



Designation: **D5835 – 95 (Reapproved 2013) D5835 – 20**

Standard Practice for Sampling Stationary Source Emissions for the Automated Determination of Gas Concentrations¹

This standard is issued under the fixed designation D5835; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope

1.1 This practice² covers procedures and equipment that will ~~permit, within certain limits, permit~~ representative sampling for the automated determination of gas concentrations of effluent gas ~~streams—streams with limitations as described below~~. The application is limited to the determination of oxygen (O_2), carbon dioxide (CO_2), carbon monoxide (CO), sulfur dioxide (SO_2), nitric oxide (NO), nitrogen dioxide (NO_2), and total oxides of nitrogen (NO_x).

1.2 Velocity measurements are required to determine the mass flow rates of gases. This is not included in this practice.

1.3 There are some combustion processes and ~~situations~~conditions that may limit the applicability of this practice. Where such conditions exist, caution and competent technical judgment are required, especially when dealing with any of the following:

1.3.1 Corrosive or highly reactive components,

1.3.2 High vacuum, high pressure, or high temperature gas streams,

1.3.3 Wet flue gases,

1.3.4 Fluctuations in velocity, temperature, or concentration due to uncontrollable variation in the process,

1.3.5 Gas stratification due to the non-mixing of gas streams,

1.3.6 Measurements made using environmental control devices, and

1.3.7 Low levels of gas concentrations.

1.4 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.5 *This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety, health, and environmental practices and determine the applicability of regulatory limitations prior to use.* For more specific safety precautions, refer to 5.1.4.8, 5.2.1.6, and 6.2.2.1.

¹ This practice is under the jurisdiction of ASTM Committee D22 on Air Quality and is the direct responsibility of Subcommittee D22.03 on Ambient Atmospheres and Source Emissions.

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² This practice is based on ISO 10396, “Stationary source emissions—Sampling for the automated determination of gas concentrations,” available from International Organization for Standardization, Casa Postale 56, CH-1211, Geneva, Switzerland.

1.6 This international standard was developed in accordance with internationally recognized principles on standardization established in the Decision on Principles for the Development of International Standards, Guides and Recommendations issued by the World Trade Organization Technical Barriers to Trade (TBT) Committee.

2. Referenced Documents

2.1 ASTM Standards:³

[D1356 Terminology Relating to Sampling and Analysis of Atmospheres](#)

[D1608 Test Method for Oxides of Nitrogen in Gaseous Combustion Products \(Phenol-Disulfonic Acid Procedures\)](#)

[D3154 Test Method for Average Velocity in a Duct \(Pitot Tube Method\)](#)

2.2 Other Document:

[40 CFR Part 60 CFR Part 60](#), Standards of Performance for Stationary Sources, Appendix A, Test Methods 2, 3, 3a, 6, 6c, 7, 7e, and 10⁴

3. Terminology

3.1 Definitions:

3.1.1 For definitions of terms used in this practice, refer to Terminology [D1356](#).

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4. Summary of Practice

4.1 This practice describes representative sampling of gases in a duct, duct or stack gas sampling, including both extractive and non-extractive sampling. In extractive sampling, these gases are conditioned to remove aerosols, particulate matter, and other interfering substances before being conveyed to the instruments. In non-extractive sampling, the measurements are made in-situ; therefore, no sample conditioning except filtering is required. is not necessary beyond filtration.

4.1.1 *Extractive Sampling*—Extractive sampling includes consists of extraction of the sample, removal of interfering materials, and maintenance retention of the gas analyte concentration without change throughout the sampling system for subsequent analysis by appropriate instrumentation (see [Fig. 1](#)).

4.2 *Non-extractive Sampling*—Non-extractive sampling does not involve removal of a sample, and sampling is sample from the duct or stack with sampling confined to the gas stream in the stack or duct (see [Figs. 2 and 3](#)).

5. Representative Factors

5.1 Nature of the Source:

5.1.1 The representativeness of the determination of gaseous analyte concentration determination in enclosed gas streams depends on several factors:

5.1.1.1 The heterogeneity of the process stream, such as variations in concentration, temperature, or velocity across the duct or stack caused by moisture or gas stratification,

5.1.1.2 Gas leakage or air infiltration and continuous gas reactions, and

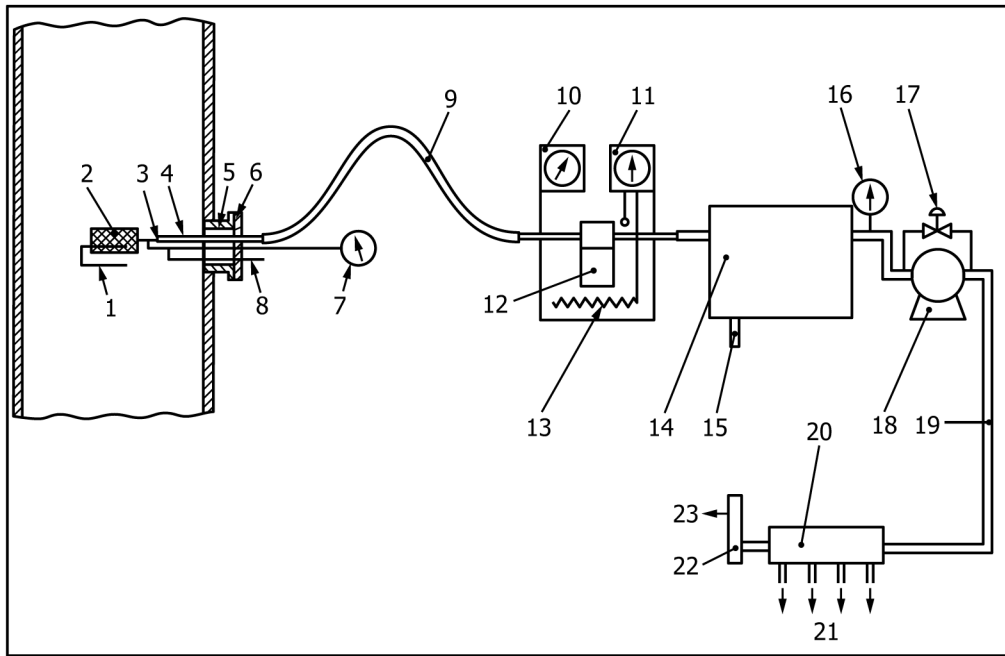
5.1.1.3 Random errors due to the finite nature of the sample and the sampling procedure adopted to obtain a representative sample.

5.1.2 Representativeness may be difficult to achieve for the following reasons:

5.1.2.1 Nature of the source (for example, cyclic, continuous, or batch),

³ For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

⁴ Available from U.S. Government Printing Office, Superintendent of Documents, 732 N. Capitol St., NW, Washington, DC 20401-0001, <http://www.access.gpo.gov>.



NOTE—Key:

1	Baffle	13	Heater
2	In-stack Filter	14	Refrigeration Unit
3	Tee	15	Water Discharge
4	Probe	16	Vacuum Gauge
4	Probe	16	Vacuum Gauge
5	Sampling Port	17	Bypass Valve
6	Cap	18	Pump
7	Pressure Gauge	19	Sampling Line (Heating Optional)
7	Pressure Gauge	19	Sampling Line (Heating Optional)
8	To Zero and Span Gases	20	Manifold
9	Heat-traced Sampling Line	21	To Analyzer(s)
10	Temperature Controller (Line)	22	Rotameter
11	Temperature Controller (Box)	23	Vent
12	Filter		

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<https://standards.iteh.ai/catalog/standards/astm-d5835-20> **FIG. 1 Extractive Sampling and Conditioning System**

5.1.2.2 Concentration level of the gas, analyte(s),

5.1.2.3 Size of the source, and

5.1.2.4 Configuration of the duct or stack network where samples are extracted.

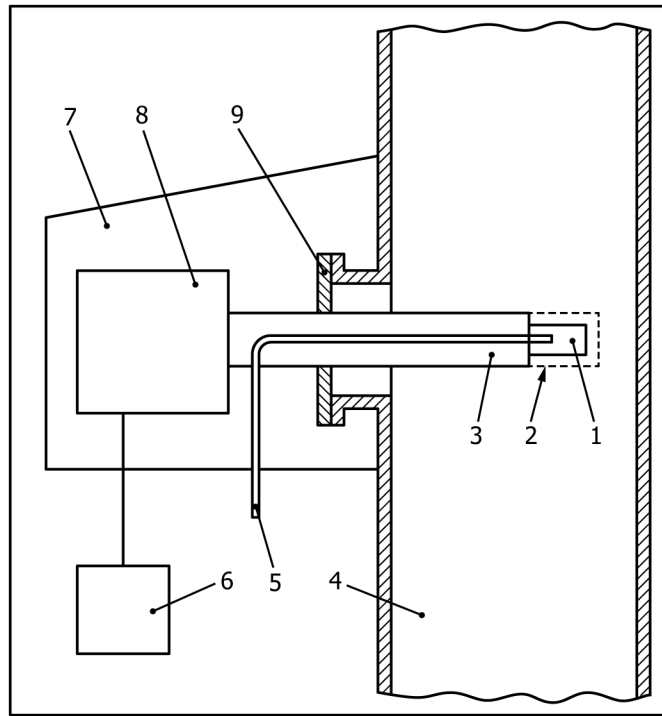
5.1.3 Where there are difficulties issues due to the nature of the source as noted in 5.1.2, establish the concentration profile for each operating condition and to determine the best sampling location.

5.1.3.1 Some sources may have more variability in process—their operating processes (for example, cyclic variation) and, consequently, any time dependent measurement may be less representative of the average concentration if a full cycle of variability is not sampled.

5.1.4 Before any measurements are carried out, it is necessary to become familiar with the pertinent/relevant operating characteristics of the process from which emissions are to be sampled and determined. These operating characteristics include, but are not necessarily limited to, the following:

5.1.4.1 Mode of process operation (cyclic, batch charging, or continuous),

5.1.4.2 Process feed rates and composition,



NOTE—Key:

- 1
- 2
- 3
- 4
- 5

- Measurement Cell 6
- Probe Filter 7
- Probe 8
- Duct or Stack 9
- Gas Calibration Line

- Data Recorder
- Protective Hood
- Transceiver
- Probe Mounting

FIG. 2 Non-Extractive Non-extractive Point Monitor

5.1.4.3 Fuel rates and composition,

5.1.4.4 Normal operating gas temperatures and pressures,

5.1.4.5 Operating and removal efficiency of the pollution control equipment,

5.1.4.6 Configuration of the ducts to be sampled leading to gas stratification,

5.1.4.7 Volumetric gas flow rates, and

5.1.4.8 Expected gas composition and likely interfering substances. (**Warning**—Exercise caution if the duct to be sampled is under pressure or vacuum, or at a high temperature.)

5.2 Location:

5.2.1 Inspection Parameters—Perform an inspection of the physical characteristics of the test site to evaluate factors such as:

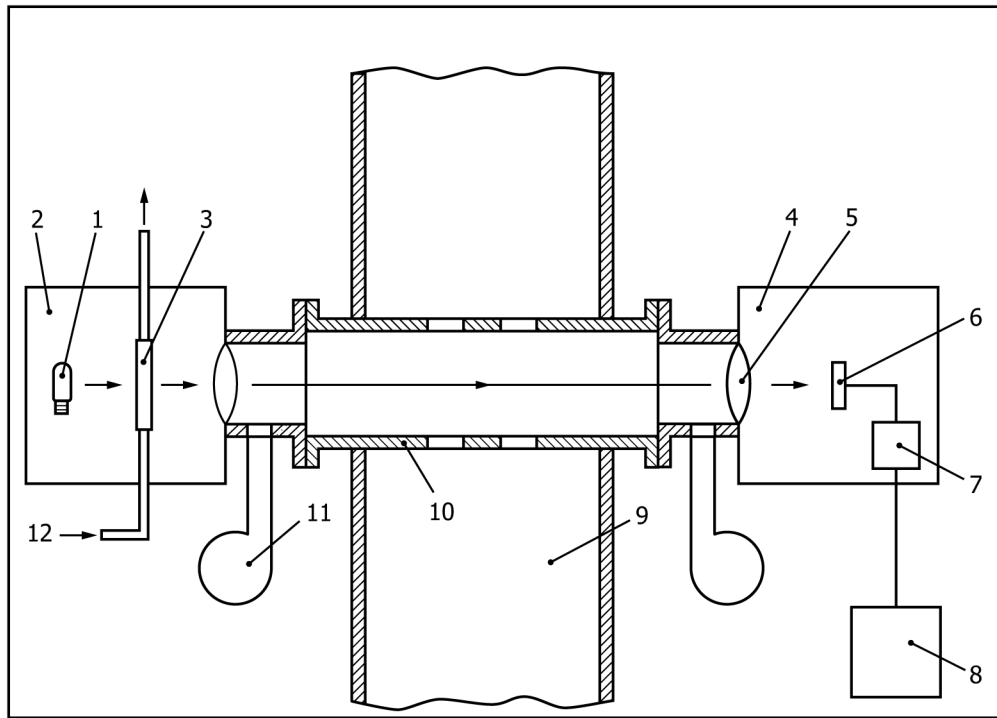
5.2.1.1 Safety of the personnel,

5.2.1.2 Location of the flow disturbances,

5.2.1.3 Accessibility of the sampling site,

5.2.1.4 Available space for the sampling equipment and instrumentation and possible scaffolding requirements,

5.2.1.5 Availability of suitable electrical power, compressed air, water, steam, etc., and



NOTE—Key:

1	Lamp	7	Electronic Module
2	Transmitter Assembly	8	Data Recorder
3	Internal Gas Calibration Cell	9	Stack or Duct
4	Receiver Assembly	10	Alignment/Calibration Pipe
5	Protective Windows	11	Purge Air Blower
6	Detector	12	Gas Calibration Line

FIG. 3 Non-Extractive Non-extractive Path Monitor

5.2.1.6 Sampling port locations. (**Warning**—Use the electrical equipment in accordance with the local safety requirements. Where a potentially explosive or hazardous atmosphere is suspected, apply particular attention and precautions to ensure the safety of the operations.)

5.2.2 Sampling Site Location:

5.2.2.1 It is necessary to ensure that the gas concentrations measured are representative of the average conditions inside the duct or stack. The requirements for ~~the~~ extractive sampling of gas may be not as ~~stringent~~ rigorous as those for particulate material. It is important that the sampling location be ~~removed from~~ free of any obstructions that will seriously disturb the gas flow in the duct or stack. ~~The~~ Since the pollutant can have cross sectional variation. ~~The~~ variation, the concentration at various points of the cross-section shall first be checked, in order to assess the homogeneity of the flow and to detect any infiltration of air or gas stratification, etc. If a preliminary analysis of ~~cross-section at~~ cross-sectional measurements ~~taken~~ indicates more than $\pm 15\%$ $\pm 15\%$ variation in concentrations, and if an alternative acceptable location is ~~not available~~, unavailable, multi-point sampling is recommended.

5.2.2.2 Multi-point sampling may be achieved either by moving the probe from point to point or having a probe with multiple access ports. Usually, the ~~cross-sectional~~ cross-sectional concentration of gaseous pollutants is uniform, because of the diffusion and turbulent mixing. ~~If so~~, In this case, it is only necessary to sample at one point within the stack or duct to determine the average concentration. Extract gas samples near the center of the stack sampling site. When using nonextractive systems, obtain a concentration as representative as possible, but ~~and~~ ensure that the instrument location is representative.

5.3 Gas Concentration, Velocity, and Temperature Profile—Before ~~commencing~~ sampling, determine if there are any spatial or temporal fluctuations in the gas concentrations by conducting a preliminary survey of the gas concentration, temperature, and velocity. Measure the concentration, temperature, and velocity at the sampling points several times to obtain their spatial and temporal profiles. Conduct this survey when the plant facility is operating under conditions that ~~will be~~ are representative of normal operation and determine whether the sampling position is suitable and whether the conditions in the duct are ~~satisfactory~~ acceptable for ensuring representative sampling (see 5.1.2).

5.3.1 The following test methods may be used to determine gas concentration, temperature, and velocity:

5.3.1.1 O_2 —Test Method **D3154**, EPA Test Methods 3 and 3a,

5.3.1.2 CO_2 —Test Method **D3154**, EPA Test Methods 3 and 3a,

5.3.1.3 CO —EPA Test Method 10,

5.3.1.4 SO_2 —EPA Test Methods 6 and 6c,

5.3.1.5 NO_x —Test Method **D1608**, EPA Test Methods 7 and 7e,

5.3.1.6 Gas Temperature—Test Method **D3154**, EPA Test Method 2, and

5.3.1.7 Gas Velocity—Test Method **D3154**, EPA Test Method 2.

5.4 Other Factors—The operational principle of operation and the components of ~~the instrument~~ instrumental systems can significantly affect the degree to which a collected sample is representative of the measured gas in the source. For example, a point sampling extractive system requires more attention to the sampling site location than an across-the-stack in-situ sampling system. Furthermore, sampling lines should not be composed of materials that have gas adsorbing properties ~~that can affect the response time of the measurement section~~ (see **Table A1.1**).

5.4.1 Exercise care to preserve the integrity of the sample taken, by a ~~good~~ selection of equipment, suitable equipment and appropriate heating, drying, and leak testing, etc. In addition, other factors such as corrosion, synergies, reaction with components, decomposition, and adsorption that might affect the integrity of a sample ~~sample~~ should be considered.

6. Equipment

6.1 Recommended construction materials are described in **Annex A1**.

6.2 Components of Extractive Sampling Equipment:

6.2.1 *Primary Filter*—The filter medium shall be constructed of an appropriate alloy (~~such~~such as a ~~specific~~ stainless steel cast ~~alloy~~alloy, quartz borosilicate, ceramics, or another suitable material. A filter that ~~retain~~traps particles greater than 10 μm in size is recommended. A secondary filter might be required as well (see 6.2.4). The filter ~~medium~~ may be ~~located~~placed outside the duct or at the tip of the sample probe (6.2.2). If placed at the tip of the probe, a deflector plate may be added to prevent particle build-up on the leading edge of the filter. This will prevent blockage of the filter. Avoid contamination of the filter with particulate matter where condensate may react with gases, resulting in ~~erroneous result~~compromised results.

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