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# Indian Standard (Reaffirmed 2015) (Reaffirmed 2020) SPECIFICATION FOR OLEUM, TECHNICAL (Second Revision)

UDC 661.251



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INDIAN STANDARDS INSTITUTION MANAK BHAVAN, 9 BAHADUR SHAH ZAFAR MARG NEW DELHI 110002

December 1986

# Indian Standard SPECIFICATION FOR OLEUM, TECHNICAL (Second Revision)

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|---|---|---|---|--|
|   | TABLE 1 REQUIREMENT   | <b>S FOR OLEUM</b> ,<br>use 3.2)  | TECHNICA  | L  |
| SL<br>No.   | Characteristic  | Grade 1   | Grade 2   | Method of<br>Test ( Ref<br>to Cl. No.  |
|   |   |   |   | in<br>Appendix A )   |
| (1)   | (2)   | (3)   | (4)   | (5)  |
| i) Fre  | e sulphur trioxide ( as SO3 ), per-<br>ent by mass, Min   | 20  | 65  | A-2  |
| ii) Re  | sidue on ignition, percent by<br>nass, Max  | 0.50  | 0.50  | A-3  |
| iii) Iro  | n (as Fe), percent by mass,<br>Max  | 0.02  | 0.02  | A-4  |
| iv) Ch  | lorides ( as Cl ), percent by mass,<br>Max  | 0.002   | 0.002   | A-5  |
| v) To   | tal nitrogen (as HNO3), per-<br>ent by mass, Max  | 0.002   | 0.002   | A-6  |
| vi) Sul   | phur dioxide (as SO <sub>2</sub> ), per-<br>ent by mass, Max  | 0.2   | 0.5   | A-7  |
| vii) Ar   | senic ( as As ), percent by mass,<br>Max  | 0.000 2   | 0.000 2   | A-9  |
| • PAC<br>• PAC<br>• 1.1 Pac<br>• 4.1.1<br>he pu<br>comply<br>vailwa<br>hereaf<br>• 4.1.2<br>value<br>value<br>• 4.1.2<br>value<br>• 4.1.2 | KING AND MARKING<br>:king<br>The material shall be packed<br>rchaser and the supplier w<br>the provisions of Red Tariff<br>y Conference Association,<br>ter.<br>When oleum is supplied in s<br>arboys, the containers shall be<br>d by the purchaser, they shall<br>s. The stoppers shall furth | d in containers<br>ben transport<br>No. 18 of 1960<br>with any alter-<br>crew-stoppered<br>e fitted with lea<br>l be provided w<br>er be sealed b | s as agreed<br>ed by rail,<br>) issued by<br>ation or ad<br>stone-ware<br>k-proof stop<br>vith asbestos<br>y a putty m. | to between<br>it shall also<br>the Indian<br>dition made<br>bottles or<br>opers, and if<br>or rubber<br>ade of china<br>mixture of |
| vasher<br>lay or  | a mixture of source since   |   |   |  |

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4.1.3 The bottles and jars shall be packed in suitable pent top packing cases. These shall be placed in an upright position on one layer of sand or ashes free from cinders or dry earth, and the empty surrounding space shall be packed in suitable iron hampers or wooden crates stuffed with chemically inert materials like clay or fullers earth.

# 4.2 Marking

4.2.1 The containers and also the packages shall be suitably marked in red letters not less than 2.5 cm high, and drums and tank wagons shall be marked in red letters not less than 5 cm high, showing the name of the acid, manufacturer's name, the grade and mass of the material, recognized trade-mark, if any, and the month and year of manufacture.

4.2.2 The packages shall be labelled with the symbol shown in Fig. 1. [see IS: 1260 (Part 1)-1973<sup>•</sup>], the words 'CORROSIVE, HANDLE WITH CARE' being prominently displayed in the lower half of the symbol.

4.2.3 All the containers and packages should be suitably marked with an arrow showing the upright position of containers.

4.2.4 The packages may also be marked with the ISI Certification Mark.

Note — The use of the ISI Certification Mark is governed by the provisions of the Indian Standards Institution (Certification Marks) Act and the Rules and Regulations made thereunder. The ISI Mark on products covered by an Indian Standard conveys the assurance that they have been produced to comply with the requirements of that standard under a well-defined system of inspection, testing and quality control which is devised and supervised by ISI and operated by the producer. ISI marked products are also continuously checked by ISI for conformity to that standard as a further safeguard. Details of conditions under which a licence for the use of the ISI Certification Mark may be granted to manufacturers or processors, may be obtained from the Indian Standards Institution.

# 5. SAMPLING

5.1 Representative test samples of the material shall be drawn as prescribed in Appendix C.

<sup>\*</sup>Pictorial markings for handling and labelling of goods: Part 1 Dangerous goods (first revision).



SYMBOL SHOWING DANGER OF CORROSION FIG. 1

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# APPENDIX A

# (Clause 3.2)

# METHODS OF TEST FOR OLEUM, TECHNICAL

# **A-1. QUALITY OF REAGENTS**

A-1.1 Unless specified otherwise, pure chemicals and distilled water (see IS: 1070-1977\*) shall be used in the tests.

Note — 'Pure chemicals' shall mean chemicals that do not contain impurities which affect the results of analysis.

# **A-2. DETERMINATION OF FREE SULPHUR TRIOXIDE**

### A-2.1 Apparatus

A-2.1.1 Spherical Glass Ampoule — of suitable shape and capacity, for example, 20 mm in diameter, having one capillary end of about 50 mm length (see Fig. 2).

A-2.1.2 Dely Weighing Tube — As shown in Fig. 3.

### A-2.2 Reagents

A-2.2.1 Hydrogen Peroxide - 60 g/1, neutral to methyl red.

A-2.2.2 Standard Sodium Hydroxide Solution — Approximately 1 N.

A-2.2.3 Methyl Red Indicator — Dissolve 0.1 g of methyl red in 95 percent ethanol (see IS : 323-1959<sup>†</sup>) and dilute to 100 ml with the same ethanol.

### A-2.3 Procedure

A-2.3.1 Test Portion — Carefully mix the sample by shaking the container. If the acid is partially crystallized, slightly heat the container until the sample is liquefied, then carefully mix again. Nearly fill a groundglass stoppered flask (capacity approximately 500 ml and neck of about 30 mm diameter) with the test sample. Slightly heat in a flame the bulb of the spherical glass ampoule previously weighed to the nearest 0.1 mg. Immerse the capillary end of the ampoule into the flask containing the test sample and ensure that the bulb is filled up to about two thirds of its volume during cooling (2 to 3 ml approximately). Withdraw the

<sup>\*</sup>Specification for water for general laboratory use (second revision).

<sup>+</sup>Specification for rectified spirit ( revised ).

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ampoule and carefully wipe the capillary and with filter paper. Seal the capillary end in the oxidizing flame, without loss of glass. Remove from the flame and allow to cool. Wash the capillary and wipe carefully with filter paper. Weigh the ampoule to the nearest 0.1 mg and calculate by difference the mass of the test portion. Alternatively, the test sample may be collected with the help of the Dely weighing tube. For this purpose, weigh the empty tube. Draw the sample into it by applying suction through a rubber tube attached to A. A mark ascertained by a previous run shall be made to indicate the point to which the acid is to be drawn to give the required quantity (3 to 4 g) of oleum. Wipe the tip B carefully with tissue paper and weigh the tube with the sample. The mass of the sample taken shall be obtained by difference.

A-2.3.2 Preparation of Sample Solution — Carefully place the ampoule containing the test portion into a thick-walled conical flask (500 ml capacity, with ground-glass stopper) containing 300 ml of cold water.

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Stopper the flask and shake to break the ampoule containing the test portion. Cool during this operation. Keep cooling and shaking until the vapours are completely absorbed. Remove the stopper and rinse it with water, collecting the washings in the conical flask. By means of a glass rod, grind the fragments of the ampoule and in particular the capillary, which may have remained intact in spite of shaking. Withdraw the glass rod and wash it with water, collecting the washings in the conical flask. Transfer quantitatively the solution to a 500-ml one-mark volumetric flask, dilute to the mark and mix thoroughly. Use this solution for test in A-2.3.3 and A-5.3.

A-2.3.3 Determination — Transfer 100.0 ml of the sample solution, prepared in A-2.3.2 to a 500-ml conical flask. Add 5 ml of hydrogen peroxide solution and genuy boil for 10 minutes. Allow to cool, add two drops of methyl red solution and titrate to the end point by means of the standard sodium hydroxide solution.

A-2.4 Calculation — For calculating the free sulphur trioxide content of oleum, it is necessary to evaluate: (a) the total sulphur trioxide content (see A-2.4.1), (b) water combined as sulphuric acid (see A-2.4.2), and (c) sulphuric acid content (see A-2.4.3).

A-2.4.1 Total Sulphur Trioxide

Total sulphur trioxide (as SO<sub>3</sub>), percent by mass =  $\frac{20.02 \times \frac{VN}{M}}{M}$ 

where

V = volume in ml of standard sodium hydroxide solution used,

N =normality of the alkali, and

M = mass in g of the oleum taken for the test.

A-2.4.2 Evaluation of Water Combined as Sulphuric Acid

Water combined as sulphuric acid, percent by mass (C) = 100-B

where

B =total sulphur trioxide content, percent by mass (A-2.4.1).

A-2.4.3 Evaluation of Sulphuric Acid Content

Sulphuric acid content, percent by mass (D) =  $C \times 5.444$ 

where

C = water combined as sulphuric acid (A-2.4.2).

NOTE - The conversion factor for water to sulphuric acid is 5.444.

### A-2.4.4 Free Sulphur Trioxide Content of Oleum

Free sulphur trioxide percent by mass = 100 - D

where

D = sulphuric acid content, as percent by mass (A-2.4.3).

# A-3. DETERMINATION OF RESIDUE ON IGNITION

A-3.1 Procedure — Weigh to the nearest 10 mg about 50 g of the test sample in a flat-bottom platinum or quartz dish previously ignited at  $800 \pm 50^{\circ}$ C cooled in a desiccator and weighed. If the sample shows any suspended impurities or any sedimentation at the bottom of the container, the contents of the container shall be thoroughly swirled to offer a uniform and homogeneous material before transferring the same to the dish. Heat the dish gradually on a sandbath until the acid is practically driven off. Then transfer to a muffle furnace maintained, at about  $800 \pm 50^{\circ}$ C and heat for 15 minutes or until there is no further fuming of sulphur trioxide. Cool the dish in a desiccator and weigh. Reserve the residue for determination of iron (see A-4) and lead (see A-8).

# A-3.2 Calculation

Residue on ignition, percent by mass =  $\frac{B-A}{M} \times 100$ 

where

B =mass in g of the dish with the residue left after ignition,

 $A = \text{mass in } \mathbf{g}$  of the empty dish, and

M = mass in g of the test sample taken for the test.

### A-4. DETERMINATION OF IRON

A-4.0 General — Two methods are prescribed. Method A is the referee method and method B is the alternative method.

### A-4.1 Method A

A-4.1.1 Apparatus

A-4.1.1.1 Nessler cylinder — 50 ml capacity.

A-4.1.2 Reagents

A-4.1.2.1 Ammonium persulphate

A-4.1.2.2 Concentrated hydrochloric acid — See IS : 265-1976\*.

\*Specification for hydrochloric acid (second revision).

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A-4.1.2.3 Butanolic potassium thiocyanate solution — Dissolve 10 g of potassium thiocyanate in 10 ml of water. Add sufficient *n*-butanol to make up to 100 ml and shake vigorously until the solution is clear.

A-4.1.2.4 Standard iron solution A — Weigh, to the nearest 1 mg, 7.022 g of iron (II) ammonium sulphate hexahydrate and place in a beaker of suitable capacity. Add 50 ml of sulphuric acid (about 2 N) and transfer quantitatively to a 1000-ml one-mark volumetric flask. Dilute to the mark and mix thoroughly. One millilitre of this solution contains 1 mg of Fe.

A-4.1.2.5 Standard iron solution B — Transfer 50.0 ml of the standard iron solution A to a 1000-ml one-mark volumetric flask. Add 5 ml of sulphuric acid ( about 2 N ), dilute to the mark and mix thoroughly. One millilitre of this solution contains 0.05 mg of Fe. The solution should be prepared just before use.

### A-4.1.3 Procedure

A-4.1.3.1 Treat the residue left on ignition in A-3.1 with 2 ml of concentrated hydrochloric acid, add 25 ml of water and heat to facilitate dissolution. Transfer quantitatively to a 500 ml flask, filtering it if necessary, dilute to the mark and mix. Use this solution for determination of iron and lead (see A-8).

A-4.1.3.2 Pipette out 10 ml of the solution into a Nessler cylinder. Add about 30 mg of ammonium persulphate, 1 ml of concentrated hydrochloric acid and 15 ml of butanolic potassium thiocyanate solution. Make up to 50 ml, shake vigorously for 30 seconds and allow the layers to separate. Carry out a control in another Nessler cylinder in the same manner, using 10 ml of standard iron solution B (or 2 ml in the case of explosives grade oleum) and compare the intensity of colour produced in the butanol layers in the two tubes.

A-4.1.3.3 The limit prescribed shall be taken as not having been exceeded, if the intensity of the red colour produced in the butanolic layer with the oleum sample is not deeper than that produced in the control test.

### A-4.2 Method B

A-4.2.1 Apparatus

A-4.2.1.1 Spectrophotometer or photoelectric colorimeter

A-4.2.2 Reagents

A-4.2.2.1 Hydrochloric acid - approximately 1 N.

A-4.2.2.2 Hydroxylammonium chloride solution — Dissolve 10 g of hydroxylammonium chloride in water and dilute to 100 ml.

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|--|---|
| A-4.2.2.3 Ammonium acetate solution acetate in water and dilute to 100 ml.   | m — Dissolve 30 g of ammonium   |
| A-4.2.2.4 Bipyridyl solution — Diss<br>of approximately 1 N hydrochloric acid  | olve 1 g of 2, 2'-bipyridyl in 10 ml<br>and dilute to 100 ml.   |
| A-4.2.2.5 Standard iron solution $C$ -<br>solution B (A-4.1.2.5) to a 1 000 ml o<br>the mark and mix thoroughly. One mill<br>of Fe. The solution should be prepared  | - Transfer 200 ml of standard iron<br>ne-mark volumetric flask, dilute to<br>ilitre of this solution contains $10\mu g$<br>l just before use.   |
| A-4.2.3 Procedure  |   |
| A-4.2.3.1 Preparation of calibration<br>eleven 100-ml one-mark volumetric flask<br>iron solution C as indicated below:   | curve — Into each of a series of s, place the quantities of standard  |
| Standard Iron Solution C   | Corresponding to Fe   |
| ml   | μĝ  |
| 0 (Compensation solution   | on) 0   |
| 5.0  | 50  |
| 10.0   | 100   |
| 15.0   | 150   |
| 20.0   | 200   |
| 30.0   | 300   |
| 35.0   | 350   |
| 40.0   | 400   |
| 45 <b>·0</b>   | 450   |
| 50.0   | 500   |
| Add to each volumetric flask an an<br>to approximately 50 ml, then 2 ml of hy<br>ammonium chloride solution and after<br>acetate solution and 1 ml of the 2, 2'-bip<br>mix thorougly and wait for 10 minutes.<br>spectrophotometer in a 1-cm cell at a wa<br>a photoelectric colorimeter using approp<br>ment to zero optical density and usi<br>solution. Prepare a calibration chart wi<br>per 100 ml of the standard matching sol<br>ponding values of optical density as ord | mount of water sufficient to dilute<br>drochloric acid, 2 ml of hydroxyl-<br>er 5 minutes, 5 ml of ammonium<br>yridyl solution. Dilute to the mark,<br>Carry out the measurement on the<br>avelength of about 522 nm (or on<br>priate filter), adjusting the instru-<br>ng as reference the compensation<br>th the iron content in micrograms<br>ution as abseissae and the corres-<br>inate. |
| A-4.2.3.2 Transfer an aliquot of th<br>containing between 50 and 500 microgra  | ne solution, prepared in A-4.1.3.1<br>ms of iron, to a 100-ml one-mark  |
|  |   |

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volumetric flask. Dilute to approximately 50 ml, add successively 2 ml of hydrochloric acid, 2 ml of the hydroxylammonium chloride solution, and after 5 minutes, 5 ml of ammonium acetate solution and 1 ml of 2, 2'-bipyridyl solution. Dilute to the mark, mix and wait for 10 minutes. Carry out the spectrophotometric (or photoelectric colorimetric) measurement according to the procedure given in. A-4.2.3.1, adjusting the instrument to zero optical density using as reference the blank test solution (see A-4.2.3.3).

A-4.2.3.3 Blank test — Simultaneously carry out a blank test using the same procedure and quantities of all reagents employed in the test.

A-4.2.4 Calculation — From the calibration chart (see A-4.2.3.1), determine the iron content corresponding to the photometric measurement.

Iron (as Fe), percent by mass =  $\frac{A \times 500 \times 100}{V \times M}$ 

where

- $A = \max s$  in g of iron present in the aliquot of the sample solution,
- V = volume in ml of the sample solution taken for the colour reaction, and

M =mass in g of the sample taken for the test in A-3.1.

# **A-5. DETERMINATION OF CHLORIDES**

# A-5.1 Apparatus

A-5.1.1 Nessler Cylinder — two, capacity 100 ml.

# A-5.2 Reagents

A-5.2.1 Dilute Nitric Acid — approximately 10 percent (m/v).

A-5.2.2 Silver Nitrate Solution — 4.2 percent (m/v).

A-5.2.3 Standard Sodium Chloride Solution — Dissolve 0.0165 g of pure ignited sodium chloride in 1 000 ml of water.

# A-5.3 Procedure

A-5.3.1 Take with the help of a burette an aliquot of the sample solution prepared in A-2.3.2 so as to represent 0.5 g of the sample and transfer it to Nessler cylinder. Add 1 ml of dilute nitric acid and 1 ml of silver nitrate solution. Make it up to the mark and shake well to get a homogeneous solution.

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A-5.3.2 In another Nessler cylinder, dilute 1 ml of standard sodium chloride solution with 80 to 90 ml of water and add 1 ml of dilute nitric acid and 1 ml of silver nitrate solution and make the volume to 100 ml. Compare the opalescence developed in the standard solution and the solution containing the sample of oleum.

A-5.3.3 The limit prescribed in Table 1 shall be taken as not having been exceeded, if the turbidity produced in the test solution is not greater than that produced in the standard solution.

### A-6. TEST FOR TOTAL NITROGEN

### A-6.1 Reagents

A-6.1.1 Sodium Hydroxide Solution — approximately 40 percent.

A-6.1.2 Devarda's Alloy — containing 45 parts. aluminium, 50 parts copper and 5 parts zinc. Heat the aluminium in a Hessian crucible in a furnace until the aluminium begins to melt. Add copper in small portions and heat until liquefied. Then plunge zinc into the molten mass. Cover the crucible and heat the mixture for a few minutes and stir with an iron rod. Allow it to cool slowly with the cover on and then pulverize the crystallized mass to particle size of 0.2 to 0.3 mm.

### A-6.1.3 Concentrated Hydrochloric Acid — See IS : 265-1976\*.

A-6.1.4 Nessler Solution — Dissolve 10 g of potassium iodide in 10 ml of ammonia-free water and add to it slowly, with stirring, a saturated aqueous solution of mercuric chloride until a slight permanent precipitate forms. Add 30 g of potassium hydroxide and, when it has dissolved, add 1 ml more of the mercuric chloride solution and dilute to 200 ml with ammonia-free water.

A-6.1.5 Standard Ammonium Chloride Solution — Prepare a solution containing 0.0849 g of ammonium chloride per litre. One millilitre of this solution contains 0.1 mg of total nitrogen (as HNO<sub>3</sub>).

# A-6.2 Procedure

A-6.2.1 Add cautiously 10 ml of the material to 100 ml of water in a 500-ml volumetric flask and, after cooling, carefully neutralize with sodium hydroxide solution till the liquid is distinctly alkalinc. Add 2 g of Devarda's alloy and fit a distillation head and condenser. After one hour, distil off 50 ml into a conical flask containing one drop of hydrochloric acid. Transfer the distillate to a Nessler cylinder, add 1 ml of sodium hydroxide solution and 2 ml of Nessler solution.

<sup>\*</sup>Specification for hydrochloric acid ( second revision ).

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A-6.2.2 The limit prescribed in Table 1 shall be taken as not having been exceeded, if the colour of the solution is not deeper than that obtained by nesslerizing 9.5 ml of a standard solution of ammonium chloride in the same total volume of the solution.

# A-7. DETERMINATION OF SULPHUR DIOXIDE

A-7.0 Methods — Two methods are prescribed. Method A is for routine analysis and method B is the referee method.

# A-7.1 Method A

A-7.1.1 Reagents

A-7.1.1.1 Potassium iodide solution — approximately 10 percent (m/v).

A-7.1.1.2 Standard iodine solution - 0.05 N.

A-7.1.1.3 Standard sodium thiosulphate solution -0.05 N.

A-7.1.1.4 Starch solution -0.5 percent m/v, freshly prepared.

A-7.1.1.5 Concentrated sulphuric acid — see IS: 266-1977\*.

A-7.1.2 Procedure — Weigh accurately about 5 g of the test sample as prescribed in A-2.3.1 and cautiously dilute in 200 ml of water contained in a 500-ml conical flask placed in a deep porcelain dish containing cold water. Add to the diluted solution 10 ml of potassium iodide solution and 20 ml of standard iodine solution. In case the iodine colour disappears altogether, another 20 ml of iodine solution may be added. Titrate the excess iodine with standard sodium thiosulphate using starch as indicator towards the end.

A-7.1.2.1 Carry out a blank test using the same reagents, taking 5 ml of concentrated sulphuric acid under similar conditions.

A-7.1.3 Calculation

Sulphur dioxide (as  $SO_2$ ), percent by mass  $= \frac{(B-A)N \times 3\cdot 2}{M}$ 

where

- B = volume in ml standard sodium thiosulphate solution used in blank test,
- A = volume in ml of standard sodium thiosulphate solution used with the test sample,

N = normality of sodium thiosulphate solution, and

M = mass in g of the test sample taken for the test.

Note — In case A is equal to B or higher than B, then there will be no sulphur dioxide in the sample of oleum.

<sup>\*</sup>Specification for sulphuric acid (second revision).

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# A-7.2 Method B

A-7.2.1 Apparatus-As shown in Fig. 4.



- A-500 ml, three-necked, round-bottomed flask.
- B 200 ml, graduated separating funnel fitting into one of the three necks of the flask.
- C Tube with cock, inserted into the central neck of the flask, extending about 1 cm from the bottom of the flask. The end of the tube is fitted with a sintered glass diaphragm.
- D Three-way cock.
- E Four wash bottles ( $E_1$ ,  $E_2$ ,  $E_3$ ,  $E_4$ ), capacity 100 ml, fitted with a sintered glass diaphragm (porosity 3 to 15 µm) at the end of the immersion tube, for gas distribution. One branch of the cock (D) is connected to the third neck of the roundbottomed flask (A); the second branch is connected to the first wash-bottle ( $E_1$ ), which can be connected and disconnected according to requirement through the cock (D); the third branch is connected to the other wash-bottles linked in series.

FIG. 4 APPARATUS FOR DETERMINATION OF SULPHUR DIOXIDE

A-7.2.2 Reagents

A-7.2.2.1 Nitrogen - Pure. See IS: 1747-1972\*.

A-7.2.2.2 Hydrogen peroxide -30 g/1, free from sulphate ions.

\*Specification for nitrogen ( first revision ).

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with boiling water until the filtrate is free from chlorides; the liquid flowing from the funnel should remain clear on addition of a few drops of silver nitrate solution. Place the filter paper and its contents in a platinum crucible, previously tared after igniting in a furnace at 600 to 800°C and cooling in a desiccator. Dry in an oven at about 110°C, and carefully char the filter paper at a low temperature to avoid flaming, then ignite in a furnace at 600 to 800°C. Remove the crucible, add a drop of sulphuric acid and again ignite. Remove the crucible, place in a desiccator, allow to cool and weigh.

A-7.2.4 Calculation

Sulphur dioxide content (SO<sub>2</sub>), percent by mass =  $\frac{A \times 0.2744 \times 100}{M}$ 

where

A = mass in g of barium sulphate precipitate, and M = mass in g of the test portion.

Note — The conversion factor from barium sulphate to SO<sub>2</sub> is 0.274 4.

# A-8. DETERMINATION OF LEAD

A-8.1 Apparatus — All glassware, including the reagent bottles, shall be of borosilicate glass or other quality free from lead, or alternatively plastics materials may be used. It shall be washed with nitric acid solution approximately 7 N, and rinsed with water three times.

A-8.1.1 Spectrophotometer or Photoelectric Colorimeter — Fitted with filters giving maximum transmission between 500 and 540 nm.

# A-8.2 Reagents

**A-8.2.1** Hydroxylammonium Chloride Solution — 10 percent (m/v).

A-8.2.2 Ammonium Citrate Solution — 10 percent (m/v).

A-8.2.3 Ammonia Solution — approximately 5 N.

Nore — A freshly prepared solution shall be used in order to avoid too high a value being obtained by the blank test, due to dissolved lead. In fact, dilute ammonia dissolves lead contained in the glass more rapidly than does concentrated ammonia.

A-8.2.4 Potassium Cyanide Solution — 5 percent (m/v).

A-8.2.5 Chloroform — redistilled in a borosilicate glass apparatus with ground joints.

A-8.2.6 Dithizone Solution - 0.025 g/1, in chloroform.

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| Volume of Lead Standard<br>Solution | Corresponding Mass<br>of Pb |
|-------------------------------------|-----------------------------|
| ml                                  | μg                          |
| 0 (compensatio                      | n solution) 0               |
| 1.0                                 | 10                          |
| 2.0                                 | 20                          |
| 3.0                                 | 30                          |
| 4.0                                 | 40                          |
| 5.0                                 | 50                          |
| 6.0                                 | 60                          |
| 7.0                                 | 70                          |
| 8· <b>O</b>                         | 80                          |
| 9.0                                 | 90                          |
| 1 <b>0·0</b>                        | 100                         |

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A-9.2 Procedure — Transfer to a series of 100 ml conical flasks, aliquots of standard arsenic solution corresponding to 0, 5, 10, 15, 20 and 25  $\mu$ g of arsenic and proceed as given in A-9.2.1.

A-9.2.1 Add 10 ml of concentrated hydrochloric acid and dilute to 50  $\pm$  5 ml with water. Add 2 ml of potassium iodide and stannous chloride solution respectively. Mix well and let it stand for 15 to 20 minutes. Pack lightly the top third of the connecting tube with impregnated absorbent cotton wool and assemble on the absorption train. Transfer 5.0 ml of silver diethyldithiocarbamate solution to absorption tube C. After 15 to 20 minutes, introduce 5 g of zinc granules into the conical flask A and quickly reassemble the apparatus. Allow the reaction to proceed for 45 to 60 minutes at room temperature.

A-9.2.2 Spectrophotometric Measurements — Disconnect the absorption tube and tilt the absorber so that the reagent solution flows back and forth between the absorber and bulb to disperse the solid contents, if any, and to mix in the solution well. Transfer the solution to a photometric cell and measure its absorbance at the wavelength of maximum absorption, 540 nm, using water as reference liquid.

NOTE — The colour of the dispersion is not very stable for long time and hence absorptiometric (nephelometric) measurement should be made within 2 hours of the development of colour. Care should also be taken to prevent the evaporation of solution as its volume is small.

In the case of fritted glass absorber raise and lower the connecting tube into the absorber several times, to allow the solution to pass through the frit back and forth effecting the dispersal of the red deposit. Let the connecting tube finally drain into the absorber.

10  $\mu$ g standard shall have an absorption of the order of 0.4.

A-9.2.3 Plotting of the Calibration Curve — Calculate corrected absorbance by subtracting the reading obtained for the solution containing no standard arsenic solution from the observed reading. Plot a graph of corrected absorbance of solution against their arsenic contents.

A-9.2.4 Blank Test — Carry out a blank test, as prescribed in A-9 to A-9.2.2 omitting the sample.

**A-9.3 Calculation** — Calculate the corrected absorbance by subtracting the value obtained for the blank solution from that obtained for the test solution and read from the calibration curve the corresponding mass of arsenic.

Arsenic content, parts per million by mass  $=\frac{M_1}{M_2}$ 

where

 $M_1 = \text{mass in g of arsenic found, and}$  $M_2 = \text{mass in g of sample in the solution tested.}$ 

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|----------------------------------|-------------------------------------|--|
| Α                                | PPENDIX B                           |  |
|                                  | ( Clause 3.4 )                      |  |
| CORRELATION BET<br>AND FREE      | WEEN RELATIVE D<br>SULPHUR TRIOXIDI | ENSITY OF OLEUM<br>E CONTENT                         |
| Relative Density at<br>35°C/15°C | Density in<br>Degree Baume          | Free Sulphur Trioxide<br>Content, Percent<br>by Mass |
| (1)                              | (2)                                 | (3)  |
| 1.8186                           | 65.3                                | 0  |
| 1.822 8                          | 65·5                                | 1  |
| 1.831 5                          | 65.8                                | 2 3  |
| 1.8360                           | 66.0                                | 4  |
| 1.839 3                          | 66·2                                | 5  |
| 1.842 5                          | 66.3                                | 6  |
| 1.846 1<br>1.840 P               | 66·5                                | 7  |
| 1.853 1                          | 66.8                                | 0<br>9   |
| 1.8565                           | 6 <b>6</b> ·9                       | 10   |
| 1.859 6                          | 67· <b>0</b>                        | 11   |
| 1.8627                           | 67.2                                | 12   |
| 1.866 0                          | 67·3                                | 13   |
| 1.872.4                          | <b>67</b> .6                        | 15   |
| 1.875 6                          | 67.7                                | 16   |
| 1.879 3                          | 67.8                                | 17   |
| 1.8830                           | 68·0                                | 18   |
| 1.8919                           | 68.4                                | 20   |
| 1.896.9                          | 68.6                                | 21   |
| 1.9020                           | 68.8                                | 22   |
| 1.905 6                          | 68.9                                | 23   |
| 1.909 2                          | 69·1                                | 24   |
| 1.912 5                          | 09.2                                | 25   |
| 1.9158                           | 69·3                                | 26   |
| 1.9220                           | 69.6                                | 27   |
| 1.9250                           | 69.7                                | 29   |
| 1·9 <b>28 0</b>                  | 69·8                                | 30   |
|                                  | 24                                  |  |
|                                  |                                     |  |

| Relative Density at<br>35°C/15°C<br>(1)<br>1.930 9<br>1.933 8<br>1.937 2<br>1.940 5<br>1.943 9<br>1.943 9<br>1.947 4<br>1.950 4<br>1.953 4<br>1.955 9<br>1.958 4 | Density in<br>Degree Baume<br>(2)<br>69·9<br>70·0<br>70·1<br>70·3<br>70·4<br>70·5<br>70·7<br>70·8<br>70·9 | Free Sulphur Trioxide<br>Content, Percent<br>by Mass<br>(3)<br>31<br>32<br>33<br>34<br>35<br>36<br>37 |
|--|---|---|
| (1)<br>1.930 9<br>1.933 8<br>1.937 2<br>1.940 5<br>1.943 9<br>1.947 4<br>1.950 4<br>1.953 4<br>1.955 9<br>1.958 4  | (2)<br>69·9<br>70·0<br>70·1<br>70·3<br>70·4<br>70·5<br>70·7<br>70·8<br>70·9                               | (3)<br>31<br>32<br>33<br>34<br>35<br>36<br>37   |
| 1.930 9<br>1.933 8<br>1.937 2<br>1.940 5<br>1.943 9<br>1.947 4<br>1.950 4<br>1.953 4<br>1.955 9<br>1.958 4   | 69·9<br>70·0<br>70·1<br>70·3<br>70·4<br>70·5<br>70·7<br>70·8<br>70·9                                      | 31<br>32<br>33<br>34<br>35<br>36<br>37  |
| 1 ·947 4<br>1 ·950 4<br>1 ·953 4<br>1 ·955 9<br>1 ·958 4   | 70 5<br>70 7<br>70 8<br>70 9  | 36<br>37  |
|  | 71.0  | 38<br>39<br>40  |
| 1-959 8<br>1-961 2<br>1-962 7<br>1-964 3<br>1-965 8  | 71.0<br>71.1<br>71.1<br>71.2<br>71.2  | 41<br>42<br>43<br>44<br>45  |
| 1·967 2<br>1·968 7<br>1·970 2<br>1·971 7<br>1·973 3  | 71·3<br>71·3<br>71·4<br>71·5<br>71·5  | 46<br>47<br>48<br>49<br>50  |
| 1·974 1<br>1·974 9<br>1·975 5<br>1·976 0<br>1·976 6  | 71.5<br>71.6<br>71.6<br>71.6<br>71.6<br>71.6  | 51<br>52<br>53<br>54<br>55  |
| 1 977 2<br>1 976 3<br>1 975 4<br>1 974 6<br>1 973 8  | 71.7<br>71.6<br>71.6<br>71.6<br>71.6<br>71.5  | 56<br>57<br>58<br>59<br>60  |
| 1·972 3<br>1·970 <b>9</b><br>1·969 1<br>1·967 2<br>1·965 4   | 71·5<br>71·4<br>71·4<br>71·3<br>71·2  | 61<br>62<br>63<br>64<br>65  |

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|----------------------------------|--|--|
| Relative Density at<br>35°C/15°C | Density <b>in</b><br>De <b>gr</b> ee Baume | Free Sulphur Trioxide<br>Content, Percent<br>by Mass |
| (1)                              | (2)  | (3)  |
| 1.963 6                          | 71.2                                       | 66   |
| 1.961 8                          | 71.1                                       | 67   |
| 1.9600                           | 71.0                                       | 68   |
| 1.958 2                          | 71.0                                       | 69   |
| 1.956 4                          | 70 9                                       | 70   |
| 1.953 3                          | 70· <b>8</b>                               | 71   |
| 1.950 2                          | 70·6                                       | 72   |
| 1.947 2                          | <b>70</b> .6                               | 73   |
| 1.944 2                          | 70 4                                       | 74   |
| 1.9411                           | 70-3                                       | 75   |
| 1.937 9                          | 7 <b>0</b> ·2                              | 76   |
| 1.9347                           | 70-1                                       | 7 <b>7</b>   |
| 1.931 5                          | 69 <b>·9</b>                               | 78   |
| 1.928 3                          | <b>6</b> 9 <b>·</b> 8                      | 79   |
| 1 925 1                          | 6 <b>9</b> ·7                              | 80   |
| 1.9217                           | 69·5                                       | 81   |
| 1.918 3                          | <b>6</b> 9 · <b>4</b>                      | 82   |
| 1.914 9                          | 6 <b>9</b> ·3                              | 83   |
| 1 <b>9</b> 11 5                  | 69.1                                       | 84   |
| 1.908 0                          | 69· <b>0</b>                               | 85   |
| 1.9046                           | 68.8                                       | 86   |
| 1.9013                           | 68.7                                       | 87   |
| 1.898 0                          | 68.6                                       | 88   |
| 1.893 4                          | 68.4                                       | 89   |
| 1 888 8                          | 68.2                                       | 90   |
| 1.8844                           | 68.1                                       | 91   |
| 1.880 0                          | 68 <b>0</b>                                | 92   |
| 1.8756                           | 67·7                                       | 93   |
| 1.871 2                          | 67· <b>5</b>                               | 94   |
| 1.865 9                          | 67.3                                       | 95   |
| 1.860 5                          | 67.1                                       | 96   |
| 1.8546                           | <b>66</b> ·8                               | 97   |
| 1 <b>84</b> 8 8                  | 6 <b>6</b> ·6                              | 98   |
| 1.8429                           | 66·3                                       | 99   |
| 1 837 0                          | 66.1                                       | 100  |
|                                  | 26   |  |

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# APPENDIX C

# ( Clause 5.1 )

# SAMPLING OF OLEUM, TECHNICAL

# **C-1. GENERAL REQUIREMENTS**

C-1.1 Samples shall not be taken in an exposed place.

C-1.2 The sampling instrument shall be clean and dry.

C-1.3 The material shall be sampled in liquid condition.

C-1.4 To draw a representative sample, the contents of each container selected for sampling shall be mixed as thoroughly as possible by suitable means.

C-1.5 Precautions shall be taken to protect the samples, the material being sampled, the sampling instrument and the containers for samples from adventitious contamination.

C-1.6 The sample shall be placed in suitable, clean, dry and air-tight glass containers, on which the material has no action.

C-1.7 The sample containers shall be of such a size that they are almost completely filled by the sample.

C-1.8 Each sample container shall be sealed air-tight with a suitable stopper after filling, and marked with full details of sampling and the month and year of manufacture of the material.

C-1.9 Samples shall be stored in a cool and dry place.

# C-2. SCALE OF SAMPLING

C-2.1 Lot — All containers in a single consignment of the material drawn from a single batch of manufacture shall constitute the lot. If a consignment is declared to consist of different batches of manufacture the batches shall be marked separately, and the groups of containers in each batch shall constitute separate lots.

C-2.2 Samples shall be tested from each lot separately for judging the conformity of the material to the requirements of the specification. For this purpose five containers shall be selected at random from each lot.

Nore — If the number of containers in the lot is five or less, the number of containers to be selected and the criteria for conformity of the lot of the specification shall be as agreed to between the purchaser and the supplier.

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