

Standard Practice for Operating the Severe Wastewater Analysis Testing Apparatus¹

This standard is issued under the fixed designation G210; 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 practice covers the basic apparatus, procedures, and conditions required to create and maintain the severe wastewater analysis testing apparatus used for testing a protective coating or lining.

1.2 This apparatus may simulate the pertinent attributes of a typical domestic severe wastewater headspace (sewer) environment. The testing chamber comprises two phases: (1) a liquid phase containing a prescribed acid and saline solution, and (2) a vapor phase consisting of air, humidity, and concentrated sewer gas (Note 1). The temperature of the test chamber is elevated to create accelerated conditions and reaction rates.

Note 1—For the purposes of this practice, sewer gas is composed of hydrogen sulfide, carbon dioxide, and methane gas.

1.3 **Caution**—This practice can be extremely hazardous. All necessary precautions need to be taken when working with sewer gas, sulfuric acid, and a glass tank. It is highly recommended that a professional testing laboratory experienced in testing with hydrogen sulfide, carbon dioxide, and methane gases perform this practice.

1.4 The values stated in inch-pound units are to be regarded as standard. The values given in parentheses are mathematical conversions to SI units that are provided for information only and are not considered standard.

1.5 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. Some specific hazards statements are given in Section 8 on Hazards.

2. Referenced Documents

2.1 ASTM Standards:²

A36 Specification for Carbon Structural Steel

- C307 Test Method for Tensile Strength of Chemical-Resistant Mortar, Grouts, and Monolithic Surfacings
- C387 Specification for Packaged, Dry, Combined Materials for Mortar and Concrete
- C580 Test Method for Flexural Strength and Modulus of Elasticity of Chemical-Resistant Mortars, Grouts, Monolithic Surfacings, and Polymer Concretes
- D610 Practice for Evaluating Degree of Rusting on Painted Steel Surfaces
- D638 Test Method for Tensile Properties of Plastics
- D660 Test Method for Evaluating Degree of Checking of Exterior Paints
- D661 Test Method for Evaluating Degree of Cracking of Exterior Paints
- D714 Test Method for Evaluating Degree of Blistering of Paints
- D790 Test Methods for Flexural Properties of Unreinforced and Reinforced Plastics and Electrical Insulating Materials
- D2370 Test Method for Tensile Properties of Organic Coatings
- D4541 Test Method for Pull-Off Strength of Coatings Using Portable Adhesion Testers
- D6677 Test Method for Evaluating Adhesion by Knife
- D7091 Practice for Nondestructive Measurement of Dry Film Thickness of Nonmagnetic Coatings Applied to Ferrous Metals and Nonmagnetic, Nonconductive Coatings Applied to Non-Ferrous Metals
- G193 Terminology and Acronyms Relating to Corrosion

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¹ This practice is under the jurisdiction of ASTM Committee G01 on Corrosion of Metals and is the direct responsibility of Subcommittee G01.05 on Laboratory Corrosion Tests.

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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.

2.2 ISO Standards:³

- ISO 16773 Paints- and Varnishes- Electrochemical Impedance Spectroscopy (EIS) on High Impedance Coated Samples. Part 1: Terms and Definitions
- ISO 16773 Paints- and Varnishes- Electrochemical Impedance Spectroscopy (EIS) on High Impedance Coated Samples. Part 2: Collection of Data
- ISO 16773 Paints- and Varnishes- Electrochemical Impedance Spectroscopy (EIS) on High Impedance Coated Samples. Part 3: Processing and Analysis of Data from Dummy Cells

3. Terminology

3.1 Definitions of Terms Specific to This Standard:

3.1.1 *domestic wastewater*, *n*—wastewater discharged from residences and from commercial, institutional, and similar facilities.

3.1.2 sewer headspace, n—the air space between the water surface and the top of the pipe (crown) or other enclosed structure.

3.2 For definitions of terms used in this practice, see Terminology G193.

4. Summary of Practice

4.1 The corrosion protection of steel, ductile iron, and concrete by a protective coating or lining may be altered by exposure to sewer gases and by the composition of the corrosive reagents found in headspace environments of domestic wastewater conveyance and treatment structures.⁴

4.2 This practice provides a controlled corrosive environment, which has been utilized to produce a simulated severe sewer headspace condition by wetting the coated samples in a cyclic fashion with a corrosive solution and then exposing the samples to air containing sewer gas. This condition is responsible for reducing the barrier properties of protective coatings and linings.

4.3 Test specimens are positioned on a carousel and placed inside an airtight testing apparatus (chamber) maintained at a temperature of $150 \pm 5^{\circ}F(65 \pm 3^{\circ}C)$. The chamber contains a prescribed aqueous solution (liquid phase) at the bottom and a headspace (vapor phase) containing sewer gas. The test specimens are immersed into liquid phase for a period of 15 min each. After immersion, the specimens are exposed to the vapor phase the balance of the time. This constitutes one complete cycle with three cycles occurring per day. This cyclic exposure continues for a period of 28 days.

4.4 The specified operating temperature, aqueous solution, sewer gases, and duration parameters are considered the standard for the purposes of this practice. The specifications may be adjusted to replicate specific environments if mutually agreed upon between the client and testing facility. Any deviations from this practice shall be reported.

5. Significance and Use

5.1 Domestic wastewater headspace environments are corrosive due to the presence of sewer gases and sulfuric acid generated during the biogenic sulfide corrosion process.⁵ This operating procedure provides an accelerated exposure to sewer gases and concentration of sulfuric acid commonly produced by bacteria within these sewer environments.⁶

5.2 The results obtained by the use of this practice can be a means for estimating the protective barrier qualities of a protective coating or lining for use in severe sewer conditions.

5.3 Some protective coatings or linings may not withstand the exposure temperature specified in this practice but have demonstrated satisfactory performance in actual sewer exposures, which are at lower temperatures.

6. Apparatus

6.1 The testing apparatus consists of the following:

6.1.1 *Glass Tank*—Minimum diameter 16 by 12 in. (40 by 30 cm) tall. The glass tank, when fitted with a polypropylene lid (tank cover) and elastomeric seal, creates an air-tight test chamber. The glass tank is inert to the aggressive reagents at the testing temperature. The glass tank shall be transparent to permit visual examination of the test specimens throughout specified testing duration.

6.1.2 *Polypropylene Lid*—Octagon or round shaped, minimum 1 in. (2.54 cm) thick by 18 in. (46 cm) span. The polypropylene lid has a 1.5 in. (3.81 cm) diameter center port to accommodate the shaft of the sample carousel. The shaft slides through an O-ring seal which is secured and tensioned with a polypropylene fitting (Note 2). The shaft slides easily up and down through the O-ring seal while preventing the release of test gases. Silicone grease lubricant can be used to facilitate movement of the shaft.

Note 2—Polypropylene has been found to be an acceptable material for this service. Other materials, such as polytetrafluoroethylene (PTFE) or polyetheretherketone (PEEK) may also provide acceptable service.

6.1.2.1 The polypropylene lid is designed with two ports for fittings, which accommodate inlet and outlet lines.

6.1.2.2 The side of the polypropylene lid which faces into the tank has a circular, shallow 1-in. (2.54-cm) wide groove. The groove accommodates a suitable corrosion resistant elastomeric seal (gasket) required to seal the lid of the glass tank.

6.1.2.3 The polypropylene lid also includes eight equallyspaced holes along the outer edge to accommodate eight threaded rod fasteners with wing nuts, nuts, and washers. The eight threaded rods connect the polypropylene lid to a solid, chemical resistant base plate made of laminated wood or

³ Available from International Organization for Standardization (ISO), 1, ch. de la Voie-Creuse, CP 56, CH-1211 Geneva 20, Switzerland, http://www.iso.org.

⁴ O'Dea, V. et al., "Testing Permeation Resistance in Coatings for Wastewater Structures," *Journal of Protective Coatings and Linings*, September 2010, pp. 16–28.

⁵ O'Dea, V., "Understanding Biogenic Sulfide Corrosion," *Materials Performance*, November 2007, pp. 36–39.

⁶ O'Dea, V. et al, "Assessing Coatings & Linings for Wastewater: Accelerated Test Evaluates Resistance to Severe Exposures," *Journal of Protective Coatings and Linings*, April 2008, pp. 44–57.

equivalent materials, located under the glass tank, hence clamping the lid to the glass tank, thereby creating an air-tight testing chamber.

6.1.3 Specimen Carousel—Constructed of polypropylene (or other suitable corrosion-resistant material) to accommodate the various types of samples. Coated steel specimens will sit in slots and rest vertically, arranged radially (Fig. 1). Coated concrete specimens will sit in slots oriented vertically. Cast shapes and free films will be oriented either vertically or horizontally, depending upon their dimensions.

6.1.3.1 The specimen carousel consists of a tray with a perpendicular central shaft, which facilitates raising and lowering of the carousel within the chamber to the liquid phase (lowered position) or vapor phase (raised position).

6.1.3.2 When the carousel is in the liquid phase (lowered position) the specimen carousel, including the test specimens, must be completely immersed in the aqueous solution.

6.1.3.3 When the carousel is in the raised position, it is locked in place with a retaining pin assembly outside the top of the oven. The aqueous solution must drain away from the test specimens through drain holes in the carousel.

6.1.4 *Gas Inlet*—A flexible and resilient polypropylene or polyethylene gas supply line connects the sewer gas supply to a polypropylene inlet fitting on the tank cover. The inlet fitting assembly accommodates a polypropylene extension tube, which runs to the bottom of the chamber. The vertical extension tube allows the sewer gas mixture to be sparged through the aqueous solution.

6.1.4.1 Gas supply line includes a gas flow controller and indicator (for example, rotameter) to measure instantaneous flow rate.

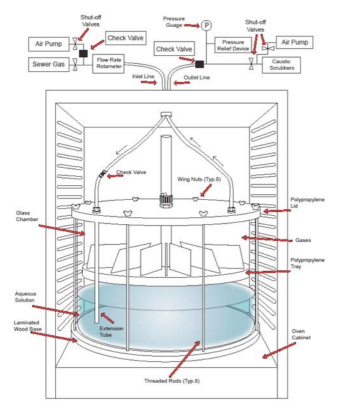


FIG. 1 Severe Wastewater Analysis Testing Apparatus

6.1.4.2 Gas supply line includes a polypropylene stopcock valve with low-friction plug of PTFE and a polypropylene gas check valve with a suitable fluoroelastomer (FKM)-coated diaphragm.

6.1.5 *Air Purge Inlet*—An external flexible and resilient polypropylene or polyethylene fresh air supply line connects the air pump to the tank through a tee into the Gas Inlet line.

6.1.5.1 Air supply line includes a polypropylene stopcock valve with low-friction plug of PTFE and a polypropylene gas check valve with FKM-coated diaphragm.

6.1.6 *Gas Outlet*—An external flexible and resilient polypropylene or polyethylene line from the tank cover with a polypropylene connector is connected to caustic scrubbers to capture H_2S from the effluent gases.

6.1.6.1 Outlet line includes a check valve, a polypropylene stopcock valve with low-friction plug of PTFE, a pressure relief valve (0.5 psi or 3.4 kPa), and a pressure gauge (0 to 1.5 psig range or 0 to 10 kPa) teed into the gas outlet line using PTFE coated isolation diaphragm. An air pump is teed into the line to the caustic scrubbers to facilitate sewer gas removal from the scrubber lines.

6.1.6.2 *Secondary Containment*—A polypropylene tray of suitable volume may be placed under the test chamber as an additional precaution against acid spillage.

6.1.7 Oven—Convection (forced air) oven of sufficient capacity to accommodate the test chamber and be capable of maintaining a temperature of $150 \pm 5^{\circ}$ F ($65 \pm 3^{\circ}$ C) throughout the duration of the test exposure. This ensures a uniform temperature throughout the chamber for the testing duration. The top of the oven must have a 3-in. (75-mm) through-wall opening to accommodate the inlet and outlet gas lines and the shaft of the specimen carousel and its movement.

6.1.8 *Air Pump*—Variable-flow air pump to purge the test chamber and outlet lines of the hydrogen sulfide gas (and other sewer gases) at the completion of the exposure time. Pump delivery pressure and pressure relief valve should be sized to avoid accidentally over-pressurizing the glass tank.

6.1.8.1 An air flow rate of at least 1.5 litres per minute (L/min) is recommended.

6.1.9 *Caustic Scrubbers*—Capable of removing H_2S from the exhaust test gas. A typical scrubber consists of a 4-L polypropylene carboy, half filled with 15 % sodium hydroxide (NaOH). Bubble dispersion media is added to the carboy.

6.1.9.1 "Percent" is defined as grams of solute per volume of solution. 15 % NaOH contains 150 g of sodium hydroxide per litre of solution, which is the same as 3.75 mol of sodium hydroxide per litre of solution.

6.1.9.2 Carboy cap is equipped with polypropylene fittings which accommodate a 0.25 in. (6.35 mm) polyethylene inlet dip tube and outlet tube (exhaust).

6.1.9.3 The use of two or more scrubbers in series improves removal efficacy and reduces the likelihood of accidental release of H_2S when scrubber capacity is exceeded.

Note 3—Carbon dioxide is also absorbed by the scrubber, reducing their capacity accordingly.