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## Animal and vegetable fats and oils — Determination of trace elements by inductively coupled plasma optical emission spectroscopy (ICP-OES)

Corps gras d'origines animale et végétale — Détermination des éléments traces dans les corps gras par spectrométrie d'émission optique à plasma induit par haute fréquence (ICP-OES)

ICS: 67.200.10

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## 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.

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

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

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ISO 21033 was prepared by Technical Committee ISO/TC 34, Food products, Subcommittee SC 11, Animal and vegetable fats and oils.

Tehn (standards and sandards an This second/third/... edition cancels and replaces the first/second/\_security (ISO/TS 21033:2011), [clause(s) / subclause(s) / table(s) / figure(s) / annex(es)] of which [has / have] been technically revised.

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# Animal and vegetable fats and oils — Determination of trace elements by inductively coupled plasma optical emission spectroscopy (ICP-OES)

## 1 Scope

This International Standard specifies an inductively coupled plasma optical emission spectroscopic method (ICP-OES) for the determination of the trace element content in oils. Depending on the dilution solvent used, most types of vegetable oils can be analysed (crude, degummed, refined, bleached, deodorized and hardened oils) and nearly all types of lecithins and phosphatides.

This procedure is only suitable when the elements are present in a solubilized form. Fine particles, such as bleaching earth, metal particles and rust, can result in poor recovery of the trace elements present as nebulization and atomization problems affect the ICP-OES analysis.

NOTE The only suitable non-ashing direct method for samples containing fine particles is graphite furnace atomic absorption spectrometry.

## 2 Normative references

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

ISO 661, Animal and vegetable fats and oils — Preparation of test sample

## 3 Terms and definitions

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

## 3.1

### trace element

element present in very low concentrations

Note 1 to entry: Trace element content is expressed as a mass fraction in milligrams per kilogram.

## 4 Principle

Solvent-diluted vegetable oils are analysed for trace elements by direct aspiration. Liquid samples are nebulized and carried into the excitation source by a flowing gas. Atoms are quantified by measuring the specific emission lines produced by atoms decaying from high energy levels.

## **5** Reagents

## WARNING — Attention is drawn to national regulations that specify the handling of hazardous substances, and users' obligations thereunder. Technical, organizational and personal safety measures shall be followed.

Unless stated otherwise, use reagents specified in ISO 6353-2<sup>[4]</sup> and ISO 6353-3,<sup>[5]</sup> if listed there, if not then use reagents of recognized analytical grade.

## **5.1 1-Butanol**, ISO 6353-3.<sup>[5]</sup>

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#### 5.2 Kerosene.

#### 5.3 **Xylene**, ISO 6353-3.<sup>[5]</sup>

5.4 **Standard elements**, present in solution as an organic soluble material<sup>1</sup>). Multi-element standards may be used.

Base oil [Base 20 Oil or Base 75 Oil from Accu-Standard<sup>1</sup>], may be used to check the blank oil 5.5 used and for the dilution of the standard solutions as needed.

5.6 Blank oil, typically, refined and bleached soya bean or other oil, which has been shown to be free of trace elements.

5.7 **Mixture of the blank oil** (5.6) with 1-butanol (5.1), mass fraction  $w_{\text{(blank oil)}} = 50 \text{ g}/100 \text{ g}$ .

#### **Apparatus** 6

Usual laboratory equipment, and, in particular, the following

- Inductively coupled plasma optical emission spectrometer. 6.1
- Analytical balance, readability 0,000 1 g, weighing precision 0,001 g 6.2
- **Oven**, capable of maintaining a temperature of  $(60 \pm 2)$ 6.3
- 6.4 Tilt table mixer.
- Volumetric flasks, capacity 100 ml. 6.5

#### Sampling 7

1.5 and ards, itel 102-41515 Sampling is not part of the method in this Technical Specification. A recommended sampling method is given in ISO 5555.[1]

It is important that the laboratory receive a truly representative sample which has not been damaged or changed during transport or storage.

#### **Preparation of test sample** 8

Prepare the test samples in accordance with ISO 661, except that the samples should not be clarified.

#### **Procedure** 9

## 9.1 General

**9.1.1** Melt solid samples at approximately 10 °C above their melting point and homogenize prior to dilution. Keep the diluted sample warm and monitor throughout the analysis to ensure it remains in solution. The maximum temperature for the analysis of hardened fats is 60 °C.

<sup>1)</sup> SPEX and Accu-Standard are examples of suitable products available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of these products.

**9.1.2** All samples, standards, and blanks are diluted (equal volume fractions) with 1-butanol (kerosene or xylene) to reduce the viscosity of the oil for better nebulization. Some samples are more soluble in 1-butanol than others.

1-Butanol is preferred because it has better moisture tolerance and allows a higher flow rate with higher pressure than kerosene without extinguishing the torch. The increased moisture tolerance permits the analysis of crude oils and lecithins without phase separation. The higher flow rate provides for improved detection limits.

If kerosene or xylene is used, all instrumental operating conditions, e.g. pump flow rate, change from those set for 1-butanol. Therefore, the analysis shall be standardized and all analyses shall be run with all standards, blank, and samples dissolved in the same solvent.

**9.1.3** The instrument is ignited and allowed to warm. Profile the instrument according to the manufacturer's instructions. Elements can be detected at the major emission lines (see <u>Table 1</u>). Additional emission lines and equipment set up instructions are given in EN 14538:2006<sup>[6]</sup> and ISO 10540-3:2012<sup>[Z]</sup>.

Flomont	Current limits of detection Major emission line(s)				
Element	mg/kg	>	nm		
Aluminium	_a	167,078	308,215		
Barium		455,404	70.		
Boron	attit	249,773			
Cadmium	a aros dato	226,502	214,441	228,802	
Calcium	0.05 stanting	315,887	393,366	317,933	
Chrome	all sta a full categor	267,716	284,325	283,563	
Copper	0,05 0,05	324,754	327,396		
Iron	0,05	259,940	238,204	261,187	
Lead	andarahora	220,353			
Magnesium	, 1/st 00,05	285,213	279,553	280,270	
Manganese	nthe of _a	257,611	259,373		
Molybdenum	a	281,615	202,095		
Nickel	0,05	231,604	221,648	341,476	
Phosphorus	0,05	213,618	178,287	177,495	
Silicon	0,1	251,611	288,158		
Silver	a	328,068			
Sodium	0,1	588,995			
Sulfur	1	180,731			
Tin	a	242,949			
Titanium	a	334,941	323,452	336,121	
Vanadium	a	309,311	311,071		
Zinc	a	202,613	213,856		
a Not reported at the time of publication					

## Table 1 — Major emission lines and limits of detection

**9.1.4** Standardize the instrument as specified in <u>9.3</u> and scan all the samples in triplicate.

NOTE Calibration drift has been noted. It can result from carbon build-up on the injector tip.

## 9.2 Preparation of standards

#### 9.2.1 Blank

Typically, refined and bleached sova bean or other oil, which has been shown to be free of trace elements. is used. Blank oil is diluted (1: 1) as described in the sampling procedure. Base 20 or Base 75 oil (5.5) is used as an absolute reference blank to determine that the blank oil is free of trace elements.

#### 9.2.2 **Standards**

The standard is prepared from commercially available single element organic-based standards. Weigh accurately the standard and add enough blank oil to total 50,00 g. Add 50,00 g of solvent (1-butanol, kerosene or xylene) to achieve a (1 + 1) dilution.

One standard concentration works; however, up to four multi-level, multi-element standards provide a better calibration for linearity and accuracy. Levels should include 2,5 mg/kg, 5 mg/kg, and 10 mg/kg standards depending on the range of values expected.

Instruments that are capable of simultaneous detection of multiple elements can achieve improved precision and accuracy by the inclusion of an internal standard in the analysis. If an internal standard is used, it should be incorporated as part of the dilution stop. Typically, the resultant dilution should contain 10 mg/kg yttrium or scandium. Thus, under the dilution sequence for sampling (see 9.1.1 and 9.4), the diluent should contain 20 mg/kg internal standard yielding 10 mg/kg yttrium or scandium in

Run the blank oil standard and diluted Base 20 or Base 75 oils at the specified wavelength for the element(s) of interest. Run the standard solutions (9.2.2) at the wavelength for the wavelength for the standard solutions (9.2.2) at the standard solutions (9.2.2) at

Blanks, samples, and standards are scanned in triplicate for trace element(s) and are averaged.

Standards and the blank are run every 10 samples or fewer and the instrument is re-standardized as needed. For accuracy, use a narrow range of standardization (0 mg/kg to 25 mg/kg of each element). Although the linearity is somewhat greater, test samples should be diluted to keep the trace element(s) content within the range of standardization.

## 9.4 Preparation of standards

Weigh  $2.5 \pm 0.02$  g of sample into an auto-sampler tube and dilute with 2.5 g of 1-butanol (kerosene or xylene) delivered from an automatic pipette. Cap the tube and invert 40 times to 50 times on a mixing table.

Dilute 0,2 g lecithins (up to 100 % acetone insoluble) to 5,0 g with blank soybean oil (dilution factor 25) and then to 10 g with 1- butanol (5.6), dilution factor 50. Mix the samples on a tilt table mixer for 1 h. Then dilute 1 part with 10 parts of the mixture of the blank oil and 1-butanol (5.7). The total dilution factor is 500.

## **10** Calculation

Computation is a feature of most instrument programs. Area counts from known standards are inserted into a linear regression formula versus the sample concentration. From this relationship, concentrations may be determined from the area counts of the samples.

Simultaneous instruments can usually accommodate internal standard calculations. If an internal standard is used, the ratio of the counts from the standard divided by the area counts of the internal standard are inserted into the linear regression formula.

It is important to include the correct dilution factor.

## **11 Precision**

## **11.1 Interlaboratory test**

Details of interlaboratory tests on the precision of the method are summarized in <u>Annex A</u>. The values derived from these interlaboratory tests may not be applicable to concentration ranges and matrices other than those given.

## **11.2 Repeatability**

The absolute difference between two independent single test results, obtained using the same method on identical test material in the same laboratory by the same operator using the same equipment within a short interval of time will not be greater than the repeatability limit *r* as shown in <u>Annex A</u>.

## **11.3 Reproducibility**

The absolute difference between two single test results, obtained using the same method on identical test material in different laboratories with different operators using different equipment, will not be greater than the reproducibility limit *R* as shown in <u>Annex A</u>.

## **12 Test report**

The test report shall contain at least the following information:

- a) all information necessary for the complete identification of the sample;
- b) the sampling method used, if known;
- c) the test method used, with reference to this International Standard (ISO 21033:2014);
- d) all operating details not specified in this Technical Specification, or regarded as optional, together with details of any incidents which may have influenced the test result(s);
- e) the test result(s) obtained;
- f) if repeatability has been checked, the final quoted result obtained.