EPA APTI Course #468
Laboratory Manual
DRAFT

Monitoring Compliance Test And Source Test Observation
Notice

This is not an official policy and standards document. The opinions and selections are those of the author and not necessarily those of the Environmental Protection Agency. Every attempt has been made to represent the present state of the art as well as subject areas still under evaluation. Any mention of products or organizations does not constitute endorsement by the United States Environmental Protection Agency (USEPA).

Usage of This Manual

The Central States Air Resource Agencies Association (CenSARA) is one of several multi-jurisdictional organizations (MJOs) operating for the U.S. Environmental Protection Agency (USEPA), through the Air Pollution Training Institute (APTI), to update more of the frequently used APTI courses. The primary objectives of the MJOs are to:

- Promote the exchange of information between the States;
- Serve as a forum to discuss regional air quality issues of common concern;
- Share resources for the common benefit of the member states; and
- Provide training services to their member air pollution control agencies.

APTI provides courses on air pollution control technology, ambient air and source monitoring, and air quality management. Historically, APTI designed courses that meet the job training needs of governmental agency personnel and others in the field of air pollution. This requires a thorough examination of both the materials for instruction and the characteristics of the student audience. Based on studies conducted by APTI of those who have participated in the various training courses, courses were developed and revised to provide training that enables every student to achieve specific course objectives. A basic goal of APTI was to provide training that will enable a student to do specific jobs in his or her home environmental agency. However, recently APTI has taken a new direction and has given money to the various MJOs, of which CenSARA is one, to update needed training course for their member states.

CenSARA meets these training needs of its member states by identifying, designing, developing and delivering needed, cost-effective, responsive, and focused educational opportunities for state and local air agency staff. Agenda and course materials are obtained from a variety of sources including EPA, colleges and universities, regional training consortia, and individual instructors. Yet, due to changes in environmental regulations, the implementation of new policies, and the advancement of technologies, agendas and course materials become out-of-date. When this happens, staffs' ability to enhance skills, knowledge and abilities are constrained, limiting their ability to excel in the dynamic field of air pollution control. So by providing up-to-date, high quality educational opportunities for staff, their chances to greatly enhance their skills, knowledge and abilities is significantly improved.
Consequently, CenSARA announced a Request for Proposals (RFP) to the environmental training community to solicit technical proposals and cost bids to review current compliance test and observation programs within the USEPA and to update as necessary the content title, agenda topics, course length, instructor and student manuals, lectures involving presentation slides, classroom and homework exercises, and other handouts and materials for EPA’s APTI Course 468 entitled: “Monitoring Compliance Test and Source Test Observation.” In response to CenSARA’s RFP, EnviroTech Solutions, William T. “Jerry” Winberry, 1502 Laughridge Drive, Cary, North Carolina 27511, jwinberry@ mindspring.com, 919-467-2785, was awarded the contract to update EPA’s APTI Course #468. Mr. Winberry is the author of this Laboratory Manual and every attempt has been made to represent the most recent advances in sampling and analytical methodology.

This material has been developed and assembled to provide training associated with EPA’s APTI Course #468 entitled: “Monitoring Compliance Test and Source Test Observation.” It is not intended to be used for regulatory purposes, or to be a substitute for, nor interpreted as official Agency policy. Every attempt was made to reflect the technical state of art and regulatory information as of the date of publication.
Table of Contents

Notice and Usage of this Material ................................................................. i
Table of Content ...................................................................................................... ii
To the User of This Material and Description of Laboratory Stations ................. iii
Laboratory Station 1: Calibration of FRM 5 Sampling Nozzle ......................... 1-1
Laboratory Station 2: Dry Gas Meter "Gamma" Calibration ............................. 2-1
Laboratory Station 3: Orifice Meter Calibration Involving ΔH@ ...................... 3-1
Laboratory Station 4: Stack Gas Velocity and Volumetric Flow Rate .......... 4-1
Laboratory Station 5: Source Simulator/Pitot Tube Calibration ....................... 5-1
Laboratory Station 6: Source Simulator/Stack Gas Moisture ......................... 6-1
Laboratory Station 7: Type S Pitot Tube Inspection ........................................... 7-1
Laboratory Station 8: Federal Reference Method 5 Sampling Train ................. 8-1
Laboratory Station 9: IsoCal Electronic Spreadsheet ........................................ 9-1
Laboratory Station 10: FRM 1 Traverse Point Determination ......................... 10-1

Appendices

Appendix A: Abbreviations and Terminology .................................................. A-1
Appendix B: Equations ......................................................................................... B-1
Appendix C: Inspector's Tool Kit ....................................................................... C-1
Introduction

To The User of This Manual and Description of Laboratory Stations

This laboratory manual (LM) has been prepared to guide the students through a series of laboratory exercises developed to provide the students a hands-on experience related to stack testing issues and concepts discussed in the lecture portion of this course. The primary objective of these exercises is for the students to gain a practical understanding of common procedures outlined in the Federal Register necessary to monitor compliance test and source test observation for determining compliance with the applicable emission regulations. This LM provides guidance and materials for completing ten (10) laboratory stations utilizing a source simulator to reinforce lecture objectives. Each station has specific objectives to be accomplished. The ten laboratory stations are:

- **Laboratory Station 1**: Calibration of Nozzle for FRM 5 Train
- **Laboratory Station 2**: Dry Gas Meter “Gamma” Calibration
- **Laboratory Station 3**: Orifice Meter Calibration Involving ΔH@ of FRM 5 Train
- **Laboratory Station 4**: Stack Gas Velocity and Volumetric Flow Rate
- **Laboratory Station 5**: Source Simulator/Pitot Tube Calibration
- **Laboratory Station 6**: Source Simulator/Stack Gas Moisture
- **Laboratory Station 7**: Type S Pitot Tube Inspection
- **Laboratory Station 8**: Federal Reference Method 5 Sampling Train
- **Laboratory Station 9**: IsoCal Electronic Spreadsheet/Isokinetic Rate Equation
- **Laboratory Station 10**: FRM 1 Traverse Point Determination

The 10 laboratory simulator stations focus on source testing activities and lecture objectives presented in the classroom involving stack gas velocity, sampling site location, determination of stack gas molecular weight, determining stack gas moisture etc. This LM contains step-by-step instructions to the participant on how to accomplish the objectives of each station exercise. Instrumentation used during each Laboratory Station exercise can be obtained from the Inspector’s Tool Kit (see Appendix C). A brief explanation for each laboratory station follows.

**Laboratory Station 1: Nozzle Diameter (D_n)**

The objective of Laboratory Station 1 is to demonstrate to the participant the difficulty in measuring a proper nozzle diameter. The participant will be given data to satisfy the estimated probe nozzle diameter equation for an upcoming stack test. Based upon the calculation, the participant will measure several nozzles using a micrometer acquired from the Inspector’s Tool Kit (see Appendix C), document the findings on standardized forms, and select the correct nozzle for the upcoming test. Probe nozzles should be inspected and calibrated in the field immediately before each use to verify that they were not damaged in transport or shipment to the test site.
Laboratory Station 2: DGM "γ" Determination

Test results from a stack emission test are meaningless without calibration of the equipment and its components. The dry gas meter (DGM) serves to record the volume of gas sampled during the test and the orifice tube on the outlet of the DGM serves to provide flow rate determination during testing. Both of these components must be calibrated. The DGM "γ" value is a calibration factor that relates volume from a primary standard (i.e., wet test meter) to the recorded dry gas meter volume. The objective of Laboratory Station 2 is to use a set of calibrated orifices obtained from the Inspector’s Tool Kit (see Appendix C) to determine the "γ" of a meter box assembly containing DGM. Once again, standardized forms are used to complete the assignment.

Laboratory Station 3: Orifice Meter "ΔH@" Determination

Test results from a stack emission test are meaningless without calibration of the equipment and its components. The dry gas meter (DGM) serves to record the volume of gas sampled during the test and the orifice tube on the outlet of the DGM serves to provide flow rate determination during testing. Both of these components must be calibrated. The orifice tube calibration factor "ΔH@" is the pressure drop across the orifice for a typical sampling flow rate of 0.75 which is the standard sampling rate for solving the isokinetic equation and setting up the nomographs (sets of equations) for testing. The objective of Laboratory Station 3 is to use a set of NIST calibrated orifices to determine the orifice tube calibration factor "ΔH@". The set of NIST calibrated orifices can be obtained from the Inspector’s Tool Kit (see Appendix C). Once again, standardized forms are used to complete the assignment.

Laboratory Station 4: Source Simulator/Stack Gas Velocity (v_s) and Volumetric Flow Rate (Q_s)

Laboratory Station 4 involves the participant to use an "S-type" pitot tube to determine simulated stack gas flow rate. In determining the stack gas flow rate, the pitot tube must be constructed to specifications identified in Federal Reference Method 2 (FRM 2), have a known pitot tube coefficient factor (C_p), and positioned in the source using proper orientation. Laboratory Station 4 will require the participant to verify that the pitot tube meets geometric specifications using a micrometer and standardized data sheet in order to assign a known C_p. In addition, Laboratory Station 4 requires the participant to divide the stack into equal area identified in Federal Reference Method 1 (FRM 1), mark the pitot tube/probe for each sample point, and determine the stack gas velocity at each of the sampling points in the centroid of the equal areas. A Type S pitot tube along with measurement tape can be acquired from the Inspector’s Tool Kit (see Appendix C). Once
Introduction

again, a standardized field test data sheet (FTDS) will be available to complete the assignment.

**Laboratory Station 5: Source Simulator/Calibration of Type S Pitot Tube**

As identified in Federal Reference Method 2, a Type S pitot tube can be calibrated using a standard type pitot tube. The objective of Laboratory Station 6 is to follow standardized procedures provided in order to calibrate a Type S pitot tube using a standard pitot tube. Entry points will be provided for inserting the Type S and standard pitot tube into the source simulator. The Type S and Standard pitot tube along with a digital manometer can be acquired from the Inspector’s Tool Kit (see Appendix C). Once again, standardized calibration forms are provided to complete this assignment.

**Laboratory Station 6: Source Simulator/Stack Gas Moisture**

One of the most important parameters to estimate correctly when selecting your nozzle diameter and setting up your isokinetic rate equation is the stack gas moisture ($B_{WS}$). For every % stack gas moisture you estimate incorrectly will cause your % isokinetics to be off the same %. The objective of Laboratory Station 6 is to determine the stack gas moisture content of the source simulator using the wet bulb/dry bulb technique. Obtain a Type K thermocouple along with the digital readout instrument from the Inspector’s Tool Kit (see Appendix C). Once again, standardized forms are provided to assist in the calculations, in addition, other techniques for determining stack gas moisture will be reviewed.

**Laboratory Station 7: Pitot Tube Inspection**

In order to assign a known pitot tube coefficient factor, $C_p$, the pitot tube must meet certain design and construction requirements as identified in Federal Reference Method 2 (FRM 2) in order to assign a value of 0.84. To complete this exercise, you will be given a Type S pitot tube along with a “bulls eye” and level indicator out of the Inspector’s Tool Kit (see Appendix C). Using the standardized data form, inspect the Type S pitot tube and see if it meets specifications by evaluating $\alpha$, $\beta$, $z$, $w$ and $A$ characteristics so a $C_p$ of 0.84 can be assigned to the Type S pitot tube.

**Laboratory Station 8: Federal Reference Method 5 (FRM 5) Sampling Train**

Laboratory Station 5 contains a complete Federal Reference Method 5 (FRM 5) sampling train, however unassembled. The task of Laboratory Station 5 is for the participant to completely assemble the FRM 5 train, including attaching a nozzle, adding silica gel and water to the impingers, and inserting a filter to the assembly. Once completed, the
Introduction

participant is to turn on the pump and leak check the complete FRM 5 sampling train to a leak rate <0.2 cfm. After leak check, the participant positions the FRM 5 sampling train at the sampling port of the “closed-looped source simulator) and commence sampling. Sampling of the source simulator gas stream should begin while traversing the source simulator diameter until at least 15 standard cubic feet have been extracted from the source through the assembled FRM 5 sampling train.

Laboratory Station 9: Isokinetic Rate Equation and Calculations

One of the major activities associated with FRM 5 is the setting and maintaining isokinetic rate conditions during a test. One can use either a nomograph to assist with this or use of applicable software. The objective of Laboratory Station 9 is to demonstrate the use of the IsoCal software spreadsheet in completing the necessary equations for proper operation of the FRM 5 sample train during sampling. The participant will be given source test data that should be entered into the lap top computer containing the IsoCal software. Based upon the data input, the participant will determine point-by-point isokinetics and average isokinetics. In addition, the participant will be able to see the variability in isokinetics as one changes the various input parameters (i.e., nozzle diameter, moisture content of stack gas etc.).

Laboratory Station 10: FRM 1 Traverse Point Determination

Federal Reference Method 1 requires that the number of traverse points for a given test be based upon the sampling port locations with reference to upstream and downstream flow disturbances. The objective of Laboratory Station 10 is to determine the number and the location of traverse points at your assigned sampling site. A measurement tape can be obtained from the Inspector's Tool Kit (see Appendix C). Once again, you will be provided with standardized procedures to assist with this task.

Source Simulator

Many of the laboratory stations outlined above require the use of a “source simulator” to complete laboratory objectives. APTI Course #468 utilizes a “closed-loop source simulator” to re-enforce lecture objectives involving Federal Reference Methods (FRMs) 1, 2, 3, 4, and 5. The source simulator is constructed of 12” diameter galvanized air conditioning duct material configured in a closed-loop containing a 360 watt 12” blower. Ambient air is circulated within the closed-loop by the blower system. Port holes are positioned throughout the duct work to provide access to the sample gas stream. Figure 1 documents the closed-loop source simulator.
Introduction

Figure 1. APTI Course #468 Closed-Loop Source Simulator

The use of the closed-loop source simulator allows each participant to acquire "field experience" associated with many of EPA's FRMs and how to observe their use to quantify emissions from stationary sources.
Nozzle Diameter (Dₙ)

The objective of Laboratory Station 1 is to demonstrate to the participant the difficulty in measuring a proper nozzle diameter. Three (3) different Federal Reference Method 5 (FRM 5) probe nozzles are provided as part of this laboratory station. The objective is to measure the nozzle diameters (Dₙ) of the three nozzles using a micrometer, document the findings on the standardized calibration data sheet, and determine if the nozzles meet the requirements specified in FRM 5. The micrometer can be obtained from the Inspector’s Tool Kit (see Appendix C). Probe nozzles should be inspected and calibrated in the field immediately before each use to verify that they were not damaged in transport or shipment.
Laboratory Station 1: Calibration of Sampling Nozzles

☐ Obtain three (3) different probe nozzles for this laboratory exercise. You are to calibrate three (3) nozzles.
☐ Before starting the calibration check, obtain the Probe Nozzle Diameter Calibration Data Sheet, Laboratory Station 1 Worksheet.
☐ Inspect the nozzle for nicks, dents and corrosion. If these are found, they should be corrected before calibration. Record the individual nozzle identification number for each nozzle on the Probe Nozzle Diameter Calibration Data Sheet.
☐ Place a reference mark on the nozzle. Place the nozzle at the center of the figure, as illustrated in Figure 1-1 (see also Calibration Data Sheet) and aligned reference mark with point P1. Mark P1 position on outside of the nozzle aligned with the diameter line for P1.

![Diagram of measurement plane]

**Figure 1-1. Measurement Plane for Measuring Inside Diameter of Nozzle**

☐ Using a vernier or dial calipers obtained from the Inspector’s Tool Kit (see Appendix C) with at least 0.025 mm (0.001 inch) tolerance, measure the inside diameter (Dn) of the nozzle from the reference mark to P1, as illustrated in Figure 1-2.
Figure 1-2. Illustration of Using Dial Caliper for Measuring Inside Diameter of Nozzle

[NOTE: Remember, the caliper must be zeroed and measurement should be just touching the inside of the nozzle.]

☐ Record your reading on the Probe Nozzle Diameter Calibration Data Sheet.

☐ Now place the same nozzle at the center of the figure once again, but this time rotate the reference point until it is at P2. Mark P2 on opposite outside of nozzle aligned with the diameter line for P2.

☐ Once again, using a vernier or dial caliper with at least 0.025 mm (0.001 inch) tolerance, measure the inside diameter of the nozzle. Remember, the calipers should just touch the inside of the nozzle.

☐ Record your reading on the Prober Nozzle Diameter Calibration Data Sheet.

☐ Now place the same nozzle at the center of the figure once again, but this time rotate the reference point until it is at P3. Mark P3 on opposite outside of nozzle aligned with the diameter line for P3.

☐ Once again, using vernier or dial caliper with at least 0.025 mm (0.001 inch) tolerance, measure the inside diameter of the nozzle. Remember, the calipers should just touch the inside of the nozzle.

☐ Record your three readings on the Probe Nozzle Diameter Calibration Data Sheet.

☐ Calculate the average ($D_{avg}$) of the three readings and the $\Delta D$ of the nozzle and record on the Probe Nozzle Diameter Calibration Data Sheet.

☐ Repeat the above procedure using two additional nozzles.

☐ If the individual readings for each nozzle do not fall within 0.1 mm (0.004 inches) of one another, the nozzle must be reshaped, re-sharpened and recalibrated.

☐ Sign and date the Probe Nozzle Diameter Calibration Data Sheet, Laboratory Station 1 Worksheet.
**Laboratory Station 1 Worksheet**

**PROBE NOZZLE DIAMETER CALIBRATION DATA SHEET**

<table>
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<th>Date</th>
<th>Calibrated By</th>
<th>Nozzle ID #</th>
<th>Nozzle Diameter (mm)</th>
<th>Hi - Lo</th>
<th>ΔD</th>
<th>D_{avg}</th>
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<td></td>
<td></td>
<td>D1</td>
<td>D2</td>
<td>D3</td>
<td></td>
</tr>
</tbody>
</table>

Where:

- D1, D2, D3 = Three different nozzle diameters at 60 degrees to each other, each measured the nearest 0.025 mm.
- ΔD = Maximum distance between any two diameters, must be ≤ 0.100 mm.
- D_{avg} = (D1 + D2 + D3) / 3

**INSTRUCTIONS**

1. Inspect the nozzle for nicks, dents and corrosion. If these are found, they should be corrected before calibration.
2. Place a reference mark on the nozzle. Place the nozzle at the center of figure, aligned with point P1. Measure and record D1.
3. Rotate the nozzle so that the reference mark is aligned with point P2. Measure and record D2.
4. Rotate the nozzle so that the reference mark is aligned with point P3. Measure and record D3.
5. Calculate ΔD and D_{avg}.

Checked By: ____________________________ Personnel (Signature/Date)  
Team Leader (Signature/Date)
Dry Gas Meter (DGM) "γ" and "ΔH@" Determination

Test results from a stack emission test are meaningless without calibration of the components of the Federal Reference Method 5 (FRM 5) equipment and its components. The dry gas meter (DGM) serves to record the volume of gas sampled during the test and the orifice tube on the outlet of the DGM serves to provide flow rate determination during testing. Both of these components must be calibrated. The DGM "γ" value is a calibration factor that relates volume from a primary standard (i.e., NIST traceable critical orifices) to the recorded dry gas meter volume. The objective of Laboratory Station 2 is to use a set of calibrated orifices from the Inspector’s Tool Kit (see Appendix C) to determine the "γ" of a meter box assembly containing the DGM. Once again, standardized forms are used to complete the assignment. The orifice tube calibration factor "ΔH@" is the pressure drop across the orifice for a sampling flow rate of 0.75 which is the standard sampling rate for solving the isokinetic equation and setting up the nomographs (sets of equations) for testing. The objective of Laboratory Station 3 is to use the same set of calibrated orifices to determine the orifice tube calibration factor "ΔH@". The Laboratory Station 2 and 3 Worksheet is provided at the end of this section.
Laboratory Stations 2 & 3

APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

Laboratory Station 2 & 3: Dry Gas Meter (DGM) “γ” Determination and “ΔH@” Determination

☐ Position your self in front of the Federal Reference Method 5 (FRM 5) meter console. Insure that the orange oil manometer (ΔH manometer) has been leveled and zeroed.

☐ Using the power toggle switch, turn on the meter box pump and adjust the ΔH using the coarse and fine adjust knobs to read~ 1.5 inches of water as illustrated by the orange oil manometer. Allow the meter box to operate for ~ 5 minutes to allow the system to warm up.

☐ Obtain Method 5 Dry Gas Meter Calibration Using Critical Orifices Form (see attached Calibration sheet) and record on the form the date, meter console number, DGM serial number, critical orifice set serial number, barometric pressure, and ambient temperature.

☐ Obtain from the Inspector’s Tool Kit (see Appendix C) the NIST traceable Calibration Orifice Set (which contains five NIST-traceable orifices) along with the manufacturer’s certification sheet (see Manufacturer Supplied Orifice Calibration Spreadsheet Example) for the five NIST-traceable orifices, as illustrated in Figure 2-1.

Figure 2-1. NIST-Traceable Calibration Orifice Set

Select one of the NIST-traceable orifices from the set to be used in the calibration of the DGM. Under the “Orifice #” column on the Laboratory Station 2 and 3 Worksheet, Method 5 Dry Gas Meter Calibration Form, record the assigned orifice number as indicated from the manufacturer’s certification sheet.

2/3-2
From the manufacturer’s certification sheet obtain for the selected NIST-traceable critical orifice the “Average K Factor Value” and record under the “K’ Factor (Avg)” column on the Laboratory Station 2 & 3 Worksheet, Method 5 Dry Gas Meter Calibration Form.

After ~5 minutes, turn off the pump.

Insert the selected NIST-traceable critical orifice male quick connect into the inlet (SAMPLE) of the Method 5 meter console, as illustrated in Figure 2-2.

Figure 2-2. NIST-Traceable Critical Orifice Inserted Into Inlet of Method 5 Meter Console

Record the Dry Gas Meter (DGM) initial reading on the Laboratory Station 2 & 3 Worksheet, Method 5 Dry Gas Meter Calibration Form, under the column “DGM Readings: Initial” along the row for Run #1. Also record ambient temperature along the row for Run #1. Set the Method 5 meter console elapsed timer to zero.

Turn on the pump and elapsed timer at the same time. Adjust the coarse and fine valves on the Method 5 meter console until a vacuum of -18 inch of mercury is indicated on the Method 5 meter box vacuum gauge. Recorded the vacuum on the Laboratory Station 2 and 3 Worksheet, Method 5 Dry Gas Meter Calibration Form under the column labeled “Test Vacuum (in. Hg).”

Continue operating the meter console for a period of 5 minutes so you achieve a minimum total volume of 5 cubic feet through the DGM for Run #1. Record ambient temperature, DGM inlet and outlet temperatures, and “ΔH” on the Laboratory Station 2 & 3 Worksheet, Method 5 Dry Gas Meter Calibration Form during the 5-minute run.

At the end of the elapsed time for sampling 5 cubic feet, stop the pump.

Record the final DGM reading on the Laboratory Station 2 & 3 Worksheet, Method 5 Dry Gas Meter Calibration Form.

Remove the first NIST-traceable orifice from the meter console and select a second NIST-traceable critical orifice and repeat the above steps.
If time permits, select your third critical orifice and once again repeat the above steps.

Using the equations found on bottom of the Laboratory Station 2 & 3 Worksheet, Method 5 Dry Gas Meter Calibration Form, calculate the \( V_m(\text{std}) \), \( V_c(\text{std}) \), \( \Delta H @ \) associated with this Method 5 meter box.
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<th>THEORETICAL CRITICAL VACUUM (in. Hg)</th>
<th>DRY GAS METER READING</th>
<th>DRY GAS METER AVG.</th>
<th>TEMP. (deg. F)</th>
<th>METER DELTA H (in. Hg)</th>
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</table>

I certify that orifice set number [J40-73] was tested in accordance with the US EPA Method 5 standards.

See the Code of Federal Regulations, title 40, part 60, Appendix A, Method 5, Item 7.3.

Signature / Name / Date 3-2-00
METHOD 5 DRY GAS METER CALIBRATION USING CRITICAL ORIFICES

1) Select three critical orifices to calibrate the dry gas meter which bracket the expected operating range.
2) Record initial manometer pressure before and after calibration procedure.
3) Run test on least vacuum (from Critical Calibration Report), for a period of time necessary to achieve minimum initial volume of 9 cubic feet.
4) Record readings in outlined boxes below, other columns are automatically calculated.

<table>
<thead>
<tr>
<th>CRITICAL ORIFICE S#</th>
<th>TESTED ORIFICE S#</th>
<th>CRITICAL ORIFICE SET SERIAL</th>
<th>BAROMETERIC PRESSURE (IN Hg)</th>
<th>INITIAL</th>
<th>FINAL</th>
<th>AVG Pราช</th>
<th>IF VARIATION EXCEEDS 2.0%, ORIFICE SHOULD BE RECALIBRATED</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
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<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>2</td>
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<tr>
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</tr>
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</table>

USING THE CRITICAL ORIFICES AS CALIBRATION STANDARDS:
The following equations are used to calculate the standard volume of gas passed through the DGM, Vm (std), with critical orifice, Vc (std), and the DGM calibration factor, Y. These equations are automatically calculated in the spreadsheet.

\[
1) \quad V_m (\text{std}) = K \cdot V_c \cdot \frac{P_{\text{meas}} + (15 \times 1.3)}{293} \\
2) \quad V_c (\text{std}) = K \cdot \sqrt{T_m - T_0} \\
3) \quad Y = \frac{V_m (\text{std})}{V_c (\text{std})} = \text{DGM calibration factor}
\]

AVERAGE DRY GAS METER CALIBRATION FACTOR, Y =

\[
\text{AVERAGE } d_H_g = \frac{V_m (\text{std})}{V_c (\text{std})}
\]
Determination of Flue Gas Velocity ($v_s$) and Volumetric Flow Rate ($Q_s$)

The objective of Laboratory Station 4 requires the participant to divide the stack into equal areas as instructed in Federal Reference Method 1 (FRM1), mark the Type S pitot tube/probe for each sample point, and determine the stack gas velocity and volumetric flow rate at each of the sampling points in the centroid of the equal areas. The port associated with the source simulator for velocity and volumetric flow rate determination will be used. Obtain the measurement tape, Type S pitot tube, digital or water manometer, and marking liquid from the Inspector's Tool Kit (see Appendix C).
APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

Laboratory Station #4: Determination of Flue Gas Velocity ($v_s$) and Volumetric Flow Rate ($Q_s$)

☐ Obtain a Type-S pitot tube with a known $C_p$ and a Method 5 meter box console containing a water manometer or a digital manometer from the Inspector's Tool Kit. Examine the Type S pitot tube top, side and end views to verify that the face openings of the tube are aligned within specifications identified in Federal Reference Method 2 (FRM 2).

☐ Level and zero the Method 5 meter box water manometer identified for use in determining stack gas velocity and volumetric flow rate or zero the digital manometer.

☐ Using the standard Method 5 umbilical cable, connect the Type-S pitot tube to the umbilical cable using the quick disconnects. Connect the other end of the umbilical cable to the pitot quick connects on the Method 5 meter box console, as illustrated in Figure 4-1.

Figure 4-1. Type-S Pitot Tube Connected to FRM 5 Meter Console Containing Manometer

[Note: One should hear a “snap” when connecting quick connects to have a leak-free connection.] If not using a Method 5 umbilical cable, then connect ¼” Teflon lines to the Type-S pitot tube and opposite end to an independent digital manometer or the Method 5 meter box console using the pitot tube quick connects.

☐ Mark the impact side of the Type-S pitot tube as $P_A$ and the other face opening as $P_B$.

☐ Leak-check the $P_A$ side of the Type-S pitot tube. Attach a small piece of rubber tubing to cover the opening of the $P_A$ side (i.e., impact side) of the Type-S pitot tube. Blow into the rubber tubing until at least 3 inches of water velocity head registers on the manometer; then, close off
the impact opening. The pressure shall remain stable for at least 15 seconds. Do the same for the static pressure side, except using suction to obtain the minimum of 3 inches water.

☐ Retrieve the data for determination of traverse point locations for the source simulator as outlined in Laboratory Station 10. Using a measurement tape and marker, mark the determined traverse point distances on the Type S pitot tube.

☐ Position the Type-S pitot tube impact side (i.e., \( P_A \) side) facing the impact of the gas stream in the source simulator at Traverse Point #1 (i.e., traverse point closest to the source simulator wall). Be sure to cover the port hole with material to prevent air entering the source simulator and affecting data.

☐ Observe the pitot tube manometer reading (i.e., \( \Delta p \)) on the Method 5 box console or digital manometer and record value on the Laboratory Station 4 Worksheet, Flue Gas Velocity Data Sheet.

☐ Move the pitot tube to the next point, obtain a manometer reading (i.e., \( \Delta p \)), and record the results on the Laboratory Station 4 Worksheet, Flue Gas Velocity Data Sheet.

☐ Repeat the above steps for all of the traverse point for a diameter. [Note: Due to the source simulator construction, only one diameter will be utilized during this laboratory exercise.]

☐ At the last traverse point, move the pitot tube to traverse point #3 and record static pressure. Then remove pitot tube from the source simulator and cover the port hole.

☐ Complete the Laboratory Station 4 Worksheet, Flue Gas Velocity Data Sheet by taking the square root of each individual \( \Delta p \)s and average the individual square root \( \Delta p \)s.

☐ Calculate the source simulator flue gas velocity (\( v_g \)) and volumetric flow rate (\( Q_v \)) utilizing the equations provided. Finally, complete the Laboratory Station 4 Worksheet found at the end of this laboratory.
## Laboratory Station 4 Worksheet

### Flue Gas Velocity Data Sheet

<table>
<thead>
<tr>
<th>Traverse Point</th>
<th>$\Delta p$</th>
<th>$\sqrt{\Delta p}$</th>
<th>$(\sqrt{\Delta p})_{avg}$</th>
</tr>
</thead>
</table>
Calculations

Flue Gas Velocity Equation

\[ v_s = K_p \times C_p \times (\sqrt{\Delta p})_{avg} \times \frac{T_{(avg)}}{P_s \times M_s} \]

Where:
- \( v_s \) = average velocity of the gas stream, ft/sec
- \( T_s \) = absolute temperature, °R (°F + 460).
  (Use dry bulb temperature from Laboratory Station 6.)
- \( P_s \) = absolute pressure, in. Hg (Assume \( P_{bar} \) for sampling location and record static pressure of source simulator.)
- \( \Delta p \) = velocity pressure, in. H2O
- \( K_p \) = constant: 85.49,
  in units of \([\text{ft}^2/\text{in. Hg}(\text{lbf/lbs-mole})/(\text{sec}^2)(\text{in. H2O})(°R)]\]
- \( C_p \) = pitot tube coefficient, dimensionless
- \( M_s \) = stack gas molecular weight
  = \( M_d(1-B_{ws}) + 18 \times B_{ws} \)
  (Since the source simulator uses ambient air, assume \( M_d = 29.0 \))
- \( B_{ws} \) = the average moisture fraction by volume of the gas stream
  (dimensionless). Use the value determined in Station #6 (~ 1-2 %).

\[ v_s = K_p \times C_p \times ( \_ ) \times ( \_ ) \times \frac{\sqrt{T_{(avg)}}}{P_s \times M_s} \]

Volumetric Flow Rate Equation

There are two equations used for calculating the volumetric flow rate of a flow gas. One determines the flow rate at actual stack temperature and pressure conditions. The other determines the volumetric flow rate corrected to standard conditions of 68 °F and 29.92 in. Hg. Both are important.

Actual Stack Gas Volumetric Flow Rate

\[ Q_a = 60 \times v_s \times A_s \]

Where: \( A_s \) = area of stack = 3.14(Diameter/2)^2

\[ Q_a = 60 \times ( \_ ) \times ( \_ ) \]
Stack Gas Volumetric Flow Rate Corrected to Standard Conditions

\[ Q_{sd} = 3600 \times (1 - B_{ws}) \times v_s \times A_s \times \frac{T_{std}}{T_s} \times \frac{P_s}{P_{std}} \]

Where: 
- \( Q_s \) = the volumetric flow rate of the gas stream on a dry basis at standard conditions (scfh)
- 3600 = conversion factor (3600 sec/hr)
- \( B_{ws} \) = the average moisture fraction by volume of the gas stream (dimensionless). Use the value determined in Station #6 (~ 1-2 %).
- \( T_s \) = absolute temperature, °R (°F + 460)
- \( P_s \) = absolute pressure, in. Hg (Assume \( P_{bar} \))
- \( T_{std} \) = the absolute temperature at standard conditions, (528 °R)
- \( P_{std} \) = the absolute pressure at standard conditions, (29.92 in. Hg)

\[ Q_{sd} = 60 \times (1 - ( ) ) \times ( ) \times ( ) \times \frac{T_{std}}{P_{std}} \]
Calculation Summary for Laboratory Station 4 Worksheet

1. No of traverse points
2. \((\sqrt{\Delta p})_{avg}\)
3. \(C_p\)
4. \(T_s(avg)\)
5. \(P_s\)
6. \(M_s\)
7. \(V_s(avg)\)
8. \(A\)
9. \(Q_s\)
10. \(Q_a\)
Type S Pitot Tube Calibration ($C_p$)

As identified in Federal Reference Method 2 (FRM 2), a Stausscheibe type (i.e., Type S) pitot tube, if used for compliance determination, must be calibrated using a standard type pitot tube. The objective of Laboratory Station #5 is to follow standardized procedures as outlined in FRM 2 to calibrate a Type S pitot tube using a standard pitot tube in developing the Type-S pitot tube $C_p$ factor. Depending upon the equipment available, you may be using either a bare Type S pitot tube or a pitot tube attached to a Method 5 sampling probe. In either case, determine the $C_p$ for both the A and B sides of the Type S pitot tube. Entry points will be provided for inserting the Type S and standard pitot tube into the source simulator. Once again, a Laboratory Station 5 Worksheet is provided to summarize your laboratory findings.
Laboratory Station 5:

Calibration of Type S Pitot Tube

- Locate the pitot tube calibration ports on the source simulator.
- At the test station, note two holes in the ductwork, as illustrated in Figure 5-1.

![Diagram of pitot tube calibration](image)

Figure 5-1. Two Sampling Ports for Insertion of Type S and Standard Pitot Tubes

[Note: The sampling port for the standard pitot tube is downstream of the port for the Type S pitot tube. The port for the standard pitot tube should be located in such a manner that the tip of the standard pitot tube will be measuring the gas pressure at the same location in the stack gas as the impact side of the Type S pitot tube.]

- If using incline manometers for documenting pressure drops across the Type S and standard pitot tubes, then set-up, level, and zero the inclined manometer. Make sure that any valves on the manometer are in the "open" position. If you are using an electronic manometer, then zero the electronic manometer without tubes attached.
- Label one leg of the Type S pitot tube as "A" and the other leg as "B."
- Determine the distance from the center of the duct to the outside wall of the duct. Insert the Type S pitot tube so that the center of the impact tube is located at the center of the duct. Place a mark on the tube such that when the mark is placed at the outside edge of the duct, the tip of the Type S pitot tube is at the center of the duct.
- Repeat the same steps for the standard pitot tube.
- Connect the Type S pitot tube to the manometer on the Method 5 meter console or to the electronic manometer by way of umbilical cord or Teflon tubing. Leak check the Type S pitot tube as performed in Laboratory Station 4.
Insert the Type S pitot tube into the duct until the mark is at the outside edge of the duct wall, as illustrated in Figure 5-2.

Align the Type S pitot tube so that the "A" leg faces directly into the flow stream. Tape the port hole with duct tape to minimize air leakage into the duct.

On the Laboratory Station 5 Worksheet, Calibration of Type S Pitot Tube, record the Δp, in inches of H₂O, indicated by either the water manometer or the electronic manometer.

Figure 5-2. Insertion of Type S Pitot Tube Into Source Simulator

Disconnect the umbilical cord or Teflon lines from the Type S pitot tube and attach them to the standard pitot tube.

Insert the standard pitot tube into the duct until the mark is at the outside edge of the duct wall.

Align the tube so that it faces directly into the flow stream and the center of the tube is in exactly the same location in the duct as the Type S pitot tube was, as illustrated in Figure 5-3.
Figure 5-3. Insertion of Standard Pitot Tube Into Source Simulator

- On the Laboratory Station 5 Worksheet, Calibration of Type S Pitot Tube, record the Δp, in inches of H₂O, indicated by either the water manometer or the electronic manometer for the standard pitot tube.
- Repeat the above steps for "B" side of Type S pitot tube. [Note: This means that you must also obtain another measurement with the standard tube.]
- Perform a second calibration run by repeating the above steps for calibration of the Type S pitot tube against a standard pitot tube.
- Perform a third calibration run by repeating the above steps for calibration of the Type S pitot tube against a standard pitot tube.
- After three (3) runs, determine the C_p for both the "A" and "B" side of the Type S pitot tube utilizing the following equation:

\[ C_{p(s)} = C_{p(\text{std})} \frac{\Delta p_{\text{std}}}{\Delta p_s} \]

Assume \( C_{p(\text{std})} = 0.99 \)

- Obtain the average \( C_p \) for leg A and leg B of the Type S pitot tube and record the result on the Laboratory Station 5 Worksheet, Type S Pitot Tube Coefficient Data Form.
Laboratory Station 5 Worksheet
Calibration of Type S Pitot Tube Form

Date: 

Time: 

Tester: 

Location: 

Type S Pitot Tube Identification Number: 

NIST Traceable Standard Pitot Tube Identification Number: 

NIST Traceable Standard Pitot Tube $C_p$ Value: 0.99

<table>
<thead>
<tr>
<th>Type S Pitot Tube Calibration Form</th>
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</thead>
<tbody>
<tr>
<td>Leg A, B of Type S Pitot Tube</td>
</tr>
<tr>
<td>---------------------------------</td>
</tr>
<tr>
<td>Test 1</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Test 2</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Test 3</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

$C_p(s) = C_{p(std)} \left( \frac{Δp_{std}}{Δp_s} \right)$
Stack Gas Moisture

One of the most important parameters to estimate correctly when selecting your nozzle diameter and setting up your isokinetic rate equation is the stack gas moisture ($B_{ws}$). For every % stack gas moisture you estimate incorrectly will cause your % isokinetics to be off the same %. The objective of Laboratory Station 6 is to determine the stack gas moisture content of the source simulator using three (3) estimation techniques. They are:

- Wet bulb/dry bulb technique;
- Nomograph technique; and
- Psychometric chart technique.

Complete the laboratory exercises and record your results on the Laboratory Station 6 Worksheet.
Moisture Determination Using Wet Bulb/dry Bulb Technique

The determination of a stack gas wet and dry bulb temperatures can be accomplished with either thermocouples or in-glass mercury thermometers, as illustrated in Figure 6-1. This experiment involves recording both the dry bulb and wet temperatures of the stack gas in order to calculate percent (%) moisture of the stack gas.

![Diagram of Wet Bulb-Dry Bulb Technique]

**Figure 6-1. Test Arrangement For Wet Bulb-Dry Bulb Technique Using In-Glass Mercury Thermometers For Moisture Determination**

During this exercise, we will use a Type K thermocouple attached to a hand held digital temperature display.

Obtain the Type K thermocouple with the hand held digital temperature display along with a cotton sleeve and a 50-mL beaker containing distilled-deionized (DI) water from the Inspector’s Tool Kit (see Appendix C).

Attach the Type K thermocouple with standard plug to the hand held digital temperature instrument. Turn on the hand held digital temperature instrument and verify that the measurement scale records in °F.

Using any port on the source simulator, insert the Type K thermocouple into the gas stream at the center of the stack. Cover the port hole with duct tape or a rag so that air does not leak in. Allow the thermocouple to come into equilibrium with the flue gas temperature. Read the temperature indicated by the hand held digital temperature instrument.

Record the dry bulb/flue gas temperature.

Dry bulb temperature: _____________ °F
Once equilibrium has been reached and the dry bulb temperature recorded, remove the Type K thermocouple from the stack gas and cover the thermocouple with a cotton sleeve. Secure the sleeve over the thermocouple with some tape. Dip the end of the thermocouple with the cotton sleeve into water to saturate the wick. Insert the sleeve covered thermocouple into the duct and allow the thermocouple to come into equilibrium with the flue gas temperature. The temperature reading (as a function of time elapsed) should change in a manner indicated by Figure 6-2.

![Graph showing Dry bulb, Wet bulb, and Inflection Area](image)

**Figure 6-2. Equilibrium and Inflection Points of Dry Bulb and Wet Bulb Thermocouples**

- The temperature of the wet bulb thermocouple will rise to an equilibrium value, and then drying out, will reach the temperature of the dry bulb temperature (i.e., flue gas) if allowed to remain for a period of time.
- Read the wet bulb temperature at the first equilibrium value (the inflection point) and record this temperature as the wet bulb temperature.

\[
\text{Wet Bulb Temperature} \quad \quad \quad \quad \quad \quad \quad \quad 0 \, ^\circ F
\]

- Obtain a value for the current barometric pressure, \( P_{\text{bar}} \), by either asking the instructor or obtaining the local barometric pressure from the National Weather Service using the computer or phoning the local weather station.
- Obtain a value for \( p_s \), the stack static pressure, by inserting a standard pitot tube into the center of the duct and measuring the pressure on the static tap of the tube. If the stack pressure is positive, attach the tube to the left-hand side of the manometer. If the stack pressure is negative, attach the tube to the right-hand side of the manometer.
- Moisture content can be calculated (using the wet bulb-dry bulb technique) from the following equations:
Laboratory Station 6

\[ B_{ws} = \frac{v.p.}{P_s} \]

Where:  
\( v.p. \) = vapor pressure of \( H_2O \)  
\( P_s \) = absolute pressure of stack gas

☐ First solve for \( v.p. \):

\[ v.p. = s.v.p. = (3.67 \times 10^{-4})(P_s)(t_d-t_w) \left( 1 + \frac{tw-32}{157.1} \right) \]

Where:  
\( s.v.p. \) = saturated \( H_2O \) vapor pressure at wet bulb temperatures taken Table 6-1.  
\( t_d \) = temperature of dry bulb measurement \( ^\circ F \)  
\( t_w \) = temperature of wet bulb measurement \( ^\circ F \)  
\( P_s \) = absolute pressure of stack gas = \( P_{bar} + P_s \)

\[ v.p. = (\_\_\_in. \text{Hg}) - (3.67 \times 10^{-4})(\_\_\_in. \text{Hg})(\_\_\_.\_\_.\_\_.\_\_.\_\_.\_\_.) \left( 1 + \frac{\_\_\_.\_\_.\_\_.\_\_.\_\_.} {157.1} \right) \]

\[ = (\_\_\_in. \text{Hg}) - (\_\_\_in. \text{Hg}) \]

\[ = \_\_\_.\_\_. \text{in. Hg} \]

☐ Next determine \( P_s \):

\[ P_s = \_\_\_.\_\_. \text{in. Hg} + \_\_\_.\_\_. \text{in. H}_2\text{O/ in. Hg} \]

\[ P_s = \_\_\_.\_\_. \text{in. Hg} \]

☐ Supplying all unknown into the equation:

\[ B_{ws} = \frac{\_\_\_.\_\_. \text{in. Hg}}{\_\_\_.\_\_. \text{in. Hg}} = \_\_\_.\_\_. \times 100 = \_\_\_.\_\_.\% \]
Table 6-1. Saturated Water Vapor Pressure (Inches of Mercury)

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<td>1.213</td>
<td>1.253</td>
<td>1.293</td>
<td>1.335</td>
<td>1.378</td>
</tr>
<tr>
<td>90</td>
<td>1.422</td>
<td>1.467</td>
<td>1.513</td>
<td>1.561</td>
<td>1.610</td>
<td>1.660</td>
<td>1.712</td>
<td>1.765</td>
<td>1.819</td>
<td>1.875</td>
</tr>
</tbody>
</table>

Record your results below and on the Laboratory Station 6 Worksheet located at the end of this laboratory instructions.

% Moisture = ______________________ %

Moisture Determination Using Nomograph Technique

Another technique for determining approximate moisture in the flue gas is by the use of a nomograph. Nomographs are mathematically constructed to solve various equations when known process information is supplied. While nomographs may not be as accurate as an actual analysis, they do provide a useful approximate moisture determination. To properly use the nomograph, obtain the wet bulb and dry bulb temperatures from the previous exercise and determine the wet bulb/dry bulb depression.

Depression = \( t_d - t_w \)

\[ \begin{align*}
\text{Depression} & = \_\_\_\_ \, ^\circ F - \_\_\_\_ \, ^\circ F \\
\end{align*} \]

6-5
On the line from absolute stack gas pressure to wet bulb depression temperature, mark the pivot point on line 1, using Figure 6-3.

Figure 6-3. Determination of % Moisture Using Nomograph Technique

- From the pivot point online 1 to the $t_w$, mark the pivot point on line 2.
- From the absolute stack gas pressure through the mark on pivot point on line 2, read % H$_2$O on scale $B_{ws}$. Record your results below and on the Laboratory Station 6 Worksheet located at the end of this laboratory instructions.

% Moisture = 

Moisture Determination Using Psychrometric Chart Technique

- Moisture concentration of a stack gas can also be determining using the psychrometric chart technique if one knows the wet bulb and dry bulb temperatures. Figure 6-4 illustrates how to use a psychrometric chart utilizing the wet bulb and dry bulb temperatures.
- Using Figure 6-5, find the dry bulb temperature on the X-axis.
- Now find the wet bulb temperature on the saturation line (wet bulb temperature line).
Draw a line "up" from the dry bulb temperature.
Draw a line sideways from the wet bulb temperature.

Figure 6-4. Using A Psychrometric Chart
Figure 6-5. Psychrometric Chart Expressing Volume of Water Vapor by Percent

The point of intersection on the two lines is read as the percent water vapor on the Y-axis. Record your results below and on the Laboratory Station 6 Worksheet located at the end of this laboratory instructions.

\[
\% \text{ Moisture} = \underline{\underline{\text{_______} \text{___}}} \% \]

Laboratory Station 6 Worksheet
Moisture Determination Using Wet Bulb/dry Bulb Technique

☐ Retrieve the results from the various methods for determination of moisture in a stack gas.
☐ Record the results below.

1. $B_{ws}$ __________________ % Calculation Method

2. $B_{ws}$ __________________ % Nomograph Method

3. $B_{ws}$ __________________ % Psychrometric Chart Method
Pitot Tube Inspection

In order to assign a known pitot tube coefficient factor, $C_p$, the pitot tube must meet certain design and construction requirements as identified in Federal Reference Method 2 (FRM 2) inorder to assign a value of 0.84. To complete this exercise, you will be given a Type S pitot tube along with a “bulls eye” and level indicator from the Inspector’s Tool Kit (see Appendix C). Using the Laboratory Station 7 Worksheet, inspect the Type S pitot tube and see if it meets specifications as defined by Federal Reference Method 2 (FRM 2) by evaluating the pitot tubes $\alpha$, $\beta$, $z$, $w$ and $A$ characteristics so a $C_p$ of 0.84 can be assigned to the Type S pitot tube.
APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

Pitot Tube Inspection

☐ Obtain a traditional Type S pitot tube to be used in this laboratory exercise. Also obtain a dial caliper, bull’s eye level, and a level indicator from the Inspector’s Tool Kit (see Appendix C).

☐ Before starting the inspection check, obtain the Laboratory Station 7 Worksheet (i.e., Type S Pitot Tube Inspection Data Sheet).

☐ Clamp the Type S pitot tube so that it is level, verify and record on Inspection Data Sheet.

☐ Verify that the Type S pitot openings are not damaged or obstructed and record on the Laboratory Station 7 Worksheet.

☐ Using an angle indicator, measure the angles ($\alpha_1$ and $\alpha_2$) between the pitot tube opening plane and the horizontal plane when viewed from the end, and record on the Laboratory Station 7 Worksheet.

☐ Measure the angles ($\beta_1$ and $\beta_2$) between the pitot tube opening plane and the horizontal plane when viewed from the side, and record on the Laboratory Station 7 Worksheet.

☐ Calculate the difference in length between the two pitot tube legs ($Z$) by measuring the angle $\alpha$ and record on the Laboratory Station 7 Worksheet.

☐ Calculate the distance that the Type S pitot tube legs are rotated ($W$) by measuring the angle $\beta$ and record on the Laboratory Station 7 Worksheet.

☐ Measure and record the vertical distances ($P_A$ and $P_B$) between each pitot tube opening plane and the center line of the Type S pitot tube and record on the Laboratory Station 7 Worksheet.

☐ Measure and record the tube external diameter ($D_T$) and calculate the minimum and maximum values of $P_A$ and $P_B$ and record on the Laboratory Station 7 Worksheet.

☐ If the Type S pitot meets the construction specifications, then sign and date the Laboratory Station 7 Worksheet (i.e., Type S Pitot Tube Inspection Data Sheet).
**Laboratory Station 7 Worksheet**

**Type S Pitot Tube Inspection Data Sheet**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Allowable Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assembly Level?</td>
<td>Yes</td>
<td></td>
</tr>
<tr>
<td>Holes Damaged?</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Obstructed?</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>$\alpha_1$</td>
<td>-10° &lt; $\alpha_1$ &lt; 10°</td>
<td></td>
</tr>
<tr>
<td>$\alpha_2$</td>
<td>-10° &lt; $\alpha_2$ &lt; 10°</td>
<td></td>
</tr>
<tr>
<td>$\beta_1$</td>
<td>-5° &lt; $\beta_1$ &lt; 5°</td>
<td></td>
</tr>
<tr>
<td>$\beta_2$</td>
<td>-5° &lt; $\beta_2$ &lt; 5°</td>
<td></td>
</tr>
<tr>
<td>$\gamma$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\delta$</td>
<td>for 1/4&quot; OD, 0.526 to 0.750</td>
<td>for 3/8&quot; OD, 0.788 to 1.125</td>
</tr>
<tr>
<td>$A$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$Z = A \sin \gamma$</td>
<td>$Z \leq 0.125^\circ$</td>
<td></td>
</tr>
<tr>
<td>$W = A \sin \delta$</td>
<td>$W \leq 0.031^\circ$</td>
<td></td>
</tr>
<tr>
<td>$P_a$</td>
<td>for 1/4&quot; OD, 0.263 to 0.375</td>
<td>for 3/8&quot; OD, 0.394 to 0.563</td>
</tr>
<tr>
<td>$P_b$</td>
<td>for 1/4&quot; OD, 0.263 to 0.375</td>
<td>for 3/8&quot; OD, 0.394 to 0.563</td>
</tr>
<tr>
<td>$P_a \cdot P_b$</td>
<td>-0.063 to 0.063°</td>
<td>0.188 to 0.375°</td>
</tr>
<tr>
<td>$D_t$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Certification**

I certify that the Type S pitot tube/probe ID # ___________ meets or exceeds all specifications, criteria and/or applicable design features and is hereby assigned a pitot tube calibration factor $C_p$ of 0.84.

Certified By:  

[Signature/Date]  

Team Leader (Signature/Date)
Federal Reference Method 5 (FRM 5) Sampling Train

Federal Reference Method 5 (FRM 5) is the principal method used for sampling particulate matter from industrial sources. However, other particulate sampling methods can also be used and may be applied for specific source categories. These methods include the FRM 8 and 17, given in 40CFR60, Appendix A and the PM-10 sampling methods given in 40CFR51, Appendix M. Equivalent methods for special purposes, or when emissions from a given facility are not capable of being measured by FRM 5, are also used. However, their use is generally subject to the approval of the agency administrator.

The objective of Laboratory Station 5 is to assemble the FRM 5 sampling train, charging the filter holder with a filter, and charging the impingers with water and silica gel. Once assembled, your objective is to “leak check” the train to see if you indeed assembled the system properly. Once the “leak check” has been completed, it is your task to use data from the other laboratory stations and determine a correct $D_n$ and “K” factor in order to set-up the isokinetic rate equation. Once the isokinetic rate equation has been established, complete all data entry to the Laboratory Station 8 Worksheet [i.e., FRM 5 Field Test Data Sheet (FTDS)] and prepare to sample the source simulator gas stream. We will only sample 4 traverse points on a single diagonal at the source simulator. Sample the 4 points for 5 minutes each maintaining isokinetic conditions. After sampling is complete, remove the sampling train from the monorail and recover the filter, water and silica gel. Use the equations in the procedure to complete the emission calculations.
Federal Reference Method 5 (FRM 5) Sampling Train

- **Pre-Test Preparation**
  - Obtain a filter from the laboratory equipment supply and check the filter visually against light for irregularities and flaws or pinhole leaks. Label the filter on the back side near the edge using numbering machine ink.
  - The filters have been desiccated at 20° ± 5.6°C and ambient pressure for ≥ 24 hr, and weighted at intervals of ≥ 6 hr to a constant weight (≤ 0.5 mg change from previous weighing). Record weight on your Laboratory Station 8 Worksheet/Field Test Data Sheet (FTDS).
  - The probe liners and probe nozzles have been internally brushed, first with tap water, then distilled/deionized water, followed by reagent-grade acetone. The probe liner and probe nozzle have been rinsed with acetone and allow to air-dry. Inspect visually for cleanliness and repeat the procedure if necessary. Cover the probe liner openings to avoid contamination. Nozzles should be kept in a case to avoid contamination or damage to the knife-edge. **Note:** Special cleaning procedures may be required for other test methods (for example, metals or dioxin).
  - The glassware (filter assemblies, impingers and connecting glassware) have been cleaned internally by wiping the joints, washing with glass cleaning detergent, rinsing with distilled/deionized water, followed by reagent-grade acetone, and then allow to air-dry. All exposed openings have been covered with parafilm, plastic caps, serum caps, ground-glass stoppers or aluminum foil (not for metals!) to avoid contamination. **Note:** Special cleaning procedures may be required for other test method (for example, metals or dioxin).

- **Preliminary Determinations**
  - Select the sampling site, measure the stack or duct dimensions, and determine the number of traverse points (Use Federal Reference Method 1 Laboratory Station 10 results).
  - Determine the stack gas pressure, range of velocity pressure heads, and temperature (Use Federal Reference Method 2 Laboratory Station 4 results).
  - Determine or estimate the dry molecular weight (Use Federal Reference Method 3 Laboratory Station 6 results).
  - Determine the moisture content of the stack gas (Use Federal Reference Method 4 Laboratory Station 6 results).
  - Select a suitable probe assembly length such that all traverse points can be sampled.
  - Using the following equation, calculate an estimated nozzle diameter \( D_{n(est)} \) using the values from other laboratory stations. Then select a nozzle from the FRM 5 nozzle set that closest measures the same diameter as the calculated value. Refer to Appendix A for definition of abbreviations and terminology.

\[
D_{n(est)} = \sqrt[3]{\frac{K_s Q_m P_m}{T_m C_p (1 - B_{sw})}} \sqrt{\frac{T_s M_s}{P_s \Delta p_{avg}}}
\]

where \( K_s = 0.6071 \) (metric units)
\( = 0.03575 \) (English units)
After selecting the appropriate nozzle, determine the K-factor for the isokinetic sampling rate using the following equation. \[ \text{[Note: Do NOT change nozzle size during the sampling run.]} \]

\[
K = \frac{\Delta H}{\Delta p} = K_6 \cdot D_n^4 \cdot \Delta H \cdot \frac{C_p^2 (1 - B_{ws})^2}{M_d T_m P_s} \cdot \frac{M_s T_s P_m}{M_x T_x P_m}
\]

where
- \(D_n\) = Nozzle diameter, mm (inches)
- \(T_m\) = Average DGM temperature, °K (°R)
- \(T_s\) = Average stack gas temperature, °K (°R)
- \(K_6\) = 0.0000804 (metric units)
- \(= 849.842\) (English units)

Select the total sampling time (we will use 20 minutes) and standard gas sample volume specified in the test procedures for the specific industry. Select equal sampling times of 5 minutes per traverse point.

C. Preparation of Sampling Train

- Mark the probe assembly with heat-resistant tape or “White-Out” to denote the proper distance into the stack or duct for each sampling point.
- Insert the probe nozzle into the probe sheath union, and finger tight the union fitting. Avoid over tightening to prevent cracking the glass probe liner. Keep the nozzle tip and the ball joint on the probe liner covered until the assembly of the train is complete and sampling is about to begin. Secure the probe assembly to the sample case by tightening the probe clamp.
- Prepare each set of impingers for a sampling run
  - Impingers 1 & 2: 100 mL water in each
  - Impinger 3: Empty
  - Impinger 4: 200 to 300 g of silica gel
- Weigh each impinger to the nearest ± 0.5 g using a top-loading electronic balance, as illustrated in Figure 8-1. Record initial weights on the Laboratory Station 8 field test data sheet (FTDS).

![Top-Loading Electronic Balance](image)

Figure 8-1. Top-Loading Electronic Balance for Weighting
Assemble the impingers in the cold box with U-tubes, double “L” adapter, and the sample case/umbilical adapter, using ball joint clamps or clips, as illustrated in Figure 8-2.

Figure 8-2. View of Assembled Impingers in Cold Box

Using tweezers or clean disposable surgical gloves, place the tared filter on the grooved side of the TFE filter support in the filter holder, as illustrated in Figure 8-3. Check the filter for tears after placement, and center on the filter support. Assemble the filter holder and tighten the clamps around the filter holder to prevent leakage around the O-ring. Record filter number on the Laboratory Station 8 FTDS.

Figure 8-3. Exploded View of Filter Assembly

Connect the filter holder and cyclone bypass (if applicable) in the hot box to the probe liner ball joint and to the “L” adapter using ball joint clamps. Close the hot box doors and fasten shut.

Connect the umbilical cable electrical and pitot tube line connections to the assembled sampling train and to the source sampler console, as illustrated in Figure 8-4.
Figure 8-4. Assembled Sampling Train Before Umbilical Hookup

- Place the assembled sampling train near the sampling port, either on the monorail or other support.
- Turn on and set probe and hot box heaters. Allow the hot box and probe to heat for at least 15 minutes before starting the test, and make periodic checks and adjustments to ensure the desired temperatures. Check all thermocouple connections by dialing through each selection and noting ambient or heated temperatures. Place crushed ice and a little water around the impingers.
- Leak-check the sampling train (see Leak-Check Procedure for Isokinetic Sampling Trains in Method 4 and Pitot Tube and Line Leak-Check in Method 2) by putting your finger over the opening of the nozzle and start the pump by pushing down on the toggle switch. Now adjust the vacuum in the sampling train to – 15 ′ Hg using both the coarse and fine adjust knobs. The dry gas meter should not rotate more than 0.02 ft³ for one minute to have a successful leak check. If passed, then slowly remove the finger until the vacuum gauge is below 5 ′ Hg vacuum, then remove the finger totally and cut off the pump. Record leak rate on Laboratory Station 8 FTDS.

D. Sampling Run Procedure
- Open and clean the portholes of dust and debris
- Level and zero the Δp and ΔH manometers on the meter box.
- Record the initial dry gas meter (DGM) reading on the Laboratory Station 8 FTDS.
- Remove the nozzle cap, verify that the hot box/filter and probe heating systems are up to temperature, and check pitot tube, temperature gauge, and probe alignments and clearances.
- Close the coarse valve and fully open the fine valve. Position the nozzle at the first traverse point. Record the clock time, read Δp on the manometer and determine ΔH from your calculations using the calculated “K” factor. Immediately start the pump, and adjust the flow to set the ΔH, first by adjusting the coarse valve and then the fine adjust valve. [Note: If necessary to overcome high negative stack pressure, turn on the pump while positioning the nozzle at the first traverse point.]
- When the probe is in position, block off the openings around the probe and porthole using duct tape, rags, gloves or towels (or flameproof materials for hot stacks), as illustrated in Figure 8-5.
Figure 8-5. Blocking off the Porthole During Sampling

- Record the ΔH, pump vacuum and temperatures for stack gas, DGM, filter box, probe, and impinger exit on the Laboratory Station 8 FTDS for Point #1. Record the ID numbers for DGM, thermocouples, pitot tube, and sample box.
- Traverse the stack cross-section for the same time (~ 5 minutes) period at each sampling point (4 sampling points) without turning off the pump except when changing ports. Do not bump the probe nozzle into the stack walls.
  - Maintain the temperature of the hot box (probe outlet or filter outlet) at the proper level.
  - Monitor the Δp during each point, and if the Δp changes by more than 20%, another set of readings should be recorded.
  - Periodically check the level and zero of the manometers, and re-adjust if necessary.
  - Record DGM readings at the beginning and end of each sampling time increment, before and after each leak-check, and when sampling is halted.
  - Take other readings (ΔH, temps, vacuum) at least once each sample point during each time increment, maintaining the ΔH/Δp isokinetic ratio.
  - Add more ice and, if necessary, salt to maintain a temperature <20°C (68°F) at the silica gel impinger exit.
- At the end of the sample run, turn off the coarse valve, remove the probe and nozzle from the stack, turn off the pump and heaters, and record the final DGM reading on the Laboratory Station 8 FTDS.
  - [Mandatory: Leak-Check the sampling train at the maximum vacuum achieved during the sample run. Record leak-check results on the Laboratory Station 8 FTDS.]
  - [Mandatory: Leak-Check the pitot lines. Record on the Laboratory Station 8 FTDS.]
- Allow the probe to cool. Wipe off all external particulate material near the tip of the probe nozzle, and cap the nozzle to prevent contamination or loss of sample. Hint: Open the hot box doors to allow the filter holder to cool.
- Before moving the sampling train to the cleanup site, disconnect the probe from the inlet and cover both ends. Do not lose any condensate that might be present. Disconnect the filter holder from the "L" Adapter and cap off the filter holder.
E. Sample Recovery

[Note: Sample Recovery is extremely important because that is where sample loss can occur (bias results low due to sampler errors or blunders) or contamination can be introduced (bias results high).]

☐ Place 200 ml of acetone from the wash bottle being used for cleanup in a glass sample container labeled “Acetone Blank”.

☐ Inspect the train prior to and during disassembly, and note any abnormal conditions on the Laboratory Station 8 FTDS.

☐ **Container No. 1 – Filter**

    ☐ Using a pair of tweezers and/or clean disposable surgical gloves, carefully remove filter from the Filter Holder, and place it in its identified petri dish container. If necessary, fold the filter such that the particulate matter cake is inside the fold.

    ☐ Using a nylon bristle brush and/or a sharp-edged blade, carefully transfer to the petri dish any PM and/or remaining pieces of filter or filter fibers that adhere to the filter support or gasket.

☐ **Container No. 2 – Acetone Rinses** – Recover any particulate matter from the internal surfaces of the probe nozzle, swaged union fitting, probe liner (use a glass funnel to aid in transferring liquid washed to the container), front half of the Filter Holder, and (if applicable) the cyclone. Recover all rinses in a single glass container, as illustrated in Figure 8-6, 8-7, 8-8 and 8-9. Perform the following steps in sample recovery:

    ☐ Before cleaning the front half of the filter holder, wipe clean all joints.

    ☐ Rinse with acetone, brush with small nylon bristle brush, and rinse with acetone until there are no visible particles. Make a final acetone rinse.

    ☐ For probe liner, repeat rinse, brush, rinse sequence at least three times for glass liners, and six times for metal liners.

---

**Figure 8-6. Acetone Rinse of Sampling Probe Liner**
Laboratory Station 8

Figure 8-7. Rinsing Probe Nozzle

Figure 8-8. Brushing Probe Nozzle

Figure 8-9. Front Half Acetone Rinse Samples
Make a final rinse of the probe brush with acetone.

For Probe Nozzle, use the nylon nozzle brush and follow the same sequence of rinse, brush, rinse as for the probe linger.

After completing the rinse, tighten the lid on the sample container. Mark the height of the fluid level. Label the container.

**Container No. 3 – Silica Gel**

- Determine whether silica gel has been completely spent, and note on the FTDS its condition and color.
- Recover the silica gel into a pre-weighted Nalgene bottle, as illustrated in Figure 8-10.

![Figure 8-10. Recovering Silica Gel Into Pre-weighted Nalgene Bottle](image)

- Weigh the silica gel for moisture determination. Either re-use in the next run, using the final weight as the initial weight for the new sampling run, or discard.

**Container No. 4 - Impinger Water**

- Note on the Laboratory Station 8 FTDS any color or film in the liquid catch.
- Measure the volume in the impingers as illustrated in Figure 8-11 or weigh to the nearest gram for calculating stack gas moisture determination.

![Figure 8-11. Measuring the Impinger Water Using Graduated Cylinder](image)
F. Calculations

At the conclusion of each sampling run, it is prudent to calculate the stack gas moisture (for the next sampling run) as well as the average isokinetic rate. To calculate the stack gas moisture content ($B_{ws}$), the following equations are used to compute the sample gas volume ($V_{m(std)}$) and gas moisture volume ($V_{wc(std)}$):

$$V_{m(std)} = K_3 Y \frac{V_m \left( P_{bar} + \frac{\Delta H}{13.6} \right)}{T_m}$$

Where:

- $\Delta H$ = Average orifice tube pressure during sampling, in. H$_2$O
- $V_m$ = Dry gas volume measured by dry gas meter, dam (dcf)
- $T_m$ = Absolute temperature at dry gas meter, °K (°R)
- $Y$ = Dry gas meter calibration factor
- $K_3$ = 0.3858 °dK/mm Hg (metric units)
  = 17.64 °R/in. Hg (English units)

$$V_{wc(std)} = K_2 (W_f - W_i)$$

Where:

- $W_f$ = Final weight of water collected, g
- $W_i$ = Initial weight of water collected, g
- $K_2$ = 0.001335 m$^3$/g (metric units)
  = 0.04715 ft$^3$/g (English units)

$$B_{ws} = \frac{V_{wc(std)}}{V_{m(std)} + V_{wc(std)}}$$

Where:

- $B_{ws}$ = Proportion of water vapor, by volume, in the gas stream

Next, the average stack gas velocity is calculated. The equation for average gas velocity in a stack or duct is:

$$V_s = K_p C_p \left( \frac{\sqrt{T_s(avg)}}{P_s M_s} \right)$$

Where:

- $V_s$ = Average stack gas velocity, m/sec (ft/sec)
- $C_p$ = Pitot tube coefficient, dimensionless
- $(\sqrt{T_s(avg)})$ = Average of the square roots of each stack gas velocity
- $T_s$ = Absolute average stack gas temperature, °K (°R)
- $P_s$ = Absolute stack gas pressure, mmHg (in. Hg)
  = $P_{bar} + P_g/13.6$
- $P_{bar}$ = Barometric pressure at measurement site, mm Hg (in. Hg)
- $P_g$ = Stack static pressure, mm H$_2$O (in. H$_2$O)
\[ M_s = \text{Molecular weight of stack on dry basis, g/g-mole (lb/lb-mole)} \]
\[ = M_b (1-B_{w_s}) + 18.0 B_{w_s} \]
\[ M_d = \text{Molecular weight of stack on dry basis, g/g-mole (lb/lb-mole)} \]
\[ K_p = \text{Constant, 34.97 for metric system (85.49 for English system)} \]

- Calculate the average percent isokinetic sampling rate using the following equation:

\[ \% I = \frac{K_s T_s V_{w(cat)}}{P_v A_n \theta (1 - B_{w_s})} \]

where
- \( A_n \) = Cross-sectional area of the nozzle, \( m^2 \) (\( ft^2 \))
- \( \theta \) = Sampling time, minutes
- \( K4 = 4.320 \) (metric units)
- \( = 0.09450 \) (English units)

G. Variations and Alternatives To Operation of FRM 5 Sampling Train
- Acceptable alternatives to glass probe liners are metal liners, for example, 316 stainless steel, Inconel or other corrosion resistant metals made of seamless tubing. These can be useful for cross-sections over 3 m (10 ft.) in diameter. Whenever practical, make every effort to use borosilicate glass or quartz probe liners. Metal liners will bias particulate matter results high.
- For large stacks, consider sampling from opposite sides of the stack to reduce the length of probe.
- Use either borosilicate or quartz glass probe liners for stack temperatures up to 480\(^\circ\)C to 900\(^\circ\)C (900 - 1,650°F). The softening temperature for borosilicate glass is 820\(^\circ\)C (1,508°F), and for quartz it is 1,500\(^\circ\)C (2,732°F).
- Rather than labeling filters, label the shipping containers (glass or plastic petri dishes), and keep the filters in these containers at all times except during sampling and weighing.
- Use more silica gel in impinger 4, if necessary, but ensure that there is no entrainment or loss during sampling. Hint: Loosely place cotton balls or glass wool in the neck of the silica gel impinger outlet stem.
- If a different type of condenser (other than impingers) is used, measure the amount of moisture condensed either volumetrically or gravimetrically.
- For moisture content, measure the impinger contents volumetrically before and after a sampling run. Use a pre-weighed amount of silica gel in a shipping container, then empty the silica gel after the run back into the container for weighing at another time.
- If the total particulate catch is expected to exceed 100 mg or more or when water droplets are present in the stack gas use a glass cyclone between the probe and filter holder.
- If high pressure drops across the filter (high vacuum on the gauge) causing difficulty in maintaining isokinetic sampling, replace the filter. Suggestion: Use another filter assembly rather than changing the filter itself. Before installing a new filter, conduct a leak-check. Add the filter assembly catches for the total particulate matter weight.
- Use a single train for the entire sampling run, except when simultaneous sampling is required in two or more separate ducts or at two or more different locations within the same duct, or in cases where equipment failure necessitates a change in trains. In all other situations, obtain approval from the regulatory agency before using two or more trains.
- When two or more trains are used, analyze separately the front-half and (if applicable) impinger catches from each train unless identical nozzle sizes were used on all trains. In this case, the front-
half catches may be combined (as may the impinger catches) and one analysis of front-half catch and one analysis of impinger catch may be performed. Consult with the regulatory agency for details concerning the calculation of results when two or more trains are used.

- If a flexible line is used between the first impinger or condenser and the filter holder, disconnect the line at the filter holder, and let any condensed water or liquid drain into the impingers or condenser.

- Do not cap off the probe tip too tightly while the sampling train is cooling down, as this would create a vacuum in the filter holder, which may draw water from the impingers into the filter holder.
### FRM 5 Particulate Field Test Data Sheet (FTDS)

<table>
<thead>
<tr>
<th>Plant</th>
<th>C</th>
<th>Ap, in. H2O</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run No.</td>
<td>$\Delta H_{\text{g}}$</td>
<td>Test Start Time</td>
</tr>
<tr>
<td>Location</td>
<td>$P_{\text{in}}$, in. Hg</td>
<td>Test Stop Time</td>
</tr>
<tr>
<td>Date</td>
<td>$D_{\text{in}}, \text{ Calculated (in.)}$</td>
<td></td>
</tr>
<tr>
<td>Operator</td>
<td>$B_{\text{in}}, \text{ (assumed)}$</td>
<td>$D_{\text{in}}, \text{ Used (in.)}$</td>
</tr>
<tr>
<td>Sample Box No.</td>
<td>$M_{B}$</td>
<td>Ambient Temp, $^\circ F$</td>
</tr>
<tr>
<td>Meter Box No.</td>
<td>$M_{i}$</td>
<td>Bar Pressure, in. Hg</td>
</tr>
<tr>
<td>Filter No.</td>
<td>$T_{\text{in}}, ^\circ F$</td>
<td>Average $\Delta H$, in. H2O</td>
</tr>
<tr>
<td>Silica Gel No.</td>
<td>$T_{\text{in}}, ^\circ F$</td>
<td>K Factor</td>
</tr>
<tr>
<td>Pre-Test Leak Rate @ 15 in. Hg</td>
<td></td>
<td>DGM “Y” Factor</td>
</tr>
<tr>
<td>Post Test Leak Rate @ 15 in. Hg</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Travel Point</th>
<th>Clock</th>
<th>DGM</th>
<th>Plot ($\Delta P$)</th>
<th>Orifice $\Delta H$ (In, H2O)</th>
<th>DGM Temp. ($^\circ F$)</th>
<th>Box Temp.</th>
<th>Impinger Temp.</th>
<th>Stack Temp.</th>
<th>Pump Vacuum</th>
<th>Stack $O_2$</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
<td>Min</td>
<td>MP</td>
<td>In, H2O</td>
<td>Desired</td>
<td>Actual</td>
<td>Inlet</td>
<td>Outlet</td>
<td>$^\circ F$</td>
<td>$^\circ F$</td>
<td>$^\circ F$</td>
<td>In, Hg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Final Volume (mL) Impingers:**

**Initial Volume (mL) Impingers:**

**Total Volume (mL) Gained:**

**Final Weight (mg) of Silica Gel:**

**Initial Weight (mg) of Silica Gel:**

**Total Weight (mg) Gained:**

**Final Weight (mg) of Filter:**

**Initial Weight (mg) of Filter:**

**Total Weight (mg) Gained:**

---

**Expert Stack Testers**
1502 Stack Testing Lane
Cary, NC 27511
919-467-2785
Email: jwinberry@mindspring.com

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**Laboratory Station 8**
**FRM 5 Field Test Data Sheet (FTDS)**
Isokinetic Rate Equation and The IsoCal Spreadsheet

One of the major activities associated with Federal Reference Method 5 is the setting and maintaining isokinetic rate conditions during the test. One can use a nomograph to assist with this or applicable software. The objective of Laboratory Station 9 is to demonstrate the use of the IsoCal software spreadsheet in completing the necessary equations for proper operation of the Federal Reference Method 5 (FRM 5) sample train during sampling. The participant will be given source test data that should be entered into the laptop computer containing the IsoCal software. Based upon the data input, the participant will determine point-by-point isokinetics and average isokinetics. In addition, the participant will be able to see the variability in isokinetics as one changes the various input parameters (i.e., stack gas moisture and nozzle diameter).
APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

Isokinetic Rate Equation and The IsoCal Spreadsheet

☐ The package before you contains a brief discussion of the IsoCal software program. Read the instructions and become familiar with the program.

☐ You have just completed a Federal Reference Method 5 (FRM 5) stack test at a brick manufacturing facility (Plant A).

☐ You have been provided with the field data from that Method 5 test on the exhaust of the process. The first several screens of the IsoCal program you have been provided with contain traditional background information and stack characterization information needed to be acquired about the facility before the actual testing begins. This information helps you to determine the proper nozzle diameter needed for the test, acquiring estimated water vapor concentration of the source to be tested, and determining average stack gas velocity, temperature and pressure. The screens which have been provided include:

- Source Sampling Title Page Information;
- Method 1: Sample and Velocity Traverses for Circular Sources;
- Method 2: Determination of Stack Gas Velocity and Volumetric Flow Rate;
- Method 3: Gas Analysis For The Determination of Dry Molecular Weight;
- Method 4: Determination of Moisture Content In Stack Gases;
- Method 5: Sample Recovery and Integrity Data Sheet; and
- Method 5: Sample Analytical Data Sheet.

☐ In addition, you have been provided the actual Field Test Data Sheet (FTDS) for the Method 5 test.

☐ Your task is to use the computer and enter the FTDS from the stack test information into the Method 5-Run 1 data screen. Only concern yourself with Run 1 from the data given to you for this exercise.

[NOTE: Before entering the data, determine with the information provided whether the correct nozzle diameter was selected based upon the nozzle diameter equation and whether the proper "K-factor" was calculated.]

☐ Now enter the data from the FTDS and generate the final screen: Method 5: Determination of Particulate Emissions-Results.

☐ Did the test steam meet overall isokinetic of 90-110%?

☐ Record your answer.

Yes  
No  

9-2
### SOURCE SAMPLING TITLE PAGE

ALARMS exist and have been acknowledged.

<table>
<thead>
<tr>
<th>Source Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Plant Name</strong></td>
</tr>
<tr>
<td><strong>Sampling Location</strong></td>
</tr>
<tr>
<td><strong>Fuel or Source Type</strong></td>
</tr>
<tr>
<td><strong>Fuel F-Factor</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Test Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Starting Test Date</strong></td>
</tr>
<tr>
<td><strong>Project #</strong></td>
</tr>
<tr>
<td><strong>Operator</strong></td>
</tr>
<tr>
<td><strong>Standard Temperature</strong></td>
</tr>
<tr>
<td><strong>Standard Pressure</strong></td>
</tr>
<tr>
<td><strong>Minimum required sample vol.</strong></td>
</tr>
<tr>
<td><strong>Run Duration</strong></td>
</tr>
<tr>
<td><strong>Base Run Number</strong></td>
</tr>
<tr>
<td><strong># of Ports Available</strong></td>
</tr>
<tr>
<td><strong># of Ports Used</strong></td>
</tr>
<tr>
<td><strong>Port Inside Diameter</strong></td>
</tr>
<tr>
<td><strong>Circular Stack?</strong></td>
</tr>
<tr>
<td><strong>Rectangular Stack?</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Test Equipment Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Meter Box Number</strong></td>
</tr>
<tr>
<td><strong>Meter Calibration Factor</strong></td>
</tr>
<tr>
<td><strong>Orifice Meter Coefficient</strong></td>
</tr>
<tr>
<td><strong>Pitot Identification</strong></td>
</tr>
<tr>
<td><strong>Pitot Tube Coefficient</strong></td>
</tr>
<tr>
<td><strong>Orsat Identification</strong></td>
</tr>
<tr>
<td><strong>Nozzle Number</strong></td>
</tr>
<tr>
<td><strong>Nozzle Diameter</strong></td>
</tr>
<tr>
<td><strong>Probe Number</strong></td>
</tr>
<tr>
<td><strong>Probe Length</strong></td>
</tr>
<tr>
<td><strong>Liner Material</strong></td>
</tr>
<tr>
<td><strong>Sample Case / Oven Number</strong></td>
</tr>
<tr>
<td><strong>Impinger Case Number</strong></td>
</tr>
<tr>
<td><strong>Acetone Lot Number</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Testing Company Information</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Company Name</strong></td>
</tr>
<tr>
<td><strong>Address</strong></td>
</tr>
<tr>
<td><strong>City, State Country Zip</strong></td>
</tr>
<tr>
<td><strong>Project Manager</strong></td>
</tr>
<tr>
<td><strong>Phone Number</strong></td>
</tr>
<tr>
<td><strong>Fax Number</strong></td>
</tr>
</tbody>
</table>

9-3
### METHOD 1 - SAMPLE AND VELOCITY TRAVERSES FOR CIRCULAR SOURCES

<table>
<thead>
<tr>
<th>Plant Name</th>
<th>Plant A</th>
<th>Date</th>
<th>12/16/03</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sampling Location</td>
<td>Outlet of Scrubber (Stack)</td>
<td>Project #</td>
<td>ENCERTEC 1</td>
</tr>
<tr>
<td>Operator</td>
<td>WTW</td>
<td># of Ports Available</td>
<td>2</td>
</tr>
<tr>
<td>Stack Type</td>
<td>Circular</td>
<td># of Ports Used</td>
<td>2</td>
</tr>
<tr>
<td>Stack Size</td>
<td>Large</td>
<td>Port Inside Diameter</td>
<td>4</td>
</tr>
</tbody>
</table>

#### Circular Stack or Duct Diameter

| Distance to Far Wall of Stack | (L_f) | 69.50 | in |
| Distance to Near Wall of Stack | (L_n) | 5.50 | in |
| Diameter of Stack | (D) | 64.00 | in |
| Area of Stack | (A) | 22.34 | ft² |

#### Distance from Port to Disturbances

| Distance Upstream | (B) | 676.00 | in |
| Diameters Upstream | (B0) | 10.56 | diameters |
| Distance Downstream | (A) | 164.00 | in |
| Diameters Downstream | (A0) | 2.56 | diameters |

#### Number of Traverse Points Required

<table>
<thead>
<tr>
<th>Number of Traverse Points Required to Flow Disturbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diameters to Particulate Velocity Points</td>
</tr>
<tr>
<td>Upstream Points</td>
</tr>
<tr>
<td>2.00-4.99</td>
</tr>
<tr>
<td>5.00-5.99</td>
</tr>
<tr>
<td>6.00-6.99</td>
</tr>
<tr>
<td>7.00-7.99</td>
</tr>
<tr>
<td>&gt;= 8.00</td>
</tr>
</tbody>
</table>

Numbers of Traverse Points Required

- **Upstream Spec:** #NAME? #NAME?
- **Downstream Spec:** #NAME? #NAME?
- **Pts Required:** #NAME? #NAME?

1 Check Minimum Number of Points for the Upstream and Downstream conditions, then use the largest.
2 8 for Circular Stacks 12 to 24 inches
12 for Circular Stacks over 24 inches

#### Traverse Point Locations

<table>
<thead>
<tr>
<th>Traverse Point Number</th>
<th>Fraction of Stack Diameter</th>
<th>Distance from Inside Wall</th>
<th>Distance Including Nipple Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.044</td>
<td>2.7/8</td>
<td>8.3/8</td>
</tr>
<tr>
<td>2</td>
<td>0.146</td>
<td>9.3/8</td>
<td>14.7/8</td>
</tr>
<tr>
<td>3</td>
<td>0.296</td>
<td>19</td>
<td>24.4/8</td>
</tr>
<tr>
<td>4</td>
<td>0.074</td>
<td>45</td>
<td>50.4/8</td>
</tr>
<tr>
<td>5</td>
<td>0.854</td>
<td>54.9/8</td>
<td>60.9/8</td>
</tr>
<tr>
<td>6</td>
<td>0.955</td>
<td>61.1/8</td>
<td>68.9/8</td>
</tr>
<tr>
<td>7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**METHOD 2 - DETERMINATION OF STACK GAS VELOCITY AND VOLUMETRIC FLOW RATE**

<table>
<thead>
<tr>
<th>Stack Dimensions</th>
<th>Velocity Travers Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diameter of Length of Stack (D)</td>
<td>Run Number 1-V1</td>
</tr>
<tr>
<td></td>
<td>Run Time</td>
</tr>
<tr>
<td>Width of Stack (W)</td>
<td></td>
</tr>
<tr>
<td>Area of Stack (A_p)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pressures</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Barometric Pressure (P_b)</td>
<td>in Hg</td>
</tr>
<tr>
<td>Static Pressure (P_s)</td>
<td>in Hg</td>
</tr>
<tr>
<td>Absolute Stack Pressure (P_a)</td>
<td>in Hg</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Stack Gas Composition</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Composition Data:</td>
<td>Actual</td>
</tr>
<tr>
<td>Carbon Dioxide Concentration (%CO2)</td>
<td>3.7</td>
</tr>
<tr>
<td>Oxygen Concentration (%O2)</td>
<td>14.8</td>
</tr>
<tr>
<td>Nitrogen Concentration (%N2)</td>
<td>81.5</td>
</tr>
<tr>
<td>Stack Moisture Content (B_w)</td>
<td>11.800</td>
</tr>
<tr>
<td>Stack Dry Molecular Weight (M_d)</td>
<td>29.18</td>
</tr>
<tr>
<td>Stack Wet Molecular Weight (M_w)</td>
<td>27.86</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Avg Stack Gas Velocity (V_g)</td>
<td>32.1</td>
</tr>
<tr>
<td>Avg Stack Dry Std Flow Rate (Q_d)</td>
<td>132.4134</td>
</tr>
<tr>
<td>Avg Stack Dry Std Flow Rate (Q_d)</td>
<td>220.899</td>
</tr>
<tr>
<td>Avg Stack Wet Std Flow Rate (Q_w)</td>
<td>403.28</td>
</tr>
</tbody>
</table>

Stack Cross Section Schematic

Average: 0.43

<table>
<thead>
<tr>
<th>Formulas Used</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A_r = DWK_r for Rectangular Stacks</td>
<td></td>
</tr>
<tr>
<td>A_c = \pi(D/2)^2/4 for Circular Stacks</td>
<td></td>
</tr>
<tr>
<td>where K_r = 1 for metric units</td>
<td></td>
</tr>
<tr>
<td>where K_c = 144 (in^2/ft^2) for English units</td>
<td></td>
</tr>
<tr>
<td>P_b = P_s + P_w/13.6</td>
<td></td>
</tr>
<tr>
<td>%N_2 = 100 - %CO_2 - %CO</td>
<td></td>
</tr>
<tr>
<td>M_d = 44(%CO2) + 32(%zw) + 18(%N2) + 28(%H2)</td>
<td></td>
</tr>
<tr>
<td>M_w = M_d(1 - B_w) + 188w</td>
<td></td>
</tr>
<tr>
<td>(sp)^{1/2}_w = 1.22(sp)^{1/2}</td>
<td></td>
</tr>
</tbody>
</table>

\[
T_{Rain} = \frac{1}{\sqrt{K_1}} + T_w \]

where \( T_w = 273 °K \) for metric units | \( T_w = 460 °R \) for English units |

\[
v_s = \frac{\Delta P}{\Delta z} \frac{1}{\rho} \frac{1}{(T_{Rain}(P,Md))^{1/2}} \]

where K_s = 34.97 for metric units | \( K_s = 85.49 \) for English units |

\[
O_{ins} = 3600(1 - B_w)\rho A_s(T_{Rain}(T_{Rain}))^{1/2} \]

where T_{Rain} = 293 °K, P = 760 mm Hg, for metric units | \( T_{Rain} = 528 °K, P_{ins} = 451.92 \) in Hg, for English units
**Method 3 - Gas Analysis for the Determination of Dry Molecular Weight**

**Plant Name:** Plant A  
**Sampling Location:** Outside of Scrubber (Slag)  
**Operator:** W/V  
**# of Pumps Used:** 2  
**Fuel Type:** NA  
**Minimum Fuel Factor:**  
**Maximum Fuel Factor:**  
**Oxidizer Leak Check:** [ ] PreTest [ ] PostTest [ ] PostIdentifacation

<table>
<thead>
<tr>
<th>Run Number</th>
<th>Run Start Time</th>
<th>Run Stop Time</th>
<th>12:05</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Analysis Time</td>
<td>Carbon Dioxide Volume (VCO)</td>
<td>Oxygen Volume (VCO)</td>
<td>Carbon Monoxide Volume (VM CO)</td>
</tr>
<tr>
<td>Run Number</td>
<td>Run Start Time</td>
<td>Run Stop Time</td>
<td>1:25</td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>---------------</td>
<td>-------</td>
</tr>
<tr>
<td>Sample Analysis Time</td>
<td>Carbon Dioxide Volume (VCO)</td>
<td>Oxygen Volume (VCO)</td>
<td>Carbon Monoxide Volume (VM CO)</td>
</tr>
<tr>
<td>Run Number</td>
<td>Run Start Time</td>
<td>Run Stop Time</td>
<td>4:00</td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>---------------</td>
<td>-------</td>
</tr>
<tr>
<td>Sample Analysis Time</td>
<td>Carbon Dioxide Volume (VCO)</td>
<td>Oxygen Volume (VCO)</td>
<td>Carbon Monoxide Volume (VM CO)</td>
</tr>
<tr>
<td>Run Number</td>
<td>Run Start Time</td>
<td>Run Stop Time</td>
<td>5:15</td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>---------------</td>
<td>-------</td>
</tr>
<tr>
<td>Sample Analysis Time</td>
<td>Carbon Dioxide Volume (VCO)</td>
<td>Oxygen Volume (VCO)</td>
<td>Carbon Monoxide Volume (VM CO)</td>
</tr>
</tbody>
</table>

**Average Calculated Fuel Factor (Fₚ/d)₀ = 1.029**  
**Average Excess Air (%EAA) = 186.5**

---

**Formulas Used**

\[ \%CO₂ = VCO₂ / VCO \]
\[ \%CO = VCO / VCO₂ \]
\[ \%N₂ = 100 - \%CO₂ - \%CO₂ - \%CO \]
\[ Mₐ = 44(\%CO₂) + 32(\%O₂) + 28(\%H₂) + \%CO₂ \]
\[ ΔMₐ = Mₐ - Mₐ₉ \]
\[ Fₚ = (20.9 - \%O₂ - 5\%CO₂)(\%CO₂ + \%CO) \]
\[ %EAA = 100(\%CO₂) - (20.9 - \%O₂)(\%CO₂ + \%CO) \]
### Method 4 - Determination of Moisture Content in Stack Gases

<table>
<thead>
<tr>
<th>Plant Name</th>
<th>Plant A</th>
<th>Date</th>
<th>12/16/03</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sampling Location</td>
<td>Outlet of Scrubber (Stack)</td>
<td>Project</td>
<td>ENCERTEC 1</td>
</tr>
<tr>
<td>Operator</td>
<td>VTVV</td>
<td>Ports Used</td>
<td>2</td>
</tr>
<tr>
<td>Stack Type</td>
<td>Circular</td>
<td>Motor Box Number 1</td>
<td></td>
</tr>
<tr>
<td>Train Leak Check</td>
<td>Yes</td>
<td>PreTest</td>
<td>PostTest</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Motor Cal Factor (V)</td>
<td>0.9903</td>
</tr>
</tbody>
</table>

#### Moisture Content Data

<table>
<thead>
<tr>
<th>Run Number</th>
<th>1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Motor Volume</strong></td>
<td>(V_D) 40.670 m³</td>
</tr>
<tr>
<td><strong>Avg Meter Temp</strong></td>
<td>61 °F</td>
</tr>
<tr>
<td><strong>Avg Stack Temp</strong></td>
<td>448 °F</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Impinger</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>g</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final Value</td>
<td>(V_D),(W_D) 150.00</td>
<td>148.00</td>
<td>0.00</td>
<td>218.70</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial Value</td>
<td>(V_D),(W_D) 100.00</td>
<td>100.00</td>
<td>0.00</td>
<td>201.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net Value</td>
<td>(V_D),(W_D) 50.00</td>
<td>48.00</td>
<td>0.00</td>
<td>18.70</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Volume</strong></td>
<td>(V_D) 98.00 m³</td>
</tr>
<tr>
<td><strong>Total Weight</strong></td>
<td>(W_D) 18.70 g</td>
</tr>
<tr>
<td><strong>Std Meter Volume</strong></td>
<td>(V_Dest) 41.017 m³</td>
</tr>
<tr>
<td>** Moisture Content**</td>
<td>(B_Dest) 11.3 %</td>
</tr>
</tbody>
</table>

#### Moisture Content Data

<table>
<thead>
<tr>
<th>Run Number</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Motor Volume</strong></td>
<td>(V_D) 37.690 m³</td>
</tr>
<tr>
<td><strong>Avg Meter Temp</strong></td>
<td>74 °F</td>
</tr>
<tr>
<td><strong>Avg Stack Temp</strong></td>
<td>451 °F</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Impinger</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>g</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final Value</td>
<td>(V_D),(W_D) 150.00</td>
<td>150.00</td>
<td>0.00</td>
<td>219.31</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial Value</td>
<td>(V_D),(W_D) 100.00</td>
<td>100.00</td>
<td>0.00</td>
<td>202.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net Value</td>
<td>(V_D),(W_D) 50.00</td>
<td>50.00</td>
<td>0.00</td>
<td>16.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Volume</strong></td>
<td>(V_D) 100.00 m³</td>
</tr>
<tr>
<td><strong>Total Weight</strong></td>
<td>(W_D) 19.31 g</td>
</tr>
<tr>
<td><strong>Std Meter Volume</strong></td>
<td>(V_Dest) 37.058 m³</td>
</tr>
<tr>
<td>** Moisture Content**</td>
<td>(B_Dest) 12.9 %</td>
</tr>
</tbody>
</table>

#### Moisture Content Data

<table>
<thead>
<tr>
<th>Run Number</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Motor Volume</strong></td>
<td>(V_D) 39.877 m³</td>
</tr>
<tr>
<td><strong>Avg Meter Temp</strong></td>
<td>73 °F</td>
</tr>
<tr>
<td><strong>Avg Stack Temp</strong></td>
<td>447 °F</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Impinger</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>g</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
<th>ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final Value</td>
<td>(V_D),(W_D) 190.00</td>
<td>110.00</td>
<td>0.00</td>
<td>213.28</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial Value</td>
<td>(V_D),(W_D) 100.00</td>
<td>100.00</td>
<td>0.00</td>
<td>201.80</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net Value</td>
<td>(V_D),(W_D) 90.00</td>
<td>10.00</td>
<td>0.00</td>
<td>11.50</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total Volume</strong></td>
<td>(V_D) 113.00 m³</td>
</tr>
<tr>
<td><strong>Total Weight</strong></td>
<td>(W_D) 11.48 g</td>
</tr>
<tr>
<td><strong>Std Meter Volume</strong></td>
<td>(V_Dest) 39.297 m³</td>
</tr>
<tr>
<td>** Moisture Content**</td>
<td>(B_Dest) 11.8 %</td>
</tr>
</tbody>
</table>
## METHOD 6 - SAMPLE RECOVERY AND INTEGRITY DATA SHEET

### Plant Name: Plant A  
### Sampling Location: Outlet of Scrubber (Stack)  
### Operator: WTW  
### Project #: ENCERTEC 1  
### Acetone Lot Number: 1  
### Date: 12/18/03

#### Run History Data

<table>
<thead>
<tr>
<th>Run Number</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run Start Time (hr:mm)</td>
<td>10:40</td>
<td>1:25</td>
<td>4:00</td>
</tr>
<tr>
<td>Run Stop Time (hr:mm)</td>
<td>12:00</td>
<td>2:40</td>
<td>8:15</td>
</tr>
<tr>
<td>Train Prepared By</td>
<td>WTW</td>
<td>WTW</td>
<td>WTW</td>
</tr>
<tr>
<td>Train Recovered By</td>
<td>WTW</td>
<td>WTW</td>
<td>WTW</td>
</tr>
<tr>
<td>Recovery Date (mm/dd/yyyy)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relinquished By</td>
<td>WTW</td>
<td>WTW</td>
<td>WTW</td>
</tr>
<tr>
<td>Received By</td>
<td>FS</td>
<td>FS</td>
<td>FS</td>
</tr>
<tr>
<td>Relinquished Date (mm/dd/yyyy)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Relinquished Time (hr:mm)</td>
<td>10:00</td>
<td>10:00</td>
<td>10:00</td>
</tr>
</tbody>
</table>

#### Equipment Identification Numbers

<table>
<thead>
<tr>
<th>Equipment</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filter</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acetone Wash</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Silica Gel</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Impinger Case</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Sample Box</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Oven</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Moisture Content Data

<table>
<thead>
<tr>
<th>Impingers 1, 2, and 3 - Water Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final Volume ($V_f$)</td>
</tr>
<tr>
<td>Initial Volume ($V_i$)</td>
</tr>
<tr>
<td>Net Volume ($V_n$)</td>
</tr>
</tbody>
</table>

#### Impinger 4 - Silica Gel Weight

<table>
<thead>
<tr>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Final Weight ($W_f$)</td>
</tr>
<tr>
<td>Initial Weight ($W_i$)</td>
</tr>
<tr>
<td>Net Weight ($W_n$)</td>
</tr>
</tbody>
</table>

#### Total Water Collected

<table>
<thead>
<tr>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Volume ($V_t$)</td>
</tr>
</tbody>
</table>

#### Formulas Used

$$V_n = V_f - V_i$$

$$W_n = W_f - W_i$$

$$V_w = \frac{V_n + W_n \cdot \rho_w}{\rho_w}$$

where $\rho_w = 0.9982$ g/ml
**METHODS - SAMPLE ANALYTICAL DATA SHEET**

<table>
<thead>
<tr>
<th>Plant A</th>
<th>Date</th>
<th>Sampling Location</th>
<th>Operator</th>
<th>Project No.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>12/16/03</td>
<td>Outlet to Sandboy (Sand)</td>
<td>WTW</td>
<td>ENCERTEC 1</td>
</tr>
</tbody>
</table>

### Analytical Data

#### Place in Desiccator 1

<table>
<thead>
<tr>
<th>Number</th>
<th>Date</th>
<th>Time</th>
<th>Run Number</th>
<th>Run Start Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>01/20/04</td>
<td>8:00</td>
<td>1</td>
<td>10:40</td>
</tr>
</tbody>
</table>

#### Acetone Wash Beaker

<table>
<thead>
<tr>
<th>Filter</th>
<th>Acetone Wash</th>
<th>Date</th>
<th>Time</th>
<th>Run Start Time</th>
<th>Leakage Evident?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Acetone</td>
<td>01/20/04</td>
<td>8:00</td>
<td>Leakage Evident?</td>
<td>No</td>
</tr>
</tbody>
</table>

#### Results

- **Final Weight** ($m_w$): 0.2694 g
- **Tare Weight** ($m_t$): 0.2568 g
- **Weight Gain** ($W_g$): 9.6 mg
- **Blank Adjustment** ($W_b$): 0.00 mg
- **Total Particulates** ($m_p$): 27.0 mg

#### Place in Desiccator 2

<table>
<thead>
<tr>
<th>Number</th>
<th>Date</th>
<th>Time</th>
<th>Run Number</th>
<th>Run Start Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>01/20/04</td>
<td>8:00</td>
<td>2</td>
<td>1:26</td>
</tr>
</tbody>
</table>

#### Acetone Wash Beaker

<table>
<thead>
<tr>
<th>Filter</th>
<th>Acetone Wash</th>
<th>Date</th>
<th>Time</th>
<th>Run Start Time</th>
<th>Leakage Evident?</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Acetone</td>
<td>01/20/04</td>
<td>8:00</td>
<td>Leakage Evident?</td>
<td>No</td>
</tr>
</tbody>
</table>

#### Results

- **Final Weight** ($m_w$): 0.2644 g
- **Tare Weight** ($m_t$): 0.2492 g
- **Weight Gain** ($W_g$): 10.2 mg
- **Blank Adjustment** ($W_b$): 0.00 mg
- **Total Particulates** ($m_p$): 25.4 mg

#### Place in Desiccator 3

<table>
<thead>
<tr>
<th>Number</th>
<th>Date</th>
<th>Time</th>
<th>Run Number</th>
<th>Run Start Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>01/20/04</td>
<td>8:00</td>
<td>3</td>
<td>4:00</td>
</tr>
</tbody>
</table>

#### Acetone Wash Beaker

<table>
<thead>
<tr>
<th>Filter</th>
<th>Acetone Wash</th>
<th>Date</th>
<th>Time</th>
<th>Run Start Time</th>
<th>Leakage Evident?</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Acetone</td>
<td>01/20/04</td>
<td>8:00</td>
<td>Leakage Evident?</td>
<td>No</td>
</tr>
</tbody>
</table>

#### Results

- **Final Weight** ($m_w$): 0.2565 g
- **Tare Weight** ($m_t$): 0.2316 g
- **Weight Gain** ($W_g$): 4.9 mg
- **Blank Adjustment** ($W_b$): 0.00 mg
- **Total Particulates** ($m_p$): 9.1 mg

### Formulas Used

- $m_w^n = \text{average of first two filter measurements}$
- $m_t = m_{w^n} - m_b^n$
- $W_g = C_{m_{w^n}}$
- $m_p = m_{w^n} + m_t - W_g$
- $m_b^n = \text{average of last two acetone measurements}$
- $m_{w^n} = m_{w^n} - m_b^n$
- $m_{w^n} = m_{w^n} + m_t - W_g$
- $m_{b^n} = \text{blank concentration}$

9-9
**Particulate Field Data Sheet**

Very important - Fill in all blanks

<table>
<thead>
<tr>
<th>Plant</th>
<th>BRIxx, Plant #4A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run no.</td>
<td>#1</td>
</tr>
<tr>
<td>Location</td>
<td>Columbia, S.C.</td>
</tr>
<tr>
<td>Date</td>
<td>12/14/03</td>
</tr>
<tr>
<td>Operator</td>
<td>N/A</td>
</tr>
<tr>
<td>Sample box no.</td>
<td>#1</td>
</tr>
<tr>
<td>Meter box no.</td>
<td>#1</td>
</tr>
<tr>
<td>Nomograph ID no.</td>
<td>#1</td>
</tr>
<tr>
<td>Filter no.</td>
<td>#1 (0.25kg)</td>
</tr>
<tr>
<td>Oral no.</td>
<td>N/A</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Point</th>
<th>Clock time (min)</th>
<th>Dry gas meter ci(MP)</th>
<th>Pitot sp. (in. H2O)</th>
<th>Office δH(in. H2O)</th>
<th>Dry gas temp. (°F)</th>
<th>Pump vac. gauge (in. Hg)</th>
<th>Box temp.</th>
<th>Impinger temp. (°F)</th>
<th>Stack pressure (in. Hg)</th>
<th>Stock Temp. (°F)</th>
<th>Pyrite SO2 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10:18</td>
<td>178.345</td>
<td>0.15</td>
<td>1.39</td>
<td>1.39</td>
<td>51</td>
<td>54</td>
<td>-4.0</td>
<td>1.48</td>
<td>51</td>
<td>44.5</td>
</tr>
<tr>
<td>2</td>
<td>10:15</td>
<td>192.826</td>
<td>0.19</td>
<td>1.76</td>
<td>1.76</td>
<td>57</td>
<td>54</td>
<td>-4.0</td>
<td>2.47</td>
<td>49</td>
<td>44.8</td>
</tr>
<tr>
<td>3</td>
<td>12:55</td>
<td>195.650</td>
<td>0.19</td>
<td>1.76</td>
<td>1.76</td>
<td>57</td>
<td>54</td>
<td>-4.0</td>
<td>2.47</td>
<td>49</td>
<td>44.8</td>
</tr>
<tr>
<td>4</td>
<td>12:55</td>
<td>199.650</td>
<td>0.20</td>
<td>2.26</td>
<td>2.26</td>
<td>61</td>
<td>61</td>
<td>-5.0</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>5</td>
<td>11:00</td>
<td>202.040</td>
<td>0.20</td>
<td>2.26</td>
<td>2.26</td>
<td>61</td>
<td>61</td>
<td>-5.0</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>6</td>
<td>11:00</td>
<td>206.400</td>
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<td>2.26</td>
<td>61</td>
<td>61</td>
<td>-5.0</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>7</td>
<td>11:00</td>
<td>209.990</td>
<td>0.19</td>
<td>1.76</td>
<td>1.76</td>
<td>61</td>
<td>59</td>
<td>-70</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>8</td>
<td>11:40</td>
<td>213.280</td>
<td>0.19</td>
<td>1.76</td>
<td>1.76</td>
<td>61</td>
<td>60</td>
<td>-70</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>9</td>
<td>11:45</td>
<td>224.650</td>
<td>0.19</td>
<td>1.76</td>
<td>1.76</td>
<td>61</td>
<td>60</td>
<td>-70</td>
<td>2.47</td>
<td>55</td>
<td>45.1</td>
</tr>
<tr>
<td>10</td>
<td>11:50</td>
<td>226.150</td>
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<td>1.58</td>
<td>68</td>
<td>61</td>
<td>-50</td>
<td>2.47</td>
<td>64</td>
<td>44.9</td>
</tr>
<tr>
<td>11</td>
<td>11:55</td>
<td>233.450</td>
<td>0.18</td>
<td>1.61</td>
<td>1.61</td>
<td>61</td>
<td>62</td>
<td>-50</td>
<td>2.47</td>
<td>64</td>
<td>44.9</td>
</tr>
<tr>
<td>12</td>
<td>12:00</td>
<td>224.820</td>
<td>0.15</td>
<td>1.39</td>
<td>1.39</td>
<td>71</td>
<td>61</td>
<td>-50</td>
<td>2.47</td>
<td>64</td>
<td>44.9</td>
</tr>
</tbody>
</table>

| Alpha, in. H2O | 0.65 |
| Test start time | 10:40 |
| Stop time | 12:00 |
| Dp, calculated (in.) | 0.254 |
| Dp, used (in.) | 0.254 |
| Ambient temp, °F | 52 |
| Bar, pressure, in. Hg | 29.93 |
| Heater box setting, °F | 250 |
| Probe heater setting, °F | 250 |

Average Eh | 410 |

Post-test | 0.02 |

**Figure C-4. Particulate field data sheet form**

Impinger No. 1: 150 mL 2nd Impinger: 148 mL
FRM 1 Traverse Point Determination

Federal Reference Method 1 (FRM 1) requires that the number of traverse points for a given test be based upon the sampling port locations with reference to upstream and downstream flow disturbances. The objective of Laboratory Station 10 is to determine the number and the location of traverse points at your assigned sampling port on the source simulator. Record your findings on the Laboratory Station 10 Worksheet, FRM 1 Field Test Data Sheet (FTDS).
Laboratory Station 10

APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

FRM 1 TRAVERSE POINT DETERMINATION

Determination of the Minimum Number of Traverse Points

☐ Obtain a tape measure from the Inspector’s Tool Kit and identify the sampling port location on the source simulator.

☐ Measure the inside dimensions of the duct at the sampling site. Sampling ports have been installed in the source simulator; use the port assigned to your laboratory group. For Laboratory Station 10 we will use only one (1) diameter. Record the information on the Laboratory Station 10 FRM 1 Field Test Data Sheet (FTDS).

☐ Determine the duct diameter distances of the sampling port from upstream and downstream flow disturbances. Keep in mind the direction of the flow of gas in the source simulator. [NOTE: Elbows and fans are considered flow disturbances in the source simulator.] Record the information on the Laboratory Station 10 FRM 1 FTDS.

☐ Determine the number of traverse points for a velocity determination using Figure 10-1.

☐ Record the information on the Laboratory Station 10 FRM 1 FTDS.

---

*Figure 10-1. Minimum Number of Traverse Points for Non-Particulate Traverses*
Determine the number of traverse points for a particulate traverse determination using Figure 10-2.

**Figure 10-2. Minimum Number of Traverse Points for Particulate Traverses**

- Record the information on the Laboratory Station 10 FRM 1 FTDS.
- Using Table 10-1, determine the locations for the particulate measurement traverse points and record your calculations on the Laboratory Station 10 FRM 1 FTDS.
- For circular stacks with diameters greater than 60 cm (24 inches), the minimum number of traverse points required is twelve (12), or six (6) in each of two directions 90° apart, when the duct diameters from disturbances are eight (8) or more upstream and two (2) or more downstream. For circular stacks with diameters between 30 and 60 cm (12 and 24 inches), the minimum number of sample points required is eight (8), or four (4) in each of two directions 90° apart. For stacks less than 30 cm (12 inches) in diameter, refer to Method 1A for calculating traverse points.
- After completing Laboratory Station 10 FRM 1 FTDS, obtain a FRM 5 sampling probe and mark the traverse point distances on the sampling probe with heat-resistant fiber tape or whiteout correction fluid, as illustrated in Figure 10-3.
<table>
<thead>
<tr>
<th>Traverse Point Number on a Diameter</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>10</th>
<th>12</th>
<th>14</th>
<th>16</th>
<th>18</th>
<th>20</th>
<th>22</th>
<th>24</th>
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</tbody>
</table>

Table 10-1. Location of Traverse Points in Circular Stacks
Figure 10-3. Marking of Probe For Traverse Point Determination
Laboratory Station 10

APTI #468: MONITORING COMPLIANCE TEST AND SOURCE TEST OBSERVATION

Laboratory Station 10 Worksheet
FRM 1 Field Test Data Sheet (FTDS)

1. Number of Traverse Points
   Port Number: ____________________________________________
   Port Location: __________________________________________
   Duct Diameter (inches): ________________________________
   Number of Duct Diameters From Upstream Disturbance: ______
   Number of Duct Diameters From Downstream Disturbance: ______
   Number of Traverse Points on a Diameter (From Table 10-1): ______

2. Traverse Point Locations

<table>
<thead>
<tr>
<th>Traverse Point Number</th>
<th>Percent of Duct Diameter From Inside Wall to Traverse Point</th>
<th>Location of Traverse Point (Inches)</th>
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<tbody>
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</tbody>
</table>
Abbreviations and Terminology

%CO  Percent CO by volume, dry basis
%CO2  Percent CO2 by volume, dry basis
%N2  Percent N2 by volume, dry basis
%O2  Percent O2 by volume, dry basis
0.280  Molecular weight of N2 or CO, divided by 100
0.320  Molecular weight of O2 divided by 100
0.440  Molecular weight of CO2 divided by 100
100  Conversion to percent.
13.6  Specific gravity of mercury.
18.0  Molecular weight of water, g/g mole (lb/lb mole).
3600  Conversion Factor, sec/hr.
60  Sec/min.
A  Cross sectional area of stack, m² (ft²).
A  Absorbance of sample.
A  Analytical detection limit, µg/ml.
A₁  Absorbance of the 100 µg NO₂ standard.
A₂  Absorbance of the 200 µg NO₂ standard.
A₃  Absorbance of the 300 µg NO₂ standard.
A₄  Absorbance of the 400 µg NO₂ standard.
Aₙ  Cross-sectional area of nozzle, m² (ft²).
B  Liquid volume of digested sample prior to aliquotting for analysis, ml.
**Appendix A**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bws</td>
<td>Water vapor in the gas stream (from Method 4 (reference method) or Method 5), proportion by volume.</td>
</tr>
<tr>
<td>C</td>
<td>Corrected to standard conditions, mg/dscm (lb/dscf).</td>
</tr>
<tr>
<td>C</td>
<td>TGNMO concentration of the effluent, ppm C equivalent.</td>
</tr>
<tr>
<td>Ca</td>
<td>Acetone blank residue concentration, mg/mg.</td>
</tr>
<tr>
<td>Ca</td>
<td>Actual concentration of SO2 in audit sample, mg/dscm.</td>
</tr>
<tr>
<td>Ca</td>
<td>Actual audit sample concentration, mg/dscm.</td>
</tr>
<tr>
<td>Ca1</td>
<td>Concentration of metal in Analytical Fraction 1A as read from the standard curve, µg/mL.</td>
</tr>
<tr>
<td>Ca2</td>
<td>Concentration of metal in Analytical Fraction 2A as read from the standard curve, (µg/mL).</td>
</tr>
<tr>
<td>Cc</td>
<td>Calculated condensible organic (condensate trap) concentration of the effluent, ppm C equivalent.</td>
</tr>
<tr>
<td>Ccm</td>
<td>Measured concentration (NMO analyzer) for the condensate trap †CV, ppm CO2.</td>
</tr>
<tr>
<td>Cd</td>
<td>Determined concentration of SO2 in audit sample, mg/dscm.</td>
</tr>
<tr>
<td>Cd</td>
<td>Determined audit sample concentration, mg/dscm.</td>
</tr>
<tr>
<td>CH2S</td>
<td>Concentration of H2S at standard conditions, mg/dscm.</td>
</tr>
<tr>
<td>Cp</td>
<td>Pilot tube coefficient, dimensionless.</td>
</tr>
<tr>
<td>Cp(s)</td>
<td>Type S pilot tube coefficient, dimensionless.</td>
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<tr>
<td>Cp(std)</td>
<td>Standard pilot tube coefficient; use 0.99 if the coefficient is unknown and the tube is designed according to the criteria of Sections 6.7.1 to 6.7.5 of FRM 2.</td>
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<tr>
<td>cs</td>
<td>Concentration of particulate matter in stack gas, dry basis, corrected to standard conditions, g/dscm (gr/dscf).</td>
</tr>
<tr>
<td>Cs</td>
<td>Concentration of a metal in the stack gas, mg/dscm.</td>
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<tr>
<td>CSO2</td>
<td>Concentration of SO2, dry basis, corrected to standard conditions, mg/dscm (lb/dscf).</td>
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<tr>
<td>Ct</td>
<td>Calculated noncondensible organic concentration (sample tank) of the effluent, ppm C equivalent.</td>
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<tr>
<td>Ct(m)</td>
<td>Measured concentration (NMO analyzer) for the sample tank, ppm NMO.</td>
</tr>
<tr>
<td>D</td>
<td>Diameter of stack, in.</td>
</tr>
<tr>
<td>De</td>
<td>Equivalent diameter.</td>
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<tr>
<td>F</td>
<td>Dilution factor. Required only if sample dilution was needed to reduce the absorbance into the range of the calibration.</td>
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</table>
Appendix A

F  Sampling flow rate, cc/min.

Fa  Aliquot factor, volume of Sample Fraction 2 divided by volume of Sample Fraction 2A

Fd  Dilution factor (Fd = the inverse of the fractional portion of the concentrated sample in the solution actually used in the instrument to produce the reading Ca1. For example, if a 2 mL aliquot of Analytical Fraction 1A is diluted to 10 mL to place it in the calibration range, Fd = 5).

ΔH  Average pressure differential across the orifice meter, mm H2O (in. H2O).

Hgbh  Total mass of Hg collected in the back

Hgbh2  Total mass of Hg collected in Sample Fraction 2, μg.

Hgbh3(A,B,C)  Total mass of Hg collected separately in Fraction 3A, 3B, or 3C, μg.

Hgbhb  Blank correction value for mass of Hg detected in back

Hgfh  Total mass of Hg collected in the front

Hgfhb  Blank correction value for mass of Hg detected in front

Hgt  Total mass of Hg collected in the sampling train, μg.

I  Percent of isokinetic sampling.

K  0.127 mm H2O (metric units); 0.005 in. H2O (English units).

K4  103 mg/μg.

Kc  Spectrophotometer calibration factor.

Kp  Velocity equation constant.

L  Length.

L  Volume of liquid injected, μl.

L1  Individual leakage rate observed during the leak-check conducted prior to the first component change, m3/min (ft3/min)

La  pretest leak-check or for a leak-check following a component change; equal to 0.00057 m3/min (0.020 cfm) or 4 percent of the average sampling rate, whichever is less.

L1  Individual leakage rate observed during the leak-check conducted prior to the "ith" component change (i = 1, 2, 3...n), m3/min or cfm.

Lp  Leakage rate observed during the post-test leak-check, m3/min (cfm).

m  Mass of NOx as NO2 in gas sample, μg.
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<th>Symbol</th>
<th>Description</th>
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<tr>
<td>M</td>
<td>Molecular weight of the liquid injected, g/g mole.</td>
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<tr>
<td>ma</td>
<td>Mass of residue of acetone after evaporation, mg.</td>
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<tr>
<td>Mbh</td>
<td>Total mass of each metal (except Hg) collected in the back</td>
</tr>
<tr>
<td>Mhb</td>
<td>Blank correction value for mass of metal detected in back</td>
</tr>
<tr>
<td>Mc</td>
<td>TGNMO mass concentration of the effluent, mg C/dscm.</td>
</tr>
<tr>
<td>Md</td>
<td>Dry molecular weight, g/g mole (lb/lb mole)</td>
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<tr>
<td>Md</td>
<td>Molecular weight of stack gas, dry basis, g/g mole (lb/lb mole).</td>
</tr>
<tr>
<td>Mfh</td>
<td>Total mass of each metal (except Hg) collected in the front half of the sampling train (Sample Fraction 1), μg.</td>
</tr>
<tr>
<td>Mfhb</td>
<td>Blank correction value for mass of metal detected in front</td>
</tr>
<tr>
<td>mn</td>
<td>Total amount of particulate matter collected, mg.</td>
</tr>
<tr>
<td>Ms</td>
<td>Molecular weight of stack gas, wet basis, g/g mole (lb/lb mole).</td>
</tr>
<tr>
<td>Mt</td>
<td>Total mass of each metal (separately stated for each metal) collected in the sampling train, μg.</td>
</tr>
<tr>
<td>Mw</td>
<td>Molecular weight of water, 18.0 g/g-mole (18.0 lb/lb-mole).</td>
</tr>
<tr>
<td>N</td>
<td>Total number of traverse points.</td>
</tr>
<tr>
<td>n</td>
<td>total number of traverse points.</td>
</tr>
<tr>
<td>N</td>
<td>Normality of barium standard titrant, meq/ml.</td>
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<tr>
<td>N</td>
<td>Carbon number of the liquid compound injected (N = 12 for decane, N = 6 for hexane).</td>
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<tr>
<td>NA</td>
<td>Normality of standard C₂H₆AsO solution, g eq/liter.</td>
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<tr>
<td>Ni</td>
<td>Normality of standard I₂ solution, g eq/liter.</td>
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<tr>
<td>NS</td>
<td>Normality of standard (0.1 N) Na₂S₂O₃ solution, g eq/liter.</td>
</tr>
<tr>
<td>NT</td>
<td>Normality of standard (0.01 N) Na₂S₂O₃ solution, assumed to be 0.1 NS, g eq/liter.</td>
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<tr>
<td>p</td>
<td>Velocity head of stack gas, mm H₂O (in. H₂O).</td>
</tr>
<tr>
<td>P</td>
<td>Allowable pressure change, cm Hg.</td>
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<tr>
<td>Pb</td>
<td>Barometric pressure, cm Hg.</td>
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<tr>
<td>Pbar</td>
<td>Barometric pressure at measurement site, mm Hg (in. Hg).</td>
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<tr>
<td>Pf</td>
<td>Final pressure of the intermediate collection vessel, mm Hg absolute.</td>
</tr>
<tr>
<td>Symbol</td>
<td>Definition</td>
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<tr>
<td>--------</td>
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<tr>
<td>Pg</td>
<td>Stack static pressure, mm Hg (in. Hg).</td>
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<tr>
<td>pi</td>
<td>Individual velocity head reading at traverse point &quot;i&quot;, mm (in.) H2O.</td>
</tr>
<tr>
<td>Pi</td>
<td>Pitch angle at traverse point i, degree.</td>
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<tr>
<td>Pi</td>
<td>Initial absolute pressure of flask, mm Hg (in. Hg).</td>
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<td>Ps</td>
<td>Absolute stack pressure (Pbar + Pg), mm Hg (in. Hg).</td>
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<tr>
<td>Δps</td>
<td>Velocity head measured by the Type S pitot tube, cm (in.) H2O.</td>
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<tr>
<td>Pstd</td>
<td>Standard absolute pressure, 760 mm Hg (29.92 in. Hg).</td>
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<tr>
<td>Δpstd</td>
<td>Velocity head measured by the standard pitot tube, cm (in.) H2O.</td>
</tr>
<tr>
<td>Pt</td>
<td>Final gas sample tank pressure after sampling, but before pressurizing, mm Hg absolute.</td>
</tr>
<tr>
<td>Ptf</td>
<td>Final gas sample tank pressure after pressurizing, mm Hg absolute.</td>
</tr>
<tr>
<td>Pl</td>
<td>Gas sample tank pressure before sampling, mm Hg absolute.</td>
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<tr>
<td>q</td>
<td>Total number of analyzer injections of intermediate collection vessel during analysis (where k = injection number, 1 ... q).</td>
</tr>
<tr>
<td>Qbh2</td>
<td>Quantity of Hg, µg, TOTAL in the ALIQUOT of Analytical Fraction 2B selected for digestion and analysis. NOTE: For example, if a 10 mL aliquot of Analytical Fraction 2B is taken and digested and analyzed (according to Section 11.1.3 and its NOTES Nos. 1 and 2), then calculate and use the total amount of Hg in the 10 mL aliquot for Qbh2.</td>
</tr>
<tr>
<td>Qbh3(A,B,C)</td>
<td>Quantity of Hg, µg, TOTAL, separately, in the ALIQUOT of Analytical Fraction 3A, 3B, or 3C selected for digestion and analysis (see NOTES in Sections 12.7.1 and 12.7.2 describing the quantity &quot;Q&quot; and calculate similarly).</td>
</tr>
<tr>
<td>Qfh</td>
<td>Quantity of Hg, µg, TOTAL in the ALIQUOT of Analytical Fraction 1B selected for digestion and analysis. NOTE: For example, if a 10 mL aliquot of Analytical Fraction 1B is taken and digested and analyzed (according to Section 11.1.3 and its NOTES Nos. 1 and 2), then calculate and use the total amount of Hg in the 10 mL aliquot for Qfh.</td>
</tr>
<tr>
<td>Qsd</td>
<td>Dry volumetric stack gas flow rate corrected to standard conditions, dscm/hr (dscf/hr).</td>
</tr>
<tr>
<td>R</td>
<td>([(\text{mm Hg})/(\text{m3})]/[(\text{K})(\text{g-mole})]) [((\text{in. Hg})(\text{ft3})]/[(\text{R})(\text{lb-mole})]].</td>
</tr>
<tr>
<td>r</td>
<td>Total number of analyzer injections of sample tank during analysis (where j = injection number, 1 ... r).</td>
</tr>
<tr>
<td>Ravg</td>
<td>Average resultant angle, degree.</td>
</tr>
</tbody>
</table>
Appendix A

RE  Relative error for QA audit samples, percent.
Ri  Resultant angle at traverse point i, degree.
Sd  Standard deviation, degree.
T  Sensitivity factor for differential pressure gauges.
Tf  Final absolute temperature of flask, K (R).
Tf  Final temperature of intermediate collection vessel, K.
Ti  Initial absolute temperature of flask, K (R).
Tm  Absolute DGM temperature.
Ts  Stack temperature, C (F).
Ts(abs)  Absolute stack temperature, K (R) = 273 + Ts for metric units, = 460 + Ts for English units.
Tstd  Standard absolute temperature, 293 K (528 R).
Tt  Sample tank temperature at completion of sampling, K.
Ttf  Sample tank temperature after pressurizing, K.
Tti  Sample tank temperature before sampling, K.
V  Sample tank volume, m3.
Va  Volume of acetone blank, ml.
Vb  Volume of sample aliquot titrated, ml.
Va  Volume of absorbing solution, 25 mL.
VA  Volume of C6H5AsO solution used for standardization, ml.
Vt  Total volume of digested sample solution (Analytical Fraction 2A), mL
VAl  Volume of standard C6H5AsO solution used for titration analysis, mL.
Vaw  Volume of acetone used in wash, ml.
Vf  Volume of flask and valve, ml.
Vf1B  Volume of aliquot of Analytical Fraction 1B analyzed, ml. NOTE: For example, if a 1 mL aliquot of Analytical Fraction 1B was diluted to 50 mL with 0.15 percent HNO3 as described in Section 11.1.3 to bring it into the proper analytical range, and then 1 mL of that 50

Vf2B  Volume of Analytical Fraction 2B analyzed, ml. NOTE: For example, if 1 mL of Analytical Fraction 2B was diluted to 10 mL with 0.15 percent HNO3 as described in Section 11.1.3 to bring it into the proper analytical range, and then 5 ml. of that 10 mL was analyzed, Vf2B would be 0.5 ml.
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>VF3(A,B,C)</td>
<td>Volume, separately, of Analytical Fraction 3A, 3B, or 3C analyzed, mL (see previous notes in Sections 12.7.1 and 12.7.2, describing the quantity &quot;V&quot; and calculate similarly).</td>
</tr>
<tr>
<td>VI</td>
<td>Volume of standard I2 solution used for standardization, ml.</td>
</tr>
<tr>
<td>VI T</td>
<td>Volume of standard I2 solution used for titration analysis, normally 50 ml.</td>
</tr>
<tr>
<td>Vlc</td>
<td>Total volume of liquid collected in impingers and silica gel (see Figure 5-6), ml.</td>
</tr>
<tr>
<td>Vm</td>
<td>Volume of gas sample as measured by dry gas meter, dcm (dcf).</td>
</tr>
<tr>
<td>Vm DGM</td>
<td>Dry gas volume as measured by the DGM, dcm (dcf).</td>
</tr>
<tr>
<td>Vm</td>
<td>Volume of gas sample at meter conditions, liters.</td>
</tr>
<tr>
<td>Vm (std)</td>
<td>Volume of gas sample measured by the dry gas meter, corrected to standard conditions, dscm (dcf).</td>
</tr>
<tr>
<td>vs</td>
<td>Stack gas velocity, calculated by Method 2,</td>
</tr>
<tr>
<td>vs</td>
<td>Average stack gas velocity, m/sec (ft/sec).</td>
</tr>
<tr>
<td>Vs</td>
<td>Gas volume sampled, dsm3.</td>
</tr>
<tr>
<td>Vsc</td>
<td>Sample volume (dry basis), ml. at standard conditions</td>
</tr>
<tr>
<td>VSI</td>
<td>Volume of 0.1 N Na2S2O3 solution used for standardization, ml.</td>
</tr>
<tr>
<td>Vsoln</td>
<td>Total volume of solution in which the SO2 sample is contained, 100 ml.</td>
</tr>
<tr>
<td>Vsoln,1</td>
<td>Total volume of digested sample solution (Analytical Fraction 1), ml.</td>
</tr>
<tr>
<td>Vsoln,2</td>
<td>Total volume of Sample Fraction 2, ml.</td>
</tr>
<tr>
<td>Vsoln,3(A,B,C)</td>
<td>Total volume, separately, of Analytical Fraction 3A, 3B, or 3C, ml.</td>
</tr>
<tr>
<td>Vt</td>
<td>Volume of barium standard titrant used for the sample (average of replicate titration), ml.</td>
</tr>
<tr>
<td>VT</td>
<td>Volume of standard (0.01 N) Na2S2O3 solution used in standardizing iodine solution (see Section 10.2.1), ml.</td>
</tr>
<tr>
<td>Vt</td>
<td>Sample train volume, cc.</td>
</tr>
<tr>
<td>Vtb</td>
<td>Volume of barium standard titrant used for the blank, ml.</td>
</tr>
<tr>
<td>VTT</td>
<td>Volume of standard (0.01 N) Na2S2O3 solution used for titration analysis, ml.</td>
</tr>
<tr>
<td>Vv</td>
<td>Intermediate collection vessel volume, m3.</td>
</tr>
</tbody>
</table>
### Appendix A

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_{w(\text{std})}$</td>
<td>Volume of gas, corrected to standard conditions, scm (scf).</td>
</tr>
<tr>
<td>$W$</td>
<td>width.</td>
</tr>
<tr>
<td>$W$</td>
<td>Weight of K$_2$Cr$_2$O$_7$ used to standardize Na$_2$S$_2$O$_3$ or C$_6$H$_5$AsO solutions, as applicable (see Sections 10.2.2 and 10.2.3), g.</td>
</tr>
<tr>
<td>$W_a$</td>
<td>Weight of residue in acetone wash, mg.</td>
</tr>
<tr>
<td>$x_i$</td>
<td>Individual measurements.</td>
</tr>
<tr>
<td>$Y$</td>
<td>Dry gas meter calibration factor.</td>
</tr>
<tr>
<td>$Y_i$</td>
<td>Yaw angle at traverse point i, degree.</td>
</tr>
</tbody>
</table>
Equations

General

\[ \Delta H = K(\Delta p) \]

\[ \text{pmr}_s = c_s Q_s \]

\[ A_s = \pi \left( \frac{D_s}{2} \right)^2 \]

\[ A_s = LW \]

\[ F_d = \frac{20.9 - \%O_2}{\%CO_2} \]

\[ \%EA = \frac{(\%O_2) + 0.5(\%CO)}{0.0264(\%N_2) - (\%O_2) + 0.5(\%CO)} \]

\[ \%N_2 = 100 - \%CO - \%O_2 - \%CO \]

Federal Reference Method I

Equation 1-1

\[ D_e = \frac{2LW}{L+W} \]

Equation 1-2

\[ R_i = \text{arccosine}[(\text{cosine}Y_i)(\text{cosine}P_i)] \]

Equation 1-3

\[ \bar{R} = \frac{\sum R_i}{n} \]
Appendix B

Equation 1-4
\[ S_d = \sqrt{\frac{\sum_{i=1}^{n} (R_i - \bar{R})^2}{(n-1)}} \]

Federal Reference Method 2

Equation 2-1
\[ D_e = \frac{2LW}{L+W} \]

Equation 2-2
\[ C_{p(s)} = C_{p(\text{std})} \sqrt{\frac{\Delta p_{\text{std}}}{\Delta p_s}} \]

Equation 2-3
\[ \text{Deviation} = C_{p(s)} - \bar{C}_p \text{ (A or B)} \]

Equation 2-4
\[ \sigma (\text{side A or B}) = \frac{\sum_{i=1}^{3} (C_{p(s)} - \bar{C}_p \text{ (A or B)})}{3} \]

Equation 2-5
\[ = M_d (1 - B_{ws}) + 18.0 B_{ws} \]

Equation 2-6
\[ = P_{\text{bar}} + P_g \]

Equation 2-7
\[ = 273 + t_s \text{ for metric.} \]

Equation 2-8
\[ = 460 + t_s \text{ for English.} \]

Equation 2-9
\[ v_s = K_p C_p (\sqrt{\Delta p})_{\text{avg}} \frac{T_{s(\text{avg})}}{P_s M_s} \]

Equation 2-10
\[ Q_{sd} = 3,600 (1 - B_{ws}) v_s A \frac{T_{\text{std}}}{T_{s(\text{avg})}} \frac{P_s}{P_{\text{std}}} \]
Appendix B

Federal Reference Method 2F

Equation 2F-1
\[ F_1 = \frac{(P_4 - P_5)}{(P_1 - P_2)} \]

Equation 2F-2
\[ F_2 = C_p \sqrt{\frac{\Delta p_{std}}{(P_1 - P_2)}} \]

Equation 2F-3
\[ \% \text{Diff} = \frac{F_{2 \text{ max}} - F_{2 \text{ min}}}{F_{2 \text{ min}}} \times 100\% \]

Equation 2F-4
\[ M_s = M_d (1 - B_{ws}) + 18.0 B_{ws} \]

Equation 2F-5
\[ P_s = P_{bar} + \frac{P_g}{13.6} \]

Equation 2F-6
\[ T_{s(i)} = 273 + t_{s(i)} \]

Equation 2F-7
\[ T_{s(i)} = 460 + t_{s(i)} \]

Equation 2F-8
\[ v_{a(i)} = K_p F_{2(i)} \sqrt{\frac{(P_1 - P_2) \cdot T_{s(i)}}{P_s \cdot M_s} (\cos \theta_{y(i)}) (\cos \theta_{p(i)})} \]

Equation 2F-9
\[ v_{a(\text{avg})} = \frac{\sum_{i=1}^{n} v_{a(i)}}{n} \]

Equation 2F-10
\[ Q_{sw} = 3,600 (v_{a(\text{avg})}) (A) \left( \frac{T_{s(\text{std})}}{T_{s(\text{avg})}} \right) \left( \frac{P_s}{P_{\text{std}}} \right) \]

Equation 2F-11
\[ Q_{sw} = 3,600 (1 - B_{ws}) (v_{a(\text{avg})}) (A) \left( \frac{T_{s(\text{std})}}{T_{s(\text{avg})}} \right) \left( \frac{P_s}{P_{\text{std}}} \right) \]
Federal Reference Method 3

Equation 3-1
\[ M_d = 0.440(\%CO_2) + 0.320(\%O_2) + 0.280(\%N_2 + \%CO) \]

Federal Reference Method 3A

Equation 3A-1
\[ C_{gas} = \frac{C_{ma} - C_{oa}}{C_m - C_o} (\bar{C} - C_m) + C_{ma} \]

Federal Reference Method 4

Equation 4-1
\[ V_{wc(std)} = \frac{(V_f - V_i)P_w RT_{std}}{P_{std}M_w} = K_1 (V_f - V_i) \]

Equation 4-2
\[ V_{wsg(std)} = \frac{(W_f - W_i)RT_{std}}{P_{std}M_w} = K_2 (W_f - W_i) \]

Equation 4-3
\[ V_{m(std)} = V_m Y \frac{(P_m)(T_{std})}{(P_{std})(T_m)} = K_3 Y \frac{V_m P_m}{T_m} \]

Equation 4-4
\[ B_{ws} = \frac{V_{wc(std)} + V_{wsg(std)}}{V_{wc(std)} + V_{wsg(std)} + V_{m(std)}} \]

Equation 4-5
\[ V_{wc} = \frac{(V_f - V_i)P_w RT_{std}}{P_{std}M_w} = K_1 (V_f - V_i) \]
Appendix B

Equation 4-6
\[ V_{m(\text{std})} = V_m \left( \frac{P_m}{P_{\text{std}}} \right) \left( \frac{T_{\text{std}}}{T_m} \right) \]
\[ = K_2 \frac{V_m P_m}{T_m} \]

Equation 4-7
\[ B_{ws} = \frac{V_{wc}}{V_{wc} + V_{m(\text{std})}} + B_{wm} \]

Method 5

Equation 5-9
\[ \Delta H_{@} = 0.0319 \Delta H \frac{T_m}{P_{\text{bar}}} \left( \frac{\bar{\theta}}{Y^2 V_m^2} \right) \]

Equation 5-1
\[ V_{m(\text{std})} = V_m Y \left( \frac{T_{\text{std}}}{T_m} \right) \left[ \frac{P_{\text{bar}} + \Delta H}{13.6} \right] \]
\[ = K_1 V_m Y \frac{P_{\text{bar}} + \Delta H}{13.6} \frac{1}{T_m} \]

Equation 5-1a
\[ \left[ V_m - (L_p - L_a) \right] \]
\[ \left[ V_m - (L_1 - L_a) \theta_1 - \sum_{i=2}^{n} (L_i - L_a) \theta_i - (L_p - L_a) \theta_p \right] \]

Equation 5-2
\[ V_{w(\text{std})} = \frac{V_{lc} \rho_w R T_{\text{std}}}{M_w P_{\text{std}}} \]
\[ = K_2 V_{lc} \]

Equation 5-3
\[ B_{ws} = \frac{V_{w(\text{std})}}{V_{m(\text{std})} + V_{w(\text{std})}} \]
Appendix B

Equation 5-4
\[ C_a = \frac{m_a}{V_a \rho_a} \]

Equation 5-5
\[ W_a = C_a V_{aw} \rho_a \]

Equation 5-6
\[ c_s = (0.001 \text{ g/mg}) \left( \frac{m_n}{V_{m(\text{std})}} \right) \]

Equation 5-7
\[ I = \frac{100 T_s}{60 \theta V_s P_s A_n} \left[ K_3 V_{ic} + \left( \frac{V_{m} Y}{T_m} \right) \left( P_{bar} + \frac{\Delta H}{13.6} \right) \right] \]

Equation 5-8
\[ I = \frac{100 T_s}{60 T_{std}} \frac{V_{m(\text{std})} P_{std}}{V_s \theta A_n P_s (1-B_{ws})} \]
\[ = \frac{K_4 T_s V_{m(\text{std})}}{P_s V_s A_n \theta (1-B_{ws})} \]

Equation 5-8a
\[ Q = K_4 \frac{P_{bar}}{t_{w} + t_{std}} \frac{V_{w}}{\theta} \]

Equation 5-9
\[ K' = \frac{K_1 V_{m} Y \left( P_{bar} + \frac{\Delta H}{13.6} \right) \sqrt{T_{amb}}}{P_{bar} T_m \theta} \]

Equation 5-10
\[ V_{n(\text{std})} = K_1 V_m \frac{P_{bar} + \frac{\Delta H}{13.6}}{T_m} \]

Equation 5-11
\[ V_{cr(\text{std})} = K' \frac{P_{bar}}{T_{amb}} \theta \]
Appendix B

Equation 5-12
\[ Y = \frac{V_{cr(std)}}{V_{m(std)}} \]

Method 6

Equation 6-1
\[ V_{m(std)} = V_m Y \left( \frac{T_{std}}{T_m} \right) \left( \frac{P_{bar}}{P_{std}} \right) \]
\[ = K_1 Y \left( \frac{V_m P_{bar}}{T_m} \right) \]

Equation 6-2
\[ (V_t - V_{tb}) N \left( \frac{V_{sola}}{V_a} \right) \]
\[ C_{SO_2} = K_3 \frac{(V_t - V_{tb}) N \left( \frac{V_{sola}}{V_a} \right)}{V_{m(std)}} \]

Equation 6-3
\[ RE = \frac{C_d - C_a}{C_a} (100) \]

Equation 6-4
\[ V_{sb(std)} = V_{sb} \left( \frac{T_{std}}{T_{amb}} \right) \left( \frac{P_{bar}}{P_{std}} \right) \]

Equation 6-5
\[ Q_{std} = \frac{V_{sb(std)}}{\theta} \]

Equation 6-6
\[ V_{m(std)} = \bar{Q}_{std} \theta_s (1 - B_{wa}) \left( \frac{P_{bar} + P_{sr}}{P_{bar} + P_c} \right) \]

Equation 6-7
\[ V_{m(std)} = \bar{Q}_{std} \theta_s (1 - B_{wa}) \left( \frac{M_a}{M_s} \left( \frac{P_{bar} + P_{sr}}{P_{bar} + P_c} \right) \right) \]
Appendix B

Method 6A

Equation 6A-1
\[ V_{\text{CO}_2(\text{std})} = 5.467 \times 10^{-4} (m_f - m_{ai}) \]

Equation 6A-2
\[ V_{w(\text{std})} = 1.336 \times 10^{-3} (m_{wf} - m_{wi}) \]

Equation 6A-3
\[ C_{\text{SO}_2} = 32.03 \frac{(V_t - V_{lb})N}{V_{\text{std}} + V_{\text{CO}_2(\text{std})}} \frac{V_{\text{soln}}}{V_s} \]

Equation 6A-4
\[ C_{\text{CO}_2} = \frac{V_{\text{CO}_2(\text{std})}}{V_{\text{m(\text{std})}} + V_{\text{CO}_2(\text{std})}} \times 100 \]

Equation 6A-5
\[ C_w = \frac{V_{w(\text{std})}}{V_{\text{m(\text{std})}} + V_{w(\text{std})} + V_{\text{CO}_2(\text{std})}} \]

Equation 6A-7
\[ m_{\text{SO}_2} = 32.03(V_t - V_{lb})N \left( \frac{V_{\text{soln}}}{V_s} \right) \]

Equation 6A-8
\[ E_{\text{SO}_2} = F_c (1.829 \times 10^9) \frac{m_{\text{SO}_2}}{(m_f - m_{ai})} \]

Method 6C

Equation 6C-1
\[ C_{\text{gas}} = \left( \frac{C - C_o}{C_m - C_o} \right) \frac{C_{ma}}{C_m - C_o} \]
Appendix B

Method 7

Equation 7-1
\[
K_c = 100 \frac{A_1 + 2A_2 + 3A_3 + 4A_4}{A_1^2 + A_2^2 + A_3^2 + A_4^2}
\]

Equation 7-2
\[
V_{sc} = \left( \frac{T_{std}}{P_{std}} \right) (V_f - V_a) \left( \frac{P_f}{T_f} - \frac{P_i}{T_i} \right)
\]
\[
= K_i (V_f - 25 ml) \left( \frac{P_f}{T_f} - \frac{P_i}{T_i} \right)
\]

Equation 7-3
\[m = 2K_c \Delta F\]

Equation 7-4
\[C = K_2 \frac{m}{V_{sc}}\]

Equation 7-5
\[R E = \frac{C_{d} - C_{a}}{C_{a}} (100)\]

Method 8

Equation 8-1
\[
V_{m(\text{std})} = V_m Y \left( \frac{T_{\text{std}}}{T_m} \right) \left( \frac{P_{\text{bar}} + \frac{\Delta H}{13.6}}{P_{\text{std}}} \right)
\]
\[
= K_4 V_m Y \left( \frac{P_{\text{bar}} + \frac{\Delta H}{13.6}}{T_m} \right)
\]

Equation 8-2
\[
C_{H_2SO_4} = K_2 \frac{N (V_f - V_{lb}) \left( \frac{V_{\text{soln}}}{V_a} \right)}{V_{m(\text{std})}}
\]
**Appendix B**

Equation 8-3

\[
C_{SO_2} = K_3 \frac{N(V_t - V_{th})}{V_{m(\text{std})}} \left( \frac{V_{soln}}{V_a} \right)
\]

Equation 8-4

\[
I = \frac{100 T_s}{60 \theta V_s P_s A_n} \left[ K_3 V_{lc} + \left( \frac{V_m Y}{T_m} \right) \left( \frac{P_{bar}}{13.6} + \frac{\Delta H}{13.6} \right) \right]
\]

Equation 8-5

\[
I = \frac{100 T_s V_{m(\text{std})} P_{\text{std}}}{60 T_{\text{std}} v_s \theta A_n P_s (1 - B_{\text{we}})} = \frac{K_4 T_s V_{m(\text{std})}}{P_s v_s A_n \theta (1 - B_{\text{ws}})}
\]

**Method 11**

Equation 11-1

\[
N_s = 2.039 \frac{W}{V_s}
\]

Equation 11-2

\[
N_A = 0.2039 \frac{W}{V_s}
\]

Equation 11-3

\[
N_i = N_f \frac{V_f}{V_i}
\]

Equation 11-4

\[
V_{m(\text{std})} = V_m Y \left[ \frac{T_{\text{std}}}{T_m} \left( \frac{P_{bar}}{P_{\text{std}}} \right) \right]
\]
Appendix B

Equation 11-5
\[ C_{H2S} = \frac{K[(V_{tN} - V_{tT} N_T)_{sample} - (V_{tN} N_T)]}{V_{m(Std)}} \]

**Method 15**

Equation 15-1
\[ C = \frac{(K)(P_f)}{ML} \]

Equation 15-2
\[ SO_2 \text{ equivalent} = \sum (H_2S, COS, 2CS_2) d \]

Equation 15-3
\[ \text{Average } SO_2 \text{ equivalent} = \frac{\sum_{i=1}^{N} SO_2 \text{ equiv}_i}{N} \]

**Method 15A**

Equation 15A-1
\[ V_{ms(Std)} = \frac{V_{ms} Y(T_{std})(P_{bar})}{(T_m)(P_{std})} = \frac{K_1 Y(V_m)(P_{bar})}{T_m} \]

Equation 15A-2
\[ V_{mc(Std)} = \frac{k_f Y_c (V_m)(P_{bar})}{T_m} \]

Equation 15A-3
\[ C_{TRS} = \frac{K_2 (V_t - V_{tb}) N \left( \frac{V_{soln}}{V_a} \right)}{V_{ms(Std)} - V_{mc(Std)}} \]

Equation 15A-4
\[ C_{RG} = \frac{C_{COS}}{Q_{COS} + Q_{N2}} \]

Equation 15A-5
\[ R = \frac{C_{TRS}}{C_{RG}} \times 100 \]
**Appendix B**

**Method 16**

Equation 16-1

\[ C = K \frac{P_r}{M_L} \]

Equation 16-2

\[ TRS = \sum (H_2S, MeSH, DMS, 2DMDS) d \]

Equation 16-3

\[ Average TRS = \frac{\sum_{i=1}^{N} TRS_i}{N(1-B_{WO})} \]

Equation 16-4

\[ C = \frac{\sum_{i=1}^{N} S_i}{N} \]

**Method 16A**

Equation 16A-1

\[ V_{m(std)} = V_m Y \frac{T_{std}}{T_m} \]

\[ P_{bar} = K_t Y \frac{V_m}{T_m} \]

Equation 16A-2

\[ C_{TRS(ppm)} = \frac{K_2 (V_{t} - V_{tb}) N \left( \frac{V_{soln}}{V_a} \right)}{V_{m(std)}} \]

Equation 16A-3

\[ C_{RG} = \frac{(Q_{H2S})(C_{H2S})}{Q_{H2S} + Q_{CG}} \]

Equation 16A-4

\[ R = \frac{C_{TRS}}{C_{RG}} \times 100 \]

Equation 16A-5

\[ N_f = \frac{1}{ml Na_2S_2O_3 Consumed} \]
Appendix B

Equation 16A-6
\[ N_i = \frac{N_c \cdot V_c}{V_i} \]

Equation 16A-7
\[ V_{m(\text{std})} = (Q_{\text{std}}) (0.1) (1-B_{wa}) \frac{M_a}{M_b} \]

Equation 16A-8
\[ C_{H2S} = \frac{K N_T (V_{TB} - V_T)}{V_{m(\text{std})}} \]

Method 16B

Equation 16B-1
\[ C_{TRS} = (C_{SO2})(d) \]

Equation 16B-2
\[ C_{TRS} = \frac{\sum_{i=1}^{n} C_{TRS}}{N} \]

Method 18

Equation 18-1
\[ C_s = \frac{10^6 (\bar{x} q_c)}{q_c + q_d} \]

Equation 18-2
\[ C_s = 106 \bar{x} \left( \frac{q_{c1}}{q_{c1} + q_{d1}} \right) \left( \frac{q_{c2}}{q_{c2} + q_{d2}} \right) \]

Equation 18-3
\[ C_s = \frac{G_v \times 10^6 \frac{293}{T_s} \frac{P_v}{760}}{V_m \ Y \ \frac{293}{T_m} \frac{P_m}{1000}} \]

\[ = \frac{G_v \times 10^2 \frac{P_v}{T_s} \frac{T_m}{P_m}}{V_m \ Y} \]
Appendix B

Equation 18-4

\[ C_s = \frac{L_v \rho (24.055 \times 10^6)}{M V_m Y_293 P_m 1000} = 6.24 \times 10^4 \frac{L_v \rho T_m}{M V_m Y P_m} \]

Equation 18-5

\[ C_c = \frac{C_s P_r T_i F_r}{P_l T_r (1 - B_{ws})} \]

Method 20

Equation 20-1

\[ C_d = \frac{C_w}{1 - B_{ws}} \]

Equation 20-2

\[ F_o = \frac{0.209 F_d}{F_c} \]

Equation 20-3

\[ X_{CO_2} = \frac{5.9}{F_o} \]

Equation 20-4

\[ C_{adj} = C_d \frac{5.9}{20.0 - %O_2} \]

Equation 20-5

\[ C_{adj} = C_d \frac{X_{CO_2}}{%CO_2} \]

Equation 20-6

\[ E = C_d F_d \frac{20.9}{20.9 - %O_2} \]

Equation 20-7

\[ E = C_d F_c \frac{100}{%CO_2} \]

Equation 20-8

\[ E = C_w F_c \frac{100}{%CO_{2w}} \]
Appendix B

Method 23

Equation 23-1
\[
RRF_i = \frac{1}{n} \sum_{j=1}^{n} \frac{A_{cij} m_{ci}^{*}}{A_{*cij} m_{ci}}
\]

Equation 23-2
\[
C_i = \frac{m_i^{*} A_i}{A_i * RRF_i V_{\text{mstd}}}
\]

Equation 23-3
\[
RRF_{rs} = \frac{A_{cl} m_{rs}^{*}}{A_{rs} m_{ci}^{*}}
\]

Equation 23-4
\[
R^{*} = \frac{A_{i} m_{rs}^{*}}{A_{rs} RRF_{rs} m_{i}^{*}} \times 100\%
\]

Equation 23-5
\[
RRF_{s} = \frac{A_{cl} m_{s}^{*}}{A_{cs} m_{ci}^{*}}
\]

Equation 23-6
\[
R_{s} = \frac{A_{s} m_{i}^{*}}{A_{i} * RRF_{s} m_{s}} \times 100\%
\]

Equation 23-7
\[
MDL = \frac{2.5 A_{al} m_{i}^{*}}{A_{cl} * RRF_{i}}
\]

Equation 23-8
\[
C_{Tr} = \sum_{i=1}^{n} C_i
\]

Method 25

Equation 25-1
\[
\Delta P = 0.01 \frac{FP_b \theta}{V_t}
\]

Equation 25-2
\[
V_s = 0.3857 V \left[ \frac{P_t}{T_t} - \frac{P_{ti}}{T_{ti}} \right]
\]
Appendix B

Equation 25-3

\[ C_t = \left[ \frac{P_t}{T_t} \right] \left[ \frac{1}{P_i} \sum_{j=1}^{r} C_{m_j} \right] \left[ \frac{1}{T_f - T_i} \right] \]

Equation 25-4

\[ C_c = 0.3857 \frac{V_r P_f}{V_s T_f} \left[ \frac{1}{q} \sum_{k=1}^{q} C_{cm_k} \right] \]

Equation 25-5

\[ C = C_t + C_c \]

Equation 25-6

\[ m_c = 0.4993 C \]

Equation 25-7

\[ \text{Percent recovery} = 1.604 \frac{M V_r P_f}{L P T_f} \frac{C_{cm}}{N} \]

Equation 25-8

\[ \text{RSD} = \frac{100 \sqrt{\frac{\sum(x_i - \bar{x})^2}{n-1}}}{\bar{x}} \]

Method 25A

Equation 25A-1

\[ C_c = K C_{\text{meas}} \]

Method 26

Equation 26-1

\[ \frac{\mu g}{ml} \text{Cl}^- = g \text{ of NaCl} \times 10^3 \times \frac{35.453}{58.44} \]

Equation 26-2

\[ \frac{\mu g}{ml} \text{Br}^- = g \text{ of NaBr} \times 10^3 \times \frac{79.904}{102.90} \]

Equation 26-3

\[ \frac{\mu g}{ml} \text{F}^- = g \text{ of NaF} \times 10^3 \times \frac{18.998}{41.99} \]
Appendix B

Equation 26-4
\[ m_{\text{HX}} = K V_s (S_x^- - B_x^-) \]

Equation 26-5
\[ m_{X_2} = V_s (S_x^- - B_x^-) \]

Equation 26-6
\[ C = K \frac{m_{\text{HX, X2}}}{V_{m(sst)}} \]

Method 26A

Equation 26A-1
\[ \mu g \frac{Cl^-}{ml} = g \text{ of } NaCl \times 10^3 \times \frac{35.453}{58.44} \]

Equation 26A-2
\[ \mu g \frac{Br^-}{ml} = g \text{ of } NaBr \times 10^3 \times \frac{79.904}{102.90} \]

Equation 26A-3
\[ \mu g \frac{F^-}{ml} = g \text{ of } NaF \times 10^3 \times \frac{18.998}{41.99} \]

Equation 26A-4
\[ m_{\text{HX}} = K V_s (S_x^- - B_x^-) \]

Equation 26A-5
\[ m_{X_2} = V_s (S_x^- - B_x^-) \]

Equation 26A-6
\[ C = K \frac{m_{\text{HX, X2}}}{V_{m(sst)}} \]

Method 29

Equation 29-1
\[ M_{m} = C_{a1} F_d V_{\text{soln,1}} \]

Equation 29-2
\[ M_{b} = C_{a2} F_a V_a \]

Equation 29-3
\[ M_{i} = (M_{m} - M_{mb}) + (M_{b} - M_{bbb}) \]
Appendix B

Equation 29-4
\[ H_g = \frac{Q_{f_{h1}}}{V_{f_{1B}}} (V_{so_{1n,1}}) \]

Equation 29-5
\[ H_{g_{bh2}} = \frac{Q_{bh2}}{V_{f_{2B}}} (V_{so_{in,2}}) \]

Equation 29-6
\[ H_{g_{bh3(A,B,C)}} = \frac{Q_{bh3(A,B,C)}}{V_{f_{3(A,B,C)}}} (V_{so_{n,3(A,B,C)}}) \]

Equation 29-7
\[ H_{g_{bh}} = H_{g_{bh2}} + H_{g_{bh3A}} + H_{g_{bh3B}} + H_{g_{bh3C}} \]

Equation 29-8
\[ H_{g_{t}} = (H_{g_{f_{h}} - H_{g_{f_{mb}}}}) + (H_{g_{bh}} - H_{g_{bhb}}) \]

Equation 29-9
\[ C_s = \frac{K_4 M_t}{V_{m_{(std)}}} \]

Method 0010

Max POHC\textsubscript{c} Mass\textsubscript{eff(std)} = Max POHC\textsubscript{c} conc

\[ \frac{LDL_{POHC} \times 10}{POHC_{c,conc}} = V_{TBC} \]

\[ C_{POHC} (\mu g/ml) \times \text{sample volume (ml)} = \text{amount (\mu g) of POHC in sample} \]

Method 0030

\[ C_g = \frac{\text{Total weight of CPD in sample, \mu g (i.e. VOST tubes & condensator)}}{\text{Volume of sample at standard conditions, dscm}} \]

Method 0050

Equation 0050-4
\[ m_{HCl} = S \times V_s \times \frac{36.46}{35.45} \]
Appendix B

Equation 0050-5
\[ m_{\text{Cl}_2} = S \times V_2 \]

Equation 0050-6
\[ C = K \times \frac{m}{V_{m(\text{std})}} \]

Method 0051
\[ m_{\text{HCl}} = S \times V_s \times \frac{36.46}{35.45} \]
\[ m_{\text{Cl}_2} = S \times V_2 \]
\[ C = K \times \frac{m}{V_{m(\text{std})}} \]

Method 0060
\[ M_{\text{fH}} = C_{a1} \times F_d \times V_{\text{solv},1} \]
\[ M_{\text{bH}} = C_{a2} \times F_a \times V_a \]
\[ M_t = (M_{\text{fH}} - M_{\text{fHb}}) + (M_{\text{bH}} - M_{\text{bHb}}) \]

Method 0061
\[ m = (S - B) \times V_s \times d \]
The Inspector's Tool Kit
Appendix C

The Inspector’s Tool Kit Items

1. **Dial Caliper**- Used for measuring nozzle diameter and inspection of pitot tubes. The dial caliper should have graduation units of 0.001 inches. Price: $38.

2. **Level Indicator**- Used for verification of proper construction dimensions and spacing requirements for Type S pitot tube. Also used in measuring yaw angle of Type S pitot tube during verification of absence of cyclonic flow at a sampling location. Price: $28.

3. **Thermocouple Simulator Source**- Used with the Method 5 meter box for checking accuracy of temperature displays and controllers. The thermocouple simulator source should be designed for Type K thermocouples with 22 precise test points. The temperature range should be from 0 to 2100 F. Price: $225.

4. **Calibration Orifice Set**- The Calibration Orifice Set is used for calibrating and auditing Method 5 metering system. The set should include 5 calibrated orifices with ½ inch quick connects. In addition, a disk with appropriate software/spreadsheet for performing calculations should also be part of the Calibration Orifice Set. Price: $395.

5. **IsoCal Software**- IsoCal Software should be a MicroSoft Excel workbook/3.5” disk designed for integrated isokinetic source sampling calculations. The IsoCal Software should provide all the worksheets necessary to perform US EPA’s Federal Reference Methods 1 through 5 test setup, data entry, and data reduction. Price: $195.

6. **Isokinetic Slide Rule or Calculator**- The Isokinetic Slide Rule or Calculator performs isokinetic stack sampling calculations to calculate a proper Delta H based upon a Delta P reading, once all the various constants/source parameters have been entered properly into the calculator or performed correctly with the slide ruler. Price: $195.

7. **Pocket Barometer**- Used to determine atmospheric pressure, readable to within 0.02 inches Hg divisions. Price: $30.

8. **Modular Pitot Tube**- A Modular Pitot Tube, capable of extension to 1 meter, used to measure stack gas velocity, static pressure, and cyclonic flow. The pitot tube would be used in conjunction with a hand-held digital manometer (0-19.99 inches). Price: $80.

9. **Hand-Held Manometer**- Used in conjunction with the Modular Pitot Tube. The hand-held digital manometer (readable to 0.01 inches of water) is a useful alternative to the standard size manometer or magnehelic gauge; and is especially ideal for field monitoring and troubleshooting. Measures positive, negative, or differential air pressure. Price: $55.
10. **Hand-held Digital Thermometers**- Used to measure stack temperature and various temperatures associated with the Method 5 sampling train. Should be a Type K thermocouple coupled to a hand-held control unit with capability of readings in the range of −58 to 1999.9°F. Price: $155.

11. **Bull's Eye Level**- Used for indicating level of pitot tube during evaluation with Level Indicator. The Bull's Eye Level may also be placed on the pitot tube (has thumb screw clamps) during sampling to indicate proper orientation. Price: $25.

12. **Tape Measure**- Used to document stack geometry and sampling port location from upstream and downstream disturbances. Material should be stainless steel and length of 100 feet. Price: $75


14. **Stack Sampling Nomographs for Field Estimations**- Field nomographs used in estimating or checking data used in stack sampling. Four basic groups of nomographs are helpful to the Agency inspector. They are:

   - Moisture nomographs (Wet bulb/dry bulb, combustion calculation, FRM 4 etc.);
   - Excess air nomographs (Temperature versus excess air, excess air versus composition etc.);
   - Velocity and volumetric flow rate nomographs (Coal-fired equipment, volume versus heat input, pitot tube velocity readings etc.); and
   - Concentration and mass emission rate nomographs (ppm emissions, pounds per million BTU, pounds per hour etc.).