# INTERNATIONAL STANDARD

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# Jewellery and precious metals — Determination of high purity gold, platinum and palladium — Difference method using ICP-OES

Joaillerie, bijouterie et métaux précieux — Dosage de l'or, du platine et du palladium à haute pureté — Méthode par différence utilisant **iTeh ST**<sup>'</sup>**CROESARD PREVIEW** 

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<u>ISO 15093:2020</u> https://standards.iteh.ai/catalog/standards/sist/ab501fa5-2a55-471c-bb23-2b32791d1f92/iso-15093-2020



Reference number ISO 15093:2020(E)

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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.

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 <a href="https://www.iso.org/directives">www.iso.org/directives</a>).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see <a href="https://www.iso.org/patents">www.iso.org/patents</a>).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see <a href="https://www.iso.org/iso/foreword.html">www.iso.org/iso/foreword.html</a>.

This document was prepared by Technical Committee ISO/TC 174, *Jewellery and precious metals*.

This third edition cancels and replaces the second edition (ISO 15093:2015), which has been technically revised. The main changes compared to the previous edition are as follows:

- a) change of title of standard;
- b) change of scope for measuring also gold, platinum and palladium with a nominal content above 999  $\%_0$  (parts per thousands);
- c) revision of this document in order to comply in structure with ISO 15096.

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

## Jewellery and precious metals — Determination of high purity gold, platinum and palladium — Difference method using ICP-OES

## **1** Scope

This document specifies the analytical procedure for the determination of gold, platinum and palladium with a nominal content of and above 999 % (parts per thousand).

This document specifies a method intended to be used as the recommended method for the determination of gold, platinum and palladium of fineness of and above 999 ‰. For the determination of fineness of and above 999,9 ‰, modifications described in <u>Annex B</u> apply.

#### Normative references 2

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 11596, Jewellery – Sampling of precious metal alloys for and in jewellery and associated products

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#### **Terms and definitions** 3

ISO 15093:2020 No terms and definitions are listed in this document. /ab501fa5-2a55-471c-bb23-

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

- ISO Online browsing platform: available at <a href="http://www.iso.org/obp">http://www.iso.org/obp</a>
- IEC Electropedia: available at <u>http://www.electropedia.org/</u>

#### Principle 4

The sample is weighed and dissolved in aqua regia to prepare a 10 g/l solution (higher concentration is used for fineness of and above 999,9 ‰). The impurities are determined by ICP-OES (see Table A.1 for wavelengths), and the precious metal content is obtained by subtraction of the total content of impurities in the sample from 1 000 ‰. For the determination of fineness of and above 999,9 ‰, modifications described in Annex B shall be applied.

#### 5 Reagents

During the analysis, unless otherwise stated, use only reagents of recognized analytical grade and only distilled water or water of equivalent purity.

5.1 Hydrochloric acid (HCl), 30 % to 37 % HCl (mass fraction).

5.2 Nitric acid  $(HNO_3)$ , 65 % to 70 %  $HNO_3$  (mass fraction).

Aqua regia (should be prepared just before use). Mix three volumes of hydrochloric acid (5.1) and 5.3 one volume of nitric acid (5.2).

**5.4 Stock solution A** (shall not contain any chloride), Ag, Al, Cd, Cr, Cu, Fe, Mg, Mn (100 mg/l each) in 3 % to 7 % HNO<sub>3</sub> (5.2) (mass fraction).

NOTE The stock solution "A" is typically prepared by mixing 10 % (of the total volume) of each 1 000 mg/l monoelemental solution with 5 % of nitric acid (5.2) (volume fraction) and making up with water. It can be kept for up to 12 months under proper storage conditions.

**5.5** Stock solution B (may contain both chlorides and nitrates), Bi, Co, Ga, In, Se, Sn, Te, Ti, Zn (100 mg/l each) in 3% to 7% aqua regia (5.3) (volume fraction).

**5.6** Stock solution C (may contain both chlorides and nitrates), As, Ni, Pb, Sb, Si, Tl, W, Zr (100 mg/l each) in 3 % to 7 % aqua regia (5.3) (volume fraction).

**5.7 Stock solution D** (may contain both chlorides and nitrates), Au, Ir, Pd, Pt, Rh, Ru (100 mg/l each) in 3 % to 7 % aqua regia (5.3) (volume fraction).

NOTE 1 Elements which do not need to be analysed can be omitted. Other elements can be added, provided they are stable and do not generate significant interferences.

NOTE 2 The stock solutions "B" to "D" are typically prepared by mixing 10 % (of the total volume) of each 1 000 mg/l monoelemental solution with 5 % of aqua regia (5.3) (volume fraction) and making up with water. They can be kept for up to 12 months under proper storage conditions.

**5.8 Reference materials: gold, platinum or palladium**, of 999,9 ‰ minimum purity. The content of each impurity shall be specified and taken into account in the calibration.

## 6 Apparatus

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#### ISO 15093:2020

6.1 Customary laboratory, apparatusch.ai/catalog/standards/sist/ab501fa5-2a55-471c-bb23-

2b32791d1f92/iso-15093-2020

**6.2 ICP-OES**, with a minimum optical resolution of 0,02 nm, a detection limit of 0,02 mg/l or better, and capability of background correction.

NOTE For preferably used wavelengths, see <u>Annex A</u>.

**6.3 Analytical balance**, with a reading accuracy of 0,1 mg.

## 7 Sampling

The sampling procedure shall be performed in accordance with ISO 11596.

## 8 Procedure

#### 8.1 General

#### WARNING — Suitable health and safety procedures should be followed.

All vessels and reagent bottles shall be teflon (e.g. PFA, FEP) or plastic.

#### 8.2 Sample solution

For each sample to be analysed, two sample solutions shall be prepared as follows and analysed.

The sample should be etched, washed and dried before use to remove any surface contamination.

Weigh  $(500 \pm 10)$  mg of the sample portion to the nearest 0,1 mg, transfer into a 50 ml beaker, and add 20 ml of aqua regia (5.3). Heat at an appropriate temperature until complete dissolution of the sample and continue to heat to expel the nitrogen oxides. Allow to cool, transfer into a 50 ml volumetric flask, make up with water to 50 ml, and mix thoroughly. Dissolution may alternatively be performed directly in chemical and temperature resistant volumetric flasks.

If an insoluble material is observed, dissolution under pressure should be performed.

#### 8.3 Calibration solutions

Weigh three portions of  $(500 \pm 10)$  mg of the reference material (5.8), and dissolve, cool and transfer into 50 ml volumetric flasks each one as specified in 8.2.

#### 8.3.1 Blank solution.

Make up with water to 50 ml and mix thoroughly.

#### 8.3.2 Calibration solution 1.

Add 5 ml of stock solution A (5.4) and 5 ml of stock solution B (5.5), make up with water to 50 ml, and mix thoroughly. This solution is unstable over time and should be prepared just before use, or its stability validated.

**8.3.3 Calibration solution 2.** Add 5 ml of stock solution C (5.6) and 5 ml of stock solution D (5.7), make up with water to 50 ml, and mix thoroughly. This solution is unstable over time and should be prepared just before use, or its stability validated.

The volumes of stock solutions shall be lowered to match the concentration of impurities in the sample. It is possible to use a multiple-point calibration by adding calibration solutions with other concentrations.

#### 8.4 Measurement

Set up the instrument in accordance with the manufacturer's instructions and choose appropriate background correction positions. A clean torch, spray chamber, and sample uptake tubes shall be used and the plasma shall be stabilized before use, following the recommendations of the instruments manufacturer.

Spray the blank and calibration solutions 1 (8.3.2) and 2 (8.3.3) in accordance with the defined instrument calibration procedure and then run the analytical procedure for the sample solutions. The result shall be displayed with enough decimal places to provide an accurate indication of concentrations at the detection limits of the relevant elements.

Each solution shall have a stabilization time of at least 30 s, followed by at least three integrations each for the determination of the net intensities (background-corrected).

The rinsing time between each measurement shall be sufficient to allow the signal of each impurity to come back to the baseline.

#### 9 **Calculation and expression of the results**

#### 9.1 Calibration curves

Set the concentration in the blank solution and the calibration solution, taking into account impurities introduced in the solution by the reference material (5.8) and calculate the calibration curve for each element using the net intensities obtained for the blank solution and the calibration solution.

#### 9.2 Calculation

By means of the calibration curves (see 9.1), convert the net intensity values into concentration values and use Formula (1) to calculate the mass portion of each relevant element ( $W_i$ ).

$$W_i = \frac{c_i \times V_{\rm s}}{m_{\rm s}} \tag{1}$$

where

- $c_i$  is the concentration of element, *i*, in the sample solution, in mg/l;
- $V_{\rm s}$  is the volume of the sample solution, in litres;
- $m_{\rm s}$  is the mass of the metallic sample, in milligrams.

The detection limit is defined as three standard deviations of the concentration of each individual element measured in the blank solution.

The precious metal fineness ( $W_{sp}$ ), expressed in parts per thousand (‰), is thus calculated using Formula (2):

$$W_{\rm sp} = 1\,000 - (\Sigma W_i \times 1\,000)$$
 (2)

where  $\sum W_i$  is the sum of the mass portion of each element found above its detection limit.

#### 9.3 Repeatability

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Duplicate determinations shall give results differing by less than 10 % for the sum of impurities. If the difference is greater than this, the assay shall be repeated.

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## **10 Test report**

The test report shall include at least the following information:

- a) identification of the sample including source, date of receipt, and form of sample;
- b) sampling procedure;
- c) method used by reference to this document, i.e. ISO 15093:2020;
- d) precious metals content of the sample, in parts per thousand (‰) by mass, as single values and mean values, with the result reported with four significant figures;
- e) if relevant, any deviations from the method specified in this document;
- f) any unusual features observed during the determination;
- g) date of test;
- h) identification of the laboratory carrying out the test;
- i) signature of the laboratory manager and operator.

# Annex A (informative)

## Wavelengths

## A.1 General

Other wavelengths than those specified in <u>Tables A.1</u>, <u>A.2</u> and <u>A.3</u> can be used. In all cases, attention shall be paid to spectral interferences. Those tables are not exhaustive. If other elements are present, they shall be analysed.

Element	Recommended	Alternative	Alternative
Element	nm	nm	nm
Ag	328,068	338,289	
Al	167,08	396,152	_
As	189,04	193,76	
Au <sup>a</sup> S	A 389,786 RI	302,920	$\mathbf{C}\mathbf{W}$ –
Bi	190,24	223,061	
Cd (S	tar <sub>214,438</sub> 05.	ITC 226,502	228,802
Со	228,616	238,892	—
Cr https://standards.iteh	205,618	020 359,349	360,533
Cu	2b32 <b>32</b> 7 <b>3</b> 96/iso-1	5093- <b>384</b> 0754	
Fe	238,204	259,941	261,187
Ga	287,424	141,44	417,206
In	303,936	325,609	158,636
Ir	215,268	183,246	
Mg	279,553	280,270	—
Mn	259,373	294,921	257,611
Ni	352,454	221,648	231,604
Pb	168,22	220,353	283,305
Pd	340,458	360,955	
Pt	177,71	214,423	203,646
Rh	343,489	233,437	—
Ru	240,272	_	
Sb	206,833	217,581	—
Se	196,09	204,050	—
Si	251,612	288,158	
Sn	189,99	147,52	
Те	214,281	225,902	—
Ti	336,121	334,941	—
Tl	190,86	—	—
<sup>a</sup> Suggested mat	trix line (only used	to ensure sample	has been properly

#### Table A.1 — Wavelenghts for measuring of impurities in gold

<sup>a</sup> Suggested matrix line (only used to ensure sample has been properly prepared and analysed, shall not be included in the calculation as described in <u>9.2</u>).