

SLOVENSKI STANDARD SIST ISO 11632:1999

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Stationary source emissions -- Determination of mass concentration of sulfur dioxide -- Ion chromatography method

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Émission de sources fixes -- Détermination de la concentration en masse de dioxyde de soufre -- Méthode par chromatographie ionique sist/6fd78493-d508-4a35-8d60-05e9f0f37d7c/sist-iso-11632-1999

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Stationary source emissions

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INTERNATIONAL STANDARD

ISO 11632

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Stationary source emissions — Determination of mass concentration of sulfur dioxide — Ion chromatography method

Émission de sources fixes — Détermination de la concentration en masse iTeh Sde dioxyde de soufre — Méthode par chromatographie ionique

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Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

International Standard ISO 11632 was prepared by Technical committee ISO/TC 146, *Air quality*, Subcommittee SC 1, *Stationary source emissions*.

Annex A of this International Standard is for information only.

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Introduction

For determining the mass concentration of sulfur dioxide in waste gas of stationary emissions sources, several integrated sampling and analysis methods exist. These include methods for independent manual sampling [ISO 7934] and for automated measuring systems [ISO 7935]. The latter have found a greater usage since a continuous measurement of sulfur dioxide, often specified by environmental authorities, can be obtained. The manual methods are required however, in calibrating automated measuring instruments.

This International Standard offers an alternative method to ISO 7934, replacing the Thorin method for analysing sulfate ions by a method based on ion chromatography. In addition, this International Standard is intended to apply to a lower emission range than ISO 7934.

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Stationary source emissions — Determination of mass concentration of sulfur dioxide — Ion chromatography method

1 Scope

This International Standard specifies a method for the determination of the mass concentration of sulfur dioxide emitted from combustion facilities and technical processes, and defines the most important performance characteristics.

The method described in this International Standard has been tested for a sulfur dioxide concentration range of 6 mg/m³ to 333 mg/m³ with sampling periods of 30 min. It is applicable to mass concentrations of sulfur dioxide exceeding this range by carrying out an appropriate dilution of the sample solutions prior to the analysis or by using larger volumes of absorption solution, and to sulfur dioxide concentrations below this range by extending the sampling period.

This International Standard is applicable to the analysis of samples containing negligible levels of sulfur trioxide and volatile sulfates (< 5 % of the expected sulfur dioxide concentration), and ammonia (< 5 mg/m³). All concentrations are based on dry gas at a temperature of 273,2 K and pressure of 101,3 kPa. uarus.iten.a.

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2 Normative references https://standards.iteh.ai/catalog/standards/sist/6fd78493-d508-4a35-8d60-

05e9f0f37d7c/sist-iso-11632-1999 The following standards contain provisions which, through reference in this text, constitute provisions of this International Standard. At the time of publication, the editions indicated were valid. All standards are subject to revision, and parties to agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent editions of the standards indicated below. Members of IEC and ISO maintain registers of currently available valid International Standards.

ISO 6879:1995, Air quality — Performance characteristics and related concepts for air quality measuring methods.

ISO 7934:1989, Stationary source emissions — Determination of the mass concentration of sulfur dioxide — Hydrogen peroxide/barium perchlorate/Thorin method.

ISO 7935:1992, Stationary source emissions — Determination of the mass concentration of sulfur dioxide — Performance characteristics of automated measuring methods.

ISO 10396:1993, Stationary source emissions — Sampling for the automated determination of gas concentrations.

ISO Guide 25:1990, General requirements for the competence of calibration and testing laboratories.

3 Principle

A representative sample of waste gas is extracted via a temperature-controlled probe, filtered and drawn through a hydrogen peroxide solution for a specified time and flow rate. The sulfur dioxide in the waste gas sample is absorbed by the solution and sulfate anions are formed. The mass concentration of sulfate in the absorption solution is subsequently determined using ion chromatography.

4 Reagents

During the analysis, use only reagents of recognized analytical grade. The sulfate-free water shall have an electrical conductivity of < 0,01 mS/m and shall not contain particulate matter of a particle size > 0,45 μ m. Normal, accepted laboratory safety practices should be followed during reagent preparation.

4.1 Absorption solution, 3 % H₂O₂

Pipette 100 cm³ of a 27 % (mole fraction) to 30 % (mole fraction) solution of hydrogen peroxide (H_2O_2) into a 1000 cm³ one-mark volumetric flask. Make up to the mark with water and mix well. Prepare this solution if possible on the day of use.

4.2 Eluent solution

The choice of eluent depends on the manufacturer's separator column and detector. For the exact composition of the eluent, refer to the instructions given by the manufacturer.

NOTE For an ion chromatograph using the suppressor technique, a typical eluent is a solution of 1.7×10^{-3} mol/l NaHCO₃ and 1.8×10^{-3} mol/l Na₂CO₃.

4.3 Standard sulfate stock solution, 10.4×10^{-3} mol/l SO₄²⁻

Dissolve 1,8141 g of analytical grade potassium sulfate (K_2SO_4) in reagent water and dilute to 1 l using a 1000 cm³ one-mark volumetric flask. 1 cm³ of stock solution corresponds to 1 mg of $SO_4^{2^2}$.

NOTE Standard sulfate stock solution is stable for at least 28 days when stored at 277 K. Calibration standards are prepared by diluting the standard stock solution with the absorption solution as specified in 7 4.2.

4.4 Regeneration solution for suppressor ISO 11632:1999

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For the exact composition of the suppressor regeneration solution, refer to the instructions given by the manufacturer of the suppressor.

NOTE An example is a solution of $12,5 \times 10^{-3}$ mol/l H₂SO₄.

5 Apparatus

5.1 Sampling equipment

5.1.1 General

Alternative variations of the sampling equipment fulfilling the specified performance requirements for each component may be used. The performance characteristics set out in clause 9 refer, however, to the examples of sampling equipment described in 5.1.1 to 5.1.16. It is important that all parts of the sampling equipment upstream of the first absorber are heated and that the components shall not react with or absorb SO₂.

NOTE In special cases an unheated gas connector line may be used between the heated filter and first absorber, but this must be thoroughly rinsed with absorption solution after sampling and the washings combined with the sample.

5.1.2 Sampling probe

The probe shall be of borosilicate glass or fused silica tube with a spherical ground joint at one end. Alternative probes of different lengths and inner diameters may be used, but the residence time of the sample gas in the probe shall be minimized. The sampling probe, which is surrounded by a heating jacket, shall be protected and positioned

using a metal outer tube. A clamp screw shall be used to adjust the probe length to reach the representative measurement point in the measurement plane of the waste gas duct.

NOTE An example of suitable probe dimensions is shown in figure 1.

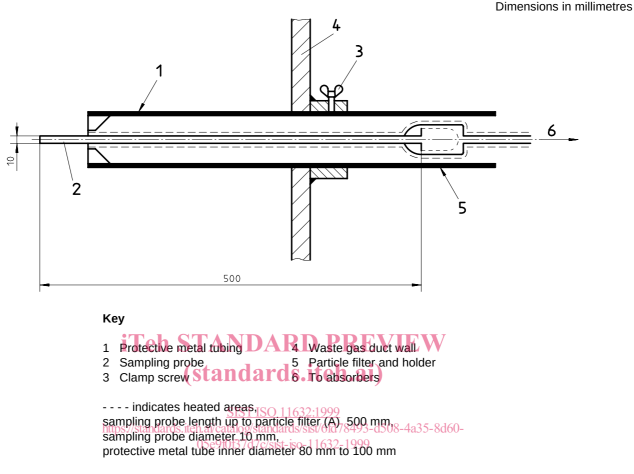


Figure 1 — Positioning of sampling probe, filter holder and protective tube

5.1.3 Filter holder

The filter holder shall be of borosilicate glass or fused silica glass with tube ends of spherical ground joints. The filter holder, which is encircled by the heating jacket, shall be connected to the sampling probe and housed within the protective metal tubing as shown in figure 1. The temperature after the filter holder is verified using a thermocouple.

NOTE 1 In special cases where the waste gas temperature is > 473 K, the heating jacket around the sampling probe, filter holder and connector line may be omitted. The temperature in the sampling line before the first absorber, however, should not fall below the acid dew point temperature of the waste gas.

NOTE 2 An example of suitable filter holder dimensions is shown in figure 2.

5.1.4 Particle filter

Alternative particle filters and filter holders of different designs may be used, but the residence time of the sample gas should be minimized. The filter material (quartz fibre or quartz wool packed progressively) shall have an efficiency better than 99,9 % for particles of cut-off diameter $0,6 \,\mu$ m for the actual sampling flow.

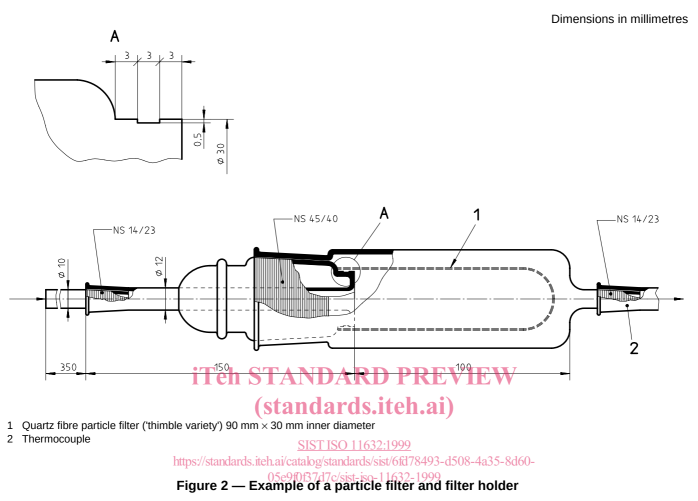
NOTE 1 An example of a suitable quartz-fibre particle filter of the "thimble" variety is shown in figure 2. The filter is held in position in the filter holder by stainless steel wire.

NOTE 2 In certain cases, a check on the possibility of a reaction between SO_2 in the sampled gas and particles retained on the filter may be warranted. This can be performed by comparing sulfate analyses on the particle fractions obtained from (i) the

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filter used when sampling according to this International Standard, and (ii) particles obtained from another source at the site, e.g. cyclone.



5.1.5 Absorbers

Two absorption bottles with spherical ground glass joints and equipped with an absorption bottle insert having a sintered filter. Alternative absorber sizes and configurations may be used, provided that the absorption efficiency criteria specified in 7.1 are met.

NOTE As an example, two 125 cm³ size absorption bottles of the Drechsel type with pore diameters in the sintered filter between 40 μ m and 90 μ m may be used (see figure 3).

5.1.6 Heating jacket

Heating jacket or bandage capable of producing a temperature of at least 473 K.

5.1.7 Temperature regulator

Temperature regulator suitable for use with the heating jacket or bandage.

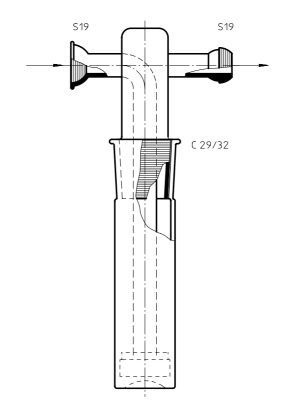


Figure 3 A Example of an absorption bottle

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5.1.8 Trap

Absorption bottle equipped with an absorption bottle insert not having a sintered filter. This bottle is connected after the second absorber and collects possible splashes of absorption solution.

NOTE The use of the trap is optional.

5.1.9 Drying tube

Glass tube or absorption bottle packed with drying agent to dry the sample gas and protect the gas metering device and pump.

NOTE As an example, silica gel (1 mm to 3 mm particle size) previously dried at 448 K for at least 2 h may be used.

5.1.10 Sampling pump

Leak-free diaphragm pump capable of drawing sample gas at a flowrate within the range 0,02 m³/h to about 0,2 m³/h during the sampling period against a pressure of -10 kPa to -30 kPa. A small surge tank between the pump and rotameter may be used to eliminate the pulsation effect of the diaphragm pump on the rotameter.

5.1.11 Rotameter

Flowmeter or rotameter capable of measuring the selected sample gas flow with limits of error < \pm 2 % of the upper limit of measurement.

5.1.12 Regulating valve

Needle valve capable of adjusting the sample gas flowrate within the range $0,02 \text{ m}^3/\text{h}$ to $0,2 \text{ m}^3/\text{h}$.