

## SLOVENSKI STANDARD oSIST prEN ISO 21392:2020

01-junij-2020

#### Kozmetika - Analizne metode - Določevanje sledov težkih kovin v končnih kozmetičnih izdelkih z masno spektrometrijo z induktivno sklopljeno plazmo (ICP/MS) (ISO/DIS 21392:2020)

Cosmetics - Analytical Methods - Measurement of traces of heavy metals in cosmetic finished products using ICP/MS technique (ISO/DIS 21392:2020)

Kosmetische Mittel - Untersuchungsverfahren - Messung von Spuren von Schwermetallen in fertigen kosmetischen Mitteln mittels ICP-MS (ISO/DIS 21392:2020) (standards.iteh.ai)

Cosmétiques - Méthodes analytiques - Détermination de traces de métaux lourds dans les produits finis cosmétiques par ICP/MS (ISO/DIS 21392:2020)

90725be532eb/osist-pren-iso-21392-2020

Ta slovenski standard je istoveten z: prEN ISO 21392

ICS:

71.100.70 Kozmetika. Toaletni pripomočki

Cosmetics. Toiletries

oSIST prEN ISO 21392:2020

en,fr,de

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# DRAFT INTERNATIONAL STANDARD ISO/DIS 21392

ISO/TC 217

Voting begins on: **2020-04-29** 

Secretariat: ISIRI

Voting terminates on: 2020-07-22

## Cosmetics — Analytical methods — Measurement of traces of heavy metals in cosmetic finished products using ICP/ MS technique

Cosmétiques — Méthodes analytiques — Détermination de traces de métaux lourds dans les produits finis cosmétiques par ICP/MS

ICS: 71.100.70

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Published in Switzerland

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### Foreword

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This document was prepared by Technical Committee ISO/TC 217 Cosmetics.

Any feedback or questions on this document should be directed to the user's hational standards body. A complete listing of these bodies can be found at <u>www.iso.org/inlembers/html</u>.

### Introduction

This standard specifies an analytical procedure for determination of trace elements (Chromium, Cobalt, Nickel, Arsenic, Cadmium, Antimony and Lead) in finished cosmetics products by Inductively Coupled Plasma Mass Spectrometry (ICP-MS) after pressure digestion of the sample. This type of analytical procedure is widely described in other areas such as environment,<sup>[1][2][3]</sup> food<sup>[1][2][3]</sup> and pharmaceutical industry <sup>[Z][8][9][10]</sup>.

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## **Cosmetics** — Analytical methods — Measurement of traces of heavy metals in cosmetic finished products using ICP/ **MS technique**

### 1 Scope

The aim of this standard is to provide a method of quantification of heavy metal trace elements in cosmetic products that consumers might be exposed to in their usage. In the sample preparation procedure, nitric acid/hydrochloric acid mixture is used and most of the cosmetic ingredients are digested allowing heavy metal trace elements to be solubilized for measurement. Some cosmetic inorganic ingredients such as silica or titanium dioxide might not be completely digested under the conditions of this standard and heavy metal trace elements confined in such ingredients might not be fully extracted. However, the heavy metal trace elements trapped in these inorganic materials are considered less relevant for the evaluation of the exposure level of consumers to unwanted trace elements. The use of ICP-MS ensures reliable measurement of trace elements due to its proven high sensitivity and selectivity.

This analytical methodology can be applied to many other elements but this standard refers only to the above listed trace elements and it is to the responsibility of the analyst to prove that it fits for that purpose. In order to obtain comparable results, it is absolutely mandatory to comply with all the conditions linked to the digestion of the samples. (standards.iteh.ai)

#### 2 Normative references

oSIST prEN ISO 21392:2020

There is no normative reference in this document.sist/f4b4f6b3-7106-49bf-bfa0-2eb/osist-pren-iso-21392-2020

#### Principle 3

Trace elements in cosmetic products are quantified by ICP-MS measurement of the solutions resulting from digestion of the cosmetic products. Digestion takes place with mineral acids in sealed vessels heated to 200°C by microwaves, producing high pressures.

#### Reagents 4

The reagents and the water used shall be free of the elements to be determined to such an extent that the results are not impaired. Unless specified otherwise, solutions are understood to be aqueous solutions.

Ultrapure water, conductivity below 0,1µS/cm-1 at 25°C according to Type 1 water specifications 4.1 defined in ISO 3696 standard<sup>[11]</sup>

4.2 Nitric acid, minimum w = 60%, density = 1,38 g/ml

**4.3 Hydrochloric acid**, minimum *w* = 30 %, density = 1,15 g/ml

#### Internal standard stock solutions 4.4

For storage and stability conditions of the internal standard stock solutions, follow the specifications of the suppliers.

#### 4.4.1 Rhodium stock solution, 1 000 mg/l

#### 4.4.2 Lutetium stock solution, 1 000 mg/l

# **4.5** Analytes stock solutions (Chromium, Cobalt, Nickel, Arsenic, Cadmium, Antimony and Lead), 1 000 mg/l for each element

Commercially available single element or mixed stock solutions can be used. For this 2 cases, the used stock solutions shall not contain other elements that could interfere with the analytes to be quantified.

**4.6** Commercially available ICP-MS Tune solution, containing e.g. Ce, Co, Li, Mg, Tl and Y ( $1 \mu g/l$ ) according to instrument manufacturer's recommendations.

#### 5 Apparatus and equipment

All apparatus and equipment that come into direct contact with sample or solutions shall be pre-cleaned with dilute nitric acid (6.1) and rinsed with ultrapure water (4.1) to ensure the lowest analyte background. To prevent contamination and adsorption, do not use lab materials made with borosilicate glass.

#### 5.1 Digestion vessels

Use commercially available, safety-tested pressure vessels and inserts made of acid-resistant, lowcontamination materials. The assembled vessels shall be able to safely withstand temperatures up to at least 200°C and pressures up to at least 40 bar.

For determination of antimony, use only digestion containers with minimal surface roughness to prevent its adhesion to the container surface <sup>(6)[12],</sup>. Quartz containers are recommended because they are usually more resistant to attrition. However, Teflon vessels without any scratch or damage on their inner surface or any deposits are appropriate. Scratched or etched containers should not be used. If there is any question regarding possible adsorption of antimony on vessels' walls, test for surface adsorption as described in section 7.3.3.

NOTE Dedicated digestion vessels are recommended for the digestion of cosmetic samples, which may have high levels of elements to be determined. To avoid memory effects, perform a blank digestion to clean vessels after digesting highly loaded samples, before digesting subsequent samples.

#### 5.2 Microwave-assisted digestion instruments

Microwave-heated systems shall be equipped with a temperature measurement unit, which simultaneously regulates the power control of the microwave. Reliable temperature measurement is obtained e.g. through measurement sensors inserted into the pressure vessel. Only use microwave-assisted digestion instruments equipped with temperature sensors and calibrate the temperature sensor before use.

#### 5.3 Membrane filter, 0,45 µm pore size

The membrane filter used shall be inert with regard to the acid concentration of the measurement solution and shall not bring any contamination into the measurement solution or adsorption of the analytes. Several types of membrane material are commercially available (PTFE, PP...) and their fit for purpose must be verified by means of appropriate measurements (blanks, QC samples...).

#### 5.4 **ICP-MS**

Mass spectrometer with inductively coupled argon plasma is composed of a sample introduction and an atomisation system, as well as an instrument control and evaluation unit. To prevent interferences with the masses of the elements of chromium, nickel, arsenic and cadmium, use of a mass spectrometer

that is capable of compensating or minimising such interferences (e.g. collision and/or reaction cell, resolution above 3000, alternatively corrective equations for higher concentrations).

#### 6 Preparation of standards solutions

For all the solutions, the terminology "part" in the standard refers to either volume or weight. That means that standards and samples can be diluted by volume or weight. However, it should be consistent for both standards and samples.

**6.1 Diluted nitric acid**, produced by mixing nitric acid (<u>4.2</u>) with pure water (<u>4.1</u>) at a ratio of approximately 1+9 parts respectively.

#### 6.2 Diluting solution

The composition of the diluting solution must have the same acid composition (total content and acid ratio) as the Analytical Solution (the diluted digest solution). This solution should contain:

- 2,5 part of nitric acid (<u>4.2</u>),
- 0,5 part of hydrochloric acid (<u>4.3</u>),
- 97 parts of water (<u>4.1</u>)

### 6.3 Internal Standard solutions. NDARD PREVIEW

The internal standards (IS) selected should cover all the mass range of the considered analytes and have similar ionisation energy to the trace element for which it is used for correction purposes. It shall also be checked that the native concentration of the internal standards in the sample to be analysed is negligible and that they are not interfered by sample constituents.

"Rhodium and Lutetium have proved to be well suited as internal standards (IS). Samples should be checked for negligible native concentrations of the IS and that the IS are not interfered with by sample constituents.

Rhodium is suitable for determination of Chromium, Cobalt, Nickel, Arsenic, Cadmium and Antimony, whereas Lutetium is suitable for determination of Lead. Alternatively, other elements may be used (for example Indium or Iridium). Scandium, however, is not suitable as an IS due to Calcium interferences. An IS with a mass (m/z) below 100 is not recommended because it may suffer from interferences from matrix components.

NOTES : - Internal standard solutions may be added in each sample and calibration solution at the same concentration or may be added through an on line Y-fitting to a pump tube.

- the concentration of the internal standard solutions must be included in the range 1 to 2 mg/l. In the following sections, a concentration of 1 mg/l has been used for all the calculations.

#### 6.3.1 Rhodium(\*) standard solution, 1 mg/l

Dilute the Rhodium stock solution (4.4.1) 1 + 999 with diluting solution (6.2). This internal standard solution is stable at room temperature for 6 months.

#### 6.3.2 Lutetium(\*) standard solution, 1 mg/l

Dilute the Lutetium stock solution (4.4.2) 1 + 999 with diluting solution (6.2). This internal standard solution is stable at room temperature for 6 months.

(\*) NOTE : Indium or Iridium may also be used as internal standards

#### 6.4 Standard solutions

The concentrations of these standard solutions are examples and shall be adjusted to the specific conditions in the laboratories.

#### 6.4.1 High concentration mixed standard solution, 10 mg/l

Dilute 100 times the analyte stock solution(s) (3.5) by adding:

- in the case of single analyte stock solutions, 1 part of each of these 7 solutions to 93 parts of the diluting solution (6.2)
- in the case of mixed stock solution, add 1 part of this solution to 99 parts of the diluting solution (6.2).

This high standard solution is stable at room temperature for 6 months.

#### $\textbf{6.4.2} \quad \textbf{Low concentration mixed standard solution, } 0.1 \ \mathrm{mg/l}$

Dilute 100 times the high concentration standard solution (5.4.1) by adding 1 part of this solution to 99 parts of the diluting solution ( $\underline{6.2}$ ). This low standard solution is stable at room temperature for 3 months.

#### 6.5 Calibration blank solution

The calibration blank solution corresponds to the matrix solution without any analyte of interest. Generally, it corresponds to the diluting solution with the suitable concentration of the appropriate internal standards if not added via a Y-fitting during the measurement.

#### 6.6 Calibration solutions

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Mixed calibration solutions are prepared by diluting the low concentration mixed standard solution (6.4.2) with the diluting solution (6.2) to levels in the linear range of the instrument and within the targeted concentration range. Include a suitable concentration of the appropriate internal standards, or add on line the internal standards by means of pumping into the sample flow through a Y-fitting. At least 3 calibration solutions with various concentrations should be prepared. These calibration solutions must be prepared daily.

Examples of preparation procedure of calibration solutions are detailed in <u>Table 1a</u> (with addition of the internal standards in all the calibration solutions) and <u>Table 1b</u> (with on line addition of the internal standards via an Y-fitting).

# Table 1a — Example of calibration solutions of the ICP-MS – addition of the internal standards in every calibration solution

Calibration solution	Part of low conc. mixed standard solution ( <u>6.4.2</u> )	Part of Rhodi- um standard solution ( <u>6.3.1</u> )	Part of Luteti- um standard solution ( <u>6.3.2</u> )	Part of the diluting solu- tion ( <u>6.2</u> )	Analyte conc. in the calibra- tion solution (μg/l)
5.5.0 Calibration blank	0	2	2	496	0
5.5.1 Calibration solution 1	2,5	2	2	493,5	0,5
5.5.2 Calibration solution 2	5	2	2	491	1
5.5.3 Calibration solution 3	10	2	2	486	2
5.5.4 Calibration solution 4	25	2	2	471	5
5.5.5 Calibration solution 5	50	2	2	446	10