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Designation: D 6350 – 98

Standard Test Method for Mercury Sampling and Analysis in Natural Gas by Atomic Fluorescence Spectroscopy¹

This standard is issued under the fixed designation D 6350; 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 covers the determination of total mercury in natural gas streams down to 0.001 μ g/m³. It includes procedures to both obtaining a representative sample and the atomic fluorescence detection of the analyte. This procedure can be applied for both organic and inorganic mercury compounds.

1.2 Both, inch-pound and SI (metric) units of measurement are used throughout this standard.

1.3 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 its use.

2. Referenced Documents

2.1 ASTM Standards:

D 3684 Test Method for Total Mercury in Coal by the Oxygen Bomb Combustion Atomic Method²

D 5954 Test Method for Mercury Sampling and Measurement in Natural Gas²

2.2 ISO Standard:

ISO 6978 Determination of Mercury in Natural Gas^{3,4–4}C

3. Summary of Test Method

3.1 Mercury from the gaseous stream is absorbed and preconcentrated onto a gold-coated silica sand trap. The analyte is desorbed by raising the temperature of the trap, and a flow of inert gas carries the mercury atoms into the cell assembly of an atomic fluorescence spectrophotometer. The cell is irradiated by a low pressure mercury vapor lamp at 253.652 nm. Excitation of mercury atoms produces resonance fluorescence which reradiates at the excitation wavelength. The fluorescence radiation is detected by a photomultiplier tube and is directly proportional to the amount of mercury in the cell. The concentration of the element in the original sample is obtained by comparison to freshly prepared standards, which are analyzed by direct injection of mercury vapor into the instrument at a specified temperature on supported gold traps.

4. Significance and Use

4.1 This test method can be used to determine the total mercury concentration of a natural gas stream down to 0.001 μ g/m³. It can be used to assess compliance with environmental regulations, predict possible damage to gas plant equipment, and monitor the efficiency of mercury removal beds.

4.2 The preferred sampling method for mercury collection is on supported gold sorbent, which allows the element to be trapped and extracted from the interfering matrix of the gas. Thermal desorption of mercury is performed by raising the temperature of the trap by means of a nichrome wire coiled around it.

4.3 Since AFS demonstrates lower detection limits approaching 0.1 pg, this test method avoids difficulties associated with prolonged sampling time. Saturation of the trap with interferants such as hydrogen sulfide (H_2S) is avoided. Average sampling can range between 15 to 30 min, or less.

5. Apparatus and Materials

5.1 Sampling Equipment:

5.1.1 Sample probe, equipped with a ball valve of the Type 316 SS, connected to the sampling point is highly recommended.

5.1.2 Pressure regulation devices, such as two-stage stainless steel pressure regulator, capable of reducing the pressure from 2000 to 30 psi.

5.1.3 On/off and micrometric-type valves capable of regulating the natural gas sample flow rate in the range of 100 to 200 mL/min.

5.1.4 Stainless steel tubing and compression-type fittings, as required.

5.1.5 Dry or wet flow meter or integrating anemometer to measure properly the total volume of the gas sample collected.

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² Annual Book of ASTM Standards, Vol 05.05

³ Available from American National Standards Institute, 11 W. 42nd St., 13th Floor New York, NY 10036.

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5.1.6 Gold-coated fused silica sand traps.

NOTE 1—For details on trap preparation refer to Test Method D 5954D 5954, the procedure of vapor deposition used in scanning electron microscopy (SEM) techniques,⁴ and, ISO 6978, 1993.

5.2 Analytical Equipment:

5.2.1 *Atomic Fluorescence Spectrophotometer*, equipped with a quartz cell and a mercury lamp capable of irradiating at 253.652-nm wavelength.

5.2.2 *Chromatography Grade Teflon*[®] *and Silicon Tubing*, for connections between the thermal desorption system and the AFS. Length, ID, and OD are selected as appropriate.

5.2.3 *Nichrome Wire* (22 gauge) coiled (20 turns/inch) around the traps for the thermal desorption of mercury.

5.2.4 *Variable Voltage Regulator*, (rheostat) used in conjunction with the nichrome wire for the rapid heating of the traps.

5.2.5 *Temperature-Resistant Rubber Tubing*, of $\frac{1}{4}$ in. (0.06 mm), connecting the trap to the temperature desorption system.

5.2.6 *GC-Grade Septa*, low bleed, made of silicone used in the injection port and mercury-sealed vial.

5.2.7 Constant Temperature Bath, capable of regulating the temperature of a sealed vial of mercury to $25 \pm 0.1^{\circ}$ C.

5.2.8 Various Stainless Steel "T" Fittings.

5.2.9 *Gastight Syringes*, fixed or variable volume, in the range of 10 to 500 μ L.

5.2.10 *A Glass Vial*, 100 mL fitted with a septum to perform as mercury container.

5.2.11 *Chart Recorder*, or integrator to process a hard copy of the data acquired by the detector.

NOTE 2—Commercially available permeation injection sources, based on the principle of permeation tubes, can be used instead of gastight syringes. Permation devices can be used in lieu of gastight syringe-based sample introduction. A permiation system can automatically introduce an accurately known amount of mercury vapor onto a gold trap. This is particularly convenient for quantifying low pg amounts of mercury.

6. Reagents

6.1 Because of the error and contamination that may be introduced from impurities in the chemicals, the use of high purity reagents is strongly recommended.

6.1.1 Mercury Analytical Grade, triple distilled.

NOTE 3—Warning: Mercury vapor is harmful. Use proper ventilation when handling.

6.2 Argon Gas, ultra high purity grade (UHP 99.999 %).

Note 4—For the permeation injection source procedure, certified mercury permeation tubes are commercially available. Tubes can also be prepared and calibrated by comparison to syringe injection or by weight loss, over time, using an analytical balance with a resolution of ± 0.01 mg.

7. Sampling Procedure

7.1 Every effort should be made to ensure that the sample is representative of the gas source from which it is taken. Select always the best and more representative sampling point for mercury trapping. Sampling will require the use of specific procedures; consult appropriate regulations.

7.2 Sampling arrangements will always use a minimum of two sampling gold tubes per location. The recommended sampling setup is shown schematically in Fig. 1.

7.3 Assemble the parts without connecting the gold traps, as depicted in Fig. 1. Open the flow of gas from the main valve and regulate the pressure down to 30 psi. Open the on/off valve and set an approximate flow of 150 mL/min with the micrometric valve adjustment. Check the flow with a dry or bubble flow meter. Let the system purge for at least 30 min. Purging is necessary, especially if the pressure regulator, tubing, and valves were used at a previous location. The longer the purging period the better.

7.4 When purging is completed, close the on/off valve and connect both gold traps to the system. Use Tygon tubing or similar to connect traps together.

7.5 Open the on/off valve again and record the time and the exact flow through the traps. Periodically check, every 15 min, that the flow remains constant throughout the duration of sampling. Best results are obtained with a 100- to 200-mL/min flow rate and an average sampling time of 15 to 30 min. Record both readings.

7.6 When sampling time has elapsed, close the on/off valve and disconnect the traps. Carefully cap and label them accordingly (Tube 1 and Tube 2). Accurately record the final time and flow data for later calculations.

8. Calibration of the Instrument (Gaseous Standard)

8.1 Calibration according to the following procedure is recommended since it is easy to perform and results in repeatability not exceeding a 10 % range between duplicate analyses. (see Footnote 5^5 and ISO 6978).

8.2 Standards are prepared by injection of different volumes of the head space from a thermostatted sealed mercury vial. Injection of the aliquots, usually in the microlitre range, should be made directly onto a mercury trapping tube, using a T-piece injection port and argon gas as carrier. See Fig. 2 for details.

8.3 All surfaces coming in contact with the mercury vapor should be passivated (except the analytical trap) before actual readings can be taken. Condition all tubing, instrument connections, as well as all syringes, by multiple injections of the gaseous mercury vapor head space contained in the temperature-controlled mercury vial.

8.4 The concentration of a particular aliquot, taken with a gastight syringe, can be calculated by the following equation of state of real gases:

$$\log(ng/mL) = (-3104/K) + 11.709$$
(1)

where:

K = Mercury temperature in Kelvins.

For instance, a $100-\mu L$ withdrawal of the head space over mercury at $24^{\circ}C$ will result in an absolute mercury concentration of 1.83 ng on the gold trap.

⁴ Fitzgerald, W.F., and Gill, G.A. "Subnanogram Determination of Mercury by Two-Stage Gold Amalgamation and Gas Phase Detection Applied to Atmospheric Analysis," *Analytical Chemistry*, 11, 1714, 1979.

⁵ Dumarey, R., Temmerman, E., Dams, R., and Hoste, J., "The Accuracy of the Vapour-Injection Calibration Method for the Determination of Mercury by Amalgamation/Cold-Vapor Atomic Absorption Spectrometry," *Analytica Chimica Acta*, 170, (1985), pp. 341-346.