SOURCE TEST PROCEDURE ST-40

STRIPPABLE VOLATILE ORGANIC COMPOUNDS

REF: Regulation 11-10

1. APPLICABILITY

1.1 This procedure is used to quantify the concentration of volatile organic compounds stripped from cooling tower water by air. It is an adapted version of the Texas Commission on Environmental Quality (TCEQ) Air Stripping Method (Modified El Paso Method) for Determination of Volatile Organic Compound Emissions from Water Sources. This procedure includes requirements that should be used in addition to the TCEQ Modified El Paso Method and is not an alternative.

1.2 The use of a flame ionization detector (FID) analyzer for measurement of organic compounds is not allowed if the compound response factors are greater than 2, relative to methane. In order to confirm that compound response factors are less than 2, relative to methane and this is the appropriate measurement technique for specific systems, an initial screening of the process water must be performed and the results reported to the Source Test Manager prior to using a FID analyzer.

1.3 This procedure may be used for the measurement of volatile organic compounds with boiling points below 140°F. Procedures for the measurement of volatile organic compounds with boiling points of 140°F or greater must be approved by the Source Test Manager.

1.4 Any changes to the procedure that require BAAQMD Source Test and/or Lab Manager approval must also be approved by U.S. EPA if data is submitted for compliance with federal regulations.

2. PRINCIPLE

2.1 A continuous sample of cooling tower water is passed through a packed column. Air is passed through the column with countercurrent flow, stripping the volatile organic compounds from the water. The stripping air is analyzed for volatile organic compound concentration. Air and water input and output points have a direct interface to prevent volatile compound loss from vaporization in ambient air.

2.2 FID Direct Measurement: Stripping air is connected to a FID analyzer with a direct interface. The sample is analyzed for total volatile organic compounds.
2.3 Integrated Sample: Stripping air is routed at a constant, measured flow into an evacuated stainless steel canister or Tedlar® bag. The sample is analyzed for total volatile organic compounds by FID. Alternatively, the sample may be speciated by gas chromatography (GC) or an equivalent methodology approved by the Source Test Manager.

3. **RANGE AND SENSITIVITY**

3.1 The minimum measurable concentration of volatile organic compounds by a FID analyzer is 1 ppmv when interferences are not present. For the FID analyzer, the minimum detection limit should be 0.3 ppmv as methane. The minimum precision should be the larger of 0.3 ppmv or 5% of reading as methane. The minimum sensitivity should be 2% of full scale as methane.

3.2 The minimum measurable concentration of volatile organic compounds by GC is 0.5 ppmv.

4. **INTERFERENCES**

4.1 FID Analysis: Response factors vary amongst hydrocarbons. A FID should not be used on compounds with response factor multipliers greater than 2, relative to methane. Use of an alternative span gas or detection method must be approved by the Source Test Manager.

4.2 FID analyzers that use air from the sample or ambient sampling environment can have an elevated baseline. Analyzers using those air sources should be calibrated in the same location as sampling.

4.3 Rubber and drying agents can absorb volatile organic compounds. Use of those materials is prohibited.

4.4 The pressure drop across the rotameter control valve can cause gas bubble formation in the water rotameter and affect the flow measurement.

4.5 Temperature changes affect compound vapor pressures and can alter results.

5. **APPARATUS**

5.1 Clear, heavy walled glass column, 36 in long with an internal diameter of 3 in.

5.2 Packing material. Use 6 mm – 8 mm beryl saddles and pack the column to a depth of 26 in.
5.3 Neoprene stoppers.

5.4 Securing clamps.

5.5 Stainless steel or Teflon tubing, ¼ in diameter and no longer than 50 ft.

5.6 Air rotameter with ± 2% accuracy. Rotameter should be calibrated according to TCEQ Modified El Paso Method specifications.

5.7 Water rotameter with ± 2% accuracy. Rotameter should be calibrated according to TCEQ Modified El Paso Method specifications.

5.8 Type K temperature probe. Temperature probe should be calibrated according to TCEQ Modified El Paso Method specifications.

5.9 Clear glass flasks – 250 ml. Leave one flask empty for moisture knockout and fill one flask with water to act as a bubbler. The bubbler shows that there is sufficient flow out of the system and no external air being drawn in.

5.10 Flame ionization detector analyzer or gas chromatograph with appropriate detectors for sample compounds.

5.11 Tedlar® bags. Use bags that are a minimum of 20 L.

5.12 Evacuated stainless steel canisters. Sample canisters should meet all requirements in the U.S. EPA Compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air, Compendium Method TO-14A.

5.13 Zero-gas. Air is used for calibration and stripping. It should be certified to contain less than or equal to 0.1 ppmv total hydrocarbon (THC).

5.14 High-level calibration gas. Sample concentration should be between 10-100% of the high-level gas concentration. Calibration gas should be certified and within 2% of the stated concentration. Methane should be used unless a surrogate is approved by the Source Test Manager.

5.15 Mid-level calibration gas. Concentration should be 20-50% of the high-level calibration gas concentration and certified within 2% of the stated concentration. Methane should be used unless a surrogate is approved by the Source Test Manager.

5.16 Orifice plate, mass flow controller, or needle valve.

5.17 For more detail and diagrams of the apparatus, refer to Figures 1 and 2 in the TCEQ Modified El Paso Method.
Figure 40-1

Air Stripping Apparatus for Strippable VOCs from Water

Figure 1 from the TCEQ Modified El Paso Method.
Figure 2 from the TCEQ Modified El Paso Method.
6. **PRE-TEST PROCEDURES**

6.1 Select a sampling point that meets the site criteria outlined in the TCEQ Modified El Paso Method. Choose a point in the return line header prior to distribution to different cells and release to atmosphere. Water should be under pressure and drawn from the vertical section near the base of the riser pipe or from the top of the horizontal section prior to the riser.

6.2 Assemble the sampling system as shown in Figure 40-1 and Figure 40-2.

6.3 Cleaning procedures should follow those outlined in the TCEQ Modified El Paso Method.

6.4 Leak test the sampling system by introducing zero-air into the system at a flow rate of 2500 ml/min and spraying Snoop solution or equivalent onto connections. No bubbling should be observed before proceeding.

6.5 Allow sample water to flush through the sample line for a minimum of five sample line volumes before connecting the water line to the apparatus. Make sure there are no air bubbles in the water rotameter.

6.6 Level the apparatus to prevent channeling.

6.7 Warm up the instruments according to manufacturers’ instructions.

6.8 Introduce zero-air and span-gas into the analyzers and calibrate the instruments according to manufacturers’ instructions. The analyzer should read $0.0 \pm 0.2$ ppmv with zero-air as the input and within $\pm 5.0\%$ of the high-level and mid-level calibration gas certified values. The mid-level calibration gas is optional if the VOC emissions are determined by GC or GC/MS speciation.

7. **SAMPLING**

7.1 Set the air flow rate to 2500 ml/min.

7.2 Fill the column with water to the top of the beryl saddle packing. Set the water flow rate to 125 ml/min.

7.3 Allow the apparatus to stabilize for at least ten minutes. Record the stabilization time, barometric pressure, ambient temperature, and cooling tower process water flow rate.

7.4 Attach the FID analyzer to the air outlet via direct interface and ensure the sample air outlet does not make contact with ambient air. Sample for at least 10 minutes. Record the air flow rate, the water flow rate, and the chamber temperature every 2 minutes.
7.5 Alternatively, a sample canister may be attached to the air outlet. A critical orifice, needle valve, or mass flow controller should be used to control the flow into the canister. Set flow to 1/10th of the canister volume per minute or less. Open the canister valve.

7.6 Sample for at least 10 minutes. Record the air flow rate, the water flow rate, the chamber temperature, and the canister vacuum every 2 minutes.

7.7 When finished, close the canister valve. Record the final sample canister vacuum, sample collection time, and sample ID. Analyze stainless steel canister samples within 5 business days of sample collection.

7.8 Before analysis, record the sample canister vacuum to determine if leaking has occurred. Pressurize the canister according to specifications in U.S. EPA Compendium Method TO-14A. If condensables have the possibility of existing in the canister, heat the canister above the stripping chamber temperature. Connect the canister to the FID analyzer and open the valve. The sample flow rate should be the same flow rate used for calibration and be held constant throughout the analysis. Ensure that there is enough sample to reach a stable output.

7.9 Tedlar® bags may be used if quality assurance and quality control guidelines are followed as described in Title 40, Code of Federal Regulations, Part 60, Appendix A, Method 18 for all target compounds. The recovery study outlined in Method 18, Section 8.4.2 must be performed and results must be corrected for percent recoveries. Tedlar® bags must be new, unused, and checked for leaks and contamination. Control the flow rate going into the bag by attaching a needle valve to the bubbler outlet and controlling the bypass. Set the bypass so that flow to the Tedlar® bag is between 500 ml/min and 1000 ml/min. Analyze Tedlar® bag samples within 72 hours of sample collection.

7.10 Alternatively, the canister or Tedlar® bag may be sent to a lab for analysis by U.S. EPA Compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air, Compendium Method TO-14A or U.S. EPA Method 18, Measurement of Gaseous Organic Compound Emissions by Gas Chromatography. Results from analysis shall be reported in THC concentration if under the limit in BAAQMD Regulation 11-10-304. When speciation is required, results shall be reported for individual compound concentrations.

8. **AUXILIARY TESTS**

*Auxiliary tests provide supplementary or additional information that may be necessary to complete required calculations or the test procedure.*

8.1 Volatile organic compound speciation and concentration. Use U.S. EPA Compendium of Methods for the Determination of Toxic Organic Compounds in
Ambient Air, Compendium Method TO-14A or U.S. EPA Method 18, Measurement of Gaseous Organic Compound Emissions by Gas Chromatography to determine VOC speciation and concentration if any compounds have a response factor greater than 2, relative to methane.

9. **CALCULATIONS**

9.1 Use Equation 1 to calculate the concentration of air strippable compounds in the sample water matrix.

\[
C = \frac{M \times (P \times 0.03342) \times b \times c}{R \times (T + 273) \times a}
\]  

(1)

C = Concentration of air strippable volatile organic compounds in the water matrix, ppmw.

M = Molecular weight of the compound, g/mol.

P = Pressure in the column, in Hg. Typically assumed the same as atmospheric pressure.

b = Stripping air flow rate, ml/min.

c = Concentration of volatile organic compounds in the stripped air, ppmv.

a = Sample water flow rate, ml/min.

R = Gas Constant, 82.054 ml-atm/mol-K.

T = Column temperature, °C.

0.03342 = Conversion from in. Hg to atm, atm/in Hg.

273 = Conversion from Celsius to Kelvin.

9.2 Use Equation 2 to calculate the mass emission rate of volatile organic compounds.

\[
E = \frac{C \times F \times 60 \times 8.329}{1,000,000}
\]  

(2)

E = Mass emission rate of volatile organic compounds, lb/hr.
C = Concentration of air strippable volatile organic compounds in the water matrix, ppmw.

F = Water circulation rate of source, gal/min.

60 = Conversion from minutes to hours, min/hr.

8.329 = Specific weight of water at 70°F, lb/gal.

1,000,000 = Conversion from ppm, ppm.
10. REPORTING

Figure 10.1

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<td>San Francisco, California 94105</td>
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<tr>
<td>Requester</td>
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SUMMARY OF SOURCE TEST RESULTS

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ST – 40 Strippable Volatile Organic Compounds, ppmv

NO COMMERCIAL USE OF THESE RESULTS IS AUTHORIZED

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<th>Air Quality Engineer</th>
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Bay Area Air Quality Management District ST-40-10 05/18/16
11. REFERENCES

