

SLOVENSKI STANDARD SIST ISO 7393-1:1996

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Water quality -- Determination of free chlorine and total chlorine -- Part 1: Titrimetric method using N,N-diethyl-1,4-phenylenediamine

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Qualité de l'eau -- Dosage du chlore libre et du chlore total -- Partie 1: Méthode titrimétrique à la N,N-diéthylphénylène-1,4 diamine

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Examination of water for chemical substances

SIST ISO 7393-1:1996

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<u>SIST ISO 7393-1:1996</u> https://standards.iteh.ai/catalog/standards/sist/cd740c46-6bd6-4f2e-977adaf4ecec99cf/sist-iso-7393-1-1996 International Standard



7393/1

INTERNATIONAL ORGANIZATION FOR STANDARDIZATION MEXACHAPOCHAR OPPAHUSALUR TO CTAHCAPTUSALUMOORGANISATION INTERNATIONALE DE NORMALISATION

Water quality — Determination of free chlorine and total chlorine — Part 1: Titrimetric method using *N,N*-diethyl-1,4-phenylenediamine

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Descriptors : water, quality, chemical analysis, determination of content, chlorine, volumetric analysis.

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.

Draft International Standards adopted by the technical committees are circulated to the member bodies for approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at the ISO Council. They are approved in approved in a least 75 % approval by the member bodies voting.

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International Standard ISO 7393/1 was prepared by Technical Committee ISO/TC 147, Water quality.

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 \bigcirc International Organization for Standardization, 1985 •

Water quality — Determination of free chlorine and total chlorine — Part 1: Titrimetric method using *N*,*N*-diethyl-1,4-phenylenediamine

0 Introduction

ISO 7393 consists of the following parts:

Part 1: Titrimetric method using *N*, *N*-diethyl-1, 4-phenylenediamine.

Part 2: Colorimetric method using *N*,*N*-diethyl-1,4-phenylenediamine, for routine control purposes.

Part 3: lodometric titration method for the determination of total chlorine.¹⁾

1 Scope and field of application

This part of ISO 7393 specifies a titrimetric method for the 393determination of free chlorine and stotal chlorine in water standards/s

daf4eccc99cf/sist-iso-Sea water and waters containing bromides and iodides comprise a group for which special procedures are required.^[2]

The method is applicable to concentrations, in terms of chlorine (Cl₂), from 0,000 4 to 0,07 mmol/I (0,03 to 5 mg/l) total chlorine and at higher concentrations by dilution of samples. For concentrations above 0,07 mmol/I, ISO 7393/3 can also be used.

In annex A a procedure is presented for the differentiation of combined chlorine of the monochloramine type, combined chlorine of the dichloramine type and combined chlorine in the form of nitrogen trichloride.

Several compounds influence the determination specified in this part of ISO 7393. Interferences are noted in clauses 7 and 9.

2 **Definitions** (see table 1)

For the purpose of this part of ISO 7393, the following definitions apply.

2.1 free chlorine: Chlorine present in the form of hypochlorous acid, hypochlorite ion or dissolved elemental chlorine.

2.2 combined chlorine: The fraction of total chlorine present in the form of chloramines and organic chloramines.

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2.4 chloramines: Derivatives of ammonia by substitution of one, two or three hydrogen atoms with chlorine atoms (monochloramine NH₂Cl, dichloramine NHCl₂, nitrogen trichloride NCl₃) and all chlorinated derivatives of organic nitrogen compounds as determined by the method specified in this part of ISO 7393.

3 Principle

3.1 Determination of free chlorine

Direct reaction with the N, N-diethyl-1,4-phenylenediamine (DPD) and formation of a red compound at pH 6,2 to 6,5. Titration by means of a standard solution of ammonium iron(II) sulfate to the disappearance of the red colour.

3.2 Determination of total chlorine

Reaction with DPD in the presence of an excess of potassium iodide then titration as in 3.1.

Table 1 - Terms and synonyms in relation to actual compounds in the solution

Term	Sy	/nonym	Compounds		
Free chlorine		Active free chlorine	Elemental chlorine, hypochlorous acid		
	Free chlorine	Potential free chlorine	Hypochlorite		
Total chlorine	Total residual chl	orine	Elemental chlorine, hypochlorous acid, hypochlorite, chloramines		

1) At present at the stage of draft.

Reagents 4

During the analysis, use only reagents of recognized analytical grade, and only water as specified in 4.1.

4.1 Water, free from oxidizing and reducing substances.

Demineralized or distilled water of which the quality is checked as follows.

Into two 250 ml chlorine-demand-free conical flasks (clause 5) place, in order,

a) in the first: 100 ml of the water to be checked and about 1 g of potassium iodide (4.4); mix and after 1 min add 5 ml buffer solution (4.2) and 5,0 ml of DPD reagent (4.3);

b) in the second: 100 ml of the water to be checked and two drops of sodium hypochlorite solution (4.8); then, after 2 min, 5,0 ml of buffer solution (4.2) and 5 ml of DPD reagent (4.3).

No coloration should appear in the first flask whereas it is essential that a light pink coloration appears in the second flask.

In the case of demineralized or distilled water not having the desired quality it must be chlorinated. After a period of contact followed by dechlorination the quality finally has to be an rechecked.

A procedure for the preparation of water free from oxidizing ISO

4.2 Buffer solution, pH 6,5.

Dissolve in water (4.1) in this order: 24 g anhydrous disodium hydrogen phosphate (Na₂HPO₄) or 60,5 g of dodecahydrate form (Na₂HPO₄·12H₂O) and 46 g of potassium dihydrogen phosphate (KH2PO4). Add 100 ml of 8 g/l disodium dihydrogenethylenedinitrilotetraacetate dihydrate (disodium EDTA dihydrate, C10H14N2O8Na2·2H2O) solution (or 0,8 g of the solid form).

If necessary, add 0,020 g of mercury(II) chloride (HgCl₂), to prevent mould growth and interference in the free available chlorine test caused by any trace amounts of iodide in the reagents.

Dilute to 1 000 ml and mix.

NOTE - Solutions containing mercury should be disposed of properly (for example a method is specified in ISO 5790, Inorganic chemical products for industrial use - General method for determination of chloride content - Mercurimetric method).

4.3 *N*,*N*-diethyl-1,4-phenylenediamine sulfate (DPD) [NH₂-C₆H₄-N(C₂H₅)₂·H₂SO₄], solution, 1,1 g/I.

Mix 250 ml water (4.1), 2 ml sulfuric acid ($\rho = 1,84$ g/ml) and 25 ml of 8 g/l disodium EDTA dihydrate solution (or 0,2 g of the solid form). Dissolve in this mixture 1,1 g of anhydrous DPD or 1,5 g of the pentahydrate form, dilute to 1 000 ml and mix.

Store the reagent in a dark bottle protected from heat.

Renew the solution after 1 month or when it becomes discoloured.

4.4 Potassium iodide, crystals.

NOTE - Reagents 4.2, 4.3 and 4.4 may be conveniently replaced by combined reagents commercially available in the form of stable powder or tablets.

4.5 Ammonium iron(II) sulfate, stock solution, $c[(NH_4)_2Fe(SO_4)_2 \cdot 6H_2O] = 0,056 \text{ mol/l}.$

4.5.1 Preparation of the solution

Dissolve 22 g of ammonium iron(II) sulfate hexahydrate (Mohr's salt) in about 250 ml water (4.1) containing 5 ml sulfuric acid ($\varrho = 1,84 \text{ g/ml}$) in a 1 000 ml one-mark volumetric flask. Make up to the mark with water and mix.

Store in a dark bottle.

Standardize this solution by means of the procedure given in 4.5.2, when required for use, or daily if large numbers of determinations have to be done.

4.5.2 Standardization of the solution

Place in a 250 ml conical flask, 50,0 ml of the stock solution (4.5.1), about 50 ml water (4.1), 5 ml orthophosphoric acid $\sqrt{p} = 1730 \,\mathrm{g/ml}$ and 4 drops of barium diphenylamine and reducing substances is given in annex. Blards. itch. ai/catalog/standards.styled.indicator. (4.9), [4.9], [Titrate, with potassium dichromate solution (4.10). The end-point is reached when one drop prodaf4ecec99cf/sis duces an intense purple coloration which remains unchanged after further addition of the potassium dichromate solution. The concentration, c_1 , expressed in millimoles of Cl₂ per litre, of this solution is given by the equation

$$c_1 = V_2 \frac{c_2}{V_1}$$

where

 c_2 is the concentration of the potassium dichromate standard reference solution (4.10), $c(1/6 \text{ K}_2 \text{Cr}_2 \text{O}_7) = 100 \text{ mmol/l};$

 V_1 is the volume, in millilitres, of ammonium iron(II) sulfate stock solution (4.5), 50,0 ml;

 V_2 is the volume, in millilitres, of the potassium dichromate standard reference solution (4.10) used in the titration.

NOTE – When V_2 becomes less than 22 ml, prepare a fresh stock solution (see 4.5.1).

4.6 Ammonium iron(II) sulfate, standard volumetric solution, $c[(NH_4)_2Fe(SO_4)_2 \cdot 6H_2O] = 2.8 \text{ mmol/l}.$

Place 50,0 ml of the freshly standardized stock solution (4.5.1) in a 1 000 ml one-mark volumetric flask. Make up to the mark with water (4.1) and mix.

Transfer to a dark bottle.

Prepare this solution when required for use, or daily if large numbers of determinations have to be done.

The concentration, c_3 , expressed in millimoles of Cl_2 per litre, of this solution is given by the equation

$$c_3 = \frac{c_1}{20}$$

where c_1 is as defined in 4.5.2.

4.7 Sodium arsenite (NaAsO₂), solution, 2 g/I; or **thioacetamide** (CH₃CSNH₂), solution, 2,5 g/I.

4.8 Sodium hypochlorite, solution $[\varrho(Cl_2) \text{ about } 0,1 \text{ g/l}]$.

Prepare by dilution of concentrated commercial sodium hypochlorite solution.

4.9 Barium diphenylamine sulfonate, indicator solution, 3 g/l.

Dissolve 0,3 g barium diphenylamine sulfonate $[(C_6H_5-NH-C_6H_4-SO_3)_2Ba]$ in 100 ml water. (standards.ie)

4.10 Potassium dichromate, standard reference solution $c(1/6 \text{ K}_2\text{Cr}_2\text{O}_7) = 100 \text{ mmol/l}, (1/6 \text{ K}_2\text{Cr}_2\text{Cr}_2\text{Cr}_2\text{Cr}_2\text{Cr}_2\text{Cr$

Weigh, to the nearest milligram, 4,904 g^{daffeccopy} of anhydrous isopotassium dichromate. Dissolve in water in a 1 000 ml one-mark volumetric flask. Make up to the mark with water and mix.

5 Apparatus

Ordinary laboratory apparatus, and

Microburette, measuring up to 5 ml and graduated in divisions of 0,02 ml.

NOTE ON THE PREPARATION OF GLASSWARE

Chlorine-demand-free glassware is obtained by filling with sodium hypochlorite solution (4.8) then, after 1 h, rinsing copiously with water (4.1). During the course of the analysis one set of glassware should be kept for the determination of free chlorine and another for the determination of total chlorine in order to avoid contamination of the free chlorine set.

6 Procedure

6.1 Test sample

Start determination immediately after taking samples. At all times avoid bright light, agitation and heat.

6.2 Test portions

Take two test portions, each of 100,0 ml. If the concentration of total chlorine exceeds 70 μ mol/I (5 mg/I) it is necessary to take a smaller volume of test sample and to dilute with water (4.1) to 100,0 ml.

6.3 Determination of free chlorine

Place rapidly in a 250 ml conical flask in this order: 5,0 ml of buffer solution (4.2), 5,0 ml of DPD reagent (4.3) and the first test portion (6.2). Mix and titrate immediately to a colourless end-point with ammonium iron(II) sulfate solution (4.6). Note the volume, V_{3} , in millilitres, used in the titration.

In the case of an unknown water, possibly being very acid, or very alkaline or with a high concentration of salts, it is advisable to verify that the volume of buffer solution (4.2) added is sufficient to bring the water to pH 6,2 to 6,5. If not, use a greater volume of the buffer solution (4.2).

6.4 Determination of total chlorine

Place rapidly in a 250 ml conical flask in this order: 5,0 ml of buffer solution (4.2), 5,0 ml of DPD reagent (4.3), the second test portion (6.2) and about 1 g of potassium iodide (4.4). Mix and after 2 min, titrate to a colourless end-point with the ammonium fron(II) sulfate solution (4.6). If within 2 min a drift back of colour is observed continue titration to a colourless end-point. Note the volume, V_4 , in millilitres, used in the titration.

In the case of an unknown water, possibly being very acid, or

7 Correction of interference due to the presence of oxidized manganese

Determine the effect of oxidized manganese by carrying out a supplementary determination on a further test portion (6.2) previously treated with the arsenite or thioacetamide solution (4.7) in order to neutralize all oxidizing compounds other than oxidized manganese.

Place this test portion in a 250 ml conical flask, add 1 ml of sodium arsenite solution (4.7) or thioacetamide solution (4.7) and mix. Again add 5,0 ml of buffer solution (4.2) and 5,0 ml of DPD reagent (4.3). Titrate immediately to a colourless endpoint against ammonium iron(II) sulfate solution (4.6). Note the volume, V_{5} , in millilitres, corresponding to the oxidized manganese.

8 Expression of results

8.1 Method of calculation

8.1.1 Calculation of the free chlorine concentration

The concentration of free chlorine, $c(Cl_2)$, expressed in millimoles per litre, is given by the equation

$$c(Cl_2) = \frac{c_3 (V_3 - V_5)}{V_0}$$

where

 c_3 is the concentration, expressed in millimoles of Cl₂ per litre, of ammonium iron(II) sulfate solution;

 V_0 is the volume, in millilitres, of test sample in the test portion (6.2);

 V_3 is the volume, in millilitres, of the ammonium iron(II) sulfate solution (4.6) used in the titration (6.3);

 V_5 is the volume, in millilitres, of the ammonium iron(II) sulfate solution (4.6) used in clause 7 ($V_5 = 0$ ml in the absence of oxidized manganese).

8.1.2 Calculation of the total chlorine concentration

The concentration of total chlorine, $c(Cl_2)$, expressed in millimoles per litre, is given by the equation

$$c(Cl_2) = \frac{c_3 (V_4 - V_5)}{V_0}$$

where

 c_3 , V_0 and V_5 are as defined in 8.1.1;

V₄ is the volume, in millilitres, of the ammonium iron(1) ISO 709_past to quantify the reproducibility of the method by sulfate solution (4.6) used in the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories have produced databased and the titration (6.4) itch ai/catalog/stand distribution of samples to different laboratories h

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8.2 Conversion of amount of substance concentration to mass concentration

The chlorine concentration expressed in moles per litre may be expressed in grams per litre by multiplying by a conversion factor of 70,91.

8.3 Repeatability and reproducibility

To obtain an indication of repeatability and reproducibility, figures are taken from measurements obtained by methods which are the same in principle as that specified in this part of ISO 7393.

The USA-EPA Environmental Monitoring and Support Laboratory^[1] evaluated the titrimetric method with the following results.

For distilled water samples at concentrations of $c(Cl_2) = 4,79$; 9,17; and 48,6 µmol/I [$\rho(Cl_2) = 0,34$; 0,65; and 3,45 mg/I] total chlorine, the relative standard deviations were 5,6 %, 0,5 % and 0,5 %, respectively. Using drinking water containing $c(Cl_2) = 13,8$ µmol/I [$\rho(Cl_2) = 0,98$ mg/I], total chlorine, the relative standard deviation was 1,2 %. With more polluted waters almost the same precision was obtained as for drinking water with the exception of raw sewage where, with a total chlorine concentration of $c(Cl_2) = 11,1$ µmol/I [$\rho(Cl_2) = 0,79$ mg/I], the relative standard deviation was 3,3 %.

Results published by the British Department of the Environment^[2] showed for total chlorine concentrations of $c(Cl_2) = 14$ and 71 µmol/l [$\varrho(Cl_2) = 1,0$ and 5,0 mg/l], relative standard deviations of 1,4 % and 0,88 %, respectively.

(standar determinations in the same laboratory and thus provide a measure of the repeatability of the method. Attempts in

The past to quantify the reproducibility of the method by distribution of samples to different laboratories have produced unreliable results because of a general instability of solutions containing free and combined chlorine. More recently it has been found by the Quality Assurance Branch of EMSL-Cincinnati^[5] that a sealed vial of sodium hypochlorite in very pure water is remarkably stable when stored in the dark inside a mailing tube. Subsequent evaluation by various US federal and state laboratories has produced the analytical parameters listed in table 2 for methods in current use.

True	value		Number of laboratories (observations)				
c(Cl ₂)	[(Cl_2)]	Method code ¹⁾		Mean		Standard deviation	
µmol/l	(mg/l)			µmol/l	(mg/l)	µmol/l	(mg/l)
7	(0,5)	A	6	6,2	(0,44)	1,3	(0,09)
		В	7	6,8	(0,48)	1,8	(0,13)
11,3	(0,80)	A	10	10,9	(0,77)	1,1	(0,08)
		В	14	11,1	(0,79)	4,1	(0,29)
		С	6	11,6	(0,82)	1,3	(0,09)
15,5	(1,10)	A	10	15,5	(1,10)	2,0	(0,14)
		В	14	16,2	(1,15)	5,5	(0,39)
		С	6	16,5	(1,17)	1,1	(0,08)
18,2	(1,29)	A	6	18,6	(1,32)	1,3	(0,09)
		В	7	19,9	(1,41)	5,4	(0,38)

Table 2 — Analytical parameters from interlaboratory analysis for free residual chlorine

1) A: lodometric titration amperometric

B: DPD colorimetric

C: DPD titrimetric

9 Interferences

Two types of interference may be noted.

9.1 Interference by other chlorine compounds

A fraction of any chlorine dioxide that might be present is measured as free chlorine. This interference may be corrected by determining the chlorine dioxide in the water.^[2, 3, 4]

9.2 Interference by compounds other than chlorine compounds

Oxidation of DPD is not specifically caused by chlorine compounds. Depending on the concentration and the chemical oxidation potential, the reaction is effected by other oxidizing agents. The following substances may be mentioned in particular: bromine, iodine, bromamines, iodoamines, ozone, hydrogen peroxide, chromate, oxidized manganese, nitrite, iron(III) ions and copper ions. The interference is suppressed by the disodium EDTA in reagents 4.2 and 4.3 in the case of copper(II) ions (< 8 mg/I) and iron(III) ions (< 20 mg/I).

Interference by chromate may be eliminated by addition of barium chloride. $^{\left[6\right] }$

10 Test report

The test report shall include the following information:

a) a reference to this part of ISO 7393;

b) all information necessary for complete identification of the sample;

c) the results and the method of expression used;

d) details of any operations not included in this part of ISO 7393, or regarded as optional, together with any circumstance that may have affected the results.

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