



SLOVENSKI STANDARD
SIST-TS ISO/TS 22176:2024

01-marec-2024

Kozmetika - Analizne metode - Razvoj globalnega pristopa za validacijo kvantitativnih analiznih metod

Cosmetics - Analytical methods - Development of a global approach for validation of quantitative analytical methods

Cosmétiques - Méthodes analytiques - Développement d'une approche globale pour la validation des méthodes analytiques quantitatives

Ta slovenski standard je istoveten z: ISO/TS 22176:2020

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>

ICS:

71.100.70	Kozmetika. Toaletni pripomočki	Cosmetics. Toiletries
-----------	--------------------------------	-----------------------

SIST-TS ISO/TS 22176:2024 **en**

TECHNICAL SPECIFICATION

ISO/TS 22176

First edition
2020-01

Cosmetics — Analytical methods — Development of a global approach for validation of quantitative analytical methods

*Cosmétiques — Méthodes analytiques — Développement d'une
approche globale pour la validation des méthodes analytiques
quantitatives*

iTeh Standards
(<https://standards.iteh.ai>)
Document Preview

[SIST-TS ISO/TS 22176:2024](https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024)

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>



Reference number
ISO/TS 22176:2020(E)

© ISO 2020

ISO/TS 22176:2020(E)

iTeh Standards
(<https://standards.iteh.ai>)
Document Preview

[SIST-TS ISO/TS 22176:2024](https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024)

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>



COPYRIGHT PROTECTED DOCUMENT

© ISO 2020

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Fax: +41 22 749 09 47
Email: copyright@iso.org
Website: www.iso.org

Published in Switzerland

Contents

Page

Foreword	v
Introduction	vi
1 Scope	1
2 Normative references	1
3 Terms, definitions and symbols	1
3.1 Terms and definitions.....	1
3.2 Symbols.....	5
4 General principles	6
4.1 Reminder.....	6
4.2 Various conditions for the estimation of precision.....	6
4.3 Accuracy profile.....	7
5 Procedure	9
5.1 Definition of the measured quantity.....	9
5.2 Definition of objectives.....	9
5.2.1 Choice of the scope of validation.....	9
5.2.2 Choice of acceptance limits.....	9
5.3 Selection of validation samples.....	10
5.3.1 Choice of the type of matrix or types of matrices.....	10
5.3.2 Methods for establishing reference values.....	10
5.4 Characterization plan for validation.....	10
5.4.1 Organization.....	10
5.4.2 Choice of the number of series, repetitions and concentrations for the characterization plan for validation.....	11
5.5 Calibration plan for the indirect methods.....	11
5.5.1 Organization.....	11
5.5.2 Choice of the number of series, repetitions and concentrations for the calibration plan.....	12
5.6 Testing.....	13
5.7 Calculation of predicted inverse concentrations for indirect methods.....	14
5.7.1 General.....	14
5.7.2 Calculation of the calibration models.....	14
5.7.3 Calculation of back-calculated concentrations by inverse prediction.....	15
5.8 Calculation of the validation criteria by concentration level.....	15
5.8.1 General.....	15
5.8.2 Trueness criteria by series.....	15
5.8.3 Trueness and precision criteria by concentration.....	16
5.8.4 Calculation of the tolerance intervals.....	17
5.9 Construction of the accuracy profile.....	18
5.10 Interpretation of the accuracy profile for validation.....	19
5.10.1 General.....	19
5.10.2 Decision rules.....	20
5.10.3 Definition of the scope of validity.....	21
5.10.4 Choice of a calibration procedure for the routine.....	21
5.10.5 Influence and significance of the β proportion.....	21
5.10.6 Identification of outliers.....	22
6 Management of the outcomes during routine use	22
Annex A (normative) Calculation of repeatability, intermediate precision and reproducibility standard deviations	23
Annex B (normative) Contents of the validation file	25
Annex C (informative) Setting-up an assay for determining the accuracy profile in the case of NDELA in cosmetic samples	27

ISO/TS 22176:2020(E)

Annex D (informative) Influence of the value of β on the tolerance interval ($R = 3$ and $s_{IP} = 1$)	37
Annex E (informative) Contribution to the uncertainty calculation	38
Bibliography	39

iTeh Standards
(<https://standards.iteh.ai>)
Document Preview

[SIST-TS ISO/TS 22176:2024](https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024)

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>

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).

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 www.iso.org/patents).

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

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 national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

[SIST-TS ISO/TS 22176:2024](https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024)

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>

Introduction

The purpose of this document is to propose a characterization protocol for the validation of a quantitative analysis method in the cosmetic field and thus responds to the requirements of ISO/IEC 17025, i.e. using the performance goals as a basis. The theoretical principles of this approach can be found in Reference [1]. This document is based on the French Standard NF V 03-110[2].

Analytical methods for analyses of cosmetics need to be validated. Validation has been long considered as a process consisting in individually verifying several different criteria, i.e. selectivity, repeatability, linearity, trueness, etc. The global approach, as proposed since 2003[1], is based on the total error concept and the term “global” means that only a single criterion should be checked to validate a method: the agreement between a future experimental result and the true value. This approach has already been applied in the domains of pharmacy[1],[9], agricultural chemistry[2], and is in agreement with quality assurance guidelines such as GLP or ISO/IEC 17025. This validation process applies generally to already developed methods and includes evaluations of the following criteria: specificity/selectivity, precision, trueness, linearity range, LOD/LOQ, stability, ruggedness.

The large number of cosmetic products and the variety of matrices present a challenge for an analytical laboratory requiring that standardized methods to be adapted for each type of samples. Additional difficulties are linked to the very low concentrations to be measured, generally of the order of the mg/kg (ppm) or µg/kg (ppb). In such context, criteria such as accuracy and uncertainty of measurement of the analytical results are of utmost importance.

When the concentration of a substance is determined by an analytical laboratory, it is important to evaluate the gap between the measured value and the known true value. This difference indicates the trueness of the analysis. If cosmetic samples are analysed several times in different conditions (laboratory, instrument, operator), the individual results will present a dispersal around the average value which represents the precision of the measurement. As for the individual measurement, it represents an error with the average value and an inaccuracy with regard to the reference value (i.e. the true value).

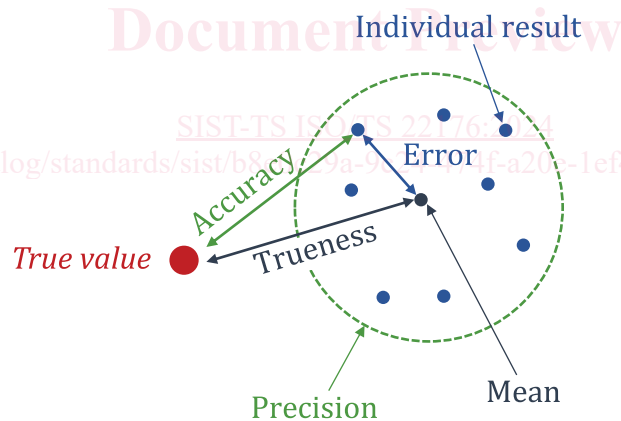


Figure 1 — Illustration of the concepts of accuracy, precision and trueness

When a laboratory measures the concentration of a given substance in a cosmetic product sample, the value which is obtained is thus characterized by a given accuracy which includes at the same time the notion of trueness and precision (see Figure 1). It can also be considered as total error. The insurance that the accuracy of a result is below acceptable limits, is thus one of the ways to make sure of the validity of a measurement.

The accuracy profile (plot of accuracy versus concentration), such as it is developed in numerous domains[3] to [9], is thus the way to know the accuracy on a result obtained with a given method applied to a type of sample in the environment of a given laboratory.

To reach this accuracy profile, it is necessary to undergo a specific assay allowing to demonstrate the validity of the analytical method, as well as the accuracy of the measurement for a given substance. In this approach, it is necessary to determine a tolerance interval^[10] which contains a given proportion (β) of future measured values inside (in average). If this tolerance interval is located inside a limit of acceptability defined a priori, taking into consideration several parameters such as the type and concentration of analyte, type of matrix, of analysis and conditions of the experiments, in this case, the method will be considered as valid, and if it goes outside this limit of acceptability, the method will be considered as non-valid (see [Figure 2](#)).

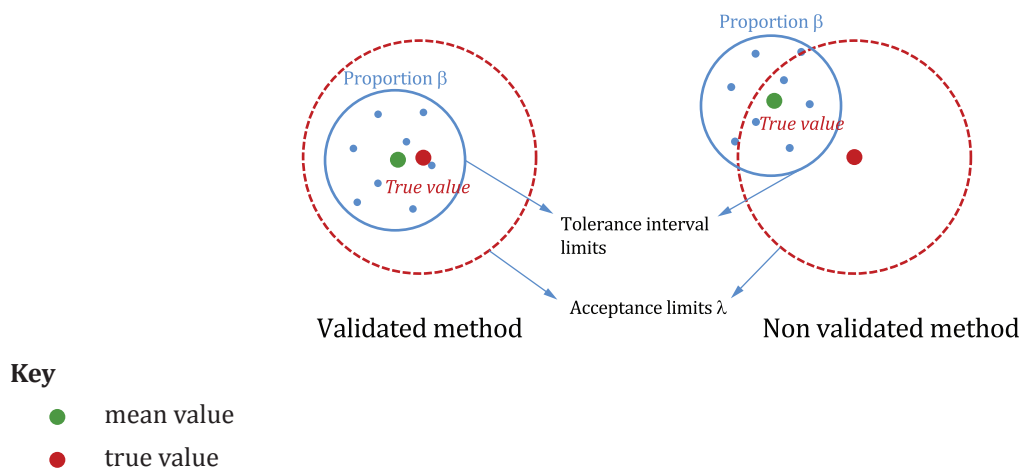


Figure 2 — Illustration of the validation principle

iTeh Standards
<https://standards.iteh.ai>
 Document Preview

[SIST-TS ISO/TS 22176:2024](https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024)

<https://standards.iteh.ai/catalog/standards/sist/b8e0c29a-9c24-474f-a20e-1ef4e592e1ab/sist-ts-iso-ts-22176-2024>

Cosmetics — Analytical methods — Development of a global approach for validation of quantitative analytical methods

1 Scope

This document defines a global approach for the validation of a quantitative analytical method, based on the construction and interpretation of an accuracy profile, and specifies its characterization procedure.

This procedure is particularly applicable for internal validation in a cosmetic testing laboratory, but its scope can be extended to the interpretation of data collected for an interlaboratory study designed according to the recommendations of the ISO 5725-1. It does not apply to microbiological trials. The present approach is particularly suited to handle the wide diversity of matrices in cosmetics. This document only applies to already fully-developed and finalized methods for which selectivity/specificity have already been studied and the scope of the method to be validated has already been defined, in terms of matrix types and measurand (for example analyte) concentrations.

2 Normative references

The following document is referred to in the text in such a way that some or all of its 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/IEC Guide 99:2007, *International vocabulary of metrology — Basic and general concepts and associated terms (VIM)* (<https://standards.iteh.ai>)

3 Terms, definitions and symbols

3.1 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO/IEC Guide 99 and the following 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 <http://www.electropedia.org/>

3.1.1

measurement

process of experimentally obtaining one or more quantity values that can reasonably be attributed to a quantity

[SOURCE: ISO/IEC Guide 99:2007, 2.1, modified — Notes to entry have been excluded.]

3.1.2

measurand

quantity intended to be measured

Note 1 to entry: The term “analyte”, employed in chemistry, is a synonym of measurand, and is used more generally.

[SOURCE: ISO/IEC Guide 99:2007, 2.3, modified — Original notes to entry have been excluded and a new note to entry has been added.]

ISO/TS 22176:2020(E)

3.1.3

measurement trueness

trueness

closeness of agreement between the average of values obtained by replicate measurements of the same or similar objects under specified conditions and a reference quantity value

[SOURCE: ISO/IEC Guide 99:2007, 2.14, modified — Notes to entry have been excluded.]

3.1.4

measurement precision

precision

closeness of agreement between indications or measured quantity values obtained by replicate measurements on the same or similar objects under specified conditions

[SOURCE: ISO/IEC Guide 99:2007, 2.15, modified — Notes to entry have been excluded.]

3.1.5

repeatability condition

condition of measurement, out of a set of conditions that includes the same measurement procedure, same operator, same measuring system, same operating conditions and same location, and replicate measurements on the same or similar objects over a short period of time

[SOURCE: ISO/IEC Guide 99:2007, 2.20, modified — Notes to entry have been excluded.]

3.1.6

measurement repeatability

repeatability

measurement precision under a set of *repeatability conditions* (3.1.5) of measurement

[SOURCE: ISO/IEC Guide 99:2007, 2.21]

3.1.7

intermediate precision condition

condition of measurement, out of a set of conditions that includes the same measurement procedure, same location, and replicate measurements on the same or similar objects over an extended period of time, but may include other conditions involving changes

[SOURCE: ISO/IEC Guide 99:2007, 2.22, modified — Notes to entry have been excluded.]

3.1.8

intermediate measurement precision

intermediate precision

measurement precision under a set of *intermediate precision conditions* (3.1.7) of measurement

[SOURCE: ISO/IEC Guide 99:2007, 2.23, modified — Notes to entry have been excluded.]

3.1.9

reproducibility condition of measurement

reproducibility condition

condition of measurement, out of a set of conditions that includes different locations, operators, measuring systems, and replicate measurements on the same or similar objects

[SOURCE: ISO/IEC Guide 99:2007, 2.24, modified — Note to entry has been excluded.]

3.1.10

measurement reproducibility

reproducibility

measurement precision under *reproducibility conditions of measurement* (3.1.9)

[SOURCE: ISO/IEC Guide 99:2007, 2.25, modified — Note to entry has been excluded.]

3.1.11**measurement accuracy
accuracy**

closeness of agreement between a measured quantity value and a true quantity value of a measurand

[SOURCE: ISO/IEC Guide 99:2007, 2.13, modified — Notes to entry have been excluded.]

3.1.12**verification**

provision of objective evidence that a given item fulfils specified requirements, taking into account any measurement uncertainty

[SOURCE: ISO/IEC Guide 99:2007, 2.44, modified — Notes to entry have been excluded.]

3.1.13**validation**

verification, where the specified requirements are adequate for an intended use

Note 1 to entry: The term “characterization” applies to the method, whereas the term “verification” applies to the outcomes. Validation of the method therefore consists of checking if the results are adequate for an intended use.

[SOURCE: ISO/IEC Guide 99:2007, 2.45, modified — Example has been excluded and a Note to entry has been added.]

3.1.14**selectivity**

property of a measuring system, used with a specified measurement procedure, whereby it provides measured quantity values for one or more measurands such that the values of each measurand are independent of other measurands or other quantities in the measuring system

Note 1 to entry: The IUPAC considers specificity as the final stage of selectivity.

[SOURCE: ISO/IEC Guide 99:2007, 4.13, modified — Examples and original notes to entry have been excluded. A new note to entry has been added.]

3.1.15**reference value**

quantity value whose associated measurement uncertainty is generally considered small enough so that the value may be used as a basis for comparison with quantity values of the same kind

[SOURCE: ISO/IEC Guide 99:2007, 5.18, modified — Notes to entry have been excluded.]

3.1.16**scope**

<of the method>all of the types of *matrix* (3.1.22) to which the method applies, taking into account the range of concentrations involved in validation

3.1.17**scope of validation**

all of the types of *matrix* (3.1.22) to which the method and range of concentrations involved in validation applies

3.1.18**scope of validity**

all of the types of *matrix* (3.1.22) to which the method and range of concentrations involved in validation applies, and for which future outcomes obtained via the method will be considered valid

3.1.19**quantitative method**

method of analysis which determines the quantity or weight fraction of an analyte so that it may be expressed as a numeric value in the appropriate units

ISO/TS 22176:2020(E)

3.1.20

reference method

method of analysis recognized by experts or used as a reference by agreement between parties, which gives, or is supposed to give the accepted reference value of the measurand

3.1.21

alternative method

method of analysis used by the laboratory instead of one or several *reference methods* ([3.1.20](#))

3.1.22

matrix

set of properties of the sample and its components other than the analyte

Note 1 to entry: The matrix effect reflects the possible influence that these properties or components can have on the instrumental response. For practical reasons, since the matrix effect can vary in the different stages of analysis (e.g. before or after mineralisation), a type of matrix is defined as a group of materials or products recognized by the analyst as having consistent behaviour with regard to the method of analysis used.

3.1.23

series

set of measurements carried out under a set of repeatability conditions

Note 1 to entry: For example, a series includes measurements carried out on the same day and/or by the same operator.

3.1.24

accuracy profile

combination, in a graphic form, of one or several *β -expectation tolerance intervals* ([3.1.25](#)) calculated at different concentrations, and of one or several *acceptance intervals* ([3.1.26](#))

3.1.25

β -expectation tolerance interval tolerance interval

interval which contains, on average, a defined proportion, β %, of future measurements, obtained according to a given procedure and for a given concentration

Note 1 to entry: The limits of the interval are calculated based on trials conducted for the purpose of validation.

Note 2 to entry: A value of 80 % for β % means that, on average, one out of five results will be outside the limits of the interval at the *limit of quantitation* ([3.1.29](#)). See [5.10](#).

3.1.26

acceptance interval

specification of the performance required for the method, expressed as an acceptable deviation around the reference value

Note 1 to entry: The limits of the interval are set by the client or by statutory requirements, sometimes according to the concentration. They are expressed as $\pm\lambda$ as absolute values and in the units of the measurand, or $(1 \pm \lambda) \times 100$ as relative values.

3.1.27

linearity

<of the method> establishment of a linear relationship between the deduced (or quantified) quantities in the samples and their reference values

Note 1 to entry: Linearity of the method is different from linearity of the response function of the measuring apparatus, which only characterizes the instrumental response during calibration and is not essential for accurate quantitation.