

Chemicals used for treatment of water intended for human consumption — Hexafluorosilicic acid



BS EN 12175:2022 BRITISH STANDARD

National foreword

This British Standard is the UK implementation of EN 12175:2022 Supersedes BS EN 12175:2013, which is withdrawn.

The UK participation in its preparation was entrusted echnical Committee CII/59, Chemicals and filtering medial water treatment.

A list of organizations represented on this emmittee can be obtained on request to its committee manager.

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English Version

Chemicals used for treatment of water intended for human consumption Rexalluorosilicic acid

Produits chimiques utilisés pour le traitement de leau Produkte zur Aufbe

destinée à la consommation hur a ne-Acide hexafluorosili

Produkte zur Aufbereitung von Wasser für den menschlichen Gebrauch - Hexafluorkieselsäure

This European Standard was approved by CEN on 13 March 2022.

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European foreword

This document (EN 12175:2022) has been prepared by Technical Committee CEN/TO Water supply", the secretariat of which is held by AFNOR.

This European Standard shall be given the status of a national standard there by publication of an identical text or by endorsement, at the latest by October 2022, and Countricing national standards shall be withdrawn at the latest by October 2022.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN shall not be held responsible for identifying any or all such patent rights.

This document supersedes EN 4 2775 2013

In comparison with the previous edition, the following technical modifications have been made:

- a) modification of 7.3 on transportation regulations and labelling, adding the sentence "The user must be aware of the incompatibilities between transported products.";
- b) modification of 7.4 on marking. The requirements of marking are also applied to the accompanying documents.

Any feedback and questions on this document should be directed to the users' national standards body. A complete listing of these bodies can be found on the CEN website.

According to the CEN-CENELEC Internal Regulations, the national standards organisations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdom.

Introduction

In respect of potential adverse effects on the quality of water intended for human consumption caused by the product covered by this document:

- a) this document provides no information as to whether the product may be used with restriction in any of the Member States of the EU or EFTA;
- b) it should be noted that, while awaiting the adoption of verifiable Forepean criteria, existing national regulations concerning the use and/or the characteristics of this product remain in force.

NOTE Conformity with this document does not confer a mply acceptance or approval of the product in any of the Member States of the EU or EFTA. The use of the product covered by this document is subject to regulation or control by National Authorities (see Annex A).

Scope

This document is applicable to hexafluorosilicic acid used for treatment of water intended for human consumption. It describes the characteristics of hexafluorosilicic acid and specifies the recomments and the corresponding test methods for hexafluorosilicic acid. It gives information on as use in water treatment. It also determines the rules relating to safe handling and use of hexaftinosilicic acid (see

Annex B).

2 Normative references

The following documents are referred to in the tex (in such a way that some or all of their content constitutes requirements of this decrease in the such as way that some or all of their content constitutes requirements of this decrease in the such as way that some or all of their content constitutes requirements of this decrease in the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that some or all of their content constitutes are referred to in the textile and the such as way that the such as way the such as way that the such as way th constitutes requirements of this document. Fixed and references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

EN ISO 3696, Water for analytical lyberatory use - Specification and test methods (ISO 3696)

ISO 3165, Sampling of chemical products for industrial use — Safety in sampling

ISO 5440, Sodium hexafluorosilicate for industrial use — Determination of phosphate content — Molybdovanadate spectrophotometric method

ISO 5993, Sodium hydroxide for industrial use — Determination of mercury content — Flameless atomic absorption spectrometric method

ISO 6206, Chemical products for industrial use — Sampling — Vocabulary

ISO 6353-1, Reagents for chemical analysis — Part 1: General test methods

Terms and definitions

No terms and definitions are listed in this document.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at https://www.iso.org/obp
- IEC Electropedia: available at https://www.electropedia.org/

Description

4.1 Identification

4.1.1 Chemical name

Hexafluorosilicic acid.

4.1.2 Synonym or common name

Hydrofluosilicic acid.

Silicate-hexafluoro-dihydrogen.

Fluorosilicic acid.

HFSA.

4.1.3 Relative molecular mass

144.09.

4.1.4 Empirical formula

H₂SiF₆.

4.1.5 Chemical formula

H₂SiF₆.

4.1.6 CAS-Registry Number ¹

16961-83-4.

4.1.7 EINECS reference ²

241-034-8.

4.2 Commercial form

The product is an aqueous solution.

4.3 Physical properties

4.3.1 Appearance and odour

The product is a clear, colourless liquid with a pungent odour.

4.3.2 Density

The density of a mass fraction 35 % solution at 25 °C is 1,35 g/mL.

The density of a mass fraction 20 % solution at 25 °C is 1,18 g/mL.

4.3.3 Solubility (in water)

The product is miscible in any proportion.

4.3.4 Vapour pressure

The vapour pressure at 20 °C is approximately 3 kPa ³.

4.3.5 Boiling point at 100 kPa

The product boils and decomposes at 110 °C.

4.3.6 Crystallization point

A solution of mass fraction of 35 % crystallizes at -30 °C and of a mass fraction of 20 % crystallizes at -11.6 °C.

4.3.7 Specific heat

Not known.

http://www.china-gauges.com/

¹ Chemical Abstracts Service Registry Number.

² European Inventory of Existing Commercial Chemical Substances.

 $^{^{3}}$ 100 kPa = 1 bar.

4.3.8 Viscosity dynamic

A solution of a mass fraction of 20 % has a dynamic viscosity of 1,4 MPa.s at 25 °C.

4.3.9 Critical temperature

Not applicable.

4.3.10 Critical pressure

Not applicable.

4.3.11 Physical hardness

Not applicable.

4.4 Chemical properties

ttp://www.china-gauges.com/ Hexafluorosilicic acid is only stable in an aqueous solution. On evaporation, it decomposes to hydrogen fluoride (HF) and silicon tetrafluoride (SiF₄).

It produces hydrogen on contact with steel and metals (e.g. nickel and aluminium). It is a strong acid and reacts violently with alkalis. It forms hydrogen fluoride (HF) on contact with concentrated acids. It attacks glass.

Purity criteria

5.1 General

This document specifies the minimum purity requirements for hexafluorosilicic acid used for the treatment of water intended for human consumption. Limits are given for impurities commonly present in the product. Depending on the raw material and the manufacturing process, other impurities may be present and, if so, this shall be notified to the user and when necessary to relevant authorities.

Users of this product should check the national regulations in order to clarify whether it is of appropriate purity for treatment of water intended for human consumption, taking into account raw water quality, required dosage, contents of other impurities and additives used in the products not stated in this document.

Limits have been given for impurities and chemicals parameters where these are likely to be present in significant quantities from the current production process and raw materials. If the production process or raw materials lead to significant quantities of impurities, by-products or additives being present, this shall be notified to the user.

5.2 Composition of commercial product

The product shall contain between a mass fraction of 10 % and 47 % hexafluorosilicic acid, the remainder being water.

The concentration of hexafluorosilicic acid shall be within ± 5 % of the manufacturer's declared value.

5.3 Impurities and main by-products

Impurities must not exceed the requirements specified in Table 1.

Table 1 — Impurities

Impurity		Limit	1
		in mass fraction in % of commercial product	comi
Phosphate as P ₂ O ₅	max.	0,75) '
Free hydrogen fluoride as HF	max.	1,5 0209	

The product shall conform to the requirements specifical in Table 2.

Partition	ŕ'	Limit	
(10)		mg/kg H ₂ SiF ₆ (100 %)	
Antimony (Sb)	max.	80	
Arsenic (As)	max.	400	
Cadmium (Cd)	max.	40	
Chromium (Cr)	max.	400	
Lead (Pb)	max.	400	
Mercury (Hg)	max.	10	
Nickel (Ni)	max.	400	
Selenium (Se)	max.	80	

NOTE Other chemical parameters and indicator parameters are not relevant in hexafluorosilicic acid because the raw materials used in the manufacturing process are free of them. For parametric values of hexafluorosilicic acid on trace metal content in drinking water, see [1].

Test methods

6.1 General

SAFETY PRECAUTIONS Hexafluorosilicic acid shall be handled with extreme care, see B.1.

All equipment in contact with hexafluorosilicic acid shall be made of plastics (for example polyethylene or polytetrafluoroethylene (PTFE)); avoid contact with glass.

6.2 Sampling

6.2.1 General

The sampling is carried out at the premises of the manufacturer of the hexafluor flicic acid unless the customer has adequate facilities to carry out this operation safely at his owner.

customer has adequate facilities to carry out this operation safely at his own profiles.

6.2.2 Sampling from drums and bottles

6.2.2.1 General

6.2.2.1.1 Mix the contents of each container to be sampled by shaking the container, by rolling it or by rocking it from side to side, taking catelogs damage the container or spill any of the liquid.

ontainer is such (for example, a narrow-necked bottle) that it is **6.2.2.1.2** If the design **6.2.2.1.2** impracticable to use a sampling implement, take a sample by pouring after the contents have been thoroughly mixed. Otherwise, proceed as described in 6.2.2.1.3.

6.2.2.1.3 Examine the surface of the liquid. If there are signs of surface contamination, take samples from the surface as described in 6.2.2.2; otherwise, take samples as described in 6.2.2.3.

6.2.2.2 Surface sampling

Take a sample using a suitable ladle. Lower the ladle into the liquid until the rim is just below the surface, so that the surface layer runs into it. Withdraw the ladle before it fills completely and allow any liquid adhering to the ladle to drain off. If necessary, repeat this operation so that, when the other selected containers have been sampled, in a similar manner, the total volume of sample required for subsequent analysis is obtained.

6.2.2.3 Procedure of sampling from a container

The samples for testing the hexafluorosilicic acid shall be taken by means of a sampling tube, for example. A tube made of polytetrafluoroethylene (PTFE), approximately 1 500 mm long, with 14,5 mm inside diameter and 1,25 mm wall thickness tapering to an inside diameter of approximately 5 mm at one end, may be used for this; fit a rubber tube approximately 200 mm long which can be closed by means of a pinch clip, to the other end. When taking the samples, insert the sampling tube as far as possible into the acid to be tested with the clip released.

Do this slowly so that the levels of liquid in the sampling tube and in the acid container are the same.

Close the clip, withdraw the sampling tube from the acid, allow any liquid adhering at the outside of the tube to drain off, and by releasing the clip discharge the contents of the sampling tube into a polytetrafluoroethylene (PTFE) bottle of 1 000 ml nominal capacity provided with a ground PTFE stopper. Stopper the bottle immediately after filling each with the content of the sampling tube. After shaking thoroughly, fill from the collective sample three PTFE bottles, each with a volume of approximately 250 ml and provided with a ground PTFE stopper. Stopper, seal and label the bottles. One of these samples is to be tested by the consignee; the other two shall be kept in case subsequent complaint requires further testing to be carried out.

6.2.3 Sampling from tanks and tankers

From each access point, take samples as follows:

- b) from the bottom of the tank or tanker, using a sampling tube as described in \$2.3 or using specially designed bottom-sampling apparatus;

 c) from one or more positions, depending on the overall depth, between the bottom and the surface using a weighted sampling can.

 6.3 Analyses

 6.3.1 Hexafluorosilicic acid (main product)

6.3.1.1.1 Cold reaction

A saturated solution of potassium nitrate is added to an aliquot of the hexafluorosilicic acid which is cooled in ice and the liberated nitric acid is titrated with standard volumetric sodium hydroxide solution using bromothymol blue as the indicator.

$$H_2SiF_6 + 2KNO_3 --> 2HNO_3 + K_2SiF_6$$

(Sample A)
$$2HNO_3 + 2NaOH --> 2NaNO_3 + 2H_2O$$

6.3.1.1.2 Hot reaction

The solution is then brought to the boil and the liberated hydrofluoric acid is titrated with standard volumetric sodium hydroxide solution.

$$K_2SiF_6 + 2H_2O --> 2KF + SiO_2 + 4HF$$

(Sample B)
$$4HF + 4NaOH --> 4NaF + 4H_2O$$

If free acid other than hexafluorosilicic acid is present, then sample A will exceed half sample B. If salts of hexafluorosilicic acid are present then sample A will be less than half sample B.

6.3.1.2 Reagents

All reagents shall be of a recognized analytical grade and the water used shall conform to the grade 3 specified in EN ISO 3696.

- **6.3.1.2.1 Ice** prepared from grade 3 water
- 6.3.1.2.2 Potassium nitrate saturated solution
- 6.3.1.2.3 **Sodium hydroxide standard volumetric solution**, c(NaOH) = 0.5 mol/l
- **6.3.1.2.4** Bromothymol blue solution, 2 g/l
- 6.3.1.3 Apparatus
- **6.3.1.3.1 Bottle**, 50 ml
- **6.3.1.3.2 Safety pipette**, 25 ml

- **6.3.1.3.3 One-mark volumetric flask**, 500 ml
- Beaker, 500 ml 6.3.1.3.4

6.3.1.3.4 Beaker, 500 ml

6.3.1.3.5 Burette, 50 ml

6.3.1.3.6 Hot plate

6.3.1.4 Procedure

Using a safety pipette (6.3.1.3.2), transfer 25 ml of santo-amount a small weighed bottle (6.3.1.3.1) and reweigh. Transfer quantitatively to a 500 ml one-mark volumetric flask (6.3.1.3.3) and dilute to the mark with water. Place about 100 g of ica (18.1.2.1) in a 500 ml beaker (6.3.1.3.4) and add 25 ml of saturated potassium nitrate solution (13.1.2.2) followed by 25 ml of the diluted solution. Titrate immediately using the burette (6.3.1.3.5), with constant stirring, with the sodium hydroxide solution (6.3.1.2.3) using bromothy (13.1.2.4) as the indicator until the blue colour persists for at least 30 s; note the volume, V₁. 30 s; note the volume, V_1 .

NOTE The indicator will turn yellow after a while.

After the titration place the beaker on the hot plate (6.3.1.3.6) and bring the contents to the boil. Titrate the hot solution with sodium hydroxide standard volumetric solution (6.3.1.2.3) until the blue colour persists for 30 s and note the volume V_2 .

6.3.1.5 Expression of results

6.3.1.5.1 Hexafluorosilicic acid

The content of hexafluorosilicic acid, W_1 , expressed as mass fraction in %, is given from the following formula:

$$W_1 = \frac{V_2 \times 0.5 \times 0.036 \times 100 \times 50}{m_1 \times 25} \tag{1}$$

where

is the volume, in millilitres, of sodium hydroxide solution used in the second titration (6.3.1.2.3);

 m_1 is the mass in grams, of the test portion.

6.3.1.5.2 Free hydrofluoric acid

The content of fluoride other than hexafluorosilicic acid, W₂, expressed as mass fraction in % of HF, is given from the following formula:

$$W_2 = \frac{\left(V_1 - 0.5V_2\right) \times 0.5 \times 0.02 \times 100 \times 500}{m_1 \times 25} \tag{2}$$

where

is the volume, in millilitres, of sodium hydroxide solution used in the first titration (6.3.1.2.3); V_1

 m_1 is the mass, in grams, of the test portion.

6.3.2 Impurities

6.3.2.1 Free hydrofluoric acid

content of a conte	free hydrofluoric acid shoate phosphate shall be deter I parameters I Chemical parameters sha	all be determined ir	accordance with 6.3.1.!	3.2. 3.2.5.COV
content of	phosphate shall be deter	mined in accordanc	e with ISO 5440	$Q_{\mathcal{O}}$
3 Chemica	l parameters		·22-90	
3.1 Genera	1	_	Chillion	
or delicit		Who		
content of	chemical parameters sha	ill be the children us	ing the procedures spec	ified in Table 3.
	Table 3 — Procedures	for the determinat	tion of chemical param	eters
Element	Reference	Method	Wavelength nm	Flame
As	See 6.3.3.3	Hydride AAS	193,7	n.a.
Sb	See 6.3.3.3	Hydride AAS	217,6	n.a.
Cd	ISO 6353-1, GM 29 See 6.3.3.2	AAS	228,8	air-acetylene
Cr	ISO 6353-1, GM 29 See 6.3.3.2	AAS	357,8	air-acetylene
Pb	ISO 6353-1, GM 29 See 6.3.3.2	AAS	217,0 or 283,3	air-acetylene
Ni	ISO 6353-1, GM 29 See 6.3.3.2	AAS	232,0	Oxidising Air-acetylene
Se	See 6.3.3.3	Hydride AAS	196,0	n.a.
Hg	In accordance with ISO 5993	Flameless AAS	253,6	n.a.

n.a. = not applicable.

AAS = Atomic Absorption Spectrometry.

6.3.3.2 Determination of cadmium (Cd), chromium (Cr), lead (Pb) and nickel (Ni)

6.3.3.2.1 Principle

The elements cadmium (Cd), chromium (Cr), lead (Pb) and nickel (Ni) are determined using atomic absorption spectrometry with standard additions method in taking account of ISO 6353-1.

6.3.3.2.2 Reagents

6.3.3.2.2.1 General

All reagents shall be of a recognized analytical grade and the water used shall conform to the grade 3 specified in EN ISO 3696.

6.3.3.2.2.2 Standard solution (100 µg/l Cd, Cr, Pb or Ni)

The standard solution shall be freshly prepared on the day of use by individual dilution of a stock solution. This stock solution with an Cd, Cr, Pb or Ni content of at least 1 mg/l shall be made by dilution of standard solutions of Cd, Cr, Pb and Ni which are available from all major suppliers of laboratory chemicals. This stock solution shall be kept in containers of tetrafluoroethylene-lexalthoropropylene copolymer (FEP), polytetrafluoroethylene (PTFE) or polyethylene (PE).

The stock solution should not be kept for longer than four weeks.

6.3.3.2.3 Apparatus

Ordinary laboratory apparatus and the following:

Atomic absorption should in should in the measurement parameters specified in

Atomic absorption spentrameter with the measurement parameters specified in edure 6.3.3.2.3.1 Table 3.

6.3.3.2.4 Procedure

6.3.3.2.4.1 Test portion

Weigh 1 g to the nearest 0,01 mg of the laboratory sample into a 100 ml one-mark volumetric flask and make up to the mark at 20 °C with water.

6.3.3.2.4.2 Determination

The reference solutions shall be made by spiking the sample with the standard solutions, which contain stepwise increasing contents of the elements to be determined.

The amount of internal standard to be added can be estimated from a preliminary investigation, determining roughly the element content of the test sample from simple calibration.

The steps in which internal standards are added shall be at least as high as the estimated content of the test sample. With the spectrometer (6.3.3.2.3.1), carry out the measurement with the parameters specified in Table 3 in accordance with the manufacturer's instructions.

Repeat the procedure with all reagents and the same volume of standard solution to be added using water in place of the sample as a blank determination.

6.3.3.2.5 Expression of results

Prepare a calibration curve using the measured absorbencies of the spiked measurement solutions.

Read the concentration of each element in the test solution by extrapolation of the correlation line to absorbance A = 0 (see Figure 1). Similarly determine the element concentration of the blank solution (see Figure 2) and subtract from the result obtained for the test solution.

Alternatively, the evaluation may be carried out by linear regression. Additional dilution steps shall be compensated in the calculation.

This interim result (y) expressed in micrograms per litre which is converted to give the final concentration according to 6.3.3.2.6.

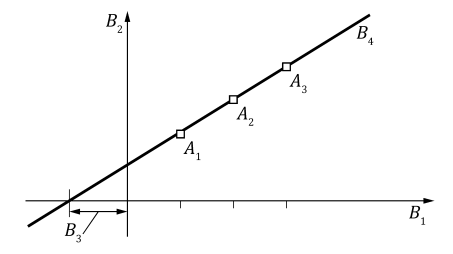
6.3.3.2.6 Calculation

From the interim result (y) determined (see 6.3.3.2.5) the content, W_3 , of each element in the laboratory

From the interim result (
$$v$$
) determined (see 6.3.3.2.5) the content, W_3 , of each element in the laboratory sample, expressed in milligrams per kilogram of 100 % hexafluorosilicic acid is given by the following formula:
$$W_3 = \frac{y \times V_3 \times 100 \times 1000}{m_2 \times W_1}$$
 where y is the interim result (6.3.3.2.5); V_3 is the volume, expressed in millilitres with test solution;

is the mass, expressed in grants of the test portion; m_2

is the content, expressed in mass fraction in % of hexafluorosilicic acid (see 6.3.1.5.1). W_1



Key

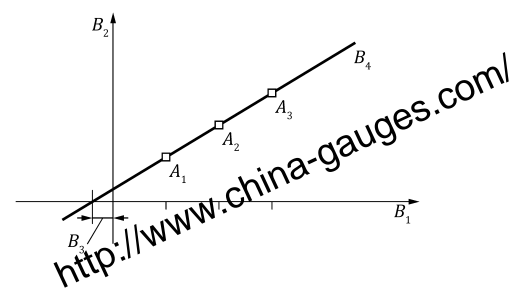
 B_1 absorbance A

spiking B_2

concentration of added standard in micrograms per litre B_3

concentration in the test solution in micrograms per litre

Figure 1 — Calculation of the element concentration in the test solution



Key

- B₁ absorbance A
- B_2 spiking
- B_3 concentration of added standard in micrograms per litre
- *B*₄ concentration in the blank solution in micrograms per litre

Figure 2 — Calculation of the element concentration in the blank solution

6.3.3.3 Determination of arsenic (As), antimony (Sb) and selenium (Se)

6.3.3.3.1 Principle

The elements arsenic, antimony, and selenium are determined by hydride-atomic absorption spectrometry. The elements are reduced by reducing agents (sodium borohydride (NaBH₄)) to form the hydrides. These volatile compounds flow through the heated measuring cuvette of an atomic absorption spectrometer where the content of the individual element is determined.

6.3.3.3.2 Reagents

6.3.3.3.2.1 Hydrochloric acid, high purity analytical grade, a mass fraction of 30 %, density $\rho = 1,15$ g/ml.

6.3.3.3.2.2 Preliminary reduction agent:

Dissolve 10 g sodium iodide and 100 g ascorbic acid, in 1 000 ml of water.

6.3.3.3.2.3 Reduction solution:

Dissolve in water NaBH₄ and NaOH in concentrations specified in the manufacturer's handbook for the spectrometer.

6.3.3.3.2.4 Standard solution (100 μ g/l As, Sb or Se):

The standard solution shall be freshly prepared on the day of use by individual dilution of a stock solution. This stock solution with an As, Sb or Se content of at least 1 mg/l shall be made by dilution of standard solutions of Se, As and Sb which are available from all major suppliers of laboratory chemicals. This stock solution shall be kept in containers of tetrafluoroethylene-hexafluoropropylene copolymer (FEP), polytetrafluorethylene (PTFE) or polyethylene (PE).

The stock solution should not be kept for longer than four weeks.

6.3.3.3.3 Apparatus

6.3.3.3.3.1 Three one-mark volumetric flasks, 1 000 ml

6.3.3.3.3.2

6.3.3.3.3.3

6.3.3.3.4

Nine one-mark volumetric flasks, 10 ml

Pipettes 5ml, 10 ml, and 100 ml

Micropipettes, volume adjustable to maximum 500 µl

Atomic absorption spectrometer with the pressurement parameters specified in 6.3.3.3.5 Table 3.

The width of the slit, the measuring time, flushing w ith argon before and after the measurement and the reaction time shall be adjusted in a condance with the manufacturer's instructions. The background measurement of As and Sb, but not for the measurement of Se. compensation shall be activated

6.3.3.3.4 Procedure

For As (procedure for Sb and Se in parentheses if different from As procedure):

Weigh a test portion of 0.5 g to the nearest 0.1 mg and transfer it into a 1 000 ml one-mark volumetric flask (6.3.3.3.3.1) and make up to the mark at 20 °C with water. Pipette 10 ml (Sb, Se:100 ml) of this solution into a 1 000 ml one-mark volumetric flask and add 5 ml of HCI (6.3.3.3.2.1) and 5 ml of preliminary reduction agent (6.3.3.3.2.2). Do not add preliminary reduction agent to the flasks for Sb and Se determination. Allow 3 h for reaction to occur and fill to the mark with water. Pipette 5 ml of this solution into three 10 ml one-mark volumetric flasks (6.3.3.3.3.2) labelled A,B,C. Add 0,8 ml of HCl (6.3.3.3.2.1) to each flask. For the purpose of internal calibration add those quantities of standard solutions (6.3.3.3.2.4) as given in Table 4 to the flasks B and C.

With the spectrometer (6.3.3.3.5), carry out the measurement with addition of the reduction agent (6.3.3.3.2.3) and the parameters of measurement in accordance with the manufacturer's instructions for the spectrometer.

Repeat the procedure with all reagents and the same volume of standard solution to be added using water in place of the sample as a blank determination.

Volume of standard solution to be added As Sb Se Flask B 200 ul 100 ul 200 ul Flask C 500 µl 200 μl 500 μl

Table 4 — Standard solution

6.3.3.3.5 Expression of results

See 6.3.3.2.5.

6.3.3.3.6 Calculation

See 6.3.3.2.6.

Labelling - Transportation - Storage 4

7.1 Means of delivery

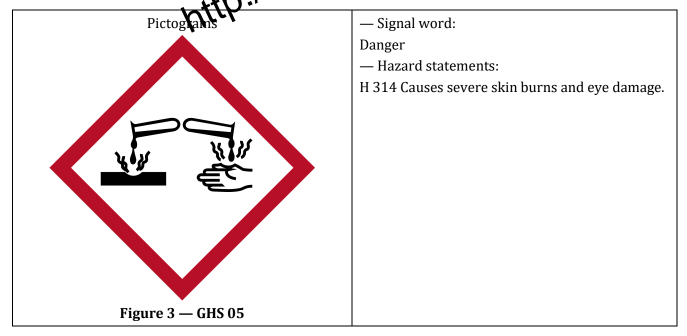
Hexafluorosilicic acid shall be delivered in suitable containers (not glass), e.g. polytetrafundo (PTFE) coated drums.

In order that the purity of the product is not affected, the means of deliver previously for any different product or it shall have been specially cleaned and prepared before use.

7.2 Labelling ⁵

The following labelling requirements shall apply to hexafluorosilicic acid at the date of publication of this document.

For concentrations of hexafluorosilicic acid above mass fraction 10 %:



The regulation [2], and its amendments for the purposes of its adaptation to technical and scientific progress, contains a list of substances classified by the EU. Substances not listed in this regulation can be classified on the basis of their intrinsic properties according to the criteria in the regulation by the person responsible for the marketing of the substance.

The supplier provides a safety data sheet in the national language in accordance with Regulation (EC) No 1907/2006, Annex II.

⁴ See Annex B.

⁵ See existing EU legislation, which foresees labelling requirements [2].

7.3 Transportation regulations and labelling

Hexafluorosilicic acid is listed as UN Number ⁶ 1778.

The user shall be aware of the incompatibilities between transported products.

7.4 Marking

The marking and the accompanying documents hall include the following:

— name "hexafluorosilicic acid" triple pame;

— net mass;

— name and "

- name and the address of supplier and/or manufacturer;
- statement "this product conforms to EN 12175".

7.5 Storage

7.5.1 Long term stability

The product is stable when stored in tightly closed containers in a cool well-ventilated place.

Hexafluorosilicic acid solutions of concentrations greater than mass fraction 22 % are less stable than those of lower concentration.

7.5.2 Storage incompatibilities

The product shall be kept away from steel and metals (e.g. nickel, aluminium), glass, acids and alkalis.

⁶ United Nations Number.

⁷ Regulations concerning International carriage of Dangerous Goods by rail.

⁸ European Agreement concerning the International carriage of Dangerous Goods by road.

⁹ International Maritime Transport of Dangerous Goods.

¹⁰ International Air Transport Association.

Annex A (informative)

Hexafluorosilicic acid is manufactured free Compounds and minerals containing bear silica (e.g. fluorite, apatite) and an acid (Mally sulfuric acid).

A.1.2 Manufacturing process

The compounds and minerals containing of the compounds and min pounds and minerals containing both fluoride and

A.2.1 Function

Hexafluorosilicic acid is used for the fluoridation of drinking water to increase the resistance of consumers to dental decay.

A.2.2 Form in which it is used

Hexafluorosilicic acid is used as an aqueous solution either as supplied or diluted with potable water.

A.2.3 Treatment dose

A typical dose of hexafluorosilicic acid of mass fraction 20 % is 6,3 mg/l to achieve a final concentration of 1 mg/l as F⁻ in the drinking water. It is important to avoid overdosing. In the EU Directive 98/83/EC, the parameter value is 1,5 mg/l of fluoride.

A.2.4 Means of application

It is usually applied using a metering pump.

A.2.5 Secondary effects

None.

A.2.6 Removal of excess product

It is practically impossible to remove excess product.

Annex B (normative)

and use

The shall provide current safety instructions.

H₂SiF₆ attacks glass; goggles are made of plastiful rubber.

Pergency procedures the staid

ly remove all and the state of the

B.1 Rules for safe handling and use

The supplier shall provide current safety instructions.

NOTE

B.2 Emergency procedures

B.2.1 First aid

Immediately remove all contaminated clothing.

In case of contact with the skin, wash immediately with plenty of water. Apply calcium gluconate gel to the affected area, rub in until locally free of pain and then continue for a further 15 min. Apply a dressing soaked with mass fraction of 20 % calcium gluconate solution.

If burns cover large areas, the patient shall be completely bathed in a calcium gluconate solution with a mass fraction of at least 1 %.

In case of contact with eyes, treat by irrigation with water with the eyelids held open. Consult a doctor (or eye specialist) immediately.

In case of inhalation, take six effervescent calcium pills (400 mg calcium per pill) dissolved in water.

In case of ingestion, immediately swallow a large quantity of calcium gluconate solution.

In all cases, obtain medical treatment as soon as possible.

B.2.2 Spillage

Confine any spillage with dry sand and cover with a suitable absorption agent. Trap escaping gases/vapours by spraying with water.

Do not empty into drains.

B.2.3 Fire

There is a risk of the container bursting if exposed to heat from a nearby fire. In case of firefighting, respirators are necessary.

There are no restrictions on extinguishing media in fire situations.

Bibliography

[1]

98/83/EC: Council Directive of 3 November 1998 on the quality of water intended of human consumption

Regulation (EC) No 1272/2008 of the European Parliament and Othe Council of 16 December 2008 on classification, labelling and packaging of substances and mixtures, amending and repealing Directives 67/548/EEC and 1999/45/EC undamending Regulation (EC) No 1907/2006 (REACH) [2]

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