ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 136

[EPA-HQ-OW-2022-0901; FRL 9346-02-OW]

RIN 2040-AG25

Clean Water Act Methods Update Rule for the Analysis of Effluent

AGENCY: Environmental Protection Agency (EPA). **ACTION:** Final rule.

SUMMARY: The U.S. Environmental Protection Agency (EPA) is finalizing changes to its test procedures required to be used by industries and municipalities when analyzing the chemical, physical, and biological properties of wastewater and other samples for reporting under the EPA's National Pollutant Discharge Elimination System permit program. The Clean Water Act requires the EPA to promulgate these test procedures (analytical methods) for analysis of pollutants. The EPA anticipates that these changes will provide increased flexibility for the regulated community in meeting monitoring requirements

while improving data quality. In addition, this update to the CWA methods will incorporate technological advances in analytical technology and make a series of minor changes and corrections to existing approved methods. As such, the EPA expects that these changes will not result in any negative economic impacts.

DATES: This final rule is effective on June 17, 2024. The incorporation by reference of certain material listed in this rule and is approved by the Director of the Federal Register as of June 17, 2024.

ADDRESSES: The EPA has established a docket for this action under Docket ID No. EPA-OW-HQ-2022-0901. All documents in the docket are listed on the http://www.regulations.gov website. Although listed in the index, some information is not publicly available, e.g., CBI or other information whose disclosure is restricted by statute. Certain other material, such as copyrighted material, is not placed on the internet and will be publicly available only in hard copy form. Publicly available docket materials are available electronically through http:// www.regulations.gov.

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I. General Information

This preamble describes the abbreviations and acronyms; reasons for the rule; and a summary of the changes and clarifications; the legal authority for the rule; methods incorporated by reference; and a summary of the changes and clarifications.

A. Does this action apply to me?

Entities potentially affected by the requirements of this action include:

Category	Examples of potentially affected entities
State, Territorial, and Indian Tribal Governments.	States authorized to administer the National Pollutant Discharge Elimination System permitting program; states, territories, and Tribes providing certification under CWA section 401; state, territorial, and Tribal- owned facilities that must conduct monitoring to comply with NPDES permits.
Industry	Facilities that must conduct monitoring to comply with NPDES permits; the environmental monitoring indus- try.
Municipalities	Publicly Owned Treatment Works or other municipality-owned facilities that must conduct monitoring to comply with NPDES permits.

This table is not intended to be exhaustive, but rather provides a guide for readers regarding entities likely to be affected by this action. This table lists types of entities that the EPA is now aware of that could potentially be affected by this action. Other types of entities not listed in the table could also be affected. To determine whether your facility is affected by this action, you should carefully examine the applicability language at 40 CFR 122.1 (NPDES purpose and scope), 40 CFR 136.1 (NPDES permits and CWA) and 40 CFR 403.1 (pretreatment standards purpose and applicability). If you have questions regarding the applicability of this action to a particular entity, consult the appropriate person listed in the preceding FOR FURTHER INFORMATION CONTACT section.

B. What action is the Agency taking?

Periodically, the EPA updates the approved methods in 40 CFR part 136.

In general, the changes in this action fall into four categories. The first category is updated versions of EPA methods currently approved in 40 CFR part 136. The second category is new or revised methods published by a voluntary consensus standard body that are similar to methods previously adopted as EPA-approved methods in 40 CFR part 136. The third category is methods the EPA has reviewed under the agency's national Alternate Test Procedure program and preliminarily concluded are appropriate for nationwide use. The fourth category is corrections or amendments to the text and tables of 40 CFR part 136. The EPA is finalizing these revisions to improve data quality, update methods to keep current with technology advances, and provide the regulated community with greater flexibility. The following paragraphs provide details on the revisions.

C. What is the Agency's authority for taking this action?

The EPA is promulgating this regulation under the authorities of sections 301(a), 304(h), and 501(a) of the CWA; 33 U.S.C. 1251, 1311(a), 1314(h) and 1361(a). Section 301(a) of the CWA prohibits the discharge of any pollutant into navigable waters unless the discharge complies with, among other provisions, an NPDES permit issued under section 402 of the CWA. Section 304(h) of the CWA requires the EPA Administrator to ". . . promulgate guidelines establishing test procedures for the analysis of pollutants that shall include the factors which must be provided in any certification pursuant to [section 401 of the CWA] or permit application pursuant to [section 402 of the CWA]." Section 501(a) of the CWA authorizes the Administrator to ". . prescribe such regulations as are necessary to carry out this function under [the CWA]." The EPA generally

has codified its test procedure regulations (including analysis and sampling requirements) for CWA programs at 40 CFR part 136, though some requirements are codified in other parts (*e.g.*, 40 CFR Chapter I, Subchapters N and O).

II. Background

Abbreviations and Acronyms Used in the Preamble

- ASTM: ASTM International¹
- ATP: Alternate Test Procedure
- BHI: Brain heart infusion
- CATC: Cyanide Amenable to Chlorination
- CFR: Code of Federal Regulations
- CNCl: Cyanogen Chloride
- CWA: Clean Water Act
- EC-MUG: EC broth with 4-
- methylumbelliferyl-β-D-glucuronide EDTA: Ethylenediaminetetraacetic acid
- EPA: the U.S. Environmental Protection Agency
- DO: Dissolved Oxygen
- GC: Gas Chromatography
- GC/MS/MS: Gas Chromatography-Tandem Mass Spectrometry
- GC/HRMS: Gas Chromatography-High Resolution Mass Spectrometry
- IBR: Incorporation by Reference
- ICP/AES: Inductively Coupled Plasma-Atomic Emission Spectroscopy
- NED: *N*-(1-naphthyl)-ethylenediamine dihydrochloride
- NPDES: National Pollutant Discharge Elimination System
- m/z: Mass to Charge Ratio
- MF: Membrane Filtration
- MPN: Most Probable Number
- nm: Nanometer
- NTTAA: National Technology Transfer and Advancement Act
- POTW: Publicly Owned Treatment Works QC: Quality Control
- TKN: Total Kjeldahl Nitrogen
- USGS: United States Geological Survey
- VCSB: Voluntary Consensus Standards Body

NPDES permits must include conditions designed to ensure compliance with the technology-based and water quality-based requirements of the CWA, including in many cases, restrictions on the quantity of specific pollutants that can be discharged as well as pollutant measurement and reporting requirements. Often, entities have a choice in deciding which approved test procedure they will use for a specific pollutant because the EPA has approved the use of more than one method.²

The procedures for the analysis of pollutants required by CWA section 304(h) are a central element of the NPDES permit program. Examples of where these EPA-approved analytical methods must be used include the following: (1) applications for NPDES permits, (2) sampling or other reports required under NPDES permits, (3) other requests for quantitative or qualitative effluent data under the NPDES regulations, (4) state CWA 401 certifications, and (5) sampling and analysis required under the EPA's General Pretreatment Regulations for Existing and New Sources of Pollution, 40 CFR 136.1 and 40 CFR 403.12(b)(5)(v).

On February 21, 2023, the EPA proposed to update the approved methods in 40 CFR part 136. The EPA received 20 comments on the proposed rulemaking (February 21, 2023, 88 FR 10724) from laboratory associations, state environmental agencies, trade associations and citizens. All commenters supported finalizing this rule and approving each proposed method.

There were some specific comments that are outside the scope of this rulemaking. As stated in the proposed rule (88 FR 10725, February 21, 2023,), the EPA only considered new or revised methods that were submitted to the EPA. Method withdrawals, methods for new parameters, methods based on new technologies (except methods approved through the alternate test procedure program) and VCSB methods not submitted from VCSBs were not considered for this routine update. Commenters requesting changes to VCSB, or vender methods should work through the method owner to revise the method and submit any supporting data to the EPA for consideration.

Commenters noted that there was a format error in the proposed rulemaking language on Table IC, footnotes 15, 16 and 17, (88 FR 10763, February 21, 2023). The new footnote 16 was inadvertently added to the end of footnote 15. The information in the new footnotes is correctly described in the preamble (88 FR 10738, February 21, 2023). This typographical error of the order and numbering of the footnotes has been corrected in the final rule. In addition, the following parameters were missing from the preamble discussions of the revision of Standard Methods method 6410B-2020: 2,2'-oxybis(1chloropropane) (also referred to as bis[2-Chloro-1-methylethyl] ether); hexachloroethane; and Nnitrosodimethylamine. The revised 6410B-2020 discussion in Section IV.C.35 of this preamble is correct.

III. Corrections or Amendments to the Text and Tables of 40 CFR Part 136

In addition to the method revision incorporated by reference as discussed

in Section IV of this preamble, Standard Methods has revised a few of their general quality control sections (2020, 3020, 4020 and 5020). The EPA is updating the year of the current references to these sections in 136.3 Table IB footnote 85. The EPA is also adding a reference to an additional Standard Methods Quality Control section: Part 6000 Individual Organic Compounds, 6020. These Quality Control Standards are available for download at www.standardmethods.org at no charge. The EPA is correcting several minor errors or inconsistencies in the tables of approved methods. The EPA is making the following changes to 40 CFR 136.3, Tables IA, IB, IC, ID or IH:

1. Table IA. Removing the units of "number per 100 mL" under parameter 1. Coliform (fecal), because parameter 1 is specifically for biosolids that are reported as "number per gram dry weight".

2. Table IA. Moving United States Geologic Survey Method "B–0050–85" from parameter 1. Coliform (fecal) number per gram dry weight to parameter 2. Coliform (fecal) number per 100 mL, to address an error from the previous rulemaking when Parameter 1 Coliform (fecal) was split into two parameters to eliminate confusion as to which methods were approved for biosolids.

3. Table IA parameter 3 and IH parameter 2. Moving the phrase "twostep" in the "Method" column from the second to the third line which returns the phrase to the proper line after having been inadvertently moved.

4. Table IB. Revising footnote 85 to remove bullet formatting.

5. Table IB. Adding footnote 86 to Method 419D. Method 419D is listed as an approved method for of determination nitrate using Colorimetric (Brucine sulfate) methodology. This addition corrects a long-standing typographical error regarding the appropriate footnote for this method in Table IB.

6. Table IB. Correcting an inadvertent error to footnote 57. The reference number was incorrectly changed to 335.4–1. The correct number is 335.4.

7. Tables IC and ID. Adding footnote 15 to the Standard Methods column header and adding footnote 15 to refer to Quality Control Section: Part 6000 Individual Organic Compounds, 6020 (2019).

8. Table IC. Changing parameter 39, dichlorodifluoromethane, to refer to Method 6200 B rather than 6200 C for the GC/MS method.

9. Table IC. Adding footnote 10 to parameters 66–72, 95, 96 and 97 which

¹Formerly known as the American Society for Testing and Materials (ASTM).

² NPDES permit regulations also specify that the approved method needs to be sufficiently sensitive. See 40 CFR 122.21(e)(3).

were inadvertently dropped in an earlier rulemaking. Footnote 10 to table IC applies to all of the 17 dioxin and furan congeners.

10. Table IH parameter 2. Moving method B–0025–85 down one row because it was inadvertently moved in an earlier rulemaking. This method is a single step membrane filtration method rather than a most probable number method.

11. Table II. Revising footnote 5 for the preservation and holding time requirements for cyanide to add the year (2015) of the ASTM method D7365–09a (15).

The recommended sampling and preservation procedures in the ASTM method have not changed since 2009, but the change to footnote 5 simplifies identification of the current method that is available from ASTM International. The 2015 reapproval date was already updated in footnote 6 to Table II in the 2021 methods update rule; however, adding the reapproval date was overlooked in the incorporated by reference section and in footnote 5 to Table II.

IV. Incorporation by Reference

Currently, hundreds of methods and ATPs are incorporated by reference within 40 CFR part 136. In most cases, 40 CFR part 136 contains multiple approved methods for a single parameter (or pollutant) and regulated entities often have a choice in selecting a method. The rule contains revisions to VCSB methods that are currently incorporated by reference, (see Sections IV.B, IV.C, and IV.E of this preamble). Two VCSBs have made such revisions: Standard Methods and ASTM. The VCSB methods are consistent with the requirements of the National Technology Transfer and Advancement Act, under which Federal agencies use technical standards developed or adopted by the VCSBs if compliance would not be inconsistent with applicable law or otherwise impracticable. This rule also includes two vendor ATPs (see Section IV.D of this preamble) and four revised EPA methods (see Section IV.A of this preamble) which the EPA is incorporating by reference.

The rule incorporates by reference the methods added in an earlier Methods Update Rule (86 FR 27226, May 19, 2021). The EPA inadvertently failed to complete the incorporation by reference review process for that final rule. The EPA proposed 68 methods for incorporation by reference into 40 CFR 136.3 (84 FR 56590, October 22, 2019). Other than ASTM D7365–09a (Reapproved 2015) and the EPA Method 1623.1, the methods are described in the 2019 proposal as well as the 2021 final rule. ASTM D7365–09a (Reapproved 2015) and Method 1623.1 are summarized in this preamble.

The EPA is also incorporating by reference an errata sheet in Table IA, footnotes 25, 26, 27. The U.S. EPA Whole Effluent Toxicity (WET) Methods Errata Sheet, EPA 821–R–02–012–ES, corrects and clarifies the WET methods referenced in those footnotes. The errata sheet was described and promulgated as part of the 2017 Clean Water Act Methods Update Rule for the Analysis of Effluent (see 82 FR 40841, August 28, 2017; docket number EPA–HQ–OW– 2014–0797). The EPA inadvertently failed to incorporate by reference the errata sheet in the 2017 final rule.

The following paragraphs provide details on the methods incorporated by reference.

A. Changes to 40 CFR 136.3 To Include New Versions of Previously Approved EPA Methods

The EPA is adding revised versions of the EPA membrane filtration methods 1103.2, 1106.2, 1600.1, and 1603.1 found in Tables IA and IH. These methods were approved from 2002 to 2014. The EPA is also summarizing method 1623.1 that was added in the earlier rule (86 FR 27226, May 19, 2021) but not summarized. The revisions include standardizing language between the related methods, updating to reflect current lab practices and clarifying edits.

These methods each describe a membrane filter procedure for the detection and enumeration of either enterococci or *Escherichia coli* bacteria by their growth after incubation on selective media. These methods provide a direct count of bacteria in water samples based on the development of colonies on the surface of the membrane filter.

1. *E. coli.* Method 1103.2 describes a MF procedure for the detection and enumeration of *Escherichia coli* bacteria in ambient (fresh) water and is currently approved in Table IH. This is a two-step method which requires transferring the membrane filter after incubation on membrane-Thermotolerant Escherichia coli Agar (mTEC) to a pad saturated with urea substrate.

2. Enterococci. Method 1106.2 describes a MF procedure for the detection and enumeration of enterococci bacteria in ambient water and is currently approved in Table IH. This is a two-step method which requires transferring the membrane filter after incubation on membraneEnterococcus (mE) agar to Esculin Iron Agar (EIA) medium.

3. Enterococci. Method 1600.1 describes a MF procedure for the detection and enumeration of enterococci bacteria in ambient (fresh and marine) water and wastewater and is currently approved in Tables IA and IH. This is a single-step method that is a modification of EPA Method 1106.1 (mE–EIA). The membrane filter containing the bacterial cells is placed on membrane-Enterococcus Indoxyl- β -D-Glucoside Agar (mEI).

4. *E. coli.* Method 1603.1 describes a MF procedure for the detection and enumeration of thermotolerant *Escherichia coli* bacteria in ambient (fresh) waters and wastewaters using Modified membrane-Thermotolerant Escherichia coli Agar (modified mTEC) and is currently approved in Table IA and IH.

5. *Cryptosporidium and Giardia.* Method 1623.1 describes a method for the detection of *Cryptosporidium* and *Giardia* in ambient water by concentration immunomagnetic separation (IMS), and immunofluorescence assay (FA) microscopy. A water sample is filtered and the oocysts, cysts, and extraneous materials are retained on the filter. EPA Method 1623.1 includes updated acceptance criteria for IPR, OPR, and MS/MSD, and clarifications and revisions based on the use of EPA Method.

The EPA methods are available free of charge on our websites (*epa.gov/cwa-methods/approved-cwa-microbiologicaltest-methods*), therefore the EPA methods incorporated by reference are reasonably available.

B. Changes to 40 CFR 136.3 To Include New Versions of Approved ASTM Methods

The EPA is adding new versions of ASTM methods previously approved in 40 CFR part 136. These changes to currently approved ASTM methods in 40 CFR part 136 include minor clarifications and editorial changes. As an example, ASTM added text to the appropriate method scope sections to indicate that the method was developed in accordance with the "Decision on Principles for the Development of International Standards, Guides and Recommendations" issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee. None of these changes will affect the performance of the method. The following describes the changes to current ASTM methods that the EPA is including in 40 CFR part 136. Each entry contains (in the following order):

the parameter, the ASTM method number (the last two digits in the method number represent the year ASTM published the method), a brief description of the analytical technique, and a brief description of any minor procedural changes (if there are any) in this revision from the last approved version of the method. Method revisions that are only formatting in nature will have no description of the changes. The methods listed below are organized according to the table at 40 CFR part 136 in the order in which they appear.

ASTM methods can be purchased from *astm.org.* The price of ASTM standards is not fixed. The price generally ranges between \$50 and \$100 per method. ASTM also offers memberships or subscriptions that allow unlimited access to their methods. The ASTM methods incorporated by reference are reasonably available.

The EPA is adding the following ASTM methods found in Table IB, and Table II at 40 CFR part 136:

1. Dissolved Oxygen. D888-18 (A, B, C), Dissolved Oxygen, Winkler, Electrode, Luminescent-based Sensor. Standard D888–18A measures dissolved oxygen using the Winkler iodometric titration procedure. The volume of titrant used is proportional to the concentration of DO in the sample. Standard D888–18B measures DO in the sample with an electrochemical probe that produces an electrical potential which is logarithmically proportional to the concentration of DO in the sample. Standard D888-18C measures DO with a luminescence-based sensor probe that employs frequency domain lifetimebased luminescence quenching and signal processing.

2. Hydrogen Ion (pH). In D1293–18 (A, B), pH, Electrometric. The activity of hydrogen ion (H+) in the sample is determined electrometrically with an ion-selective electrode in comparison to at least two standard reference buffers and pH is reported as the negative log of that activity.

3. Metals Series. In D1976–20, Elements in Water by Inductively-Coupled Plasma Atomic Emission Spectroscopy for determination of aluminum, antimony, arsenic, beryllium, boron, cadmium, chromium, cobalt, copper, iron, lead, magnesium, manganese, molybdenum, nickel, selenium, silver, thallium, vanadium, and zinc. The sample is acid digested and analyzed by ICP/AES for the simultaneous or sequential determination of 29 elements. The changes include changing the initial instrument calibration from using four standards as the first option to using

only one standard and a calibration blank.

4. Surfactants. In D2330–20, Methylene Blue Active Substances, the sample is mixed with an acidic aqueous solution of methylene blue reagent, which forms a blue-colored ion pair with any anionic surfactants which is subsequently extracted with chloroform and washed with an acidic solution to remove interferences. The intensity of the blue color is measured using a photometer at 650 nanometers. The concentration of methylene blue active substances is determined in comparison to a standard curve.

5. Residue, filterable and nonfilterable. In D5907–18 (A and B), Filterable Matter (Total Dissolved Solids) and Nonfilterable Matter (Total Suspended Solids) under Test Method A, an aliquot of the sample is filtered through a glass fiber filter and the solids trapped on the filter are dried at 105 °C and weighed to determine the nonfilterable material (total suspended solids) by difference. Under Test Method B, the filtrate from Test Method A, or a separate filtrate, is evaporated to dryness at 180 °C and the residue weighed to determine the total dissolved solids.

6. Cyanide—Free. In D7237-18, Free Cyanide, Flow Injection, followed by Gas Diffusion Amperometry an aliquot of the sample is introduced into a flow injection analysis instrument, where it mixes with a phosphate buffer to release hydrogen cyanide which diffuses through a hydrophobic gas diffusion membrane into an alkaline solution and is detected amperometrically with a silver electrode. This version also added new information about sulfide interferences and potential mitigation strategies that the EPA anticipates will improve data quality. There are no other procedural changes.

7. Cyanide—Total. In D7284–20, Total Cyanide, Manual Distillation with MgCl₂ followed by Flow Injection, Gas Diffusion Amperometry, the sample is distilled with acid and a magnesium chloride catalyst to release cyanide to a sodium hydroxide solution. An aliquot of the sodium hydroxide solution is introduced into a flow injection analysis instrument, where it is acidified, and the hydrogen cyanide diffuses through a hydrophobic gas diffusion membrane into an alkaline solution and is detected amperometrically with a silver electrode.

8. Cyanide. D7365–09a (Reapproved 2015) is applicable for the collection and preservation of water samples for the analysis of cyanide. Samples are collected in appropriate containers and mitigated for known interferences either in the field during sample collection or in the laboratory prior to analysis. The sampling, preservation and mitigation of interference procedures described in this practice are recommended for the analysis of total cyanide, available cyanide, weak acid dissociable cyanide, and free cyanide by ASTM Methods D2036, D4282, D4374, D6888, D6994, D7237, D7284, and D7511.

9. Organic Carbon. In D7573–18a^{e1}, Total Organic Carbon, Combustion, the sample is sparged with an inert gas to remove dissolved inorganic carbon, acidified, and then combusted at high temperature to convert organic carbon to carbon dioxide. The carbon dioxide is measured with an infra-red detector. This version also adds data from an interlaboratory method validation study and new method detection limit values, but there are no procedural changes.

C. Changes to 40 CFR 136.3 To Include New Versions of Approved "Standard Methods" Methods

The EPA is approving new versions of methods developed by the Standard Methods Committee that were previously approved in 40 CFR part 136. Standard Methods has reviewed many of their methods in preparation for releasing the next edition of "Standard Methods for the Examination of Water & Wastewater." The newer versions provide clarifications and make editorial corrections. These edits include removal of referents to specific brand names and trademarks, incorporation of footnotes into the text, a reformatting of figures, tables and reference lists, removal of bibliographical references that are no longer available, small editorial changes based on current style guides and changes to scientific publishing standards, and minor clarifications to procedures based on input from users. For example, the revisions replace distilled water with reagent water in all methods.

Each entry contains the Standard Method number and date, the parameter, and a brief description of the analytical method. The EPA lists only one version of a method. The date indicates the specific version approved for use under the CWA. The methods listed below are organized according to the table at 40 CFR part 136 in the order in which they appear.

Methods approved under Standard Methods can be purchased from *standardmethods.org.* The price generally ranges between from \$60 to \$80 per method. Standard Methods also offers memberships or subscriptions that allow unlimited access to their methods. The methods incorporated by reference are reasonably available.

The EPA is adding the following methods to Tables IB, IC, and ID at 40 CFR part 136 for the following parameters:

1. Color. 2120 B–2021, Visual Comparison Method, is a platinumcobalt method of measuring color, the unit of color being that produced by one mg platinum per liter in the form of the chloroplatinate ion. The 1:2 ratio of cobalt to platinum resulting from the preparation of the standard platinumcobalt solution matches the color of natural waters.

2120 F–2021, American Dye Manufacturers Institute (ADMI) Weighted-Ordinate Spectrophotometric Method. This method calculates singlenumber color difference values (*i.e.*, uniform color differences) in accordance with the Adams-Nickerson chromatic value formula. Values are independent of chroma and hue. Transmittance of light is measured

spectrophotometrically at multiple wavelengths and converted to a set of abstract numbers, which then are converted to a single number that indicates color value. This number is expressed on a scale used by the ADMI.

2. Turbidity. 2130 B–2020, Nephelometric Method is based on a comparison of the intensity of light scattered by the sample under defined conditions with the intensity of light scattered by a standard reference suspension under the same conditions. The higher the intensity of scattered light, the higher the turbidity. Formazin polymer is used as the primary standard reference suspension.

3. Acidity. 2310 B-2020, Titration Method measures the hydrogen ions present in a sample as a result of dissociation or hydrolysis of solutes that react with additions of standard alkali. Acidity thus depends on the endpoint pH or indicator used. The construction of a titration curve by recording a sample's pH after successive small, measured additions of titrant permits identification of inflection points and buffering capacity, if any, and allows the acidity to be determined with respect to any pH of interest. Samples of industrial wastes, acid mine drainage, or other solutions that contain appreciable amounts of hydrolyzable metal ions such as iron, aluminum, or manganese are treated with hydrogen peroxide to ensure the oxidation of any reduced forms of polyvalent cations and are boiled to hasten hydrolysis. Acidity results may be highly variable if this procedure is not followed exactly.

4. Alkalinity. 2320 B–2021 Titration Method, measures the hydroxyl ions

present in a sample resulting from dissociation or hydrolysis of solutes that react with additions of standard acid. Alkalinity thus depends on the endpoint pH used. For samples of low alkalinity (less than 20 mg/L CaCO₃) an extrapolation technique based on the near proportionality of concentration of hydrogen ions to excess of titrant beyond the equivalence point is used. The amount of standard acid required to reduce the pH exactly 0.30 pH unit is measured carefully. Because this change in pH corresponds to an exact doubling of the hydrogen ion concentration, a simple extrapolation can be made to the equivalence point.

5. Hardness.

a. In 2340 B–2021, Hardness by Calculation is the preferred method for determining hardness by calculating it from the results of separate determinations of calcium and magnesium by any approved method provided that the sum of the lowest point of quantitation for Ca and Mg is below the NPDES permit requirement for hardness.

b. In 2340 C–2021,

Ethylenediaminetetraacetic acid Titrimetric Method, EDTA forms a chelated soluble complex when added to a solution of certain metal cations. If a small amount of a dye such as eriochrome black T or calmagite is added to an aqueous solution containing calcium and magnesium ions at a pH of 10.0 ± 0.1 , the color of the solution becomes wine red. If EDTA is added as a titrant, the calcium and magnesium will be complexed, and when all of the magnesium and calcium has been complexed, the solution turns from wine red to blue, marking the endpoint of the titration. The volume of titrant used is proportional to hardness in the sample. Magnesium ion must be present to yield a satisfactory endpoint. To ensure this, a small amount of complexometrically neutral magnesium salt of EDTA is added to the buffer; this automatically introduces sufficient magnesium and obviates the need for a blank correction.

6. Specific Conductance. 2510 B–2021 measures conductance (or resistance) in the laboratory using a standard potassium chloride solution and from the corresponding conductivity, a cell constant is calculated. Most conductivity meters do not display the actual solution conductance, or resistance, rather, they generally have a dial that permits the user to adjust the internal cell constant to match the conductivity of a standard. Once the cell constant has been determined, or set, the conductivity of an unknown solution is displayed by the meter.

7. Residue-Total.

a. In 2540 B–2020 an aliquot of a wellmixed sample is evaporated in a preweighed evaporating dish at 103–105 °C to constant weight in a 103 to 105 °C oven. The increase compared to the empty pre-weighed dish weight represents total solids.

b. In 2540 C–2020, Total Dissolved Solids Dried at 180 °C (Residuefilterable in Table IB) a measured volume of a well-mixed sample is filtered through a glass fiber filter with applied vacuum. The entire exposed surface of the filter is washed with at least three successive volumes of reagent-grade water with continued suction until all traces of water are removed. The total filtrate (with washings) is then transferred to a preweighed dish and evaporated to dryness. Successive volumes of sample are added to the same dish after evaporation if necessary to yield between 2.5 and 200 mg of dried residue. The evaporated residue is then dried for one hour or more in an oven at 180 °C, cooled in a desiccator to ambient temperature, and weighed until the weight change is less than 0.5 mg.

c. In 2540 D–2020, Total Suspended Solids Dried from 103 to 105 °C (Residue—non-filterable total suspended solids (TSS) in Table IB) a well-mixed sample is filtered through a pre-weighed standard glass-fiber filter. The filter and the retained residue are then dried to a constant weight in a 103 to 105 °C oven. The increase in filter weight represents TSS.

d. In 2540 E–2020, Fixed and Volatile Solids Ignited at 550 °C (Residuevolatile in Table IB) the residue obtained from the determination of total (Method 2540 B), filterable (Method 2540 C), or non-filterable residue (Method 2540 D) is ignited at 550 ± 50 °C in a muffle furnace, cooled in a desiccator to ambient temperature and weighed. Repeated successive cycles of drying, cooling, desiccating, and weighing are performed until the weight change is less than 0.5 mg. The remaining solids are fixed total, dissolved, or suspended solids, while those lost to ignition are volatile total, dissolved, or suspended solids.

e. In 2540 F–2020, Settleable Solids (aka, Residue—settleable in Table IB), a well-mixed sample is used to fill an Imhoff cone or graduated cylinder to the one liter mark. The sample is allowed to settle for 45 minutes, then gently agitated near the sides of the cone (or graduated cylinder) with a rod or by spinning. The sample is then allowed to settle for another 15 minutes and the volume of settleable solids in the cone (or graduated cylinder) is recorded as mL/L. When applicable, the recorded volume is corrected for interference from pockets of liquid volume.

8. Multiple metals by flame atomic absorption spectrometry.

a. 3111 B–2019, Direct Air-Acetylene Flame Method. The method is approved in Table IB for determination of antimony, cadmium, calcium, chromium, cobalt, copper, gold, iridium, iron, lead, magnesium, manganese, nickel, palladium, platinum, potassium, rhodium, ruthenium, silver, sodium, thallium, tin, and zinc. A sample is aspirated into a flame and the metals are atomized. A light beam is directed through the flame, into a monochromator, and onto a detector that measures the amount of light absorbed by the atomized metal in the flame. Because each metal has its own characteristic absorption wavelength, a source lamp composed of that element is used. The amount of energy at the characteristic wavelength absorbed in the flame is proportional to the concentration of the element in the sample over a limited concentration range.

b. 3111 C–2019, Extraction and Air-Acetylene Flame Method consists of chelation with ammonium pyrrolidine dithiocarbamate (APDC) and extraction into methyl isobutyl ketone (MIBK), followed by aspiration into an airacetylene flame and is suitable for the determination of low concentrations of cadmium, chromium, cobalt, copper, iron, lead, manganese, nickel, silver, and zinc. The method is approved in Table IB for determination of cadmium, chromium, cobalt, copper, iron, lead, nickel, silver, and zinc. The EPA is also approving method 3111 C for manganese. This parameter was inadvertently left off in an earlier rulemaking approving method 3111 C.

c. 3111 D–2019, Direct Nitrous Oxide-Acetylene Flame Method. A sample is aspirated into a flame produced using a mixture of nitrous oxide and acetylene and the metals are atomized. A light beam is directed through the flame, into a monochromator, and onto a detector that measures the amount of light absorbed by the atomized metal in the flame. The method is approved in Table IB for determination of aluminum, barium, beryllium, molybdenum, osmium, titanium, and vanadium. In addition, the EPA is approving method 3111 D for calcium. This parameter was inadvertently left off in an earlier rulemaking approving method 3111 D. d. 3111 E–2019, Extraction and

d. 3111 E–2019, Extraction and Nitrous Oxide-Acetylene Flame Method. The method consists of chelation with 8-hydroxyquinoline, extraction with MIBK, and aspiration into a nitrous oxide-acetylene flame and is suitable for the determination of aluminum at concentrations less than 900 μ g/L and beryllium at concentrations less than 30 μ g/L. The method is approved in Table IB for determination of aluminum, and beryllium.

9. Mercury—Total. 3112 B–2020, Metals by Cold-Vapor Atomic Absorption Spectrometric Method is a flameless AA procedure based on the absorption of radiation at 253.7 nm by mercury vapor. The mercury in a sample is reduced to the elemental state and aerated from solution in a closed system. The mercury vapor passes through a cell positioned in the light path of an atomic absorption spectrophotometer. Absorbance is measured as a function of mercury concentration. The method is approved in Table IB for determination of mercury.

10. Metals by AA Furnace. In 3113 B-2020, Electrothermal Atomic Absorption Spectrometric Method, a discrete sample volume is dispensed into the graphite sample tube (or cup). Typically, determinations are made by heating the sample in three or more stages. First, a low current heats the tube to dry the sample. The second, or charring, stage destroys organic matter and volatilizes other matrix components at an intermediate temperature. Finally, a high current heats the tube to incandescence and, in an inert atmosphere, atomizes the element being determined. Additional stages frequently are added to aid in drying and charring, and to clean and cool the tube between samples. The resultant ground-state atomic vapor absorbs monochromatic radiation from the source. A photoelectric detector measures the intensity of transmitted radiation. The inverse of the transmittance is related logarithmically to the absorbance, which is directly proportional to the number density of vaporized ground-state atoms (the Beer-Lambert law) over a limited concentration range. The method is approved in Table IB for determination of aluminum, antimony, arsenic, barium, beryllium, cadmium, chromium, cobalt, copper, iron, lead, manganese, molybdenum, nickel, selenium, silver, and tin. Although not specifically listed as target analytes in 3113 B, the 2010 version of the method is also approved in Table IB for determination of gold, thallium, and vanadium, as these elements may also be determined using the method.

11. Arsenic and Selenium by AA Gaseous Hydride.

a. 3114 Å–2020, Manual Hydride Generation/Atomic Absorption Spectrometric Method is a manual hydride generation method that is applicable to the determination of arsenic and selenium by conversion to their hydrides by sodium borohydride reagent and transport into an atomic absorption atomizer. The method is approved in Table IB for determination of arsenic and selenium.

b. 3114 C–2020, Continuous Hydride Generation/Atomic Absorption Spectrometric Method is a continuousflow hydride generation method that is applicable to the determination of arsenic and selenium by conversion to their hydrides by sodium borohydride reagent and transport into an atomic absorption atomizer. The continuous hydride generator offers the advantages of simplicity in operation, excellent reproducibility, low detection limits, and high sample volume throughput for selenium analysis following preparations as described in 3500-Se B or 3114 B.4c and d. The method is approved in Table IB for determination of arsenic and selenium.

12. Multiple Metals by ICP/AES (Plasma Emission Spectroscopy). In 3120 B-2020, an Inductively Coupled Plasma source consists of a flowing stream of argon gas ionized by an applied radio frequency field typically oscillating at 27.1 MHz. This field is inductively coupled to the ionized gas by a water-cooled coil surrounding a quartz torch that supports and confines the plasma. A sample aerosol is generated in an appropriate nebulizer and spray chamber and is carried into the plasma through an injector tube located within the torch. The sample aerosol is injected directly into the ICP, subjecting the constituent atoms to temperatures of about 6000 to 8000 °K. Because this results in almost complete dissociation of molecules, significant reduction in chemical interferences is achieved. The high temperature of the plasma excites atomic emission efficiently. Ionization of a high percentage of atoms produces ionic emission spectra. The ICP provides an optically thin source that is not subject to self-absorption except at very high concentrations. Total metals are determined after appropriate digestion. The method is approved in Table IB for determination of aluminum, antimony, arsenic, barium, beryllium, boron, cadmium, calcium, chromium, cobalt, copper, iron, lead, magnesium, manganese, molybdenum, nickel, potassium, selenium, silica, silver, sodium, thallium, vanadium, and zinc. Although not specifically listed as a target analyte in method 3120 B, the 2011 version of the method is also approved in Table IB for determination

of phosphorus because this element may also be determined using the method.

13. Multiple Metals by Inductively Coupled Plasma-Mass Spectrometry. In this method, 3125 B-2020, Inductively Coupled Plasma-Mass Spectrometry-Method, a sample is introduced into an argon-based, high-temperature radiofrequency plasma, usually via pneumatic nebulization. As energy transfers from the plasma to the sample stream, the target element undergoes desolvation, atomization, and ionization. The resulting ions are extracted from the plasma through a differential vacuum interface and separated based on their mass-to-charge (m/z) ratio by a mass spectrometer. Typically, either a quadrupole (with or without collision cell technology or dynamic reaction cell) or magnetic sector (high-resolution) mass spectrometer is used. An electron multiplier detector counts the separated ions, and a computer-based datamanagement system processes the resulting information. The method is approved in Table IB for determination of aluminum, antimony, arsenic, barium, beryllium, cadmium, chromium, cobalt, copper, lead, manganese, molybdenum, nickel, potassium, selenium, silver, thallium, vanadium, and zinc. Although not specifically listed as a target analyte in method 3125 B, the 2011 version of the method is also approved in Table IB for determination of boron, calcium, gold, iridium, iron, magnesium, palladium, platinum, potassium, rhodium, ruthenium, silica, sodium, tin, and titanium as these elements may also be determined using the method.

14. 3500 Colorimetric Series for Multiple Metals.

a. Aluminum. In 3500-Al B–2020, Eriochrome Cyanine R Method with Eriochrome cyanine R dye, dilute aluminum solutions buffered to a pH of 6.0 produce a red to pink complex that exhibits maximum absorption at 535 nm. The intensity of the developed color is influenced by the aluminum concentration, reaction time, temperature, pH, alkalinity, and concentration of other ions in the sample. To compensate for color and turbidity, the aluminum in one portion of a sample is complexed with EDTA to provide a blank. The interference of iron and manganese, two elements commonly found in water when aluminum is present, is eliminated by adding ascorbic acid. The method is approved in Table IB for determination of aluminum.

b. Arsenic. In 3500-As B–2020, Silver Diethyldithiocarbamate Method, arsenite, containing trivalent arsenic, is reduced selectively by aqueous sodium borohydride solution to arsine, AsH₃, in an aqueous medium of pH 6. Arsenate, methylarsonic acid, and dimethylarsinic acid are not reduced under these conditions. The generated arsine is swept by a stream of oxygen-free nitrogen from the reduction vessel through a scrubber containing glass wool or cotton impregnated with lead acetate solution into an absorber tube containing silver diethyldithiocarbamate and morpholine dissolved in chloroform. The intensity of the red color that develops is

measured at 520 nm. The method is approved in Table IB for determination of arsenic. c. Calcium. In 3500-Ca B–2020, EDTA Titrimetric Method, EDTA is added to water containing both calcium and

water containing both calcium and magnesium, where it combines first with the calcium. Calcium can be determined directly, with EDTA, when the pH is made sufficiently high that the magnesium is largely precipitated as the hydroxide and an indicator is used that combines with calcium only. Several indicators give a color change when all the calcium has been complexed by the EDTA at a pH of 12 to 13. The method is approved in Table IB for determination of calcium.

d. Chromium. 3500-Cr B-2020, Colorimetric Method measures total chromium and dissolved hexavalent chromium, (chromium VI). For total chromium, an unfiltered sample must first be digested using an approved digestion procedure (see Table IB, footnote 4). For dissolved hexavalent chromium, a sample is filtered, and the hexavalent chromium is determined colorimetrically by reaction with diphenylcarbazide in acid solution. A red-violet colored complex of unknown composition is produced. The method is approved in Table IB for determination of total chromium after digestion of the sample, and for dissolved hexavalent chromium (chromium VI).

In 3500-Cr C–2020, Ion Chromatographic Method. This method is applicable to determination of dissolved hexavalent chromium in drinking water, groundwater, and industrial wastewater effluents. An aqueous sample is filtered, and its pH adjusted to between 9 and 9.5 with a concentrated buffer. This pH adjustment reduces the solubility of trivalent chromium and preserves the hexavalent chromium oxidation state. The sample is introduced into the instrument's eluent stream of ammonium sulfate and ammonium hydroxide. Trivalent chromium in solution is separated from the hexavalent chromium by the column. After separation, hexavalent

chromium reacts with an azide dye to produce a chromogen that is measured at 530 or 540 nm. Hexavalent chromium is identified based on retention time. The method is approved in Table IB for determination of dissolved hexavalent chromium (chromium VI).

e. Copper Colorimetric. In 3500-Cu B-2020, Neocuproine Method, the sample is treated with hydroxylamine hydrochloride to reduce any cupric ions (Cu2+) to cuprous ions (Cu+). Sodium citrate is used to complex metallic ions that might precipitate when the pH is raised. The pH is adjusted to between 4 and 6 with ammonium hydroxide (NH₄OH), a solution of neocuproine (2,9-dimethyl-1,10-phenanthroline) in methanol is added, and the resultant complex is extracted into chloroform (CHCl₃). After dilution of the CHCl₃ to an exact volume with methanol (CH₃OH), the absorbance of the solution is measured at 457 nm. The method is approved in Table IB for determination of copper.

In 3500-Cu C–2020, Bathocuproine Method, cuprous ion forms a watersoluble orange-colored chelate with disodium bathocuproine disulfonate (sodium 4,4'-(2,9-dimethyl-1,10phenanthroline-4,7-

diyl)dibenzenesulfonate). While the color forms over the pH range 3.5 to 11.0, the recommended pH range is between 4 and 5. The sample is buffered at a pH of about 4.3 and reduced with hydroxylamine hydrochloride. The absorbance is measured at 484 nm. The 2011 editorial revision currently is approved in Table IB for determination of copper.

f. Potassium. In 3500-K B–2020, Flame Photometric Method, trace amounts of potassium can be determined in either a direct-reading or internal-standard type of flame photometer at a wavelength of 766.5 nm. The method is approved in Table IB for determination of potassium.

In 3500-K C–2020, Potassium-Selective Electrode Method, potassium ions are measured potentiometrically by using a potassium ion-selective electrode and a double-junction, sleevetype reference electrode. The analysis is performed with either a pH meter having an expanded millivolt scale capable of being read to the nearest 0.1 mV or a specific-ion meter having a direct concentration scale for potassium. Before measurement, an ionic strength adjustor reagent is added to both standards and samples to maintain a constant ionic strength. The electrode response is measured in standard solutions with potassium concentrations spanning the range of interest using a calibration line derived either by the

instrument meter or manually. The electrode response in sample solutions is measured following the same procedure and potassium concentration determined from the calibration line or instrument direct readout. The 2011 editorial revision currently is approved in Table IB for determination of potassium.

g. Manganese. In 3500-Mn B–2020, Persulfate Method, persulfate oxidation of soluble manganous compounds to form permanganate is carried out in the presence of silver nitrate. The resulting color is stable for at least 24 hours if excess persulfate is present and organic matter is absent. The method is approved in Table IB for determination of manganese.

h. Sodium. In 3500-Na B-2020, Flame Emission Photometric Method, a sample is nebulized into a gas flame under carefully controlled, reproducible excitation conditions. The sodium resonant spectral line at 589 nm is isolated by interference filters or by light-dispersing devices such as prisms or gratings. Emission light intensity is measured by a phototube, photomultiplier, or photodiode. The light intensity at 589 nm is approximately proportional to the sodium concentration. The method is approved in Table IB for determination of sodium.

i. Lead. In 3500-Pb B–2020, Dithizone Method, an acidified sample containing microgram quantities of lead is mixed with ammoniacal citrate-cyanide reducing solution and extracted with dithizone in chloroform (CHCl₃) to form a cherry-red lead dithizonate. The color of the mixed color solution is measured photometrically. The method is approved in Table IB for determination of lead.

j. Zinc. 3500-Zn B-2020, Zincon Method. Zinc forms a blue complex with zincon (2-carboxy-2'-hydroxy-5'sulfoformazyl benzene) in a solution buffered to pH 9.0. Other heavy metals likewise form colored complexes with zincon. Cyanide is added to complex zinc and heavy metals. Cyclohexanone is added to selectively free zinc from its cyanide complex so that it can be complexed with zincon to form a blue color which is measured spectrophotometrically at 620 nm. Sodium ascorbate reduces manganese interference. The developed color is stable except in the presence of copper. The method is approved in Table IB for determination of zinc.

15. 4110 Series, Ion Chromatography. a. In 4110 B–2020, Ion Chromatography with Chemical Suppression of Eluent Conductivity, a water sample is injected into a stream of

eluent and passed through a series of ion exchangers. The anions of interest are separated based on their relative affinities for a low-capacity, strongly basic anion exchanger (guard and analytical columns). The separated anions are directed through a suppressor device that provides continuous suppression of eluent conductivity and enhances analyte response. In the suppressor, the separated anions are converted to their highly conductive acid forms while the conductivity of the eluent is greatly decreased. The separated anions in their acid forms are measured by conductivity. They are identified based on retention time as compared to standards. Quantitation is by measurement of peak area or peak height. The method is approved in Table IB for determination of bromide, chloride, fluoride, nitrate, combined nitrate-nitrite, nitrite, orthophosphate, and sulfate.

b. 4110 C-2020, Single-Column Ion Chromatography with Direct Conductivity Detection. An aqueous sample is injected into an ion chromatograph consisting of an injector port, analytical column, and conductivity detector. The sample merges with the eluent stream and is pumped through the analytical column where the anions are separated based on their affinity for the active sites of the column packing material. Concentrations are determined by direct conductivity detection without chemical suppression. The method is approved in Table IB for determination of bromide, chloride, fluoride, nitrate, combined nitrate-nitrite, nitrite, orthophosphate, and sulfate.

c. 4110 D–2020, Ion Chromatographic Determination of Oxyhalides and Bromide. The sample is analyzed in a manner similar to that in 4110 B-2020. However, bromate has been shown to be subject to positive interferences in some matrices. The interference is noticeable usually as a flattened peak. It often can be eliminated by passing the sample through an H+ off-line solid-phase extraction (SPE) cartridge, by selection of a different column-eluent combination, or by diluting the eluent, which will increase retention times and spread the chromatogram. Additionally, chloride or a nontarget analyte present in unusually high concentration may overlap with a target analyte sufficiently to cause problems in quantitation or may cause retention-time shifts. Dilution of the sample may resolve this problem. The method is approved in Table IB for determination of bromide.

16. Inorganic Anions by CIE/UV (Capillary Ion Electrophoresis). In 4140

B-2020, Capillary Ion Electrophoresis with Indirect UV Detection, the sample is introduced at the cathodic end of the capillary and anions are separated based on their differences in mobility in the electric field as they migrate through the capillary. Cations migrate in the opposite direction and are not detected. Water and neutral organics are not attracted toward the anode. They migrate after the anions and thus do not interfere with anion analysis. Anions are detected as they displace charge-forcharge the UV-absorbing electrolyte anion (chromate), causing a net decrease in UV absorbance in the analyte anion zone compared to the background electrolyte. Detector polarity is reversed to provide positive millivolt response to the data system. As in chromatography, the analytes are identified by their migration time and quantitated by using time-corrected peak area relative to standards. The method is approved in Table IB for determination of bromide, chloride, fluoride, nitrate, combined nitrate-nitrite, nitrite, orthophosphate, and sulfate.

17. 4500 Series, Chloride. a. 4500-Cl⁻ B–2021, Titrimetric Method. In a neutral or slightly alkaline solution, potassium chromate can indicate the endpoint of the silver nitrate titration of chloride. Silver chloride is precipitated quantitatively before red silver chromate is formed. In this version of the method approved by the Standard Methods Committee in 2021, additional information regarding removal of interferences caused by sulfide, thiosulfate, and sulfite ions by digestion of the sample with hydrogen peroxide prior to titration has been added to the sample preparation procedures. A tighter pH range of 8 to 10, as opposed to 7 to 10, is specified for adjustment of the pH of the sample prior to titration. A reference has been added for the 2021 Standard Methods Joint Task Group validation report titled: "Interlaboratory validation study for the use of H_2O_2 with boiling for determining Cl⁻." The method is approved in Table IB for determination of chloride.

b. 4500-Cl⁻ C-2021, Mercuric Nitrate Method. Chloride can be titrated with mercuric nitrate, Hg(NO₃)₂, because of the formation of soluble, slightly dissociated mercuric chloride. In the pH range 2.3 to 2.8, diphenylcarbazone indicates the titration endpoint by formation of a purple complex with the excess mercuric ions. Xylene cyanol FF serves as a pH indicator and endpoint enhancer. Increasing the strength of the titrant and modifying the indicator mixtures extends the range of measurable chloride concentrations. The method is approved in Table IB for determination of chloride.

c. 4500-Cl⁻ D-2021, Potentiometric Method. Chloride is determined by potentiometric titration with silver nitrate solution with a glass and silversilver chloride electrode system. During titration, an electronic voltmeter is used to detect the change in potential between the two electrodes. The endpoint of the titration is that instrument reading at which the greatest change in voltage has occurred for a small and constant increment of silver nitrate added. The method is approved in Table IB for determination of chloride.

d. 4500-Cl⁻ E–2021, Automated Ferricyanide Method. Thiocyanate ion is liberated from mercuric thiocyanate by the formation of soluble mercuric chloride. In the presence of ferric ion, free thiocyanate ion forms a highly colored ferric thiocyanate, of which the intensity is proportional to the chloride concentration. The is approved in Table IB for determination of chloride.

18. 4500 Series Cyanide Total or Available.

a. 4500-CN⁻ B-2021, Manual Distillation (as Preliminary Treatment of Samples). Total cyanides are measured after preliminary treatment of samples for preservation and to remove interferences. The preliminary treatment required depends on which interfering substances the samples contain. Distillation removes many interfering substances, but other pretreatment procedures will be needed for samples containing sulfides, fatty acids, oxidizing agents, nitrites, and nitrates. The method is approved in Table IB for preliminary treatment of samples to be used for determination of cyanide.

b. 4500–CN⁻ C–2021, Total Cyanide after Distillation. Hydrogen cyanide (HCN) is liberated from an acidified sample by distillation and purging with air, with the HCN gas collected in a NaOH scrubbing solution. The cyanide concentration in the scrubbing solution is determined via titrimetric, colorimetric, or potentiometric procedures. The method is approved in Table IB for preliminary treatment of samples to be used for determination of cyanide.

c. $4500-CN^-$ D-2021, Titrimetric Method. CN⁻ in the alkaline distillate from the preliminary treatment procedures ($4500-CN^-$ B and C) is titrated with standard silver nitrate (AgNO₃) to form the soluble cyanide complex Ag(CN)²⁻. As soon as all CN⁻ has been complexed and a small excess of Ag⁺ has been added, the silversensitive indicator, p-

dimethylaminobenzalrhodanine, detects

the excess Ag⁺ and immediately changes color from yellow to salmon. The method is approved in Table IB for determination of cyanide.

d. 4500–CN⁻ E–2021, Spectrophotometric Method. Total CN⁻ in the alkaline distillate from the preliminary treatment procedures (4500–CN⁻ B and C) is converted to cyanogen chloride (CNCl) by reaction with chloramine-T at pH less than 8 without hydrolyzing to cyanate (CNO⁻). After the reaction is complete, adding a pyridine-barbituric acid reagent turns CNCl a red-blue color. Maximum color absorbance in aqueous solution is between 575 and 582 nm. The method is approved in Table IB for determination of cyanide.

e. 4500– CN^- F–2021, Ion Selective Electrode Method. Total CN^- in the alkaline distillate from the preliminary treatment procedures (4500- CN^- B and C) is determined potentiometrically by using a CN^- -ion selective electrode. The 2016 version of the method currently is approved in Table IB for determination of cyanide.

f. 4500–CN[–] G–2021, Cyanides Amenable to Chlorination after Distillation. Available cvanide, or cyanide amenable to chlorination (CATC), can be determined when a portion of the sample is chlorinated at high pH and cvanide levels in the chlorinated sample are determined after manual distillation followed by titrimetric or spectrophotometric measurement. CATC is calculated by the difference between the results for cyanide in the unchlorinated sample and the results for the chlorinated sample. The method is approved in Table IB for preliminary treatment of samples to be used for determination of available cvanide.

g. 4500– $\check{C}N^-$ N–2021, Total Cyanide after Distillation by Flow Injection Analysis. Total cyanides are digested and steam-distilled from the sample (4500– CN^- C). The cyanide in this distillate is converted to CNCl by reaction with chloramine-T at pH less than 8. The CNCl then forms a red-blue dye by reacting with pyridine-barbituric acid reagent. The absorbance of this red dye is measured at 570 nm and is proportional to the total or weak acid dissociable cyanide in the sample. The method is approved in Table IB for determination of cyanide.

19. 4500 Total Fluoride Series. a. 4500– F^- B–2021, Preliminary Distillation Step. Fluoride is separated from other nonvolatile constituents in water by conversion to hydrofluoric or fluosilicic acid and subsequent distillation. The conversion is accomplished by using a strong, highboiling acid. To protect against glassware etching, hydrofluoric acid is converted to fluosilicic acid by using soft glass beads. Quantitative fluoride recovery is accomplished by using a relatively large sample. Acid and sulfate carryover are minimized by distilling over a controlled temperature range. The method is approved in Table IB for preliminary treatment of samples to be used for determination of fluoride.

b. 4500-F⁻ C-2021, Ion-Selective Electrode Method. The fluoride electrode is an ion-selective sensor that measures the ion activity of fluoride in solution rather than concentration. The key element in the fluoride electrode is the laser-type doped lanthanum fluoride crystal across which a potential is established by fluoride solutions of different concentrations. The crystal contacts the sample solution at one face and an internal reference solution at the other. Fluoride ion activity depends on the solution total ionic strength and pH, and on fluoride complexing species. Adding an appropriate buffer provides a nearly uniform ionic strength background, adjusts pH, and breaks up complexes. In effect, the electrode measures concentration. The method is approved in Table IB for determination of fluoride.

c. 4500–F⁻ D–2021, SPADNS Method. The SPADNS colorimetric method is based on the reaction between fluoride and a "lake" of zirconium-dye. Fluoride reacts with the dye lake, dissociating a portion of it into a colorless complex anion (ZrF₆^{2–}) and the dye. As the amount of fluoride increases, the color produced becomes progressively lighter and absorbance is measured colorimetrically at 570 nm. The method is approved in Table IB for determination of fluoride.

d. 4500–F⁻ E–2021, Complexone Method. The sample is distilled in the automated system, and the distillate is reacted with alizarin fluorine bluelanthanum reagent to form a blue complex that is measured colorimetrically at 620 nm. method is approved in Table IB for determination of fluoride.

20. 4500 Hydrogen ion (pH). In 4500– H⁺ B–2021, Electrometric Method, the basic principle of electrometric pH measurement is determination of the activity of the hydrogen ions by potentiometric measurement using a standard hydrogen electrode and a reference electrode. The hydrogen electrode consists of a platinum electrode across which hydrogen gas is bubbled at a pressure of 101 kilopascal. Because of difficulty in its use and the potential for poisoning the hydrogen electrode, the glass electrode commonly is used. The electromotive force produced in the glass electrode system varies linearly with pH. This linear relationship is described by plotting the measured emf against the pH of different buffers. A sample's pH is determined by extrapolation. This version of the method adds information to Section 2—Apparatus, regarding equipment that may be used for manual or automatic temperature compensation. The 2011 editorial revision currently is approved in Table IB for determination of pH.

21. 4500 Kjeldahl Nitrogen Total (TKN) Series.

a. 4500–Norg B–2021, Macro-Kjeldahl Method. In the presence of sulfuric acid (H_2SO_4) , potassium sulfate (K_2SO_4) , and a cupric sulfate (CuSO₄) catalyst, amino nitrogen of many organic materials is converted to ammonium. Free ammonia also is converted to ammonium. After the addition of base, the ammonia is distilled from an alkaline medium and absorbed in boric or sulfuric acid. The ammonia may be determined colorimetrically, by ammonia-selective electrode, or by titration with a standard mineral acid. The method is approved in Table IB for preliminary treatment of samples to be used for determination of total Kjeldahl nitrogen (TKN)

b. 4500– N_{org} C–2021, Semí-Micro-Kjeldahl Method. This is a reducedvolume version of 4500 N_{org} B that specifies use of Kjeldahl flasks with a capacity of 100 mL in a semi-micro-Kjeldahl digestion apparatus equipped with heating elements to accommodate Kjeldahl flasks and a suction outlet to vent fumes. The method is approved in Table IB for preliminary treatment of samples to be used for determination of total Kjeldahl nitrogen (TKN).

c. 4500–N_{org} D–2021, Block Digestion and Flow Injection Analysis. Samples are digested in a block digestor with sulfuric acid and copper sulfate as a catalyst. The digested sample is injected onto the FIA manifold, where its pH is controlled by raising it to a known, basic pH by neutralization with a concentrated buffer. This in-line neutralization converts the ammonium cation to ammonia, and also prevents undue influence of the sulfuric acid matrix on the pH-sensitive color reaction that follows. The ammonia thus produced is heated with salicylate and hypochlorite to produce a blue color that is proportional to the ammonia concentration. The color is intensified by adding sodium nitroprusside. The presence of EDTA in the buffer prevents the precipitation of calcium and magnesium. The resulting peak's absorbance is measured at 660 nm. The peak area is proportional to the

concentration of TKN in the original sample. The method is approved in Table IB for determination of TKN.

22. 4500–NH $_3$ Nitrogen (Ammonia as nitrogen) Series.

a. 4500–NH₃ B–2021, Preliminary Manual Distillation Step. The sample is buffered at pH 9.5 with a borate buffer to decrease hydrolysis of cyanates and organic nitrogen compounds. It is distilled into a solution of boric acid when titration is to be used, or into H_2SO_4 , when the phenate method is used as the determinative step. The ammonia in the distillate can be determined either colorimetrically by the phenate method or titrimetrically with standard H₂SO₄ and a mixed indicator or a pH meter. Ammonia in the distillate also can be determined by the ammonia-selective electrode method, using $0.04 \text{ N} \text{H}_2\text{SO}_4$ to trap the ammonia. This revision replaces instructions for storage of ammonia-free water with instructions for preparation of ammonia-free water using an ion exchange resin and simply says that if high blank values are produced, the analyst should prepare fresh ammoniafree water. The method is approved in Table IB for preliminary treatment of samples to be used for determination of ammonia.

b. 4500–NH₃ C–2021, Titration Method. The titrimetric method is used only on samples that have been carried through preliminary distillation. Ammonia is titrated with a standardized sulfuric acid titrant using a mixed indicator of methyl red and methylene blue. The method is approved in Table IB for determination of ammonia as well as for determination of TKN after appropriate digestion/distillation of the sample.

c. 4500-NH₃ D-2021, Electrode Method. The ammonia-selective electrode uses a hydrophobic gaspermeable membrane to separate the sample solution from an electrode internal solution of ammonium chloride. Dissolved ammonia (NH_{3(aq)} and NH₄⁺) is converted to NH_{3(aq)} by raising the pH to above 11 with a strong base. $NH_{3(aq)}$ diffuses through the membrane and changes the internal solution pH that is sensed by a pH electrode. The fixed level of chloride in the internal solution is sensed by a chloride ion-selective electrode that serves as the reference electrode of the sample. Potentiometric measurements are made with a pH meter having an expanded millivolt scale or with a specific ion meter. The method is approved in Table IB for determination of ammonia, as well as for determination of TKN after appropriate digestion/distillation of the sample.

d. 4500-NH3 E-2021, Electrode Method. Ammonia is determined using an ammonia-selective electrode. When a linear relationship exists between concentration and response, known addition is convenient for measuring occasional samples because no calibration is needed. Because an accurate measurement requires that the concentration at least double as a result of the addition, sample concentration must be known within a factor of three. The total concentration of ammonia can be measured in the absence of complexing agents down to 0.8 mg/L NH₃-N or in the presence of a large excess (50 to 100 times) of complexing agent. The method is approved in Table IB for determination of ammonia, as well as for determination of TKN after appropriate digestion/distillation of the sample.

e. 4500–NH₃ F–2021, Phenate Method. An intensely blue compound, indophenol, is formed by the reaction of ammonia, hypochlorite, and phenol catalyzed by sodium nitroprusside. The color is measured spectrophotometrically at 640 nm. The method is approved in Table IB for determination of ammonia, as well as for determination of TKN after appropriate digestion/distillation of the sample.

f. 4500–NH₃ G–2021, Semi-Automated Phenate Method. Alkaline phenol and hypochlorite react with ammonia to form indophenol blue that is proportional to the ammonia concentration. The blue color formed is intensified with sodium nitroprusside. The color is measured spectrophotometrically at 630 to 660 nm. The method is approved in Table IB for determination of ammonia, as well as for determination of TKN after appropriate digestion/distillation of the sample.

g. 4500-NH₃ H-2021, Semi-Automated Phenate Method. A water sample containing ammonia or ammonium cation is injected into an FIA carrier stream to which a complexing buffer (alkaline phenol) and hypochlorite are added. This reaction, the Berthelot reaction, produces the blue indophenol dye. The blue color is intensified by the addition of nitroferricyanide. The resulting peak's absorbance is measured at 630 nm. The peak area is proportional to the concentration of ammonia in the original sample. The method is approved in Table IB for determination of ammonia, as well as for determination of TKN after appropriate digestion/distillation of the sample.

23. $4500-NO_2^-$ Nitrite as Nitrogen. $4500-NO_2^-$ B-2021, Spectrophotometric Method. Nitrite (NO2⁻) in a sample is determined through formation of a reddish-purple azo dye produced at pH 2.0 to 2.5 by coupling diazotized sulfanilamide with N-(1-naphthyl)-ethylenediamine dihydrochloride (NED) and absorbance is measured spectrophotometrically at 543 nm. The method is approved in Table IB for determination of nitrite.

24. 4500–NO3 [–] Nitrogen (Nitrite/ Nitrate as Nitrogen Series).

a. 4500–NO3⁻ D–2019, Nitrate Electrode Method. Nitrate is measured using an ion-selective electrode that develops a potential across a thin, inert membrane holding in place a waterimmiscible liquid ion exchanger. The method is approved in Table IB for determination of nitrate.

b. 4500–NO₃⁻ E–2019, Cadmium Reduction Method. Nitrate (NO₃⁻) is reduced almost quantitatively to nitrite (NO_2^{-}) in the presence of cadmium (Cd). This method uses commercially available Cd granules treated with copper sulfate (CuSO₄) and packed in a glass column. The NO_2^- is then diazotized with sulfanilamide and coupled with NED to form a highly colored azo dye that is measured spectrophotometrically. To correct for any NO_2^- present in the sample before NO₃⁻ reduction, samples also must be analyzed without the reduction step. The method is approved in Table IB for determination of nitrate (by subtraction), as well as for determination of combined nitrate + nitrite, and for determination of nitrite singly when bypassing the reduction step.

c. $4500-NO_3^{-}$ F–2019, Automated Cadmium Reduction Method. This is an automated version of the cadmium reduction method $4500 NO3^{-}$ E. Nitrate in a sample is reduced to nitrite using cadmium reduction and then diazotized with sulfanilamide and coupled with NED to form a highly colored azo dye that is measured

spectrophotometrically. To correct for any NO_2^- present in the sample before NO_3^- reduction, samples also must be analyzed without the reduction step. The method is approved in Table IB for determination of nitrate (by subtraction), as well as for determination of combined nitrate + nitrite, and for determination of nitrite singly when bypassing the reduction step.

d. $4500-NO_3^{-}$ H–2019, Automated Hydrazine Reduction Method. Nitrate in a sample is reduced to nitrite using hydrazine sulfate then diazotized with sulfanilamide and coupled with NED to form a highly colored azo dye that is measured spectrophotometrically. The method is approved in Table IB for determination of combined nitrate and nitrite.

e. 4500-NO₃- I-2019, Cadmium Reduction Flow Injection Method. A sample is passed through a copperized cadmium column to quantitatively reduce its nitrate content to nitrite. The nitrite is diazotized with sulfanilamide and coupled with NED to yield a watersoluble dye with a magenta color whose absorbance at 540 nm is proportional to the nitrate + nitrite in the sample. Nitrite concentrations may be determined by bypassing the cadmium column and nitrate concentration may be calculated by subtraction of the result for the nitrite concentration from the result for the combined nitrate + nitrite concentration. The method is approved in Table IB for determination of nitrate, as well as for determination of combined nitrate + nitrite, and for determination of nitrite singly by bypassing the reduction step.

25. 4500–O Oxygen (Dissolved) Series.

a. 4500-O B-2021, Iodometric Methods. A divalent manganese solution is added and then a strong alkali is added to a sample in a glassstoppered bottle and dissolved oxygen (DO) rapidly oxidizes an equivalent amount of the dispersed divalent manganous hydroxide precipitate into higher-valency hydroxides. Oxidized manganese reverts to the divalent state in the presence of iodide ions in an acidic solution, liberating an amount of iodine equivalent to the original DO content. The iodine is then titrated with a standard thiosulfate solution. The method is approved in Table IB for determination of DO.

b. 4500-O C-2021, Azide Modification. The sample is treated with manganous sulfate, potassium hydroxide, and potassium iodide (the latter two reagents combined in one solution) and finally sulfuric acid. The initial precipitate of manganous hydroxide, Mn(OH)₂, combines with the DO in the sample to form a brown precipitate, manganic hydroxide, $MnO(OH)_2$. Upon acidification, the manganic hydroxide forms manganic sulfate, which acts as an oxidizing agent to release free iodine from the potassium iodide. The iodine, which is stoichiometrically equivalent to the DO in the sample, is then titrated with sodium thiosulfate or phenylarsine oxide (PAO). The azide modification effectively removes nitrite interference, which is the most common interference in biologically treated effluents and incubated biochemical oxygen demand (BOD) samples. The method is approved in Table IB for determination of DO.

c. 4500-O D-2021, Permanganate Modification. The permanganate modification is used only on samples containing Fe(II) (e.g., acid mine water). Concentrated sulfuric acid, potassium permanganate in solution and potassium fluoride in solution are added to the sample. Enough KMnO₄ solution is added to obtain a violet tinge that persists for 5 minutes. 0.5 to 1.0 mL potassium oxalate solution is then added only until permanganate color is removed completely. From this point, the procedure closely parallels that in 4500–O C. The method is approved in Table IB for determination of DO.

d. 4500-O E-2021, Alum Flocculation Modification. Samples high in suspended solids may consume appreciable quantities of iodine in acid solution. The interference due to solids may be removed by alum flocculation. Concentrated ammonium hydroxide and aluminum potassium sulfate solution are added to a sample. The sample is allowed to settle for about 10 min and the clear supernatant is siphoned into a 250- to 300-mL DO bottle until it overflows. From this point, the procedure closely parallels that in 4500–O C. The method is approved in Table IB for determination of DO.

e. 4500–O F–2021, Copper Sulfate-Sulfamic Acid Flocculation Modification. This modification is used for biological flocs (*e.g.*, activated sludge mixtures), which have high oxygen utilization rates. A copper sulfatesulfamic acid inhibitor solution is added to the sample. The suspended solids are allowed to settle, and the relatively clear supernatant liquor is siphoned into a 250- to 300-mL DO bottle. From this point, the procedure closely parallels that in 4500–O C. The method is approved in Table IB for determination of DO.

f. 4500–O G–2021, Electrode Method. Oxygen-sensitive polarographic or galvanic membrane electrodes are composed of two solid metal electrodes in contact with supporting electrolyte separated from the test solution by a selective membrane. Polyethylene and fluorocarbon membranes are commonly used because they are permeable to molecular oxygen and are relatively rugged. The diffusion current is linearly proportional to the molecular-oxygen concentration. The measured current can be converted easily to concentration units (e.g., mg/L) by a number of calibration procedures. The method is approved in Table IB for determination of DO.

g. 4500–O H–2021, Luminescencebased Method. The optical probe uses luminescence-based oxygen sensors to measure the light-emission characteristics of a luminescent reaction; oxygen quantitatively quenches the luminescence. The change in the luminescence signal's lifetime correlates to the DO concentration. The method is approved in Table IB for determination of DO.

26. 4500–P Phosphorus Total and Ortho Phosphorus Series.

a. 4500–P B–2021, Digestion Sample Preparation. Because phosphorus may occur in combination with organic matter, a digestion method to determine total phosphorus must be able to oxidize organic matter effectively to release phosphorus as orthophosphate. Three digestion methods are given in 4500–P B.3, 4, and 5. The perchloric acid method in B.5 is the most vigorous and time-consuming method, and is recommended for particularly difficult samples, such as sediments. The nitric acid-sulfuric acid method is recommended for most samples. The simplest digestion method that may be used for determination of total phosphorus is the persulfate oxidation technique in which 50 mL of an unfiltered sample is boiled with sulfuric acid and either ammonium persulfate or potassium persulfate for approximately 30–40 minutes or until a final volume of about 10 mL is reached. The method is approved in Table IB for preliminary treatment of samples to be used for determination of total phosphorus as orthophosphorus using manual or automated versions of the ascorbic acid reduction, colorimetric methods.

b. 4500–P E–2021, Manual Method. Ammonium molybdate and antimony potassium tartrate react in an acid medium with orthophosphate to form phosphomolybdic acid, a heteropoly acid that is reduced to intensely colored molybdenum blue by ascorbic acid and is measured spectrophotometrically. This revision adds that possible interference from silicate should be evaluated when reporting concentrations less than 10 μ g/L. The method is approved in Table IB for determination of total phosphorus after digestion of the sample, as well as for determination of orthophosphorus in a filtered, undigested sample.

c. 4500–P F–2021, Automated Ascorbic Acid Reduction Method. Ammonium molybdate and antimony potassium tartrate react with orthophosphate in an acid medium to form an antimony-phosphomolybdate complex, which on reduction with ascorbic acid yields an intense blue color suitable for photometric measurement using continuous flow analytical equipment. The method is approved in Table IB for determination of total phosphorus after digestion of the sample, as well as for determination of orthophosphorus in a filtered, undigested sample.

d. 4500–P G–2021, Automated. Ammonium molybdate and antimony potassium tartrate react with orthophosphate in an acid medium to form an antimony-phosphomolybdate complex, which on reduction with ascorbic acid yields an intense blue color suitable for photometric measurement using flow injection analysis. The method is approved in Table IB for determination of total phosphorus after digestion of the sample, as well as for determination of orthophosphorus in a filtered, undigested sample.

e. 4500-P H-2021, Automated Total Phosphorus. Samples are manually digested using the approved procedure for preliminary treatment of samples to be used for determination of total phosphorus. When the resulting solution is injected onto the manifold, the orthophosphate ion reacts with ammonium molybdate and antimony potassium tartrate under acidic conditions to form a complex. This complex is reduced with ascorbic acid to form a blue complex suitable for photometric measurement using flow injection analysis. The method is approved in Table IB for determination of total phosphorus.

27. $4500-\dot{S}_2$ Sulfide Series. a. $4500-\dot{S}_2$ B-2021, Sample Pretreatment. Dissolved sulfide is measured by first removing insoluble matter. This is done by adding sodium hydroxide and aluminum chloride solutions producing an aluminum hydroxide floc that is settled, leaving a clear supernatant for analysis. The method is approved in Table IB for preliminary treatment of samples to be used for determination of sulfide.

b. 4500–S₂⁻ C–2021, Sample Pretreatment. Interferences due to sulfite, thiosulfate, iodide, and many other soluble substances, but not ferrocyanide, are eliminated by first precipitating zinc sulfide (ZnS) by addition of sodium hydroxide and zinc acetate solutions, removing the supernatant, and replacing it with reagent water. The same procedure is used even when not needed for removal of interferences, to concentrate sulfide prior to analysis. The method is approved in Table IB for preliminary treatment of samples to be used for determination of sulfide.

c. $4500-S_2^-$ D-2021, Colorimetric Method. The methylene blue method is based on the reaction of sulfide, ferric chloride, and dimethyl-pphenylenediamine to produce methylene blue. Ammonium phosphate is added after color development to remove ferric chloride color, which is measured photometrically. The procedure is applicable at sulfide concentrations between 0.1 and 20.0 mg/L. The method is approved in Table IB for determination of sulfide.

d. $4500-S_2^-$ F-2021, Titrimetric. Iodine oxidizes sulfide in acid solution. A titration based on this reaction is an accurate method for determining sulfide at concentrations above one mg/L if interferences are absent and if loss of H₂S is avoided. The method is approved in Table IB for determination of sulfide.

e. 4500– S_2^- G–2021, Ion-Selective Electrode Method. The potential of a sulfide ion-selective electrode (ISE) is related to the sulfide ion activity. An alkaline antioxidant reagent (AAR) is added to samples and standards to inhibit oxidation of sulfide by oxygen and to provide a constant ionic strength and pH. Use of the AAR allows calibration in terms of total dissolved sulfide concentration. All samples and standards must be at the same temperature. Sulfide concentrations between 0.032 mg/L and 100 mg/L can be measured without preconcentration. For lower concentrations, preconcentration is necessary. The method is approved in Table IB for determination of sulfide.

28. 4500-SiO₂ Silica Series. a. 4500–SiO₂ C–2021, Colorimetric Method. Ammonium molybdate at pH approximately 1.2 reacts with silica and any phosphate present to produce heteropoly acids. Oxalic acid is added to destroy the molybdophosphoric acid, but not the molybdosilicic acid. Even if phosphate is known to be absent, the addition of oxalic acid is highly desirable and is a mandatory step. The intensity of the yellow color produced is proportional to the concentration of molybdate-reactive silica and is measured photometrically. The method is approved in Table IB for determination of silica.

b. 4500–SiO₂ E–2021, Automated Method for Molybdate-Reactive Silica. Ammonium molybdate at pH approximately 1.2 reacts with silica and any phosphate present to produce heteropoly acids. Oxalic acid is added to destroy the molybdophosphoric acid, but not the molybdosilicic acid. The yellow molybdosilicic acid is reduced by means of amino naphthol sulfonic acid to heteropoly blue. The blue color is more intense than the yellow color of $4500-SiO_2$ C and provides increased sensitivity. The method is approved in Table IB for determination of silica.

c. $4500-SiO_2$ F–2021, Automated Method for Molybdate-Reactive Silicate. Silicate reacts with molybdate under acidic conditions to form yellow betamolybdosilicic acid. This acid is subsequently reduced with stannous chloride to form a heteropoly blue complex that is measured photometrically. Oxalic acid is added to reduce the interference from phosphate. The method is approved in Table IB for determination of silica.

29. $4500-SO_{42}^{-}$ Sulfate Series. a. $4500-SO_{42}^{-}$ C-2021, Gravimetric Method with Ignition of Residue. Sulfate is precipitated in a hydrochloric acid (HCl) solution as barium sulfate (BaSO₄) by the addition of barium chloride (BaCl₂). The precipitation is carried out near the boiling temperature, and after a period of digestion, the precipitate is filtered, washed with water until free of Cl⁻, ignited at 800 °C for an hour and weighed as BaSO₄. The method is approved in Table IB for determination of sulfate.

b. $4500-SO_{42}$ ⁻ D-2021, Gravimetric Method with Drying of Residue. Sulfate is precipitated in a hydrochloric acid (HCl) solution as barium sulfate (BaSO₄) by the addition of barium chloride (BaCl₂). The precipitation is carried out near the boiling temperature, and after a period of digestion the precipitate is filtered, washed with water until free of Cl⁻, dried to a constant weight in an oven at 105 °C or higher, and weighed as BaSO₄. The method is approved in Table IB for determination of sulfate.

c. $4500-SO_{42}^{-}$ E-2021, Turbidimetric Method. Sulfate ion (SO₄₂⁻) is precipitated in an acetic acid medium with barium chloride (BaCl₂) to form barium sulfate (BaSO₄) crystals of uniform size. Light absorbance of the BaSO₄ suspension is measured by a photometer and the SO₄₂⁻ concentration is determined by comparison of the reading with a standard curve. The method is approved in Table IB for determination of sulfate.

d. 4500–SO₄₂⁻ F–2021, Automated Colorimetric Method. Barium sulfate is formed by the reaction of the SO₄₂⁻ with barium chloride (BaCl₂) at a low pH. At high pH, excess barium reacts with methylthymol blue (MTB) to produce a blue chelate. The uncomplexed methylthymol blue is gray. The intensity of gray (uncomplexed methylthymol blue) is measured photometrically and is proportional to concentration of sulfate. The method is approved in Table IB for determination of sulfate.

e. $4500-SO_{42}$ ⁻ G-2021, Automated Colorimetric Method. At pH 13.0, barium forms a blue complex with methylthymol blue (MTB). The sample is injected into a low, but known, concentration of sulfate. The sulfate from the sample then reacts with the ethanolic barium-MTB solution and displaces the MTB from the barium to give barium sulfate and uncomplexed MTB. Uncomplexed MTB has a grayish color. The pH is raised with NaOH and the gray color of the uncomplexed MTB is measured photometrically. The intensity of the gray color is proportional to the sulfate concentration. The method is approved in Table IB for determination of sulfate.

30. Sulfite $4500-SO_{32}$ ⁻ B-2021, Titrimetric Iodometric Method. An acidified sample containing sulfite (SO₃₂⁻) is titrated with a standardized potassium iodide-iodate titrant. Free iodine, liberated by the iodide-iodate reagent, reacts with SO₃₂⁻. The titration endpoint is signaled by the blue color resulting from the first excess of iodine reacting with a starch indicator. The method is approved in Table IB for determination of sulfite.

31. 5520 Oil and Grease Series. a. 5520 B-2021, Liquid-Liquid, Partition-Gravimetric Method. Dissolved or emulsified oil and grease is extracted from water by intimate contact with an extracting solvent (n-hexane). The extract is dried over sodium sulfate. The solvent is then distilled from the extract and the hexane extractable material is desiccated and weighed. Some extractables, especially unsaturated fats and fatty acids, oxidize readily; hence, special precautions regarding temperature and solvent vapor displacement are included to minimize this effect. Organic solvents shaken with some samples may form an emulsion that is very difficult to break. This method includes a means for handling such emulsions. Recovery of solvents is discussed. Solvent recovery can reduce both vapor emissions to the atmosphere and costs. The method is approved in Table IB for determination of oil and grease (hexane extractable material or HEM).

b. 5520 F-2021, Hydrocarbons. The oil and grease extracted by 5520 B is used for this test. When only hydrocarbons are of interest, this procedure is introduced before final measurement. When hydrocarbons are to be determined after total oil and grease has been measured, redissolve the extracted oil and grease in n-hexane. Silica gel has the ability to adsorb polar materials. The solution of extracted hydrocarbons and fatty materials in nhexane is mixed with silica gel, and the fatty acids are removed selectively from solution. The solution is filtered to remove the silica gel, the solvent is distilled, and the silica gel treated hexane extractable material (SGT-HEM) is weighed. The materials not eliminated by silica gel adsorption are

designated hydrocarbons by this test. The method is approved in Table IB for determination of oil and grease (hexane extractable material or HEM).

32. 5530 Phenols Series.

a. 5530 B–2021, Manual Distillation. Phenols, defined as hydroxy derivatives of benzene and its condensed nuclei, may occur in domestic and industrial wastewaters, natural waters, and potable water supplies. Phenols are distilled from nonvolatile impurities. Because the volatilization of phenols is gradual, the distillate volume must ultimately equal that of the original sample. The method is approved in Table IB for preliminary treatment of samples to be used for determination of phenols.

b. 5530 D–2021, Colorimetric Method. Steam-distillable phenolic compounds react with 4-aminoantipyrine at pH 7.9 \pm 0.1 in the presence of potassium ferricyanide to form a colored antipyrine dye. This dye is kept in aqueous solution and the absorbance is measured photometrically at 500 nm. The method is approved in Table IB for determination of phenol. Note that for regulatory compliance monitoring required under the Clean Water Act, the colorimetric reaction must be performed at a pH of 10.0 \pm 0.2 as stated in 40 CFR 136.3, Table IB, footnote 27.

33. 5540 Surfactants. In 5540 C–2021 this colorimetric method comprises three successive extractions from an acid aqueous medium containing excess methylene blue into chloroform (CHCl₃), followed by an aqueous backwash and measurement of the blue color in the CHCl₃ by spectrophotometry at 652 nm. The method is applicable to methylene blue active substances at concentrations down to about 0.025 mg/L. The method is approved in Table IB for determination of surfactants.

34. 6200 Volatile Organic Compounds Series.

a. 6200 B-2020, Purge and Trap Capillary-Column Gas Chromatographic/Mass Spectrometric (GC/MS) Method. Volatile organic compounds are transferred efficiently from the aqueous to the gaseous phase by bubbling an inert gas (*e.g.*, helium) through a water sample contained in a specially designed purging chamber at ambient temperature. The vapor is swept through a sorbent trap that adsorbs the analytes of interest. After purging is complete, the trap is heated and backflushed with the same inert gas to desorb the compounds onto a gas chromatographic column. The gas chromatograph is temperatureprogrammed to separate the compounds. The detector is a mass spectrometer. The method is approved in Table IC for determination of

benzene, bromodichloromethane, bromoform, bromomethane, carbon tetrachloride, chlorobenzene, chloroethane, chloroform, chloromethane, dibromochloromethane, 1,2-dichlorobenzene, 1,3dichlorobenzene, 1,4-dichlorobenzene, dichlorodifluoromethane, 1,1dichloroethane, 1,2-dichloroethane, 1,1dichloroethene, trans-1,2dichloroethene, 1,2-dichloropropane, cis-1,3-dichloropropene, trans-1,3dichloropropene, ethylbenzene, methylene chloride, 1,1,2,2tetrachloroethane, tetrachloroethene, toluene, 1,1,1-trichloroethane, 1,1,2trichloroethane, trichloroethene, trichlorofluoromethane, and vinyl chloride.

b. 6200 C–2020, Purge and Trap Capillary-Column Gas Chromatographic (GC) Method. Volatile organic compounds are transferred efficiently from the aqueous to the gaseous phase by bubbling an inert gas (*e.g.*, helium) through a water sample contained in a specially designed purging chamber at ambient temperature. The vapor is swept through a sorbent trap that adsorbs the analytes of interest. After purging is complete, the trap is heated and backflushed with the same inert gas to desorb the compounds onto a gas chromatographic column. The gas chromatograph is temperatureprogrammed to separate the compounds and detected using a photoionization detection and an electrolytic conductivity detection in series. The method is approved in Table IC for determination of benzene, bromodichloromethane, bromoform, bromomethane, carbon tetrachloride, chlorobenzene, chloroethane, chloroform, chloromethane, dibromochloromethane, 1,2dichlorobenzene, 1,3-dichlorobenzene, 1,4-dichlorobenzene, 1,1dichloroethane, 1,2-dichloroethane, 1,1dichloroethene, trans-1,2dichloroethene, 1,2-dichloropropane, cis-1,3-dichloropropene, trans-1,3dichloropropene, ethylbenzene, methylene chloride, 1,1,2,2tetrachloroethane, tetrachloroethene, toluene, 1,1,1-trichloroethane, 1,1,2trichloroethane, trichloroethene, trichlorofluoromethane, and vinyl chloride.

35. 6410 Extractable Base/Neutrals and Acids 6410 B–2020, Liquid–Liquid Extraction Gas Chromatographic/Mass Spectrometric Method. This method is applicable to the determination of organic compounds that are partitioned into an organic solvent and are amenable to gas chromatography in municipal and industrial discharges. A measured volume of sample is extracted

serially with methylene chloride at a pH of approximately 2 and again at pH 11. The extract is dried, concentrated, and analyzed by GC/MS. Qualitative compound identification is based on retention time and relative abundance of three characteristic masses (m/z). Quantitative analysis uses internalstandard techniques with a single characteristic m/z. This revision adds a note that although the method was validated extracting base-neutrals first and then acids, performance may be improved by extracting acids first and then base-neutrals. In addition, the EPA is approving method 6410–B for endrin aldehyde in Table ID. This parameter was inadvertently left off the 2007 MUR rulemaking (72 FR 11200, March 12, 2007). The method is approved in Table IC for determination of acenaphthene, acenaphthylene, anthracene, benzidine, benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(g,h,i)perylene, benzo(k)fluoranthene, butyl benzyl phthalate, bis(2-chloroethoxy) methane, bis(2-chloroethyl) ether, bis(2ethylhexyl) phthalate, bromodichloromethane, 4-bromophenvl phenyl ether, 4-chloro-3-methyl phenol, 2-chloronaphthalene, 2-chlorophenol, 4chlorophenyl phenyl ether, chrysene, dibenzo(a,h)anthracene, 3,3'dichlorobenzidine, 2,4-dichlorophenol, diethyl phthalate, 2,4-dimethylphenol, dimethyl phthalate, di-n-butyl phthalate, di-n-octyl phthalate, 2, 4dinitrophenol, 2,4-dinitrotoluene, 2,6dinitrotoluene, fluoranthene, fluorene, hexachlorobenzene, hexachlorobutadiene, hexachlorocyclopentadiene, hexachloroethane, indeno(1,2,3-c,d) pyrene, isophorone, 2-methyl-4,6dinitrophenol, naphthalene, nitrobenzene, 2-nitrophenol, 4nitrophenol, N-nitrosodimethylamine, n-nitrosodi-n-propylamine, nnitrosodiphenylamine, 2,2'-oxybis(1chloropropane), PCB-1016, PCB-1221, PCB-1232, PCB-1242, PCB-1248, PCB-1254, PCB-1260, pentachlorophenol, phenanthrene, phenol, pyrene, 1,2,4trichlorobenzene, and 2,4,6trichlorophenol and in Table ID for determination of aldrin, α -BHC, β -BHC, δ-BHC, γ-BHC (lindane), chlordane, 4,4'-DDD, 4,4'-DDE, 4,4'-DDT, dieldrin, endosulfan I, endosulfan II, endosulfan sulfate, endrin, heptachlor, heptachlor epoxide, and toxaphene.

36. 6420 Phenols. 6420 B–2020, Liquid–Liquid Extraction Gas Chromatographic Method. A measured volume of sample is acidified and extracted with methylene chloride. The extract is dried and exchanged to 2propanol during concentration. Target analytes in the extract are separated by gas chromatography and are identified by retention time and measured with a flame ionization detector, or derivatized and measured with an electron capture detector. This revision of the method replaces distilled, deionized water with reagent water, adds that the packed columns used for validation of the method are no longer available or recommended, and includes information on alternative capillary columns that may be used. The method is approved in Table IC for determination of 4-chloro-3methylphenol, 2-chlorophenol, 2,4dichlorophenol, 2,4-dimethylphenol, 2,4-dinitrophenol, 2-methyl-4,6dinitrophenol, 2-nitrophenol, 4nitrophenol, pentachlorophenol, phenol, and 2,4,6-trichlorophenol.

37. 6440 Polynuclear Aromatic Hydrocarbons. 6440 B-2021, Liquid-Liquid Extraction Chromatographic Method. A measured volume of sample is extracted with methylene chloride. The extract is dried, concentrated, and separated by the high-performance liquid chromatographic (HPLC) or gas chromatographic (GC) method. Ultraviolet (UV) and fluorescence detectors are used with HPLC to identify and measure the polynuclear aromatic hydrocarbons. A flame ionization detector is used with GC. The method is approved in Table IC for determination of acenaphthene, acenaphthylene, anthracene, benzo(a)anthracene, benzo(a)pyrene, benzo(b)fluoranthene, benzo(g,h,i)perylene, benzo(k)fluoranthene, chrysene, dibenzo(a,h)anthracene, fluoranthene, fluorene, indeno(1,2,3-c,d)pyrene, naphthalene, phenanthrene, and pyrene. 38. 6630 Organochlorine Pesticides

Series.

a. 6630 B-2021, Liquid-Liquid Extraction Gas Chromatographic Method I. In this procedure, the pesticides are extracted with a mixed solvent, diethyl ether-hexane or methylene chloride-hexane, by either liquid-liquid extraction using a separatory funnel or by continuous liquid-liquid extraction. The extract is concentrated by evaporation and, if necessary, is cleaned up by column adsorption chromatography. The individual pesticides then are separated by gas chromatography and the compounds are measured with an electron capture detector (ECD). This revision of the method adds information regarding alternative capillary columns that may be used in place of the packed columns that were used for validation of the method, removes information

regarding preparation of packed columns, replaces information regarding the manual injection technique with use of an autosampler and states that gas chromatography/mass spectrometry (GC/MS) may be used for confirmatory analyses in place of a second column and ECD detection. There are no other procedural changes. The method is approved in Table ID for determination of aldrin, α -BHC, β -BHC, δ -BHC, γ -BHC (lindane), captan, carbophenothion, chlordane, 4,4'-DDD, 4,4'-DDE, 4,4'-DDT, dichloran, dieldrin, endosulfan I, endosulfan II, endrin, heptachlor, heptachlor epoxide, isodrin, malathion, methoxychlor, mirex, parathion methyl, parathion ethyl, PCNB, strobane, toxaphene, and trifluralin.

b. În 6630 C–2021, Liquid–Liquid Extraction Gas Chromatographic Method II. In this procedure, a measured volume of sample is extracted with methylene chloride either by liquid-liquid extraction using separatory funnels or by continuous liquid–liquid extraction. The extract is dried and exchanged to hexane during concentration. The target analytes are separated by gas chromatography and the compounds are measured with an electron capture detector (ECD). This revision of the method adds information regarding alternative capillary columns that may be used in place of the packed columns that were used for validation of the method, and states that gas chromatography/mass spectrometry (GC/MS) may be used for confirmatory analyses in place of a second column and ECD detection. There are no other procedural changes. The method is approved in Table ID for determination of aldrin, α -BHC, β -BHC, δ -BHC, γ -BHC (lindane), chlordane, 4,4'-DDD, 4,4'-DDE, 4,4⁻DDT, dieldrin, endosulfan I, endosulfan II, endosulfan sulfate, endrin, endrin aldehyde, heptachlor, heptachlor epoxide, isodrin, methoxychlor, mirex, PCNB, strobane, and toxaphene.

39. 6640 Acidic Herbicide Compounds. 6640 B-2021, Micro Liquid–Liquid Extraction Gas Chromatographic Method. A 40-mL sample is adjusted to pH ≥12 with 4 N sodium hydroxide and is kept for 1 hour at room temperature to hydrolyze derivatives. Because the chlorophenoxy acid herbicides are formulated as a variety of esters and salts, the hydrolysis step is required and may not be skipped. The aqueous sample then is acidified with sulfuric acid to pH ≤1 and extracted with 4 mL of methyl tert-butyl ether (MtBE) that contains the internal standard. The chlorinated acids, which have been partitioned into the MtBE, then are converted to methyl esters by

derivatization with diazomethane. The target esters are separated and detected by capillary column gas chromatography using an electron capture detector (GC/ECD). Analytes are quantified using an internal-standardbased calibration curve. The method is approved in Table IC for determination of 2,4–D, 2,4,5–T, and 2,4,5–TP (Silvex).

D. Changes to 40 CFR 136.3 To Include Alternate Test Procedures in Table IC

To promote method innovation, the EPA maintains a program that allows method developers to apply for the EPA review and potential approval of an alternative method to an existing approved method. This alternate test procedure (ATP) program is described for CWA applications at 40 CFR 136.4 and 136.5. The EPA is approving two ATPs for nationwide use. Based on EPA's review, the performance of these ATPs is equally effective as other methods already approved for measurement of 2,3,7,8-substituted tetra- through octa-chlorinated dibenzop-dioxins and dibenzofurans (PCDDs/ PCDFs) in wastewater. The ATP applicants supplied the EPA with study reports that contain the data from their validation studies. These study reports, the final methods, and the letters documenting EPA's review are included as supporting documents in the docket for this rule.

These new methods are: SGS AXYS Method 16130, "Determination of 2,3,7,8-Substituted Tetra- through Octa-Chlorinated Dibenzo-p-Dioxins and Dibenzofurans (CDDs/CDFs) Using Waters and Agilent Gas Chromatography- Mass Spectrometry (GC-MS/MS), Revision 1.0" and Pace Analytical Method PAM-16130-SSI, "Determination of 2,3,7,8-Substituted Tetra- through Octa-Chlorinated Dibenzo-p-Dioxins and Dibenzofurans (CDDs/CDFs) Using Shimadzu Gas Chromatography Mass Spectrometry (GC-MS/MS), Revision 1.1." These ATPs are the results of separate collaborative efforts between SGS AXYS Analytical Services Ltd, and the instrument manufacturers Waters Corporation and Agilent Technologies, and between Pace Analytical Services LLC and the instrument manufacturer Shimadzu Scientific Instruments, Inc. These final methods are heavily adapted from EPA Method 1613B. Neither ATP makes changes to the extraction or cleanup procedures specified in Method 1613B. All required quality control tests (or analogous tests) and associated QC acceptance criteria have been included in both SGS AXYS 16130 and PAM-16130-SSI.

To minimize costs to both the applicants and the Agency where possible, SGS AXYS, Pace Analytical, and the instrument manufacturers who collaborated on these methods worked closely with EPA's CWA ATP Coordinator to design single-laboratory validation studies for these methods. The goal of these validation studies was to demonstrate that all of the performance criteria specified in Method 1613B could be met and that comparable performance could be achieved when using GC-MS/MS instrumentation for determination of PCDDs/PCDFs in extracts from real-

world samples. The ATP methods are available free of charge on their respective websites (sgsaxys.com or pacelabs.com), therefore the ATP methods incorporated by reference are reasonably available.

In these two methods, referred to in the rule as SGS AXYS 16130 and PAM 16130–SSI, each sample is spiked with the same suite of carbon-13 labeled standards prior to extraction and those standards are used for isotope dilution quantitation in the same way as is done in EPA Method 1613B. All of the relevant QC acceptance criteria are the same in the methods as well. The difference between these methods and the approved EPA method (1613B) is the use of an MS/MS detector system that uses Multiple Reaction Monitoring (MRM) in place of a high-resolution mass spectrometer (HRMS) detector system. The GC portions of the methods did not change.

E. Changes to 40 CFR 136.3 To Include New Standard Methods Committee Methods Based on Previously Approved Technologies

The EPA is adding five new methods in furtherance of the National Technology Transfer and Advancement Act of 1995 (NTTAA), Public Law 104– 113, that provides that Federal agencies and departments shall use technical standards developed or adopted by the VCSBs if compliance would not be inconsistent with applicable law or otherwise impracticable. These methods were submitted by Standard Methods and are consistent with other already approved methods. As discussed in Section IV. B and C of this preamble, these methods are reasonably available.

The EPA is adding $4500-CN^- P-2021$, $4500-CN^- Q-2021$, $4500 - CN^- Q-2021$, $4500 - CN^- R-2021$, and $4500-F^- G-2021$ to Table IB for cyanide and fluoride and is adding 5520 G-2021 to Table IB for oil and grease, based on the following reasons:

1. Cyanide. Although method 4500– CN⁻ P–2021, Total Cyanide by Segmented Flow Injection, UV- Irradiation with Gas Diffusion, and Amperometric Measurement is new to Standard Methods for the Examination of Water and Wastewater, it is based on ASTM D7511-12(17), which is approved in Table IB for determination of total cyanide and relies on the same underlying chemistry and determinative technique to determine total cyanide. Total cyanide consists of dissolved HCN, sodium cyanide (NaCN), and various metal-cyanide complexes, which a continuous flow analyzer converts to aqueous HCN by mixing it with sulfuric acid, irradiating with UV light, and precipitating potentially interfering sulfides with bismuth ion. The aqueous HCN is captured in a donor stream that is passed across a hydrophobic gas-permeable membrane, which selectively diffuses the gaseous HCN into a parallel acceptor stream of dilute sodium hydroxide forming dissolved CN⁻. The cyanide ion in this acceptor stream is measured using an amperometric detector, where the cyanide ion dissolves the silver electrode, resulting in a proportional current.

2. 4500–CN[–] Q–2021, Weak and Dissociable Cyanide by Flow Injection, Gas Diffusion, and Amperometric Measurement. Weak and dissociable cyanide consists of dissolved HCN, NaCN, and various metal-cyanide complexes and includes the same forms of cvanide as those measured using other methods approved in Table IB for determination of available cyanide. Analysts pretreat for weak and dissociable cyanide by mixing a sample with ligand reagents. They then inject the sample into a sulfuric acid and bismuth nitrate solution to produce a donor stream containing aqueous dissolved HCN and precipitated sulfide, if sulfide is present. The donor stream is passed across a hydrophobic gaspermeable membrane, which selectively diffuses gaseous HCN into a parallel acceptor stream of dilute sodium hydroxide, forming dissolved CN⁻. The cyanide ion in this acceptor stream is measured using an amperometric detector, where the cyanide ion dissolves the silver electrode, resulting in a proportional current. Although this method is new to Standard Methods for the Examination of Water and Wastewater, it is based on ASTM D6888–16, which is approved in Table IB for determination of available cyanide and relies on the same underlying chemistry and determinative technique to determine available cyanide.

3. 4500–CN[–] R–2021, Free Cyanide by Flow Injection, Gas Diffusion, and Amperometric Measurement. Free

cvanide (FCN) consists of dissolved HCN, NaCN, and the soluble fraction of various metal-cyanide complexes. To determine FCN, analysts pretreat a sample by mixing it with a buffered solution in the pH range of 6 to 8 that simulates the receiving water resulting in a donor stream containing aqueous dissolved HCN in equilibrium with the cyanide anion. The donor stream is passed across a hydrophobic gaspermeable membrane, which selectively diffuses gaseous HCN into a parallel acceptor stream that consists of dilute sodium hydroxide, forming dissolved CN⁻. The cyanide ions in this acceptor stream are measured when it is passed through an amperometric detector, where the cyanide ion dissolves the silver electrode, resulting in a proportional current. Although this method is new to Standard Methods for the Examination of Water and Wastewater, it is based on ASTM D7237-15, which is approved in Table IB for determination of free cvanide and relies on the same underlying chemistry and determinative technique to determine free cvanide.

4. Fluoride. 4500–F⁻ G–2021, Ion-Selective Electrode Flow Injection Analysis is an automated version of method 4500-F⁻ C and relies on the same underlying chemistry and determinative technique as USGS Method I-4237-85, which currently is approved in Table IB for determination of fluoride. Fluoride is determined potentiometrically by using a combination fluoride ion selective electrode (ISE) in a flow cell. The fluoride electrode consists of a lanthanum fluoride crystal across which a potential is developed by fluoride ions

5. Oil and Grease. In 5520 G–2021, Solid-Phase, Partition-Gravimetric Method, dissolved or emulsified oil and grease is extracted from water by passing a sample through a solid-phase extraction (SPE) disk where the oil and grease are adsorbed by the disk and subsequently eluted with n-hexane. SPE is a modification allowed under EPA Methods 1664 A and B and relies on the same underlying chemistry and determinative technique as Methods 1664 A and B. Some extractables, especially unsaturated fats and fatty acids, oxidize readily; hence, special precautions regarding temperature and solvent vapor displacement are provided. This method is not applicable to materials that volatilize at temperatures below 85 °C, or crude and heavy fuel oils containing a significant percentage of material not soluble in nhexane. This method may be a satisfactory alternative to liquid-liquid

extraction techniques, especially for samples that tend to form difficult emulsions during the extraction step.

V. Statutory and Executive Order Reviews

A. Executive Order 12866: Regulatory Planning and Review and Executive Order 14094: Modernizing Regulatory Review

This action is not a significant regulatory action as defined in Executive Order 12866, as amended by Executive Order 14094, and was therefore not subject to a requirement for Executive Order 12866 review.

B. Paperwork Reduction Act

This action does not impose an information collection burden under the Paperwork Reduction Act. This rule does not impose any information collection, reporting, or recordkeeping requirements. This rule merely revises or adds alternate CWA test procedures.

C. Regulatory Flexibility Act (RFA)

I certify that this action will not have a significant economic impact on a substantial number of small entities under the RFA. This action will not impose any requirements on small entities. This action would approve new alternate and revised versions of CWA testing procedures. Generally, these changes would have a positive impact on small entities by increasing method flexibility, thereby allowing entities to reduce costs by choosing more costeffective methods. In general, the EPA expects the revisions would lead to few, if any, increased costs. The changes clarify or improve the instructions in the method, update the technology used in the method, improve the QC instructions, make editorial corrections, or reflect the most recent approval year of an already approved method. In some cases, the rule adds alternatives to currently approved methods for a particular analyte (e.g., ASTM Method D7511). Because these methods would be alternatives rather than requirements, there are no direct costs associated with the methods approved by the EPA and incorporated by reference. If a permittee elected to use these methods, they could incur a small cost associated with obtaining these methods from the listed sources. See Sections IV. A through D of this preamble.

D. Unfunded Mandates Reform Act

This action does not contain any unfunded mandate as described in the Unfunded Mandates Reform Act, 2 U.S.C. 1531–1538, and does not significantly or uniquely affect small governments. The action imposes no enforceable duty on any state, local or tribal governments or the private sector.

E. Executive Order 13132: Federalism

This final rule does not have federalism implications. It will not have substantial direct effects on the states, on the relationship between the national government and the states, or on the distribution of power and responsibilities among the various levels of government.

F. Executive Order 13175: Consultation and Coordination With Indian Tribal Governments

This action does not have tribal implications as specified in Executive Order 13175. This rule would merely approve new alternate and revised versions of test procedures. The EPA has concluded that the final rule would not lead to any costs to any tribal governments, and if incurred, the EPA projects they would be minimal. Thus, Executive Order 13175 does not apply to this action.

G. Executive Order 13045: Protection of Children From Environmental Health Risks and Safety Risks

The EPA interprets Executive Order 13045 as applying only to those regulatory actions that concern environmental health or safety risks that the EPA has reason to believe may disproportionately affect children, per the definition of "covered regulatory action" in section 2–202 of the Executive Order.

Therefore, this action is not subject to Executive Order 13045 because it does not concern an environmental health risk or safety risk. Since this action does not concern human health, EPA's Policy on Children's Health also does not apply.

H. Executive Order 13211: Actions That Significantly Affect Energy Supply, Distribution, or Use

This action is not subject to Executive Order 13211 because it is not a significant regulatory action under Executive Order 12866.

I. National Technology Transfer and Advancement Act of 1995

This action involves technical standards. The EPA is approving the use of technical standards developed and recommended by the Standard Methods Committee and ASTM International for use in compliance monitoring where the EPA determined that those standards meet the needs of CWA programs. As described above, this final rule is consistent with the NTTAA.

J. Executive Order 12898: Federal Actions To Address Environmental Justice in Minority Populations and Low-Income Populations

The EPA believes that this type of action does not concern human health or environmental conditions and therefore cannot be evaluated with respect to potentially disproportionate and adverse effects on communities with environmental justice concerns. This action has no effect on communities because this action will approve new alternate and revised versions of CWA testing procedures. These changes would provide increased flexibility for the regulated community in meeting monitoring requirements while improving data quality. In addition, this update to the CWA methods will incorporate technological advances in analytical technology. Although this action does not concern human health or environmental conditions, the EPA identifies and addresses environmental justice concerns by identifying and addressing, as appropriate, disproportionately high and adverse human health or environmental effects of their programs, policies, and activities on minority populations (people of color) and lowincome populations.

K. Congressional Review Act

This action is subject to the Congressional Review Act and the EPA will submit a rule report to each House of the Congress and to the Comptroller General of the United States. This is not a "major rule" as defined by 5 U.S.C. 804(2).

List of Subjects in 40 CFR Part 136

Environmental protection, Incorporation by reference, Reporting and recordkeeping requirements, Test procedures, Water pollution control.

Michael S. Regan,

Administrator.

For the reasons set forth in the preamble, the EPA amends 40 CFR part 136 as follows:

PART 136—GUIDELINES ESTABLISHING TEST PROCEDURES FOR THE ANALYSIS OF POLLUTANTS

■ 1. The authority citation for part 136 continues to read as follows:

Authority: Secs. 301, 304(h), 307 and 501(a), Pub. L. 95–217, 91 Stat. 1566, *et seq.* (33 U.S.C. 1251, *et seq.*) (the Federal Water Pollution Control Act Amendments of 1972 as amended by the Clean Water Act of 1977).

■ 2. Amend § 136.3 by:

■ a. In paragraph (a), revising tables IA, IB, IC, ID, and IH;

■ b. Revising paragraph (b) introductory text;

■ c. Revising and republishing paragraphs (b)(8), (10), (15), (19), (26), and (27);

■ d. Redesignating paragraphs (b)(33) through (39) as paragraphs (b)(35) through (41);

■ e. Adding new paragraphs (b)(33) and (34);

■ f. Revising the newly redesignated paragraphs (b)(40) introductory text, (b)(40)(ii), (ix), and (xiv); and

■ g. In paragraph (e), table II, revising Footnote "5".

The revisions and additions read as follows:

§ 136.3 Identification of test procedures.

(a) * * *

TABLE IA—LIST OF APPROVED BIOLOGICAL METHODS FOR WASTEWATER AND SEWAGE SLUDGE

Parameter and units	nd units Method 1		Standard methods	AOAC, ASTM, USGS	Other				
	Bacteria								
 Coliform (fecal), number per gram dry weight. 	Most Probable Number (MPN), 5 tube, 3 dilution, or. Membrane filter (MF), ²⁵ single step	p. 132, ³ 1680, ^{11 15} 1681 ^{11 20} p. 124 ³	9221 E-2014. 9222 D-2015. ²⁹						
2. Coliform (fecal), number per 100 mL	MPN, 5 tube, 3 dilution, or		9221 E–2014, 9221 F–2014. ³³						
	Multiple tube/multiple well, or				Colilert- 18 [®] . ^{13 18 28}				
3. Coliform (total), number per 100 mL	MF, ²⁵ single step ⁵ MPN, 5 tube, 3 dilution, or		9222 D–2015 ²⁹ 9221 B–2014.	B-0050-85.4					
	MF, ²⁵ single step or MF, ²⁵ two step with enrichment			B-0025-85.4					

TABLE IA—LIST OF APPROVED BIOLOGICAL METHODS FOR WASTEWATER AND SEWAGE SLUDGE—Continued

Parameter and units	Method ¹	EPA	Standard methods	AOAC, ASTM, USGS	Other
4. <i>E. coli,</i> number per 100 mL	MPN 6816 multiple tube, or		9221 B2014/9221 F-		
	multiple tube/multiple well, or		2014. ^{12 14 33} 9223 B–2016 ¹³	991.15 ¹⁰	Colilert®. ^{13 18}
					Colilert-
	MF, ²⁵⁶⁷⁸ two step, or		9222 B-2015/9222 I-		18 [®] . ^{13 17 18}
	Single step	1603.1 ²¹	2015. ³¹		m-ColiBlue24 [®] . ¹⁹
5. Fecal streptococci, number per 100	MPN, 5 tube, 3 dilution, or	p. 139 ³	9230 B–2013.		111-COllBlde24
mL.	MF, ² or	p. 136 ³	9230 C–2013 ³²	B-0055-85.4	
	Plate count	p. 143. ³			
 Enterococci, number per 100 mL 	MPN, 5 tube, 3 dilution, or MPN, ^{6 8} multiple tube/multiple well, or.	p. 139 ³	9230 B–2013. 9230 D–2013	D6503–99 ⁹	Enterolert [®] . ^{13 23}
	MF ²⁵⁶⁷⁸ single step or Plate count	1600.1 ²⁴ p. 143. ³	9230 C-2013.32		
7. Salmonella, number per gram dry	MPN multiple tube	1682. ²²			
weight ¹¹ .					
	Aqua	tic Toxicity			
3. Toxicity, acute, fresh water orga-	Water flea, Cladoceran,	2002.0.25			
nisms, LC_{50} , percent effluent.	Ceriodaphnia dubia acute. Water flea, Cladocerans, Daphnia	2021.0.25			
	pulex and Daphnia magna acute.	0000 0 25			
	Fish, Fathead minnow, <i>Pimephales promelas,</i> and Bannerfin shiner,	2000.0.25			
	<i>Cyprinella leedsi,</i> acute. Fish, Rainbow trout, <i>Oncorhynchus</i>	2019.0. ²⁵			
	mykiss, and brook trout,	2010.0.			
 Toxicity, acute, estuarine and ma- rine organisms of the Atlantic Ocean and Gulf of Mexico, LC₅₀, percent 	Salvelinus fontinalis, acute. Mysid, Mysidopsis bahia, acute	2007.0. ²⁵			
effluent.	Fish Chaspeheed minney	0004.0.25			
	Fish, Sheepshead minnow, <i>Cyprinodon variegatus,</i> acute.	2004.0.25			
	Fish, Silverside, <i>Menidia beryllina,</i> <i>Menidia menidia</i> , and <i>Menidia</i> <i>peninsulae</i> , acute.	2006.0.25			
10. Toxicity, chronic, fresh water orga- nisms, NOEC or IC ₂₅ , percent efflu-	Fish, Fathead minnow, <i>Pimephales</i> promelas, larval survival and	1000.0. ²⁶			
ent.	growth. Fish, Fathead minnow, <i>Pimephales</i> <i>promelas</i> , embryo-larval survival	1001.0. ²⁶			
	and teratogenicity. Water flea, Cladoceran,	1002.0.26			
	Ceriodaphnia dubia, survival and	1002.0.20			
	reproduction. Green alga, <i>Selenastrum</i> <i>capricornutum</i> , growth.	1003.0. ²⁶			
11. Toxicity, chronic, estuarine and marine organisms of the Atlantic Ocean and Gulf of Mexico, NOEC or	<i>Cyprinodon variegatus,</i> larval survival and growth	1004.0.27			
IC ₂₅ , percent effluent.	C C	1005 0 37			
	Fish, Sheepshead minnow, <i>Cyprinodon variegatus</i> , embryo-	1005.0.27			
	larval survival and teratogenicity. Fish, Inland silverside, Menidia	1006.0.27			
	<i>beryllina,</i> larval survival and growth. Mysid, <i>Mysidopsis bahia,</i> survival,	1007.0.27			
	growth, and fecundity. Sea urchin, <i>Arbacia punctulata,</i> fer-	1008.0.27			

Table IA notes:

¹ The method must be specified when results are reported. ² A 0.45-µm membrane filter (MF) or other pore size certified by the manufacturer to fully retain organisms to be cultivated and to be free of extractables which could interfere with their growth. ³ Microbiological Methods for Monitoring the Environment, Water and Wastes, EPA/600/8–78/017. 1978. US EPA. ⁴ U.S. Geological Survey Techniques of Water-Resource Investigations, Book 5, Laboratory Analysis, Chapter A4, Methods for Collection and Analysis of Aquatic Biological and Microbiological Samples. 1989. USGS. ⁵ Breaven the Met tophene unreliable word works are environment to method works and the resource the Methods for Collection and Analysis of Aquatic

⁵ Because the MF technique usually yields low and variable recovery from chlorinated wastewaters, the Most Probable Number method will be required to resolve

^a Because the MiF technique usually yields low and variable recovery from chlorinated wastewaters, the Most Probable Number method will be required to resolve any controversies. ^a Tests must be conducted to provide organism enumeration (density). Select the appropriate configuration of tubes/filtrations and dilutions/volumes to account for the quality, character, consistency, and anticipated organism density of the water sample. ⁷ When the MF method has been used previously to test waters with high turbidity, large numbers of noncoliform bacteria, or samples that may contain organisms stressed by chlorine, a parallel test should be conducted with a multiple-tube technique to demonstrate applicability and comparability of results. ⁸ To assess the comparability of results obtained with individual methods, it is suggested that side-by-side tests be conducted across seasons of the year with the water samples routinely tested in accordance with the most current *Standard Methods for the Examination of Water and Wastewater* or EPA alternate test procedure (ATD) middlinge

(ATP) guidelines.

 ⁹Annual Book of ASTM Standards—Water and Environmental Technology, Section 11.02. 2000, 1999, 1996. ASTM International.
 ¹⁰Official Methods of Analysis of AOAC International. 16th Edition, 4th Revision, 1998. AOAC International.
 ¹¹Recommended for enumeration of target organism in sewage sludge.
 ¹²The multiple-tube fermentation test is used in 9221B.2–2014. Lactose broth may be used in lieu of lauryl tryptose broth (LTB), if at least 25 parallel tests are conducted between this broth and LTB using the water samples normally tested, and this comparison demonstrates that the false-positive rate and false-negative rate for total coliform using lactose broth is less than 10 percent. No requirement exists to run the completed phase on 10 percent of all total coliform-positive tubes on a sea-oracle between the served base. sonal basis

¹³These tests are collectively known as defined enzyme substrate tests. ¹⁴After prior enrichment in a presumptive medium for total coliform using 9221B.2–2014, all presumptive tubes or bottles showing any amount of gas, growth or acidity within 48 h ± 3 h of incubation shall be submitted to 9221F–2014. Commercially available EC–MUG media or EC media supplemented in the laboratory with

¹⁵ Method 1680: Fecal Coliforms in Sewage Sludge (Biosolids) by Multiple-Tube Fermentation Using Lauryl-Tryptose Broth (LTB) and EC Medium, EPA-821-R-14-009. September 2014. U.S. EPA.

¹⁶Samples shall be enumerated by the multiple-tube or multiple-well procedure. Using multiple-tube procedures, employ an appropriate tube and dilution configura-tion of the sample as needed and report the Most Probable Number (MPN). Samples tested with Colilert[®] may be enumerated with the multiple-well procedures, Quanti-Tray[®] or Quanti-Tray[®]/2000 and the MPN calculated from the table provided by the manufacturer.

¹⁷Colilert-18[®] is an optimized formulation of the Colilert[®] for the determination of total coliforms and *E. coli* that provides results within 18 h of incubation at 35°C ¹⁷ Collert-18[®] is an optimized formulation of the Colliert[®] for the determination of total coliforms and *E. coli* that provides results within 18 h of incubation at 35°C rather than the 24 h required for the Colliert[®] test and is recommended for marine water samples.
 ¹⁸ Descriptions of the Colliert[®], Colliert-18[®], Quanti-Tray[®], and Quanti-Tray[®]/2000 may be obtained from IDEXX Laboratories, Inc.
 ¹⁹ A description of the mColliBlue24[®] test is available from Hach Company.
 ²⁰ Method 1681: Fecal Coliforms in Sewage Sludge (Biosolids) by Multiple-Tube Fermentation Using A–1 Medium, EPA–821–R–06–013. July 2006. U.S. EPA.
 ²¹ Method 1603.1: *Escherichia coli* (*E. coli*) in Water by Membrane Filtration Using Modified membrane-Thermotolerant *Escherichia coli* Agar (Modified mTEC), EPA–821–R–23–008. September 2023. U.S. EPA.
 ²² Method 1682: *Salmonella* in Sewage Sludge (Biosolids) by Modified Semisolid Rappaport-Vassiliadis (MSRV) Medium, EPA–821–R–14–012. September 2014.

U.S. EPA.

A description of the Enterolert[®] test may be obtained from IDEXX Laboratories Inc. Method 1600.1: Enterococci in Water by Membrane Filtration Using Membrane-Enterococcus Indoxyl-β-D-Glucoside Agar (mEl), EPA-821-R-23-006. Sep-24 Method 1600.1: tember 2023. U.S. EPA.

²⁵Methods for Measuring the Acute Toxicity of Effluents and Receiving Waters to Freshwater and Marine Organisms, EPA–821–R–02–012. Fifth Edition, October 2002. U.S. EPA; and U.S. EPA Whole Effluent Toxicity Methods Errata Sheet, EPA 821–R–02–012–ES. December 2016.
 ²⁶Short-term Methods for Estimating the Chronic Toxicity of Effluents and Receiving Waters to Freshwater Organisms, EPA–821–R–02–013. Fourth Edition, October 2002. U.S. EPA; and U.S. EPA Whole Effluent Toxicity Methods Errata Sheet, EPA 821–R–02–012–ES. December 2016.

 ²⁷Short-term Methods for Estimating the Chronic Toxicity of Effluents and Receiving Waters to Marine and Estuarine Organisms, EPA-821–R–02–014. Third Edition, October 2002. U.S. EPA; and U.S. EPA Whole Effluent Toxicity Methods Errata Sheet, EPA 821–R–02–012–ES. December 2016.
 ²⁸To use Colliert-18® to assay for fecal coliforms, the incubation temperature is 44.5 ± 0.2 °C, and a water bath incubator is used.
 ²⁹On a monthly basis, at least ten blue colonies from positive samples must be verified using Lauryl Tryptose Broth. Where possible, verifications should be done from randomic accurate accurate. domized sample sources.

30 On a monthly basis, at least ten sheen colonies from positive samples must be verified using lauryl tryptose broth and brilliant green lactose bile broth, followed by count adjustment based on these results; and representative non-sheen colonies should be verified using lauryl tryptose broth. Where possible, verifications should be done from randomized sample sources.

³¹ Subject coliform positive samples determined by 9222 B–2015 or other membrane filter procedure to 9222 I–2015 using NA–MUG media. ³² Verification of colonies by incubation of BHI agar at 10 ± 0.5 °C for 48 ± 3 h is optional. As per the Errata to the 23rd Edition of *Standard Methods for the Exam-ination of Water and Wastewater* "Growth on a BHI agar plate incubated at 10 ± 0.5 °C for 48 ± 3 h is further verification that the colony belongs to the genus

Enterococcus." 339221F. 2-2014 allows for simultaneous detection of *E. coli* and thermotolerant fecal coliforms by adding inverted vials to EC-MUG; the inverted vials collect gas produced by thermotolerant fecal coliforms.

Parameter	Methodology 58	EPA 52	Standard methods 84	ASTM	USGS/AOAC/Other
1. Acidity (as CaCO ₃), mg/L.	Electrometric endpoint or phenolphthalein endpoint.		2310 B-2020	D1067–16	I-1020-85. ²
 Alkalinity (as CaCO₃), mg/L. 	Electrometric or Colorimetric titration to pH 4.5. Manual.		2320 B–2021	D1067–16	973.43, ³ I-1030-85. ²
 Aluminum—Total,⁴ mg/L. 	Automatic Digestion, ⁴ followed by any of the fol- lowing:	310.2 (Rev. 1974) ¹			I–2030–85. ²
	AA direct aspiration ³⁶		3111 D–2019 or 3111 E–2019.		I–3051–85. ²
	AA furnace STGFAA	200.9 Rev. 2.2	3113 B–2020.		
	ICP/AES ³⁶	(1994). 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994).	3120 B–2020	D1976–20	I–4471–97. ⁵⁰
	ICP/MS	200.8, Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4472– 97. ⁸¹
	Direct Current Plasma (DCP) ³⁶ Colorimetric (Eriochrome cyanine R)		3500–Al B–2020.	D4190–15	
4. Ammonia (as N), mg/L.	Manual distillation ⁶ or gas diffusion (pH > 11), followed by any of the following:	350.1 Rev. 2.0 (1993).	4500–NH ₃ B–2021		973.49. ³
3	Nesslerization			D1426–15 (A)	973.49, ³ I–3520–85. ²
	Electrode		4500–NH ₃ D–2021 or E–2021.	D1426–15 (B)	
	Manual phenate, salicylate, or other sub- stituted phenols in Berthelot reaction- based methods.		4500–NH ₃ F–2021		See footnote.60
	Automated phenate, salicylate, or other substituted phenols in Berthelot reaction- based methods.	350.1, ³⁰ Rev. 2.0 (1993).	4500–NH ₃ G–2021, 4500–NH ₃ H–2021.		I–4523–85, ² I–2522– 90. ⁸⁰
	Automated electrode			 D6919–17.	See footnote.7
	Automated gas diffusion, followed by con- ductivity cell analysis.				Timberline Ammonia- 001.74
	Automated gas diffusion followed by fluo- rescence detector analysis.				FIAlab100.82
 Antimony—Total,⁴ mg/L. 	Digestion, ⁴ followed by any of the fol- lowing:				

2	7	3	0	7

Parameter	Methodology 58	EPA 52	Standard methods ⁸⁴	ASTM	USGS/AOAC/Ot
	AA direct aspiration ³⁶		3111 B-2019.		
	AA furnace		3113 B-2020.		
	STGFAA	200.9 Rev. 2.2			
		(1994).			
	ICP/AES ³⁶	200.5 Rev 4.2	3120 B-2020	D1976–20.	
		(2003),68 200.7			
		Rev. 4.4 (1994).			
	ICP/MS	200.8, Rev. 5.4	3125 B–2020	D5673–16	993.14, ³ I–4472–
		(1994).			97. ⁸¹
. Arsenic—Total,4 mg/	Digestion, ⁴ followed by any of the fol-	206.5 (Issued 1978). ¹			
L.	lowing:		0114 D 0000 or 0114	D0070 15 (D)	I-3062-85.2
	AA gaseous hydride		3114 B–2020 or 3114 C–2020.	D2972–15 (B)	1-3062-65.4
	AA furnace		3113 B-2020	D2972–15 (C)	I–4063–98.49
	STGFAA	200.9 Rev. 2.2	5115 D=2020	D2372-13 (0)	1-4000-30.
		(1994).			
	ICP/AES ³⁶	200.5, Rev 4.2	3120 B-2020	D1976–20.	
		(2003),68 200.7			
		Rev. 4.4 (1994).			
	ICP/MS	200.8, Rev. 5.4	3125 B-2020	D5673–16	993.14,3 I-4020-
		(1994).			05.70
	Colorimetric (SDDC)		3500–As B–2020	D2972–15 (A)	I-3060-85. ²
. Barium—Total,4 mg/	Digestion, ⁴ followed by any of the fol-				
L.	lowing:				
	AA direct aspiration ³⁶		3111 D-2019		I-3084-85. ²
	AA furnace		3113 B-2020	D4382–18.	
	ICP/AES ³⁶	200.5, Rev 4.2	3120 B-2020		I–4471–97. ⁵⁰
		(2003),68 200.7			
		Rev. 4.4 (1994).			
	ICP/MS		3125 B–2020	D5673–16	993.14, ³ I–4472–
	DOD 26	(1994).			97. ⁸¹
D # T 14	DCP ³⁶				See footnote.34
. Beryllium—Total, ⁴	Digestion, ⁴ followed by any of the fol-				
mg/L.	lowing:		0111 D 0010 at		I-3095-85.2
	AA direct aspiration		3111 D-2019 or	D3645–15 (A)	1-3095-65.4
	AA furnace		3111 E-2019. 3113 B-2020	D3645–15 (B).	
	STGFAA	200.9, Rev. 2.2	3113 D=2020	D3043-13 (D).	
		(1994).			
	ICP/AES	200.5 Rev 4.2	3120 B-2020	D1976–20	I-4471-97. ⁵⁰
		(2003),68 200.7	0120 0 2020	D1070 20	1 1 1 1 07.
		Rev. 4.4 (1994).			
	ICP/MS	200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14,3 I-4472-
		(1994).			97. ⁸¹
	DCP			D4190–15	See footnote.34
	Colorimetric (aluminon)		See footnote 61		
. Biochemical oxygen	Dissolved Oxygen Depletion		5210 B-2016 ⁸⁵		973.44 ³ p. 17, ⁹ l
demand (BOD ₅), mg/					1578–78, ⁸ see
			4500 D D 0011		footnote. ¹⁰⁶³
0. Boron—Total, ³⁷	Colorimetric (curcumin)		4500–B B–2011		I–3112–85. ²
mg/L.	ICP/AES	200 5 Poy 4 2	2120 B 2020	D1976–20	1 4471 07 50
		(2003),68 200.7	3120 B-2020	D1976-20	1-4471-97.00
		Rev. 4.4 (1994).			
	ICP/MS	200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14. ³
		(1994).	0120 0 2020	20070 10	000.14.
	DCP			D4190–15	See footnote.34
1. Bromide, mg/L	Electrode			D1246–16	I-1125-85.2
, <u> </u>	Ion Chromatography	300.0 Rev 2.1	4110 B-2020, C-	D4327–17	993.30, ³ I–2057–
		(1993), and 300.1	2020 or D-2020.		85. ⁷⁹
		Rev 1.0 (1997).			
	CIE/UV		4140 B–2020	D6508–15	D6508 Rev. 2.54
2. Cadmium—Total,4	Digestion, ⁴ followed by any of the fol-				
2. Cadmium—Total, ⁴ mg/L.	lowing:				
,			3111 B–2019 or 3111	D3557–17 (A or B)	
,	lowing:		3111 B–2019 or 3111 C–2019.	D3557–17 (A or B)	3135-85 ² or I-
,	lowing: AA direct aspiration ³⁶		C–2019.		3135–852 or I– 3136–85.2
,	lowing: AA direct aspiration ³⁶			D3557–17 (A or B) D3557–17 (D)	3135-85 ² or I-
,	lowing: AA direct aspiration ³⁶		C–2019.		3135–852 or I– 3136–85.2
2. Cadmium—Total,4 mg/L.	lowing: AA direct aspiration ³⁶ AA furnace STGFAA	 200.9 Rev. 2.2 (1994)	C–2019. 3113 B–2020	D3557–17 (D)	I–4138–89. ⁵¹
,	lowing: AA direct aspiration ³⁶	200.9 Rev. 2.2 (1994). 200.5 Rev 4.2	C–2019.		3135–85 ² or I– 3136–85. ² I–4138–89. ⁵¹
,	lowing: AA direct aspiration ³⁶ AA furnace STGFAA	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003), ⁶⁸ 200.7	C–2019. 3113 B–2020	D3557–17 (D)	3135–85 ² or l- 3136–85. ² I–4138–89. ⁵¹
,	lowing: AA direct aspiration ³⁶ AA furnace STGFAA ICP/AES ³⁶	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003). ⁶⁸ 200.7 Rev. 4.4 (1994).	C-2019. 3113 B-2020 3120 B-2020	D3557–17 (D) D1976–20	3135–85 ² or I- 3136–85. ² I–4138–89. ⁵¹ I–1472–85 ² or I– 4471–97. ⁵⁰
,	lowing: AA direct aspiration ³⁶ AA furnace STGFAA	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8, Rev. 5.4	C–2019. 3113 B–2020	D3557–17 (D)	3135–85 ² or I- 3136–85. ² I–4138–89. ⁵¹ I–1472–85 ² or I- 4471–97. ⁵⁰ 993.14, ³ I–4472–
,	lowing: AA direct aspiration ³⁶ AA furnace STGFAA ICP/AES ³⁶	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003). ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8, Rev. 5.4 (1994).	C–2019. 3113 B–2020 3120 B–2020 3125 B–2020	D3557–17 (D) D1976–20 D5673–16	3135-85 ² or I- 3136-85. ² I-4138-89. ⁵¹ I-1472-85 ² or I- 4471-97. ⁵⁰ 993.14, ³ I-4472- 97. ⁸¹
,	Iowing: AA direct aspiration ³⁶ AA furnace STGFAA ICP/AES ³⁶ DCP ³⁶	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8, Rev. 5.4 (1994).	C-2019. 3113 B-2020 3120 B-2020 3125 B-2020	D3557–17 (D) D1976–20 D5673–16 D4190–15	3135-85 ² or I- 3136-85. ² I-4138-89. ⁵¹ I-1472-85 ² or I- 4471-97. ⁵⁰ 993.14, ³ I-4472-
,	Iowing: AA direct aspiration ³⁶ AA furnace STGFAA ICP/AES ³⁶ ICP/MS DCP ³⁶ Voltammetry ¹¹	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8, Rev. 5.4 (1994).	C-2019. 3113 B-2020 3120 B-2020 3125 B-2020	D3557–17 (D) D1976–20 D5673–16	3135-85 ² or I- 3136-85. ² I-4138-89. ⁵¹ I-1472-85 ² or I- 4471-97. ⁵⁰ 993.14, ³ I-4472- 97. ⁸¹
,	Iowing: AA direct aspiration ³⁶ AA furnace STGFAA ICP/AES ³⁶ DCP ³⁶	200.9 Rev. 2.2 (1994) 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8, Rev. 5.4 (1994).	C-2019. 3113 B-2020 3120 B-2020 3125 B-2020	D3557–17 (D) D1976–20 D5673–16 D4190–15	3135-85 ² or I- 3136-85. ² I-4138-89. ⁵¹ I-1472-85 ² or I- 4471-97. ⁵⁰ 993.14, ³ I-4472- 97. ⁸¹

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 Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². Chemical oxygen 	AA direct aspiration ICP/AES ICP/MS	200.5 Rev 4.2 (2003), ⁶⁸ 200.7	3111 B–2019 or 3111 D–2019. 3120 B–2020	D511–14 (B)	I–3152–85. ²
 Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². Chemical oxygen 	ICP/MS	(2003),68 200.7			
 Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². Chemical oxygen 	ICP/MS	(2003),68 200.7			I-4471-97. ⁵⁰
4. Carbonaceous bio- chemical oxygen de- mand (CBOD ₅), mg/ L ¹² . 5. Chemical oxygen		D 4 4 / 100 4			
4. Carbonaceous bio- chemical oxygen de- mand (CBOD ₅), mg/ L ¹² . 5. Chemical oxygen		Rev. 4.4 (1994). 200.8, Rev. 5.4	3125 B–2020	D5673–16	993.14. ³
 Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². Chemical oxygen 	DCD	(1994).	3123 D=2020	D3073-10	355.14.*
 Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². Chemical oxygen 					See footnote.34
 4. Carbonaceous bio- chemical oxygen de- mand (CBOD₅), mg/ L¹². 5. Chemical oxygen 	Titrimetric (EDTA) Ion Chromatography		3500-Ca B–2020	D511–14 (A). D6919–17.	
mand (CBOD ₅), mg/ L ¹² . 5. Chemical oxygen	Dissolved Oxygen Depletion with nitrifica-		5210 B-2016 ⁸⁵		See footnotes.35 63
	tion inhibitor.				
demand (COD), mg/L.	Titrimetric	410.3 (Rev. 1978) ¹	5220 B-2011 or C- 2011.	D1252–06(12) (A)	973.46 ³ p. 17, ⁹ l– 3560–85. ²
	Spectrophotometric, manual or automatic	410.4 Rev. 2.0 (1993).	5220 D-2011	D1252–06(12) (B)	See footnotes, ^{13 14 83} I–3561–85. ²
	Titrimetric: (silver nitrate)		4500-Cl ⁻ B-2021 4500-Cl ⁻ C-2021	D512–12 (B) D512–12 (A)	l–1183–85. ² 973.51, ³ l–1184–85.
	Colorimetric: manual		+000 01 0 2021	DOTE TE (A)	I–1187–85. ²
	Automated (ferricyanide)		4500-CI- E-2021		I-2187-85.2
	Potentiometric Titration		4500-CI- D-2021.		
	Ion Selective Electrode			D512–12 (C).	
	Ion Chromatography	300.0 Rev 2.1 (1993), and 300.1	4110 B–2020 or 4110	D4327–17	993.30, ³ I–2057– 90. ⁵¹
		Rev 1.0 (1997).	C–2020.		90.51
	CIE/UV		4140 B-2020	D6508–15	D6508, Rev. 2. ⁵⁴
	Amperometric direct		4500-CI D-2011	D1253–14.	
sidual, mg/L.	Amperometric direct (low level)		4500-CI E-2011.		
	Iodometric direct		4500-CI B-2011.		
	Back titration ether end-point 15		4500-CI C-2011.		
	DPD-FAS		4500-CI F-2011.		
	Spectrophotometric, DPD		4500-Cl G–2011.		Can fastasta 16
	Electrode Amperometric direct		 4500-Cl D–2011	D1253–14	See footnote. ¹⁶
	Amperometric direct (low level)		4500-CI E-2011.		
	DPD-FAS		4500-CI F-2011.		
	Spectrophotometric, DPD 0.45-micron filtration followed by any of the following:		4500-Cl G–2011.		
	AA chelation-extraction Ion Chromatography		3111 C–2019 3500-Cr C–2020	D5257–17	I–1232–85. ² 993.23. ³
	Colorimetric (diphenyl-carbazide)	(1994).	3500-Cr B–2020	D1687–17 (A)	I-1230-85.2
	Digestion, ⁴ followed by any of the fol-		5500-01 D-2020	D1007-17 (A)	1-1230-03.
mg/L.	lowing:				
	AA direct aspiration ³⁶		3111 B-2019	D1687–17 (B)	974.27, ³ I–3236–85.
	AA chelation-extraction		3111 C–2019. 3113 B–2020	D1687–17 (C)	1-3233-03 46
	STGFAA	200.9 Rev. 2.2	0110 D 2020	D1007 17 (0)	1 0200 30.
		(1994).			
	ICP/AES ³⁶	200.5 Rev 4.2 (2003), ⁶⁸ 200.7	3120 B–2020	D1976–20.	
	ICP/MS	Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4020– 05 ⁷⁰ I–4472–97. ⁸¹
	DCP ³⁶			D4190–15	See footnote.34
	Colorimetric (diphenyl-carbazide)		3500-Cr B–2020.		
). Cobalt—Total,4 mg/ L.	Digestion, ⁴ followed by any of the fol- lowing:				
	AA direct aspiration		3111 B–2019 or 3111 C–2019.	D3558–15 (A or B)	p. 37, ⁹ I–323985. ²
	AA furnace		3113 B-2020	D3558–15 (C)	I-4243-89.51
	STGFAA	200.9 Rev. 2.2			
	ICP/AES	(1994). 200.7 Rev. 4.4 (1994).	3120 B–2020	D1976–20	I-4471-97. ⁵⁰
	ICP/MS	(1994). 200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4020– 05 ⁷⁰ I–4472–97. ⁸¹
	DCP	(1994).		D4190–15	See footnote. ³⁴
1. Color, platinum co- balt units or domi- nant wavelength,	Colorimetric (ADMI)		2120 F-2021. ⁷⁸	2	
hue, luminance purity.	Platinum cobalt visual comparison		2120 B-2021		I-1250-85.2
	Spectrophotometric				See footnote.18
	Digestion, ⁴ followed by any of the fol- lowing:				

Parameter	Methodology 58	EPA 52	Standard methods 84	ASTM	USGS/AOAC/Other
	AA direct aspiration ³⁶		3111 B-2019 or 3111 C-2019.	D1688–17 (A or B)	974.27, ³ p. 37, ⁹ l– 3270–85 ² or l– 3271–85. ²
	AA furnace STGFAA		3113 B-2020	D1688–17 (C)	I-4274-89. ⁵¹
		(1994).			
	ICP/AES ³⁶	200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994).	3120 B-2020	D1976–20	I–4471–97. ⁵⁰
	ICP/MS	200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4020– 05, ⁷⁰ I–4472–97. ⁸¹
	DCP ³⁶		3500-Cu B–2020.	D4190–15	See footnote.34
23. Cyanide—Total,	Colorimetric (Neocuproine) Colorimetric (Bathocuproine) Automated UV digestion/distillation and		3500-Cu B-2020. 3500-Cu C-2020		See footnote. ¹⁹ Kelada-01. ⁵⁵
mg/L.	Colorimetry. Segmented Flow Injection, In-Line Ultra- violet Digestion, followed by gas diffusion		4500-CN- P-2021	D7511–12 (17)	
	amperometry. Manual distillation with MgCl ₂ , followed by	335.4 Rev. 1.0	4500–CN [–] B–2021	D2036–09(15)(A),	10–204–00–1–X. ⁵⁶
	any of the following: Flow Injection, gas diffusion amperometry	(1993) ⁵⁷ .	and C-2021.	D7284–20. D2036–09(15)(A)	
	Titrimetric		4500–CN [–] D–2021	D7284–20. D2036–09(15)(A)	See footnote 9 p. 22.
	Spectrophotometric, manual		4500–CN [–] E–2021	D2036–09(15)(A)	I-3300-85. ²
	Semi-Automated ²⁰	335.4 Rev. 1.0 (1993) ⁵⁷ .	4500-CN ⁻ N-2021		10-204-00-1-X, ⁵⁶ I- 4302-85. ²
	Ion Chromatography			D2036-09(15)(A).	4302-05
	Ion Selective Electrode		4500-CN - F-2021	D2036–09(15)(A).	
24. Cyanide—Avail- able, mg/L.	Cyanide Amenable to Chlorination (CATC); Manual distillation with MgCl ₂ , followed by Titrimetric or Spectrophotometric.		4500–CN [–] G–2021	D2036–09(15)(B).	
	Flow injection and ligand exchange, fol- lowed by gas diffusion amperometry 59.		4500–CN [–] Q–2021	D6888–16	OIA-1677-09.44
	Automated Distillation and Colorimetry (no UV digestion).				Kelada-01.55
24A. Cyanide—Free, mg/L.	Flow Injection, followed by gas diffusion amperometry.		4500–CN [–] R–2021	D7237–18 (A)	OIA-1677-09.44
25. Fluoride—Total, mg/L.	Manual micro-diffusion and colorimetry Manual distillation, ⁶ followed by any of the following:.		4500–F ⁻ B–2021	D4282–15. D1179–16 (A).	
	Electrode, manual		4500-F - C-2021	D1179–16 (B).	
	Electrode, automated Colorimetric, (SPADNS)		4500–F ⁻ G–2021 4500–F ⁻ D–2021.		I-4327-85. ²
	Automated complexone		4500-F ⁻ E-2021.		
	Ion Chromatography	300.0 Rev 2.1 (1993) and 300.1 Rev 1.0 (1997).	4110 B–2020 or C– 2020.	D4327–17	993.30. ³
26. Gold—Total,⁴ mg/L	CIE/UV Digestion, ⁴ followed by any of the fol-		4140 B-2020	D6508–15	D6508, Rev. 2. ⁵⁴
	lowing: AA direct aspiration		3111 B-2019.		
	AA furnace	231.2 (Issued 1978) ¹	3113 B-2020.		
	ICP/MS	200.8, Rev. 5.4 (1994).	3125 B-2020	D5673–16	993.14. ³
	DCP				See footnote.34
 Hardness—Total (as CaCO₍₃₎, mg/L. 	Automated colorimetric	130.1 (Issued 1971). ¹			
	Titrimetric (EDTA)		2340 C-2021	D1126–17	973.52B, ³ I–1338– 85. ²
	Ca plus Mg as their carbonates, by any ap- proved method for Ca and Mg (See Pa- rameters 13 and 33), provided that the sum of the lowest point of quantitation for Ca and Mg is below the NPDES per-		2340 B-2021.		
00 huden en (, 1)	mit requirement for Hardness				070 44 31 4500 65 3
 Hydrogen ion (pH), pH units. 	Electrometric measurement		4500–H+ B–2021	D1293–18 (A or B)	973.41, ³ I–1586–85. ² See footnote ²¹ I–
29. Iridium—Total,4 mg/	Digestion, ⁴ followed by any of the fol-				2587–85. ²
L.	lowing:. AA direct aspiration		3111 B-2019.		
	AA furnace	235.2 (Issued 1978).1			
30. Iron—Total,4 mg/L	ICP/MS Digestion, ⁴ followed by any of the fol- lowing:		3125 B–2020.		
	AA direct aspiration ³⁶		3111 B-2019 or 3111	D1068–15 (A)	974.27,3 I-3381-85.2
	AA furnace		C–2019. 3113 B–2020	D1068-15 (B)	
	STGFAA	200.9, Rev. 2.2	2020	D1068–15 (B).	
		(1994).			

Parameter	Methodology 58	EPA ⁵²	Standard methods 84	ASTM	USGS/AOAC/Other
	ICP/AES ³⁶	200.5 Rev. 4.2 (2003), ⁶⁸ 200.7	3120 B–2020	D1976–20	I-4471-97. ⁵⁰
		Rev. 4.4 (1994).			
	ICP/MS	200.8, Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14. ³
	DCP ³⁶			D4190–15	See footnote.34
	Colorimetric (Phenanthroline)		3500-Fe B-2011	D1068–15 (C)	
 Kjeldahl Nitro- 	Manual digestion 20 and distillation or gas		4500-N _{org} B-2021 or	D3590–17 (A)	I-4515-91.45
gen 5-Total (as N),	diffusion, followed by any of the fol-		C-2021 and 4500-		
mg/L.	lowing:		NH ₃ B–2021.		070 40 2
	Titration Nesslerization		4500–NH ₃ C–2021	 D1426–15 (A).	973.48. ³
	Electrode		4500–NH ₃ D–2021 or	D1426–15 (R).	
			E-2021.	()	
	Semi-automated phenate	350.1 Rev. 2.0	4500-NH3 G-2021 or		
	Manual phanata, activulata, ar athar aut	(1993).	4500–NH ₃ H–2021.		Cas fastasta 60
	Manual phenate, salicylate, or other sub- stituted phenols in Berthelot reaction		4500–NH ₃ F–2021		See footnote.60
	based methods.				
	Automated gas diffusion, followed by con-				Timberline Ammonia
	ductivity cell analysis.				001.74
	Automated gas diffusion followed by fluo-				FIAlab 100.82
	rescence detector analysis.				
	Automated Methods for TKN that do not require manual distillation				
	Automated phenate, salicylate, or other	351.1 (Rev. 1978) ¹			I-4551-78.8
	substituted phenols in Berthelot reaction-				
	based methods colorimetric (auto diges-				
	tion and distillation)	_			
	Semi-automated block digestor colorimetric	351.2 Rev. 2.0	4500–N _{org} D–2021	D3590–17 (B)	I-4515-91.45
	(distillation not required). Block digester, followed by Auto distillation	(1993).			See footnote.39
	and Titration.				See lootilote.
	Block digester, followed by Auto distillation				See footnote.40
	and Nesslerization.				
	Block Digester, followed by Flow injection				See footnote.41
	gas diffusion (distillation not required).				Upph 10040 76
	Digestion with peroxdisulfate, followed by Spectrophotometric (2,6-dimethyl phenol).				Hach 10242.76
	Digestion with persulfate, followed by Col-				NCASI TNTP
	orimetric.				W10900.77
32. Lead—Total,4 mg/L	Digestion, ⁴ followed by any of the fol-				
	lowing:				074 07 8 4 0000 05 6
	AA direct aspiration ³⁶		3111 B–2019 or 3111 C–2019.	D3559–15 (A or B)	974.27, ³ I–3399–85. ²
	AA furnace		3113 B-2020	D3559–15 (D)	I-4403-89.51
	STGFAA	200.9 Rev. 2.2		20000 10 (2)	1 1100 00.
		(1994).			
	ICP/AES ³⁶	200.5 Rev. 4.2	3120 B-2020	D1976–20	I-4471-97. ⁵⁰
		(2003),68 200.7			
	ICP/MS	Rev. 4.4 (1994). 200.8 Rev. 5.4	3125 B_2020	D5673–16	993.14, ³ I–4472–
		(1994).	5125 D=2020	05075-10	97. ⁸¹
	DCP ³⁶			D4190–15	See footnote.34
	Voltammetry 11			D3559–15 (C).	
	Colorimetric (Dithizone)		3500-Pb B–2020.		
33. Magnesium—	Digestion, ⁴ followed by any of the fol-				
Total, ⁴ mg/L.	lowing: AA direct aspiration		3111 B-2019	D511–14 (B)	974.27,3 I-3447-85.2
	ICP/AES	200.5 Rev. 4.2	3120 B-2020	D1976–20	I-4471-97. ⁵⁰
		(2003),68 200.7		2.0.0 20	
		Rev. 4.4 (1994).			
	ICP/MS	200.8 Rev. 5.4	3125 B–2020	D5673–16	993.14. ³
	DCP	(1994).			See footnote.34
	Ion Chromatography			D6919–17.	See loolilole.
34. Manganese—	Digestion, ⁴ followed by any of the fol-			20010 111	
Total, ⁴ mg/L.	lowing:				
	AA direct aspiration ³⁶		3111 B-2019 or 3111	D858–17 (A or B)	974.27,3 I-3454-85.2
			C-2019.	D050 17 (O)	
	AA furnace STGFAA	200.9 Rev. 2.2	3113 B–2020	D858–17 (C).	
		(1994).			
	ICP/AES ³⁶	200.5, Rev. 4.2	3120 B-2020	D1976–20	I-4471-97.50
		(2003); 68 200.7,			
	100/110	Rev. 4.4 (1994).		D5070 10	000 44 0 4 1
	ICP/MS	200.8 Rev. 5.4	3125 B–2020	D5673–16	993.14, ³ I–4472– 97. ⁸¹
	DCP ³⁶	(1994).		D4190–15	See footnote.34
			3500-Mn B–2020	D4190-15	920.203. ³
	Colorimetric (Persulfate)		3300-IVIII D-2020		520.200.

Parameter	Methodology 58	EPA 52	Standard methods 84	ASTM	USGS/AOAC/Other
35. Mercury—Total,	Cold vapor, Manual	245.1 Rev. 3.0	3112 B-2020	D3223–17	977.22,3 I-3462-85.
mg/L.		(1994).			
	Cold vapor, Automated	245.2 (Issued 1974). ¹			1 4404 01 71
	Cold vapor atomic fluorescence spectrom- etry (CVAFS).	245.7 Rev. 2.0 (2005) ¹⁷ .			I-4464-01. ⁷¹
	Purge and Trap CVAFS	1631E. ⁴³			
36. Molybdenum—	Digestion, ⁴ followed by any of the fol-				
Total,4 mg/L.	lowing:				
	AA direct aspiration		3111 D-2019		I-3490-85. ²
	AA furnace		3113 B-2020	 D1976–20	I-3492-96.47 I-4471-97.50
	ICP/AES	200.7 Rev. 4.4 (1994).	3120 B-2020	D1976-20	1-4471-97.00
	ICP/MS		3125 B-2020	D5673–16	993.14, ³ I–4472–
		(1994).			97. ⁸¹
	DCP				See footnote.34
37. Nickel—Total,4 mg/	Digestion, ⁴ followed by any of the fol-				
L.	lowing: AA direct aspiration ³⁶		3111 B-2019 or 3111	D1886–14 (A or B)	I-3499-85. ²
			C-2019.	D1000-14 (A 01 D)	1-3499-03
	AA furnace		3113 B-2020	D1886–14 (C)	I-4503-89.51
	STGFAA	200.9 Rev. 2.2			
		(1994).			
	ICP/AES ³⁶	200.5 Rev. 4.2	3120 B-2020	D1976–20	I-4471-97. ⁵⁰
		(2003), ⁶⁸ 200.7 Rev. 4.4 (1994).			
	ICP/MS		3125 B-2020	D5673–16	993.14, ³ I–4020–
		(1994).	0120 0 2020	20070 10	05, ⁷⁰ I–4472–97. ⁸
	DCP 36			D4190–15	See footnote.34
38. Nitrate (as N), mg/L	Ion Chromatography	300.0 Rev. 2.1	4110 B-2020 or C-	D4327–17	993.30. ³
		(1993) and 300.1	2020.		
	CIE/UV	Rev. 1.0 (1997).	4140 B–2020	D6508–15	D6508, Rev. 2.54
	Ion Selective Electrode		4500–NO ₃ ⁻ D–2019	D0500-15	D0000, nev. 2.91
	Colorimetric (Brucine sulfate)	352.1 (Issued 1971) ¹			973.50, ³ 419D, ⁸⁶ p.
					28. ⁹
	Spectrophotometric (2,6-dimethylphenol)				Hach 10206.75
	Nitrate-nitrite N minus Nitrite N (see pa-				
39. Nitrate-nitrite (as	rameters 39 and 40). Cadmium reduction, Manual		4500–NO ₃ - E–2019	D3867–16 (B).	
N), mg/L.			4500-NO3 L-2019	D3007-10 (D).	
· · ,, · · · · ·	Cadmium reduction, Automated	353.2 Rev. 2.0	4500-NO3- F-2019	D3867–16 (A)	I-2545-90.51
		(1993).	or 4500–NO ₃ - I–		
			2019.		
	Automated hydrazine Reduction/Colorimetric		4500–NO ₃ – H–2019.		See footnote.62
	Ion Chromatography		4110 B–2020 or C–	D4327–17	993.30. ³
		(1993) and 300.1	2020.		000.00.
		Rev. 1.0 (1997).			
	CIE/UV		4140 B–2020	D6508–15	D6508, Rev. 2.54
	Enzymatic reduction, followed by auto-			D7781–14	I-2547-11, ⁷² I-
	mated colorimetric determination.				2548–11, ⁷² N07– 0003. ⁷³
	Enzymatic reduction, followed by manual		4500–NO ₃ – J–2018.		0000.
	colorimetric determination.				
	Spectrophotometric (2,6-dimethylphenol)				Hach 10206.75
40. Nitrite (as N), mg/L	Spectrophotometric: Manual		4500-NO ₂ - B-2021		See footnote.25
	Automated (Diazotization)				I-4540-85 ² see foot note, ⁶² I-2540-
					90. ⁸⁰
	Automated (*bypass cadmium reduction)	353.2 Rev. 2.0	4500–NO ₃ – F–2019,	D3867–16 (A)	I-4545-85.2
		(1993).	4500-NO ₃ - I-		
			2019.		
	Manual (*bypass cadmium or enzymatic re-		4500–NO ₃ – E–2019,	D3867–16 (B).	
	duction).		4500–NO ₃ [–] J– 2018.		
	Ion Chromatography	300.0 Rev. 2.1	4110 B-2020 or C-	D4327–17	993.30. ³
		(1993) and 300.1	2020.	5 1021 17 1111	
		Rev. 1.0 (1997).			
	CIE/UV		4140 B–2020	D6508–15	D6508, Rev. 2.54
	Automated (*bypass Enzymatic reduction)			D7781–14	I-2547-11, ⁷² I- 2548-11, ⁷² N07-
					0003. ⁷³
41. Oil and grease—	Hexane extractable material (HEM): n-	1664 Rev. A 1664	5520 B or G-2021.38.		
Total recoverable,	Hexane extraction and gravimetry.	Rev. B ⁴² .			
mg/L.		1004 D 1 1	5500 D 0		
	Silica gel treated HEM (SGT-HEM): Silica	1664 Rev. A, 1664	5520 B or G-2021 38		
	gel treatment and gravimetry.	Rev. B ⁴² .	and 5520 F- 2021. ³⁸		
42. Organic carbon—	Combustion		5310 B-2014	D7573–18a ^{e1}	973.47. ³ p. 14. ²⁴
					· · · · · · · · · · · · · · · · · · ·

Parameter	Methodology 58	EPA 52	Standard methods 84	ASTM	USGS/AOAC/Other
	Heated persulfate or UV persulfate oxida- tion.		5310 C–2014, 5310 D–2011.	D4839–03(17)	973.47, ^{3,} p. 14. ²⁴
 43. Organic nitrogen (as N), mg/L. 44. Ortho-phosphate (as P), mg/L. 	Total Kjeldahl N (Parameter 31) minus am- monia N (Parameter 4). Ascorbic acid method:				
(),	Automated	365.1 Rev. 2.0 (1993).	4500–P F–2021 or G–2021.		973.56, ³ I–4601–85, ² I–2601–90. ⁸⁰
	Manual, single-reagent Manual, two-reagent		4500–P E–2021	D515–88 (A)	973.55. ³
	Ion Chromatography	300.0 Rev. 2.1 (1993) and 300.1 Rev. 1.0 (1997).	4110 B–2020 or C– 2020.	D4327–17	
15 Opmium Total	CIE/UV Digestion, ⁴ followed by any of the fol-		4140 B-2020	D6508–15	D6508, Rev. 2.54
45. Osmium—Total⁴, mg/L.	lowing: AA direct aspiration		3111 D–2019.		
46. Oxygen, dissolved,	AA furnace	252.2 (Issued 1978). ¹	4500–O (B–F)–2021		973.45B, ³ I–1575–
mg/L.			4500-0 (B-F)-2021	D888–18 (A)	78. ⁸
	Electrode Luminescence-Based Sensor		4500–O G–2021 4500–O H–2021	D888–18 (B) D888–18 (C)	
47. Palladium—Total, ⁴ mg/L.	Digestion, ⁴ followed by any of the fol- lowing:			D000-10 (C)	See looliloles.
	AA direct aspiration AA furnace	253.2 (Issued 1978). ¹	3111 B–2019.		
	ICP/MS		3125 B–2020.		See footnote.34
48. Phenols, mg/L	Manual distillation, ²⁶ followed by any of the following:.	420.1 (Rev. 1978) ¹	5530 B–2021	D1783–01(12).	
	Colorimetric (4AAP) manual	420.1 (Rev. 1978) ¹	5530 D-2021 27	D1783–01(12) (A or B).	
10. Dhaashamus (ala	Automated colorimetric (4AAP)	420.4 Rev. 1.0 (1993).			0
49. Phosphorus (ele- mental), mg/L.	Gas-liquid chromatography				See footnote.28
50. Phosphorus—Total, mg/L.	Digestion, ²⁰ followed by any of the fol- lowing:.		4500–P B (5)–2021		973.55. ³
	Manual Automated ascorbic acid reduction	365.3 (Issued 1978) ¹ 365.1 Rev. 2.0 (1993).	4500–P E–2021 4500–P (F–H)–2021	D515–88 (A).	973.56, ³ I–4600–85. ²
	ICP/AES ⁴³⁶ Semi-automated block digestor (TKP di- gestion).	200.7Rev. 4.4 (1994) 365.4 (Issued 1974) ¹	3120 B–2020	 D515–88 (B)	I–4471–97. ⁵⁰ I–4610–91. ⁴⁸
	Digestion with persulfate, followed by Col- orimetric.				NCASI TNTP W10900.77
51. Platinum—Total, ⁴ mg/L.	Digestion, ⁴ followed by any of the fol- lowing:				
	AA direct aspiration AA furnace		3111 B–2019.		
	ICP/MS		3125 B-2020.		
50 Determinen Tetel 4					See footnote.34
52. Potassium—Total, ⁴ mg/L.	Digestion, ⁴ followed by any of the fol- lowing:				
Ū.	AA direct aspiration		3111 B-2019		973.5, ³ I-3630-85. ²
	ICP/AES	200.7 Rev. 4.4 (1994).	3120 B–2020.		
	ICP/MS	200.8, Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14. ³
	Flame photometric		3500–K B–2020. 3500–K C–2020.		
53. Residue—Total, mg/L.	Ion Chromatography Gravimetric, 103–105°		2540 B–2020	D6919–17.	I-3750-85. ²
54. Residue—filterable, mg/L.	Gravimetric, 180°		2540 C-2020	D5907–18 (B)	I–1750–85. ²
55. Residue—non-filter- able (TSS), mg/L.	Gravimetric, 103–105° post-washing of res- idue.		2540 D-2020	D5907–18 (A)	I-3765-85. ²
56. Residue—settle- able, mg/L.	Volumetric (Imhoff cone), or gravimetric		2540 F-2020.		
 57. Residue—Volatile, mg/L. 58. Rhodium—Total,⁴ 	Gravimetric, 550° Digestion, ⁴ followed by any of the fol-	160.4 (Issued 1971) ¹	2540 E-2020		I–3753–85. ²
mg/L.	lowing: AA direct aspiration, or		3111 B–2019.		
	AA furnace	265.2 (Issued 1978).1			
59. Ruthenium—Total,4	ICP/MS Digestion, ⁴ followed by any of the fol- lowing:		3125 B–2020.		
mg/L.					

n	-	n	1	n
2	1	5	т	5

60. Selenium—Total, ⁴ D mg/L. A 61. Silica—Dissolved, ³⁷ 0 mg/L. C	Methodology ⁵⁸ CP/MS Digestion, ⁴ followed by any of the fol- lowing: AA furnace STGFAA CP/AES ³⁶ CP/MS AA gaseous hydride	EPA ⁵² 200.9 Rev. 2.2 (1994). 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	Standard methods 84 3125 B-2020. 3113 B-2020	ASTM D3859–15 (B)	USGS/AOAC/Other I–4668–98.49
60. Selenium—Total, ⁴ D mg/L. A 61. Silica—Dissolved, ³⁷ 0 mg/L. C	Digestion, ⁴ followed by any of the fol- lowing: AA furnace STGFAA CP/AES ³⁶ CP/MS	200.9 Rev. 2.2 (1994). 200.5 Rev 4.2 (2003). ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	3113 B-2020		I–4668–98. ⁴⁹
A S IC A S 61. Silica—Dissolved, ³⁷ mg/L. C	AA furnace STGFAA CP/AES ³⁶ CP/MS AA gaseous hydride	200.9 Rev. 2.2 (1994). 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).			I–4668–98. ⁴⁹
61. Silica—Dissolved, ³⁷ 0 mg/L. C	CP/AES ³⁶ CP/MS	(1994). 200.5 Rev 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	3120 B-2020	D4070 00	
61. Silica—Dissolved, ³⁷ 0 mg/L. C	CP/MS	(2003), ⁶⁸ 200.7 Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	3120 B–2020	D1070 00	
61. Silica—Dissolved, ³⁷ 0 mg/L. C	AA gaseous hydride	200.8 Rev. 5.4 (1994).		D1976–20.	
61. Silica—Dissolved, ³⁷ 0 mg/L.			3125 B–2020	D5673–16	993.14, ³ I–4020– 05 ⁷⁰ I–4472–97. ⁸¹
mg/L.	45-micron filtration followed by any of the		3114 B–2020, or 3114 C–2020.	D3859–15 (A)	I-3667-85. ²
	following:				
	Colorimetric, Manual		4500–SiO ₂ C–2021 4500–SiO ₂ E–2021	D859–16	I–1700–85. ² I–2700–85. ²
10	CP/AES	200.5 Rev. 4.2	or F–2021. 3120 B–2020		I-4471-97. ⁵⁰
		(2003), ⁶⁸ 200.7 Rev. 4.4 (1994).			
	CP/MS	200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14. ³
mg/L.	Digestion, ^{4 29} followed by any of the fol- lowing:				
A	AA direct aspiration		3111 B–2019 or 3111 C–2019.		974.27, ³ p. 37, ⁹ l– 3720–85. ²
	AA furnace		3113 B-2020		I-4724-89. ⁵¹
	STGFAA	200.9 Rev. 2.2 (1994).			
10	CP/AES	200.5 Rev. 4.2 (2003), ⁶⁸ 200.7	3120 B-2020	D1976–20	I–4471–97. ⁵⁰
10	CP/MS	Rev. 4.4 (1994). 200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4472– 97. ⁸¹
	DCP				See footnote.34
63. Sodium—Total, ⁴ D mg/L.	Digestion, ⁴ followed by any of the fol- lowing:				
A	AA direct aspiration		3111 B-2019		973.54, ³ I–3735–85. ²
	CP/AES	200.5 Rev. 4.2 (2003), ⁶⁸ 200.7 Rev. 4.4 (1994).	3120 B-2020		I–4471–97. ⁵⁰
10	CP/MS	200.8 Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14. ³
	DCP Flame photometric				See footnote.34
lo	on Chromatography			D6919–17.	
64. Specific conduct- ance, micromhos/cm at 25 °C.	Wheatstone bridge	120.1 (Rev. 1982) ¹	2510 B-2021	D1125–95(99) (A)	973.40, ³ I–2781–85. ²
	Automated colorimetric	375.2 Rev. 2.0 (1993).	4500–SO₄ ^{2.} F–2021 or G–2021.		
	Gravimetric		4500–SO ₄ ² C–2021 or D–2021.		925.54. ³
	Furbidimetric		4500–SO4 ^{2.} E–2021	D516–16.	000 00 3 1 4000
	on Chromatography	300.0 Rev. 2.1 (1993) and 300.1 Rev. 1.0 (1997).	4110 B–2020 or C– 2020.	D4327–17	993.30, ³ I–4020– 05. ⁷⁰
-	CIE/UV Sample Pretreatment		4140 B–2020 4500–S ^{2.} B, C– 2021.	D6508–15	D6508 Rev. 2.54
	Fitrimetric (iodine)		4500–S ^{2.} F–2021		I-3840-85. ²
	Colorimetric (methylene blue)on Selective Electrode		4500–S ^{2.} D–2021. 4500–S ^{2.} G–2021	D4658–15.	
	Fitrimetric (iodine-iodate)		4500–SO ₃ ² · B– 2021.	54000 10.	
68. Šurfactants, mg/L C	Colorimetric (methylene blue)		5540 C-2021	D2330–20.	
70. Thallium-Total,4 D	Thermometric Digestion, ⁴ followed by any of the fol-		2550 B–2010		See footnote.32
mg/L.	lowing: AA direct aspiration		3111 B-2019.		
	AA furnace	279.2 (Issued 1978) ¹ 200.9 Rev. 2.2	3113 B–2020.		
10	CP/AES	(1994). 200.7 Rev. 4.4 (1994).	3120 B–2020	D1976–20.	
10	CP/MS	200.8, Rev. 5.4 (1994).	3125 B–2020	D5673–16	993.14, ³ I–4471– 97 ⁵⁰ I–4472–97. ⁸¹
	Digestion, ⁴ followed by any of the fol- lowing: AA direct aspiration		2111 B 2010		

Parameter	Methodology 58	EPA 52	Standard methods 84	ASTM	USGS/AOAC/Other
	AA furnace		3113 B-2020.		
	STGFAA	200.9 Rev. 2.2			
	100/450	(1994).			
	ICP/AES	200.5 Rev. 4.2			
		(2003), ⁶⁸ 200.7 Rev. 4.4 (1994).			
	ICP/MS	200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14. ³
		(1994).	5125 D-2020	D3073-10	333.14.
72. Titanium—Total, ⁴ mg/L.	Digestion, ⁴ followed by any of the fol- lowing:				
	AA direct aspiration		3111 D-2019.		
	AA furnace	283.2 (Issued 1978).1			
	ICP/AES	200.7 Rev. 4.4			
		(1994).			
	ICP/MS	200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14. ³
	DOD	(1994).			0
73. Turbidity, NTU 53	DCP Nephelometric		2130 B–2020	D1889–00	See footnote. ³⁴ I–3860–85, ² see foot-
73. Turbidity, NTO		(1993).	2130 D=2020	D1009-00	notes. ⁶⁵ 66 67
74. Vanadium—Total,4	Digestion, ⁴ followed by any of the fol-	(1000).			110100.
mg/L.	lowing:				
C C	AA direct aspiration		3111 D-2019.		
	AA furnace		3113 B-2020		
	ICP/AES	200.5 Rev. 4.2	3120 B-2020	D1976–20	I-4471-97. ⁵⁰
		(2003),68 200.7			
	ICP/MS	Rev. 4.4 (1994). 200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14. ³ I–4020–
	ICP/MIS	(1994).	3125 6-2020	D5073-10	993.14,° 1–4020– 05. ⁷⁰
	DCP			D4190–15	See footnote.34
	Colorimetric (Gallic Acid)		3500-V B-2011.		
75. Zinc—Total,4 mg/L	Digestion, ⁴ followed by any of the fol-				
	lowing:				
	AA direct aspiration ³⁶		3111 B–2019 or 3111 C–2019.	D1691–17 (A or B)	974.27 ³ p. 37, ⁹ l– 3900–85. ²
	AA furnace	289.2 (Issued 1978).1			
	ICP/AES ³⁶	200.5 Rev. 4.2	3120 B-2020	D1976–20	I-4471-97.50
		(2003),68 200.7,			
	100/110	Rev. 4.4 (1994).	0405 B 0000	D5070 40	000 4 4 3 4 4000
	ICP/MS	200.8 Rev. 5.4	3125 B-2020	D5673–16	993.14, ³ I–4020–
	DCP ³⁶	(1994).		D4190–15	05, ⁷⁰ I–4472–97. ⁸¹ See footnote. ³⁴
	Colorimetric (Zincon)		3500 Zn B–2020	D4190-15	See footnote.33
76. Acid Mine Drainage		1627. ⁶⁹	0000 211 0-2020		
. e iola mino Brainage					

Table IB Notes:

Table IB Notes: 1 Methods for Chemical Analysis of Water and Wastes, EPA-600/4–79–020. Revised March 1983 and 1979, where applicable. U.S. EPA.
² Methods for Chemical Analysis of Inorganic Substances in Water and Fluvial Sediments, Techniques of Water-Resource Investigations of the U.S. Geological Survey, Book 5, Chapter A1, unless otherwise stated. 1989. USGS.
³ Official Methods of Analysis of the Association of Official Analytical Chemists, Methods Manual, Sixteenth Edition, 4th Revision, 1998. AOAC International.
⁴ For the determination of total metals (which are equivalent to total recoverable metals) the sample is not filtered before processing. A digestion procedure is required to solubilize analytes in suspended material and to break down organic-metal complexes (to convert the analyte to a detectable form for colorimetric analysis) for analysis. The procedure used should subject the sample to gentle acid refluxing, and at no time should the sample be taken to dryness. For direct aspiration ffame atomic absorption (FLAA) determinations, a combination acid (nitric and hydrochloric acids) digestion is preferred, prior to analysis. The procedure used should 200.2 in Supplement 1 of "Nethods for the Determination of Metals in Environmental Samples" EPA/600R-94/111. May 1994, and is reproduced in EPA Methods 200.7, 200.8, and 200.9 from the same Supplement. However, when using the gaseous hydride technique or for the determination of certain elements such as antimony, arsenic, selenium, silver, and tin by non-EPA graphite furnace atomic absorption, the noble metals and titanium by FLAA, a specific or modified sample digestion procedure may be required, and, in all sectometry by cold vapor atomic absorption, the noble metals and titanium by FLAA.
Scopper sulfate or other catalysis that have been found sultable may be used for those respective methods. Regardless of the digestion procedure an EPA Methods 200.7, 200.8, and 200.9 may be used for those respective methods. R

nologies Inc. ⁸ The approved method is that cited in Methods for Determination of Inorganic Substances in Water and Fluvial Sediments, Techniques of Water-Resources Inves-

tigations of the U.S. Geological Survey, Book 5, Chapter A1. 1979. USGS. ⁹American National Standard on Photographic Processing Effluents. April 2, 1975. American National Standards Institute.

¹⁰ In-Situ Method 1003–8–2009, Biochemical Oxygen Demand (BOD) Measurement by Optical Probe. 2009. In-Situ Incorporated.

¹⁰ In-Situ Method 1003–8–2009, Biochemical Oxygen Demand (BOD) Measurement by Optical Probe. 2009. In-Situ Incorporated.
 ¹¹ The use of normal and differential pulse voltage ramps to increase sensitivity and resolution is acceptable.
 ¹² Carbonaceous biochemical oxygen demand (CBOD₅) must not be confused with the traditional BOD₅ test method which measures "total 5-day BOD." The addition of the nitrification inhibitor is not a procedural option but must be included to report the CBOD₅ parameter. A discharger whose permit requires reporting the traditional BOD₅ may not use a nitrification inhibitor.
 ¹³ OIC Chemical Oxygen Demand Method. 1978. Oceanography International Corporation.
 ¹⁴ Method 8000, Chemical Oxygen Demand, Hach Handbook of Water Analysis, 1979. Hach Company.
 ¹⁵ The back-titration method will be used to resolve controversy.
 ¹⁶ Orion Research Instruction Manual, Residual Chlorine Electrode Model 97–70. 1977. Orion Research Incorporated. The calibration graph for the Orion residual chlorine method must be derived using a reagent blank and three standard solutions, containing 0.2, 1.0, and 5.0 mL 0.00281 N potassium iodate/100 mL solution, respectively.

respectively.

¹⁷ Method 245.7, Mercury in Water by Cold Vapor Atomic Fluorescence Spectrometry, EPA–821–R–05–001. Revision 2.0, February 2005. US EPA.
 ¹⁸ National Council of the Paper Industry for Air and Stream Improvement (NCASI) Technical Bulletin 253 (1971) and Technical Bulletin 803, May 2000.
 ¹⁹ Method 8506, Bicinchoninate Method for Copper, Hach Handbook of Water Analysis. 1979. Hach Company.
 ²⁰ When using a method with block digestion, this treatment is not required.

²¹ Industrial Method Number 378–75WA, Hydrogen ion (pH) Automated Electrode Method, Bran & Luebbe (Technicon) Autoanalyzer II. October 1976. Bran &

²² Method 8008, 1,10-Phenanthroline Method using FerroVer Iron Reagent for Water. 1980. Hach Company.
 ²³ Method 8034, Periodate Oxidation Method for Manganese, Hach Handbook of Wastewater Analysis. 1979. Hach Company.
 ²⁴ Methods for Analysis of Organic Substances in Water and Fluvial Sediments, Techniques of Water-Resources Investigations of the U.S. Geological Survey, Book
 5, Chapter A3, (1972 Revised 1987). 1987. USGS.
 ²⁵ Method 8057, Nitrogen, Nitrite-Low Range, Diazotization Method for Water and Wastewater. 1979. Hach Company.
 ²⁶ Inst Prior to distillation adjust the suffurio-adjudrated approxed sample to pt 4 with 1 + 9 No.H

²⁶ Just prior to distillation, adjust the sulfuric-acid-preserved sample to pH 4 with 1 + 9 NaOH. ²⁷ The colorimetric reaction must be conducted at a pH of 10.0 ± 0.2.

28 Addison, R.F., and R.G. Ackman. 1970. Direct Determination of Elemental Phosphorus by Gas-Liquid Chromatography, Journal of Chromatography, 47(3):421-426

²⁹ Approved methods for the analysis of silver in industrial wastewaters at concentrations of 1 mg/L and above are inadequate where silver exists as an inorganic halide. Silver halides such as the bromide and chloride are relatively insoluble in reagents such as nitric acid but are readily soluble in an august buffer of solutions of the buffer of the buffer of solutions of the buffer of the

³⁰ The use of EDTA decreases method sensitivity. Analysis may omit EDTA or replace with another suitable complexing reagent provided that all method-specified quality control acceptance criteria are met. ³¹ For samples known or suspected to contain high levels of silver (*e.g.*, in excess of 4 mg/L), cyanogen iodide should be used to keep the silver in solution for analysis. Prepare a cyanogen iodide solution by adding 4.0 mL of concentrated NH₄OH, 6.5 g of KCN, and 5.0 mL of a 1.0 N solution of I₂ to 50 mL of reagent water in a volumetric flask and dilute to 100.0 mL. After digestion of the sample, adjust the pH of the digestate to <7 to prevent the formation of HCN under acidic conditions. Add 1 mL of the cyanogen iodide solution to the sample digestate and adjust the volume to 100 mL with reagent water (NOT acid). If cyanogen iodide is added to sample digestates, then silver standards must be prepared that contain cyanogen iodide as well. Prepare working standards by diluting a small volume of a silver stock solution with water and adjusting the pH \leq 7 with NH $_4$ OH. Add 1 mL of the cyanogen iodide solution and let stand 1 hour. Transfer to a 100-mL volumetric flask and dilute to volume with water. 32 "Water Temperature-Influential Factors, Field Measurement and Data Presentation," Techniques of Water-Resources Investigations of the U.S. Geological Sur-

vey, Book 1, Chapter D1. 1975. USGS.

 Method 8009, Zincon Method for Zinc, Hach Handbook of Water Analysis, 1979. Hach Company.
 ³⁴ Method AES0029, Direct Current Plasma (DCP) Optical Emission Spectrometric Method for Trace Elemental Analysis of Water and Wastes. 1986—Revised 1991. Thermo Jarrell Ash Corporation.

³⁶ In-Situ Method 1004–8-2009, Carbonaceous Biochemical Oxygen Demand (CBOD) Measurement by Optical Probe. 2009. In-Situ Incorporated. ³⁶ Microwave-assisted digestion may be employed for this metal, when analyzed by this methodology. Closed Vessel Microwave Digestion of Wastewater Samples

³⁷When determining boron and silica, only plastic, PTFE, or quartz laboratory ware may be used from start until completion of analysis. ³⁸Only use *n*-hexane (*n*-Hexane—85% minimum purity, 99.0% min. saturated C6 isomers, residue less than 1 mg/L) extraction solvent when determining Oil and Grease parameters—Hexane Extractable Material (HEM), or Silica Gel Treated HEM (analogous to EPA Methods 1664 Rev. A and 1664 Rev. B). Use of other ex-traction provide the provide the second solvent is the second solvent in the second solvent is the se traction solvents is prohibited.

³⁹ Method PAI–DK01, Nitrogen, Total Kjeldahl, Block Digestion, Steam Distillation, Titrimetric Detection. Revised December 22, 1994. OI Analytical.
 ⁴⁰ Method PAI–DK02, Nitrogen, Total Kjeldahl, Block Digestion, Steam Distillation, Colorimetric Detection. Revised December 22, 1994. OI Analytical.
 ⁴¹ Method PAI–DK03, Nitrogen, Total Kjeldahl, Block Digestion, Automated FIA Gas Diffusion. Revised December 22, 1994. OI Analytical.
 ⁴² Method PAI–DK03, Nitrogen, Total Kjeldahl, Block Digestion, Automated FIA Gas Diffusion. Revised December 22, 1994. OI Analytical.
 ⁴² Method 1664 Rev. B is the revised version of EPA Method 1664 Rev. A. U.S. EPA. February 1999, Revision A. Method 1664, *n*-Hexane Extractable Material (HEM; Non-polar Material) by Extraction and Gravimetry. EPA-821–R–98–002.
 U.S. EPA. February 2010, Revision B. Method 1664, *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (HEM;

U.S. EPA. February 2010, Revision B. Method 1664, *n*-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (SGT-HEM; Non-polar Material) by Extraction and Gravimetry. EPA-821-R-10-001.
 ⁴³Method 1631, Revision E, Mercury in Water by Oxidation, Purge and Trap, and Cold Vapor Atomic Fluorescence Spectrometry, EPA-821-R-02-019. Revision E. August 2002, U.S. EPA. The application of clean techniques described in EPA's Method 1669: *Sampling Ambient Water for Trace Metals at EPA Water Quality Criteria Levels*, EPA-821-R-96-011, are recommended to preclude contamination at low-level, trace metal determinations.
 ⁴⁴Method OIA-1677-09, Available Cyanide by Ligand Exchange and Flow Injection Analysis (FIA). 2010. OI Analytical.
 ⁴⁵Open File Report 00-170, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Ammonium Plus Organic Nitrogen by a Kjeldahl Digestion Method and an Automated Photometric Finish that Includes Digest Cleanup by Gas Diffusion. 2000. USGS.
 ⁴⁷Open File Report 93-449, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Chromium in Water by Graphite Furnace Atomic Absorption Spectrophotometry. 1993. USGS.
 ⁴⁷Open File Report 97-188, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Molybdenum by Graphite Furnace Atomic Absorption Spectrophotometry. 1997. USGS.
 ⁴⁸Open File Report 97-186, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Molybdenum by Graphite Furnace Atomic Absorption Spectrophotometry. 1997. USGS.
 ⁴⁹Open File Report 92-146, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Total Phosphorus by Kieldahl

Furnace Atomic Absorption Spectrophotometry. 1997. USGS. ⁴⁹ Open File Report 92–146, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Total Phosphorus by Kjeldahl Digestion Method and an Automated Colorimetric Finish That Includes Dialysis. 1992. USGS. ⁴⁹ Open File Report 98–639, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Arsenic and Selenium in Water and Sediment by Graphite Furnace-Atomic Absorption Spectrometry. 1999. USGS. ⁵⁰ Open File Report 98–165, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Elements in Whole-water Di-gests Using Inductively Coupled Plasma-Optical Emission Spectrometry and Inductively Coupled Plasma-Mass Spectrometry. 1998. USGS. ⁵¹ Open File Report 93–125, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Inorganic and Organic Con-stituents in Water and Fluvial Sediments. 1993. USGS. ⁵² Unless otherwise indicated, all EPA methods excluding EPA Method 300.1 are published in U.S. EPA May 1994. Methods for the Determination of Methols in ⁵² Methods and LEPA methods excluding EPA Method 300.1 are published in U.S. EPA May 1994. Methods for the Determination of Methols in ⁵⁴ Methods and Herbories and Fluvial Sediments. 1993. USGS.

⁵² Unless otherwise indicated, all EPA methods, excluding EPA Method 300.1, are published in U.S. EPA. May 1994. Methods for the Determination of Metals in Environmental Samples, Supplement I, EPA/600/R–94/111; or U.S. EPA. August 1993. Methods for the Determination of Inorganic Substances in Environmental Samples, EPA/600/R–93/100. EPA Method 300.1 is U.S. EPA. Revision 1.0, 1997, including errata cover sheet April 27, 1999. Determination of Inorganic Ions in Drinking Water by Ion Chromatography. ⁵³ Styrene divinyl benzene beads (*e.g.*, AMCO-AEPA-1 or equivalent) and stabilized formazin (*e.g.*, Hach StablCalTM or equivalent) are acceptable substitutes for

formazin.

³⁴Waters Corp. Now included in ASTM D6508-15, Test Method for Determination of Dissolved Inorganic Anions in Aqueous Matrices Using Capillary Ion Electro-

⁵⁵Kelada-01, Kelada Automated Test Methods for Total Cyanide, Acid Dissociable Cyanide, and Thiocyanate, EPA 821–B–01–009, Revision 1.2, August 2001. US EPA. Note: A 450–W UV lamp may be used in this method instead of the 550–W lamp specified if it provides performance within the quality control (QC) acceptance criteria of the method in a given instrument. Similarly, modified flow cell configurations and flow conditions may be used in the method, provided that the QC accept

⁵⁶ QuikChem Method In/a given instrument.
 ⁵⁶ QuikChem Method 10–204–00–1–X, Digestion and Distillation of Total Cyanide in Drinking and Wastewaters using MICRO DIST and Determination of Cyanide by Flow Injection Analysis. Revision 2.2, March 2005. Lachat Instruments.
 ⁵⁷ When using sulfide removal test procedures described in EPA Method 335.4, reconstitute particulate that is filtered with the sample prior to distillation.
 ⁵⁸ Unless otherwise stated, if the language of this table specifies a sample digestion and/or distillation "followed by" analysis with a method, approved digestion

⁵⁹ Unley during stilled reinovariest proceedings described in EPA Metricol 353.4, reconstitute particulate marks intered with the sample proved digestion and/or distillation are required prior to analysis.
 ⁵⁹ Samples analyzed for available cyanide using OI Analytical method OIA–1677–09 or ASTM method D6888–16 that contain particulate matter may be filtered only after the ligand exchange reagents have been added to the samples, because the ligand exchange process converts complexes containing available cyanide to free cyanide, which is not removed by filtration. Analysts are further cautioned to limit the time between the addition of the ligand exchange reagents and sample filtration to no more than 30 minutes to preclude settling of materials in samples.
 ⁶⁰ Analysts should be aware that pH optima and chromophore absorption maxima might differ when phenol is replaced by a substituted phenol as the color reagent, pH optimum and wavelength of maximum absorbance are about 11.5 and 635 nm, respectively—see, Patton, C.J. and S.R. Crouch. March 1977. *Anal. Chem.* 49:464–469.
 ⁶¹ If atomic absorption or ICP instrumentation is not available, the aluminon colorimetric method detailed in the 19th Edition of *Standard Methods for the Examination of Water and Wastewater* may be used. This method has poorer precision and bias than the methods of choice.
 ⁶² Easy (1-Reagent) Nitrate Method, Revision November 12, 2011. Craig Chinchilla.
 ⁶³ Hach Method 10360, Luminescence Measurement of Dissolved Oxygen in Water and Wastewater and for Use in the Determination of BOD₅ and CBOD₅. Revision 1.2, October 2011. Hach Company. This method may be used to measure dissolved oxygen when performing the methods approved in this table IB for measurement of Discolved Oxygen (DO) Measurement by Optical Probe. 2009. In-Situ Incorporated.
 ⁶⁴ In-Situ Method 1002–8–2009, Dissolved Oxygen (DO) Measurement by Optical Probe. 2009. In-Situ Incorp

⁶⁶ Mitchell Method M5271, Determination of Turbidity by Nephelometry. Revision 1.0, July 31, 2008. Leck Mitchell.
 ⁶⁷ Orion Method AQ4500, Determination of Turbidity by Nephelometry. Revision 5, March 12, 2009. Thermo Scientific.
 ⁶⁸ EPA Method 200.5, Determination of Trace Elements in Drinking Water by Axially Viewed Inductively Coupled Plasma-Atomic Emission Spectrometry, EPA/600/

⁶⁶ EPA Method 200.5, Determination of Trace Elements in Drinking Water by Axially Viewed Inductively Coupled Plasma-Atomic Emission Spectrometry, EPA/600/ R-06/115. Revision 4.2, October 2003. US EPA.
 ⁶⁹ Method 1627, Kinetic Test Method for the Prediction of Mine Drainage Quality, EPA-821–R-09-002. December 2011. US EPA.
 ⁷⁰ Techniques and Methods Book 5–B1, Determination of Elements in Natural-Water, Biota, Sediment and Soil Samples Using Collision/Reaction Cell Inductively
 ⁷¹ Water-Resources Investigations Report 01–4132, Methods of the National Water Quality Laboratory, Book 5, Laboratory Analysis, 2006. USGS.
 ⁷² USGS Techniques and Methods 5–B8, Chapter 8, Section B, Methods of the National Water Quality Laboratory Book 5, Laboratory Analysis, 2011 USGS.
 ⁷³ NECE Method 1007–0003, "Nitrate Reductase Nitrate-Nitrogen Analysis," Revision 9.0, March 2014, The Nitrate Elimination of. Ic., Inc.
 ⁷⁴ Timberline Instruments, LLC Method Ammonia-001, "Determination of Inorganic Ammonia by Continuous Flow Gas Diffusion and Conductivity Cell Analysis," June 2011, Timberline Instruments, LLC.

⁷⁵ Hach Company Method 10206, "Spectrophotometric Measurement of Nitrate in Water and Wastewater," Revision 2.1, January 2013, Hach Company.
⁷⁶ Hach Company Method 10242, "Simplified Spectrophotometric Measurement of Total Kjeldahl Nitrogen in Water and Wastewater," Revision 1.1, January 2013,

⁷⁸ The PH adjusted sample is to be adjusted to 7.6 for NPDES reporting purposes.
 ⁷⁸ The PH adjusted Sample Super S

 ⁷⁸ The pH adjusted sample is to be adjusted to 7.6 for NPDES reporting purposes.
 ⁷⁹ I-2057-85 in U.S. Geological Survey Techniques of Water-Resources Investigations, Book 5, Chap. A1, Methods for Determination of Inorganic Substances in Water and Fluvial Sediments, 1989.
 ⁸⁰ Methods I-2522-90, I-2540-90, and I-2601-90 in U.S. Geological Survey Open-File Report 93-125, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Inorganic and Organic Constituents in Water and Fluvial Sediments, 1993.
 ⁸¹ Method I-4472-97 in U.S. Geological Survey Open-File Report 98-165, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Inorganic Constituents in Water and Fluvial Sediments, 1993.
 ⁸² FIAlab 100, "Determination of Inorganic Constituents in Water and Fluvial Sediments, 1998.
 ⁸² FIAlab 100, "Determination of Inorganic Ammonia by Continuous Flow Gas Diffusion and Fluorescence Detector Analysis", April 4, 2018, FIAlab Instruments, Inc.
 ⁸⁴ Please refer to the following applicable Quality Control Sections: Part 2000 Methods, Physical and Aggregate Properties 2020 (2021); Part 3000 Methods, Metals, 3020 (2021); Part 4000 Methods, Inorganic Nonmetallic Constituents, 4020 (2022); Part 5000 Methods, and Aggregate Organic Constituents, 5020 (2022). These Quality Control Standards are available for download at *www.standardmethods.org* at no charge.
 ⁸⁵ Each laboratory may establish its own control limits by performing at least 25 glucose-glutamic acid (GGA) checks over several weeks or months and calculating the mean and standard deviation. The laboratory may then use the mean ± 3 standard deviations as the control limit for future GGA checks. However, GGA acceptance criteria can be no wider than 198 ± 30.5 mg/L for BDD₅. GGA acceptance criteria for CBOD must be either 198 ± 30.5 mg/L, or the lab may develop control ch than 250 mg/L

⁸⁶ The approved method is that cited in Standard Methods for the Examination of Water and Wastewater, 14th Edition, 1976.

TABLE IC—LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS

Parameter ¹	Method	EPA ²⁷	Standard methods ¹⁷	ASTM	Other
1. Acenaphthene	GC	610.			
· · · · · · · · · · · · · · · · · · ·	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2021	D4657–92 (98).	
2. Acenaphthylene	GC	610.	0110 2 2021 11111	2.007 02 (00).	
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2021	D4657-92 (98).	
3. Acrolein	GC	603.	0110 2 2021 11111	2.007 02 (00).	
	GC/MS	624.1 ⁴ , 1624B.			
4. Acrylonitrile	GC	603.			
	GC/MS	624.1 ⁴ , 1624B			O-4127-96. ¹³
5. Anthracene	GC	610.			0 1127 00.
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2021	D4657–92 (98).	
6. Benzene	GC	602	6200 C-2020.	D 1007 02 (00).	
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
7. Benzidine	Spectro-photo-		0200 0 2020		See footnote ³ p.1.
	metric.				
	GC/MS	625.1 ⁵ , 1625B	6410 B–2020.		
	HPLC	605.	0410 D 2020.		
8. Benzo(a)anthracene	GC	610.			
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2021	D4657–92 (98).	See loothole * p. 27.
9. Benzo(a)pyrene	GC	610.	0440 D-2021	D4037-92 (90).	
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2020	D4657–92 (98).	See loothole - p. 27.
10. Benzo(b)fluoranthene	GC	610.	0440 D-2021	D4037-92 (90).	
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	610	6440 B-2020	 D4657–92 (98).	See loothole * p. 27.
11. Benzo(g,h,i)perylene	GC	610.	0440 D-2021	D4057-92 (90).	
TT. Benzo(g,n,i)perviene	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	HPLC	/	6440 B-2020	 D4657–92 (98).	See loothole ° p. 27.
10 Denze (k) fluerenthene	GC	610 610.	6440 D-2021	D4057-92 (98).	
12. Benzo(k)fluoranthene	GC		6410 B 2020		Saa faataata 9 p. 07
		625.1, 1625B	6410 B-2020	D 4057 00 (00)	See footnote ⁹ p. 27.
10. Departul ablarida	HPLC	610	6440 B–2021	D4657–92 (98).	Cap fastasta 3 p. 100
13. Benzyl chloride	GC				See footnote ³ p. 130.
					See footnote ⁶ p. S102.
14. Butyl benzyl phthalate	GC	606.	0.440 D 0000		0 ()) 0 07
	GC/MS	625.1, 1625B	6410 B-2020		See footnote 9 p. 27.
15. bis(2-Chloroethoxy) methane	GC	611.			a ()) a a
	GC/MS	625.1, 1625B	6410 B-2020		See footnote 9 p. 27.
16. bis(2-Chloroethyl) ether	GC	611.			
	GC/MS	625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
17. bis(2-Ethylhexyl) phthalate	GC	606.	.		
	GC/MS	625.1, 1625B			See footnote ⁹ p. 27.
18. Bromodichloromethane	GC	601	6200 C–2020.		
	GC/MS	624.1, 1624B	6200 B-2020		0–4127–96 ¹³ , 0–4436–16. ¹⁴

TABLE IC-LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS-Continued

Parameter ¹	Method	EPA ²⁷	Standard methods 17	ASTM	Other
19. Bromoform	. GC GC/MS	601 624.1, 1624B	6200 C–2020. 6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
20. Bromomethane		601 624.1. 1624B	6200 C–2020. 6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
21. 4-Bromophenyl phenyl ether		611. 625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
22. Carbon tetrachloride	. GC	601	6200 C–2020		See footnote ³ p. 130.
23. 4-Chloro-3-methyl phenol		624.1, 1624B 604	6200 B–2020 6420 B–2021.		O-4127-96 ¹³ , O-4436-16. ¹⁴
24. Chlorobenzene		625.1, 1625B 601, 602	6410 B–2020 6200 C–2020		See footnote ⁹ p. 27. See footnote ³ p. 130.
25. Chloroethane	GC/MS GC	624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ O-4436-16. ¹⁴
26. 2-Chloroethylvinyl ether	GC/MS	624.1, 1624B 601.	6200 B–2020		O-4127-96. ¹³
27. Chloroform	GC/MS	624.1, 1624B.	6200 C–2020		See footnote ³ p. 130.
28. Chloromethane	GC/MS	624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ , O-4436-16. ¹⁴
29. 2-Chloronaphthalene	GC/MS	624.1, 1624B 612.	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
30. 2-Chlorophenol	GC/MS	625.1, 1625B	6410 B–2020 6420 B–2021.		See footnote ⁹ p. 27.
·	GC/MS	625.1, 1625B 611.	6410 B–2020		See footnote ⁹ p. 27.
31. 4-Chlorophenyl phenyl ether	GC/MS	625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
32. Chrysene	GC/MS	610. 625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
33. Dibenzo(a,h)anthracene		610 610.	6440 B–2021	D4657–92 (98).	
	GC/MS HPLC	625.1, 1625B 610	6410 B–2020 6440 B–2021	 D4657–92 (98).	See footnote ⁹ p. 27.
34. Dibromochloromethane	. GC GC/MS	601 624.1, 1624B	6200 C–2020. 6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
35. 1,2-Dichlorobenzene	. GC GC/MS	601, 602 624.1, 1625B	6200 C–2020. 6200 B–2020		See footnote ⁹ p. 27, O–4127–96 ¹³ ,
36. 1,3-Dichlorobenzene	. GC	601, 602	6200 C–2020.		O-4436-16. ¹⁴
37. 1,4-Dichlorobenzene	GC/MS	624.1, 1625B 601, 602	6200 B–2020 6200 C–2020.		See footnote ⁹ p. 27, O-4127-96. ¹³
	GC/MS	624.1, 1625B	6200 B–2020		See footnote ⁹ p. 27, O–4127–96 ¹³ , O–4436–16. ¹⁴
38. 3,3'-Dichlorobenzidine	. GC/MS	625.1, 1625B	6410 B–2020.		
39. Dichlorodifluoromethane		601.	6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
40. 1,1-Dichloroethane		601 624.1, 1624B	6200 C–2020. 6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
41. 1,2-Dichloroethane	. GC	601	6200 C–2020.		
42. 1,1-Dichloroethene		624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ , O-4436-16. ¹⁴
43. trans-1,2-Dichloroethene		624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ , O-4436-16. ¹⁴
44. 2,4-Dichlorophenol	GC/MS	624.1, 1624B 604	6200 B–2020 6420 B–2021.		O-4127-96 ¹³ , O-4436-16. ¹⁴
45. 1,2-Dichloropropane	GC/MS	625.1, 1625B 601	6410 B–2020 6200 C–2020.		See footnote ⁹ p. 27.
46. cis-1,3-Dichloropropene	GC/MS	624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ O-4436-16. ¹⁴
47. trans-1,3-Dichloropropene	GC/MS	624.1, 1624B 601	6200 B–2020 6200 C–2020.		O-4127-96 ¹³ , O-4436-16. ¹⁴
48. Diethyl phthalate	GC/MS	624.1, 1624B 606.	6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
49. 2,4-Dimethylphenol	GC/MS	625.1, 1625B 604	6410 B–2020 6420 B–2021.		See footnote 9 p. 27.
50. Dimethyl phthalate	GC/MS	625.1, 1625B 606.	6410 B–2020		See footnote ⁹ p. 27.
51. Di- <i>n</i> -butyl phthalate	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
	GC/MS	625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
52. Di- <i>n</i> -octyl phthalate	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
53. 2, 4-Dinitrophenol	GC/MS	604 625.1, 1625B	6420 B–2021 6410 B–2020.		See footnote ⁹ p. 27.
54. 2,4-Dinitrotoluene	GC/MS	609. 625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
55. 2,6-Dinitrotoluene	GC/MS	609. 625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
56. Epichlorohydrin	. GC GC/MS				See footnote ³ p. 130. See footnote ⁶ p. S102.

TABLE IC-LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS-Continued

Parameter ¹	Method	EPA ²⁷	Standard methods ¹⁷	ASTM	Other
57. Ethylbenzene	GC GC/MS	602 624.1, 1624B	6200 C–2020. 6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
58. Fluoranthene	GC GC/MS	610. 625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
9. Fluorene	HPLC GC	610 610.	6440 B–2021		
	GC/MS	625.1, 1625B 610	6410 B–2020 6440 B–2021		See footnote ⁹ p. 27.
 1,2,3,4,6,7,8-Heptachloro- dibenzofuran. 	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130- SSI. ¹⁶
1. 1,2,3,4,7,8,9-Heptachloro- dibenzofuran.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130- SSI. ¹⁶
2. 1,2,3,4,6,7,8- Heptachloro- dibenzo- <i>p</i> -dioxin.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130- SSI. ¹⁶
3. Hexachlorobenzene	GC GC/MS	612. 625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
4. Hexachlorobutadiene	GC GC/MS	612.	6410 B–2020		See footnote ⁹ p. 27, O-4127-96. ¹³
5. Hexachlorocyclopentadiene	GC GC/MS	612. 625.1 ⁵ , 1625B	6410 B–2020		See footnote ⁹ , p. 27, O–4127–96. ¹³
6. 1,2,3,4,7,8-Hexachloro- dibenzofuran.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130– SSI. ¹⁶
dibenzofuran.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130– SSI. ¹⁶
dibenzofuran. 8. 1,2,3,7,8,9-Hexachloro- dibenzofuran.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130– SSI. ¹⁶
dibenzofuran. 9. 2,3,4,6,7,8-Hexachloro- dibenzofuran.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130– SSI. ¹⁶
dioenzordian. 70. 1,2,3,4,7,8-Hexachloro-dibenzo- <i>p</i> - dioxin.	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130– SSI. ¹⁶
1. 1,2,3,6,7,8-Hexachloro-dibenzo-p-	GC/MS	1613B ¹⁰			SGS AXYS 16130 15, PAM 16130-
dioxin. 2. 1,2,3,7,8,9-Hexachloro-dibenzo- <i>p</i> -	GC/MS	1613B ¹⁰			SSI. ¹⁶ SGS AXYS 16130 ¹⁵ , PAM 16130–
dioxin. 3. Hexachloroethane	GC	612.	0440 0 0000		SSI. ¹⁶
4. Indeno(1,2,3-c,d) pyrene	GC/MS	625.1, 1625B 610.	6410 B-2020		See footnote ⁹ p. 27, O–4127–96. ¹³
	GC/MS	625.1, 1625B 610	6410 B–2020 6440 B–2021		See footnote ⁹ p. 27.
5. Isophorone	GC GC/MS	609. 625.1, 1625B	6410 B–2020		See footnote 9 p. 27.
6. Methylene chloride	GC GC/MS	601 624.1, 1624B	6200 C–2020 6200 B–2020		See footnote ³ p. 130. O-4127-96 ¹³ , O-4436-16. ¹⁴
7. 2-Methyl-4,6-dinitrophenol	GC GC/MS	604 625.1, 1625B	6420 B–2021. 6410 B–2020		See footnote ⁹ p. 27.
8. Naphthalene	GC GC/MS	610. 625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27.
9. Nitrobenzene	HPLC GC	610 609.	6440 B–2021.		
	GC/MS	625.1, 1625B			See footnote ⁹ p. 27.
0. 2-Nitrophenol	GC GC/MS	604 625.1, 1625B	6420 B–2021. 6410 B–2020		See footnote ⁹ p. 27.
1. 4-Nitrophenol	GC GC/MS	604 625.1, 1625B	6420 B–2021. 6410 B–2020		See footnote ⁹ p. 27.
2. N-Nitrosodimethylamine	GC GC/MS	607. 625.1 ⁵ , 1625B	6410 B–2020		See footnote ⁹ p. 27.
3. N-Nitrosodi-n-propylamine	GC GC/MS	607. 625.1 ⁵ , 1625B	6410 B–2020		See footnote ⁹ p. 27.
4. N-Nitrosodiphenylamine	GC GC/MS	607. 625.1 ⁵ , 1625B	6410 B–2020		See footnote ⁹ p. 27.
5. Octachlorodibenzofuran	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130- SSI. ¹⁶
6. Octachlorodibenzo-p-dioxin	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130- SSI. ¹⁶
7. 2,2'-oxybis(1-chloropropane) ¹² [also known as bis(2-Chloro-1- methylethyl) ether].	GC	611.			
8. PCB–1016	GC/MS GC	625.1, 1625B 608.3	6410 B–2020		See footnote ⁹ p. 27. See footnote ³ p. 43, see footnote. ⁸
9. PCB-1221	GC/MS GC	625.1 608.3	6410 B–2020.		See footnote ³ p. 43, see footnote. ⁸
0. PCB-1232	GC/MS	625.1 608.3	6410 B–2020.		See footnote ³ p. 43, see footnote. ⁸
1. PCB-1242	GC/MS	625.1 608.3	6410 B–2020.		See footnote ³ p. 43, see footnote. ⁸
2. PCB-1248	GC/MS	625.1 608.3	6410 B–2020.		See footnote ³ p. 43, see footnote. ⁸
2.1 00-1240	GC/MS	625.1	6410 B–2020.		

TABLE IC—LIST OF APPROVED TEST PROCEDURES FOR NON-PESTICIDE ORGANIC COMPOUNDS—Continued

Parameter ¹	Method	EPA ²⁷	Standard methods 17	ASTM	Other
	GC/MS	625.1	6410 B-2020.		
94. PCB-1260	GC	608.3			See footnote 3 p. 43, see footnote.8
	GC/MS	625.1	6410 B–2020.		
					AND AND ANALASIS BANK ANALAS
95. 1,2,3,7,8-Pentachloro-	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130-
dibenzofuran.					SSI. ¹⁶
6. 2,3,4,7,8-Pentachloro-	GC/MS	1613B ¹⁰			SGS AXYS 16130 15, PAM 16130-
dibenzofuran.					SSI. ¹⁶
07. 1,2,3,7,8-Pentachloro-dibenzo-p-	GC/MS	1613B ¹⁰			SGS AXYS 16130 15, PAM 16130-
	GC/NIS	10130 10			
dioxin.					SSI. ¹⁶
8. Pentachlorophenol	GC	604	6420 B-2021		See footnote ³ p. 140.
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
9. Phenanthrene	GC	610.			
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
					See 10011101e * p. 27.
	HPLC	610	6440 B–2021	D4657–92 (98).	
00. Phenol	GC	604	6420 B–2021.		
	GC/MS	625.1, 1625B	6410 B-2020		See footnote 9 p. 27.
01. Pyrene	GC	610.			
	GC/MS	625.1, 1625B	6410 B-2020		See footnote ⁹ p. 27.
					3ee 10011101e ° p. 27.
	HPLC	610	6440 B–2021	D4657–92 (98)	
02. 2,3,7,8-Tetrachloro-dibenzofuran	GC/MS	1613B ¹⁰			SGS AXYS 16130 ¹⁵ , PAM 16130-
					SSI. ¹⁶
03. 2,3,7,8-Tetrachloro-dibenzo-p-	GC/MS	613, 625.1 ⁵ ,			SGS AXYS 16130 15, PAM 16130-
dioxin.		1613B.			SSI. ¹⁶
	00		0000 0 0000		
04. 1,1,2,2-Tetrachloroethane	GC	601	6200 C–2020		See footnote ³ p. 130.
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96.13
05. Tetrachloroethene	GC	601	6200 C–2020		See footnote ³ p. 130.
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
06. Toluene	GC	602	6200 C-2020.		
					0 4107 0018 0 4400 4014
	GC/MS	624.1, 1624B	6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
107. 1,2,4-Trichlorobenzene	GC	612			See footnote ³ p. 130.
	GC/MS	625.1, 1625B	6410 B–2020		See footnote ⁹ p. 27, O–4127–96 ¹³ O–4436–16. ¹⁴
108. 1,1,1-Trichloroethane	GC	601	6200 C-2020.		
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
00 4 4 0 T : I I II		, ,			
09. 1,1,2-Trichloroethane	GC	601	6200 C–2020		See footnote ³ p. 130.
	GC/MS	624.1, 1624B	6200 B–2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
10. Trichloroethene	GC	601	6200 C-2020.		
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
11. Trichlorofluoromethane	GC	601	6200 C-2020.		
					0 4407 00 12
	GC/MS	624.1	6200 B–2020		O-4127-96. ¹³
12. 2,4,6-Trichlorophenol	GC	604	6420 B–2021.		
	GC/MS	625.1, 1625B	6410 B-2020		See footnote 9 p. 27.
13. Vinyl chloride	GC	601	6200 C-2020.		
	GC/MS	624.1, 1624B	6200 B-2020		O-4127-96 ¹³ , O-4436-16. ¹⁴
		, ,			0-4127-9010, 0-4430-10.11
14. Nonylphenol	GC/MS			D7065–17.	
15. Bisphenol A (BPA)	GC/MS			D7065–17.	
16. <i>p-tert</i> -Octylphenol (OP)	GC/MS			D7065–17.	
17. Nonylphenol Monoethoxylate	GC/MS			D7065-17.	
(NP1EO).					
 Nonylphenol Diethoxylate (NP2EO). 	GC/MS			D7065–17.	
19. Adsorbable Organic Halides	Adsorption and	1650.11			
(AOX).	Coulometric Ti- tration.				
120. Chlorinated Phenolics	In Situ Acetylation and GC/MS.	1653.11			

Table IC notes:

¹ All parameters are expressed in micrograms per liter (μg/L) except for Method 1613B, in which the parameters are expressed in picograms per liter (pg/L). ² The full text of Methods 601–613, 1613B, 1624B, and 1625B are provided at appendix A, Test Procedures for Analysis of Organic Pollutants. The standardized test procedure to be used to determine the method detection limit (MDL) for these test procedures is given at appendix B of this part, Definition and Procedure for the Determination of the Method Detection Limit. These methods are available at: *https://www.epa.gov/cwa-methods* as individual PDF files. ³ Methods for Benzidine: Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater. September 1978. U.S. EPA.

⁴Method 624.1 may be used for quantitative determination of acrolein and acrylonitrile, provided that the laboratory has documentation to substantiate the ability to detect and quantify these analytes at levels necessary to comply with any associated regulations. In addition, the use of sample introduction techniques other than simple purge-and-trap may be required. QC acceptance criteria from Method 603 should be used when analyzing samples for acrolein and acrylonitrile in the absence of such criteria in Method 624.1.

of such criteria in Method 624.1. ⁵Method 625.1 may be extended to include benzidine, hexachlorocyclopentadiene, N-nitrosodimethylamine, N-nitrosodi-*n*-propylamine, and N-nitrosodiphenylamine. However, when they are known to be present, Methods 605, 607, and 612, or Method 1625B, are preferred methods for these compounds. Method 625.1 may be ap-plied to 2,3,7,8-Tetrachloro-dibenzo-*p*-dioxin for screening purposes only. ⁶Selected Analytical Methods Approved and Cited by the United States Environmental Protection Agency, Supplement to the 15th Edition of *Standard Methods for the Examination of Water and Wastewater*. 1981. American Public Health Association (APHA). ⁷Each analyst must make an initial, one-time demonstration of their ability to generate acceptable precision and accuracy with Methods 601–603, 1624B, and 1625B in accordance with procedures in Section 8.2 of each of these methods. Additionally, each laboratory, on an on-going basis must spike and analyze 10% (5% for Methods 624.1 and 625.1 and 100% for methods 1624B and 1625B) of all samples to monitor and evaluate laboratory data quality in accordance with Sections 8.3 and 8.4 of these methods. When the recovery of any parameter falls outside the quality control (QC) acceptance criteria in the pertinent method, analytical results for that parameter in the unspiked sample are suspect. The results should be reported but cannot be used to demonstrate regulatory compliance. If the method does not contain QC acceptance criteria, control limits of ±three standard deviations around the mean of a minimum of five replicate measurements must be used. These ⁸Organochlorine Pesticides and PCBs in Wastewater Using EmporeTM Disk. Revised October 28, 1994. 3M Corporation. ⁹Method O–3116–87 is in Open File Report 93–125, Methods of Analysis by U.S. Geological Survey National Water Quality Laboratory—Determination of Inorganic and Organic Constituents in Water and Fluvial Sediments. 1993. USGS.

¹⁰ Analysts may use Fluid Management Systems, Inc. Power-Prep system in place of manual cleanup provided the analyst meets the requirements of Method 1613B (as specified in Section 9 of the method) and permitting authorities. Method 1613, Revision B, Tetra- through Octa-Chlorinated Dioxins and Furans by Isotope Dilution HRGC/HRMS. Revision B, 1994. U.S. EPA. The full text of this method is provided in appendix A to this part and at *https://www.epa.gov/cwa-methods/ap-*

¹¹Method 1650, Adsorbable Organic-compounds. ¹¹Method 1650, Adsorbable Organic Halides by Adsorption and Coulometric Titration. Revision C, 1997 U.S. EPA. Method 1653, Chlorinated Phenolics in Waste-water by In Situ Acetylation and GCMS. Revision A, 1997 U.S. EPA. The full text for both of these methods is provided at appendix A in part 430 of this chapter, The

Method 1900, Provided to Span and GCMS. Revision A, 1997 U.S. EPA. The full text for both of these methods is provided at appendix A in part 430 of this chapter, The Pulp, Paper, and Paperboard Point Source Category.
 ¹² The compound was formerly inaccurately labeled as 2,2'-oxybis(2-chloropropane) and bis(2-chloroisopropyl) ether. Some versions of Methods 611, and 1625 in-accurately list the analyte as "bis(2-chloroisopropyl) ether." but use the correct CAS number of 108–60–1.
 ¹³ Method O–4127–96, U.S. Geological Survey Open-File Report 97–829, Methods of analysis by the U.S. Geological Survey National Water Quality Laboratory— Determination of 86 volatile organic compounds in water by gas chromatography/mass spectrometry, including detections less than reporting limits, 1998, USGS.
 ¹⁴ Method O–4436–16 U.S. Geological Survey Techniques and Methods, book 5, chap. B12, Determination of heat purgeable and ambient purgeable volatile organic compounds in water by gas chromatography/mass spectrometry, 2016, USGS.
 ¹⁵ SGS AXYS Method 16130, "Determination of 2,3,7,8-Substituted Tetra- through Octa-Chlorinated Dibenzo-*p*-Dioxins and Dibenzofurans (CDDs/CDFs) Using Waters and Agilent Gas Chromatography JTandem-Mass Spectrometry (GC/MS/MS), Revision 1.0" is available at: https://www.sgsaxys.com/wp-content/uploads/2022/09/SGS-AXYS-Method-16130-Rev-10.pdf.
 ¹⁶ Pace Analytical Method PAM–16130–SSI, "Determination of 2,3,7,8-Substituted Tetra- through Octa-Chlorinated Dibenzo-*p*-Dioxins and Dibenzofurans (CDDs/CDFs) Using Shimadzu Gas Chromatography Mass Spectrometry (GC-MS/MS), Revision 1.1," is available at: *https://www.sgsaxys.com/wp-content/uploads/2022/*09/SGS-AXYS-Method-16130-Rev-10.pdf.
 ¹⁶ Pace Analytical Method PAM–16130–SSI, "Determination of 2,3,7,8-Substituted Tetra- through Octa-Chlorinated Dibenzo-*p*-Dioxins and Dibenzofurans (CDDs/CDFs) Using Shimadzu Gas Chromatography Mass Spectrometry (GC-MS/MS), Revisio

Parameter	Method	EPA 27 10	Standard methods ¹⁵	ASTM	Other
1. Aldrin	GC	617, 608.3	6630 B-2021 & C- 2021.	D3086–90, D5812–96 (02).	See footnote ³ p. 7, see footnote ⁴ O- 3104–83, see footnote ⁸ 3M0222.
2. Ametryn	GC/MS GC	625.1 507, 619	6410 B–2020.		See footnote ³ p. 83, see footnote ⁹ O– 3106–93, see footnote ⁶ p. S68.
3. Aminocarb	GC/MS	525.2, 625.1			See footnote 3 p. 94, see footnote 6 p. S60.
4. Atraton	HPLC GC	632. 619			See footnote ³ p. 83, see footnote ⁶ p. S68.
5. Atrazine	GC/MS GC	625.1. 507, 619, 608.3			See footnote ³ p. 83, see footnote ⁶ p.
	HPLC/MS GC/MS				S68, see footnote 9 O=3106=93. See footnote 12 O=2060=01. See footnote 11 O=1126=95.
6. Azinphos methyl	GC	614, 622, 1657			See footnote ³ p. 25, see footnote ⁶ p. S51.
7. Barban	GC–MS TLC	625.1			See footnote ¹¹ O–1126–95. See footnote ³ p. 104, see footnote ⁶ p. S64.
8. α-BHC	HPLC GC/MS GC	632. 625.1. 617, 608.3		D3086–90,	See footnote ³ p. 7, see footnote ⁸
9. β-BHC	GC/MS GC	625.1 ⁵ 617, 608.3	2021. 6410 B–2020 6630 B–2021 & C– 2021.	D5812–96(02). D3086–90, D5812–96(02).	3M0222. See footnote ¹¹ O–1126–95. See footnote ⁸ 3M0222.
10. δ-BHC	GC/MS GC	625.1 617, 608.3	6410 B–2020. 6630 B–2021 & C– 2021.	D3086–90, D5812–96(02).	See footnote ⁸ 3M0222.
11. γ-BHC (Lindane)	GC/MS GC	625.1 617, 608.3	6410 B-2020 6630 B-2021 & C- 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ , O– 3104–83, see footnote ⁸ 3M0222.
12. Captan	GC/MS GC	625.1 ⁵ 617, 608.3	6410 B–2020 6630 B–2021	D3086–90, D5812–96(02).	See footnote ¹¹ , O–1126–95. See footnote ³ p. 7.
13. Carbaryl	TLC				See footnote ³ p. 94, see footnote ⁶ p. S60.
14. Carbophenothion	HPLC HPLC/MS GC/MS GC	531.1, 632. 553 625.1 617, 608.3	 6630 B–2021		See footnote ¹² O–2060–01. See footnote ¹¹ O–1126–95. See footnote ⁴ page 27, see footnote ⁶ p. S73.
15. Chlordane	GC/MS GC	625.1. 617, 608.3	6630 B-2021 & C- 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ O– 3104–83, see footnote ⁸ 3M0222.
16. Chloropropham	GC/MS TLC	625.1	6410 B–2020.		See footnote ³ p. 104, see footnote ⁶ p. S64.
17. 2,4–D	HPLC GC/MS GC	632. 625.1. 615	6640 B–2021		See footnote ³ p. 115, see footnote ⁴ O-3105-83.
18. 4,4'-DDD	HPLC/MS GC	 617, 608.3		D3086–90, D5812–96(02).	See footnote ¹² O–2060–01. See footnote ³ p. 7, see footnote ⁴ O– 3105–83, see footnote ⁸ 3M0222.
19. 4,4'-DDE	GC/MS GC	625.1 617, 608.3	6410 B–2020. 6630 B–2021 & C– 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ , O- 3104–83, see footnote ⁸ 3M0222.
	GC/MS	625.1	6410 B–2020	D5812-90(02).	See footnote ¹¹ O–1126–95.

TABLE ID—LIST OF APPROVED TEST PROCEDURES FOR PESTICIDES¹

TABLE ID—LIST OF APPROVED TEST PROCEDURES FOR PESTICIDES 1—Continued

Parameter	Method	EPA ²⁷¹⁰	Standard methods ¹⁵	ASTM	Other
20. 4,4'-DDT	GC	617, 608.3	6630 B-2021 & C-	D3086–90,	See footnote ³ p. 7, see footnote ⁴ O-
	GC/MS	625.1	2021. 6410 B–2020.	D5812–96(02).	3104-83, see footnote ⁸ 3M0222.
21. Demeton-O	GC	614, 622			See footnote ³ p. 25, see footnote ⁶ p. S51.
22. Demeton-S	GC/MS	625.1 614, 622			See footnote ³ p. 25, see footnote ⁶ p.
	GC/MS	,			S51.
23. Diazinon	GC	507, 614, 622,			See footnote ³ p. 25, see footnote ⁴ O-
	GC/MS	1657. 525.2, 625.1			3104–83, see footnote ⁶ p. S51. See footnote ¹¹ O–1126–95.
24. Dicamba	GC HPLC/MS	615			See footnote ³ p. 115. See footnote ¹² O–2060–01.
25. Dichlofenthion	GC	622.1			See footnote ⁴ page 27, see footnote ⁶ p. S73.
26. Dichloran 27. Dicofol		608.2, 617, 608.3 617, 608.3	6630 B–2021		See footnote ³ p. 7. See footnote ⁴ O–3104–83.
28. Dieldrin		617, 608.3	6630 B-2021 & C-	D3086–90,	See footnote 3 p. 7, see footnote 4 O-
	GC/MS	625.1	2021. 6410 B–2020	D5812–96(02).	3104–83, see footnote ⁸ 3M0222. See footnote ¹¹ O–1126–95.
29. Dioxathion	GC	614.1, 1657			See footnote ⁴ page 27, see footnote ⁶ p. S73.
30. Disulfoton	GC	507, 614, 622, 1657.			See footnote ³ p. 25, see footnote ⁶ p. S51.
31. Diuron	GC/MS	525.2, 625.1			See footnote ¹¹ O–1126–95. See footnote ³ p. 104, see footnote ⁶
	-				p. S64.
	HPLC HPLC/MS				See footnote ¹² O–2060–01.
32. Endosulfan I	GC	617, 608.3	6630 B-2021 & C- 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ O- 3104–83, see footnote ⁸ 3M0222.
33. Endosulfan II	GC/MS	625.1 ⁵ 617, 608.3	6410 B-2020 6630 B-2021 & C-	D3086–90,	See footnote ¹³ O–2002–01. See footnote ³ p. 7, see footnote ⁸
	GC/MS		2021. 6410 B–2020	D5812–96(02).	3M0222. See footnote ¹³ O–2002–01.
34. Endosulfan Sulfate	GC	617, 608.3	6630 C-2021		See footnote ⁸ 3M0222.
35. Endrin	GC/MS GC	505, 508, 617,	6410 B-2020. 6630 B-2021 & C-	D3086–90,	See footnote 3 p. 7, see footnote 4 O-
	GC/MS	1656, 608.3. 525.1, 525.2, 625.1 ⁵ .	2021. 6410 B–2020.	D5812–96(02).	3104–83, see footnote ⁸ 3M0222.
36. Endrin aldehyde		617, 608.3	6630 C-2021		See footnote 8 3M0222.
37. Ethion	GC/MS GC	625.1 614, 614.1, 1657	6410 B–2020		See footnote 4 page 27, see foot-
	GC/MS	625.1			note ⁶ , p. S73. See footnote ¹³ O–2002–01.
38. Fenuron	TLC				See footnote ³ p. 104, see footnote ⁶ p. S64.
	HPLC HPLC/MS				See footnote ¹² O–2060–01.
39. Fenuron-TCA					See footnote ³ p. 104, see footnote ⁶ p. S64.
0. Heptachlor	HPLC	632. 505, 508, 617,	6630 B-2021 & C-	D3086–90,	See footnote ³ p. 7, see footnote ⁴ O–
		1656, 608.3.	2021.	D5812–96(02).	3104–83, see footnote ⁸ 3M0222.
	GC/MS	625.1.	6410 B–2020.		
11. Heptachlor epoxide	GC	617, 608.3	6630 B-2021 & C- 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ O– 3104–83, see footnote ⁶ p. S73, see footnote ⁸ 3M0222.
12. Isodrin	GC/MS GC	625.1 617. 608.3	6410 B–2020. 6630 B–2021 & C–		See footnote ⁴ O-3104-83, see foot-
F2. 19001111	GC/MS	625.1.	2021.		note ⁶ p. S73.
13. Linuron	GC				See footnote ³ p. 104, see footnote ⁶
	HPLC	632.			p. S64.
	HPLC/MS GC/MS	553			See footnote ¹² O–2060–01. See footnote ¹¹ O–1126–95.
14. Malathion	GC	614, 1657	6630 B–2021		See footnote ³ p. 25, see footnote ⁶ p. S51.
15. Methiocarb	GC/MS	625.1			See footnote ¹¹ O–1126–95. See footnote ³ p. 94, see footnote ⁶ p.
		632.			S60.
	HPLC HPLC/MS				See footnote ¹² O–2060–01.
16. Methoxychlor		505, 508, 608.2, 617, 1656, 608.3.	6630 B-2021 & C- 2021.	D3086–90, D5812–96(02).	See footnote ³ p. 7, see footnote ⁴ O- 3104–83, see footnote ⁸ 3M0222.
	GC/MS	525.1, 525.2, 625.1.			See footnote ¹¹ O–1126–95.

TABLE ID—LIST OF APPROVED TEST PROCEDURES FOR PESTICIDES 1—Continued

Parameter	Method	EPA 27 10	Standard methods ¹⁵	ASTM	Other
47. Mexacarbate	TLC				See footnote ³ p. 94, see footnote ⁶ p.
	HPLC	632.			S60.
	GC/MS				
48. Mirex	GC			D3086–90,	See footnote ³ p. 7, see footnote ⁴ O-
	GC/MS	625.1.	2021.	D5812–96(02).	3104–83.
49. Monuron	TLC				See footnote ³ p. 104, see footnote ⁶ p. S64.
	HPLC	632.			p. 364.
50. Monuron-TCA	TLC				See footnote ³ p. 104, see footnote ⁶
	HPLC	632.			p. S64.
51. Neburon	TLC				See footnote ³ p. 104, see footnote ⁶
	HPLC	632.			p. S64.
	HPLC/MS				See footnote ¹² O–2060–01.
52. Parathion methyl	GC	614, 622, 1657	6630 B–2021		See footnote ⁴ page 27, see footnote ³ p. 25.
	GC/MS	625.1			See footnote ¹¹ O–1126–95.
53. Parathion ethyl	GC	614	6630 B-2021		See footnote ⁴ page 27, see footnote ³
	GC/MS				p. 25. See footnote ¹¹ O–1126–95.
54. PCNB	GC	608.1, 617, 608.3	6630 B-2021 & C-	D3086–90,	See footnote ³ p. 7.
55. Perthane	GC	617, 608.3	2021.	D5812–96(02). D3086–90,	See footnote ⁴ O-3104-83.
55. Fertilane	GC	017, 000.5		D5812–96(02).	See 10011101e · 0-3104-63.
56. Prometon	GC	507, 619			See footnote ³ p. 83, see footnote ⁶ p. S68, see footnote ⁹ O–3106–93.
	GC/MS	525.2, 625.1			See footnote ¹¹ O–1126–95.
57. Prometryn	GC	507, 619			See footnote ³ p. 83, see footnote ⁶ p.
	GC/MS	525.1, 525.2, 625.1.			S68, see footnote ⁹ O–3106–93. See footnote ¹³ O–2002–01.
58. Propazine	GC	507, 619, 1656,			See footnote ³ p. 83, see footnote ⁶ p.
	GC/MS	608.3. 525.1, 525.2,			S68, see footnote ⁹ O–3106–93.
50 Bronhom	TLC	625.1			Can fastrata 3 n. 10. and fastrata 6 n.
59. Propham					See footnote ³ p. 10, see footnote ⁶ p. S64.
	HPLC HPLC/MS	632.			See footnote ¹² O–2060–01.
60. Propoxur	TLC				See footnote ³ p. 94, see footnote ⁶ , p.
		632.			S60.
61. Secbumeton	HPLC				See footnote ³ p. 83, see footnote ⁶ p.
					S68.
62. Siduron	GC				See footnote ³ p. 104, see footnote ⁶
					p. S64.
	HPLC HPLC/MS				See footnote ¹² O–2060–01.
63. Simazine	GC	505, 507, 619,			See footnote ³ p. 83, see footnote ⁶ p.
	GC/MS	1656, 608.3. 525.1, 525.2,			S68, see footnote ⁹ O–3106–93. See footnote ¹¹ O–1126–95.
	GC/M3	625.1.			See 1001101e ··· 0-1126-95.
64. Strobane	GC	617, 608.3	6630 B-2021 & C- 2021.		See footnote ³ p. 7.
65. Swep	TLC		2021.		See footnote ³ p. 104, see footnote ⁶
	HPLC	632.			p. S64.
66. 2,4,5–T	GC	615	6640 B-2021		See footnote ³ p. 115, see footnote ⁴
67. 2,4,5-TP (Silvex)	GC	615	6640 B–2021		O-3105-83. See footnote ³ p. 115, see footnote ⁴
68. Terbuthylazine	GC	619, 1656, 608.3			O–3105–83. See footnote ³ p. 83, see footnote ⁶ p.
	GC/MS				S68. See footnote ¹³ O–2002–01.
69. Toxaphene	GC	505, 508, 617,	6630 B-2021 & C-	D3086–90,	See footnote 3 p. 7, see footnote 8, see
	GC/MS	1656, 608.3. 525.1, 525.2,	2021. 6410 B–2020.	D5812–96(02).	footnote ⁴ O-3105-83.
		625.1.			
		LENO 617 607	6630 B-2021		See footnote ³ p. 7, see footnote ⁹ O-
70. Trifluralin	GC	508, 617, 627, 1656, 608.3.	0030 B-2021		3106–93.

 Table ID notes:
 1

 1
 Pesticides are listed in this table by common name for the convenience of the reader. Additional pesticides may be found under table IC of this section, where entries are listed by chemical name.

 2
 The standardized test procedure to be used to determine the method detection limit (MDL) for these test procedures is given at appendix B to this part, Definition and Procedure for the Determination of the Method Detection Limit.

³Methods for Benzidine, Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater. September 1978. U.S. EPA. This EPA publication includes thin-layer chromatography (TLC) methods. ⁴ Methods for the Determination of Organic Substances in Water and Fluvial Sediments, Techniques of Water-Resources Investigations of the U.S. Geological Sur-

vey, Book 5, Chapter A3. 1987. USGS. 5 The method may be extended to include α -BHC, γ -BHC, endosulfan I, endosulfan II, and endrin. However, when they are known to exist, Method 608 is the pre-

ferred method. Selected Analytical Methods Approved and Cited by the United States Environmental Protection Agency, Supplement to the 15th Edition of Standard Methods for the Examination of Water and Wastewater. 1981. American Public Health Association (APHA).

Teach analyst must make an initial, one-time, demonstration of their ability to generate acceptable precision and accuracy with Methods 608.3 and 625.1 in accord-ance with procedures given in Section 8.2 of each of these methods. Additionally, each laboratory, on an on-going basis, must spike and analyze 10% of all samples analyzed with Method 608.3 or 5% of all samples analyzed with Method 625.1 to monitor and evaluate laboratory data quality in accordance with Sections 8.3 and 8.4 of these methods. When the recovery of any parameter falls outside the warning limits, the analytical results for that parameter in the unspiked sample are sus-pect. The results should be reported, but cannot be used to demonstrate regulatory compliance. These quality control requirements also apply to the Standard Meth-ods, ASTM Methods, and other methods cited.

ods, ASTM Methods, and other methods cited. ⁸ Organochlorine Pesticides and PCBs in Wastewater Using Empore[™] Disk. Revised October 28, 1994. 3M Corporation. ⁹Method O–31 is in Open File Report 94–37, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Tri-azine and Other Nitrogen-Containing Compounds by Gas Chromatography with Nitrogen Phosphorus Detectors. 1994. USGS. ¹⁰ EPA Methods 608.1, 608.2, 614, 614.1, 615, 617, 619, 622, 622.1, 627, and 632 are found in Methods for the Determination of Nonconventional Pesticides in Municipal and Industrial Wastewater, EPA 821–R-92–002, April 1992, U.S. EPA. EPA Methods 505, 507, 508, 525.1, 531.1 and 553 are in Methods for the Deter-mination of Nonconventional Pesticides in Municipal and Industrial Wastewater, Volume II, EPA 821–R–93–010B, 1993, U.S. EPA. EPA Method 525.2 is in Deter-mination of Organic Compounds in Drinking Water by Liquid-Solid Extraction and Capillary Column Gas Chromatography/Mass Spectrometry, Revision 2.0, 1995, U.S. EPA. EPA methods 1656 and 1657 are in Methods for The Determination of Nonconventional Pesticides In Municipal and Industrial Wastewater, Volume I, EPA 821–R–93–010A, 1993, U.S. EPA. Methods 608.3 and 625.1 are available at: cwa-methods/approved-cwa-test-methods-organic-compounds. ¹¹Method O–1126–95 is in Open-File Report 95–181 Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of pes-¹¹Method O–1126–95 is In Open-File Report 95–181 Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of pes-¹¹Method O–1126–95 is In Open-File Report 95–181 Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of pes-¹¹Method O–1126–95 is In Open-File Report 95–181 Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of pes-¹¹Method O–1126–95 is In Open-File Report 95–181 Methods of Ana

¹¹Method O–1126–95 is in Open-File Report 95–181, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of pes-ticides in water by C–18 solid-phase extraction and capillary-column gas chromatography/mass spectrometry with selected-ion monitoring. 1995. USGS. ¹²Method O–2060–01 is in Water-Resources Investigations Report 01–4134, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Pesticides in Water by Graphitized Carbon-Based Solid-Phase Extraction and High-Performance Liquid Chromatography/Mass Spectrometry.

2001. USGS.

¹³Method O-2002-01 is in Water-Resources Investigations Report 01-4098, Methods of Analysis by the U.S. Geological Survey National Water Quality Labora-

¹⁴Method 0–2002–011s in Vater-Nestrices investigations investigations investigations of analysis by the U.S. Geological Survey National Vater Quality Laboratory—Determination of moderate-use pesticides in vater by S0Id-phase extraction and capillary-column gas chromatography/mass spectrometry. 2001. USGS. ¹⁴Method 0–1121–91 is in Open-File Report 91–519, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of organonitrogen herbicides in water by solid-phase extraction and capillary-column gas chromatography/mass spectrometry. 2001. USGS. ¹⁵Please refer to the following applicable Quality Control Section: Part 6000 Methods, Individual Organic Compounds 6020 (2019). These Quality Control Standards are available for download at *www.standardmethods.org* at no charge.

TABLE IH—LIST OF APPROVED MICROBIOLOGICAL METHODS FOR AMBIENT WATER

Parameter and units	Method ¹	EPA	Standard methods	AOAC, ASTM, USGS	Other
		Bacteria	·	·	
1. Coliform (fecal), number per 100 mL.	Most Probable Number (MPN), 5 tube, 3 dilution, or.	p. 132 ³	9221 E-2014, 9221 F-2014. ³²		
2. Coliform (total), num- ber per 100 mL.	Membrane filter (MF) ² , single step MPN, 5 tube, 3 dilution, or	p. 124 ³ p. 114 ³		B-0050-85.4	
·	MF ² , single step or MF ² , two step with enrichment	p. 108 ³ p. 111 ³	9222 B-2015.27	B-0025-85.4	
3. <i>E. coli,</i> number per 100 mL.	MPN ⁵⁷¹³ , multiple tube, or		9221 B.3–2014/9221 F–2014. ^{10 12 32}		
	Multiple tube/multiple well, or		9223 B-2016 ¹¹	991.15 ⁹	Colilert ^{® 11 15} , Colilert-18 [®] . ^{11 14}
	MF ²⁵⁶⁷ , two step, or	1103.2 ¹⁸	9222 B-2015/9222 I- 2015 ¹⁷ , 9213 D- 2007.	D5392–93.8	
	Single step	1603.1 ¹⁹ , 1604 ²⁰			m-ColiBlue24 ^{® 16} , KwikCount [™] FC ^{28 29}
 Fecal streptococci, number per 100 mL. 	MPN, 5 tube, 3 dilution, or	p. 139 ³	9230 B–2013.		
	MF ² , or Plate count	p. 136 ³ p. 143. ³	9230 C-2013 30	B-0055-85.4	
i. Enterococci, number per 100 mL.	MPN ⁵⁷ , multiple tube/multiple well, or		9230 D–2013	D6503–99 ⁸	Enterolert®.11 21
20. 100 mL.	MF ²⁵⁶⁷ two step, or Single step, or Plate count	1106.2 ²² 1600.1 ²³ p. 143. ³	9230 C-2013 ³⁰ 9230 C-2013. ³⁰	D5259–92. ⁸	
		Protozoa	·		
		1000.24 1000.25			

1622²⁴, 1623²⁵, 6. Cryptosporidium Filtration/IMS/FA 1623.1.25 31 1623²⁵, 1623.1.^{25 31} 7. Giardia Filtration/IMS/FA

Table 1H notes:

¹ The method must be specified when results are reported.

²A 0.45-µm membrane filter (MF) or other pore size certified by the manufacturer to fully retain organisms to be cultivated and to be free of extractables which could interfere with their growth.

³ Microbiological Methods for Monitoring the Environment, Water and Wastes. EPA/600/8–78/017. 1978. US EPA. ⁴ U.S. Geological Survey Techniques of Water-Resource Investigations, Book 5, Laboratory Analysis, Chapter A4, Methods for Collection and Analysis of Aquatic Biological and Microbiological Samples. 1989. USGS.

⁵Tests must be conducted to provide organism enumeration (density). Select the appropriate configuration of tubes/filtrations and dilutions/volumes to account for the quality, character, consistency, and anticipated organism density of the water sample. ⁶When the MF method has not been used previously to test waters with high turbidity, large numbers of noncoliform bacteria, or samples that may contain orga-nisms stressed by chlorine, a parallel test should be conducted with a multiple-tube technique to demonstrate applicability and comparability of results ⁷To assess the comparability of results obtained with individual methods, it is suggested that side-by-side tests be conducted across seasons of the year with the water samples routinely tested in accordance with the most current *Standard Methods for the Examination of Water and Wastewater* or EPA alternate test procedure (ATD) middlingo

(ATP) guidelines.

 ⁸ Annual Book of ASTM Standards—Water and Environmental Technology. Section 11.02. 2000, 1999, 1996. ASTM International.
 ⁹ Official Methods of Analysis of AOAC International, 16th Edition, Volume I, Chapter 17. 1995. AOAC International.
 ¹⁰ The multiple-tube fermentation test is used in 9221B.3–2014. Lactose broth may be used in lieu of lauryl tryptose broth (LTB), if at least 25 parallel tests are conducted between this broth and LTB using the water samples normally tested, and this comparison demonstrates that the false-positive rate and false-negative rate for total coliform using lactose broth is less than 10 percent. No requirement exists to run the completed phase on 10 percent of all total coliform-positive tubes on a seasonal basis.

These tests are collectively known as defined enzyme substrate tests.

¹² After prior enrichment in a presumptive medium for total coliform using 9221B.3–2014, all presumptive tubes or bottles showing any amount of gas, growth or acidity within 48 h ± 3 h of incubation shall be submitted to 9221F–2014. Commercially available EC–MUG media or EC media supplemented in the laboratory with

¹³ Samples shall be enumerated by the multiple-tube or multiple-well procedure. Using multiple-tube procedures, employ an appropriate tube and dilution configura-tion of the sample as needed and report the Most Probable Number (MPN). Samples tested with Colilert® may be enumerated with the multiple-well procedures, Quanti-Tray® or Quanti-Tray® 2000, and the MPN calculated from the table provided by the manufacturer. ¹⁴ Colilert-18® is an optimized formulation of the Colilert® for the determination of total coliforms and *E. coli* that provides results within 18 h of incubation at 35 °C,

rather than the 24 h required for the Colliert® test and is recommended for marine water samples. ¹⁵Descriptions of the Colliert®, Colliert-18®, Quanti-Tray^{supreg}, and Quanti-Tray[®]/2000 may be obtained from IDEXX Laboratories Inc. ¹⁶A description of the mColiBlue24® test may be obtained from Hach Company.

¹⁷ Subject coliform positive samples determined by 9222B–2015 or other membrane filter procedure to 9222I–2015 using NA–MUG media. ¹⁸ Method 1103.2: *Escherichia coli* (*E. coli*) in Water by Membrane Filtration Using membrane-Thermotolerant *Escherichia coli* Agar (mTEC), EPA–821–R–23–009. September 2023. US EPA.

¹⁹Method 1603.1: Escherichia coli (E. coli) in Water by Membrane Filtration Using Modified membrane-Thermotolerant Escherichia coli Agar (Modified mTEC), EPA-821-R-23-008. September 2023. US EPA.

20 Method 1604: Total Coliforms and Escherichia coli (E. coli) in Water by Membrane Filtration by Using a Simultaneous Detection Technique (MI Medium), EPA 821-R-02-024. September 2002. US EPA. ²¹ A description of the Enterolert[®] test may be obtained from IDEXX Laboratories Inc.

106.2: Enterococci in Water by Membrane Filtration Using membrane-Enterococcus-Esculin Iron Agar (mE-EIA), EPA-821-R-23-007. September 22 Method 2023. US EPA. ²³Method 1600.1: Enterococci in Water by Membrane Filtration Using membrane-Enterococcus Indoxyl-β-D-Glucoside Agar (mEl), EPA-821-R-21-006. Sep-

tember 2023. US EPA.

²⁴Method 1622 uses a filtration, concentration, immunomagnetic separation of oocysts from captured material, immunofluorescence assay to determine concentra-tions, and confirmation through vital dye staining and differential interference contrast microscopy for the detection of *Cryptosporidium*. Method 1622: *Cryptosporidium* in Water by Filtration/IMS/FA, EPA-821-R-05-001. December 2005. US EPA.

²⁵Methods 1623 and 1623.1 use a filtration, concentration, immunomagnetic separation of oocysts and cysts from captured material, immunofluorescence assay to determine concentrations, and confirmation through vital dye staining and differential interference contrast microscopy for the simultaneous detection of *Cryptosporidium* and *Giardia* oocysts and cysts. Method 1623: *Cryptosporidium* and *Giardia* in Vater by Filtration/IMS/FA. EPA–821–R–05–002. December 2005. US EPA.
EPA. Method 1623: 1: *Cryptosporidium* and *Giardia* in Water by Filtration/IMS/FA. EPA–821–R–05–002. December 2005. US EPA.
²⁶ On a monthly basis, at least ten blue colonies from positive samples must be verified using Lauryl Tryptose Broth and EC broth, followed by count adjustment

based on these results; and representative non-blue colonies should be verified using Lauryl Tryptose Broth. Where possible, verifications should be done from randomized sample sources. ²⁷On a monthly basis, at least ten sheen colonies from positive samples must be verified using Lauryl Tryptose Broth and brilliant green lactose bile broth, followed

by count adjustment based on these results; and representative non-sheen colonies should be verified using Lauryl Tryptose Broth. Where possible, verifications should be done from randomized sample sources. ²⁸ A description of KwikCount[™] EC may be obtained from Roth Bioscience, LLC

²⁹ Approved for the analyses of *E. coli* in freshwater only. ³⁰ Verification of colonies by incubation of BHI agar at 10 \pm 0.5 °C for 48 \pm 3 h is optional. As per the Errata to the 23rd Edition of *Standard Methods for the Examination of Water and Wastewater* "Growth on a BHI agar plate incubated at 10 \pm 0.5 °C for 48 \pm 3 h is further verification that the colony belongs to the genus Enterococcus

³¹ Method 1623.1 includes updated acceptance criteria for IPR, OPR, and MS/MSD and clarifications and revisions based on the use of Method 1623 for years and technical support questions. 329221 F.2-2014 allows for simultaneous detection of *E. coli* and thermotolerant fecal coliforms by adding inverted vials to EC-MUG; the inverted vials collect gas

produced by thermotolerant fecal coliforms.

(b) The material listed in this paragraph (b) is incorporated by reference into this section with the approval of the Director of the Federal Register under 5 U.S.C. 552(a) and 1 CFR part 51. All approved incorporation by reference (IBR) material is available for inspection at the EPA and at the National Archives and Records Administration (NARA). Contact the EPA at: EPA's Water Docket, EPA West, 1301 Constitution Avenue NW, Room 3334, Washington, DC 20004; telephone: 202-566-2426; email: docketcustomerservice@epa.gov. For information on the availability of this material at NARA, visit www.archives.gov/federal-register/cfr/ ibr-locations or email fr.inspection@ nara.gov. The material may be obtained from the following sources in this paragraph (b).

(8) Office of Water, U.S. Environmental Protection Agency (U.S. EPA), mail code 4303T, 1301 Constitution Avenue NW, Washington, DC 20460; website: www.epa.gov/cwamethods.

(i) Method 245.7, Mercury in Water by Cold Vapor Atomic Fluorescence Spectrometry. Revision 2.0, February 2005. EPA-821-R-05-001. Table IB, Note 17.

(ii) Method 1103.2: Escherichia coli (E. coli) in Water by Membrane Filtration Using membrane-Thermotolerant Escherichia coli Agar (mTEC), EPA-821-R-23-009. September 2023. Table IH, Note 18.

(iii) Method 1106.2: Enterococci in Water by Membrane Filtration Using membrane-Enterococcus-Esculin Iron Agar (mE-EIA), EPA-821-R-23-007. September 2023. Table IH, Note 22.

(iv) Method 1600.1: Enterococci in Water by Membrane Filtration Using membrane-Enterococcus Indoxyl-β-D-Glucoside Agar (mEI), EPA-821-R-23-006, September 2023. Table 1A, Note 24: Table IH. Note 23.

(v) Method 1603.1: Escherichia coli (E. coli) in Water by Membrane Filtration Using Modified membrane-Thermotolerant Escherichia coli Agar (Modified mTEC), EPA-821-R-23-008, September 2023. Table IA, Note 21; Table IH, Note 19.

(vi) Method 1604: Total Coliforms and Escherichia coli (E. coli) in Water by

Membrane Filtration Using a Simultaneous Detection Technique (MI Medium). September 2002. EPA-821-R-02-024. Table IH, Note 21.

(vii) Whole Effluent Toxicity Methods Errata Sheet, EPA 821-R-02-012-ES. December 2016, Table IA, Notes 25, 26, and 27.

(viii) Method 1623: Cryptosporidium and Giardia in Water by Filtration/IMS/ FA. December 2005. EPA-821-R-05-002. Table IH, Note 26.

(ix) Method 1623.1: Cryptosporidium and Giardia in Water by Filtration/IMS/ FA. EPA 816-R-12-001. January 2012. U.S. EPA, Table IH, Notes 25 and 31.

(x) Method 1627, Kinetic Test Method for the Prediction of Mine Drainage Quality. December 2011. EPA-821-R-09–002. Table IB, Note 69.

(xi) Method 1664. n-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated *n*-Hexane Extractable Material (SGT-HEM; Nonpolar Material) by Extraction and Gravimetry. Revision A, February 1999. EPA-821-R-98-002. Table IB, Notes 38 and 42.

(xii) Method 1664, n-Hexane Extractable Material (HEM; Oil and Grease) and Silica Gel Treated n-Hexane Extractable Material (SGT–HEM; Nonpolar Material) by Extraction and Gravimetry, Revision B, February 2010. EPA–821–R–10–001. Table IB, Notes 38 and 42.

(xiii) Method 1669, Sampling Ambient Water for Trace Metals at EPA Water Quality Criteria Levels. July 1996. Table IB, Note 43.

(xiv) Method 1680: Fecal Coliforms in Sewage Sludge (Biosolids) by Multiple-Tube Fermentation using Lauryl Tryptose Broth (LTB) and EC Medium. September 2014. EPA–821–R–14– 009.Table IA, Note 15.

(xv) Method 1681: Fecal Coliforms in Sewage Sludge (Biosolids) by Multiple-Tube Fermentation using A–1 Medium. July 2006. EPA 821–R–06–013. Table IA, Note 20.

(xvi) Method 1682: *Salmonella* in Sewage Sludge (Biosolids) by Modified Semisolid Rappaport-Vassiliadis (MSRV) Medium. September 2014. EPA 821–R–14–012. Table IA, Note 23.

(10) American Public Health Association, 800 I Street, NW, Washington, DC 20001; phone: (202)777–2742, website: www.standardmethods.org.

(i) Standard Methods for the Examination of Water and Wastewater.
14th Edition, 1975. Table IB, Notes 27 and 86.

(ii) Standard Methods for the Examination of Water and Wastewater. 15th Edition, 1980, Table IB, Note 30; Table ID.

(iii) Selected Analytical Methods Approved and Cited by the United States Environmental Protection Agency, Supplement to the 15th Edition of Standard Methods for the Examination of Water and Wastewater. 1981. Table IC, Note 6; Table ID, Note 6.

(iv) Standard Methods for the Examination of Water and Wastewater. 18th Edition, 1992. Tables IA, IB, IC, ID, IE, and IH.

(v) Standard Methods for the Examination of Water and Wastewater. 19th Edition, 1995. Tables IA, IB, IC, ID, IE, and IH.

(vi) Standard Methods for the Examination of Water and Wastewater. 20th Edition, 1998. Tables IA, IB, IC, ID, IE, and IH.

(vii) Standard Methods for the Examination of Water and Wastewater. 21st Edition, 2005. Table IB, Notes 17 and 27.

(viii) 2120, Color. Revised September 4, 2021. Table IB.

- (ix) 2130, Turbidity. Revised 2020. Table IB.
- (x) 2310, Acidity. Revised 2020. Table IB.

(xi) 2320, Alkalinity. Revised 2021. Table IB.

- (xii) 2340, Hardness. Revised 2021. Table IB.
- (xiii) 2510, Conductivity. Revised 2021. Table IB.
- (xiv) 2540, Solids. Revised 2020. Table IB.
- (xv) 2550, Temperature. 2010. Table IB.
- (xvi) 3111, Metals by Flame Atomic Absorption Spectrometry. Revised 2019. Table IB.
- (xvii) 3112, Metals by Cold-Vapor
- Atomic Absorption Spectrometry.

Revised 2020. Table IB.

- (xviii) 3113, Metals by Electrothermal Atomic Absorption Spectrometry. Revised 2020. Table IB.
- (xix) 3114, Arsenic and Selenium by Hydride Generation/Atomic Absorption
- Spectrometry. Revised 2020, Table IB.
- (xx) 3120, Metals by Plasma Emission Spectroscopy. Revised 2020. Table IB.
- (xxi) 3125, Metals by Inductively Coupled Plasma-Mass Spectrometry.
- Revised 2020. Table IB.
- (xxii) 3500-Al, Aluminum. Revised 2020. Table IB.
- (xxiii) 3500-As, Arsenic. Revised 2020. Table IB.
- (xxiv) 3500-Ca, Calcium. Revised 2020. Table IB.
- (xxv) 3500-Cr, Chromium. Revised 2020. Table IB.
- (xxvi) 3500-Cu, Copper. Revised 2020. Table IB.
- (xxvii) 3500-Fe, Iron. 2011. Table IB. (xxviii) 3500-Pb, Lead. Revised 2020. Table IB.
- (xxix) 3500-Mn, Manganese. Revised 2020. Table IB.
- (xxx) 3500–K, Potassium. Revised 2020. Table IB.
- (xxxi) 3500-Na, Sodium. Revised 2020. Table IB.
- (xxxii) 3500–V, Vanadium. 2011. Table IB.
- (xxxiii) 3500-Zn, Zinc. Revised 2020. Table IB.
- (xxxiv) 4110, Determination of Anions by Ion Chromatography. Revised 2020. Table IB.
- (xxxv) 4140, Inorganic Anions by Capillary Ion Electrophoresis. Revised 2020. Table IB.
- (xxxvi) 4500–B, Boron. 2011. Table IB.
- (xxxvii) 4500 Cl⁻, Chloride. Revised 2021. Table IB.
- (xxxviii) 4500-Cl, Chlorine (Residual). 2011. Table IB.
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- (xlv) 4500–N_(org), Nitrogen (Organic). Revised 2021. Table IB.
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- (xlvii) 4500–P, Phosphorus. Revised 2021. Table IB.
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- (xlix) 4500–S $^{2-}$, Sulfide. Revised 2021. Table IB.
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Table II—Required Containers, **Preservation Techniques, and Holding** Times

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⁵ ASTM D7365–09a (15) specifies treatment options for samples containing oxidants (e.g., chlorine) for cyanide analyses. Also, Section 9060A of Standard Methods for the Examination of Water and Wastewater (23rd edition) addresses dechlorination procedures for microbiological analyses. * * *

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