

## SLOVENSKI STANDARD SIST-TS ISO/TS 17379-2:2015

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Nadomešča: SIST ISO 9965:1998

# Kakovost vode - Določevanje selena - 2. del: Metoda z uporabo atomske absorpcijske spektrometrije s hidridno tehniko (HG/AAS)

Water quality - Determination of selenium - Part 2: Method using hydride generation atomic absorption spectrometry (HG-AAS)

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Qualité de l'eau - Dosage du sélénium - Partie 2: Méthode par spectrométrie d'absorption atomique à génération d'hydrures (HG-AAS) https://standards.iteh.ai/catalog/standards/sist/a6ceac47-be63-44c6-ac0c-5c12591cac94/sist-ts-iso-ts-17379-2-2015

Ta slovenski standard je istoveten z: ISO/TS 17379-2:2013

## <u>ICS:</u>

13.060.50

Preiskava vode na kemične snovi

Examination of water for chemical substances

SIST-TS ISO/TS 17379-2:2015

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#### **SIST-TS ISO/TS 17379-2:2015**

# TECHNICAL SPECIFICATION

# ISO/TS 17379-2

First edition 2013-07-15

# Water quality — Determination of selenium —

Part 2:

Method using hydride generation atomic absorption spectrometry (HG-

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(Standard de l'eau + Dosage du sélénium — Partie 2: Méthode par spectrométrie d'absorption atomique à génération d'hydrures (HG-AAS)

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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.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2, www.iso.org/directives.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received, www.iso.org/patents.

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The committee responsible for this document is ISO/TC 147, *Water quality*, Subcommittee SC 2, *Physical, chemical and biochemical methods*eh STANDARD PREVIEW

This first edition of ISO/TS 17379-2 cancels and replaces ISO 9965:1993, which has been technically revised.

ISO/TS 17379 consists of the following parts, under the general title *Water quality — Determination of* selenium: <u>SIST-TS ISO/TS 17379-2:2015</u>

- Part 1: Method using hydride generation atomic fluorescence spectrometry (HG-AFS)

— Part 2: Method using hydride generation atomic absorption spectrometry (HG-AAS)

## Introduction

This part of ISO/TS 17379 is intended for use by analysts experienced with the handling of trace elements at very low concentrations.

Inorganic selenium normally occurs in two oxidation states; Se(VI) and Se(IV). It is essential to convert all selenium species to the Se(IV) state prior to generating the hydrides. Selenium(VI) does not form a hydride.

In natural water sources, selenium compounds generally occur in very small quantities, typically less than 1  $\mu$ g/l. Higher concentrations may be found, e.g. in industrial waste water. Selenium occurs naturally in organic and inorganic compounds and may have oxidation states –II, 0, IV, and VI.

In order to fully decompose all of the selenium compounds, a digestion procedure is necessary. Digestion can only be omitted if it is certain that the selenium in the sample can form a covalent hydride without the necessity of a pre-oxidation digestion step.

The user should be aware that particular problems could require the specification of additional marginal conditions.

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## Water quality — Determination of selenium —

## Part 2: Method using hydride generation atomic absorption spectrometry (HG-AAS)

WARNING — Persons using this document should be familiar with normal laboratory practice. This document does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user to establish appropriate safety and health practices and to ensure compliance with any national regulatory conditions.

**IMPORTANT** — It is absolutely essential that tests conducted in accordance with this document be carried out by suitably trained and experienced staff.

#### 1 Scope

This part of ISO/TS 17379 specifies a method for the determination of selenium. The method is applicable to drinking water, surface water, ground water, and rain water. The dynamic range of this part of ISO/TS 17379 is approximately 0.5  $\mu$ g/l to 20  $\mu$ g/l. Samples containing selenium at higher concentrations than the application range can be analysed following appropriate dilution. The method is unlikely to detect organoselenium compounds.

The sensitivity of this method is dependent on the selected operating conditions. <u>SIST-TS ISO/TS 17379-2:2015</u>

It is important to use high purity reagents in all cases with minimum levels of selenium.

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#### 2 Normative references

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 3696, Water for analytical laboratory use — Specification and test methods

ISO 5667-1, Water quality — Sampling — Part 1: Guidance on the design of sampling programmes and sampling techniques

ISO 5667-3, Water quality — Sampling — Part 3: Preservation and handling of water samples

ISO 5667-5, Water quality — Sampling — Part 5: Guidance on sampling of drinking water from treatment works and piped distribution systems

ISO 5667-6, Water quality — Sampling — Part 6: Guidance on sampling of rivers and streams

ISO 5667-8, Water quality — Sampling — Part 8: Guidance on the sampling of wet deposition

ISO 5667-11, Water quality — Sampling — Part 11: Guidance on sampling of groundwaters

ISO 8466-1, Water quality — Calibration and evaluation of analytical methods and estimation of performance characteristics — Part 1: Statistical evaluation of the linear calibration function

ISO 8466-2, Water quality — Calibration and evaluation of analytical methods and estimation of performance characteristics — Part 2: Calibration strategy for non-linear second order calibration functions

ISO 15587-1, Water quality — Digestion for the determination of selected elements in water — Part 1: Aqua regia digestion

### **3** Principle

An aliquot of sample is treated with concentrated hydrochloric acid (5.2). Se(VI) is pre-reduced to Se(IV) by gently refluxing in 6 mol/l HCl for 1 h. Care is necessary to avoid any losses of volatile selenium components. A suitable apparatus is shown in Figure B.3. The sample solutions are then treated with sodium tetrahydroborate to generate the covalent gaseous hydride (SeH<sub>2</sub>). The hydride and excess hydrogen are swept out of the generation vessel in the batch mode and out of the gas/liquid separator in the continuous mode into a heated silica cell. After atomization the absorbance of selenium is determined at a wavelength,  $\lambda = 196,0$  nm. The procedure is automated by means of an auto sampler and control software.

Other measurement techniques may be applicable provided that the performance criteria can be adequately demonstrated or exceeded by the user laboratory (see <u>Annex C</u>).

#### **4** Interferences

The hydride generation technique is prone to interferences by transition and easily reducible metals. For the majority of natural water samples, this type of interference should not be significant. The user should carry out recovery tests on typical waters and also determine the maximum concentrations of potentially interfering elements, using appropriate methods. If such interferences are indicated, the level of interferences should be assessed by performing spike recoveries.

The reaction conditions set out in this part of ISO/TS 17379 have been chosen so that any interferences are reduced to a minimum. (standards.iteh.ai)

See References [1][2] for further information concerning these interferences and the technique. <u>SIST-TS ISO/TS 17379-2:2015</u>

 $These elements do not cause interferences provided the concentrations {\cite{specified} in {\cite{Table 1}} are not exceeded.} {\cite{5c12591cac94/sist-ts-iso-ts-17379-2-2015}} are not exceeded.$ 

# Table 1 — Maximum mass concentrations of hydride-forming or volatile elements causing no interferences

Element									
As	Sb	Sn	Те	Hg					
Mass concentration of interfering elements in test solution									
mg/l									
1	1	0,1	1	0,1					

If the concentrations in <u>Table 1</u> are exceeded, it may be necessary to use the standard addition method (9.3). Assuming that the selenium content is high enough, an appropriate dilution of the water sample is preferred.

Metals which are readily reduced by sodium tetrahydroborate may also cause interferences. In particular, these include chromium, iron, copper, nickel, and lead. If the concentrations of these elements specified in <u>Table 2</u> are exceeded, a significant decrease of absorption may occur.

# Table 2 — Maximum mass concentrations in test solution of interfering heavy metals(valid for flow systems)

Interfering element									
Cr	Fe	Cu	Ni	Pb					
Mass concentration in test solution									
mg/l									
500	500	500	250	100					

NOTE If batch systems are used, mass concentrations which are appreciably lower than those specified in Table 1 and Table 2 can cause interferences.

## 5 Reagents and standards

During the analysis, unless otherwise stated, use only reagents of recognized analytical grade.

Reagents may contain selenium as an impurity. All reagents shall have selenium concentrations below that which would result in a selenium blank value for the method being above the lowest level of interest.

Reagents shall be prepared to manufacturer's recommendations using the following series as an example.

- **5.1 Water**, complying with grade 1 as defined in ISO 3696, for all sample preparation and dilutions.
- **5.2** Hydrochloric acid,  $\rho(HCl) = 1,16$  g/ml.
- 5.3 Hydrochloric acid, c(HCl) = 1 mol/l.
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- **5.4 Sodium hydroxide**, NaOH.

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5.5 Sodium tetrahydroborate solution, ρ(NaBH4) = 13.8/1-be63-44c6-ac0c-

Dissolve an appropriate amount of sodium tetrahydroborate (13 g has proven suitable for the system illustrated in Annex B) in 500 ml water (5.1) and add (4,0  $\pm$  0,1) g of sodium hydroxide (5.4). Dilute to 1 000 ml with water (5.1). Prepare on day of use and do not keep in a closed container because of pressure build-up due to hydrogen evolution.

NOTE 1 The concentration of NaBH $_4$  is dependent on the hydride generator manifold and flow rate conditions. See recommendations of the manufacturer.

NOTE 2 Suitably stored sodium tetrahydroborate pellets have a shelf-life of 6 months.

**5.6** Nitric acid,  $\rho$ (HNO<sub>3</sub>) = 1,40 g/ml.

NOTE Nitric acid is available both as  $\rho(HNO_3) = 1,40 \text{ g/ml} [w(HNO_3) = 650 \text{ g/kg}]$  and  $\rho(HNO_3) = 1,42 \text{ g/ml} [w(HNO_3) = 690 \text{ g/kg}]$ .

Prepare a nitric acid cleaning mixture by diluting nitric acid [ $\rho$ (HNO<sub>3</sub>) = 1,40 g/ml] with an equal volume of water (5.1) by carefully adding the acid to the water.

#### 5.7 Reagent blank.

For each 1 000 ml, prepare a solution containing  $(300 \pm 3)$  ml of hydrochloric acid (5.2). Dilute to volume with water (5.1).

On the continuous flow system, the reagent blank solution is run as background. Since the blank solution may contain detectable trace levels of selenium, it is important that the same reagents be used for both sample and standard preparation, as well as for preparation of the reagent blank. The analyte signal is superimposed on this signal once the sample is introduced into the measurement cycle. The selenium concentration of the blank solution should be less than the lower level of interest.