

## **SLOVENSKI STANDARD** SIST EN 15199-2:2006

01-december-2006

### Naftni proizvodi - Določanje porazdelitve območja vrelišč z metodo plinske kromatografije - 2. del: Težki destilati in goriva iz destilacijskih preostankov

Petroleum products - Determination of boiling range distribution by gas chromatography method - Part 2: Heavy distillates and residual fuels

Mineralölerzeugnisse - Gaschromatographische Bestimmung des Siedeverlaufes - Teil 2: Schweröle und Rückstandsöle TANDARD PREVIEW

Produits pétroliers - Détermination de la répartition dans l'intervalle de distillation par méthode de chromatographie en phase gazeuse - Partie 2: Fiouls lourds et fiouls résiduels

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### ICS:

75.080	Naftni proizvodi na splošno	Petroleum products in general
75.160.01	Goriva na splošno	Fuels in general

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en



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#### SIST EN 15199-2:2006

## EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

## EN 15199-2

October 2006

ICS 75.080

**English Version** 

### Petroleum products - Determination of boiling range distribution by gas chromatography method - Part 2: Heavy distillates and residual fuels

Produits pétroliers - Détermination de la répartition dans l'intervalle de distillation par méthode chromatographie en phase gazeuse - Partie 2: Distillats severes et residuals Mineralölerzeugnisse - Gaschromatographische Bestimmung des Siedeverlaufes - Teil 2: Schweröle und Rückstandsöle

This European Standard was approved by CEN on 28 August 2006.

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 any alteration. 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.

This European Standard exists 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.

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EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

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#### SIST EN 15199-2:2006

### EN 15199-2:2006 (E)

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### Foreword

This document (EN 15199-2:2006) has been prepared by Technical Committee CEN/TC 19 "Gaseous and liquid fuels, lubricants and related products of petroleum, synthetic and biological origin", the secretariat of which is held by NEN.

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 2007, and conflicting national standards shall be withdrawn at the latest by April 2007.

EN 15199 consists of the following parts, under the general title *Petroleum products* — *Determination of boiling range distribution by gas chromatography method*:

- Part 1: Middle distillates and lubricating base oils
- Part 2: Heavy distillates and residual fuels
- Part 3: Crude oil

This part of the standard describes the determination of boiling range distribution of materials with initial boiling points (IBP) above 100 °C and final boiling points (FBP) above 750 °C. For testing materials with initial boiling points (IBP) above 100 °C and final boiling point (FBP) below 750 °C, Part 1 of the standard may be used. For testing materials with initial boiling points (IBP) below 100 °C and final boiling points (FBP) below 750 °C, Part 1 of the standard may be used. For testing materials with initial boiling points (IBP) below 100 °C and final boiling points (FBP) above 750 °C, such as crude oils, Part 3 is applicable.

This part of the standard is a joint development between the EI [1], ASTM [2] and CEN.

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

#### 1 Scope

This European Standard specifies a method for the determination of the boiling range distribution of petroleum products by capillary gas chromatography using flame ionisation detection. The standard is applicable to materials having a vapour pressure low enough to permit sampling at ambient temperature, and which have a boiling range of at least 100 °C. The standard is applicable to materials with initial boiling points (IBP) above 100 °C and final boiling points (FBP) above 750 °C, for example, heavy distillate fuels and residuals. The method is not applicable to bituminous samples.

The test method is not applicable for the analysis of petroleum or petroleum products containing low molecular weight components (for example naphthas, reformates, gasolines, diesel). Components containing hetero atoms (for example alcohols, ethers, acids, or esters) or residue are not to be analyzed by this test method.

NOTE For the purposes of this European Standard, the terms "% (*m/m*)" and "% (*V/V*)" are used to represent respectively the mass fraction and the volume fraction.

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

#### 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.

EN ISO 3170, Petroleum liquids - Manual sampling (ISO 3170:2004)

https://standards.iteh.ai/catalog/standards/sist/07eba177-4022-44c4-9247-EN ISO 3171, Petroleum liquids — Automatic\_pipeline\_sampling; (ISO 3171:1988)

#### 3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

NOTE Explanation of some of the terms is given in Figure 1.

### 3.1

### initial boiling point

IBP

temperature corresponding to the retention time at which a net area count equal to 0,5 % of the total sample area under the chromatogram is obtained

#### 3.2

final boiling point

#### FBP

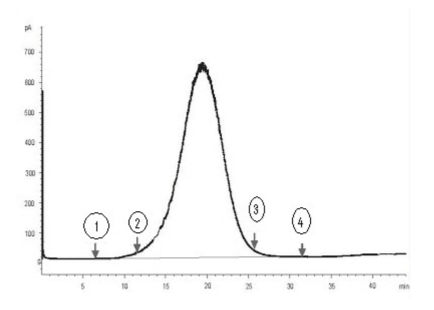
temperature corresponding to the retention time at which a net area count equal to 99,5 % of the total sample area under the chromatogram is obtained

#### 3.3

#### area slice

area resulting from the integration of the chromatographic detector signal within a specified retention time interval

NOTE In area slice mode peak detection parameters are bypassed and the detector signal integral is recorded as area slices of consecutive, fixed duration time interval.



Key:

1

2

**iTeh STANDARD** final boiling point (FBP) end of elution initial boiling point (IBP) (standards.iteh

#### Figure 1 - Typical chromatogram

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#### corrected area slice

start of elution

area slice corrected for baseline offset by subtraction of the exactly corresponding area slice in a previously recorded blank (non-sample) analysis

#### 3.5

3.4

#### cumulative corrected area

accumulated sum of corrected area slices from the beginning of the analysis through a given retention time, ignoring any non-sample area for example of solvent

#### 3.6

#### slice rate

time interval used to integrate the continuous (analogue) chromatographic detector response during an analysis

NOTE The slice rate is expressed in Hz (for example integrations per second or slices per second).

#### 3.7

#### slice time

analysis time associated with each area slice throughout the chromatographic analysis

NOTE The slice time is the time at the end of each contiguous area slice.

#### 3.8

#### total sample area

cumulative corrected area, from the initial area point to the final area point, where the chromatographic signal has returned to baseline after complete sample elution

#### 3.9

#### net area

cumulative area counts for the sample minus the cumulative area count for the blank

#### 3.10

#### recovery

ratio of the cumulative area count of the sample to that of the reference material (external standard) corrected for dilution and material weights combined with the percentage of light ends, if applicable

#### 4 Principle

A test portion is introduced into a gas chromatographic column, which separates hydrocarbons in the order of increasing boiling point. The column temperature is raised at a linear reproducible rate and the area under the chromatogram is recorded throughout the analysis. Boiling points are assigned to the time-axis from a calibration curve, which is obtained by running a mixture of known normal alkanes covering the test portion boiling range, under the same conditions. From these data, the boiling range distribution is obtained. The recovery at a specified temperature is determined by comparing the area under the chromatogram with that of a reference standard which has been completely eluted. The temperature at which the recovery was measured is recorded.

NOTE If the found recovery is less than 100 %, the final boiling point is reported as 720 °C or 750 °C at that recovery.

# 5 Reagents and materials the STANDARD PREVIEW

Unless otherwise stated, only chemicals of recognized analytical quality shall be used.

5.1 Liquid stationary phase, a methyl silicone stationary phase for the column.

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**5.2** Carrier gases, helium#itrogen or hydrogehgof at least 99,999 % (VV) putity 4Any oxygen present is removed by a chemical resin filter. 42cbfe864681/sist-en-15199-2-2006

Warning Follow the safety instructions from the filter supplier.

5.3 Hydrogen, grade suitable for flame ionisation detectors.

5.4 Compressed air, regulated for flame ionisation detectors.

**5.5** Alkanes, normal alkanes of at least 98 % (m/m) purity from C<sub>5</sub> to C<sub>10</sub>, C<sub>12</sub>, C<sub>14</sub>, C<sub>16</sub>, C<sub>18</sub>, C<sub>20</sub>, C<sub>24</sub> and C<sub>28</sub> to be used with Polywax 655 or 1000 (5.6).

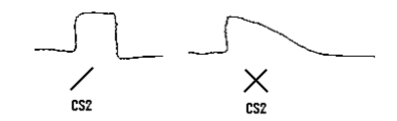
NOTE The calibration mixture from ISO 3924 [3] is also suitable.

#### 5.6 Polywax 655 or 1000

**5.7** Carbon disulfide, with a minimum purity of 99,7 % (*V*/*V*).

#### WARNING — Extremely flammable and toxic.

NOTE To confirm the suitability of the carbon disulfide as a solvent, it is recommended to check elution profiles (see Figure 2).



Key:

/ good X bad

#### Figure 2 — Solvent peak shape

#### 5.8 Calibration mixture

The mixture shall contain at least one normal alkane with a boiling point lower than the IBP of the sample, and at least one normal alkane with a boiling point close to the temperature at which the recovery is measured.

Dissolve 0,1 g of Polywax (5.6) in 7 ml carbon disulfide (5.7), warming gently if necessary. Prepare an equal volume mixture of alkanes (5.5) and add 10  $\mu$ l to the Polywax solution.

NOTE 1 Commercially available alkane standards are suitable for column performance checks.

NOTE 2 The calibration mix is used to determine the column resolution, skewness of the C<sub>20</sub> peak, and retention time versus boiling point calibration curve.

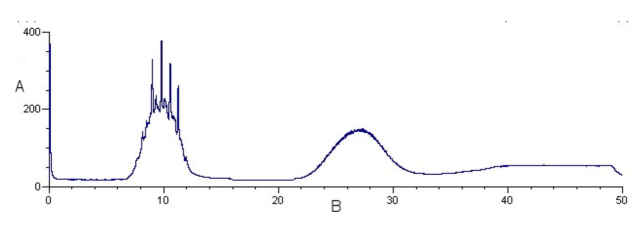
5.9 Reference materials (RM) SIST EN 15199-2:2006 https://standards.iteh.ai/catalog/standards/sist/07eba177-4022-44c4-9247-

- **5.9.1** A reference material has two functions.
- External Standard: to determine the recovery of samples by comparing the total sample area (3.8) of the reference material with the total sample area of the unknown sample.
- Boiling Point Distribution Standard: to check the proper functioning of the system by comparing the results with a known boiling point distribution on a routine basis. Typical example is given in (5.9.2).

**5.9.2** Reference Material 5010, a reference sample that has been analyzed by laboratories participating in the test method cooperative study. Consensus values for the boiling range distribution of this sample are given in Table 1.

**5.9.3** Cyclohexane.  $(C_6H_{12})$ —(99+ % pure) may be used in place of  $CS_2$  for the preparation of the calibration mixture.

**5.9.4 Binary gravimetric blend**, a binary distillate mixture with boiling points ranges that gives a baseline at the start, a baseline between the two peaks and an end time that is as close to the end of the chromatogram as possible (see Figure 3 and B.3). This mixture is used to check the relative response of the two distillates and to check the baselines at the start, middle and end of the chromatogram.



Key:

A response

B retention time (min)

Figure 3 — Typical chromatogram of binary gravimetric blend distillate

% recovered	Reference temperature tandards	Maximum allowable range ite95,5 % Cl	V
IBP	428 SIST EN 1510	9	
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10 42	cbfe8644931/sist-en	-15199-2-2306	
15	502	3	
20	510	3	
25	518	4	
30	524	4	
35	531	4	
40	537	4	
45	543	4	
50	548	5	
55	554	4	
60	560	4	
65	566	4	
70	572	4	
75	578	5	
80	585	4	
85	593	4	
90	602	4	
95	616	4	
FBP	655	18	

Table 1 — Reference Material 5010

#### 6 Apparatus

6.1 Gas chromatograph, with the following performance characteristics.

**6.1.1** Flame ionisation detector, connected to the column so as to avoid any cold spots. The detector shall be capable of operating at a temperature at least equivalent to the maximum column temperature employed in the method.

NOTE The capillary column should sit just below the flame tip and it is recommended that the orifice of the jet should be 0,6 mm minimum to prevent frequent blocking with silicones.

**6.1.2** Column temperature programmer, capable of linear programmed temperature operation over a range of 10 °C above ambient to 450 °C.

**6.1.3 Sample inlet system**, consisting of a programmable temperature vaporizer (PTV) or cold on-column (COC) injection port. The maximum temperature of the injection device shall be equal to, or higher than, the final oven temperature. The minimum temperature shall be low enough to prevent sample or solvent flashback, but high enough to allow sample focusing at the front of the column. Table 2 contains the typical operating conditions.

#### 6.2 Column

Use a metal column with 0,53 mm internal diameter and coated with methyl silicone (5.1). Commercially available columns with film thickness ( $d_f$ ) = 0,09 µm (for analysis up to C<sub>120</sub>) and ( $d_f$ ) = 0,17 µm (for analysis up to C<sub>100</sub>) have been found to be satisfactory.

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NOTE 1 It is recommended that the column resolution, *R*, is at least 2 and not more than 4 (see B.2).

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Use some form of column bleed compensation to obtain a stable baseline.

NOTE 2 This may be carried out by subtraction of a column bleed profile previously obtained using exactly the same conditions as used for the sample analysis by injecting the same volume, using solvent for the blank run and sample dilution from one batch taken at the same time, to avoid differences due to contamination.

Column length, m	5
Column internal diameter, mm	0,53
Column material	Ultimetal
Stationary phase	Methyl silicone
Film thickness, µm	0,09 or 0,17
Initial column temperature, °C	35
Final column temperature, °C	430
Program rate, °C/min	10
Injector initial temperature, °C	100
Injector final temperature, °C	430
Program rate, °C/min	15
Hold time, min	5
Detector temperature, °C	450
Carrier gas	Не
Carrier gas flow rate, ml/min	19
Sample size, µl	1,0
Sample concentration, % (m/m)	2
Injector	PTV or COC

#### Table 2 — Typical operating conditions for gas chromatograph