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Workplace air — Analysis of airborne water immiscible mineral oil droplets and vapor with Fourier — Transform infrared spectroscopy

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Foreword

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The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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Introduction

Mineral oils are by-products from the distillation of crude oil. The mineral oils produced in this way are used in a wide range of modern applications. Their composition varies according to their application. Mineral oils contain saturated hydrocarbons, cyclic saturated hydrocarbons, aromatic compounds, and unsaturated hydrocarbons.

They are used as fuels, for example in the form of petroleum, diesel and kerosene for internal combustion engines, as heating oil, and as lubricants in the metal processing and application industries. Mineral oils are also found in cosmetics and medical applications. This wide range of applications gives rise to exposure for human beings and the natural environment to mineral oils in the form of vapours and droplets. Since mineral oils and their constituents are harmful to health, it is important for quantification of the exposure to be possible and where necessary for corrective measures to be taken.

The method described in this document is intended to support swift, cost-effective and straightforward quantification of the concentration of mineral oil in a large number of samples. This method has been validated in accordance with the provisions of EN 13936 and ISO 20581 and has an expanded measurement uncertainty in the region of 20 % to 30 % within the measurement range stated.

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Workplace air — Analysis of airborne water immiscible mineral oil droplets and vapor with Fourier — Transform infrared spectroscopy

1 Scope

The method described in this document quantifies the absolute exposure to mineral oil vapours and droplets, within a concentration range from 0.5 mg/m^3 to 125 mg/m^3 , in the inhalable fraction of the workplace air.

This document contains comprehensive information and instructions on the equipment and chemicals to be used.

This method is applicable for water soluble oils and metal working fluids.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements 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 1042, Laboratory glassware Sone-mark volumetric flasks

ISO 7708, Air quality — Particle size fraction definitions for health-related sampling

ISO 13137:2013, Workplace atmospheres pumps for personal sampling of chemical and biological agents — Requirements and test methods 7 dec01/iso-tdis-23506

ISO 18158, Workplace air — Terminology

ISO 20581, Workplace air — General requirements for the performance of procedures for the measurement of chemical agents

EN 13936, Workplace exposure — Procedures for measuring a chemical agent present as a mixture of airborne particles and vapour — Requirements and test methods

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 18158 apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at https://www.iso.org/obp
- IEC Electropedia: available at http://www.electropedia.org/

4 Principle

A defined airflow is drawn over the sampler with the aid of a pump. The droplets are deposited on the glass fibre filter, the vapours on the polymer resin. By extraction of the glass fibre filter and the polymer resin, the adsorbed mineral oils are desorbed and can be analysed by means of FTIR and evaluated.

The Gesamtstaub-Gas-Probenahme (GGP) sampling system enables both droplets and vapours to be collected but any other personal sampler capable of collecting the inhalable droplets and vapours

simultenously can be used. A glass fibre filter is used to collect the droplets. A downstream sampling tube containing polymer resin is used to collect the vapours and more volatile components in the mineral oil. The GGP system cannot be used for distinction between exposure to gases and droplets, since droplets collected during sampling can vaporize from the filter and collect on the polymer resin.

Following extraction, the samples are analysed by Fourier transform infrared spectroscopy (FTIR) in the wavenumber range from 3 $000~\rm cm^{\text{-}1}$ to 2 $800~\rm cm^{\text{-}1}$. This method is not specific for mineral oil, since it detects all substances with C-H bonds collected on the sampler.

The principle of the method described here for the quantification of mineral oil is based upon activation of the C-H bonds of the hydrocarbons in the mineral oil. Hydrocarbons absorb light in the wavenumber range from 3 000 cm⁻¹ to 2 800 cm⁻¹. This enables the concentration of mineral oil at the site of sampling to be determined by external calibration against a defined mineral oil at known concentrations.

The method described in this document is based upon the "MAK Collection for Occupational Health and Safety 2016, Vol 1, No 1", "Cooling lubricants and other complex hydrocarbon mixtures, immiscible with water – droplets and vapours" [2].

5 Apparatus and reagents

5.1 Collecting materials for analysis and other chemicals

5.1.1 Glass fibre filter.

The droplets shall be collected by a binder-free 85/90 glass fibre filter with a diameter of 37 mm.

Glass fibre filters generally have no blank value. This shall however be verified beforehand.

5.1.2 Polymer resin.

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Poly[styrene-divinylbenzene], a mesoporous polymer residenthe particle size range from 0,5 mm to 0,9 mm shall be used for collection of the vapour.

Polymer resins are well suited as a collection phase for mineral oils. This has been demonstrated with reference to Amberlite® XAD-2¹). XAD-2 requires cleaning prior to use in case it exhibits a blank value.

Any other polymer resin equivalent to XAD-2 can be accepted if it is validated according to Annex A.

5.1.3 Tetrachloroethene.

Tetrachloroethene, also know as perchloroethylene, in the purest grade (≥99,0 %) is used to extract the samples and to prepare the calibration standards, and to clean the polymer resin if necessary.

Any other solvent can be used if it is validated according to Annex A.

5.1.4 Mineral oil, for calibration purposes.

The mineral oil used for evaluation of the air samples shall be that used at the workplace.

Fresh unused mineral oil is usually preferred for practical reasons.

If there are several mineral oil products used, it is recommended to use the one with the lower mineral oil content. This leads to a conservative result.

¹⁾ Amberlite® XAD-2 is the trade name of a product supplied by Sigma Aldrich This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of the product named. Equivalent products may be used if they can be shown to lead to the same results.

5.1.5 Liquid paraffin standard.

In case it not be possible to obtain a site-specific oil sample or if this sample is not soluble in tetrachloroethene, a mineral oil standard for spectroscopy, namely liquid paraffin, CAS number 8012-95-1, should be used for evaluation.

The different response factors against the mineral oils of the workplaces which were used for this validation are listed in <u>Table A.13</u> and <u>Table A.14</u>.

5.2 Sampler equipment

The following equipment is required for sampling.

The performance of the samplers used shall match the criteria for inhalable fraction as specified in ISO 7708.

- **5.2.1 Sampling pump**, conforming with the requirements of ISO 13137.
- **5.2.2 Personal sampler**, shall be used for the simultaneous collection of inhalable droplets and vapours. Any personal sampler capable of collecting the inhalable droplets and vapours simultaneously may be used; for the validation experiments written in this document, the GGP sampling system, with matching cone for the specified flow rate (3.5 l/min) was used[4].

The GGP sampler is a modification of the GSP sampler (see CEN/TR 15230^[11]), which match the criteria of ISO 7708. Sampling efficiency of this modification was determined by George C. Dragan et al. (2015)^[9].

5.2.2.1 Various lengths of **tubes** (e.g. outer diameter: 9 mm) or 10 mm; inner diameter: 6 mm), for connecting the personal sampler to the pump.

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5.2.2.2 Matching **filter capsule** for the 37 mm glass fibre filter, for insertion into the personal sampler. e84a857dec01/iso-fdis-23506

5.2.2.3 Plastic cartridge manufactured from polyvinylidene fluoride (PVDF).

Matching plastic tube (outer diameter: 20 mm; inner diameter: 16 mm, length: 70 mm) for the polymer resin for insertion into the personal sampling system, with high-grade steel sieve stopper and plastic sealing caps.

- **5.2.2.4 Silicone adapter**, which shall fit onto the sampler head to connect the flow meter for measuring/setting the air flow.
- **5.2.3 Flow meter**, conforming with the requirements for test instruments in ISO 13137:2013, 6.2.
- **5.2.4 Glass metering aid**, for filling the plastic cartridge with 8 ml polymer resin.
- **5.2.5** A 500 ml glass **Erlenmeyer flask** for the tilting dispenser head.

5.3 Analytical equipment

- **5.3.1 Dispenser**, for organic solvents, with a volume of 1 ml to 10 ml, shall be used for extraction of the samples and preparation of the calibration.
- **5.3.2 Piston pipettes with tips for organic solvents**, with a volume range of 30 μ l to 300 μ l and of 0,5 ml to 5,0 ml are required for preparation of the calibration.

30 ml screw-cap glass vials. 5.3.3

The filter and the polymer resin shall be transferred separately to the 30 ml extraction vessels for extraction.

5.3.4 Screw cap with septum, for sealing extraction vessels.

The extraction vessels shall be sealed by means of plastic screw caps and a PTFE septa.

Refrigerator, at 4 °C to 8 °C. 5.3.5

5.3.6 Fourier transform infrared spectrometer (FTIR)

Spectral range 7 800 cm⁻¹ – 350 cm⁻¹

Spectral resolution: Better than 0,8 cm⁻¹

Sensitivity: S/N ratio 2 200 cm⁻¹ – 2 100 cm⁻¹

8 000: 1 peak to peak in 5 s

22 000: 1 peak to peak in 1 min

Deuterated triglycine sulfate (DTGS) detector

- 5.3.7 A suitable quartz-glass cuvette for infrared spectrometry with 10 mm path length and polytetrafluoroethylene (PTFE) cuvette stoppers. (Standards.iteh.ai)
- One 5 ml glass syringe with Luer-lock cone and cannula for filling the cuvette. 5.3.8

- Glass beakers, 150 ml and 50 ml. as/2428574-2017 Street St 5.3.9
- **5.2.10** Glass volumetric flasks with glass stoppers, 100 ml ± 0,1 ml, A, ISO 1042, in 20 °C, with glass stopper.
- 5.3.11 100 ml glass Erlenmeyer flask with ground joint, with glass stopper.
- **5.3.12** Analytical balance, capable of weighing at least to 320 g with a precision of 0,1 mg.

5.4 Other equipment

- 1 000 ml Erlenmeyer flask with ground joint, for treating the polymer resin with tetrachloroethene and ethanol.
- **5.4.2 Ultrasonic bath**, with sufficient capacity to hold two 1 000 ml erlenmeyer flasks (cross reference to be included).
- 5.4.3 **Drying cabinet or oven with** fan-assisted circulation and **extractor**, set at 60 °C.
- 1 l glass **vacuum flask** with matching rubber stopper. 5.4.4
- Porcelain Büchner funnel, for round filters (diameter: 125 mm; volume: approximately 1 000 ml) for vacuum filtration of the polymer resin following treatment with tetrachloroethene and ethanol.

5.4.6 Round glass fibre filters, for vacuum filtration (diameter: 125 mm).

5.4.7 Water-jet pump or vacuum pump.

Where a water-jet pump is used, an absorption bottle shall be placed between the vacuum flask and the water-jet pump in order to prevent tetrachloroethene from entering the waste water.

5.4.8 Rubber hose, for connecting the vacuum flask to the water-jet pump/vacuum pump.

5.4.9 Sieve

Mesh: > 0.5 mm

- **5.4.10** A large porcelain **Petri dish** (diameter: 30 cm) for drying the polymer resin.
- **5.4.11** A glass powder **funnel** is required for polymer resin transfer.

6 Polymer resin preparation

6.1 Drying

Polymer resin is generally supplied in the damp state and shall therefore first be dried. The polymer resin shall be dried in a large Petri dish at 60 °C in the drying cabinet (5.4.3).

6.2 Sieving (standards.iteh.ai)

Polymer resin in the particle size range from 0,5 mm to 0,9 mm shall be used for sampling; the dried polymer resin shall therefore be sieved talog/standards/sist/3abfd522-7e6b-4200-b287-

e84a857dec01/iso-fdis-23506 NOTE If the cartridges are filled with polymer resin with a particle size of <0,5 mm, the pump can fail to reach or maintain the desired flow rate owing to excessive pressure drop.

6.3 Purification of unused polymer resin

6.3.1 General

In the as-delivered state, polymer resin is not free of a blank value in the measurement range of 2 800 cm⁻¹ to 3 000 cm⁻¹, and shall therefore be purified prior to use.

Ethanol (\geq 99,5 %) is required for initial purification of the polymer resin.

6.3.2 Step 1: Preparatory purification with ethanol

Fresh polymer resin shall first be treated with ethanol in order to eliminate the blank value. For this purpose, approximately 400 ml of the sieved polymer resin shall be placed in a 1000 ml Erlenmeyer flask and ethanol added until the polymer resin floats. This shall then be treated in the ultrasonic bath at $50 \,^{\circ}\text{C}$ for $1 \,^{\circ}\text{h}$ and then filtered using the Buchner funnel. This process shall be performed a total of four times using fresh solvent. The vapours shall be routed into the extractor. The polymer resin shall then be dried for $24 \,^{\circ}\text{h}$ at $60 \,^{\circ}\text{C}$ in the drying cabinet (5.4.3).

6.3.3 Step 2: Purifying with tetrachloroethene

Polymer resin that has been purified with ethanol shall also be treated subsequently with tetrachloroethene. The procedure is the same as for purification with ethanol. Polymer resin purified

with tetrachloroethene shall be dried for at least 7 days in the drying cabinet with fan-assisted circulation.

NOTE The purification step with tetrachloroethene described here is also be employed for used polymer resin in preparation for use in further analyses.

The transmission spectrum of the blank value of the polymer resin is generally a negative spectrum. If this is not the case, purification shall be repeated. The laboratory should document purified polymer resin shelf-life if it is not used directly after purification.

7 Sampling

7.1 Sampler preparation

7.1.1 Glass fibre filter (droplets)

The glass fibre filter shall be placed in the filter capsule, fixed, and sealed for transport with the caps.

7.1.2 Polymer resin (vapours)

Approximately 8 ml polymer resin, corresponding approximately to 3,2 g, shall be placed in the plastic cartridge. The cartridge shall be closed at both ends with the high-grade steel sieve stoppers. The cartridges shall be closed with plastic stoppers for transport. For alternative personal samplers, the amount of polymer resin may be altered as appropriate following the requirements of ISO 20581.

7.1.3 Marking

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The samplers used should be marked unambiguously in order to prevent confusion with other samplers. This also assures clear assignment of the filter capsule to the polymer resin cartridge.

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7.1.4 Cleaning

All reusable equipment, such as the filter capsule and cartridge, shall be cleaned each time before use with detergent and water or suitable solvent such as ethanol. Any oil residues or other soiling shall be removed. Contaminated equipment can lead to incorrect results.

7.1.5 Assembly

The filter capsule and the polymer resin cartridge shall be placed in the sampler and the latter checked for leaks. For alternative personal samplers, connect the two components appropriately.

7.2 Sampling

7.2.1 Parameters

The method described in this document was validated with these parameters for a GGP sampler.

Sampling duration: minimum 2 h, up to 8 h

Flow rate: 3,5 l/min

Temperature: 24 °C to 28 °C (ambient)

NOTE 1 The atmospheric humidity does not exert an influence.

NOTE 2 The influence of higher or lower temperatures was not tested.