) |

Additional Information: The values listed below are based on measurements made in one laboratory and while no reason exists to suspect systematic bias in these numbers, no attempt was made to evaluate such bias attributable to either the method or the laboratory. The method used for each set of measurements is also listed. The indicated uncertainties are two times the standard deviation of the mean. These values are included for information only.

Table 3

| <u>Constituent</u> | <u>Content wt. percent</u> |
|-------------------------------------|--------------------------------|
| Kjeldahl Nitrogen | (0.0797% ± 0.0048) |
| Total Phosphorus | (0.051% ± 0.001) [1] |
| Loss on Ignition (800 °C) | (10.72% ± 0.28) |
| Oil and Grease (Freon) | (1.71% ± 0.26) [3] |
| Chemical Oxygen Demand (Dichromate) | (149,400 mg/kg ± 9,000)[2] |

References

1. ASTM Method E-350
2. Standards Methods for the Examination of Water and Waste Water, 14th Edition (1975), Section 508, pp 550.
3. Ibid., Section 502, pp 518.

Table 3A Methods and Analysts

| Method/ Element | A | B | C | D | E | F |
|--------------------|---|---|---|---|---|---|
| Aluminum | • | | • | | • | |
| Arsenic | | | • | | | |
| Antimony | • | | | | | |
| Cadmium | | | • | • | | |
| Calcium | | | • | | | |
| Chromium | | • | • | | | |
| Cobalt | | | • | | • | |
| Copper | | • | • | | | |
| Fluorine | | | | | | • |
| Iron | • | | • | | | |
| Lanthanum | | | • | | | |
| Lead | | • | | • | | |
| Magnesium | • | | | | • | |
| Manganese | | • | • | | | |
| Mercury | • | | • | | | |
| Nickel | | • | | • | | |
| Potassium | • | | • | | | |
| Scandium | | | • | | | |
| Selenium | • | | | | | |
| Sodium | • | | • | | | |
| Sulfur | | | | | | • |
| Thallium | | • | | | | |
| Thorium | | • | | | | |
| Uranium | | • | | | | |
| Vanadium | • | | • | | | |
| Zinc | • | | | • | | |

Analytical Methods

- A. Atomic Absorption Spectrometry
- B. Isotope Dilution Mass Spectrometry
- C. Neutron Activation Analysis
- D. Polarography
- E. D. C. Plasmas Atomic Emission Spectrometry
- F. Ion Chromatography

Analysts

NBS Center for Analytical Chemistry

- | | |
|--------------------|---------------------|
| 1. T. J. Brady | 10. R. R. Greenberg |
| 2. E. R. Deardorff | 11. S. H. Harrison |
| 3. L. P. Powell | 12. G. J. Lutz |
| 4. M. S. Epstein | 13. L. A. Machlan |
| 5. R. Filby | 14. E. J. Maienthal |
| 6. M. Gallorini | 15. T. C. Rains |
| 7. E. L. Garner | 16. H. L. Rook |
| 8. T. E. Gills | 17. T. A. Rush |
| 9. J. W. Gramlich | 18. W. P. Schmidt |

Cooperators

19. L. Kosta (Nuclear Chemistry Section, Josef Stefan Institute, Ljubljana, Yugoslavia)

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1646

Estuarine Sediment

This Standard Reference Material is intended primarily for calibrating instrumentation and evaluating the reliability of analytical methods for the determination of major, minor, and trace elements in sediments, and similar matrices.

Values of Constituent Elements: The *certified* values for the constituent elements are shown in Table 1. They are based on results obtained either by definitive methods or by two or more independent, reliable analytical methods. *Non-certified values*, which are given for information only, appear in Table 2. All values are based on a minimum sample size of 500 mg of the material dried as indicated under "Instructions for Drying".

Notice to Users:

Expiration of Certification: The certification of this SRM will be invalid 5 years after date of shipping.

Use: The material should be kept in its original bottle and shaken well before each use. A minimum sample of 500 mg of the dried material (see Instructions for Drying) should be used for any analytical determination to be related to a certified value of this certificate.

Statistical consultation was provided by K. R. Eberhardt of the Statistical Engineering Division.

The overall direction and coordination of the technical measurements leading to certification were performed in the Inorganic Analytical Research Division, E. L. Garner, Chief.

The technical and support aspects involved in the preparation, certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by R. Alvarez.

Washington, D.C. 20234
June 7, 1982
(Revision of Certificate
dated 1-6-82)

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

Table 1. Certified Concentration of Constituent Elements

| Element | Concentration, weight % | Element | Concentration, weight % |
|-------------------------------|----------------------------|------------------------------|----------------------------|
| Aluminum ^{2b,c;6} | 6.25 ± 0.20 | Magnesium ^{1c;2c} | 1.09 ± 0.08 |
| Calcium ^{2b,c;6} | 0.83 ± 0.03 | Phosphorus ^{2a,6} | 0.054 ± 0.005 |
| Iron ^{2c;4a;6} | 3.35 ± 0.10 | | |
| Element | Concentration, µg/g | Element | Concentration, µg/g |
| Arsenic ^{1d;4b} | 11.6 ± 1.3 | Manganese ^{1c;2c} | 375 ± 20 |
| Cadmium ^{1b,3a,b;4b} | 0.36 ± 0.07 | Mercury ^{1a;4b} | 0.063 ± 0.012 |
| Chromium ^{1c;3b;4a} | 76 ± 3 | Nickel ^{1b;2c;5} | 32 ± 3 |
| Cobalt ^{1b;4a} | 10.5 ± 1.3 | Vanadium ^{2a,3a} | 94 ± 1 |
| Copper ^{1c;2c;4b} | 18 ± 3 | Zinc ^{1b,c;2c;3b;5} | 138 ± 6 |
| Lead ^{1b;3a;5} | 28.2 ± 1.8 | | |

1. Atomic absorption spectrometry

- a. cold vapor
- b. graphite furnace
- c. flame
- d. hydride generation

2. Atomic emission spectrometry

- a. dc plasma
- b. flame
- c. inductively coupled plasma

3. Isotope dilution mass spectrometry

- a. thermal ionization
- b. spark source

4. Neutron activation

- a. instrumental
- b. radiochemical

5. Polarography

6. X-ray fluorescence spectrometry

Notes: (1.) Analytical values are based on the "dry-weight" of material (see Instructions for Drying). Mercury should be determined on samples without drying and the results adjusted to a "dry-weight" basis by determining the moisture content of separate samples.

(2.) The estimated uncertainty for an element is based on judgment and represents an evaluation of the combined effects of method imprecision, possible systematic errors among methods, and material variability for samples 500 mg or more.

Table 2. Non-certified Concentrations of Constituent Elements

Note: The values shown in this table are not certified because they are not based on the results of either a definitive method or two or more independent analytical methods. These values are included, for information only, to provide additional information on the composition.

| <u>Element</u> | <u>Concentration, Weight %</u> | <u>Element</u> | <u>Concentration, Weight %</u> |
|----------------|--|----------------|--|
| Potassium | (1.4) | Sulfur | (0.96) |
| Silicon | (31) | Titanium | (0.51) |
| Sodium | (2.0) | | |
| <u>Element</u> | <u>Concentration, $\mu\text{g/g}$</u> | <u>Element</u> | <u>Concentration, $\mu\text{g/g}$</u> |
| Antimony | (0.4) | Molybdenum | (2.0) |
| Beryllium | (1.5) | Rubidium | (87) |
| Cerium | (80) | Scandium | (10.8) |
| Cesium | (3.7) | Selenium | (0.6) |
| Europium | (1.5) | Tellurium | (0.5) |
| Germanium | (1.4) | Thallium | (0.5) |
| Lithium | (49) | Thorium | (10) |

Analysts:

Inorganic Analytical Research Division, National Bureau of Standards. I. L. Barnes, M. B. Blackburn, C. G. Blundell, T. A. Butler, M. S. Epstein, T. E. Gills, J. W. Gramlich, R. R. Greenberg, S. Hanamura, W. R. Kelly, H. M. Kingston, L. Machlan, E. J. Maienthal, J. D. Messman, T. J. Murphy, T. C. Rains, T. A. Rush, R. Sedivy, and R. L. Watters, Jr.

Cooperating Analysts:

University of Tokyo, Tokyo, Japan; present address: Meteorological Research Institute; Tsukuba, Ibaraki, Japan; Y. Dokiya (NBS Guest Worker).

Division of Chemistry, National Research Council of Canada, Ottawa, Canada; S. Berman, A. Desaulniers, R. Sturgeon, A. Mykytuik, J. McLaren, V. Boyko, and P. Semeniuk.

Instructions for Drying: Except for mercury, elements should be determined on samples that have been dried at 110 °C for 2 hours.

Mercury should be determined on undried samples. However, because the certified concentration is reported on a "dry-weight" basis, the concentration determined on undried samples should be adjusted for the moisture content of the samples.

Source and Preparation of Material: The material for this SRM was supplied by R. Huggett, Virginia Institute of Marine Sciences, Gloucester Point, Va. It had been dredged from the Chesapeake Bay at a location: 37° 11.1' N, 76° 17.1' W. The material was freeze-dried at Eastern Freeze-Dry Corporation, Lancaster, Pa., and radiation sterilized at Neutron Products Inc., Dickerson, Md. At NBS, the sediment was sieved through a screen with openings of 1.00 mm (No. 18) to remove coarse contaminants; ball-milled to pass a sieve with openings of 150 μm (No. 100); thoroughly mixed in a V-blender; placed in polyethylene bags; and bottled.

Homogeneity Assessment: A preliminary evaluation of homogeneity was made by instrumental neutron activation using samples of approximately 250 mg taken from various locations of the bulk materials. The samples were irradiated and the activities from radionuclides of Ce, Co, Cr, Cs, Eu, Fe, Rb, Sc and Th were counted. Except for Ce and Th, the observed sample-to-sample variations for the elements were approximately the same as the counting statistics indicating satisfactory homogeneity for these elements within approximately 2%. The homogeneity of the material for As, Cd, Hg, N, and Zn was evaluated by various analytical techniques using samples weighting 250 to 300 mg and found to be satisfactory. The homogeneity of the remaining certified elements was determined using sample weights not exceeding one gram.

The uncertainties of the elemental concentrations in Table 1 take into account possible material inhomogeneity for samples weighing 500 mg.

National Bureau of Standards

Certificate of Analysis

Standard Reference Material 1648

Urban Particulate Matter

This Standard Reference Material (SRM) is intended for use in the calibration of apparatus and evaluation of methods used in the analysis of atmospheric particulate matter and materials with a similar matrix.

Certified Values of Constituent Elements: The certified values for the constituent elements are shown in Table 1. The analytical techniques used in the certification are shown in Table 3. The certified values are based on measurements of 6 to 30 samples by each of the analytical techniques indicated. Noncertified values are given for information only in Table 2.

Notice and Warnings to Users: This material is a naturally occurring urban dust to be used for analytical purposes only. It may contain a number of chemicals of unknown toxicities, therefore, the utmost caution and care must be exercised in its use.

Expiration of Certification: This certification is invalid after 5 years from date of purchase. Should it be shown to be invalid prior to that time, users will be notified by NBS.

Stability: This material should be kept in its original bottle and stored at temperatures between 10-30 °C. It should not be exposed to intense source of radiation, including ultraviolet lamps or sunlight. Ideally, the bottle should be kept in a desiccator at the recommended temperature.

Use: A minimum of 100 mg of the dried material (See Drying Instructions) should be used for any analytical determination to be related to the certified values of this certificate.

Source and Preparation of Material: This SRM was prepared from urban particulate matter collected in the St. Louis, Missouri, area in a baghouse specially designed for this purpose. The material was removed from the filter bags and combined in a single lot. This product was screened through a fine-mesh sieve to remove extraneous materials and thoroughly blended in a V-blender. The material was then bottled and sequentially numbered. The material was collected over a period in excess of 12 months and, therefore, is a time-integrated sample. While not represented to be typical of the area in which it was collected, its use should typify the analytical problems of atmospheric samples obtained from industrialized urban areas.

Homogeneity Assessment: Randomly selected bottles were used for the analytical measurements. Each analyst examined at least 6 bottles, some of them measuring replicates from each bottle. No correlation was found between measured values and the bottling sequence. Also, the results of measurements of samples from different bottles were not significantly different than the measurements of replicate samples from single bottles. Accordingly, all bottles of this SRM have been assigned the same certified values of constituent elements.

Instructions for Drying: This material should be dried at 105 °C for 8 hours before use. Because the certified concentrations are reported on a "dry-weight" basis, the concentrations determined on undried samples should be adjusted for the moisture content of the samples.

The overall direction and coordination of the technical measurements leading to certification were performed under the chairmanship of J.K. Taylor.

The technical and support aspects involved in the preparation, current and previous certification, and issuance of this Standard Reference Material were coordinated through the Office of Standard Reference Materials by T.E. Gills and W.P. Reed.

Washington, D.C. 20234
May 11, 1982
(Revision of Certificate
dated 11-16-78)

George A. Uriano, Chief
Office of Standard Reference Materials

(over)

Table 1. Certified Values of Constituent Elements

| <u>Major Constituents</u> | | <u>Minor Constituents</u> | |
|---------------------------|--|---------------------------|--|
| <u>Element</u> | Content ^a <u>Wt. Percent</u> | <u>Element</u> | Content ^a <u>Wt. Percent</u> |
| Aluminum ^b | 3.42 ± 0.11 | Lead | 0.655 ± 0.008 |
| Iron | 3.91 ± 0.10 | Sodium ^b | 0.425 ± 0.002 |
| Potassium ^b | 1.05 ± 0.01 | Zinc | 0.476 ± 0.014 |

Trace Constituents

| <u>Element</u> | Content ^a <u>µg/g</u> | <u>Element</u> | Content ^a <u>µg/g</u> |
|----------------|-------------------------------------|-----------------------|-------------------------------------|
| Arsenic | 115 ± 10 | Nickel | 82 ± 3 |
| Cadmium | 75 ± 7 | Selenium ^b | 27 ± 1 |
| Chromium | 403 ± 12 | Uranium | 5.5 ± 0.1 |
| Copper | 609 ± 27 | Vanadium ^b | 140 ± 3 |

^aThe uncertainties shown for the elements except those noted by superscripts include errors associated with both measurement and material variability. They represent the 95 percent tolerance limits for individual subsamples, i.e., 95 percent of the subsamples from a single unit of this SRM would be expected to have a composition within the indicated range of values 95 percent of the time.

^bThese elements were recently certified as a part of the NBS update certification program. The values for the indicated constituent are the "best value" based on all measurement methods used and the associated uncertainty is expressed as the standard error considering variability within and between analytical methods.

Table 2. Noncertified Values for Constituent Elements

Note: The following values are not certified because they are not based on the results of either a reference method or two or more independent methods. These values are included for information only.

| <u>Major Constituents</u> | | <u>Minor Constituents</u> | |
|---------------------------|-------------------------------|---------------------------|-------------------------------|
| <u>Element</u> | Content <u>Wt. Percent</u> | <u>Element</u> | Content <u>Wt. Percent</u> |
| Sulfur | (5.0) | Chlorine | (0.45) |
| Magnesium | (0.8) | Titanium | (0.40) |

| <u>Trace Constituents</u> | | | |
|---------------------------|------------------------|----------------|------------------------|
| <u>Element</u> | Content <u>µg/g</u> | <u>Element</u> | Content <u>µg/g</u> |
| Antimony | (45) | Lanthanum | (42) |
| Barium | (737) | Rubidium | (52) |
| Bromine | (500) | Manganese | (860) |
| Cerium | (55) | Samarium | (4.4) |
| Cesium | (3) | Scandium | (7) |
| Cobalt | (18) | Silver | (6) |
| Europium | (0.8) | Thorium | (7.4) |
| Hafnium | (4.4) | Tungsten | (4.8) |
| Indium | (1.0) | | |
| Iodine | (20) | | |

Supplemental Information

The values listed below are based on measurements made in a single laboratory and are given for information only. While no reason exists to suspect systematic bias in these numbers, no attempt was made to evaluate such bias attributable to either the method or the laboratory. The method used for each set of measurements is also listed. The uncertainties indicated are two times the standard deviation of the means.

| <u>Constituent</u> | <u>Content</u> <u>Wt. Percent</u> |
|-----------------------------|--------------------------------------|
| Nitrogen (NO ₃) | (1.07 ± 0.06) |
| Nitrogen (NH ₄) | (2.01 ± 0.08) |
| Sulfate | (15.42 ± 0.14) |
| SiO ₂ | (26.8 ± 0.38) |
| Freon Soluble | (1.19 ± 0.47) |

Methods Used:

Nitrate - Extraction with water and measurement by ASTM Method D992

Ammonia - NaOH addition followed by steam distillation and titration

Sulfate - Extraction with water and measurement by ASTM D516

SiO₂ - Solution and measurement by ASTM Method E350

Freon Soluble - Extraction with Freon 113, using the Method described in "Standard Methods in Examination of Water and Waste Water," 14th Edition, p. 518, American Public Health Association, Washington, D.C.

Analysts

Inorganic Analytical Chemistry Division

- | | |
|-----------------------|---------------------|
| 1. C. G. Blundell | 9. J. W. Gramlich |
| 2. R. W. Burke | 10. R. R. Greenberg |
| 3. T. A. Butler | 11. L. A. Machlan |
| 4. E. R. Deardorff | 12. E. J. Maienthal |
| 5. B. I. Diamondstone | 13. T. J. Murphy |
| 6. M. S. Epstein | 14. L. P. Powell |
| 7. M. Gallorini | 15. T. C. Rains |
| 8. E. L. Garner | 16. T. A. Rush |

Table 3 Methods and Analysis

| Method/ Element | A | B | C | D | E | F | G | H | I |
|--------------------|---|---|---|---|---|---|---|---|---|
| Ag | | | • | | | | | | |
| Al | | | • | | | | | • | |
| As | | | • | | • | | | | |
| Ba | | | • | | | | | | |
| Br | | | • | | | | | | |
| Cd | • | • | • | • | | | | | |
| Ce | | | • | | | | | | |
| Cl | | | | | | | • | | |
| Co | | | • | | | | | | |
| Cr | | • | • | | | | | | |
| Cs | | | • | | | | | | |
| Cu | • | • | | | • | | | | |
| Eu | | | • | | | | | | |
| Fe | • | • | • | | • | | | | |
| Hf | | | • | | | | | | |
| I | | | • | | | • | | | |
| In | | | • | | | | | | |
| K | • | | • | | | | | | |
| La | | | • | | | | | | |
| Mg | | | • | | | | | | |
| Mn | • | | • | | | | | | • |
| Na | • | | • | | | | | | • |
| Ni | • | • | | • | | | | | |
| Pb | • | • | | • | | | | | |
| Rb | | | • | | | | | | |
| S | | | | | | | • | | |
| Sb | | | • | | | | | | |
| Sc | | | • | | | | | | |
| Se | • | | • | | | | | | • |
| Sm | | | • | | | | | | |
| Th | | | • | | | | | | |
| Ti | | | • | | | | | | |
| U | | • | | | | | | | |
| V | • | | • | | | | | | |
| W | | | • | | | | | | |
| Zn | • | • | • | • | | | | | |

Analytical Methods

- A. Atomic Absorption Spectrometry
- B. Isotope Dilution Mass Spectrometry
- C. Neutron Activation Analysis
- D. Polarography
- E. Spectrophotometry
- F. Photon Activation Analysis
- G. Ion Chromatography
- H. D.C. Plasma Atomic Emission Spectrometry
- I. Flame Emission Spectrometry

Guide for Requesting Development of Standard Reference Materials

The National Bureau of Standards develops Standard Reference Materials (SRM's) to provide a basis for comparison of measurements on materials and to aid in the control of production processes. The Office of Standard Reference Materials evaluates the requirements of science, industry, and government for carefully characterized reference materials, then directs the production and distribution of these materials.

NBS currently has over 1000 SRM's available, about 100 new ones in preparation, and requests for the development of many more. The demand for new SRM's greatly exceeds the Bureau's capacity to produce and certify these materials. Consequently, requests for new SRM's of limited use are deferred in favor of those that serve a substantial area of interest. In determining which requests receive top priority, NBS relies heavily upon information supplied by industry and interested organizations.

The Bureau welcomes all requests for SRM's. Both the Bureau and potential users would be helped if these requests included as much of the information below as possible.

- 1.** Short title of the proposed Standard Reference Material.
- 2.** Purpose for which this SRM is intended.
- 3.** Reason why the SRM would be useful.
- 4.** Special characteristics and/or requirements of the material. Include necessary additional information, if more than one SRM is needed for standardization in an area.
- 5.** An estimate of the possible present and future (6-10 years) demand for such an SRM in your operations and elsewhere. (National and international estimates are very useful).

6. Facts about whether such an SRM (or a similar one) could be produced by, or obtained from a source other than NBS. If so, justify preparation by NBS.

7. Other pertinent information to justify the SRM, such as: (a) an estimate of the range of application, economic significance of the measurement affected, and scientific and/or technological significance, including estimates of the impact upon industrial productivity or growth, and (b) supporting letters from industry leaders, trade organizations, interested committees, and others.

In developing an NBS-SRM, the candidate material must meet one or more of the criteria listed below:

- 1.** The SRM must permit users to attain more accurate measurements.
- 2.** The production of the SRM must not be economically or technically feasible elsewhere,
- 3.** The SRM must serve as an industry-wide standard for commerce, provided by a unique neutral source,
- 4.** NBS production of the SRM would provide readily available, highly characterized material useful to science, industry, or government.

NBS has recognized the need to enlarge the scope of the SRM program to include all types of well-characterized materials that can be used to calibrate a measurement system, or to produce scientific data that can be widely used. Input from science, industry, and government assists NBS in continuing to provide Standard Reference Materials that will be valuable in many areas.

GUIDE TO ORDERING STANDARD REFERENCE MATERIALS

ORDERING

Orders should be addressed to:
Office of Standard Reference Materials
Room B311, Chemistry Building
National Bureau of Standards
Gaithersburg, MD 20899
Telephone: (301) 921-2045

Orders should give number of units, catalog number, and name of the material requested. For example, 1 each, No. 11h, Basic-Open-Hearth Steel, 0.2 percent C. The materials described in this Catalog are distributed only in the units listed or in multiples thereof.

Acceptance of an order does not imply acceptance of any provision set forth in the order contrary to the policy, practice, or regulations of the National Bureau of Standards or the U.S. Government.

Orders received for "out-of-stock" materials are cancelled if only out-of-stock items are ordered. On other orders, shipment is made of available materials and out-of-stock items are cancelled. Back-orders are not accepted for out-of-stock materials; if a renewal lot of material is available, it will be furnished automatically.

TERMS

Prices quoted are in U.S. dollars, and are published in the SRM Price List. When SRM Price Lists are issued they are sent to persons or organizations who have requested them. These prices are subject to revision without notice and orders will be billed for the prices in effect at the time of shipment. No discounts are given on purchases of SRM's, RM's, or GM's.

Remittances of the purchase price need not accompany purchase orders. Payment of invoices is expected within 30 days of receipt of an invoice. Payment on foreign orders may be made by any of the following:

- a. banker's draft against U.S.A. bank
- b. bank to bank transfer to U.S.A. bank
- c. cash against documents
- d. sight draft
- e. International Money Order
- f. UNESCO coupons

Letters of Credit cannot be accepted. If a Letter of Credit or any method of payment other than those listed above is to be used, you must secure the services of an agent in the United States to act in your behalf. Your agent would purchase the material and our invoice would indicate that he is the purchaser. The material would be shipped to your agent, who would transship in accordance with your instructions.

NBS cannot "prepay and add" shipping charges to the invoice. Restricted categories such as hydrocarbons, organic sulfur compounds, compressed gasses, rubber compounding materials, radioactive standards, and similar materials are shipped FOB Gaithersburg, MD.

LATE CHARGES

Unless otherwise notified, payment for SRM's is due within 30 days of shipment of the order to the customer. For non-Federal customers, the U.S. Treasury regulations require late charges, based on the current value of funds to Treasury, be assessed for each 30-day period or portion thereof that the payment is overdue.

PROFORMA INVOICE (PRICE QUOTATION)

Proforma invoice service will frequently require three to four weeks to process, and will be furnished only to those requiring such service.

DOMESTIC SHIPMENTS

Shipments of material (except for certain restricted categories) intended for the United States and Canada are normally shipped prepaid (providing that the parcel does not exceed the weight limitations as prescribed by postal laws and regulations).

FOREIGN SHIPMENTS

The regulations of various nations covering the importation of SRM's, GM's, and RM's differ widely; any attempt to list all possible variations would be impractical. Therefore, where the shipping practices outlined below do not apply, purchasers will be informed of the best method of shipment for their countries.

Most orders will be shipped by prepaid International Air Parcel Post. Exceptions are items in restricted categories and those shipments that exceed parcel post weight limitations. These exceptions will be shipped FOB Gaithersburg, MD, unless an agent (shipping or brokerage firm) located in the United States is required. Where an agent is required, the purchaser will be so notified and will be requested to designate an agent of his/her choice. In this case, the material will be packaged for overseas shipment and will be forwarded to the agent FOB Gaithersburg, MD.

DOCUMENTATION

Listed below are the only documents that we will furnish. All documents are printed in English.

- a. six commercial invoices
- b. two sight drafts
- c. two packing slips
- d. customs invoices for Canada, New Zealand, Australia, and South Africa
- e. Certificate of Origin
- f. parcel post receipts for parcel post shipments
- g. air waybill for air shipments

If documents other than those listed above are required, the services of an agent in the United States will be needed to purchase and ship the materials.

Note: Orders and inquiries submitted in English will be processed more rapidly than those requiring translations.

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