

SLOVENSKI STANDARD oSIST prEN ISO 6143:2023

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Analiza plinov - Primerjalne metode za določevanje in preverjanje sestave kalibrirnih plinskih zmesi (ISO/DIS 6143:2023)

Gas analysis - Comparison methods for determining and checking the composition of calibration gas mixtures (ISO/DIS 6143:2023)

Gasanalyse - Vergleichsverfahren zur Bestimmung und Überprüfung der Zusammensetzung von Kalibriergasgemischen (ISO/DIS 6143:2023)

Analyse des gaz - Méthodes de comparaison pour la détermination et la vérification de la composition des mélanges de gaz d'étalonnage (ISO/DIS 6143:2023)

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Chemical analysis

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DRAFT INTERNATIONAL STANDARD ISO/DIS 6143

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Gas analysis — Comparison methods for determining and checking the composition of calibration gas mixtures

Analyse des gaz — Méthodes comparatives pour la détermination et la vérification de la composition des mélanges de gaz pour étalonnage

ICS: 71.040.40

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Contents

Page

Foreword				
Intro	ductio)n	vi	
1	Scor	De		
2	Nor	native references		
3	Terms and definitions			
4		bols and abbreviated terms		
5	Principle			
6	General procedure			
	6.1 6.2	Determination of the analysis function Validation of the analysis function 6.2.1 Purpose 6.2.2 Validation of the response model 6.2.3 Examining compliance with uncertainty requirements		
	6.3 6.4	 6.2.4 Drift control of the measuring system. 6.2.5 Validation of applicability to mismatching calibration gases Determination of the composition of a calibration gas mixture Supplementary instructions	10 11 12 13 13	
7	Spec 7.1 7.2	cial procedures Checking of a pre-assigned composition for the second se		
8		ort of results		
Anne	x A (n	ormative) Procedures for data evaluation Media 666-6910-49d5-6666-		
Anne	x B (ir	oformative) Examples		
		formative) Computer implementation of recommended methods		
Annex D (informative) Additional information on data evaluation				
Bibliography				
21011	or up	.,		

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 www.iso.org/directives).

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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 www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 158, Analysis of Gases.

This third edition cancels and replaces the second edition (ISO 6143:2001), which has been technically revised.

The main changes compared to the previous edition are as follows:

- update of definitions, in particular those taken from the VIM
- update of the bibliography and the corresponding references in the text
- update of the information in <u>Annex C</u> on the computer programme B_LEAST; information on alternative software (<u>Annex D</u>)
- amendment of <u>section 6.2</u> (now <u>7.2</u>) "Comparison of several calibration gas mixtures" and related statements in other parts of the Document
- amendment of the recommendations concerning the number of replicate measurements per sample
- revision of the requirements for the report of results ("Test report")
- new <u>Annex D</u> (informative) "Additional information on data evaluation", including a comparison between the regression method used in ISO 6143 and ordinary least-squares regression, providing information on relevant documents and software, and dealing with the evaluation of replicate response measurements
- deletion of <u>section A.1</u> "Uncertainty specifications for reference gas mixtures" that is no longer required
- additional references to relevant ISO standards (ISO 12963, ISO 14912, ISO 15796)
- correction of the formula for the power function

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>.

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Introduction

In gas analysis, calibration of analytical systems is most often confined to the determination of a straight line through the origin, or of a straight-line segment, using only the minimum number of calibration standards (one for a straight line through the origin, two for a line segment). This approach was also adopted in the first edition (1981) of ISO 6143. However, ISO 6143 is devoted to a very special task: the derivation of calibration gases from appropriate reference gases. Consequently, the multiplier effect of errors in calibration gases – an error in a calibration gas may cause errors in thousands of analytical results – implies high demands on the metrological quality of the analysis of calibration gases. In the development of the second edition (2001) it was therefore decided to use the best available measurement strategy and data evaluation method. The main changes in the revision of the (1981) edition related to calibration as well as to uncertainty evaluation:

- including non-linear response curves and/or functions,
- replacing interpolation by regression,
- taking into account the uncertainty on the calibration standards,
- including validation of calculated response curves and/or functions,
- calculating uncertainties by uncertainty propagation.

After twenty years, the principles and procedures specified in the second edition are still fit for purpose. The current revision therefore mainly concerns additional supporting information.

As a consequence of adopting non-linear response models, advanced regression techniques (errors in both variables) and uncertainty propagation, the main calculation procedures can only be performed on a computer, using a specific program. A dedicated program (B_LEAST) is available and will be provided without cost (see <u>Annex C</u>). Information on other publicly available software that can be used for at least the vast majority of the calculations required by ISO 6143 is given in <u>Annex D</u>. As an alternative, sufficient information is given in the Document to enable the user to develop a program on his own.

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DRAFT INTERNATIONAL STANDARD

Gas analysis — Comparison methods for determining and checking the composition of calibration gas mixtures

1 Scope

This document provides methods for:

- determining the composition of a calibration gas mixture by comparison with appropriate reference gas mixtures,
- calculating the uncertainty of the composition of a calibration gas mixture in relation to the known uncertainty of the composition of the reference gas mixtures with which it was compared,
- checking the composition attributed to a calibration gas mixture by comparison with appropriate reference gas mixtures,
- consistency testing and outlier search in suites of calibration gas mixtures of closely related composition.

NOTE In principle, the method described in this document is also applicable to the analysis of (largely) unknown samples instead of prospective calibration gas mixtures (i.e. gas mixtures which are intended for use as calibration gas mixtures). Such applications, however, require appropriate care and consideration of additional uncertainty components, for example concerning the effect of matrix differences between the reference gases used for calibration and the analysed sample. On the other hand, in many applications one- and two-point calibration is preferred over multi-point calibration. ISO 12963^[5] is designed for such applications.

Normative references

2

There are no normative references in this Document. iso-6143-2023

Terms and definitions 3

For the purposes of this Document, the following terms and definitions 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 <u>https://www.electropedia.org/</u>

3.1

composition

characteristic of a gas mixture given by the kind and content of each specified mixture component (analyte) and the composition of the complementary gas (matrix)

Note 1 to entry: In this Document, the analyte content is specified as an amount fraction, exclusively. Amount fractions have the advantage of being perfectly independent of the pressure and the temperature of the gas mixture. Therefore their use is recommended. However, for specific measuring systems, other composition measures (e.g. mass concentrations) may be more appropriate. Their use then requires due care concerning the dependence on pressure and temperature. Methods for conversion between different quantities of composition are specified in ISO 14912.

3.2

comparison method

method for determining the content of a specified gas mixture component (analyte) by measuring an instrumental response

Note 1 to entry: Comparison of measuring systems requires calibration, in which the relationship between response and analyte content is established. This is achieved by measuring the response to known values of analyte content provided by reference gas mixtures.

3.3

calibration

operation that, under specified conditions, in a first step, establishes a relation between the quantity values with measurement uncertainties provided by measurement standards and corresponding indications with associated measurement uncertainties and, in a second step, uses this information to establish a relation for obtaining a measurement result from an indication

[SOURCE: VIM 3, 2.39]

3.4

response function

functional relationship between instrumental response and analyte content

Note 1 to entry: The response function can be expressed in two different ways as a calibration function or an analysis function, depending on the choice of the dependent and the independent variable.

Note 2 to entry: The response function is conceptual and cannot be determined exactly. It is determined approximately through calibration. **DIANDARD**

3.4.1

calibration function

instrumental response expressed as a function of analyte content

3.4.2

analysis function https://standards.iteh.ai/catalog/standards/sist/fcf16bf6-f910-49d5-bb6b-

analyte content expressed as a function of instrumental response $^{143-2023}$

3.5

measurement uncertainty

uncertainty of measurement

uncertainty

non-negative parameter characterizing the dispersion of the quantity values being attributed to a measurand, based on the information used

[SOURCE: VIM 3, 2.26]

3.6

metrological traceability

traceability

property of a measurement result whereby the result can be related to a reference through a documented unbroken chain of calibrations, each contributing to the measurement uncertainty

[SOURCE: VIM 3, 2.41]

3.7

measurement standard

realization of the definition of a given quantity, with stated quantity value and associated measurement uncertainty, used as a reference

[SOURCE: VIM 3, 5,1]

3.8 reference measurement standard reference standard

measurement standard designated for the calibration of other measurement standards for quantities of a given kind in a given organization or at a given location

[SOURCE: VIM 3, 5.6]

3.9

working measurement standard working standard

measurement standard that is used routinely to calibrate or verify measuring instruments or measuring systems

[SOURCE: VIM 3, 5.7]

3.10

reference material

material, sufficiently homogeneous and stable with reference to specified properties, which has been established to be fit for its intended use in measurement or in examination of nominal properties

[SOURCE: VIM 3, 5.13]

3.11

calibration gas mixture

gas mixture of known stability and homogeneity whose composition is well established for use in the calibration or verification of a measuring instrument or for the validation of a measurement

[SOURCE: ISO 7504, 5.1]

3.12

reference gas mixture

calibration gas mixture whose composition is well established and stable to be used as a reference standard of composition from which other composition data measurements are derived

[SOURCE: ISO 7504, 5.2]

4 Symbols and abbreviated terms

a_j	parameters of the calibration function $F(j = 0, 1,, N)$
b_j	parameters of the analysis function $G(j = 0, 1,, N)$
D	sensitivity matrix
F	calibration function, $y = F(x)$, for the specified analyte
G	analysis function, $x = G(y)$, for the specified analyte
k	coverage factor
L	limit of detection
M _{cal}	(sample of) calibration gas mixture
M _{ref}	(sample of) reference gas mixture
Q	transform matrix
S	standard deviation of a data set (sample standard deviation)

S	sum of weighted squared deviations			
S _{res}	residual sum of weighted squared deviations			
$t_{\nu;(1-\alpha)}$	quantile of the <i>t</i> -distribution (ν degrees of freedom, confidence level (1- α))			
U(q)	expanded uncertainty of an estimated quantity q , $U(q) = ku(q)$			
u(q)	uncertainty of an estimated quantity <i>q</i> , expressed as a standard deviation (standard uncer- tainty)			
u(p,q)	covariance of two estimated quantities p and q			
$u^{2}(q)$	variance of an estimated quantity q			
V	variance/covariance matrix			
X	amount fraction of the specified analyte			
(x_i, y_i)	calibration points (<i>i</i> = 1, 2,, <i>n</i>)			
$(\hat{x}_{i,}\hat{y}_{i})$	adjusted calibration points (<i>i</i> = 1, 2,, <i>n</i>)			
у	instrumental response of the specified analyte			
$\chi^2_{\nu;(1-\alpha)}$	quantile of the Chi ² -distribution (ν degrees of freedom, confidence level (1- α))			
γ	dilution factor (standards.iteh.ai)			
σ	standard deviation of a probability distribution			
Г	measure of goodness-of-fit <u>oSIST prEN ISO 6143:2023</u> https://standards.iteh.ai/catalog/standards/sist/fcf16bf6-f910-49d5-bb6b-			
2ed2a81b5d07/osist-pren-iso-6143-2023				

5 Principle

The composition of a gas mixture is determined by separate determination of the amount fraction of every specified analyte. Therefore, the procedure for determining the amount fraction of only one specified analyte is described. Possible interferences of other components on the measurement of the analyte under consideration should be considered by the user and taken into account. However, this subject is not addressed in this Document.

This Document is also applicable if other composition quantities than amount fraction are used. However, it is recommended that the final result be expressed as an amount fraction. Methods for conversion between different quantities of composition are specified in ISO 14912^[6].

The general procedure for determining the amount fraction *x* of a specified analyte in a sample of a calibration gas mixture, or in a series of such samples, is performed in a sequence of steps summarized below.

- a) Specify the analytical range of interest, i.e. the range of the amount fractions x to be determined, and the acceptable uncertainty level (see <u>6.1</u>, step A).
- b) Specify the analytical method and the measuring system to be used (see <u>6.1</u>, step B).
- c) Examine the available information on the relevant response characteristics of the measuring system (e.g. linearity and sensitivity), paying attention to possible interferences. If necessary, carry out a performance evaluation to check the suitability of the system. Specify the type of mathematical function to be considered for description of the response in the specified range (see <u>6.1</u>, step C).

- d) Set up a design for the calibration experiment in which the relevant experimental parameters are specified, such as:
 - calibration range (to include the analytical range),
 - composition, including uncertainty, of the reference gas mixtures for calibration,
 - parameters of the analytical method,
 - conditions of measurement, if relevant,
 - number and sequence of calibration measurements (see <u>6.1</u>, steps D, E, F).
- e) Perform the calibration experiment, i.e. measure the response, *y*, for samples of the chosen reference gas mixtures, and estimate the uncertainty u(y) of these response values (see <u>6.1</u>, step G).
- f) Calculate the analysis function, x = G(y), from the calibration data, using regression analysis (see <u>6.1</u>, step H).
- g) Examine whether the calculated analysis function is consistent with the calibration data within the relevant uncertainties. If the result is acceptable, proceed to h). If not, revise the calibration design (see <u>6.2.2</u>).
- h) Determine the uncertainty level of the prospective results based on the analysis function for the relevant ranges of responses and analyte contents. If the result is acceptable, proceed to i). If not, revise the calibration design (see <u>6.2.3</u>).
- i) Prior to analysing a prospective calibration gas sample, test for instrument drift to ensure that the analysis function is still valid for the specified analytical task (see <u>6.2.4</u>). If the result is acceptable, proceed to j). If not, recalibrate the measuring system.

If the prospective calibration gas contains other components than the reference gas mixtures used for calibration, validate the applicability of the analysis function using at least one additional reference gas mixture of appropriate composition (see <u>6.2.5</u>).

NOTE It is not necessary to test for drift in conjunction with every analysis of a calibration gas sample. The frequency should be based on experience concerning the stability of the measuring system.

Similarly, the composition of additional reference gas mixtures used for validation should be based on experience concerning the cross-sensitivities of the measuring system.

- j) Determine the composition of the prospective calibration gas as follows:
 - measure the response *y*,
 - determine the uncertainty u(y) of the response y,
 - calculate the amount fraction x = G(y) using the analysis function determined in f),
 - calculate the uncertainty u(x) of the amount fraction x by propagation of uncertainty on the measured response and on the parameters of the analysis function (see <u>6.3</u>).
- k) State the result of the entire analysis (see <u>clause 8</u>).

In addition to determining the composition of a (prospective) calibration gas mixture, the general procedure may be used to check a pre-established composition. To this end, the mixture under consideration is analysed using the procedure outlined above, and the composition obtained is compared with the pre-established composition. <u>Clause 7</u> specifies a procedure where, for each analyte concerned, the difference between the content obtained by the confirmation analysis and the pre-established content is examined against the uncertainty on this difference for significant departure from zero.