

SLOVENSKI STANDARD **SIST EN 22719:1998**

01-maj-1998

Nafni proizvodi in maziva - Določanje plamenišča - Metoda z zaprto posodo po Pensky-Martensu (ISO 2719:1988)

Petroleum products and lubricants - Determination of flash point - Pensky-Martens closed cup method (ISO 2719:1988)

Mineralölerzeugnisse und Schmierstoffe - Bestimmung des Flammpunktes - Verfahren nach Pensky-Martens im geschlossenen Tiegel (ISO 2719:1988) W

Produits pétroliers et lubrifiants - Détermination du point d'éclair - Méthode Pensky-Martens en vase clos (ISO 2719:1988)_{IST EN 22719:1998}

https://standards.iteh.ai/catalog/standards/sist/5148479e-9d00-4aaf-8e11-

Ta slovenski standard je istoveten z: EN 22719-1998

ICS:

75.080 Naftni proizvodi na splošno Petroleum products in

general

75.100 Maziva Lubricants, industrial oils and

related products

SIST EN 22719:1998 en SIST EN 22719:1998

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN 22719:1998

https://standards.iteh.ai/catalog/standards/sist/5148479e-9d00-4aaf-8e11-b9a85ade91e3/sist-en-22719-1998

EUROPEAN STANDARD

EN 22719

NORME EUROPÉENNE

EUROPÄISCHE NORM

October 1993

UDC 665.7:543.873

Descriptors:

Petroleum products, lubricants, tests, determination, flash point, test equipment, Pensky-Martens apparatus

English version

Petroleum products and lubricants - Determination of flash point - Pensky-Martens closed cup method (ISO 2719:1988)

Produits Produits pétroliers et lubrifiants -Détermination du point d'éclair - Méthode Pensky-Martens en vase clos (ISO 2719:1988) pétroliers Mineralölerzeugnisse und Schmierstoffe – Bestimmung des Flammpunktes – Verfahren nach Pensky-Martens im geschlossenen Tiegel (ISO 2719:1988)

> N 22719:1970 S L O V E N 48479e-9d00-4aaf-8e11-REPUBLIKA MINISTRSTVO ZA ZNANOST IN TEHNOLOGIJO

Urad RS za standardizacijo in meroslovje

LJUBLJANA EN 22719

PREVZET PO METODI RAZGLASITVE

This European Standard was approved by CEN on 1993-10-20. CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without

Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the Central Secretariat or to any CEN member.

The European Standards exist in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the Central Secretariat has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Denmark, Finland, France, Germany, Greece, Iceland, Ireland, Italy, Luxembourg, Netherlands, Norway, Portugal, Spain, Sweden, Switzerland and United Kingdom.

CEN

European Committee for Standardization Comité Européen de Normalisation Europäisches Komitee für Normung

Central Secretariat: rue de Stassart, 36 B-1050 Brussels

SIST EN 22719:1998

Page 2 EN 22719:1993

Foreword

This European Standard is the endorsement of ISO 2719:1988. Endorsement of ISO 2719 was recommended by Technical Committee CEN/TC 19 "Methods of test and specifications for petroleum products" under whose competence this European Standard will henceforth fall.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by April 1994, and conflicting national standards shall be withdrawn at the latest by April 1994.

The standard was approved and in accordance with the CEN/CENELEC Internal Regulations, the following countries are bound to implement this European Standard: Austria, Belgium, Denmark, Finland, France, Germany, Greece, Iceland, Ireland, Italy, Luxembourg, Netherlands, Norway, Portugal, Spain, Sweden, Switzerland, United Kingdom.

Endorsement notice

The text of the International Standard ISO 2719:1988 was approved by CEN as a European Standard without any modification. STANDARD PREVIEW

(standards.iteh.ai)

<u>SIST EN 22719:1998</u> i/catalog/standards/sist/5148479e-9d0

https://standards.iteh.ai/catalog/standards/sist/5148479e-9d00-4aaf-8e11-b9a85ade91e3/sist-en-22719-1998

INTERNATIONAL STANDARD

ISO 2719

Second edition 1988-12-01



INTERNATIONAL ORGANIZATION FOR STANDARDIZATION ORGANISATION INTERNATIONALE DE NORMALISATION МЕЖДУНАРОДНАЯ ОРГАНИЗАЦИЯ ПО СТАНДАРТИЗАЦИИ

 $\begin{array}{lll} \textbf{Petroleum products and lubricants} & -\textbf{Determination} \\ \textbf{of flash point} & -\textbf{Pensky-Martens closed cup method} \end{array}$

Produits pétroliers et lubrifiants Détermination du point d'éclair Méthode W Pensky-Martens en vase clos (standards.iteh.ai)

SIST EN 22719:1998

https://standards.iteh.ai/catalog/standards/sist/5148479e-9d00-4aaf-8e11-b9a85ade91e3/sist-en-22719-1998

ISO 2719: 1988 (E)

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 approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at VIEW least 75 % approval by the member bodies voting.

(standards.iteh.ai)

International Standard ISO 2719 was prepared by Technical Committee ISO/TC 28, Petroleum products and lubricants.

https://standards.iteh.ai/catalog/standards/sist/5148479e-9d00-4aaf-8e11-

This second edition cancels and replaces the first edition (ISO 2719; 1973), of which it constitutes a technical revision.

Annexes A, B and C form an integral part of this International Standard.

ISO 2719 : 1988 (E)

Petroleum products and lubricants — Determination of flash point — Pensky-Martens closed cup method

1 Scope

This International Standard specifies methods, using the Pensky-Martens closed cup apparatus, for determining the flash point of combustible liquids, liquids with suspended solids, lubricating oils, liquids that tend to form a surface film under the test conditions, and other liquids.

Open cup flash and fire points of petroleum products may be determined by the use of ISO 2592: 1973, Petroleum products — Determination of flash and fire points — Cleveland open cup method. Flash points of paints and varnishes and drying oils may be determined by the use of ISO 1523: 1983, Paints, varnishes, petroleum and related products — Determination of flashpoint — Closed cup equilibrium method.

through an opening (kept closed at other times) into the cup at regular temperature intervals with simultaneous interruption of stirring. The flash point is the lowest temperature at which application of the test flame causes the vapour above the test portion to ignite.

4 Apparatus

- **4.1** Thermometer, partial immersion, conforming to the appropriate specification in annex A:
 - low range, for samples giving a flash point between
 10 °C and 110 °C;
- medium range, such as ASTM 88C or IP 101C therstandards.ite mometers;

high range, for samples giving a flash point between

NOTES

1 The method described in this International Standard may be — high range, for sa employed for the detection of contamination of lubricating oils by 719:1998 110 °C and 370 °C. minor amounts of volatile material, which also often occur in heat described transfer oils due to partial cracking. However, the lowest temperature

transfer oils due to partial cracking. However, the lowest temperature at which such a liquid is capable of producing an ignitable atmosphere may be lower than that found by this method. (See also clause 6, second paragraph, and 7.3, note.)

2 This International Standard should be used to measure and describe the properties of materials, products or systems in response to heat and flame under controlled laboratory conditions. Under actual fire conditions the response to heat and flame may be different.

n-22 4 2-1 Pensky-Martens closed cup apparatus, as described in annex B.

Automatic flash point testers are available and in use which may be advantageous in the saving of testing time, in permitting the use of smaller samples, and in other factors which may merit their use. If automatic testers are used, the user shall be sure that all the manufacturer's instructions for calibrating, adjusting and operating the instrument are followed.

In any cases of dispute, the flash point as determined manually shall be considered the referee test.

4.3 Adapter, for use with low-range thermometer (see annex C).

2 Definition

flash point: The lowest temperature, corrected to a barometric pressure of 101,3 kPa, at which application of a test flame causes the vapour of a test portion to ignite under specified conditions of test.

NOTES

- 1 The sample is deemed to have flashed when a large flame appears and instantaneously propagates itself over the surface of the sample.
- 2 Occasionally, particularly near the actual flash point, the application of the test flame will cause a blue halo or an enlarged flame; this is not a flash and should be ignored.

3 Principle

The test portion is heated at a slow, constant rate with continual stirring in a cup closed by a lid. A small flame is directed

5 Preparation of apparatus

Support the apparatus on a level, steady table. Unless tests are made in a draught-free room or compartment, it is good practice, but not obligatory, to surround the tester on three sides with a shield approximately 400 mm wide and 600 mm high.

6 Preparation of test sample

Test samples shall not be stored in polyethylene, polypropylene or other plastics bottles, because volatile material may diffuse through the walls of the bottle.

ISO 2719: 1988 (E)

If it is suspected that a test sample contains volatile contaminants, the treatment described in 6.1 and 6.2 shall be omitted.

6.1 Test samples of very viscous materials may be warmed until they are reasonably fluid before they are tested. However, no test sample shall be heated more than is absolutely necessary. A test sample shall never be heated above a temperature 17 °C below its expected flash point.

If the temperature difference between the melting point and flash point of a liquid is less than 20 °C, heat the sample to 3 °C \pm 1 °C above its melting point and immediately transfer a test portion into the cup and proceed as in clause 8, neglecting minimum temperatures specified in this clause.

6.2 Test samples containing dissolved or free water may be dehydrated with calcium chloride or by filtering through a qualitative filter paper or a loose plug of dry absorbent cotton. Warming the test sample is permitted, but it shall not be heated for prolonged periods or above a temperature 17 °C below its expected flash point.

On removing water, any water-soluble flammable material present is also likely to be removed.

the heating chamber (or cooling bath in the case of an expected flash point below ambient temperature — see 6.3). Ensure that the cup is properly seated. Insert the thermometer. Light the test flame and adjust it to a diameter of 4 mm \pm 0,5 mm. Heat at a rate such that the temperature as indicated by the thermometer increases 5 °C/min to 6 °C/min. Turn the stirrer at 90 r/min to 120 r/min, so that the test portion is made to flow from top to bottom.

- 7.2 For products whose flash point is expected to be equal to or below 110 °C, apply the flame at each degree up to 110 °C, beginning at a temperature between 18 °C and 28 °C below the expected flash point. Apply the test flame by operating the mechanism on the lid which controls the shutter and test flame burner so that the flame is lowered into the vapour space of the cup in 0,5 s, left in its lowered position for 1 s, and quickly retracted. Do not stir the test portion while applying the test flame. (See also 6.1, second paragraph.)
- **7.3** If the test portion is expected to have a flash point above 110 $^{\circ}$ C, apply the test flame in the manner just described at each temperature that is a multiple of 2 $^{\circ}$ C, beginning at a temperature between 17 $^{\circ}$ C and 28 $^{\circ}$ C below the expected flash point.

NOTE — The procedure described in 7.2 and 7.3 is not necessarily applicable when the test is run to determine the possible presence of volatile contaminants.

6.3 For expected flash points below ambient temperature, a prepare the apparatus as follows.

ileh SIA

- **6.3.1** Remove the test cup assembly (including lid, thermometer and stirrer) from the apparatus and ards itch av catalog/standards.
- **6.3.2** Place the test cup assembly in a suitable cooling bath (water or a 1+1 mixture of water and ethylene glycol may be used). The bath shall include a stirrer and cover. Provide support for the test cup assembly in the bath so that the lid and upper edge are horizontal and the cup is immersed in direct contact with the bath liquid in such a position that the level of the test portion in the cup is the same as that of the liquid in the water bath.

NOTE — If acetone and dry ice are used to cool the bath, do not use in direct contact with the bath liquid.

6.3.3 When the thermometer, in contact with the test portion, reaches a temperature at least 5 °C below the expected flash point, remove the bath. Apply the test flame every 1 °C as the higher ambient temperature causes the test portion to rise in temperature.

7 Procedure

7.1 Thoroughly clean and dry all parts of the cup and its accessories before starting the test. Ensure that all traces of solvent used to clean the equipment have been completely removed. Fill the cup with the test portion to the level indicated by the filling mark. Place the lid on the cup and set the latter in

7.49 Record as the flash point the temperature read on the thermometer at the time the second or subsequent application of the test flame, up to the twentieth application, causes a distinct flash in the interior of the cup. Do not confuse the true flash point with the bluish halo that sometimes surrounds the test flame at applications preceding the one that causes the actual flash.

If a flash is observed on the initial test flame application, or if no flash has been observed by the twentieth application, the procedure shall be started again with a new test portion, this time fixing a lower or a higher expected flash point, respectively. If a flash is observed at the initial test flame application but no flash is observed at a temperature lower than that of the initial test flame application, then the temperature of the initial test flame application is taken as the flash point.

8 Alternative procedure for highly viscous products

8.1 Bring the material to be tested and the tester to a temperature of 15 °C \pm 5 °C or 11 °C lower than the expected flash point, whichever is the lower. Turn the stirrer at 250 r/min \pm 10 r/min, stirring so that the test portion is made to flow from top to bottom. Raise the temperature throughout the duration of the test at a rate of not less than 1 °C/min and not more than 1,5 °C/min. With the exception of these requirements for rates of stirring and heating, proceed as prescribed in clause 7.

ISO 2719 : 1988 (E)

- 8.2 If the temperature difference between the melting point and flash point of a test material is less than 14 °C, heat the sample to 3 °C ± 1 °C above its melting point and immediately transfer a test portion into the cup and proceed as in clause 7, neglecting minimum temperatures specified in the clause.
- 8.3 Examples of highly viscous materials are heavy oils, polymeric solutions, adhesives, etc. If the results obtained with such materials, following the directions in clauses 6 and 7, are in doubt, repeat using the alternative procedure given in this clause. The higher flash temperature shall be considered the flash point of the material.

Calibration

- 9.1 Determine the flash point of p-xylene following the directions in clauses 6 and 7. When the tester is operating properly, a value of 27,2 °C ± 1,1 °C will be obtained.
- **9.2** If the flash point obtained with p-xylene is not within the limits stated in 9.1, check the condition and operation of the apparatus to ensure conformity with the details listed in annex B, especially with regard to the tightness of the lid, the action of the shutter and the position of the test flame. After adjustment, if necessary, repeat the test. p-Xylene having a flash point of 27,2 °C ± 1,1 °C is not a suitable reference material in the high-temperature range of the Pensky-Martens Closed Tester, which may be as high as 370 °C standards.if
- **9.3** The p-xylene shall conform to the following requirements: **SIST EN 22719:1**

ds.iteh.ai/catalog/standards/sist/5148479c-9d00-4aaf-8c11relative density at 15,56 °C / 15,56 °C.... e91e3/sist-en-22719-1998 0,860 min./ 0,866 max.

boiling range..... 2 °C from start to dry (the range shall include the boiling point of pure p-xylene, which is 138,35 °C)

purity..... 95 % min. (freezing point 11,23 °C min.)

Expression of results

10.1 Correction for barometric pressure

Observe and record the ambient barometric pressure at the time of the test. When the pressure differs from 101,3 kPa, correct the flash point using the following formula:

Corrected flash point = C + 0.25(101.3 - p)

where

- C is the observed flash point, in degrees Celsius;
- p is the ambient barometric pressure, in kilopascals.

The barometric pressure used in this calculation shall be the ambient pressure in the laboratory at the time of test. Many aneroid barometers, such as those used at weather stations and airports, are precorrected to give sea-level readings. These shall not be used.

NOTE - If the pressure is measured in millimetres of mercury, use the formula

corrected flash point = C + 0.033 (760 - p')

where p' is the pressure in millimetres of mercury.

Record the corrected flash point to the nearest 0,5 °C.

10.3 Precision

10.3.1 Basic procedure

The precision of this method, as obtained by statistical examination of interlaboratory test results, is as follows.

10.3.1.1 Repeatability

The difference between successive test results, obtained by the same operator with the same apparatus under constant operating conditions on identical test material, would, in the long run, in the normal and correct operation of the test method, exceed the values shown in table 1 only in one case in 20.

Table 1 — Repeatability

P Material R	Flash point range °C	Repeatability °C
Suspension of solids	35 to 43,5	2
All others	104 and under Above 104	2 6

10.3.1.2 Reproducibility

The difference between two single and independent results, obtained by different operators working in different laboratories on identical test material, would, in the long run, in the normal and correct operation of the test method, exceed the values shown in table 2 only in one case in 20.

Table 2 — Reproducibility

Material	Flash point range °C	Reproducibility °C
Suspension of solids	35 to 43,5	3,5
All others	104 and under Above 104	3,5 8,5

10.3.2 Alternative procedure

The following criteria shall be used for judging the acceptability (95 % confidence) of results obtained on viscous materials, which tend to form a surface film.

10.3.2.1 Repeatability

The average of two results obtained on the same sample on the same day by the same operator and that of two results on a different day should be considered suspect if they differ by more than 5 °C.