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DEPARTMENT OF NATURAL RESOURCES

NR 219.04

Chapter NR 219

ANALYTICAL TEST METHODS AND PROCEDURES

NR 219.01	Purpose.	NR 219.033	Alternate test procedures.
NR 219.02	Applicability.	NR 219.037	Laboratory certification or registration.
NR 219.03	Definitions.	NR 219.04	Identification of test procedures.

Note: A number of the references cited in this chapter are no longer in print. Copies of references which are out–of–print are available at any public library by inter–library loan.

NR 219.01 Purpose. The purpose of this chapter is to establish analytical test methods, preservation procedures, requirements for laboratories, and procedures applicable to effluent limitations for discharges from point sources as authorized by ss. 299.11 and 283.55 (1), Stats.

History: Cr. Register, August, 1976, No. 248, eff. 9–1–76; am. Register, April, 1986, No. 364, eff. 8–28–86; am. Register, June, 1986, No. 366, eff. 7–1–86; am. Register, April, 1988, No. 388, eff. 5–1–88; corrections made under s. 13.93 (2m) (b) 7., Stats., Register, November, 1996, No. 491.

NR 219.02 Applicability. (1) The procedures prescribed herein shall, except as provided in s. NR 219.037, be used in the determination of concentrations and quantities of pollutant parameters as required for:

(a) An application submitted to the department for a permit under ch. 283, Stats.

(b) Reports required to be submitted by dischargers in accordance with the conditions of issued permits.

(2) Section NR 219.037 requires that laboratories conducting tests under this chapter be certified, registered, or approved under ch. NR 149.

History: Cr. Register, August, 1976, No. 248, eff. 9–1–76; am. Register, April, 1986, No. 364, eff. 8–28–86; am. (1) (intro.), Register, June, 1986, No. 366, eff. 7–1–86; correction in (1) (a) made under s. 13.93 (2m) (b) 7., Stats., Register, November, 1996, No. 491; correction in (2) made under s. 13.93 (2m) (b) 7., Stats., Register October 2002 No. 562; correction in (2) made under s. 13.93 (2m) (b) 7., Stats., Register November 2004 No. 587; CR 13–112: am. (1), (2) Register May 2015 No. 713, eff. 6–1–15.

NR 219.03 Definitions. As used in this chapter:

(1) "EPA" means the U.S. environmental protection agency.

(2) "Department" means the department of natural resources.

(3) "Sludge" is defined in ss. NR 204.03 (55) and 214.03 (34). History: Cr. Register, August, 1976, No. 248, eff. 9–1–76; am. (1), (2), (3) and (4m), Register, January, 1978, No. 265, eff. 2–1–78; r. and recr. Register, June, 1986, No. 366, eff. 7–1–86; r. and recr. (1), r. (3) and (4), Register, November, 1992, No. 443, eff. 12–1–92: CR 04–033: cr. (3) Register November 2004 No. 587, eff. 12–1–94.

NR 219.033 Alternate test procedures. Approvals of alternate test procedures for nationwide use and specific discharges are granted by EPA. The department may approve the use of an alternate test procedure on a case–by–case basis if the criteria for approval of the alternate procedure established in s. NR 149.42 are met. If the department or the EPA approves an alternate test procedure, it shall be considered equivalent to the approved method.

Note: The federal requirements for alternate test procedure approval are given in 40 CFR 136.5.

History: Cr. Register, August, 1976, No. 248, eff. 9–1–76; r. and recr. January, 1978, No. 265, eff. 2–1–78; renum. from NR 219.04 and am. Register, June, 1986, No. 366, eff. 7–1–86; r. and recr. Register, November, 1992, No. 443, eff. 12–1–92; am. Register, February, 1996, No. 482, eff. 3–1–96; correction made under s. 13.92 (4) (b) 7., Stats., Register May 2009 No. 641, eff. 6–1–09; CR 13–112: renum. from 219.05 Register May 2015 No. 713, eff. 6–1–15.

NR 219.037 Laboratory certification or registration. Bacteriological analyses of groundwater samples, and all radiological analyses shall be performed by the state laboratory of hygiene or at a laboratory certified or approved by the department of agriculture, trade and consumer protection. Other laboratory test results, including effluent toxicity, submitted to the department under a WPDES permit shall be performed by a laboratory certified or registered under ch. NR 149. The following tests are excluded from this requirement:

(1) Temperature,

- (2) Turbidity,
- (3) Bacteria tests in wastewater effluent and sludges,
- (4) pH,
- (5) Chlorine residual,
- (6) Specific conductance,
- (7) Physical properties of soils and sludges,
- (8) Nutrient tests of soils and sludges,
- **(9)** Flow measurements.

History: Cr. Register, April, 1986, No. 364, eff. 8–28–86; renum. from NR 219.07 and am. (intro.) Register, November, 1992, No. 443, eff. 7–1–93; am. Register February, 1996, No. 482, eff. 3–1–96; correction in (intro.) made under s. 13.93 (2m) (b) 6., Stats., Register November 2004 No. 587; CR 13–112: renum. from 219.06 Register May 2015 No. 713, eff. 6–1–15.

NR 219.04 Identification of test procedures. (1) ANALYTICAL TEST PROCEDURES. Parameters or pollutants, for which wastewater analytical methods are approved, are listed together with test procedure descriptions and references in tables A to H. Parameters or pollutants, for which sludge analytical methods are approved, are listed together with test procedure descriptions and references in table EM. The discharge values for the listed parameters shall be determined by one of the standard analytical test procedures identified in a table under this subsection or by an alternate test procedure established under ss. NR 219.033 and 149.12.

(2) SAMPLE PRESERVATION PROCEDURES. Sample preservation techniques, container materials, and maximum allowable holding times for parameters identified in tables A to H are prescribed in table F. Sludge samples shall be preserved at the time of collection by cooling to less than or equal to 65° C where required. All samples requiring thermal preservation at less than or equal to 65° C shall be cooled immediately after collection, and the required temperature maintained during shipping. Any person may apply for a variance from the prescribed preservation procedures applicable to samples taken from a specific discharge. Applications for variances may be made by letters to the regional administrator and shall provide sufficient data to assure that the variance does not adversely affect the integrity of the sample. The regional administrator will make a decision on whether to approve or deny a variance within 90 days of receipt of the application.

(3) TEMPERATURE REPORTING PROCEDURES. Samples cooled with ice packs or not in direct contact with ice during shipping shall be cooled to less than or equal to 6° C prior to shipping, and a temperature blank shall be submitted with the samples. Samples cooled during shipping with ice packs may not be recorded as received on ice. Samples may be recorded as received on ice only if solid ice is present in the cooler at the time the samples are received. If the samples are not received on ice, the laboratory shall record one of the following at the time of receipt:

(a) The temperature of an actual sample.

(b) The temperature of a temperature blank shipped with the samples.

(c) The temperature of the melt water in the shipping container.

(4) INCORPORATION BY REFERENCE. The materials in this section are incorporated by reference for the purposes of the permit program under ch. 283, Stats.

Note: Copies of the publications referenced in Tables A–H are available for inspection at the offices of the department of natural resources and the legislative reference bureau. Many of these materials are also available through inter–library loan.

History: Cr. Register, June, 1986, No. 366, eff. 7–1–86; r. and recr. Tables B and E, Register, April, 1988, No. 388, eff. 5–1–88; am; r. and recr. Tables A to F, Register, November, 1992, No. 443, eff. 12–1–92; am. (1), am. Tables A to F, Register, April, 1994, No. 460, eff. 5–1–94; am. (1) and (2), Tables A to F, cr. (3), Register, February,

1996, No. 482, eff. 3–1–96; CR 02–019: am. Table B Register October 2002 No. 562, eff. 11–1–02; CR 04–033: r. and recr. Table A, Table B, Table BM, Table C, Table E, Table E, Table EM, and Table F, cr. Table ES Register November 2004 No. 587, eff. 12–1–04; CR 04–101: am. Table A Note 29 Register May 2005 No. 593, eff. 6–1–05; CR 08–076: am. (2) and (3) (intro.), cr. (4), r. and recr. Table A, B, C to EM and F Register May 2009 No. 641, eff. 6–1–09; correction to Table B Parameter No. 41 made under s. 13.92 (4) (b) 7., Stats., Register September 2009 No. 645; correction to Table B Parameter No. 15 made under s. 13.92 (4) (b) 7., Stats., Register April 2010 No. 652; CR 13–112: am. (1) and (2), r. and recr. Table A, B, r. Table BM, r. and recr. Table C to F, cr. Tables G and H, Register May 2015 No. 713, eff. 6–1–15; CR 19–014: r. and recr. Table A and Notes, am. Table EM, r. and recr. Table EM Notes 8, 11, cr. Table EM Notes 16 to 23, r. and recr. Table H and Notes Register April 2020 No. 772, eff. 5–1–20; correction in (2) made under s. 35.17, Stats., Register July 2020 No. 775.

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Parameter and units	Analytical Technology ¹	EPA	Standard Methods 25,26	AOAC, ASTM, USGS	Other
	В	acteria			
1. Coliform (fecal), number per 100 mL or number per gram dry weight	Most Probable Number (MPN), 5 tube, 3 dilution, or	p. 132 ³ 1680 ^{11,15} 1681 ^{11,20}	9221 E-2014		
	Membrane filter (MF), ^{2,5} single step	p. 124 ³	9222 D-2015 ²⁹	B-0050-854	
2. Coliform (fecal), number per 100 mL	MPN, 5 tube, 3 dilution, or	p. 132 ³	9221 E-2014; 9221 F.2-2014 ³³		
	Multiple tube/multiple well, or				Colilert-18 ^{®13,18,28}
	MF, ^{2,5} single step ⁵	p. 124 ³	9222 D-2015 ²⁹		
3. Coliform (total), number per 100 mL	MPN, 5 tube, 3 dilution, or	p. 114 ³	9221 B-2014		
	MF, ^{2,5} single step or two step	p. 108 ³	9222 B-2015 ³⁰	B-0025-854	
	MF ^{2,5} with enrichment	p. 111 ³	9222 (B+B.4e) -2015 ³⁰		
4. E. coli, number per 100 mL	MPN ^{6,8,16} multiple tube, or		9221 B.3–2014/9221 F–2014 ^{12,14,33}		
	Multiple tube/multiple well, or		9223 B-2016 ¹³	991.15 ¹⁰	Colilert ^{® 13,18} Colilert–18 ^{® 13,17,18}
	MF,2,5,6,7,8 two step, or		9222 B-2015/ 9222 I-2015 ³¹		
	Single step	1603 ²¹			m-ColiBlue24®19
5. Fecal streptococci, number per 100 mL	MPN, 5 tube, 3 dilution, or	p. 139 ³	9230 B-2013		
-	MF ² , or	p. 136 ³	9230 C-2013 ³²	B-0055-854	
	Plate count	p. 143 ³			
6. Enterococci, number per 100 mL	MPN, 5 tube, 3 dilution, or	p. 139 ³	9230 B-2013		
	MPN, ^{6,8} multiple tube/multiple well, or		9230 D-2013	D6503-99 ⁹	Enterolert ^{® 13,23}
	MF ^{2,5,6,7,8} single step or	1600 ²⁴	9230 C-2013 ³²		
	Plate count	p. 143 ³			
7. Salmonella number per gram dry weight ¹¹	MPN multiple tube	1682 ²²			
	1	tic Toxicity			
8. Toxicity, acute, fresh water organisms, percent effluent	Daphnia, <i>Ceriodaphnia dubia</i> , 48–h static–renewal mortality				Note 27
	Fathead Minnow, <i>Pimephales promelas</i> , 96–h static–renewal mortality, or 96–h flow–through mortality				Note 27
9. Toxicity, chronic, fresh water organisms, percent effluent	Daphnia, <i>Ceriodaphnia dubia</i> , survival and reproduction				Note 27
	Fathead minnow, <i>Pimephales promelas</i> , larval survival and growth				Note 27

Table A
List of Approved Biological Methods for Wastewater and Sewage Sludge

¹ The method must be specified when results are reported.

 2 A 0.45- μ m 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. U.S. 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 Because the MF technique usually yields low and variable recovery from chlorinated wastewaters, the Most Probable Number method will be required to resolve any controversies.

⁶ 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 (ATP) guidelines.

⁹ Annual Book of ASTM Standards–Water and Environmental Technology, Section 11.02. 2000, 1999, 1996. ASTM International.

¹⁰ Official Methods of Analysis of AOAC International. 16th Edition, 4th Revision, 1998. AOAC International.

¹¹ Recommended for enumeration of target organism in sewage sludge.

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- ¹² The 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 seasonal basis.
- ¹³ These tests are collectively known as defined enzyme substrate tests.
- ¹⁴ After 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 50 µg/mL of MUG may be used.
- 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.
- ¹⁶ Samples shall be enumerated by the multiple-tube or multiple-well procedure. Using multiple-tube procedures, employ an appropriate tube and dilution configuration 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[®], Quanti-Tray[®]/2000 and the MPN calculated from the table provided by the manufacturer.
- ¹⁷ 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 Colilert[®] test and is recommended for marine water samples.
- ¹⁸ Descriptions of the Colilert[®], Colilert–18[®], Quanti–Tray[®], and Quanti–Tray[®]/2000 may be obtained from IDEXX Laboratories, Inc.
- ¹⁹ A description of the mColiBlue24[®] 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: Escherichia coli (E. coli) in Water by Membrane Filtration Using Modified Membrane–Thermotolerant Escherichia coli Agar (modified mTEC), EPA–821–R–14–010. September 2014. 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.
- ²³ A description of the Enterolert[®] test may be obtained from IDEXX Laboratories Inc.
- ²⁴ Method 1600: Enterococci in Water by Membrane Filtration Using membrane–Enterococcus Indoxyl–β–D–Glucoside Agar (mEI), EPA–821–R–14–011. September 2014. U.S. EPA.
- ²⁵ Standard Methods for the Examination of Water and Wastewater, Joint Editorial Board, American Public Health Association, American Water Works Association, and Water Pollution Control Federation, 23rd Edition (2017), 22nd Edition (2012), 21st Edition (2005), 20th Edition (1998), 19th Edition (1995), and 18th Edition (1992).
- ²⁶ Standard Methods for the Analysis of Water and Wastewater. With the promulgation of Federal Register /Vol. 77, No. 97 / Friday, May 18, 2012, the EPA lists only the most recently EPA–approved version of a Standard Method (regardless of the printed or online edition) in 40 CFR Part 136, with few exceptions, to identify the method with the year of Standard Methods approval or adoption designated by the last four digits in the method number (e.g., Standard Method 3113B–2004). This approach clearly identifies the version of the standard method approved under Part 136 and no longer ties it to a particular compendium printing or edition of Standard Methods. Methods can be purchased at www.standardmethods.org.
- ²⁷ Compliance monitoring must be performed in accordance with the specifications in the "State of Wisconsin Aquatic Life Toxicity Testing Methods Manual, 2nd Edition," Wisconsin Department of Natural Resources, 2004. This publication is available for inspection at the offices of the Department of Natural Resources and the Legislative Reference Bureau. Copies are available from the Department of Natural Resources, Bureau of Science Services, P.O. Box 7921, Madison, WI 53707.
- ²⁸ To use Colilert–18[®] to assay for fecal coliforms, the incubation temperature is 44.5 ± 0.2 °C, and a water bath incubator is used.
- ²⁹ 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.
- 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 Examination 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.*"
- ³³ 9221 F. 2–2014: This procedure allows for simultaneous detection of *E. coli* and thermotolerant coliforms by adding inverted vials to EC–MUG; the inverted vials collect gas produced by thermotolerant coliforms.

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Parameter Units	Analytical Technology ⁵⁸	EPA 52	Standard methods	ASTM	USGS AOAC Other
1. Acidity, as	Electrometric endpoint or		2310 B-1997	D1067-06	I-1020-85 ²
CaCO ₃ , mg/L	phenolphthalein endpoint				
2. Alkalinity, as CaCO ₃ , mg/L	Electrometric or Colorimetric titration to pH 4.5, Manual		2320 B-1997	D1067-06	I-1030-85 ² 973.43 ³
	Automatic	310.2 (Rev. 1974) ¹			I-2030-85 ²
3. Aluminum— Fotal, ⁴ mg/L	Digestion, ⁴ followed by any of the fol- lowing:				
	AA direct aspiration (FLAA) ³⁶		3111 D–1999 or 3111 E–1999		I-3051-85 ²
	Graphite furnace AA (GFAA)		3113 B-2004		
	Stabilized temperature graphite furnace AA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP-AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999		I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct Current Plasma (DCP) 36			D4190-08	Note ³⁴
	Colorimetric (Eriochrome cyanine R)		3500–A1 B–2001		
. 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– 1997		973.49 ³
	Titration		4500–NH ₃ C– 1997		
	Electrode		4500–NH ₃ D– 1997 or E–1997	D1426-08 (B)	
	Manual phenate, salicylate, or other substituted phenols in Berthelot reac- tion based methods		4500-NH ₃ F- 1997		Note ⁶⁰
	Automated phenate, salicylate, or other substituted phenols in Berthelot reaction based methods	350.1, ³⁰ Rev. 2.0 (1993)	4500–NH ₃ G– 1997 4500–NH ₃ H– 1997.		I-4523-85 ²
	Automated electrode				Note ⁷
	Ion Chromatography			D6919-09	
. Anti- nony—Total, ⁴ mg/	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999		
	Graphite furnace AA (GFAA)		3113 B-2004		
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I–4471–97 ⁵⁰
					2

200.8, Rev. 5.4

(1994)

3125 B-2009 D5673-05

Inductively coupled plasma- mass

spectrometry (ICP/MS)

Table B
List of Approved Inorganic Test Procedures For Wastewater

993.14,³ I–4471–97 ⁵⁰

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Parameter Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
6. Arsenic–Total, ⁴	Digestion, ⁴ followed by any of the	206.5 (Issued	methous	ASTM	Ouler
mg/L	following:	1978) ¹			
	AA gaseous hydride		3114 B–2009 or	D2972-08 (B)	I-3062-85 ²
			3114 C-2009		
	Graphite furnace AA (GFAA)		3113 B-2004	D2972-08 (C)	I-4063-98 49
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I-4020-05 ⁷⁰
	Colorimetric (SDDC)		3500–As B–1997	D2972-08 (A)	I-3060-85 ²
7. Barium–Total, ⁴ mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 D-1999	D 1000 00 00	I-3084-85 ²
	Graphite furnace AA (GFAA)	200 5 B 4 2	3113 B-2004	D4382-02(07)	1 4471 07 50
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003) ⁶⁸ ; 200.7, Rev. 4.4 (1994)	3120 B-1999		I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP) 36				Note ³⁴
8. Beryl- lium—Total, ⁴ mg/ L	Digestion, ⁴ followed by any of the following: AA direct aspiration (FLAA)		3111 D-1999	D3645-08 (A)	I-3095-85. ²
			or 3111 E–1999		
	Graphite furnace AA (GFAA)		3113 B-2004	D3645-08 (B)	
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma– atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I–4471–97 ⁵⁰
	Inductively coupled plasma– mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP)			D4190-08	Note ³⁴
	Colorimetric (Aluminon)		Note ⁶¹		
9. Biochemical oxygen demand (B0 mg/L	Dissolved Oxygen Depletion OD5),		5210 B-2001		973.44, ³ p. 17, ⁹ I–1578–78 ⁸ Notes ^{10,63}
10. Boron—Total, ³⁷ m- g/L	Colorimetric (Curcumin)		4500-В В -2000		I-3112-85 ²
	Inductively coupled plasma– atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma– mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I-4471-97 ⁵⁰

(1994)

Table B
List of Approved Inorganic Test Procedures For Wastewater

mass spectrometry (ICP/MS)

I-4471-97 ⁵⁰

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Direct current plasma (DCP)			D4190-08	Note ³⁴
11. Bromide, mg/L	Titrimetric				I–1125–85 ²
	Ion selective electrode (ISE)			D1246-05	
	Ion Chromatography	300.0, Rev 2.1 (1993) 300.1–1, Rev 1.0 (1997)	4110 B-2000, C-2000, D-2000	D4327-03	993.30 ³
	CIE/UV		4140 B-1997	D6508-00(05)	D6508, Rev. 2 54
12. Cad- mium—Total, ⁴ mg/ L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999 or 3111 C-1999	D3557–02(07) (A or B)	974.27, ³ p. 37, ⁹ I–3135–85 ² or
			5111 € 1777		I-3136-85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D3557-02(07) (D)	I-4138-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I–1472–85 ² or I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP) ³⁶			D4190-08	Note ³⁴
	Voltametry ¹¹			D3557-02(07) (C)	
	Colorimetric (Dithizone)		3500-Cd-D- 1990		
13. Cal- cium—Total, ⁴ mg/ L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999	D511-08(B)	I-3152-85 ²
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999		I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Direct current plasma (DCP)				Note ³⁴
	Titrimetric (EDTA)		3500–Ca B–1997	D511-08 (A)	
	Ion Chromatography			D6919-09	
14. Carbonaceous biochemical oxygen demand (CBOD ₅), mg/L ¹²	Dissolved Oxygen Depletion with nitrification inhibitor		5210 B-2001		Note ^{35,63}
15. Chemical oxygen demand (COD), mg/L	Titrimetric	410.3 (Rev. 1978) ¹	5220 B-1997 or C-1997	D1252-06 (A)	I-3560-85, ² 973.46, ³ p. 17 ⁹
	Spectrophotometric, manual or	410.4, Rev. 2.0	5220 D-1997	D1252-06 (B)	Note ^{13,14} I-3561-85. ²

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Parameter Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Titrimetric: (silver nitrate)	EIA	4500-Cl ⁻ B-1997	D512-04 (B)	I-1183-85 ²
	Colorimetric: manual				I–1187–85 ²
	Colorimetric, Automated (Ferricyanide)		4500–Cl [–] E–1997		I-2187-85 ²
	Potentiometric Titration		4500–Cl [–] D–1997		
	Ion Selective Electrode			D512-04 (C)	
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0	4110 B-2000 or 4110 C-2000	D4327-03	993.30, ³ I–2057–90 ⁵¹
	Capillary ion electrophoresis (CIE/UV)	(1997)	4140 B-1997	D6508-00(05)	D6508, Rev. 2 54
17. Chlorine–Total residual, mg/L	Amperometric direct		4500–Cl D–2000	D1253-08	
	Amperometric direct (low level)		4500–Cl E–2000		
	Iodometric direct		4500–Cl B–2000		
	Back titration ether end-point ¹⁵		4500–C1 C–2000		
	Colorimetric, DPD-FAS		4500–Cl F–2000		
	Spectrophotometric, DPD		4500–C1 G–2000		
	Ion selective electrode (ISE)				Note ¹⁶
17A. Chlorine– Free Available, mg/L	Amperometric direct		4500–C1 D–2000	D1253-08	
	Amperometric direct (low level)		4500–Cl E–2000		
	DPD-FAS		4500–Cl F–2000		
	Spectrophotometric, DPD		4500–C1 G–2000		
18. Chromium VI dissolved, mg/L	0.45-micron Filtration followed by any of the following:				
	AA chelation-extraction		3111 C-1999		I-1232-85 ²
	Ion Chromatography	218.6, Rev. 3.3 (1994)	3500–Cr C–2009	D5257-03	993.23
	Colorimetric (Diphenyl-carbazide)		3500–Cr B–2009	D1687–02(07) (A)	I-1230-85 ²
19. Chromium—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999	D1687-02(07) (B)	974.27, ³ I–3236–85 ²
	AA chelation-extraction		3111 C-1999		

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
,	Graphite furnace AA (GFAA)		3113 B-2004	D1687–02(07) (C)	I-3233-93 ⁴⁶
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003), ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	Direct current plasma (DCP) 36			D4190-08	Note ³⁴
	Colorimetric (Diphenyl-carbazide)		3500–Cr B–2009		
20. Cobalt—Total, ⁴ mg /L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999 or 3111 C-1999	D3558-08 (A or B)	p. 37, ⁹ I–3239–85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D3558-08 (C)	I-4243-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	Direct current plasma (DCP)			D4190-08	Note ³⁴
21. Color, platinum cobalt units or dominant wave- length, hue, luminance purity	Colorimetric (ADMI)				Note ¹⁸
	Colorimetric (Platinum cobalt)		2120 B-2001		I-1250-85 ²
22. Cop- per—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B–1999 or	D1688–07 (A or B)	974.27, ³ p. 37, ⁹
			3111 C-1999		I-3270-85 ² or I-3271-85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D1688-07 (C)	I-4274-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	Direct current plasma (DCP) 36			D4190-08	Note ³⁴
	Colorimetric (Neocuproine)		3500–Cu B–1999		
	Colorimetric (Bathocuproine)		3500–Cu C–1999		Note ¹⁹

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
23. Cyanide—Total, mg/L	Automated UV digestion/distillation and Colorimetry				Kelada–01. ⁵⁵
-	Segmented Flow Injection, In–Line Ultraviolet Digestion, followed by gas diffusion amperometry			D7511-09	
	Manual distillation with MgCl ₂ , followed by any of the following:	335.4, Rev. 1.0 (1993) ⁵⁷	4500–CN [–] B–1999 or C–1999	D2036–09(A), D7284–08	10-204-00-1-X ⁵⁶
	Flow Injection, gas diffusion ampero- metry			D2036-09(A) D7284-08	
	Titrimetric		4500-CN ⁻ D-1999	D2036-09(A)	p. 22 ⁹
	Colorimetry; Spectrophotometric, manual		4500-CN ⁻ E-1999	D2036-09(A)	I-3300-85 ²
	Colorimetry; Semi-Automated ²⁰	335.4, Rev. 1.0 (1993) ⁵⁷			10-204-00-1-X, ⁵⁶ I-4302-85 ²
	Ion Chromatography			D2036-09(A)	
	Ion Selective Electrode		4500-CN ⁻ F-1999	D2036-09(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-1999	D2036-09(B)	
	Flow injection and ligand exchange, followed by gas diffusion amperometry ⁵⁹			D6888-09	OIA-1677-09 ⁴⁴
	Automated Distillation and Colorime- try (no UV digestion)				Kelada-01 55
24.A Cyanide– Free, mg/L	Flow Injection, followed by gas diffusion amperometry			D7237-10	OIA-1677-09 ⁴⁴
	Manual micro-diffusion and colori- metry			D4282-02	
25. Fluo- ride—Total, mg/L	Manual distillation, ⁶ followed by any of the following:		4500–F [–] B–19 97		
	Electrode, manual (ISE)		4500-F ⁻ C-19 97	D1179-04 (B)	
	Electrode, automated (ISE)				I-4327-85 ²
	Colorimetric, (SPADNS)		4500-F ⁻ D-19 97	D1179-04 (A)	
	Automated complexone		4500-F ⁻ E-19 97		
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	4110 B-2000 or C-2000	D4327-03	993.30 ³
	Capillary ion electrophoresis (CIE/ UV)		4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
26. Gold—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		
	Graphite furnace AA (GFAA)	231.2 (Issued 1978) ¹	3113 B-2004		

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Parameter. Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
Turumeter, emits	Inductively coupled plasma–mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Direct current plasma (DCP)				Note ³⁴
27. Hard- ness—Total, as CaCO ₃ , mg/L	Automated colorimetric	130.1 (Issued 1971) ¹			
	Titrimetric (EDTA)		2340 C-1997	D1126-02(07)	973.52B, ³ I–1338–85 ²
	Ca plus Mg as their carbonates, by inductively coupled plasma or AA direct aspiration. (See Parameters 13 and 33).		2340 B-1997		
28. Hydrogen ion (pH), pH units	Electrometric measurement		4500-H ⁺ B-20 00	D1293–99 (A or B)	973.41, ³ I–1586–85 ²
	Automated electrode	150.2 (Dec. 1982) ¹			See footnote, ²¹ I–2587–85 ²
29. Irid- ium—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		
	Graphite furnace AA (GFAA)	235.2 (Issued 1978) ¹			
	Inductively coupled plasma-mass spectrometry (ICP/MS)		3125 B-2009		
30. Iron—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B–1999 or 3111 C–1999	D1068–05 (A or B)	974.27, ³ I–3381–85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D1068-05 (C)	
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003) ⁶⁸ ; 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14. ³
	Direct current plasma (DCP) ³⁶			D4190-08	Note ³⁴
	Colorimetric (Phenanthroline)		3500–Fe B–1997	D1068-05 (D)	Note ²²
31. Kjeldahl Nitro- gen ⁵ —Total, (as N), mg/L	Manual digestion ²⁰ and distillation or gas diffusion, followed by any of the following:		4500-N _{org} - B-1997 or C-1997 and 4500-NH ₃ B-1 997	D3590-02(06) (A)	I-4515-91 ⁴⁵
	Titration		4500-NH ₃ C-1 997		973.48 ³
	Electrode		4500–NH ₃ D– 1997 or E–1997	D1426-08 (B)	
	Semi-automated phenate	350.1 Rev 2.0 1993	4500–NH ₃ G–1997, 4500–NH ₃ H–1997		

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Manual phenate, salicylate, or other substituted phenols in Berthelot reac- tion based methods		4500–NH ₃ F–1 997		Note ⁶⁰
Automated Metho	ds for TKN that do not require manua	al distillation			
	Automated phenate, salicylate, or other substituted phenols in Berthelot reaction based methods colorimetric (auto digestion and distillation)	351.1 (Rev. 1978) ¹			I–4551–78. ⁸
	Semi-automated block digestor colo- rimetric (distillation not required)	351.2, Rev. 2.0 (1993)	4500–N _{org} - D–1997	D3590-02(06) (B)	I-4515-91 ⁴⁵
	Block digester, followed by Auto dis- tillation and Titration				Note ³⁹
	Block Digester, followed by Flow injection gas diffusion (distillation not required)				Note ⁴¹
32. Lead—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999 or 3111 C-1999.	D3559–08 (A or B)	974.27, ³ I–3399–85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D3559-08 (D)	I-4403-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP) 36			D4190-08	Note ³⁴
	Voltametry ¹¹			D3559-08 (C)	
	Colorimetric (Dithizone)		3500–Pb B–1997		
33. Magne- sium—Total, ⁴ mg/ L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999	D511-08 (B)	974.27, ³ I–3447–85 ²
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Direct current plasma (DCP)				Note ³⁴
	Ion Chromatography			D6919-09	
34. Man- ganese—Total ⁴ , mg/L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999	D858–07 (A or B)	974.27, ³ I–3454–85 ²
	Graphite furnace AA (GFAA)		3113 B-2004	D858-07 (C)	
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I–4471–97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I-4471-97 ⁵⁰
	Direct current plasma (DCP) 36			D4190-08	Note ³⁴
	Colorimetric (Persulfate)		3500–Mn B–1999		920.203 ³
	Colorimetric (Periodate)				Note ²³
35. Mer- cury—Total, ⁴ mg/L	Cold vapor, Manual	245.1, Rev. 3.0 (1994)	3112 B-2009	D3223-02(07)	977.22, ³ I–3462–85 ²
	Cold vapor, Automated	245.2 (Issued 1974) ¹			
	Cold vapor atomic fluorescence spec- trometry (CVAFS)	245.7 Rev. 2.0 (2005) ¹⁷			I-4464-01 ⁷¹
	Purge and Trap CVAFS	1631E ⁴³			
36. Molybde- num—Total, ⁴ mg/L	Digestion, ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 D-1999		I-3490-85 ²
	Graphite furnace AA (GFAA)		3113 B-2004		I-3492-96 47
	Inductively coupled plasma–atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP)				Note ³⁴
37. Nickel—Total, ⁴ mg /L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B–1999 or	D1886–08 (A or B)	I-3499-85 ²
			3111 C-1999		
	Graphite furnace AA (GFAA)	200.0. P 2.2	3113 B-2004	D1886-08 (C)	I-4503-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	Direct current plasma (DCP) 36			D4190-08	Note ³⁴
38. Nitrate (as N), mg/L	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	4110 B-2000 or C-2000	D4327-03	993.30 ³
	Capillary ion electrophoresis (CIE/ UV)		4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
	Ion Selective Electrode		4500–NO ₃ -D- 2000	-	

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Parameter. Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
1	Nitrate–nitrite N minus Nitrite N (See parameters 39 and 40)		literious		Note ⁶²
39. Nitrate + nitrite (as N), mg/L	Cadmium reduction, Manual		4500–NO ₃ -E– 2000	D3867-04 (B)	
	Cadmium reduction, Automated	353.2, Rev. 2.0 (1993)	4500-NO ₃ -F- 2000	D3867-04 (A)	I-2545-90 ⁵¹
	Automated hydrazine		4500–NO ₃ -H– 2000		
	Reduction/Colorimetric				Note ⁶²
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	4110 B-2000 or C-2000	D4327-03	993.30 ³
	Capillary ion electrophoresis (CIE/ UV)		4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
40. Nitrite (as N), mg/L	Spectrophotometric: Manual		4500–NO ₂ -B– 2000		Note ²⁵
	Automated (Diazotization)				I-4540-85, ² Note ⁶²
	Automated (*bypass cadmium reduction)	353.2, Rev. 2.0 (1993)	4500-NO ₃ -F- 2000	D3867-04 (A)	I-4545-85 ²
	Manual (*bypass cadmium reduction)		4500–NO ₃ -E– 2000	D3867-04 (B)	
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	4110 B-2000 or C-2000	D4327-03	993.30 ³
	Capillary ion electrophoresis (CIE/ UV)	(1))))	4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
41. Oil and grease—Total recoverable, mg/L	Hexane extractable material (HEM): n–Hexane extraction and gravimetry	1664 Rev. A; 1664 Rev. B ⁴²	5520 B-2001 38		
	Silica gel treated HEM (SGT-HEM): Silica gel treatment and gravimetry	1664 Rev. A; 1664 Rev. B ⁴²	5520 B-2001 ³⁸ and 5520 F-2001 ³⁸		
42. Organic car- bon—Total (TOC), mg/L	Combustion		5310 B-2000	D7573-09	973.47, ³ p. 14 ²⁴
	Heated persulfate or UV persulfate oxidation		5310 C- 2000 5310 D-2000	D4839-03	973.47, ³ p. 14 ²⁴
43. Organic nitro- gen (as N), mg/L	Total Kjeldahl N (Parameter 31) minus ammonia N (Parameter 4)				
44. Ortho-phos- phate (as P), mg/L	Colorimetry, Ascorbic acid, Auto- mated	365.1, Rev. 2.0 (1993)	4500–P F–1999 or G–1999		973.56, ³ I–4601–85 ²
	Colorimetry, Ascorbic Acid, Manual single reagent		4500-Р Е-1999	D515-88(A)	973.55 ³
	Colorimetry, Ascorbic Acid, Manual two reagent	365.3 (Issued 1978) ¹			
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	4110 B-2000 or C-2000	D4327-03	993.30 ³

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Parameter. Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Capillary ion electrophoresis (CIE/ UV)		4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
45. Osmium—Total ⁴ , mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 D-1999		
	Graphite furnace AA (GFAA)	252.2 (Issued 1978) ¹			
46. Oxygen, dis- solved, mg/L	Winkler (Azide modification)		4500–O B–2001, C–2001, D–2001, E–2001, F–2001	D888–09 (A)	973.45B, ³ I–1575–78 ⁸
	Electrode		4500–O G–2001	D888-09 (B)	I-1576-78 ⁸
	Luminescence Based Sensor			D888-09 (C)	Note ⁶³ Note ⁶⁴
47. Palla- dium—Total, ⁴ mg/ L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		
	Graphite furnace AA (GFAA)	253.2 ¹ (Issued 1978)			
	Inductively coupled plasma-mass spectrometry (ICP/MS)		3125 B-2009		
	Direct current plasma (DCP)				Note ³⁴
48. Phenols, mg/L	Manual distillation ²⁶ , followed by any of the following:	420.1 ¹ (Rev. 1978)	5530 B-2005	D1783-01	
	Colorimetric (4AAP) manual	420.1 ¹ (Rev. 1978)	5530 D-2005 ²⁷	D1783–01 (A or B)	
	Colorimetric (4AAP), Automated	420.4 Rev. 1.0 (1993)			
49. Phosphorus (elemental), mg/L	Gas-liquid chromatography				Note ²⁸
50. Phospho- rus—Total, mg/L	Digestion ²⁰ , followed by any of the following:		4500–P B(5)–1999		973.55 ³
	Colorimetric, Manual ascorbic acid	365.3 ¹ (Issued 1978)	4500-P E-1999	D515–88 (A)	
	Colorimetric, Automated ascorbic acid reduction	365.1 Rev. 2.0 (1993)	4500–P F–1999, G–1999, H–1999		973.56, ³ I–4600–85 ²
	Colorimetric, Semi–automated block digestor (TKP digestion)	365.4 ¹ (Issued 1974)		D515-88 (B)	I-4610-91 ⁴⁸
51. Plat- inum—Total, ⁴ mg/ L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		
	Graphite furnace AA (GFAA)	255.2 (Issued 1978) ¹			

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	List of Approved Inor	ganic Test Proc	edures For Wa	stewater	
Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
1 41 4110001, 0 1110	Inductively coupled plasma-mass spectrometry (ICP/MS)		3125 B-2009		0.00
	Direct current plasma (DCP)				Note ³⁴
52. Potas- sium—Total, ⁴ mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		973.53, ³ I–3630–85 ²
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.7, Rev. 4.4 (1994)	3120 B-1999		
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Flame photometric		3500-К В-1997		
	Electrode		3500-К С-1997		
	Ion Chromatography		0510 -	D6919-09	1 0750 070
53. Residue—Total, mg/L	Gravimetric, 103–105°C		2540 B-1997		I-3750-85 ²
54. Residue—fil- terable (TDS), mg/ L	Gravimetric, 180°C		2540 C-1997	D5907-03	I-1750-85 ²
55. Residue—non- filterable (TSS), mg/L	Gravimetric, 103–105°C post washing of residue		2540 D-1997	D5907-03	I-3765-85 ²
56. Residue—set- tleable, mg/L	Volumetric, (Imhoff cone), or gravimetric		2540 F-1997		
atile, mg/L	Gravimetric, 550°C	160.4 (Issued 1971) ¹	2540-Е-1997		I-3753-85 ²
58. Rhodium—Total, ⁴ mg/L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA), or		3111 B-1999		
	Graphite furnace AA (GFAA)	265.2 (Issued 1978) ¹			
	Inductively coupled plasma-mass spectrometry (ICP/MS)		3125 B-2009		
59. Ruthe- nium—Total, ⁴ mg/ L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA), or		3111 B-1999		
	Graphite furnace AA (GFAA)	267.2 ¹			
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8	3125 B-2009		
60. Sele- nium—Total, ⁴ mg/ L	Digestion ⁴ , followed by any of the following:				
	Graphite furnace AA (GFAA)		3113 B-2004	D3859–08 (B)	I-4668-98 ⁴⁹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			

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				stewater	USGS
Parameter. Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	AOAC Other
	Inductively coupled plasma-atomic	200.5, Rev 4.2	3120 B-1999	D1976-07	Guiti
	emission spectrometry (ICP/AES) ³⁶	(2003); ⁶⁸ 200.7, Rev. 4.4 (1994)			
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	AA gaseous hydride		3114 B–2009, or 3111 C–2009	D3859–08 (A)	I-3667-85 ²
61. Silica—Dis- solved, ³⁷ mg/L	0.45-micron filtration followed by any of the following:				
	Colorimetric, Manual		4500–SiO ₂ C– 1997	D859-05	I-1700-85 ²
	Colorimetric, Automated (Molybdo- silicate)		4500–SiO ₂ E– 1997 or F–1997		I-2700-85 ²
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999		I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
62. Silver—Total, ^{4,} ³¹ mg/L	Digestion ^{4, 29} , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B–1999 or		974.27, ³ p. 37, ⁹
			3111 C-1999		I-3720-85 ²
	Graphite furnace AA (GFAA)		3113 B-2004		I-4724-89 ⁵¹
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976–07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
	Direct current plasma (DCP)				Note ³⁴
63. Sodium—Total, ⁴ m- g/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		973.54 ³ , I–3735–85 ²
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003) ⁶⁸ ; 200.7, Rev. 4.4 (1994)	3120 B-1999		I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Direct current plasma (DCP)				Note ³⁴
	Flame photometric		3500–Na B–1997		
	Ion Chromatography			D6919-09	
64. Specific con- ductance, microm- hos/cm at 25 °C	Wheatstone bridge	120.1 ¹ (Rev. 1982)	2510 B-1997	D1125–95(99) (A)	973.40, ³ I–2781–85 ²
65. Sulfate (as SO ₄), mg/L	Colorimetric, Automated	375.2, Rev. 2.0 (1993)	4500–SO ₄ ^{2–} F –1997 or G–1997		

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Parameter. Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Gravimetric		4500–SO ₄ ^{2–} C –1997 or D–1997		925.54 ³
	Turbidimetric		4500–SO ₄ ^{2–} E –1997	D516-07	
	Ion Chromatography	300.0, Rev 2.1 (1993) and 300.1–1, Rev 1.0 (1997)	or	D4327-03	993.30, ³ I–4020–05 ⁷⁰
	Capillary ion electrophoresis (CIE/ UV)		4140 B-1997	D6508-00(05)	D6508, Rev. 2 ⁵⁴
66. Sulfide (as S), mg/L	Sample Pretreatment		4500–S ^{2–} B, C–2000		
	Titrimetric (iodine)		4500-S ²⁻ F-20 00		I-3840-85 ²
	Colorimetric (methylene blue)		4500–S ^{2–} D–2 000		
	Ion Selective Electrode		4500–S ^{2–} G–2 000	D4658-08	
67. Sulfite (as SO ₃), mg/L	Titrimetric (iodine-iodate)		4500–SO ₃ ^{2–} B –2000		
68. Surfactants, mg/L	Colorimetric (methylene blue)		5540 C-2000	D2330-02	
69. Temperature, °C	Thermometric		2550 B-2000		Note ³²
70. Thallium–To- tal, ⁴ mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		
	Graphite furnace AA (GFAA)	279.2 ¹ (Issued 1978)	3113 B-2004		
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.7, Rev. 4.4 (1994); 200.5 Rev. 4.2 (2003) ⁶⁸	3120 B-1999	D1976–07	
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4471–97 ⁵⁰
71. Tin–Total, ⁴ mg/ L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 B-1999		I-3850-78 ⁸
	Graphite furnace AA (GFAA)		3113 B-2004		
	Stabilized temperature GFAA (STGFAA)	200.9, Rev. 2.2 (1994)			
	Inductively coupled plasma–atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)			
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
72. Titanium–To- tal, ⁴ mg/L	Digestion ⁴ followed by any of the following:				
	AA direct aspiration (FLAA)		3111 D-1999		

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Parameter, Units	Analytical Technology ⁵⁸	EPA ⁵²	Standard methods	ASTM	USGS AOAC Other
	Graphite furnace AA (GFAA)	283.2 ¹ (Issued 1978)	intendus		
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.7, Rev. 4.4 (1994)			
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³
	Direct current plasma (DCP)				Note ³⁴
73. Turbidity, NTU ⁵³	Nephelometric	180.1, Rev. 2.0 (1993)	2130 B-2001	D1889-00	I-3860-85 ² Note ⁶⁵ Note ⁶⁶ Note ⁶⁷
74. Vanadium–To- tal, ⁴ mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA)		3111 D-1999		
	Graphite furnace AA (GFAA)		3113 B-2004	D3373-03(07)	
	Inductively coupled plasma-atomic emission spectrometry (ICP/AES)	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14, ³ I–4020–05 ⁷⁰
	Direct current plasma (DCP)			D4190-08	Note ³⁴
	Colorimetric (Gallic Acid)		3500-V B-1997		
75. Zinc–Total ⁴ , mg/L	Digestion ⁴ , followed by any of the following:				
	AA direct aspiration (FLAA) ³⁶		3111 B-1999 or 3111 C-1999	D1691–02(07) (A or B)	974.27, ³ p. 37, ⁹ I–3900–85 ²
	Graphite furnace AA (GFAA)	289.2 ¹ (Issued 1978)			
	Inductively coupled plasma–atomic emission spectrometry (ICP/AES) ³⁶	200.5, Rev 4.2 (2003); ⁶⁸ 200.7, Rev. 4.4 (1994)	3120 B-1999	D1976-07	I-4471-97 ⁵⁰
	Inductively coupled plasma-mass spectrometry (ICP/MS)	200.8, Rev. 5.4 (1994)	3125 B-2009	D5673-05	993.14 ³ I-4020-05 ⁷⁰
	Direct current plasma (DCP) ³⁶			D4190-08	Note ³⁴
	Colorimetric (Zincon)		3500 Zn B-1997		Note ³³
76. Acid Mine Drainage		1627 ⁶⁹			

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1 Methods for Chemical Analysis of Water and Wastes, EPA-600/4-79-020. Revised March 1983 and 1979, where applicable. U.S. EPA. Available from: National Technical Information Service, 5285 Port Royal Road, Springfield, Virginia 22161.

2 Methods for 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 non-platform graphite furnace atomic absorption determinations a digestion using nitric acid (as specified in Section 4.1.3 of Methods for the Chemical Analysis of Water and Wastes) is required prior to analysis. The procedure used should subject the sample to gentle, acid refluxing and at no time should the sample be taken to dryness.

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For direct aspiration flame atomic absorption determinations (FLAA) a combination acid (nitric and hydrochloric acids) digestion is preferred prior to analysis. The approved total recoverable digestion is described as Method 200.2 in Supplement I of "Methods 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 methods, mercury by cold vapor atomic absorption, the noble metals and titanium by FLAA, a specific or modified sample digestion procedure may be required and in all cases the referenced method write–up should be consulted for specific instruction and/or cautions.

For analyses using inductively coupled plasma-atomic emission spectrometry (ICP-AES), the direct current plasma (DCP) technique or the EPA spectrochemical techniques (platform furnace AA, ICP-AES, and ICP-MS) use EPA Method 200.2 or an approved alternate procedure (e.g., CEM microwave digestion, which may be used with certain analytes as indicated in Table IB); the total recoverable digestion procedures in EPA Methods 200.7, 200.8, and 200.9 may be used for those respective methods. Regardless of the digestion procedure, the results of the analysis after digestion procedure are reported as "total" metals.

5 Copper sulfate or other catalysts that have been found suitable may be used in place of mercuric sulfate.

- 6 Manual distillation is not required if comparability data on representative effluent samples are on file to show that this preliminary distillation step is not necessary: however, manual distillation will be required to resolve any controversies. In general, the analytical method should be consulted regarding the need for distillation. If the method is not clear, the laboratory may compare a minimum of 9 different sample matrices to evaluate the need for distillation. For each matrix, a matrix spike and matrix spike duplicate are analyzed both with and without the distillation step. (A total of 36 samples, assuming 9 matrices). If results are comparable, the laboratory may dispense with the distillation step for future analysis. Comparable is defined as < 20% RPD for all tested matrices). Alternatively the two populations of spike recovery percentages may be compared using a recognized statistical test.
- 7 Industrial Method Number 379–75 WE Ammonia, Automated Electrode Method, Technicon Auto Analyzer II. February 19, 1976. Bran & Luebbe Analyzing Technologies Inc.
- 8 The approved method is that cited in Methods for Determination of Inorganic Substances in Water and Fluvial Sediments, Techniques of Water-Resources Investigations of the U.S. Geological Survey, Book 5, Chapter A1. 1979. USGS.
- 9 American National Standard on Photographic Processing Effluents. April 2, 1975. American National Standards Institute (ANSI), 25 West 43rd St., New York, NY 10036.
- 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 BOD₅test method which measures "total BOD." The addition of the nitrification inhibitor is not a procedural option, but must be included to report the CBOD₅parameter. A discharger whose permit requires reporting the traditional BOD₅may not use a nitrification inhibitor in the procedure for reporting the results. Only when a discharger's permit specifically states CBOD₅is required can the permittee report data using a nitrification inhibitor.
- ¹³ OIC Chemical Oxygen Demand Method. 1978. Oceanography International Corporation. 512 West Loop, P.O. Box 2980, College Station, TX 77840.
- ¹⁴ Method 8000, Chemical Oxygen Demand, Hach Handbook of Water Analysis, 1979. Hach Company. P.O. Box 389, Loveland, CO 80537. Available on–line at http://www.hach.com.
- ¹⁵ The back titration method will be used to resolve controversy.
- ¹⁶ Orion Research Instruction Manual, Residual Chlorine Electrode Model 97–70. Thermo Scientific, 81 Wyman Street, Waltham, MA 02454. 1977. Orion Research Incorporated. The calibration for the Orion residual chlorine method must be derived using at least three standard solutions, prepared from a 0.00281 N potassium iodate solution.
- 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., available from the U.S. EPA Sample Control Center (operated by CSC), 6101 Stevenson Avenue, Alexandria, VA 22304.
- ¹⁸ National Council of the Paper Industry for Air and Stream Improvement (NCASI) Technical Bulletin 253, December 1971.
- ¹⁹ Method 8506, Biocinchoninate Method for Copper, Hach Handbook of Water Analysis. 1979. Hach Company. P.O. Box 389, Loveland, CO 80537. Available on-line at http://www.hach.com.
- 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 & Luebbe Analyzing Technologies. Elmsford, NY 10523.
- ²² Method 8008, 1,10–Phenanthroline Method using FerroVer Iron Reagent for Water. 1980. Hach Company P.O. Box 389, Loveland, CO 80537. Available on–line at http://www.hach.com.
- ²³ Method 8034, Periodate Oxidation Method for Manganese, Hach Handbook of Wastewater Analysis. 1979. Hach Company Loveland, CO 80537. Available on-line at http://www.hach.com.
- 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) p. 14. 1987. USGS. Available from: U.S. Geological Survey, 604 S. Pickett Street, Alexandria, VA 22304.
- ²⁵ Method 8507, Nitrogen, Nitrite–Low Range, Diazotization Method for Water and Wastewater. 1979. Hach Company P.O. Box 389, Loveland, CO 80537. Available on–line at http://www.hach.com.
- 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 Chromatog-raphy*, 47(3):421–426. Available in most public libraries. Back volumes of the Journal of Chromatography are available from: Elsevier/North–Holland, Inc., Journal Information Centre, 52 Vanderbilt Avenue, New York, NY 10164.

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- 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 aqueous buffer of sodium thiosulfate and sodium hydroxide to pH of 12. Therefore, for levels of silver above 1 mg/L, 20 mL of sample should be diluted to 100 mL by adding 40 mL each of 2 M Na₂S₂O₃ and NaOH. Standards should be prepared in the same manner. For levels of silver below 1 mg/L the approved method is satisfactory.
- ³⁰ The use of EDTA decreases method sensitivity. Analysts 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>7 with NH₄OH. 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 Survey, Book 1, Chapter D1. 1975. USGS. Available from: U.S. Geological Survey, 604 S. Pickett Street, Alexandria, VA 22304.
- ³³ Method 8009, Zincon Method for Zinc, Hach Handbook of Water Analysis, 1979. Hach Company. Loveland, CO 80537. Available on–line at http://www.hach.com.
- 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. Available from: Thermo Scientific, 81 Wyman Street, Waltham, MA 02454.
- ³⁵ 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 for Determination of Metals. April 16, 1992. CEM Corporation, P.O. Box 200, Matthews, NC 28106–0200.
- ³⁷ When determining boron and silica, only plastic, PTFE, or quartz laboratory ware may be used from start until completion of analysis.
- ³⁸ Only 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 extraction solvents is prohibited.39 Method PAI–DK01, Nitrogen, Total Kjeldahl, Block Digestion, Steam Distillation, Titrimetric Detection. Revised December 22, 1994. OI Analytical/ALP– KEM, P.O. Box 9010, College Station, TX 77842.
- 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/ALP-KEM, P.O. Box 9010, College Station, TX 77842.
- 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; Oil and Grease) and Silica Gel Treated n–Hexane Extractable Material (SGT–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 (SGT–HEM; Non–polar Material) by Extraction and Grease) and Silica Gel Treated n–Hexane Extractable Material (SGT–HEM; Non–polar Material) by Extraction and Grease) and Silica Gel Treated n–Hexane Extractable Material (SGT–HEM; Non–polar Material) by Extraction and Grease) and Silica Gel Treated n–Hexane Extractable Material (SGT–HEM; Non–polar Material) by Extraction and Gravimetry. EPA–821–R–10–001. Available at NTIS, PB–121949, U.S. Department of Commerce, 5285 Port Royal, Springfield, VA 22161.
- ⁴³ Method 1631, 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. Available at NTIS, PB-121949, U.S. Department of Commerce, 5285 Port Royal, Springfield, Virginia 22161.
- ⁴⁴ Method OIA–1677–09, Available Cyanide by Ligand Exchange and Flow Injection Analysis (FIA). 2010. OI Analytical/ALPKEM, P.O. Box 9010, College Station, TX 77842.
- ⁴⁵ Open 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.
- ⁴⁶ Open 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.
- ⁴⁷ Open File Report 97–198, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Molybdenum by Graphite Furnace Atomic Absorption Spectrophotometry. 1997. USGS.
- ⁴⁸ 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 Digests 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 Constituents in Water and Fluvial Sediments. 1993. USGS.
- ⁵² Unless otherwise indicated, all EPA methods, excluding EPA Method 300.1–1, 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 US 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.
- ⁵⁴ Method D6508, Test Method for Determination of Dissolved Inorganic Anions in Aqueous Matrices Using Capillary Ion Electrophoresis and Chromate Electrolyte. December 2000. Waters Corp., 34 Maple St., Milford, MA, 01757, Telephone: 508/482–2131, Fax: 508/482–3625.
- ⁵⁵ Kelada–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. National Technical Information Service (NTIS), 5285 Port Royal Road, Springfield, VA 22161 [Order Number PB 2001–108275]. The toll free telephone number is: 800–553–6847.

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 acceptance criteria are met.

- 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. Available from Hach Company, P.O. Box 389, Loveland, CO 80537.
- ⁵⁷ When using sulfide removal test procedures described in EPA Method 335.4–1, 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 and/or distillation are required prior to analysis.
- ⁵⁹ Samples analyzed for available cyanide using OI Analytical method OIA–1677–09 or ASTM method D6888–09 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 in Berthelot Reaction ("phenol-hypochlorite reaction") colorimetric ammonium determination methods. For example when phenol is used 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. These reaction parameters increase to pH > 12.6 and 665 nm when salicylate is used as the color reagent—see, Krom, M.D. April 1980. The Analyst 105:305–316.
- ⁶¹ If atomic absorption or ICP instrumentation is not available, the aluminon colorimetric method detailed in the 19th Edition of *Standard Methods* 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 Table IB for measurement of biochemical oxygen demand (BOD) and carbonaceous biochemical oxygen demand (CBOD).
- ⁶⁴ In-Situ Method 1002-8-2009, Dissolved Oxygen (DO) Measurement by Optical Probe. 2009. In-Situ Incorporated.
- ⁶⁵ Mitchell Method M5331, Determination of Turbidity by Nephelometry. Revision 1.0, July 31, 2008. Leck Mitchell.
- 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/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 Coupled Plasma–Mass Spectrometry, Chapter 1, Section B, Methods of the National Water Quality Laboratory, Book 5, Laboratory Analysis, 2006. USGS.
- ⁷¹ Water–Resources Investigations Report 01–4132, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Organic Plus Inorganic Mercury in Filtered and Unfiltered Natural Water With Cold Vapor–Atomic Fluorescence Spectrometry, 2001. USGS.
- ⁷² Quality control requirements for low level mercury are found in s. NR 106.145 (9) and (10), Wis. Adm. Code. Low–level mercury methods are performance based so some method modifications are allowable, provided quality control requirements are met. If an atomic absorption detector is substituted for atomic fluorescence detector, the appropriate method citation is 245.1 (manual) or 245.2 (automated). If method 1631E is modified to eliminate the purge and trap step, the appropriate method citation is 245.7.

Table C List of Approved Test Procedures for Non–Pesticide Organic Compounds in Wastewater						
Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Other	
1. Acenaphthene	GC	610				
	GC/MS	625,1625B	6410 B-2000		Note, ⁹ p. 27	

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Parameter ¹	Analytical	EPA ^{2,7}	Standard	ASTM	Other
	Technology		methods		
	HPLC	610	6440 B-2000	D4657-92 (98)	
2. Acenaphthylene	GC	610			
	GC/MS	625,1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
3. Acrolein	GC	603			
	GC/MS	624 ⁴ ,1624B			
4. Acrylonitrile	GC	603			
	GC/MS	624 ⁴ , 1624B			
5. Anthracene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440B-2000	D4657-92 (98)	
6. Benzene	GC	602	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
7. Benzidine	Spectro- photometric				Note, ³ p.1
	GC/MS	625, ⁵ 1625B	6410 B-2000		
	HPLC	605			
8. Benzo(a)anthracene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
9. Benzo(a)pyrene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
10. Benzo(b)fluoranthene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
11. Benzo(g,h,i)perylene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
12. Benzo(k)fluoranthene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
13. Benzyl chloride	GC				Note, ³ p. 130
	GC/MS				Note, ⁶ p. S102
14. Butyl benzyl phthalate	GC	606			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
15. bis(2–Chloroethoxy) methane	GC	611			-
	GC/MS	625, 1625B	6410 B-2000		Note ⁹ , p. 27

,	Table C		
List of Approved Test Procedures for No	n–Pesticide Or	ganic Compounds in	Wast
Analytical	EPA ^{2,7}	Standard	Α

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Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Othe
16. bis(2–Chloroethyl) ether	GC	611			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
17. bis(2–Ethylhexyl) phthalate	GC	606			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
18. Bromodichloromethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
19. Bromoform	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
20. Bromomethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
21. 4–Bromophenyl phenyl ether	GC	611			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
22. Carbon tetrachloride	GC	601	6200 C-1997		Note, ³ p 130
	GC/MS	624, 1624B	6200 B-1997		150
23. 4–Chloro–3–methyl phenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
24. Chlorobenzene	GC	601, 602	6200 C-1997		Note, ³ p 130
	GC/MS	624, 1624B	6200 B-1997		
25. Chloroethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
26. 2–Chloroethylvinyl ether	GC	601			
	GC/MS	624, 1624B			
27. Chloroform	GC	601	6200 C-1997		Note, ³ p 130
	GC/MS	624, 1624B	6200 B-1997		
28. Chloromethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
29. 2–Chloronaphthalene	GC	612			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
30. 2–Chlorophenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
31. 4–Chlorophenyl phenyl ether	GC	611			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27

Table C
List of Approved Test Procedures for Non-Pesticide Organic Compounds in Wastewate

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Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Other
32. Chrysene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p
	HPLC	610	6440 B-2000	D4657-92 (98)	27
33. Dibenzo(a,h)anthracene	GC	610	0110 12 2000	21037 72 (70)	
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
34. Dibromochloromethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
35. 1,2–Dichlorobenzene	GC	601, 602	6200 C-1997		
	GC/MS	624, 1625B	6200 B-1997		Note, ⁹ p 27
36. 1,3–Dichlorobenzene	GC	601, 602	6200 C-1997		27
	GC/MS	624, 1625B	6200 B-1997		Note, ⁹ p 27
37. 1,4–Dichlorobenzene	GC	601, 602	6200 C-1997.		27
	GC/MS	624, 1625B	6200 B-1997		Note, ⁹ p 27
38. 3,3'–Dichlorobenzidine	GC/MS	625, 1625B	6410 B-2000		
	HPLC	605			
39. Dichlorodifluoromethane	GC	601			
	GC/MS		6200 C-1997		
0. 1,1–Dichloroethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
1. 1,2–Dichloroethane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
2. 1,1–Dichloroethene	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
43. <i>trans</i> -1,2-Dichloroethene	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
14. 2,4–Dichlorophenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
45. 1,2–Dichloropropane	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
46. <i>cis</i> –1,3–Dichloropropene	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		

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Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Other
47. trans-1,3-Dichloropropene	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
48. Diethyl phthalate	GC	606			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
49. 2,4–Dimethylphenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
50. Dimethyl phthalate	GC	606			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
51. Di–n–butyl phthalate	GC	606			-
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
52. Di-n-octyl phthalate	GC	606			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
53. 2, 3–Dinitrophenol	GC	604	6420 B-2000]	Note, ⁹ p 27
	GC/MS	625, 1625B	6410 B-2000		
54. 2, 4–Dinitrophenol	GC	604	6420 B-2000		Note, ⁹ p 27
	GC/MS	625, 1625B	6410 B-2000		
55. 2, 6–Dinitrophenol	GC	604	6420 B-2000		Note, ⁹ p 27
	GC/MS	625, 1625B	6410 B-2000		
56. 2,3–Dinitrotoluene	GC	609			0
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
57. 2,4–Dinitrotoluene	GC	609			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
58. 2,6–Dinitrotoluene	GC	609			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
59. Epichlorohydrin	GC				Note, ³ p 130
	GC/MS]	Note, ⁶ p
60. Ethylbenzene	GC	602	6200 C-1997		S102
	COMS	604 1604D	6000 B 1007		
61. Fluoranthene	GC/MS GC	624, 1624B 610	6200 B-1997		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
	HPLC	610	6440 B-2000	D4657-92 (98)	

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Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Other
62. Fluorene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
63. 1,2,3,4,6,7,8–Heptachloro– dibenzofuran	HRGC/MS	1613B			
64. 1,2,3,4,7,8,9–Heptachloro– dibenzofuran	HRGC/MS	1613B			
65. 1,2,3,4,6,7,8– Heptachloro– dibenzo–p–dioxin	HRGC/MS	1613B			
66. Hexachlorobenzene	GC	612			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
67. Hexachlorobutadiene	GC	612			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
68. Hexachlorocyclopentadiene	GC	612			
	GC/MS	625, ⁵ 1625B	6410 B-2000		Note, ⁹ p 27
69. 1,2,3,4,7,8–Hexachloro– dibenzofuran	HR GC/MS	1613B			
70. 1,2,3,6,7,8–Hexachloro– dibenzofuran	HR GC/MS	1613B			
71. 1,2,3,7,8,9–Hexachloro– dibenzofuran	HR GC/MS	1613B			
72. 2,3,4,6,7,8–Hexachloro– dibenzofuran	HR GC/MS	1613B			
73. 1,2,3,4,7,8–Hexachloro– dibenzo–p–dioxin	HR GC/MS	1613B			
74. 1,2,3,6,7,8–Hexachloro– dibenzo–p–dioxin	HR GC/MS	1613B			
75. 1,2,3,7,8,9–Hexachloro– dibenzo–p–dioxin	HR GC/MS	1613B			
76. Hexachloroethane	GC	612			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
77. Indeno(1,2,3–c,d) pyrene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	21
78. Isophorone	GC	609			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
79. Methylene chloride	GC	601	6200 C-1997		Note, ³ p 130
	GC/MS	624, 1624B	6200 B-1997		
80. 2-Methyl-4,6-dinitrophenol	GC	604	6420 B-2000		

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			rganic Compounds i		<u> </u>
Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Othe
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
31. Naphthalene	GC	610			
	GC/MS	625, 1625B	6410 B-2000.		Note, ⁹ p
	HPLC	610	6440 B-2000.		27
82. Nitrobenzene	GC	609			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
	HPLC			D4657-92 (98))
83. 2–Nitrophenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
84. 4–Nitrophenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
85. N–Nitrosodimethylamine	GC	607			
	GC/MS	625, ⁵ 1625B	6410 B-2000		Note, ⁹ p 27
6. N–Nitrosodi–n–propylamine	GC	607			
	GC/MS	625, ⁵ 1625B	6410 B-2000		Note, ⁹ p 27
87. N–Nitrosodiphenylamine	GC	607			
	GC/MS	625, ⁵ 1625B	6410 B-2000		Note, ⁹ p 27
88. Octachlorodibenzofuran	HR GC/MS	1613B ¹⁰			
89. Octachlorodibenzo-p-dioxin	HR GC/MS	1613B ¹⁰			
90. 2,2'-Oxybis(2-chloro-propane) [prev.: bis(2-Chloroisopropyl) ether]	GC	611			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p 27
91. PCB–1016, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p 43; Note ⁸
	GC/MS	625	6410 B-2000		
	HR GC/MS	1668A ¹⁴			
92. PCB–1221, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p 43; Note ⁸
	GC/MS	625	6410 B-2000		
	HRGC/ MS	1668A ¹⁴			
93. PCB–1232, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p 43; Note ⁸
	GC/MS	625	6410 B-2000		
	HRGC/ MS	1668A ¹⁴			

Table C

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Parameter ¹	Analytical Technology	EPA ^{2,7}	Standard methods	ASTM	Other
94. PCB–1242, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p. 43; Note ⁸
	66046	(25	(410 D 2000		Note ⁶
	GC/MS HRGC/ MS	625 1668A ¹⁴	6410 B-2000		
95. PCB-1248, (Aroclor or	GC GC	608			
$(Albeloi of congeners)^{12,13}$		008			
	GC/MS	625	6410 B-2000		
	HRGC/ MS	1668A ¹⁴			
96. PCB–1254, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p. 43; Note ⁸
	GC/MS	625	6410 B-2000		
	HRGC/ MS	1668A ¹⁴			
97. PCB–1260, (Aroclor or congeners) ^{12,13}	GC	608			Note, ³ p. 43; Note ⁸
	GC/MS	625	6410 B-2000		
	HRGC/ MS	1668A ¹⁴			
98. 1,2,3,7,8–Pentachloro–dibenzofuran	GC/MS	1613B			
99. 2,3,4,7,8–Pentachloro–dibenzofuran	GC/MS	1613B			
00. 1,2,3,7,8,–Pentachloro–dibenzo–p– lioxin	GC/MS	1613B			
101. Pentachlorophenol	GC	604	6420 B-2000		Note, ³ p. 140
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
102. Phenanthrene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
03. Phenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
104. Pyrene	GC	610			
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
	HPLC	610	6440 B-2000	D4657-92 (98)	
105. 2,3,7,8–Tetrachloro–dibenzofuran	HR GC/MS	1613B ¹⁰	2.1.2 2000		
106. 2,3,7,8-Tetrachloro-dibenzo-p- lioxin	GC/MS	613, 625, ^{5a}			
107. 1,1,2,2–Tetrachloroethane	GC	601	6200 C-1997		Note, ³ p. 130
	GC/MS	624, 1624B	6200 B-1997		
08 Tetrachlorocatechol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
109. Tetrachloroethene	GC	601	6200 C-1997		Note, ³ p. 130
	GC/MS	624, 1624B	6200 B-1997		

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Parameter ¹	Analytical	EPA ^{2,7}	Standard	ASTM	Other
	Technology	LIA	methods	ASIM	Other
10. Tetrachloroguaicol	GC		6420 B-2000		
	GC/MS	1653 ¹¹	6410 B-2000		
111. 2,3,4,6– Tetrachlorophenol	GC		6420 B-2000		
	GC/MS		6410 B-2000		
112. Toluene	GC	602	6200 C-1997		
		(21.1(2))	(200 5 4005		
113. 1,2,4–Trichlorobenzene	GC/MS GC	624, 1624B	6200 B-1997		Nets 3 m
115. 1,2,4–1richlorobenzene	GC	612			Note, ³ p. 130
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
114. 3,4,5–Trichlorocatechol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
115. 3,4,6–Trichlorocatechol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
16. 1,1,1–Trichloroethane	GC	601	6200 C-1997		
	uc .	001	0200 C-1777		
	GC/MS	624, 1624B	6200 B-1997		
117. 1,1,2–Trichloroethane	GC	601	6200 C-1997		Note, ³ p. 130
	GC/MS	624, 1624B	6200 B-1997		
18. Trichloroethene	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
119. Trichlorofluoromethane	GC	601	6200 C-1997		
	GC/MS	624	6200 B-1997		
120. 3,4,5–Trichloroguaicol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
21. 3,4,6–Trichloroguaicol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
122. 4,5,6–Trichloroguaicol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
123. 2,4,5–Trichlorophenol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
124. 2,4,6–Trichlorophenol	GC	604	6420 B-2000		
	GC/MS	625, 1625B	6410 B-2000		Note, ⁹ p. 27
25. Trichlorosyringol	GC		6420 B-2000		
	GC/MS	165311	6410 B-2000		
26. Vinyl chloride	GC	601	6200 C-1997		
	GC/MS	624, 1624B	6200 B-1997		
127. Nonylphenol	GC/MS			D7065-06	
28. Bisphenol A (BPA)	GC/MS			D7065-06	
29. p-tert-Octylphenol (OP)	GC/MS			D7065-06	
30. Nonylphenol Monoethoxylate NP1EO)	GC/MS			D7065-06	
131. Nonylphenol Diethoxylate NP2EO)	GC/MS			D7065-06	
32. Adsorbable Organic Halides (AOX)	Adsorption and Coulometric Titra- tion	1650 11			
133. Chlorinated Phenolics	In Situ Acetylation	1653 11			

Table C

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- ¹All parameters are expressed in micrograms per liter (μ g/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, 624, 625, 1613B, 1624B, and 1625B are provided at Appendix A, Text Procedures for Analysis of Organic Pollutants, of 40 CFR Part 136. The standardized test procedure to be used to determine the method detection limit (MDL) for these test procedures is given at 40 CFR Part136, Appendix B, Definition and Procedure for the Determination of the Method Detection Limit.
- ³ Methods for Benzidine: Chlorinated Organic Compounds, Pentachlorophenol and Pesticides in Water and Wastewater. September 1978. U.S. EPA.
- ⁴ Method 624 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.
- ⁵ Method 625 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.
- ^{5a} Method 625, screening only.
- ⁶ 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).
- ⁷ Each analyst must make an initial, one-time demonstration of their ability to generate acceptable precision and accuracy with Methods 601–603, 624, 625, 1624B, and 1625B in accordance with procedures each 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 and 625 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 warning limits, the analytical results for that parameter in the unspiked sample are suspect. The results should be reported, but cannot be used to demonstrate regulatory compliance. These quality control requirements also apply to the Standard Methods, ASTM Methods, and other methods cited.
- ⁸ Organochlorine Pesticides and PCBs in Wastewater Using EmporeTMDisk. Revised October 28, 1994. 3M Corporation.
- ⁹ Method 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 40 CFR Part 136 and at http://water.epa.gov/scitech/methods/cwa/index.cfm.
- ¹¹ Method 1650, Adsorbable Organic Halides by Adsorption and Coulometric Titration. Revision C, 1997. U.S. EPA. Method 1653, Chlorinated Phenolics in Wastewater 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, "Methods 1650 and 1653", in Part 430, The Pulp, Paper, and Paperboard Point Source Category. Also available on–line at http://www.gpo.gov/.
- ¹² EPA Method 1668A may be used to test for all PCB congeners. If this method is employed, all PCB congeners shall be delineated. Non-detects shall be treated as zero. The values that are between the limit of detection and the limit of quantitation shall be used when calculating the total value of all congeners. All results shall be added together and the total PCB concentration reported. It is recognized a number of congeners will co-elute with others, so there will not be 209 results to sum.
- ¹³ If congener specific analysis is performed, the list of congeners tested shall include at least congener numbers 5, 18, 31, 44, 52, 66, 87, 101, 110, 138, 141, 151, 153, 170, 180, 183, 187, and 206 plus any other additional congeners which might be reasonably expected to occur in the particular sample. If Aroclor analysis is performed, clean up steps of the extract shall be performed as necessary to remove interference. If congener specific analysis is done, clean up steps of the extract shall be performed as necessary to remove interference. If desired limits of detection cannot be achieved after using the appropriate clean up techniques, a reporting limit that is achievable for the Aroclors or each congener for sample shall be determined. This report limit should be reported and qualified indicating the presence of an interference. The laboratory conducting the analysis shall perform as many the following methods as necessary to remove interference:

Florisil, Gel Permeation, Silica Gel, Alumina, Sulfur Clean Up, Sulfuric Acid Clean Up.

¹⁴ "Method 1668A, Revision A: Chlorinated Biphenyl Congeners in Water, Soil, Sediment, and Tissue by HRGC/HRMS", EPA-821-R-00-002, Environmental Protection Agency, Office of Water, Washington, D.C., December 1999. Available from: the National Technical Information Service, 5285 Port Royal Road, Springfield, Virginia 22161.

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D			Procedures for F		
Parameter	Analytical Technology	EPA ^{2,7,10}	Standard Methods	ASTM	Other
1. Aldrin	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96 (02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		
2. Ametryn	GC	507, 619			Note, ³ p. 83; Note, ⁹ O–3106–93; Note, ⁶ p. S68
	GC/MS	525.2			Note, ¹⁴ O-1121-91
3. Aminocarb	TLC				Note, ³ p. 94; Note, ⁶ p. S60
	HPLC	632			
4. Atraton	GC	619			Note, ³ p. 83; Note, ⁶ p. S68
5. Atrazine	GC	507, 619			Note, ³ p. 83; Note, ⁶ p. S68; Note, ⁹ O-3106-93
	HPLC/MS				Note, ¹² O-2060-01
	GC/MS	525.1, 525.2			Note, ¹¹ O-1126-95
6. Azinphos methyl	GC	614, 622, 1657			Note, ³ p. 25; Note, ⁶ p. S51
	GC-MS				Note, ¹¹ O-1126-95
7. Barban	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			
3. α–BHC	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁸ 3M0222
	GC/MS	625 ⁵	6410 B-2000		Note, ¹¹ O–1126–95
9. β–ВНС	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		
10. δ–BHC	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		
11. γ–BHC (Lindane)	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	625 ⁵	6410 B-2000		Note, ¹¹ O-1126-95
2. Captan	GC	617	6630 B-2000	D3086–90, D5812–96(02)	Note, ³ p. 7
13. Carbaryl	TLC				Note, ³ p. 94, Note, ⁶ p. S60
	HPLC	531.1, 632			
	HPLC/MS	553			Note, ¹² O-2060-01
	GC/MS				Note, ¹¹ O–1126–95
14. Carbophenothion	GC	617	6630 B-2000		Note, ⁴ page 27; Note, ⁶ p. S73
15. Chlordane	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
		(25	(410 D 2000		
	GC/MS	625	6410 B-2000		

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	List	Ta of Approved Test	able D Procedures for H	Pesticides ¹	
Parameter	Analytical Technology	EPA 2,7,10	Standard Methods	ASTM	Other
	HPLC	632			
17. 2,4–D	GC	615	6640 B-2001		Note, ³ p. 115; Note, ⁴ O–3105 –83
	HPLC/MS				Note, ¹² O-2060-01
18. 4,4'-DDD	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3105–83; Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000.		
19. 4,4'-DDE	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		Note, ¹¹ O-1126-95
20. 4,4'-DDT	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		
21. Demeton–O	GC	614, 622			Note, ³ p. 25; Note, ⁶ p. S51
22. Demeton–S	GC	614, 622			Note, ³ p. 25; Note, ⁶ p. S51
3. Diazinon	GC	507, 614, 622, 1657			Note ³ , p. 25; Note ⁴ , O–3104–83; Note ⁶ , p. S51
	GC/MS	525.2			Note ¹¹ , O-1126-95
24. Dicamba	GC	615			Note ³ , p. 115
	HPLC/MS				Note ¹² , O-2060-01
25. Dichlofenthion	GC	622.1			Note ⁴ , page 27; Note ⁶ , p. S73
26. Dichloran	GC	608.2, 617	6630 B-2000		Note ³ , p. 7
27. Dicofol	GC	617			Note ⁴ , O-3104-83
28. Dieldrin	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note ³ , p. 7; Note ⁴ , O–3104–83; Note ⁸ , 3M0222
	GC/MS	625	6410 B-2000		Note ¹¹ , O-1126-95
29. Dioxathion	GC	614.1, 1657			Note ⁴ , page 27; Note ⁶ , p. S73
30. Disulfoton	GC	507, 614, 622, 1657			Note ³ , p. 25; Note, ⁶ p. S51
	GC/MS	525.2			Note, ¹¹ O–1126–95
31. Diuron	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			12 0 2010 01
22 Endo16 1	HPLC/MS	553	((20 D 2000	D2086 00	Note, ¹² O–2060–01
32. Endosulfan I	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	625 ⁵	6410 B-2000		Note, ¹³ O-2002-01
33. Endosulfan II	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁸ 3M0222
	GC/MS	625 ⁵	6410 B-2000		Note, ¹³ O-2002-01
34. Endosulfan Sulfate	GC	608, 617	6630 C-2000		Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000		

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Parameter	Analytical	EPA ^{2,7,10}	Standard	ASTM	Other
	Technology		Methods		
5. Endrin	GC	505, 508, 608, 617,		D3086–90,	Note, 3 p. 7;
		1656	6630 C-2000	D5812-96(02)	Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	525.1, 525.2, 625 ⁵	6410 B-2000		
36. Endrin aldehyde	GC	608, 617	6630 C-2000		Note, ⁸ 3M0222
	GC/MS	625			
37. Ethion	GC	614, 614.1,1657			Note, ⁴ page 27; Note, ⁶ p. S73
	GC/MS				Note, ¹³ O-2002-01
38. Fenuron	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			· •
	HPLC/MS				Note, ¹² O-2060-01
39. Fenuron–TCA	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			-
40. Heptachlor	GC	505, 508, 608, 617, 1656	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁸ 3M0222
	GC/MS	525.1, 525.2, 625	6410 B-2000		
41. Heptachlor epoxide	GC	608, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83; Note, ⁶ p. S73; Note, ⁸ 3M0222
	GC/MS	625	6410 B-2000.		
42. Isodrin	GC	617	6630 B-2000 6630 C-2000		Note, ⁴ O–3104–83; Note, ⁶ p. S73
43. Linuron	GC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			
	HPLC/MS	553			Note, ¹² O–2060–01
	GC/MS				Note, ¹¹ O–1126–95
44. Malathion	GC	614, 1657	6630 B-2000		Note, ³ p. 25; Note, ⁶ p. S51
	GC/MS				Note, ¹¹ O-1126-95
45. Methiocarb	TLC				Note, ³ p. 94; Note, ⁶ p. S60
	HPLC	632			
	HPLC/MS				Note, ¹² O-2060-01
46. Methoxychlor	GC	505, 508, 608.2, 617, 1656	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104 –83 Note, ⁸ 3M0222
	GC/MS	525.1, 525.2			Note, ¹¹ O-1126-95
47. Mexacarbate	TLC				Note, ³ p. 94; Note, ⁶ p.S60
	HPLC	632			
48. Mirex	GC	617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3104–83
49. Monuron	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			-

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	List o	of Approved Test	ible D Procedures for I	Pesticides ¹	
Parameter	Analytical Technology	EPA ^{2,7,10}	Standard Methods	ASTM	Other
	HPLC	632			
51. Neburon	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			
	HPLC/MS				Note, ¹² O–2060–01
52. Parathion methyl	GC	614, 622, 1657	6630 B-2000		Note, ³ p. 25; Note, ⁴ page 27
	GC/MS				Note, ¹¹ O-1126-95
53. Parathion ethyl	GC	614	6630 B-2000		Note, ³ p. 25; Note, ⁴ page 27
	GC/MS				Note, ¹¹ O-1126-95
54. PCNB	GC	608.1, 617	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7.
55. Perthane	GC	617		D3086–90, D5812–96(02)	Note, ⁴ O-3104-83
56. Prometon	GC	507, 619			Note, ³ p. 83; Note, ⁶ p. S68; Note, ⁹ O-3106-93
	GC/MS	525.2			Note, ¹¹ O-1126-95
57. Prometryn	GC	507, 619			Note, ³ p. 83; Note, ⁶ p. S68; Note, ⁹ O-3106-93
	GC/MS	525.1, 525.2			Note, ¹³ O-2002-01
58. Propazine	GC	507, 619, 1656			Note, ³ p. 83; Note, ⁶ p. S68; Note, ⁹ O-3106-93
	GC/MS	525.1, 525.2.			
59. Propham	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			*
	HPLC/MS				Note, ¹² O-2060-01
60. Propoxur	TLC				Note, ³ p. 94; Note, ⁶ p. S60
	HPLC	632			*
61. Secbumeton	TLC				Note, ³ p. 83; Note, ⁶ p. S68
	GC	619			
62. Siduron	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			
	HPLC/MS				Note, ¹² O–2060–01
63. Simazine	GC	505, 507, 619, 1656			Note, ³ p. 83; Note, ⁶ p. S68; Note, ⁹ O–3106–93
	GC/MS	525.1, 525.2			Note, ¹¹ O-1126-95
64. Strobane	GC	617	6630 B-2000 6630 C-2000		Note, ³ p. 7
65. Swep	TLC				Note, ³ p. 104; Note, ⁶ p. S64
	HPLC	632			· •
66. 2,4,5–T	GC	615	6640 B-2001		Note, ³ p. 115; Note, ⁴ O–3105–83
67. 2,4,5–TP (Silvex)	GC	615	6640 B-2001		Note, ³ p. 115; Note, ⁴ O–3105–83

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	List o	of Approved Test Pr	ocedures for I	Pesticides ¹	
Parameter	Analytical Technology	EPA ^{2,7,10}	Standard Methods	ASTM	Other
68. Terbuthylazine	GC	619, 1656			Note, ³ p. 83; Note, ⁶ p. S68
	GC/MS				Note, ¹³ O-2002-01
59. Toxaphene GC	GC	505, 508, 608, 617, 1656	6630 B-2000 6630 C-2000	D3086–90, D5812–96(02)	Note, ³ p. 7; Note, ⁴ O–3105–83 Note, ⁸ 3M0222
	GC/MS	525.1, 525.2, 625	6410 B-2000		
70. Trifluralin	GC	508, 617, 627, 1656	6630 B-2000		Note, ³ p. 7; Note, ⁹ O–3106–93
	GC/MS	525.2			Note, ¹¹ O-1126-95

Table D
List of Annroved Test Procedures for Pesticides

¹ Pesticides are listed in this table by common name for the convenience of the reader. Additional pesticides may be found under Table C, 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 40 CFR Part 136, Appendix B, Definition and Procedure for the Determination of the Method Detection Limit.

³ Methods 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 Survey, Book 5, Chapter A3. 1987. USGS.
- ⁵ The method may be extended to include á–BHC, ã–BHC, endosulfan I, endosulfan II, and endrin. However, when they are known to exist, Method 608 is the preferred method.

⁶ 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).

⁷ Each analyst must make an initial, one-time, demonstration of their ability to generate acceptable precision and accuracy with Methods 608 and 625 in accordance 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 or 5% of all samples analyzed with Method 625 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 suspect. The results should be reported, but cannot be used to demonstrate regulatory compliance. These quality control requirements also apply to the Standard Methods, ASTM Methods, and other methods cited.

⁸ Organochlorine Pesticides and PCBs in Wastewater Using Empore TM Disk. Revised October 28, 1994. 3M Corporation.

- ⁹ Method O-3106-93 is in Open File Report 94-37, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of Triazine 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.

The full text of Methods 608 and 625 are provided at 40 CFR Part 136, Appendix A, Test Procedures for Analysis of Organic Pollutants.

EPA Methods 505, 507, 508, 525.1, 531.1 and 553 are in Methods for the Determination 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 Determination 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.

- ¹¹ Method 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 pesticides in water by C-18 solid-phase extraction and capillary-column gas chromatography/mass spectrometry with selected-ion monitoring. 1995. USGS.
- ¹² Method 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.
- ¹³ Method O-2002-01 is in Water-Resources Investigations Report 01-4098, Methods of Analysis by the U.S. Geological Survey National Water Quality Laboratory—Determination of moderate-use pesticides in water by C-18 solid-phase extraction and capillary-column gas chromatography/mass spectrometry. 2001. USGS.
- ¹⁴ Method O–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 with selected–ion monitoring. 1992. USGS.

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Table E List of Approved Radiological Analytical Methods for Wastewater						
			Reference (me	ethod number or page	e)	
Parameter and units	Method	EPA ¹	Standard Methods 18th, 19th, 20th Ed.	Standard Methods Online	ASTM	USGS ²
1. Alpha–Total, pCi per liter	Proportional or scintillation counter	900.0	7110 B	7110 B-00	D1943–90, 96	pp. 75 and 78 ³
2. Alpha–Counting error, pCi per liter	Proportional or scintillation counter	Appendix B	7110 B	7110 B-00	D1943-90, 96	p. 79
3. Beta–Total, pCi per liter	Proportional counter	900.0	7110 B	7110 B-00	D1890–90, 96	pp. 75 and 78 ³
4. Beta–Counting error, pCi	Proportional counter	Appendix B	7110 B	7110 B-00	D1890–90, 96	p. 79
5. (a) Radium Total pCi per liter	Proportional counter	903.0	7500–Ra B	7500–Ra B–01	D2460–90, 97	
(b) Radium, pCi per liter	Scintillation counter	903.1	7500–Ra C	7500-Ra C-01	D3454–91, 97	p. 81

¹ Prescribed Procedures for Measurement of Radioactivity in Drinking Water, EPA-600/4-80-032 (1980), U.S. Environmental Protection Agency, August 1980.

² Fishman, M. J. and Brown, Eugene, "Selected Methods of the U.S. Geological Survey of Analysis of Wastewaters," U.S. Geological Survey, Open–File Report 76–177 (1976).

³ The method found on p. 75 measures only the dissolved portion while the method on p. 78 measures only the suspended portion. Therefore, the two results must be added to obtain the "total."

		Sample —— Preparation ——		Determinative Method			
Parameter	Analytical Technology	SW-846 ¹	EPA ⁴	SW-846 ¹	EPA ^{2,3}	Standard Methods [ed.] ^{8,9}	Other
Metals							
Arsenic	Gaseous Hydride ⁵	7061A		7061A			
	Graphite Furnace Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B–99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		
Beryllium	Flame Atomic Absorption	3050B, 3051A	200.2	7000B		3111 D [18,19,21], 3111 D–99	
	Graphite Furnace Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B–99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		

Table EM List of Approved Analytical Methods for Sludge

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		Sample —— Preparation ——		Determinative			
Parameter	Analytical Technology	SW-846 ¹	EPA ⁴	SW-846 ¹	EPA 2,3	Standard Methods [ed.] ^{8,9}	Other
Cadmium	Flame Atomic Absorption	3050B, 3051A	200.2	7000B			
	Graphite Furnace Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B–99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B–99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		
Chromium	Flame Atomic Absorption	3050B, 3051A	200.2	7000B		3111 B [18,19,21], 3111 B-99	
	Graphite Furnace Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B–99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		
Copper	Flame Atomic Absorption	3050B, 3051A	200.2	7000B		3111 B or C [18,19·21], 3111 B–99 or C–99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B-99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		
Lead	Flame Atomic Absorption	3050B, 3051A	200.2	7000B			
	Graphite Furnace ⁶ Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B–99	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8		
Mercury	Cold Vapor Atomic Absorption	7471A, 7471B		7471A, 7471B			
	Cold vapor atomic Fluorescence Spectrometry	7474					
Molybdenum	Graphite Furnace ⁶ Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99	
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B-99	

Table EM List of Approved Analytical Methods for Sludge (Continued)

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		Samp —— Prepara		Determinative ——— Method ————				
Parameter	Analytical Technology	SW-846 ¹	EPA ⁴	SW-846 ¹	EPA ^{2,3}	Standard Methods [ed.] ^{8,9}	Other	
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8			
Nickel	Flame Atomic Absorption	3050B, 3051A	200.2	7000B		3111 B or C [18,19,21], 3111 B–99 or C–99		
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B-99		
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8			
Selenium	Gaseous Hydride ⁵	7741A		7741A				
	Graphite Furnace Atomic Absorption	3050B, 3051A	200.2	7010	200.9	3113 B [18,19,21], 3113 B-99		
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B-99		
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8			
Zinc	Flame Atomic Absorption	3050B, 3051A	200.2	7000B		3111 B or C [18,19,21], 3111 B–99 or C–99		
	Inductively Coupled Plasma Emission	3050B, 3051A	200.2	6010B, 6010C	200.7	3120 B [20,21], 3120 B-99		
	Inductively Coupled Plasma/ Mass Spectrometry	3050B, 3051A		6020A	200.8			
Organics Dioxins and Furans	Gas Chromatography/ Mass Spectrometry	8290A	1613B	8290A	1613B			
PCBs Aroclor or Congeners)	Gas Chromatography	3540B, 3540C, 3545A		8082, 8082A ¹²				
PCB (Congeners)	Gas Chromatography/ Mass Spectrometry	1668A ^{13,} 14,15			1668A ^{13, 14, 15}			
Biological								
Enteric Viruses	Centrifuge Concentration						D 4994-89 (02), ⁷ Appendix H ¹⁰	
Coliform (fecal), number per gram dry weight	Most Probable Number (MPN), 5 tube, 3 dilution, or		p. 132, ¹⁶ 1680, ^{11,17} 1681 ^{11,18}		p. 132, ¹⁶ 1680, ^{11,17} 1681 ^{11,18}	9221 E-2014	Appendix F ¹⁰	
-	Membrane filter (MF) ^{21,22} single step		p. 124 ¹⁶	1	p. 124 ¹⁶	9222 D-2015 ¹⁹		

Table EM List of Approved Analytical Methods for Sludge (Continued)

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		Sample ————————————————————————————————————		Determinative Method				
Parameter	Analytical Technology	SW-846 ¹	EPA ⁴	SW-846 ¹	EPA ^{2,3}	Standard Methods [ed.] ^{8,9}	Other	
Helminth ova	Density Gradient Flotation						Note ⁹ or Appendix I ⁹	
Specific Oxygen Uptake Rate	Respirometer					2710 B [18,19, 20,21], 2710 B-04	Appendix D.2. ¹⁰	
Salmonella number per gram dry weight ¹¹	MPN multiple tube		1682 ²⁰		1682 ²⁰			
Physical								
Solids	Gravimetric					2540 G [18,19,20,21], 2540 G–97		
Percent Volatiles Solids Reduction	Calculation						Appendix D.1. and 3 ¹⁰	

 Table EM

 List of Approved Analytical Methods for Sludge (Continued)

- ¹ "Test Methods for Evaluating Solid Waste", Physical/Chemical Methods," SW–846, Environmental Protection Agency, Office of Solid Waste and Emergency Response, 401 M Street, S.W., Washington D.C. 20460, September 1986 (Third edition), including July 1992 (Update I), September 1994 (Update II), August 1993 (Update IIA), January 1995 (Update IIB), December 1996 (Update III), April 1998 (Update IIIA), November 2004 (Update IIIB), February 2007 (Update IV) updates. Available from: The Superintendent of Documents, U.S. Government Printing Office, Room 190, Federal Building, P.O. Box 371954, Pittsburgh, PA 15250–7954. Available online at http://www.epa.gov/epaoswer/hazwaste/test/ sw846.htm.
- ² If an alternative digestion procedure is specified in the analytical method, the digestion in this table shall be used. In all cases, consult the analytical method for special requirements and cautions. SW–846 method 3051A is an acceptable alternate digestion procedure to SW–846 method 3050B.
- ³ "Methods for the Determination of Metals in Environmental Samples", EPA-600/4-91-010, Environmental Protection Agency, Environmental Monitoring Systems Laboratory, Cincinnati, OH 45268, June 1991. Available from: the National Technical Information Service (NTIS), 5258 Port Royal Road, Springfield, Virginia 22161.
- ⁴ "Sample Preparation Procedure for Spectrochemical Determination of Total Recoverable Elements", Method 200.2, Revision 2.8, Environmental Protection Agency, Environmental Monitoring Systems Laboratory, Cincinnati, OH 45268, 1994. Available from: the National Technical Information Service (NTIS), 5258 Port Royal Road, Springfield, Virginia 22161.
- ⁵ High levels of chromium, copper, mercury, silver, cobalt, or molybdenum may interfere with the analysis. Consult Method 3114, of "Standard Methods for the Examination of Water and Wastewater", 18th, 19th, 20th, or 21st edition, for more information.
- ⁶ Concentrations of lead in municipal sludge may exceed the working range of graphite furnace.
- ⁷ 1993 Annual Book of ASTM Standards, Section 11.02, Water and Environmental Technology", American Society for Testing and Materials, 1993, 1916 Race Street, Philadelphia, PA 19103. Available from: the American Society for Testing and Materials, 1916 Race Street, Philadelphia, PA 19103.
- ⁸ Standard Methods for the Examination of Water and Wastewater, Joint Editorial Board, American Public Health Association, American Water Works Association, and Water Pollution Control Federation, 23rd Edition (2017), 22nd Edition (2012), 21st Edition (2005), 20th Edition (1998), 19th Edition (1995), and 18th Edition (1992).
- ⁹ "Standard Methods for the Examination of Water and Wastewater", Joint Editorial Board, American Public Health Association, American Water Works Association, and Water Pollution Control Federation, 2006. On–line subscription service available at http://www.standardmethods.org.
- ¹⁰ "Occurrence of Pathogens in Distribution and Marketing Municipal Sludges", EPA 600/1–87–014, Environmental Protection Agency, 1987. Available from: the National Technical Information Service, order # PB 88–154273/AS, 5285 Port Royal Road, Springfield, Virginia 22161.
- ¹¹ Recommended for enumeration of target organism in sewage sludge.
- ¹² Analysts may use Fluid Management Systems, Inc. PowerPrep system in place of manual cleanup provided that the analysis meet the requirements of Method 1613B (as specified in Section 9 of the method) and permitting authorities.
- ¹³ EPA Method 1668A may be used to test for all PCB congeners. If this method is employed, all PCB congeners shall be delineated. Non-detects shall be treated as zero. The values that are between the limit of detection and the limit of quantitation shall be used when calculating the total value of all congeners. All results shall be added together and the total PCB concentration by dry weight reported. It is recognized that a number of the congeners will co-elute with others, so there will not be 209 results to sum.

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¹⁴ EPA Method 8082A shall be used for PCB–Aroclor analysis and may be used for congener specific analysis as well. If congener specific analysis is performed using Method 8082A, the list of congeners tested shall include at least congener numbers 5, 18, 31, 44, 52, 66, 87, 101, 110, 138, 141, 151, 153, 170, 180, 183, 187, and 206 plus any other additional congeners which might be reasonably expected to occur in the particular sample. For either type of analysis, the sample shall be extracted using Soxhlet extraction Method 3540C or Pressurized Fluid Extraction Method 3545A. If Aroclor analysis is performed using Method 8082A, clean up steps of the extract shall be performed as necessary to remove interference and achieve as close to a limit of detection of 0.11 mg/kg as possible. If congener specific analysis is done using Method 8082A, clean up steps of the extract shall be performed as necessary to remove interference and to achieve as close to a limit of detection of 0.003 mg/kg as possible for each congener. If the aforementioned limits of detection cannot be achieved after using the appropriate clean up techniques, a reporting limit that is achievable for the Aroclors or each congener for sample shall be determined. This report limit should be reported and qualified indicating the presence of an interference. The laboratory conducting the analysis shall perform as many the following methods as necessary to remove interference:

3620C - Florisil

- 3640A Gel Permeation
- 3630C Silica Gel
- 3611B Alumina
- 3660B Sulfur Clean Up
- 3665A Sulfuric Acid Clean Up.
- ¹⁵ "Method 1668A, Revision A: Chlorinated Biphenyl Congeners in Water, Soil, Sediment, and Tissue by HRGC/HRMS", EPA-821-R-00-002, Environmental Protection Agency, Office of Water, Washington, D.C., December 1999. Available from: the National Technical Information Service, 5285 Port Royal Road, Springfield, Virginia 22161.
- 16 Microbiological Methods for Monitoring the Environment, Water, and Wastes, EPA/600/8-78/017. 1978. U.S. EPA.
- 17 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.
- 18 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.
- 19 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.
- 20 Method 1682: Salmonella in Sewage Sludge (Biosolids) by Modified Semisolid Rappaport–Vassiliadis (MSRV) Medium, EPA–821–R–14–012. September 2014. U.S. EPA.
- 21 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.
- 22 Because the MF technique usually yields low and variable recovery from chlorinated wastewaters, the Most Probable Number method will be required to resolve any controversies.

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Table FS

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Pharmaceuticals pollutants	CAS registry No.	Analytical method number ^{1,2}
acetonitrile	75-05-8	1666 / 1671 / D3371 / D3695
n–amyl acetate	628-63-7	1666 / D3695
n-amyl alcohol	71-41-0	1666 / D3695
Benzene	71-43-2	D4763 / D3695 / 502.2 / 524.2
n-butyl-acetate	123-86-4	1666 / D3695
tert–butyl alcohol	75-65-0	1666
chlorobenzene	108-90-7	502.2 / 524.2
chloroform	67-66-3	502.2 / 524.2 / 551
o–dichlorobenzene	95-50-1	1625C / 502.2 / 524.2
1,2-dichloroethane	107-06-2	D3695 / 502.2 / 524.2
diethylamine	109-89-7	1666 / 1671
dimethyl sulfoxide	67-68-5	1666 / 1671
Ethanol	64-17-5	1666 / 1671 / D3695
ethyl acetate	141-78-6	1666 / D3695
n-heptane	142-82-5	1666 / D3695
n-hexane	110-54-3	1666 / D3695
isobutyraldehyde	78-84-2	1666 / 1667
isopropanol	67-63-0	1666 / D3695
isopropyl acetate	108-21-4	1666 / D3695
isopropyl ether	108-20-3	1666 / D3695
methanol	67-56-1	1666 / 1671 / D3695
Methyl Cellosolve Δ	109-86-4	1666 / 1671
methylene chloride	75-09-2	502.2 / 524.2
methyl formate	107-31-3	1666
4-methyl-2-pentanone (MIBK)	108-10-1	1624C / 1666 / D3695 / D4763 / 524.2
Phenol	108-95-2	D4763
n–propanol	71-23-8	1666 / 1671 / D3695
2-propanone (acetone)	67-64-1	D3695 / D4763 / 524.2
Tetrahydrofuran	109-99-9	1666 / 524.2
Toluene	108-88-3	D3695 / D4763 / 502.2 / 524.2
Triethlyamine	121-44-8	1666 / 1671
Xylenes	(see Note 3)	1624C / 1666

¹ For compounds that also appear in Table C, test methods listed in Table C may also be used.

² EPA Methods 1666, 1667, and 1671 listed in the table above are published in the compendium titled Analytical Methods for the Determination of Pollutants in Pharmaceutical Manufacturing Industry Wastewaters (EPA 821–B–98–016).

EPA Methods 502.2 and 524.2 have been incorporated by reference into 40 CFR 141.24 and are in Methods for the Determination of Organic Compounds in Drinking Water, EPA–600/4–88–039, December 1988, Revised, July 1991, and Methods for the Determination of Organic Compounds in Drinking Water–Supplement II, EPA–600/R–92–129, August 1992, respectively. These EPA test method compendia are available from the National Technical Information Service, NTIS PB91–231480 and PB92–207703, U.S. Department of Commerce, 5285 Port Royal Road, Springfield, Virginia 22161. The toll–free number is 800–553–6847. ASTM test methods D3371, D3695, and D4763 are available from the American Society for Testing and Materials, 100 Barr Harbor Drive, West Conshohocken, PA 19428–2959.

³ 1624C: m-xylene 108-38-3, o,p-xylene E-14095 (Not a CAS number; this is the number provided in the Environmental Monitoring Methods Index (EMMI) database.); 1666: m,p-xylene 136777-61-2, o-xylene 95-47-6

Table F Required Containers, Preservation Techniques, and Holding Times for wastewater			
Parameter Number/Name	Container ¹	Preservation ^{2,3}	Maximum Holding Time ⁴
Table A — Bacterial Tests			
1–5. Coliform, total, fecal and E. coli	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ^{22,23}
6. Fecal streptococci	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ²²
7. Enterococci	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ²²
8. Salmonella	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ²²
Table A — Aquatic Toxicity Tests			
9-12. Toxicity, acute and chronic	P, FP, G	Cool, ≤6°C ¹⁶	36 hours

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Parameter Number/Name	Container ¹	Preservation ^{2,3}	Maximum Holding Time ⁴
Table B — Inorganic Tests			
1. Acidity	P, FP, G	Cool, ≤6°C ¹⁸	14 days
2. Alkalinity	P, FP, G	Cool, ≤6°C ¹⁸	14 days
4. Ammonia	P, FP, G	Cool, ≤6°C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
9. Biochemical oxygen demand	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
11. Bromide	P, FP, G	None required	28 days
14. Biochemical oxygen demand, carbonaceous	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
15. Chemical oxygen demand	P, FP, G	Cool, ≤6°C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
16. Chloride	P, FP, G	None required	28 days
17. Chlorine, total residual	P, G	None required	Analyze within 15 minutes
21. Color	P, FP, G	Cool, $\leq 6^{\circ}C^{18}$	48 hours
23–24. Cyanide, total or available (or CATC) free	P, FP, G	Cool, ≤6°C, ¹⁸ NaOH to pH>10, ⁶ reducing agent if oxidizer present	14 days
25. Fluoride	Р	None required	28 days
27. Hardness	P, FP, G	HNO ₃ or H ₂ SO ₄ to pH<2	6 months
28. Hydrogen ion (pH)	P, FP, G	None required	Analyze within 15 minutes
31, 43. Kjeldahl and organic N	P, FP, G	Cool, $\leq 6^{\circ}$ C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
38. Nitrate	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
39. Nitrate – nitrite	P, FP, G	Cool, $\leq 6^{\circ}$ C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
40. Nitrite	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
41. Oil and grease	G	Cool, ≤6°C, ¹⁸ HCl or H ₂ SO ₄ to pH<2	28 days
42. Organic carbon	P, FP, G	Cool, ≤6°C, ¹⁸ HCl, H ₂ SO ₄ or H ₃ PO ₄ to pH<2	28 days
44. Orthophosphate	P, FP, G	Cool, to $\leq 6^{\circ}C^{18,24}$	Filter within 15 minutes; Analyze within 48 hours.
46. Oxygen, dissolved (Probe or Luminescence)	G, Bottle and top	None required	Analyze within 15 minutes
47. Oxygen, Dissolved Winkler	G, Bottle and top	Fix on site and store in dark	8 hours
48. Phenols	G	Cool, $\leq 6^{\circ}$ C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
49. Phosphorus (elemental)	G	Cool, ≤6°C ¹⁸	48 hours
50. Phosphorus, total	P, FP, G	Cool, $\leq 6^{\circ}$ C, ¹⁸ H ₂ SO ₄ to pH<2	28 days
53. Residue, total	P, FP, G	Cool, ≤6°C ¹⁸	7 days
54. Residue, Filterable (TDS)	P, FP, G	Cool, ≤6°C ¹⁸	7 days
55. Residue, Nonfilterable (TSS)	P, FP, G	Cool, ≤6°C ¹⁸	7 days
56. Residue, Settleable	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
57. Residue, Volatile	P, FP, G	Cool, ≤6°C ¹⁸	7 days
61. Silica	P or Quartz	Cool, $\leq 6^{\circ}C^{18}$	28 days
64. Specific conductance	P, FP, G	Cool, ≤6°C ¹⁸	28 days
65. Sulfate	P, FP, G	Cool, ≤6°C ¹⁸	28 days
66. Sulfide	P, FP, G	Cool, ≤6°C, ¹⁸ add zinc acetate plus sodium hydroxide to pH>9	7 days
67. Sulfite	P, FP, G	None required	Analyze within 15 minutes
68. Surfactants	P, FP, G	Cool, ≤6°C ¹⁸	48 hours
69. Temperature	P, FP, G	None required	Analyze
73. Turbidity	P, FP, G	Cool, ≤6°C ¹⁸	48 hours

Table F (Continued)
Required Containers, Preservation Techniques, and Holding Times for wastewater

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Required Container	s, Preservation Technique	s, and Holding Times for waste	
Parameter Number/Name	Container ¹	Preservation ^{2,3}	Maximum Holding Time ⁴
Table B — Metals ⁷			
10. Boron	P, FP, or Quartz	HNO ₃ to pH<2	6 months
18. Chromium VI	P, FP, G	Cool, $\leq 6^{\circ}$ C, ¹⁸ pH = 9.3 – 9.7 ²⁰	28 days
35. Mercury (CVAA)	P, FP, G	HNO ₃ to pH<2	28 days
35. Mercury (CVAFS)	FP, G; and FP-lined cap ¹⁷	5 mL/L 12N HCl or 5 mL/L BrCl ¹⁷	90 days ¹⁷
3, 5–8, 12, 13, 19, 20, 22, 26, 29, 30, 32–34, 36, 37, 45, 47, 51, 52, 58–60, 62, 63, 70–72, 74, 75. Metals, except boron, chromium VI, and mercury.	P, FP, G	HNO ₃ to pH<2, or at least 24 hours prior to analysis ¹⁹	6 months
Table C — Organic Tests ⁸			
3, 4. Acrolein and acrylonitrile	G, FP-lined septum	Cool, \leq 6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃ , pH to 4–5 ¹⁰	14 days ¹⁰
19. Adsorbable Organic Halides (AOX)	G	Cool, <6°C, 0.008% Na ₂ S ₂ O ₃ HNO ₃ to pH <2	Hold <i>at least</i> 3 days, but not more than 6 months
114–118. Alkylated phenols	G	Cool, <6°C, H ₂ SO ₄ to pH <2	28 days until extraction,40 days after extraction
7, 38. Benzidines ^{11, 12}	G, FP-lined cap	Cool, ≤6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃ ⁵	7 days until extraction ¹³
29, 35–37, 63–65, 107. Chlorinated hydrocarbons ¹¹	G, FP–lined cap	Cool, ≤6°C ¹⁸	7 days until extraction, 40 days after extraction
20. Chlorinated Phenolics		Cool, <6°C, 0.008% Na ₂ S ₂ O _{3,} H ₂ SO ₄ to pH <2	30 days until acetylation, 30 days after acetylation.
15, 16, 21, 31, 87. Haloethers ¹¹	G, FP-lined cap	Cool, ≤6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃ ⁵	7 days until extraction, 40 days after extraction
54, 55, 75, 79. Nitroaromatics and Isophorone ¹¹	G, FP-lined cap	Cool, ≤6°C, ¹⁸ store in dark, 0.008% Na ₂ S ₂ O ₃ ⁵	7 days until extraction, 40 days after extraction
82–84. Nitrosamines ^{11, 14}	G, FP-lined cap	Cool, ≤6°C, ¹⁸ store in dark, 0.008% Na ₂ S ₂ O ₃ ⁵	7 days until extraction, 40 days after extraction
88–94. PCBs ¹¹	G, FP-lined cap	Cool, ≤6°C ¹⁸	1 year until extraction, 1 year after extraction
60–62, 66–72, 85, 86, 95–97, 102, 103. PCDDs/PCDFs ¹¹			
Aqueous Samples: Field and Lab Preservation	G	Cool, ≤6°C ¹⁸ , 0.008% Na ₂ S ₂ O ₃ ⁵ , pH<9	1 year
Solids and Mixed–Phase Samples: Field Preservation	G	Cool, ≤6°C ¹⁸	7 days
Fissue Samples: Field Preservation	G	Cool, ≤6°C ¹⁸	24 hours
Solids, Mixed–Phase, and Tissue Samples: Lab Preservation	G	Freeze, $\leq -10^{\circ}$ C	1 year
23, 30, 44, 49, 53, 77, 80, 81, 98, 100, 112. Phenols ¹¹	G, FP-lined cap	Cool, ≤6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃	7 days until extraction, 40 days after extraction
14, 17, 48, 50–52. Phthalate esters ¹¹	G, FP-lined cap	Cool, ≤6°C ¹⁸	7 days until extraction, 40 days after extraction
1, 2, 5, 8–12, 32, 33, 58, 59, 74, 78, 99, 101. Polynuclear aromatic hydrocarbons ¹¹	G, FP-lined cap	Cool, $\leq 6^{\circ}$ C, ¹⁸ store in dark, 0.008% Na ₂ S ₂ O ₃ ⁵	7 days until extraction, 40 days after extraction
6, 57, 106. Purgeable aromatic hydrocarbons	G, FP-lined septum	Cool, ≤6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃ ⁵ , HCl to pH 2 ⁹	14 days ⁹
13, 18–20, 22, 24–28, 34–37, 39–43, 45–47, 56, 76, 104, 105, 108–111, 113. Purgeable halocarbons.	G, FP-lined septum	Cool, ≤6°C, ¹⁸ 0.008% Na ₂ S ₂ O ₃ ⁵	14 days

Table F (Continued)	
Required Containers, Preservation Techniques, and Holding Times for wastewater	

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Parameter Number/Name	Container ¹	Preservation ^{2,3}	Maximum Holding Time ⁴	
Table D — Pesticides Tests:				
1–70. Pesticides ¹¹	G, FP–lined cap	Cool, $\leq 6^{\circ}$ C, ¹⁸ pH 5–9 ¹⁵	7 days until extraction, 40 days after extraction	
Table E — Radiological Tests:				
1–5. Alpha, beta and radium	P, FP, G	HNO ₃ to pH<2	6 months	
Table H — Bacterial Tests:				
1. E. coli	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ²²	
2. Enterococci	PA, G	Cool, <10°C, 0.0008% Na ₂ S ₂ O ₃ ⁵	8 hours. ²²	
Table H — Protozoan Tests:				
8.Cryptosporidium	LDPE; field filtration	1–10 °C	96 hours. ²¹	
9.Giardia	LDPE; field filtration	1–10 °C	96 hours. ²¹	

Table F (Continued)
Required Containers, Preservation Techniques, and Holding Times for wastewater

¹ "P" is for polyethylene; "FP" is fluoropolymer (polytetrafluoroethylene (PTFE); Teflon[), or other fluoropolymer, unless stated otherwise in this Table F; "G" is glass; "PA" is any plastic that is made of a sterilizable material (polypropylene or other autoclavable plastic); "LDPE" is low density polyethylene.

² Except where noted in this table and the method for the parameter, preserve each grab sample within 15 minutes of collection. For a composite sample collected with an automated sample (e.g., using a 24-hour composite sampler, refrigerate the sample at ≤ 6 °C during collection unless specified otherwise in this table or in the method(s). For a composite sample to be split into separate aliquots for preservation and/or analysis, maintain the sample at ≤ 6 °C, unless specified otherwise in this table or in the method(s), until collection, splitting, and preservation is completed. Add the preservative to the sample container prior to sample collection when the preservative will not compromise the integrity of a grab sample, a composite sample, or aliquot split from a composite sample within 15 minutes of collection.

The temperature of the samples shall be documented upon receipt at the laboratory. If the samples are shipped in crushed or cube ice (not "blue ice" packs) and solid ice is still present in the cooler, the lab may simply report the samples as "received on ice". If the ice has melted, the lab must report the either the temperature of the melt– water or of a temperature blank. A temperature blank is defined as an aliquot of deionized water, in an appropriate sample container, which is transported along with the samples. Since shipping simply with "blue ice" packs does not insure that samples are maintained at the appropriate temperatures, the sample collector must submit a temperature blank when using these ice packs for shipping.

- ³ When any sample is to be shipped by common carrier or sent via the U.S. Postal Service, it must comply with the Department of Transportation Hazardous Materials Regulations (49 CFR part 172). The person offering such material for transportation is responsible for ensuring such compliance. For the preservation requirement, the Office of Hazardous Materials, Materials Transportation Bureau, Department of Transportation has determined that the Hazardous Materials Regulations do not apply to the following materials: Hydrochloric acid (HCl) in water solutions at concentrations of 0.04% by weight or less (pH about 1.96 or greater; Nitric acid (HNO₃) in water solutions at concentrations of 0.15% by weight or less (pH about 1.62 or greater); Sulfuric acid (H₂SO₄) in water solutions at concentrations of 0.35% by weight or less (pH about 1.15 or greater); and Sodium hydroxide (NaOH) in water solutions at concentrations of 0.080% by weight or less (pH about 12.30 or less).
- ⁴ Samples should be analyzed as soon as possible after collection. The times listed are the maximum times that samples may be held before the start of analysis and still be considered valid. Samples may be held for longer periods only if the permittee or monitoring laboratory has data on file to show that, for the specific types of samples under study, the analytes are stable for the longer time, and has received a variance from the EPA Regional Administrator under s. NR219.05). For a grab sample, the holding time begins at the time of collection. For a composite sample collected with an automated sampler (e.g., using a 24–hour composite sampler); the holding time begins at the time of the end of collection of the composite sample. For a set of grab samples composited in the field or laboratory, the holding time begins at the time of collection of the last grab sample in the set. Some samples may not be stable for the maximum time period given in the table. A permittee or monitoring laboratory is obligated to hold the sample for a shorter time if it knows that a shorter time is necessary to maintain sample stability. See 40 CFR 136.3(e) for details.
- ⁵ ASTM D7365–09a specifies treatment options for samples containing oxidants (e.g.,chlorine). Also, Section 9060A of Standard Methods for the Examination of Water and Wastewater (20th and 21st editions) addresses dechlorination procedures.
- ⁶ Sample collection and preservation: Collect a volume of sample appropriate to the analytical method in a bottle of the material specified. If the sample can be analyzed within 48 hours and sulfide is not present, adjust the pH to >12 with sodium hydroxide solution (e.g., 5% w/v), refrigerate as specified, and analyze within 48 hours. Otherwise, to extend the holding time to 14 days and mitigate interferences, treat the sample immediately using any or all of the following techniques, as necessary, followed by adjustment of the sample pH to >12 and refrigeration as specified.

There may be interferences that are not mitigated by approved procedures. Any procedure for removal or suppression of an interference may be employed, provided the laboratory demonstrates that it more accurately measures cyanide. Particulate cyanide (e.g., ferric ferrocyanide) or a strong cyanide complex (e.g., cobalt cyanide) are more accurately measured if the laboratory holds the sample at room temperature and pH >12 for a minimum of 4 hours prior to analysis, and performs UV digestion or dissolution under alkaline (pH=12) conditions, if necessary.

Sulfur: To remove elemental sulfur (S8), filter the sample immediately. If the filtration time will exceed 15 minutes, use a larger filter or a method that requires a smaller sample volume (e.g., EPA Method 335.4 or Lachat Method 01). Adjust the pH of the filtrate to >12 with NaOH,

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refrigerate the filter and filtrate, and ship or transport to the laboratory. In the laboratory, extract the filter with 100 mL of 5% NaOH solution for a minimum of 2 hours. Filter the extract and discard the solids. Combine the 5% NaOH–extracted filtrate with the initial filtrate, lower the pH to approximately 12 with concentrated hydrochloric or sulfuric acid, and analyze the combined filtrate. Because the detection limit for cyanide will be increased by dilution by the filtrate from the solids, test the sample with and without the solids procedure if a low detection limit for cyanide is necessary. Do not use the solids procedure if a higher cyanide concentration is obtained without it. Alternatively, analyze the filtrates from the sample and the solids separately, add the amounts determined (in ?g or mg), and divide by the original sample volume to obtain the cyanide concentration.

(1) Sulfide: If the sample contains sulfide as determined by lead acetate paper, or if sulfide is known or suspected to be present, immediately con- duct one of the volatilization treatments or the precipitation treatment as follows: Volatilization-Headspace expelling. In a fume hood or well-ventilated area, transfer 0.75 liter of sample to a 4.4-L collapsible container (e.g., CubitainerTM). Acidify with concentrated hydrochloric acid to pH <2. Cap the container and shake vigorously for 30 seconds. Remove the cap and expel the headspace into the fume hood or open area by collapsing the container without expelling the sample. Refill the headspace by expanding the container. Repeat expelling a total of five headspace volumes. Adjust the pH to >12, refrigerate, and ship or transport to the laboratory. Scaling to a smaller or larger sample volume must maintain the air to sample volume ratio. A larger volume of air will result in too great a loss of cyanide (> 10%). Dynamic stripping: In a fume hood or well ventilated area, transfer 0.75 liter of sample to a container of the material specified and acidify with concentrated hydrochloric acid to pH <2. Using a calibrated air sampling pump or flowmeter, purge the acidified sample into the fume hood or open area through a fritted glass aerator at a flow rate of 2.25 L/min for 4 minutes. Adjust the pH to >12, refrigerate, and ship or transport to the laboratory. Scaling to a smaller or larger sample volume must maintain the air to sample volume ratio. A larger volume of air will result in too great a loss of cyanide (>10%). Precipitation: If the sample contains particulate matter that would be removed by filtration, filter the sample prior to treatment to assure that cyanide associated with the particulate matter is included in the measurement. Ship or transport the filter to the laboratory. In the laboratory, extract the filter with 100 mL of 5% NaOH solution for a minimum of 2 hours. Filter the extract and discard the solids. Combine the 5% NaOH-extracted filtrate with the initial filtrate, lower the pH to approximately 12 with concentrated hydrochloric or sulfuric acid, and analyze the combined filtrate. Because the detection limit for cyanide will be increased by dilution by the filtrate from the solids, test the sample with and without the solids procedure if a low detection limit for cyanide is necessary. Do not use the solids procedure if a higher cyanide concentration is obtained without it. Alternatively, analyze the filtrates from the sample and the solids separately, add the amounts determined (in µg or mg), and divide by the original sample volume to obtain the cyanide concentration. For removal of sulfide by precipitation, raise the pH of the sample to >12 with NaOH solution, then add approximately 1 mg of powdered cadmium chloride for each mL of sample. For example, add approximately 500 mg to a 500-mL sample. Cap and shake the container to mix. Allow the precipitate to settle and test the sample with lead acetate paper. If necessary, add cadmium chloride but avoid adding an excess. Finally, filter through 0.45 micron filter.

Cool the sample as specified and ship or transport the filtrate and filter to the laboratory. In the laboratory, extract the filter with 100 mL of 5% NaOH solution for a minimum of 2 hours. Filter the extract and discard the solids. Combine the 5% NaOH–extracted filtrate with the initial filtrate, lower the pH to approximately 12 with concentrated hydrochloric or sulfuric acid, and analyze the combined filtrate. Because the detection limit for cyanide will be increased by dilution by the filtrate form the solids, test the sample with and without the solids procedure if a low detection limit for cyanide is necessary. Do not use the solids procedure if a higher cyanide concentration is obtained without it. Alternatively, analyze the filtrates from the sample and the solids separately, add the amounts determined (in g or mg), and divide by the original sample volume to obtain the cyanide concentration. If a ligand–exchange method is used (e.g., ASTM D6888), it may be necessary to increase the ligand exchange reagent to offset any excess of cadmium chloride.

(2) Sulfite, thiosulfate, or thiocyanate: If sulfite, thiosulfate, or thiocyanate is known or suspected to be present, use UV digestion with a glass coil (Method Kelada–01) or ligand exchange (Method OIA–1677) to preclude cyanide loss or positive interference.

(3) Aldehyde: If formaldehyde, acetaldehyde, or another water-soluble aldehyde is known or suspected to be present, treat the sample with 20 mL of 3.5% ethylenediamine solution per liter of sample.

(4) Carbonate: Carbonate interference is evidenced by noticeable effervescence upon acidification in the distillation flask, a reduction in the pH of the absorber solution, and incomplete cyanide spike recovery. When significant carbonate is present, adjust the pH to ≥ 12 using calcium hydroxide instead of sodium hydroxide. Allow the precipitate to settle and decant or filter the sample prior to analysis (also see Standard Method 4500–CN.B.3.d).

(5) Chlorine, hypochlorite, or other oxidant: Treat a sample known or suspected to contain chlorine, hypochlorite, or other oxidant as directed in footnote 5⁷ For dissolved metals, filter grab samples within 15 minutes of collection and before adding preservatives. For a composite sample collected with an automated sampler, filter the sample within 15 minutes after completion of collection and before adding preservatives. If it is known or suspected that dissolved sample integrity will be compromised during collection of a composite sample collected automatically over time (e.g., by interchange of a metal between dissolved and suspended forms), collect and filter grab samples to be composited (footnote 2) in place of a composite sample collected automatically.

⁷ For dissolved metals, filter grab samples within 15 minutes of collection and before adding preservatives. For a composite sample collected with an automated sampler, filter the sample within 15 minutes after completion of collection and before adding preservatives. If it is known or suspected that dissolved sample integrity will be compromised during collection of a composite sample collected automatically over time (e.g., by interchange of a metal between dissolved and suspended forms), collect and filter grab samples to be composited (footnote 2) in place of a composite sample collected automatically.

⁸ Guidance applies to samples to be analyzed by GC, LC, or GC/MS for specific compounds.

 9 If the sample is not adjusted to pH < 2, then the sample must be analyzed within seven days of sampling.

- ¹⁰ The pH adjustment is not required if acrolein will not be measured. Samples for acrolein receiving no pH adjustment must be analyzed within 3 days of sampling.
- ¹¹ When the extractable analytes of concern fall within a single chemical category, the specified preservative and maximum holding times should be observed for optimum safeguard of sample integrity (*i.e.*, use all necessary preservatives and hold for the shortest time listed). When the analytes of concern fall within two or more chemical categories, the sample may be preserved by cooling to ≤ 6 °C, reducing residual chlorine with 0.008% sodium thiosulfate, storing in the dark, and adjusting the pH to 6–9; samples preserved in this manner may be held for seven days before extraction and for forty days after extraction. Exceptions to this optional preservation and holding time procedure are noted in footnote 5 (regarding the requirement for thiosulfate reduction), and footnotes 12, 13 (regarding the analysis of benzidine).

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 12 If 1,2–diphenylhydrazine is likely to be present, adjust the pH of the sample to 4.0 ± 0.2 to prevent rearrangement to benzidine.

¹³ Extracts may be stored up to 30 days at < 0 °C.

¹⁴ For the analysis of diphenylnitrosamine, add 0.008% Na₂S₂O₃ and adjust pH to 7–10 with NaOH within 24 hours of sampling.

- ¹⁵ The pH adjustment may be performed upon receipt at the laboratory and may be omitted if the samples are extracted within 72 hours of collection. For the analysis of aldrin, add 0.008% Na₂S₂O₃.
- ¹⁶ Place sufficient ice with the samples in the shipping container to ensure that ice is still present when the samples arrive at the laboratory. However, even if ice is present when the samples arrive, immediately measure the temperature of the samples and confirm that the preservation temperature maximum has not been exceeded. In the isolated cases where it can be documented that this holding temperature cannot be met, the permittee can be given the option of on–site testing or can request a variance. The request for a variance should include supportive data which show that the toxicity of the effluent samples is not reduced because of the increased holding temperature. Aqueous samples must not be frozen. Hand–delivered samples used on the day of collection do not need to be cooled to 0 to 6 °C prior to test initiation.
- ¹⁷ Samples collected for the determination of trace level mercury (<100 ng/L) using EPA Method 1631 must be collected in tightly-capped fluoro-polymer or glass bottles and preserved with BrCl or HCl solution within 48 hours of sample collection. The time to preservation may be extended to 28 days if a sample is oxidized in the sample bottle. A sample collected for dissolved trace level mercury should be filtered in the laboratory within 24 hours of the time of collection. However, if circumstances preclude overnight shipment, the sample should be filtered in a designated clean area in the field in accordance with procedures given in Method 1669. If sample integrity will not be maintained by shipment to and filtration in the laboratory, the sample must be filtered in a designated clean area in the field within the time period necessary to maintain sample integrity. A sample that has been collected for determination of total or dissolved trace level mercury must be analyzed within 90 days of sample collection.</p>
- ¹⁸ Aqueous samples must be preserved at ≤6 °C, and should not be frozen unless data demonstrating that sample freezing does not adversely impact sample integrity is maintained on file and accepted as valid by the regulatory authority. Also, for purposes of NPDES monitoring, the specification of "≤6 °C" is used in place of the "4 °C" and "<4 °C" sample temperature requirements listed in some methods. It is not necessary to measure the sample temperature to three significant figures (1/100th of 1 degree); rather, three significant figures are specified so that rounding down to 6 °C may not be used to meet the ≤6 °C requirement. The preservation temperature does not apply to samples that are analyzed immediately (less than 15 minutes).</p>
- ¹⁹ An aqueous sample may be collected and shipped without acid preservation. However, acid must be added at least 24 hours before analysis to dissolve any metals that adsorb to the container walls. If the sample must be analyzed within 24 hours of collection, add the acid immediately (see footnote 2). Soil and sediment samples do not need to be preserved with acid. The allowances in this footnote supersede the preservation and holding time requirements in the approved metals methods.
- ²⁰ To achieve the 28-day holding time, use the ammonium sulfate buffer solution specified in EPA Method 218.6. The allowance in this footnote supersedes preservation and holding time requirements in the approved hexavalent chromium methods, unless this supersession would compromise the measurement, in which case requirements in the method must be followed.
- ²¹ Holding time is calculated from time of sample collection to elution for samples shipped to the laboratory in bulk and calculated from the time of sample filtration to elution for samples filtered in the field.
- ²² Sample analysis should begin as soon as possible after receipt; sample incubation must be started no later than 8 hours from time of collection.
- ²³ For fecal coliform samples for sewage sludge (biosolids) only, the holding time is extended to 24 hours for the following sample types using either EPA Method 1680 (LTB–EC) or 1681 (A–1): Class A composted, Class B aerobically digested, and Class B anaerobically digested.
- ²⁴ The immediate filtration requirement in orthophosphate measurement is to assess the dissolved or bio–available form of orthophosphorus (*i.e.*,that which passes through a 0.45–micron filter), hence the requirement to filter the sample immediately upon collection (*i.e.*,within 15 minutes of collection).

Test Methods for Pesticide Active Ingredients				
EPA survey code	Pesticide name	CAS No.	EPA analytical method No.(s) ³	
8	Triadimefon	43121-43-3	1656, 507, 633, 525.1, 525.2	
12	Dichlorvos	62-73-7	1657, 507, 525.1, 525.2, 622	
16	2,4–D; 2,4–D Salts and Esters [2,4–Dichloro–phenoxyacetic acid]	94–75–7	1658, 515.1, 515.2, 555, 615	
17	2,4–DB; 2,4–DB Salts and Esters [2,4–Dichlorophenoxybutyric acid]	94-82-6	1658, 515.1, 515.2, 555, 615	
22	Mevinphos	7786-34-7	1657, 507, 525.1, 525.2, 622	
25	Cyanazine	21725-46-2	507, 629	
26	Propachlor	1918-16-7	1656, 508, 608.1, 525.1, 525.2	
27	MCPA; MCPA Salts and Esters [2–Methyl–4–chlorophenoxyacetic acid]	94–74–6	1658, 555, 615	
30	Dichlorprop; Dichlorprop Salts and Esters [2–(2,4–Dichlorophenoxy) propionic acid]	120-36-5	1658, 515.1, 515.2, 555, 615	
31	MCPP; MCPP Salts and Esters [2–(2–Methyl–4–chlorophenoxy) propionic acid]	93-65-2	1658, 555, 615	

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ED4	Test Methods for Pesticide Active Ingredients					
EPA survey code	Pesticide name	CAS No.	EPA analytical method No.(s) ³			
35	ТСМТВ	21564-17-0	637			
	[2–(Thiocyanomethylthio) benzo–thiazole]					
39	Pronamide	23950-58-5	507, 525.1, 525.2, 633.1			
41	Propanil	709-98-8	1656, 632.1			
45	Metribuzin	21087-64-9	1656, 507, 525.1, 525.2, 633			
52	Acephate	30560-19-1	1656, 1657			
53	Acifluorfen	50594-66-6	515.1, 515.2, 555			
54	Alachlor	15972-60-8	1656, 505, 507, 525.1, 525.2, 645			
55	Aldicarb	116-06-3	531.1			
58	Ametryn	834-12-8	507, 525.2, 619			
50	Atrazine	1912-24-9	1656, 505, 507, 525.1, 525.2, 619			
52	Benomyl	17804-35-2	631			
58	Bromacil; Bromacil Salts and Esters	314-40-9	1656, 507, 525.1, 525.2, 633			
59 59	Bromoxynil	1689-84-5	1625, 1661			
59 59	Bromoxynil octanoate	1689-99-2	1656			
70	Butachlor	23184-66-9	1656, 507, 525.1, 525.2, 645			
73	Captafol	2425-06-1	1656			
75	Carbaryl [Sevin]	63-25-2	531.1, 553, 632			
76	Carbofuran	1563-66-2	531.1, 535, 652			
30	Chloroneb	2675-77-6	1656, 508, 525.1, 525.2, 608.1			
30 32	Chlorothalonil	1897-45-6	1656, 508, 525.1, 525.2, 608.2			
82 34						
	Stirofos	961-11-5	1657, 507, 525.1, 525.2, 622			
36	Chlorpyrifos	2921-88-2	1657, 508, 622			
90	Fenvalerate	51630-58-1	1660			
103	Diazinon	333-41-5	1657, 507, 525.2, 614, 622			
107	Parathion methyl	298-00-0	1657, 614, 622			
110	DCPA [Dimethyl 2,3,5,6-tetrachloro-terephthalate]	1861-32-1	1656, 508, 525.1, 525.2, 515.1 ² , 515.2 ² , 608.2			
112	Dinoseb	88-85-7	1658, 515.1, 515.2, 555, 615			
113	Dioxathion	78-34-2	1657, 614.1			
118	Nabonate [Disodium cyanodithioimidocarbonate]	138-93-2	630.1			
119	Diuron	330-54-1	553, 632			
123	Endothall	145-73-3	548, 548.1			
124	Endrin	72-20-8	1656, 505, 508, 525.1, 525.2, 608, 617			
125	Ethalfluralin	55283-68-6	1656, 627 See footnote 1			
126	Ethion	563-12-2	1657, 614, 614.1			
127	Ethoprop	13194-48-4	1657, 507, 525.1, 525.2, 622			
132	Fenarimol	60168-88-9	1656, 507, 525.1, 525.2, 633.1			
133	Fenthion	55-38-9	1657, 622			
138	Glyphosate [N-(Phosphonomethyl) glycine]	1071-83-6	547			
140	Heptachlor	76-44-8	1656, 505, 508, 525.1, 525.2, 608, 617			
144	Isopropalin	33820-53-0	1656, 627			
148	Linuron	330-55-2	553, 632			
150	Malathion	121-75-5	1657, 614			
154	Methamidophos	10265-92-6	1657			
156	Methomyl	16752-77-5	531.1, 632			
158	Methoxychlor	72-43-5	1656, 505, 508, 525.1, 525.2, 608.2, 617			
172	Nabam	142-59-6	630, 630.1			
172	Naled	300-76-5	1657, 622			
175	Norflurazon	27314-13-2	1656, 507, 525.1, 525.2, 645			
175	Benfluralin	1861-40-1	1656, 627 See footnote 1			

Table G	
Test Methods for Pesticide Active Ingredie	nt

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EPA	Pesticide name	CAS No.	redients EPA analytical method No.(s) ³	
survey code				
182	Fensulfothion	115-90-2	,	
183	Disulfoton	298-04-4	1657, 507, 525,2, 614, 622	
185	Phosmet	732-11-6	1657, 622.1	
186	Azinphos Methyl	86-50-0	1657, 614, 622	
192	Organo-tin pesticides	12379-54-3	200.7, 200.9, Ind-01	
197	Bolstar	35400-43-2	1657, 622	
203	Parathion	56-38-2	1657, 614	
204	Pendimethalin	40487-42-1	1656	
205	Pentachloronitrobenzene	82-68-8	1656, 608.1, 617	
206	Pentachlorophenol	87-86-5	1625, 515.2, 555, 515.1, 525.1, 525.2, 625	
208	Permethrin	52645-53-1	1656, 1660, 508, 525.1, 525.2, 608.2	
212	Phorate	298-02-2	1657, 622	
218	Busan 85 [Potassium dimethyldithiocarbamate]	128-03-0	630, 630.1	
219	Busan 40 [Potassium N-hydroxymethyl-N-methyldithiocarbamate]	51026-28-9	630, 630.1	
220	KN Methyl [Potassium N–methyl–dithiocarbamate]	137-41-7	630, 630.1	
223	Prometon	1610-18-0	507, 525.2, 619	
224	Prometryn	7287-19-6	507, 525.1, 525.2, 619	
226	Propazine	139-40-2	1656, 507, 525.1, 525.2, 619	
230	Pyrethrin I	121-21-1	1660	
232	Pyrethrin II	121-29-9	1660	
236	DEF [S,S,S–Tributyl phosphorotrithioate]	78-48-8	1657	
239	Simazine	122-34-9	1656, 505, 507, 525.1, 525.2, 619	
241	Carbam–S [Sodium dimethyldithio–carbamate]	128-04-1	630, 630.1	
243	Vapam [Sodium methyldithiocarbamate]	137-42-8	630, 630.1	
252	Tebuthiuron	34014-18-1	507, 525.1, 525.2	
254	Terbacil	5902-51-2	1656, 507, 525.1, 525.2, 633	
255	Terbufos	13071-79-9	1657, 507, 614.1, 525.1, 525.2	
256	Terbuthylazine	5915-41-3	1656, 619	
257	Terbutryn	886-50-0	507, 525.1, 525.2, 619	
259	Dazomet	533-74-4	1659, 630, 630.1	
262	Toxaphene	8001-35-2	1656, 505, 508, 525.1, 525.2, 608, 617	
263	Merphos [Tributyl phosphorotrithioate]	150-50-5	1657, 507, 525.1, 525.2, 622	
264	Trifluralin ¹	1582-09-8	1656, 508, 525.2, 617, 627	
268	Ziram [Zinc dimethyldithiocarbamate]	137-30-4	630. 630.1	

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Test Methods for Pesticide Active Ingredien

¹ Monitor and report as total Trifluralin.

² Applicable to the analysis of DCPA degradates.

³ EPA Methods 608.1 through 645, 1645 through 1661, and Ind-01 are available in Methods For The Determination of Nonconventional Pesticides In Municipal and Industrial Wastewater, Volume I, EPA 821-R-93-010A, Revision I, August 1993, U.S. EPA. EPA Methods 200.9 and 505 through 555 are available in Methods For The Determination of Nonconventional Pesticides In Municipal and Industrial Wastewater, Volume II, EPA 821-R-93-010B, August 1993, U.S. EPA. The full text of Methods 608, 625 and 1625 are provided at Appendix A of this Part 136. The full text of Method 200.7 is provided at appendix C of this part 136.

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	List of Approved M	0			0.4
Parameter and units	Method ¹	EPA	Standard Methods ³³	AOAC, ASTM, USGS	Other
Bacteria:					
1. Coliform (fecal), number per 100 mL	Most Probable Number (MPN), 5 tube, 3 dilu- tion, or	p. 132 ³	9221 E-2014, 9221 F.2-2014 ³²		
	Membrane filter (MF), ² single step	p. 124 ³	9222 D-2015 ²⁶	B-0050-85 ⁴	
2. Coliform (total), number per 100 mL	MPN, 5 tube, 3 dilution, or	p. 114 ³	9221 B-2014		
	MF, ² single step or two step	p. 108 ³	9222 B-2015 ²⁷	B-0025-85 ⁴	
	MF ² with enrichment	p. 111 ³	9222 (B + B.4e)–2015 ²⁷		
3. <i>E. coli</i> , number per 100 mL	MPN, ^{5,7,13} 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®11,14,15
	MF, ^{2, 5, 6, 7} two step, or	1103.1 ¹⁸	9222 B-2015/ 9222 I-2015, ¹⁷ 9213 D-2007	D5392–93 ⁸	
	Single step	1603, ¹⁹ 1604 ²⁰			m–ColiBlue24, ^{®16} KwikCount TM EC ^{28,29}
4. Fecal streptococci, num- ber 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	
5. Enterococci, number per 100 mL	MPN ^{5,7} , multiple tube/ multiple well, or	p. 1 10	9230 D-2013	D6503-99 ⁸	Enterolert ^{® 11,21}
	MF, ² , ⁵ , ⁶ , ⁷ two step, or Single step, or	$\frac{1106.1^{22}}{1600^{23}}$	9230 C-2013 ³⁰ 9230 C-2013 ³⁰	D5259–92 ⁸	
	Plate count	p. 143 ³			
Protozoa:					
6. Cryptosporidium	Filtration/IMS/FA	1622, ²⁴ 1623, ²⁵ 1623,1 ^{25,31}			
7. Giardia	Filtration/IMS/FA	1623, ²⁵ 1623.1 ^{25,31}			

¹ The method must be specified when results are reported.

 2 A 0.45- μ m 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. U.S. 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 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.
- ⁶ When the MF method has not 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.
- ⁷ 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 (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.

¹¹ 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 50 µg/mL of MUG may be used.
- ¹³ Samples shall be enumerated by the multiple-tube or multiple-well procedure. Using multiple-tube procedures, employ an appropriate tube and dilution configuration 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 Colilert[®] test, and is recommended for marine water samples.
- ¹⁵ Descriptions of the Colilert®, Colilert–18®, Quanti–Tray®, and Quanti–Tray®/2000 may be obtained from IDEXX Laboratories Inc.
- ¹⁶ A 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 1103.1: Escherichia coli (E. coli) in Water by Membrane Filtration Using membrane–Thermotolerant Escherichia coli Agar (mTEC), EPA-821–R-10–002. March 2010. U.S. EPA.
- ¹⁹ Method 1603: Escherichia coli (E. coli) in Water by Membrane Filtration Using Modified membrane–Thermotolerant Escherichia coli Agar (Modified mTEC), EPA–821–R–14–010. September 2014. U.S. 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. U.S. EPA.
- ²¹ A description of the Enterolert[®] test may be obtained from IDEXX Laboratories Inc.
- ²² Method 1106.1: Enterococci in Water by Membrane Filtration Using membrane–Enterococcus–Esculin Iron Agar (mE–EIA), EPA–821–R–09–015. December 2009. U.S. EPA.
- ²³ Method 1600: Enterococci in Water by Membrane Filtration Using membrane–Enterococcus Indoxyl–β–D–Glucoside Agar (mEI), EPA–821–R–14–011. September 2014. U.S. EPA.
- ²⁴ Method 1622 uses a filtration, concentration, immunomagnetic separation of oocysts from captured material, immunofluorescence assay to determine concentrations, 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. U.S. EPA.
- ²⁵ Methods 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 Water by Filtration/IMS/FA. EPA-821-R-05-002. December 2005. US EPA. Method 1623.1: *Cryptosporidium* and *Giardia* in Water by Filtration/IMS/FA. EPA 816-R-12-001. January 2012. U.S. 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.
- ²⁷ 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.
- ²⁸ A description of KwikCount [™] EC may be obtained from Micrology Laboratories LLC.
- ²⁹ Approved for the analyses of *E. coli* in freshwater only.
- ³⁰ 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 Examination 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."
- 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.
- ³² 9221 F.2–2014: This procedure allows for simultaneous detection of *E. coli* and thermotolerant coliforms by adding inverted vials to EC–MUG; the inverted vials collect gas produced by thermotolerant coliforms.
- ³³ Standard Methods for the Examination of Water and Wastewater, Joint Editorial Board, American Public Health Association, American Water Works Association, and Water Pollution Control Federation, 23rd Edition (2017), 22nd Edition (2012), 21st Edition (2005), 20th Edition (1998), 19th Edition (1995), and 18th Edition (1992).