ALUMINUM and compounds, as Al

AI MW: 26.98 (AI); 101.96 (AI₂0₃) CAS: 7429-90-5 (AI); 1344-28-1 (AI₂0₃) RTECS:

METHOD: 7013, Issue 2 CLASS B Issue 1: 15 February 1984

Issue 2: 15 August 1992

OSHA: no standard PROPERTIES: ductile metal; valence 3; MP 660 °C;

VP not significant

NIOSH: no standard ACGIH: 10 mg/m³ (metal, oxide);

5 mg/m³ (pyro powder, welding fume);

2 mg/m³ (soluble salts, alkyls)

SYNONYMS: vary depending upon the compound; Alumina (Al ₂O₃)

	SAMPLING		MEASUREMENT
SAMPLER:	FILTER	TECHNIQUE:	ATOMIC ABSORPTION, FLAME
	(0.8-µm cellulose ester membrane)	ANALYTE:	aluminum
FLOW RATE:	1 to 3 L/min	ASHING:	conc. HNO ₃ , 6 mL; 140 °C
-MAX:	10 L @ 5 mg/m ³ 400 L	FINAL SOLUTION:	: 10% HNO₃, 10 mL 1000 μg/mL Cs
SHIPMENT: SAMPLE	routine	FLAME:	nitrous oxide-acetylene, reducing
STABILITY:	stable	WAVELENGTH:	309.3 nm
BLANKS:	2 to 10 field blanks per set	BACKGROUND COLLECTION: none used	
		CALIBRATION:	Al*** in 10% HNO ₃
	ACCURACY	RANGE:	50 to 5000 μg per sample [1]
RANGE STUDIED: not studied		ESTIMATED LOD:	2 µg per sample [4]
		PRECISION (s,):	0.03 [1,4]
ACCURACY:			
BIAS:	none identified		
OVERALL PR	ECISION (s,): not evaluated		

APPLICABILITY: The working range is 0.5 to 10 mg/m³ for a 100-L sample. This is an elemental analysis, not compound specific. Verify that the types of compounds in the samples are soluble with this ashing procedure. Aliquots of the samples can be analyzed separately for approximately four additional metals.

INTERFERENCES: Cesium at 1000 μ g/mL controls ionization in the nitrous oxide-acetylene flame. Iron and HCl at greater than 0.2% (w/w) decrease the sensitivity. Vanadium or H $_2$ SO $_4$ require 1% (w/w) La as a releasing agent.

OTHER METHODS: This is Method P&CAM 173 for Al [1] in a revised format. Method 7300 (ICP-AES) is an alternate analytical method.

REAGENTS:

- 1. Nitric acid, conc.
- 2. Nitric acid, 10% (w/v). Add 100 mL conc. HNO₃ to 500 mL water; dilute to 1 L.
- Calibration stock solution, 1000 μg Al/mL. Commercially available or dissolve 1.000 g Al wire in minimum volume of (1+1) HCl using small drop of Hg as catalyst. Dilute to 1 L with 1% (v/v) HCl.
- 4. Cs solution, 50 mg/mL. Dissolve 73.40 g CsNO₃ in 100 mL water; dilute to 1 L.
- 5. Nitrous oxide.
- 6. Acetylene.
- 7. Distilled or deionized water.

EQUIPMENT:

- Sampler: cellulose estr membrane filter, 0.8
 µm pore size, 37-mm diameter; in cassette
 filter holder.
- 2. Personal sampling pump, 1 to 3 L/min, with flexible connecting tubing.
- 3. Atomic absorption spectrophotometer with a nitrous oxide-acetylene burner head and aluminum hollow cathode lamp.
- 4. Regulators, two-stage, for N ₂O and acetylene.
- 5. Beakers, Phillips, 125-mL, or Griffin, 50-mL, with watchglass covers.*
- 6. Volumetric flasks, 10- and 100-mL.*
- 7. Micropipets, 5 to 500 µL.*
- 8. Hotplate, surface temperture 100 to 140 °C.
 - * Clean with conc. nitric acid and rinse thoroughly with distilled or deionized water before use.

SPECIAL PRECAUTIONS: Perform all acid digestions in a fume hood.

SAMPLING:

- 1. Calibrate each personal sampling pump with a representative sampler in line.
- 2. Sample at an accurately known flow rate between 1 and 3 L/min for a total sample size of 10 to 400 L. Do not exceed 2 mg total dust loading on the filter.

SAMPLE PREPARATION:

NOTE: The following sample preparation gave quantitative recovery (see EVALUATION OF METHOD). Steps 4 through 9 of Method 7300 or other quantitative ashing techniques may be substituted, especially if several metals are to be determined on a single filter.

- 3. Open the cassettes and transfer the samples and blanks to clean beakers.
- 4. Add 6 mL conc. HNO 3 and cover with a watchglass. Start reagent blanks at this point.
- 5. Heat on hotplate (140 °C) until sample dissolves and a slightly yellow solution is produced. Add acid as needed to completely destroy organic material.
- When the sample solution is clear, remove watchglass and rinse into the beaker with 10% HNO 3.
- 7. Place the beakers on a hotplate and allow to go to dryness.
- 8. When sample is dry, rinse walls of beaker with 3 to 5 mL 10% HNO 3. Reheat for 5 min to dissolve the residue, then allow to air cool.
- 9. Transfer the solution quantitatively to a 10-mL volumetric flask containing 0.2 mL 50 mg/mL Cs solution. Dilute to volume with 10% HNO $_3$.
 - NOTE: If vanadium or sulfuric acid are present, add 1% (w/w) La as a releasing agent [1,3].

CALIBRATION AND QUALITY CONTROL:

10. Add known amounts, covering the range 0 to 500 mg Al per sample, of calibration stock solution to 100-mL volumetric flasks containing 2.0 mL 50 mg/mL Cs solution and dilute to volume with 10% HNO₃.

- 11. Analyze the working standards together with the samples and blanks (steps 16 and 17).
- 12. Prepare a calibration graph of absorbance vs. solution concentration (µg/mL).
- 13. Aspirate a standard for every 10 samples to check instrument drift.
- 14. Check recoveries with at least one spiked media blank per 10 samples.
- 15. Use method of additions occasionally to check for interferences.

MEASUREMENT:

- 16. Set spectrophotometer according to manufacturer's instructions and to conditions on page 7013-1.
- 17. Aspirate standards and samples. Record absorbance readings. NOTE: If the absorbance values for the samples are above the linear range of the standards, dilute the solutions with 10% HNO $_{\rm 3}$ and an appropriate amount of the 50 mg/mL Cs solution, reanalyze, and apply the appropriate dilution factor in the calculations.

CALCULATIONS:

- 18. Using the measured absorbances, calculate the corresponding concentrations (mg/mL) of aluminum
- in the sample, C _s, and average media blank, C _b, from the calibration graph.

 19. Using the solution volumes (mL) of the sample, V _s, and media blanks, V _b, calculate the concentration, C (mg/m ³), of aluminum in the volume of air sampled, V (L):

 $C \sim = \sim \{(C \text{ SUB s V SUB s} \sim - \sim C \text{ SUB b V SUB b})\} \text{ OVER V }, \sim \text{ mg/m SUP 3}.$