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Designation: D5464 - 11 <u>D5464 - 16</u>

Standard Test Method for pH Measurement of Water of Low Conductivity¹

This standard is issued under the fixed designation D5464; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ε) indicates an editorial change since the last revision or reapproval.

1. Scope

1.1 This test method is applicable to determine the pH of water samples with a conductivity of 2 to 100μ S/cm over the pH range of 3 to 11-11 and is frequently used in power generation low conductivity samples. pH measurements of water of low conductivity are problematic. Specifically, this test method avoids contamination of the sample with atmospheric gases and prevents volatile components of the sample from escaping. This test method provides for pH electrodes and apparatus that address additional considerations discussed in Annex A2. This test method also minimizes problems associated with the sample's pH temperature coefficient when the operator uses this test method to calibrate an on-line pH monitor or controller (see Appendix X1).

1.2 This test method covers the measurement of pH in water of low conductivity with a lower limit of 2.0 μ S/cm, utilizing a static grab-sample procedure where it is not practicable to take a real-time flowing sample.

NOTE 1—Test Method D5128 for on-line measurement is preferred over this method whenever possible. Test Method D5128 is not subject to the limited conductivity range, temperature interferences, potential KCl contamination, and time limitations found with this method.

1.3 For on-line measurements in water with conductivity of 100 µS/cm and higher, see Test Method D6569.

1.4 For laboratory measurements in water with conductivity of 100 μ S/cm and higher, see Test Method $\frac{D1193D1293}{D1293}$.

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

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

2. Referenced Documents

2.1 ASTM Standards:²

D1129 Terminology Relating to Water

D1193 Specification for Reagent Water

D1293 Test Methods for pH of Water and and sist/07322395-ed76-4c60-8de1-5be410a6a497/astm-d5464-16 D2777 Practice for Determination of Precision and Bias of Applicable Test Methods of Committee D19 on Water

D3370 Practices for Sampling Water from Closed Conduits

D4453 Practice for Handling of High Purity Water Samples

D5128 Test Method for On-Line pH Measurement of Water of Low Conductivity

D6569 Test Method for On-Line Measurement of pH

3. Terminology

3.1 *Definitions—<u>Definitions of Terms</u>*. For definitions of terms used in these this test methods, method, refer to Terminology D1129.

3.2 *Definitions*:

3.3 Definitions of Terms Specific to This Standard:

3.3.1 *liquid-junction potential*, *n*—a dc potential that appears at the point of contact between the reference electrode's salt bridge (also known as reference junction or diaphragm) and the sample solution.

¹ These test methods are under the jurisdiction of ASTM Committee D19 on Water and are the direct responsibility of Subcommittee D19.03 on Sampling Water and Water-Formed Deposits, Analysis of Water for Power Generation and Process Use, On-Line Water Analysis, and Surveillance of Water

Current edition approved May 1, 2011 June 1, 2016. Published May 2011 June 2016. Originally approved in 1993. Last previous edition approved in $\frac{20072011}{10.1520/D5464-11.10.1520/D5464-16}$ as

² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For Annual Book of ASTM Standards volume information, refer to the standard's Document Summary page on the ASTM website.

3.3.1.1 Discussion-

Ideally, this potential is near zero and is stable. However, in low-conductivity water this potential may change from its value in buffer solution by an unknown amount, and is a zero offset (1).³

4. Reagents

4.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society where such specifications are available. ⁴ Other grades may be used, provided it is first ascertained that the reagent is of sufficiently high purity to permit its use without lessening the accuracy of the determination.

4.2 *Purity of Water*—References to water that is used for reagent preparation, rinsing or dilution shall be understood to mean water that conforms to the quantitative specifications of Type II reagent water of Specification D1193.

4.3 Commercial Buffer Solutions —Commercially available prepared buffers traceable to NIST standards should be adequate to perform the calibration procedures in 10.1 - 10.4. The exact pH of the buffer will change with temperature and this pH versus temperature data will be provided by the purveyor of the specific buffer. Refer to Test Methods D1293 for the preparation of reference buffer solutions if desired.

4.4 Buffer A—Commercially available 7.0 pH buffer.

4.5 Buffer B-Commercially available 4.0 pH buffer.

4.6 Buffer C-Commercially available 9.0 or 10.0 pH buffer.

5. Summary of Test Method

5.1 The pH meter and associated electrodes are first standardized with two calibration pH buffer solutions.

5.2 A grab sample of high purity water is taken by means of rinsing and filling two narrow mouth bottles at the sample point. Once each container is filled to the top with a representative sample excluding any air, the container is containers are capped and the samples are transported quickly to a laboratory for analysis.

5.3 pH measurement of the sample is made with high purity water pH calibration apparatus comprised of pH and reference electrodes, and automatic temperature compensator (if used). The first container is used to rinse the sensors and begin temperature equilibration and the second container is used for measurement.

5.4 A trace amount of KCl electrolyte enters calibration buffer solutions and samples via the controlled leakage rate of the reference electrode liquid junction (diaphragm) to stabilize the liquid junction potential. Excessive KCl introduction from the electrode liquid junction into low ionic strength samples will increase solution conductivity, and may alter solution pH, and should be avoided.

5.5 Temperature must be measured and both Solution Temperature Coefficient (STC) and <u>NernstianNernst</u> electrode effects compensated, either manually to the measured value or automatically by the pH meter. See Appendix X1 for a discussion of temperature effects.

6. Significance and Use

6.1 pH measurement of low conductivity water is frequently applied to power plant water and condensed steam samples for corrosion and scale prevention. It is sometimes used in pure water treatment systems between multiple pass membranes to optimize performance.

6.2 High purity water is highly unbuffered and small amounts of contamination can change the pH significantly. Specifically, high purity water rapidly absorbs CO_2 gas from the atmosphere, which lowers the pH of the sample. The sample container and accompanying pH measurement technique minimize exposure of the high purity water sample to the atmosphere.

6.3 The high purity water sample may contain volatile trace components that will dissipate from the sample if exposed to the atmosphere. The sample container used in this test method will prevent these losses.

6.4 High purity water has a significant solution temperature coefficient. For greatest accuracy the sample to be measured should be close to the temperature of the sample stream and appropriate compensation should be applied.

6.5 When the preferred Test Method D5128, which requires a real-time, flowing sample, cannot be utilized for practical reasons such as physical plant layout, unacceptable loss of water, location of on-line equipment sample points, or availability of dedicated

³ The boldface numbers in parentheses refer to the list of references at the end of this standard.

⁴ Reagent Chemicals, American Chemical Society Specifications, American Chemical Society, Washington, DC. For Suggestions on the testing of reagents not listed by the American Chemical Society, see Annual Standards for Laboratory Chemicals, BDH Ltd., Poole, Dorset, U.K., and the United States Pharmacopeia and National Formulary, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.



test equipment, this method offers a viable alternative. The most significant difference between the two test methods is that Test Method D5128 obtains a real-time pH measurement from a flowing sample and this method obtains a time delayed pH measurement from a static grab sample.

6.6 pH measurements of low conductivity water are always subject to interferences (7.1 - 7.5) and Test Method D5128 is more effective in eliminating these interferences especially with regard to contamination. This static grab sample method is more prone to contamination and temperature-induced errors because of the time lag between the sampling in the plant and sample pH reading which is taken in the laboratory.

7. Interferences

7.1 High purity, low conductivity samples are especially sensitive to contamination from atmospheric gases, from sample containers, from sample handling techniques and from excessive electrolyte (KCl) contamination from reference electrode or sample preparation such as a KCl "dosing" technique. Refer to Practice D4453 and *ASTM STP 823*(2) for discussions of sample handling and avoidance of sample contamination.

7.2 High purity water will rapidly absorb CO_2 from the atmosphere and this will lower the pH of the sample at a rate depending on the buffer capacity of the sample, the surface area of the sample exposed to air, movement of the sample, and the concentration of CO_2 at the surface of the sample which may increase if the operator exhales over the container during sampling or measurement. See Appendix X3, Table X3.1, and Fig. X3.1.

7.3 The temperature stability of the sample and how closely the sample's temperature matches the sample stream's temperature will have a direct effect on accuracy of the pH determination since temperature compensation is not perfect.

7.4 If pH is to be referenced to 25°C as required by most specifications, temperature compensation must be provided for both the <u>NernstianNernst</u> response of the electrode output (provided in most pH meters) and solution ionization effects (provided only with some on-line pH meters or by calculation with lab meters). For a discussion of temperature effects on pH measurements of high purity water see Appendix X1.

7.5 The reference junction potential can vary with ionic strength of the sample and provide an undetectable zero offset between the high ionic strength of the buffer solution and the low ionic strength of the sample. A flowing junction reference electrode (one which requires periodic refilling with electrolyte solution or that has internal electrolyte pressurization or both) minimizes this effect.

8. Apparatus

8.1 *pH Meter*—Meter, See 10.1 in Test Methodscapable of reading to 0.01 D1293.pH. The meter should preferably have automatic temperature compensation for the Nernst response of pH electrodes which provides the conversion of the electrode millivolt signal to the pH value at the measured temperature. Some on-line meters also provide an input for a solution temperature coefficient for ionization effects of the particular sample. This allows direct readout of pH referenced to 25°C. Otherwise a calculation is required.

8.2 *Sample Containers*—Two clean, narrow-mouth 250 to 500 mL bottles with cap are required. The mouth diameter should be the minimum necessary to allow insertion of the electrode(s), and temperature compensator or thermometer. A 3-hole stopper may be used to hold these sensors. The container and cap minimize exposure to atmospheric gases.

8.3 Combination pH Electrode—A probe incorporating the measuring, reference and temperature compensator functions in a single unit is recommended for its ease of insertion into a very narrow mouth sample container. Each function should conform to the characteristics in 8.4 - 8.6. Where this is not available, individual electrodes and compensator (8.4 - 8.6) may be used with a 3-hole stopper to hold them and seal the container during measurement.

8.4 *pH Glass Electrode*—The pH response of the glass electrode shall conform to the requirements set forth in 12.1 through 12.5 of Test Methods D1293. New glass electrodes and those that have been stored dry shall be conditioned and maintained as recommended by the manufacturer.

8.5 *Reference Electrode*—Double junction design, having a <u>refillable</u> flowing junction with a positive electrolyte leakage rate not to exceed 10 μ L/h. <u>A sealed reference electrode is suitable only if it is internally pressurized to force an electrolyte flow outward</u>. Prepare and maintain the reference electrode according to the manufacturer's instructions.

8.6 *Temperature Compensator*—See paragraph 10.4 in Test Methods D1293. The automatic temperature compensator must adapt for use with <u>fit into</u> the sample container to measure the temperature of the water within the container.

8.7 *Temperature Indicator*—A direct temperature indicating device must be used to measure sample water temperature within the sample container if an automatic temperature compensator is not used.

8.8 Equipment for this test method should be dedicated for high purity water use only.

9. Sampling and Sample Handling

9.1 Equipment described in Section 8 should be dedicated for high purity water use only.



9.1 The sample containers should be rinsed three times before use, with sample or reagent water. The electrode(s) and temperature device should be rinsed three times, using the same procedure as with the sample containers after each calibration in pH buffer solutions and before they are inserted into the first sample container.

9.2 Static grab samples are taken via vinylclean tubing attached to the sample take-off point. The other end of the tubing is inserted to the bottom of the sample container. The sample flows through the vinyl tube, into the bottom of the container, and then over the top to trough or sink drain. After a 5-min flush of the container with the sample water, the tubing is removed and the cap immediately installed, excluding any air. This seals the container and isolates the sample from the atmosphere. The second container is filled and capped in the same way. In the lab, the first container is used to rinse the electrodes and temperature device and the second container is used for measurement.

10. Calibration

10.1 Turn on the pH meter and allow it to warm up according to the manufacturer's instructions. If an on-line meter is used that has a solution temperature coefficient setting, be sure it is turned off or is set to 0 pH/°C for calibration in buffer solutions. Conventional Nernst temperature compensation should be active at all times and requires no setting.

10.2 Remove the electrode(s) and temperature compensator (if used) from storage. Check the reference electrode for proper electrolyte level as recommended by the manufacturer.

10.3 Calibrate the electrode(s) and pH meter at two points according to manufacturer's instructions. Also, refer to Section 12 of Test Methods D1293 for guidelines on the calibration of a pH meter and electrode assembly. Use a quiescent sample of Buffer A and Buffer B if the sample point of interest is below 7.0 pH. Use a quiescent sample of Buffer A and Buffer C if the sample point of interest is above 7.0 pH. Use laboratory glassware dedicated for this service only. Thoroughly rinse the electrode(s) and glassware with reagent water three times between immersion in each buffer solution.

10.4 Obtain calibration precision of the pH electrode(s) and the pH meter by repeating the two-point calibration described in 10.3, making any necessary readjustments to the pH meter. If the electrode slope (efficiency) is less than 94 % or greater than 101 %, refer to manufacturer's instructions for repair or replacement of electrode(s).

Note 2—The pH electrodes in use may pass the above calibration procedures (10.1 - 10.4), but caution should be taken. pH electrodes that are not specifically designed for use in high purity water may develop problems with liquid junction potential during actual test measurements.

10.5 Determine the frequency of the two-point calibration of the electrode(s) and the pH meter based on experience. Perform calibration at least daily when pure water sample testing is performed daily. For less frequent pure water sample testing, perform calibration procedures just prior to a consecutive series of sample tests.

10.6 Thoroughly rinse the electrode(s) and the temperature compensator or temperature reading device and the sample container three times with sample or reagent water after calibration and before measurement.

11. Procedurendards.iteh.ai/catalog/standards/sist/07322395-ed76-4c60-8de1-5be410a6a497/astm-d5464-16

11.1 Before starting the procedure, make certain the two sample containers are clean and empty. See Section 9.

NOTE 3—If on-line pH sensors are to be calibrated by this test method (refer to Test Method D5128), steps mustshould be taken to prevent the disturbance of the on-line pressure and flow rate while the grab sample is being taken. The <u>Appropriate</u> pressure and flow control regulation equipment shown inshould be Fig. 1 will control and stabilize this on-line pressure and flow rate by installed at the grab sample point upstream of the on-line sensor. See D3370means of rotameter. Appendix X2 Reprovides_T and secondary pressure regulatorguidance on grab sample PR calibration_T even when a sample is taken at grab sample point of on-line instrumentation.S₂ (see <u>Appendix X2</u>).

11.2 Open the sample valve \forall_z and pull sample without interrupting the sample flow-rate or pressure of the on-line pH sensor assembly <u>if used</u> (see on-line pH sensor manufacturer's instructions for its optimum flowrate). The grab sample flowrate should be at least 200 mL/min.

11.3 With the sample flowing, rinse the outside surfaces of the vinyl sample tubing for at least 15 seconds by holding it with the open discharge end upward like a fountain, allowing the sample to flow over the outside tubing surfaces for more than the length that will be later immersed in the sample container, allowing the flow to drain into the sampling trough or sink.

11.4 Run the end of the vinyl sample tubing to the bottom of the sample container letting the sample overflow into the sink.

11.5 Thoroughly flush the sample container with sample water for at least 5 min at a flow rate of at least 200 mL/min.

11.6 Remove the vinyl-sample tubing and immediately cap the container. Repeat for the second container and close sample valve-valve. V_2 .

11.7 Transport the sample containers to the laboratory without delay and immediately immerse the sensors (which have already been calibrated and rinsed thoroughly) in the first container for 1 min to acclimate them to the sample composition and temperature.

11.8 Move the sensors to the second container. Take the pH measurement and temperature measurements within 3 min in order to minimize KCl contamination from the reference electrode and change in the temperature of the grab sample.

11.9 Report the pH value with 0.01 pH resolution and the temperature with 0.1°C resolution.



<u>11.10 If the pH must be referenced to 25°C and the sample is not already at 25.0 \pm 0.5°C, determine the applicable solution temperature coefficient and calculate the pH at 25.0°C in accordance with Appendix X1 and report it with resolution of 0.01 pH.</u>

11.11 To use this pH measurement to calibrate an on-line pH sensor, call this reading the value from 11.9 or 11.10 R_1 and refer to the procedure addendum in Appendix $\frac{X3}{2}$.

NOTE 4—If no further samples are to be taken, the calibrated pH electrode(s) may be kept stored in the sample container containing the last pure water sample until the next calibration or sample requirement. For long term storage of the pH electrode(s), replace them in their respective soaker bottles and appropriate storage solutions (see manufacturer's instructions).

12. Quality Control

12.1 Two-point instrument calibration must be performed according to the manufacturer's instructions within 12 h of making measurements.

12.2 With power plant samples containing predominately ammonia and negligible carbon dioxide, an accurate correlation with specific conductivity can be made. See Fig. X3.1. Under these conditions, periodically verify pH readings corrected to 25°C by solution temperature compensation with the conductivity also referenced to 25°C. Some two-channel conductivity instrumentation includes an option for the calculation of pH based on specific and cation conductivity and will make a correction for the presence of minor amounts of carbon dioxide and other contaminants. If available, use this calculated value to verify the measured pH readings under normal plant operating conditions.

13. Precision and Bias

13.1 Neither precision nor bias data can be obtained for this test method from a collaborative study designed in accordance with the requirements of Practice D2777 since this test method relies on the capture of a flowing sample, and the sample must remain captive until after the measurement is taken due to its high vulnerability to contamination. This inability to obtain precision and bias data for continuous determinations is recognized and stated in the scope of Practice D2777.

14. Keywords

14.1 automatic temperature compensator; controlled leakage rate; flowing liquid junction; high purity water; liquid junction potential; low conductivity water; pH glass electrode; pH temperature coefficient; reference electrode; solution temperature coefficient

Document Preview

(Mandatory Information)

https://standards A1. DISCUSSION OF THE CONDUCTIVITY RANGE FOR THIS TEST METHOD astm-d5464-6

A1.1 The scope of this test method provides the user with a guideline conductivity range (<100 μ S/cm) for what is considered low conductivity or pure water. This conductivity range is not a specific cut-off or limit, but is intended to provide the user with an indication of when to apply this test method for reliable pH measurements of waters with low conductivity. A number of known interferences and problems apply to pH measurement apparatus and methods when attempting pH measurement of waters with low conductivities. These interferences (see measurements. 7.1 – Tables A1.1 and A1.27.5) are not necessarily applicable to highly buffered waters typified by conductivity levels above 100 μ S/cm. Table A1.1 and Table A1.2 illustrate the high pH sensitivity of low ionic strength water samples containing low concentrations of contaminants. The sources of these contaminants can include the atmosphere (see Table X3.1), the pH measurement apparatus, and improper procedures. The user of these test methods is advised to use the specialized apparatus and procedures given in this test method when attempting pH measurements of waters with a conductivity <100 μ S/cm.



TABLE A1.1 Calculated Conductivity and pH Values at 25°C of Low Concentrations of NaOH in Pure Water A, B

Note 1—This table tabulates the theoretical conductivity and pH values of low levels of NaOH in pure water as calculated from available thermodynamic data.

Note 2—To illustrate the high sensitivity of the sample pH at these low concentrations to contaminants, the last column lists errors that would result if the sample were contaminated with an additional 1 mg/L through sample or equipment handling errors.

Sample Concen- tration, mg/L	Sample Conduc- tivity, µS/cm	Sample pH	∆pH Error from Additional 1 mg/L NaOH Contami- nate
0.001	0.055	7.05	∆2.35
0.010	0.082	7.45	∆1.95
0.100	0.625	8.40	∆1.03
1.0	6.229	9.40	∆0.30
8.0	49.830	10.30	∆0.05

^{*A*} Data courtesy of R. C. Hunt, "Calculated pH and Conductivity Values," Sensor Development, Inc., June 1986.

^B This data was developed from algorithms originally published in *Ultrapure Water* as: Hunt, Robert C., "A Review of pH and Conductivity Measurement Techniques in High Purity Water, with Bibliography," *Ultrapure Water*, Vol 4, 1987, p. 26.

TABLE A1.2 Calculated Conductivity and pH Values at 25°C of Low Concentrations of HCl in Pure Water A. B

NOTE 1—This table tabulates the theoretical conductivity and pH values of low levels of HCl in pure water as calculated from available thermodynamic data.

NOTE 2—To illustrate the high sensitivity of the sample pH at these low concentrations to contaminants, the last column lists errors that would result if the sample were contaminated with an additional 1 mg/L through sample or equipment handling errors.

Sample Concen-	Sample Conduc-		∆pH Error from	
Concentration,	Conductivity,	Sample pH	Additional 1 mg/L	
tration, mg/L	tivity, μS/cm		HCI Contaminate	
0.001	0.060	5466.946	Δ2.38	
0.010	0.134	6.51	∆1.95	
talog/0.100 dard	s/sist/1.166223	95-e5.56)-4c6	0-8da1.03 be4	
1.0	11.645	4.56	∆0.30	
8.0	93.163	3.66	∆0.05	

^A Data courtesy of R. C. Hunt, "Calculated pH and Conductivity Values," Sensor Development, Inc., June 1986.

^B This data was developed from algorithms originally published in *Ultrapure Water* as: Hunt, Robert C., "A Review of pH and Conductivity Measurement Techniques in High Purity Water, with Bibliography," *Ultrapure Water*, Vol 4, 1987, p. 26.

A2. CONSIDERATIONS FOR THE REQUIREMENT OF SPECIALIZED pH MEASUREMENT APPARATUS

A2.1 pH electrodes are available in many different configurations containing a wide variety of membrane and liquid junction designs as well as a variety of internal electrolyte formulations. Selection of the appropriate pH electrode features required for pH measurement of low conductivity water is not necessarily obvious.

A2.1.1 The temperature coefficient of the electrodes will affect the accuracy and repeatability of the measurement. Electrodes which quickly equilibrate to each other and the sample temperature must be selected for this service. Refer to X1.2 in Test Method D1293 and <u>ASTM STP 190 (3)</u>.