

SLOVENSKI STANDARD SIST ISO 5275:2014

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Naftni proizvodi in ogljikovodikova topila - Ugotavljanje prisotnosti tiolov in drugih žveplovih spojin - "Doctor test"

Petroleum products and hydrocarbon solvents - Detection of thiols and other sulfur species - Doctor test

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Produits pétroliers et solvants hydrocarbonés - Détection des thiols (mercaptans) et autres espèces soufrées - Méthode au plombite de sodium ("Doctor test")

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Ta slovenski standard je istoveten z 7628e/ISO 5275;2003

ICS:

71.080.15 Aromatski ogljikovodiki Aromatic hydrocarbons Naftni proizvodi na splošno 75.080 Petroleum products in

general

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INTERNATIONAL STANDARD

ISO 5275

Second edition 2003-06-01

Petroleum products and hydrocarbon solvents — Detection of thiols and other sulfur species — Doctor test

Produits pétroliers et solvants hydrocarbonés — Détection des thiols (mercaptans) et autres espèces soufrées — Méthode au plombite de sodium («Doctor test»)

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Foreword

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International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. 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.

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.

ISO 5275 was prepared by Technical Committee ISO/TC 28, Petroleum products and lubricants.

This second edition cancels and replaces the first edition (ISO 5275 1979), which has been technically revised.

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Petroleum products and hydrocarbon solvents — Detection of thiols and other sulfur species — Doctor test

WARNING — The use of this International Standard may involve hazardous materials, operations and equipment. This International Standard does not purport to address all of the safety problems associated with its use. It is the responsibility of the user of this International Standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

1 Scope

This International Standard specifies a procedure (Doctor test) for the detection of thiols (mercaptans), hydrogen sulfide and elemental sulfur in hydrocarbon solvents and distillate petroleum feedstocks and products. Preliminary procedures also detect the presence of peroxides and phenolic substances, which, when present in more than trace quantities, make the application of this International Standard inappropriate. Carbon disulfide, at relatively high concentrations [above 0.4 % (m/m) sulfur], also interferes with the interpretation of the test by causing darkening of the aqueous layer.

NOTE For the purposes of this International Standard, the term "% (m/m)" is used to represent the mass fraction of a material.

The test is a go/no-go procedure with a threshold value of thiol concentration dependent on the material under test. It is frequently used as a substitute for the quantitative determination of thiol content.

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2 Normative references

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

ISO 1995:1981, Aromatic hydrocarbons — Sampling

ISO 3170:—1), Petroleum liquids — Manual sampling

ISO 3171:1988, Petroleum liquids — Automatic pipeline sampling

ISO 3696:1987, Water for analytical laboratory use — Specification and test methods

3 Principle

A test portion is shaken with sodium plumbite solution and the mixture observed. From its appearance, the presence or absence of thiols, hydrogen sulfide, elemental sulfur or peroxides may be deduced. The presence of thiols may be confirmed by the addition of sublimed sulfur, further shaking, and observation of the appearance of the final mixture.

¹⁾ To be published. (Revision of ISO 3170:1988)

4 Reagents and materials

4.1 General

Unless otherwise specified, the reagents specified in 4.2 to 4.11 shall be of analytical reagent grade, and water shall conform to the requirements of grade 3 of ISO 3696.

- **4.2** Lead acetate trihydrate [(CH₃COO)₂Pb·3H₂O], crystals.
- 4.3 Sodium hydroxide (NaOH), solid.

4.4 Sodium plumbite (Doctor) solution

Dissolve 25 g of lead acetate (4.2) in 200 ml of water, filter and add to a solution of 60 g of sodium hydroxide (4.3) in 100 ml of water. Heat the mixture in a boiling water bath for 30 min \pm 5 min, cool and dilute to 1 000 ml with water.

Store the solution in a tightly closed stoppered bottle and filter before use if it is not clear.

4.5 Cadmium chloride (CdCl₂)

See the warning in 4.7.

4.6 Hydrochloric acid (HCl), concentrated, approximately 36 % (*m/m*) HCl (11 mol/l).

4.7 Cadmium chloride solution

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Dissolve 100 g of cadmium chloride (4.5) in water, add 10 ml of hydrochloric acid (4.6) and dilute to 1 000 ml.

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WARNING — Cadmium chloride is toxic, and shall be disposed of as environmental toxic waste.

e45a8e7fe28e/sist-iso-5275-2014 For routine analysis, sodium hydrogen carbonate (NaHCO₃) solution at 50 g/l in water can be used, but since sodium sulfide is colourless, lead acetate paper is required to confirm the removal of hydrogen sulfide (see 7.3).

4.8 Sulfur, sublimed ("flowers of sulfur"), dry.

Store in a closed container.

4.9 Potassium iodide solution (KI)

Freshly prepare (daily) before use, a solution of 100 g/l of potassium iodide in water.

4.10 Acetic acid solution (CH₃COOH)

Freshly prepare (daily) before use, a solution of 100 g/l of glacial acetic acid in water.

4.11 Starch solution

Freshly prepare (daily) before use, a solution of 5 g/l of starch in water.

5 Apparatus

- **5.1 Mixing cylinders**, made of glass, stoppered, of capacity 50 ml.
- **5.2 Measuring cylinders**, made of glass, of capacity 5 ml and 10 ml.
- **5.3 Separating funnel**, made of glass, stoppered, of capacity 50 ml.

6 Samples and sampling

- **6.1** Unless otherwise specified, obtain laboratory samples in accordance with the procedures described in ISO 1995, ISO 3170 or ISO 3171, appropriate to the type of material being analysed and the sample source.
- **6.2** Mix laboratory samples by thoroughly shaking before withdrawing the test portion. Take care to avoid pressure build-up in the container if the sample has a vapour pressure exceeding 30 kPa at the laboratory temperature. Release pressure frequently and safely as required.

7 Procedure

7.1 Preliminary tests

7.1.1 Phenolic substances STANDARD PREVIEW

If the material under test is suspected of containing phenolic substances used as oxidation inhibitors, and which may interfere with the interpretation of test results (coloration of the aqueous layer), shake vigorously for 15 s in a mixing cylinder (5.1), a 10 ml test portion with 5 ml of 10 % (m/m) sodium hydroxide (4.3) in water solution. Observe the degree of coloration, and use this for comparison when assessing the results from 7.1.2. If the coloration is significant, the test shall be discontinued: 950808-f87e-47c0-8883-

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NOTE Any coloration beyond pale yellow is significant. If yellow is produced, the fourth assessment in Table 1 may need to be modified, dependent upon the intensity.

7.1.2 Sulfur species and peroxides

Place a 10 ml test portion of the sample and 5 ml of sodium plumbite solution (4.4) in a mixing cylinder (5.1) and shake vigorously for 15 s. Observe the appearance of the mixture and continue the procedure as indicated in Table 1.

Table 1 — Observations from preliminary test

Observation	Inference	Continue test as described in subclause
Black precipitate forms immediately	Hydrogen sulfide present	7.3
Brown precipitate forms slowly	Peroxides probably present	7.2
During the shaking period, the solution becomes opalescent and then darkens in colour	Thiols (mercaptans) and/or elemental sulfur present	7.4
No change occurs or a yellow colour is produced	Thiols may be present	7.4