
**Nickel alloys — Determination of
molybdenum content — Inductively
coupled plasma/atomic emission
spectrometric method**

*Alliages de nickel — Détermination de la teneur en molybdène —
Méthode par spectrométrie d'émission atomique à plasma induit par
haute fréquence*



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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.

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.

ISO 11435 was prepared by Technical Committee ISO/TC 155, *Nickel and nickel alloys*, Subcommittee SC 3, *Analysis of nickel, ferronickel and nickel alloys*.

This second edition cancels and replaces the first edition (ISO 11435:2005), of which it constitutes a minor revision.

Nickel alloys — Determination of molybdenum content — Inductively coupled plasma/atomic emission spectrometric method

1 Scope

This International Standard specifies an inductively coupled plasma/atomic emission spectrometric method for the determination of the mass fraction of molybdenum between 0,05 % and 20 % in nickel alloys.

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 648:2008, *Laboratory glassware — Single-volume pipettes*

ISO 1042:1998, *Laboratory glassware — One-mark volumetric flasks*

ISO 3696:1987, *Water for analytical laboratory use — Specification and test methods*

ISO 5725-1:1994, *Accuracy (trueness and precision) of measurement methods and results — Part 1: General principles and definitions*

ISO 5725-2:1994, *Accuracy (trueness and precision) of measurement methods and results — Part 2: Basic method for the determination of repeatability and reproducibility of a standard measurement method*

ISO 5725-3:1994, *Accuracy (trueness and precision) of measurement methods and results — Part 3: Intermediate measures of the precision of a standard measurement method*

3 Principle

Dissolution of a test portion in a mixture of hydrochloric, nitric and phosphoric acid and fuming with a mixture of phosphoric and perchloric acids. Addition of hydrofluoric acid and, if desired, of an internal reference element and dilution of the solution to known volume. Nebulization of the solution into an inductively coupled plasma/atomic emission spectrometer and measurement of the intensity of the emitted light from molybdenum, and eventually from the internal reference element, simultaneously.

Examples of the analytical lines for molybdenum are given in Table 1.

Table 1 — Examples of analytical lines together with interfering elements

Element	Analytical line nm	Interferences
Molybdenum	202,03	Ta
	281,61	Al, Hf

The method uses a calibration based on a very close matrix-matching of the calibration solutions to the sample and bracketing of the contents between 0,75 and 1,25 of the approximate concentration of molybdenum in the sample to be analysed. The concentration of all elements in the sample has, therefore, to be approximately known. If the concentrations are not known, the sample has to be analysed by some semi-quantitative method. The advantage with this procedure is that all possible interferences from the matrix will be automatically compensated, which will result in high accuracy. This is most important for spectral interferences, which can be severe in very highly alloyed metals. All possible interferences shall be kept at a minimum level. Therefore, it is essential that the spectrometer used meets the performance criteria specified in the method for the selected analytical lines.

Two lines have been carefully investigated (see Annex B). The strongest possible interferences are given in Table 1. If other lines are used, they shall be carefully checked so that interferences are not higher than the values given in Annex B. The analytical line for the internal standard should be selected carefully. It is recommended to use scandium 363,07 nm. This line is interference-free for the elements and concentrations generally found in nickel alloys.

NOTE The use of an internal standard is not essential since no relevant differences between laboratories operating with or without internal standards were found.

4 Reagents

During the analysis, unless otherwise stated, use only reagents of recognized analytical grade and only grade 2 water as specified in ISO 3696:1987.

4.1 Hydrofluoric acid, HF, 40 % (mass fraction), $\rho = 1,14$ g/ml, or 50 % (mass fraction), $\rho = 1,17$ g/ml.

WARNING — Hydrofluoric acid is extremely irritating and corrosive to skin and mucous membranes, producing severe skin burns which are slow to heal. In the case of contact with skin, wash well with water, apply a topical gel containing 2,5 % (mass fraction) calcium gluconate, and seek immediate medical treatment.

4.2 Hydrochloric acid, HCl, $\rho = 1,19$ g/ml.

4.3 Nitric acid, HNO₃, $\rho = 1,40$ g/ml.

4.4 Phosphoric acid, H₃PO₄, $\rho = 1,70$ g/ml.

4.5 Perchloric acid, HClO₄, 60 % (mass fraction), $\rho = 1,54$ g/ml or 70 %, $\rho = 1,67$ g/ml.

4.6 Internal standard solution, 100 mg/l.

Choose a suitable element to be added as internal reference and prepare a 100 mg/l solution.

4.7 Molybdenum standard solution, 10 g/l.

Weigh, to the nearest 0,000 5 g, 1 g of high-purity molybdenum [min 99,95 % (mass fraction)], place it in a 250 ml beaker and dissolve in a mixture of 10 ml hydrochloric acid (4.2) and 10 ml nitric acid (4.3).

Cool and transfer to a calibrated 100 ml one-mark volumetric flask. Dilute to the mark with water and mix.

NOTE 1 ml of this solution contains 10 mg of molybdenum.

4.8 Molybdenum standard solution, 1 g/l.

Using a calibrated pipette (or burette), transfer 10 ml of the molybdenum standard solution (4.7) into a calibrated 100 ml one-mark volumetric flask. Add 10 ml of hydrochloric acid (4.2). Dilute to the mark with water and mix.

NOTE 1 ml of this solution contains 1 mg of molybdenum.

4.9 Molybdenum standard solution, 100 mg/l.

Using a calibrated pipette (or burette), transfer 10 ml of the molybdenum standard solution (4.8) into a calibrated 100 ml one-mark volumetric flask. Add 10 ml of hydrochloric acid (4.2). Dilute to the mark with water and mix.

NOTE 1 ml of this solution contains 0,1 mg of molybdenum.

4.10 Standard solutions of interfering elements.

Prepare standard solutions for each element whose mass fraction is higher than 1 % of that contained in the test sample. Use pure metal or chemical substances with mass fractions of molybdenum less than 10 µg/g.

5 Apparatus

All volumetric glassware shall be class A and calibrated in accordance with ISO 648:2008 or ISO 1042:1998, as appropriate.

Ordinary laboratory apparatus and the following.

5.1 Polytetrafluoroethylene (PTFE) beakers.**5.2 Polypropylene volumetric flasks**, of capacity 100 ml, calibrated according to ISO 1042.**5.3 Atomic emission spectrometer (AES).**

The spectrometer shall be equipped with an inductively coupled plasma (ICP) and a nebulization system resistant to hydrofluoric acid. The ICP/AES used will be satisfactory if, after optimizing according to 7.3, it meets the following performance criteria.

The spectrometer can be either the simultaneous or the sequential type. If a sequential spectrometer can be equipped with an extra arrangement for simultaneous measurement of the internal standard line, it can be used with the internal reference technique. If the sequential spectrometer is not equipped with this arrangement, an internal reference cannot be used and an alternative technique without an internal standard should be used.

5.3.1 Practical resolution of the sequential spectrometer

Calculate the bandwidth (full width at half maximum), according to A.2, for the analytical line used, including the line for internal reference. The bandwidth shall be less than 0,030 nm.

5.3.2 Short-term stability

Calculate the standard deviation of ten measurements of the absolute intensity or intensity ratio of the emitted light of the most concentrated calibration solution for molybdenum according to A.3. The relative standard deviation shall not exceed 0,4 %.

5.3.3 Background equivalent concentration

Calculate the background equivalent concentration (BEC) according to A.4, for the analytical line, using a solution containing only the analyte element. The maximum values of BEC obtained should be 0,4 mg/l.

6 Sampling and sample preparation

6.1 Sampling and preparation of the laboratory sample shall be carried out by normal agreed procedures or, in case of dispute, by the relevant International Standard.

6.2 The laboratory sample is normally in the form of millings or drillings and no further mechanical preparation is necessary.

6.3 The laboratory sample shall be cleaned by washing with pure acetone and drying in air.

6.4 If brazed alloy tools are used in the preparation of the laboratory sample then the sample shall be further cleaned by pickling in 15 % (mass fraction) nitric acid for a few minutes. It shall then be washed several times with distilled water, followed by washing in acetone and drying in air.

7 Procedure

7.1 Test portion

Weigh, to the nearest 0,000 5 g, 0,25 g of the test sample.

7.2 Preparation of test solution T_{Mo}

7.2.1 Place the test portion in a 200 ml Pyrex Erlenmeyer flask.

NOTE A PTFE or PFA beaker with a graphite base and a cover can also be used. In this case, the solution does not need to be transferred to another PTFE beaker after fuming (7.2.2).

7.2.2 Add 30 ml of HCl (4.2), 3 ml of HNO₃ (4.3) and 2,5 ml of H₃PO₄ (4.4). Let the dissolution begin at room temperature. If necessary, heat to complete dissolution. Add 7,5 ml of HClO₄ (4.5) and heat until the perchloric acid starts to fume. Continue to fume for 2 to 3 min (the white smoke shall be on the top of the Erlenmeyer flask).

In the case where a complete dissolution is difficult to obtain, 5 ml of HF (4.1) may be added, together with the HCl (4.2), HNO₃ (4.3) and H₃PO₄ (4.4). In this case, a PTFE or PFA beaker should be used.

7.2.3 Cool the solution and add 10 ml of water to dissolve the salts. Some residues can remain undissolved. Transfer the solution and possible undissolved residues quantitatively to a 100 ml PTFE beaker. Add 2 ml of HF (4.1). Heat gently for 20 min until the residues dissolve completely.

7.2.4 Cool the solution to room temperature and transfer the solution quantitatively to a 100 ml volumetric polypropylene flask (5.2). If an internal standard is used, add an adapted quantity with a calibrated pipette (or burette), for example 10 ml of the internal standard solution (4.6).

7.2.5 Dilute to the mark with water and mix.

7.3 Optimization of spectrometer

7.3.1 Start the ICP/AES and let it run for at least 30 min before taking any measurements.

7.3.2 Optimize the instrument according to the manufacturer's instructions.

7.3.3 Prepare the software to measure the intensity, mean value and relative standard deviation of the analytical lines.

7.3.4 If an internal standard is used, prepare the software to calculate the ratio between analyte intensity and internal standard intensity. The intensity of the internal standard shall be measured simultaneously with the analyte intensity.

7.3.5 Check the instrument performance requirements given in 5.3.1 to 5.3.3.

7.4 Predetermination of the test solution

Prepare a calibration solution K_{20} , corresponding to a molybdenum mass fraction of 20 % and matrix-matched to the test sample solution as follows.

7.4.1 Using a calibrated pipette (or burette), add 5 ml of the molybdenum standard solution (4.7) to a 100 ml volumetric polypropylene flask (5.2) marked K_{20} .

7.4.2 To this volumetric flask K_{20} , add volumes of standard solutions (4.10) necessary to match the sample matrix to be tested, for each element whose mass fraction is above 1 %.

The matrix should be matched to the nearest percent.

7.4.3 Add 2,5 ml of H_3PO_4 (4.4), 7,5 ml of $HClO_4$ (4.5) and 10 ml of the internal standard solution (4.6). Dilute with water and mix.

7.4.4 Also prepare a blank calibration solution, K_0 , prepared in the same way as the calibration solution K_{20} omitting molybdenum.

7.4.5 Measure the absolute intensities I_0 and I_{20} for the solutions K_0 and K_{20} .

7.4.6 Measure the absolute intensity $I_{T_{Mo}}$ for the test solution T_{Mo} .

7.4.7 Calculate the approximate mass fraction of molybdenum, $w_{T_{Mo}}$, in percentage, in the test solution by means of the following equation:

$$w_{T_{Mo}} = \frac{I_{T_{Mo}} (K_{20} - K_0)}{I_{20} - I_0}$$

7.5 Preparation of calibration solutions for bracketing, $K_{l,Mo}$ and $K_{h,Mo}$

For each test solution T_{Mo} , prepare two matrix-matched calibration solutions, $K_{l,Mo}$ and $K_{h,Mo}$ with molybdenum concentrations in $K_{l,Mo}$ slightly below, and in $K_{h,Mo}$ slightly above, the concentration in the unknown test solution as follows

7.5.1 Using calibrated pipettes (or burettes), add molybdenum standard solution (4.8 or 4.9) to one 200 ml Pyrex Erlenmeyer marked $K_{l,Mo}$ so that the mass fraction of molybdenum $w_{l,Mo}$, in percentage, is approximately $w_{T_{Mo}} \times 0,75 < w_{l,Mo} < w_{T_{Mo}} \times 0,95$. If a pipette is used, select $w_{l,Mo}$ in such a way as to take a volume easily.

7.5.2 Using calibrated pipettes (or burettes), add molybdenum standard solution (4.8 or 4.9) to one 200 ml Pyrex Erlenmeyer marked $K_{h,Mo}$ so that the mass fraction of molybdenum $w_{h,Mo}$ in percentage, is approximately $w_{T_{Mo}} \times 1,05 < w_{h,Mo} < w_{T_{Mo}} \times 1,25$. If a pipette is used, select $w_{h,Mo}$ in such a way as to take a volume easily.

7.5.3 Add to the calibration solutions $K_{l,Mo}$ and $K_{h,Mo}$ all matrix elements whose mass fractions are above 1 % in the test sample, using the appropriate amount of standard solutions (4.10) to match the equivalent matrix concentration to the nearest percent.

7.5.4 Proceed as directed in 7.2.2 to 7.2.5.

7.6 Measurement of test solutions

Firstly, measure the absolute or ratioed intensity of the analytical line of the lowest calibration solution $K_{l,Mo}$, then test sample solution, T_{Mo} , and finally the highest calibration solution $K_{h,Mo}$. Repeat this sequence three times and calculate the mean intensities $I_{l,Mo}$ and $I_{h,Mo}$ for the low and high calibration solution and $I_{T_{Mo}}$ for the test solution respectively.

8 Expression of results

8.1 Method of calculation

Calculate the mass fraction of molybdenum, w_{Mo} , in percentage, in the test solution T_{Mo} , by means of the equation:

$$w_{\text{Mo}} = w_{\text{I,Mo}} + \frac{(I_{T_{\text{Mo}}} - I_{\text{I,Mo}})(w_{\text{h,Mo}} - w_{\text{I,Mo}})}{I_{\text{h,Mo}} - I_{\text{I,Mo}}}$$

8.2 Precision

8.2.1 Laboratory tests

Twelve laboratories in six countries participated in an inter-laboratory test programme under the auspices of ISO/TC 155/SC 3/WG 8, involving three determinations of molybdenum at nine levels. Each laboratory did two determinations under repeatability conditions as defined in ISO 5725-1, i.e. one operator, same apparatus, identical operating conditions, same calibration and a minimum period of time. The third determination was done on a different day using the same apparatus with a different calibration.

8.2.2 Wavelength for measurements

Concerning the wavelength taken for measurements which have been statistically evaluated, only one laboratory operated at 281,62 nm, whereas the other eleven operated at 202,03 nm. No relevant difference between laboratories operating with or without an internal standard was found.

8.2.3 Statistical analysis

Statistical analysis was done in accordance with ISO 5725-1, ISO 5725-2 and ISO 5725-3. Results from one laboratory were discarded as the laboratory did not operate under the repeatability conditions specified and the results from a second laboratory were rejected as consistent outliers.

Results of the evaluation were used to calculate the smoothed values for repeatability, r , and within-laboratory reproducibility, R_w , and between-laboratory reproducibility, R , that are given in Table 2.

The smoothed values are approximately equivalent to the results reported in the precision study for the determination of molybdenum in steels by ICP.

Table 2 — Repeatability and reproducibility limits

Molybdenum mass fraction %	Repeatability limit r	Within-laboratory reproducibility limit R_w	Between-laboratory reproducibility limit R
0,05	0,002 3	0,004 1	0,006 2
0,1	0,003 9	0,006 6	0,010 5
0,2	0,006 6	0,010 8	0,017 8
0,5	0,013 1	0,020 4	0,035 7
1,0	0,022 3	0,033 0	0,060 4
2,0	0,037 6	0,053 6	0,102 1
5,0	0,075 1	0,101 5	0,204 5
10,0	0,126 8	0,164 7	0,345 9
20,0	0,214 0	0,267 0	0,584 9

8.3 Trueness

The determined mean mass fractions in the test samples (see Annex C) are given in Table 3 together with the accepted values. Two of the values are certified. Comparing both sets of values allow to conclude that trueness is satisfactory.

Table 3 — Evaluation of trueness

Sample No.	Name	Accepted value % (mass fraction)	Value found % (mass fraction)
8-7-Mo	ETI 449 ^a	0,063	0,055 ₇
8-6-Mo	ETI 407 ^a	0,116	0,108 ₄
8-5-Mo	ETI 618 ^a	0,620	0,598 ₅
8-4-Mo	ETI 600 ^a	1,71	1,69 ₇
8-8-Mo	BCS 351	3,06	3,05 ₉
8-3-Mo	ETI 542 ^a	3,95	3,96 ₅
8-2-Mo	ETI 421 ^a	5,10	5,01 ₉
8-9-Mo	EMRC 377-1	8,94	8,93 ₆
8-1-Mo	ETI 443 ^a	16,94	17,11 ₅
^a Value not certified.			

9 Test report

The test report shall include the following information:

- all information necessary for the identification of the sample, the laboratory and the date of analysis or of the test report;
- the method used by reference to this International Standard;
- the results and the unit in which they are expressed;
- any unusual features noted during the determination;
- any operation not specified in this International Standard, or any optional operation which may have influenced the results.

Annex A (normative)

Checking the performance of an ICP instrument

A.1 Introduction

A joint working group (ISO/TC 47/SC 1) involving representatives from ISO/TC 47, ISO/TC 17 and ISO/TC 155 was formed in 1995 to establish guidelines for inductively coupled plasma spectrometry. The project reached the stage of a committee draft (ISO/CD 12235) but the work was not completed. This annex is abstracted from this committee draft and was used in the tests of this International Standard.

A.2 Resolution of a spectrometer

The resolution of a spectrometer can be defined as the wavelength difference $\Delta\lambda$, between two lines which can still just be observed separately. In practice, the parameter FWHM (Full Width at Half Maximum) is used as a resolution measure.

Ideally, the resolution should be of the same order as the physical line width in ICP/AES spectra, i.e. 2 pm to 5 pm (1 pm = 10^{-12} m). In practice, however, the observed width of the emission lines and, consequently, the resolution, will often be determined by the bandwidth (r_{bp}) of the spectrometer being used. As long as broadening resulting from aberrations can be neglected, this bandwidth is given by:

$$r_{bp} = \text{FWHM} = (d\lambda/dx)(w_i + w_u)/2$$

where

w_i and w_u are the widths of the entrance slit and exit slit, respectively;

$d\lambda/dx$ is the reciprocal linear dispersion which is given by:

$$d\lambda/dx = d(\cos \beta)/nL$$

where

L is the focal length of the spectrometer;

n is the order number;

d is the reciprocal of the groove density in the grating;

β is the diffraction angle.

Normally, commercial spectrometers present resolutions in the range of 4 pm to 30 pm. A good resolution is of great importance to cope with the frequent spectral interferences which occur in ICP/OES. Since a line with a wavelength in the second order will have the same diffraction angle β as a line with a wavelength 2λ in the first order, a spectrometer must either have an order-sorting possibility or an optical filter to avoid an order overlap.