ETHYL BROMIDE

CH₃CH₂Br  MW: 108.97  CAS: 74-96-4  RTECS: KH6475000

METHOD: 1011, Issue 2  EVALUATION: PARTIAL


OSHA: 200 ppm
NIOSH: no REL
ACGIH: 5 ppm (skin); suspected carcinogen
       (1 ppm = 4.46 mg/m³ @ NTP)

PROPERTIES:
liquid; d 1.461 g/mL @ 20 °C;
BP 38 °C; MP -119 °C;
VP 51 kPa (380 mm Hg; 50% v/v)
@ 20 °C; explosive range 6.8 to
11% v/v in air

SYNONYMS: bromoethane

SAMPLING

SAMPLER: SOLID SORBENT TUBE
(coconut shell charcoal, 100 mg/50 mg)
FLOW RATE: 0.01 to 0.2 L/min
VOL-MIN: 0.5 L @ 200 ppm
         -MAX: 4 L
SAMPLE STABILITY: not known
FIELD BLANKS: 2 to 10 field blanks per set

MEASUREMENT

TECHNIQUE: GAS CHROMATOGRAPHY, FID
ANALYTE: ethyl bromide
DESORPTION: 1 mL 2-propanol; overnight
INJECTION VOLUME: 5 µL
TEMPERATURE-INJECTION: 200 °C
         -DETECTOR: 300 °C
         -COLUMN: 60 °C
CARRIER GAS: N₂ or He, 30 mL/min
COLUMN: 3 m x 3-mm OD stainless steel; 10%
         FFAP; on 80/100 mesh Chromosorb
         W-AW
CALIBRATION: standard solutions of ethyl bromide in
2-propanol
RANGE: 0.4 to 10 mg per sample
ESTIMATED LOD: 0.02 mg per sample [2]
PRECISION (S_r): 0.025 @ 1.8 to 8.5 mg per sample [1]

ACCURACY

RANGE STUDIED: 550 to 2200 mg/m³ [1]
(4-L samples)
BIAS: - 6.4%
OVERALL PRECISION (S_r): 0.054 [1]
ACCURACY: ± 15.0%

APPLICABILITY: The working range is 22 to 560 ppm (100 to 2500 mg/m³) for a 4-L air sample. Ethyl bromide is used in organic synthesis as a solvent and as a refrigerant, anesthetic, and grain and fruit fumigant.

INTERFERENCES: None reported. An alternate GC column is 30 m x 0.32-mm fused silica capillary coated with 0.5-µm DBWAX at 50 °C isothermal [2].

OTHER METHODS: This revises Method S106 [3].

REAGENTS:
1. 2-Propanol, chromatographic grade.*
2. Ethyl bromide, reagent grade.*
3. Calibration stock solution, 0.073 mg/µL. Dilute 0.5 mL ethyl bromide to 10 mL with 2-propanol.
4. Nitrogen or helium, purified.
5. Hydrogen, prepurified.
6. Air, filtered, compressed.

* See SPECIAL PRECAUTIONS.

EQUIPMENT:
1. Sampler: glass tube, 7 cm long, 6-mm OD, 4-mm ID, flame-sealed ends with plastic caps, containing two sections of activated (600 °C) coconut shell charcoal (front =100 mg; back = 50 mg) separated by a 2-mm urethane foam plug. A silylated glass wool plug precedes the front section and a 3-mm urethane foam plug follows the back section. Pressure drop across the tube at 1 L/min airflow must be less than 3.4 kPa. Tubes are commercially available.
2. Personal sampling pump, 0.01 to 0.2 L/min, with flexible connecting tubing.
3. Gas chromatograph, FID, integrator and column (page 1011-1).
4. Vials, glass, 2-mL, PTFE-lined caps.
5. Syringes, 10-µL and other convenient sizes for preparing standards, readable to 0.1 µL.
6. Volumetric flasks, 10-mL.
7. Pipets, 1- and 5-mL, with pipet bulb.

SPECIAL PRECAUTIONS: Ethyl bromide is flammable, a dangerous fire hazard (flash point = 26 °C), respiratory irritant, narcotic, and a hepato- and renal toxin [4].

2-Propanol is flammable and a dangerous fire risk with a flash point of 11 °C.

SAMPLING:
1. Calibrate each personal sampling pump with a representative sampler in line.
2. Break the ends of the sampler immediately before sampling. Attach sampler to personal sampling pump with flexible tubing.
3. Sample at an accurately known flow rate between 0.01 and 0.2 L/min for a total sample size of 0.5 to 4 L.

SAMPLE PREPARATION:
5. Place the front and back sorbent sections of the sampler tube in separate vials. Discard the glass wool and foam plugs.
6. Add 1.0 mL 2-propanol to each vial. Attach crimp cap to each vial.
7. Allow to stand overnight with occasional agitation.

CALIBRATION AND QUALITY CONTROL:
8. Calibrate daily with at least six working standards over the range 0.02 to 10 mg ethyl bromide per sample.
   a. Add known amounts of ethyl bromide or calibration stock solution to 2-propanol in 10-mL volumetric flasks and dilute to the mark. Make serial dilutions as necessary to obtain solutions in the range 0.02 to 10 mg ethyl bromide/mL.
b. Analyze together with samples and blanks (steps 11 and 12).
c. Prepare calibration graph (peak area vs. mg ethyl bromide).

9. Determine desorption efficiency (DE) at least once for each lot of charcoal used for sampling in the calibration range (step 8). Prepare three tubes at each of five levels plus three media blanks.
   a. Remove and discard back sorbent section of a media blank sampler.
   b. Inject a known amount (1 to 20 µL) of ethyl bromide or calibration stock solution directly onto front sorbent section with a microliter syringe.
   c. Cap the tube. Allow to stand overnight.
   d. Desorb (steps 5 through 7) and analyze together with working standards (steps 11 and 12).
   e. Prepare a graph of DE vs. mg ethyl bromide recovered.

10. Analyze three quality control blind spikes and three analyst spikes to ensure that the calibration graph and DE graph are in control.

MEASUREMENT:

11. Set gas chromatograph according to manufacturer’s recommendations and to conditions given on page 1011-1. Inject sample aliquot manually using solvent flush technique or with autosampler.
   NOTE: If peak area is above the linear range of the working standards, dilute with 2-propanol, reanalyze, and apply the appropriate dilution factor in calculations.

12. Measure peak area.

CALCULATIONS:

13. Determine the mass, mg (corrected for DE) of ethyl bromide found in the sample front \( (W_f) \) and back \( (W_b) \) sorbent sections, and in the average media blank front \( (B_f) \) and back \( (B_b) \) sorbent sections.
   NOTE: If \( W_b > W_f/10 \), report breakthrough and possible sample loss.

14. Calculate concentration, \( C \), of ethyl bromide in the air volume sampled, \( V \) (L):

\[
C = \frac{(W_f + W_b - B_f - B_b) \cdot 10^3}{V}, \text{ mg/m}^3
\]

EVALUATION OF METHOD:

Method S106 was issued on April 11, 1975 [3], and validated over the range 550 to 2200 mg/m \(^3\) using 4-L air samples [1]. The concentrations of ethyl bromide were independently determined by gas chromatographic analysis of the synthetic atmospheres using a 5-mL sampling loop. The average recovery of generated samples was 93.8% with a relative standard deviation of 6.3%. No storage stability study was done. The breakthrough volume was 6.4 L when dry air containing 2213 mg/m \(^3\) of ethyl bromide was sampled at 0.2 L/min. Desorption efficiencies averaged 0.84 over the range 1.8 to 7.1 mg per sample. At 0.03 mg per sample, a desorption efficiency of 0.63 was found [2].

REFERENCES:

METHOD REVISED BY:

Y. T. Gagnon, NIOSH/DPSE; S106 originally validated under NIOSH Contract CDC-99-74-45.