



SLOVENSKI STANDARD
SIST EN ISO 10993-18:2020/oprA1:2021

01-oktober-2021

Biološko ovrednotenje medicinskih pripomočkov - 18. del: Kemična opredelitev lastnosti materialov za medicinske pripomočke znotraj procesov obvladovanja tveganja - Dopolnilo 1: Določitev faktorja negotovosti (ISO 10993-18:2020/DAM 1:2021)

Biological evaluation of medical devices - Part 18: Chemical characterization of medical device materials within a risk management process - Amendment 1: Determination of the uncertainty factor (ISO 10993-18:2020/DAM 1:2021)

iTeh STANDARD PREVIEW

Biologische Beurteilung von Medizinprodukten - Teil 18: Chemische Charakterisierung von Werkstoffen für Medizinprodukte im Rahmen eines Risikomanagementsystems - Änderung 1: Bestimmung des Unsicherheitsfaktors (ISO 10993-18:2020/DAM 1:2021)

[https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-oprA1-2021)

Évaluation biologique des dispositifs médicaux - Partie 18: Caractérisation chimique des matériaux des dispositifs médicaux au sein d'un processus de gestion du risque - Amendement 1: Détermination du coefficient d'incertitude (ISO 10993-18:2020/DAM 1:2021)

Ta slovenski standard je istoveten z: EN ISO 10993-18:2020/prA1

ICS:

11.100.20	Biološko ovrednotenje medicinskih pripomočkov	Biological evaluation of medical devices
-----------	---	--

SIST EN ISO 10993-18:2020/oprA1:2021 en,fr,de

iTeh STANDARD PREVIEW
(standards.iteh.ai)

[SIST EN ISO 10993-18:2020/oprA1:2021](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021)

<https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021>

DRAFT AMENDMENT

ISO 10993-18:2020/DAM 1

ISO/TC 194

Secretariat: DIN

Voting begins on:
2021-08-16Voting terminates on:
2021-11-08

Biological evaluation of medical devices —

Part 18: Chemical characterization of medical device materials within a risk management process

AMENDMENT 1: Determination of the uncertainty factor

*Évaluation biologique des dispositifs médicaux —**Partie 18: Caractérisation chimique des matériaux des dispositifs médicaux au sein d'un processus de gestion du risque*

AMENDEMENT 1

iTeh STANDARD PREVIEW
(standards.iteh.ai)

ICS: 11.100.20

[SIST EN ISO 10993-18:2020/oprA1:2021](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021)<https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021>

THIS DOCUMENT IS A DRAFT CIRCULATED FOR COMMENT AND APPROVAL. IT IS THEREFORE SUBJECT TO CHANGE AND MAY NOT BE REFERRED TO AS AN INTERNATIONAL STANDARD UNTIL PUBLISHED AS SUCH.

IN ADDITION TO THEIR EVALUATION AS BEING ACCEPTABLE FOR INDUSTRIAL, TECHNOLOGICAL, COMMERCIAL AND USER PURPOSES, DRAFT INTERNATIONAL STANDARDS MAY ON OCCASION HAVE TO BE CONSIDERED IN THE LIGHT OF THEIR POTENTIAL TO BECOME STANDARDS TO WHICH REFERENCE MAY BE MADE IN NATIONAL REGULATIONS.

RECIPIENTS OF THIS DRAFT ARE INVITED TO SUBMIT, WITH THEIR COMMENTS, NOTIFICATION OF ANY RELEVANT PATENT RIGHTS OF WHICH THEY ARE AWARE AND TO PROVIDE SUPPORTING DOCUMENTATION.

This document is circulated as received from the committee secretariat.

ISO/CEN PARALLEL PROCESSING

Reference number
ISO 10993-18:2020/DAM 1:2021(E)

iTeh STANDARD PREVIEW (standards.iteh.ai)

[SIST EN ISO 10993-18:2020/oprA1:2021
https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021)



COPYRIGHT PROTECTED DOCUMENT

© ISO 2021

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
Email: copyright@iso.org
Website: www.iso.org

Published in Switzerland

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 the following URL: www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 194, *Biological and clinical evaluation of medical devices*.

<https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-10993-18:2020/oprA1:2021>

A list of all parts in the ISO 10993 series can be found on the ISO website.

iTeh STANDARD PREVIEW
(standards.iteh.ai)

[SIST EN ISO 10993-18:2020/oprA1:2021](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021)

<https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021>

Biological evaluation of medical devices —

Part 18:

Chemical characterization of medical device materials within a risk management process

AMENDMENT 1: Determination of the uncertainty factor

5.6, paragraph below Figure 3, last sentence

Replace Table 3 by Table 4

6.2, Table 3,

In the column “Qualitative” for the example method “Gravimetric”, insert “—”

iTeh STANDARD PREVIEW
(standards.iteh.ai)

6.4, Table 4,

In the columns “Qualitative” and “Quantitative” for the example methods “HPLC, with UV, CAD, ELSD and/or MS*” insert “X” in both columns

<https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-oprA1-2021>

Table D.2, Table footnote ^a

Replace the text in the table footnote ^a with the following:

^a Abbreviations include:

ABS	poly(acrylonitrile-butadiene-styrene);
ACN	acetonitrile;
AE	ethyl acetate;
DCM	dichloromethane;
DMF	dimethylformamide;
HFIP	hexafluoroisopropanol;
PET	poly(ethylene terephthalate);
TCB	trichlorobenzene;
THF	tetrahydrofuran;
MeOH	methanol;
EtOH	ethanol;

ISO 10993-18:2020/DAM 1:2021(E)

iPrOH = isopropyl alcohol.

Table D.2,

In the column "Anti-solvents" for "Polymer" Polystyrene and Styrenics (ABS), replace "can" by "ACN"

Page 52, E.3

Replace E.3 with the following:

Quantification in extractables profiling is achieved by various means which differ with respect to the accuracy of the estimated and reported concentration, where the accuracy can vary significantly depending on the quantification means employed. For example, quantification could involve the use of an internal standard to normalize the responses obtained for all relevant analytes. In such an approach, one estimates the concentration of each analyte based on the simplifying assumption that all analytes respond similarly, among themselves and with respect to the internal standard (i.e. all substances have the same response factor). Depending on the validity of this simplifying assumption, the concentration estimates thus obtained can have widely differing uncertainties and degrees of accuracy. If the simplifying assumption is true and response factors are constant, then the resulting concentration estimates for all analytes will be highly accurate. If the simplifying assumption is false and the response factors vary widely, then the resulting concentration estimates for the analytes will have widely varying accuracies, and the accuracy of the concentration estimate for each analyte will vary in proportion to the difference between the analyte's response factor and the internal standard's response factor.

(standards.iteh.ai)

Other quantitation means can produce highly accurate concentration estimates. For example, if quantification is achieved via the use of calibration curves generated via the analysis of authentic standards employed in qualified analytical methods, the concentration estimates obtained for the qualified analytes will be highly accurate. As noted above, if response factors are constant, then quantitation with an internal standard will also be highly accurate.

Other quantification strategies could produce concentration estimates whose accuracy is somewhere between these two extremes; greater accuracy than use of an internal standard's response factor but lesser accuracy than use of a calibration curve generated with an authentic reference standard. For example, relative response factors can be obtained for extractables, where the relative response factor is the ratio of the response of the extractable versus that of an internal standard at equal concentrations of extractable and internal standard. Use of relative response factors in quantification accounts and adjusts for differences in response factors, extractable versus internal standard.

Recognizing that response factors for extractables and internal standards can vary, the AET is adjusted to account for more poorly responding analytes. Such an adjustment increases the likelihood that even a poorly responding analyte can be recognized as being above the AET when it is present in a sample at levels greater than the AET. The adjustment is accomplished by adding an uncertainty factor (UF) to the calculation of the AET to account for response factor variation. Use of a UF is the same principle as calculation of a final AET from an estimated AET (e.g. see Reference [45]). In essence, use of the UF adjusts the AET down to a lower value, ensuring that poorly responding compounds are properly flagged as being at or above the AET and therefore being reportable.

In cases where the response factor variation is known to be acceptably low, a UF value of 1 can be justified. Examples of these cases are methods with comparable response factors between expected extractables and applied internal standards and qualified methods for targeted extractables. Otherwise, the value of the uncertainty factor is based on an assessment of the analytical methodology to which the AET is applied. For example, a UF value of 2 has been proposed [39] [45] as being appropriate, in certain situations, to the screening of extracts for organic extractables via GC-FID or GC-MS, as analytical FID or MS response factors for extractables are somewhat consistent, extractable to extractable. Alternatively, the UF for other analytical methods used for extractables screening, such as HPLC-MS, may be higher, given the frequently wide variation in response factors among extractables by this methodology. At the

current time, there is no available general guidance which recommends a specific value for the UF for these methods; however, the user should justify the UF values selected.

One approach to establishing and justifying a particular UF is statistical analysis of a database of response factors specific to the analytical method being considered and the population of extractables for which that method is applicable. In this approach, the value of the UF is linked to the relative standard deviation of the response factors according to [Formula \(E.2\)](#):

$$UF = \frac{1}{(1-RSD)} \quad (E.2)$$

where, RSD is the relative standard deviation of the response factors from the reference database

[Formula E.2](#). presumes a more or less normal distribution of response factors, which is not exhibited for all chromatographic detection methods. The database of response factors used to calculate a UF according to this equation should be described and reviewed to establish whether the resulting UF is sufficiently conservative to properly account for low response factor analytes. In certain circumstances, alternate means of establishing the UF can be considered and justified if adopted.

[Formula \(E.2\)](#) is equivalent to equations proposed by PQRI and Jordi (see References [41] and [46]).

When the variation in responses factors is large relative to the mean response factor (e.g. std = 0,9 X mean), the variation in response factors is so large that although a UF can be calculated, its scientific validity becomes questionable. For example, although a UF > 10 can be calculated, the reality of a UF as large as 10 (or larger) is that the quantification method being used is inherently inaccurate and thus might not be appropriate for the purpose of producing the data that is the foundation of a toxicological risk assessment. Additionally, the use of a large value for UF could produce an adjusted AET that is so small that it cannot be achieved by the specified analytical method; that is, the method's limit of detection (LoD) is greater than the AET. In these cases, while it is possible to establish an adjusted AET, it is inappropriate to do so. The AET concept should not be applied in these cases and consideration should be given to further improvement of the method before it is used for the purpose of quantification supporting toxicological risk assessment.

In cases where the standard deviation is greater than or equal to the mean (i.e. the RSD ≥ 1), a UF cannot be calculated via [Formula E.2](#), as the result is either infinity or a negative number. Clearly an analytical method with this much variation in response factors is not optimal for the purpose of reporting data that is the foundation of a toxicological risk assessment. Optimization of the method to reduce response factor variation should be considered.

In cases where the variation in response factors among extractables cannot be established or where the variation is established to be large, the value of UF can be so large (e.g. UF values of 10 or greater) that the adjusted AET is so low that the AET concept has little practical value (e.g. the analytical method's LoD or LoQ are greater than the AET). In such cases, it is necessary that all the compounds associated with all observed analytical responses obtained by the screening analyses be identified and quantified, as all the observed analytical responses could be greater than the AET. Optimization of the method to reduce response factor variation should be considered in such cases.

It is noted that screening for extractables is typically accomplished via the use of orthogonal and complementary analytical methods, for example, GC-MS and LC-MS. The use of multiple analytical methods can reduce response factor variation and may be considered in the determination of the necessary UF that is then applied to all the complementary methods.

In any event and in all circumstances, the use of the uncertainty factor, the value of the uncertainty factor that is used, and the means by which the uncertainty factor was established should always be justified.

E.4, Example C.2, paragraph after fourth indent

Replace the paragraph with the following:

ISO 10993-18:2020/DAM 1:2021(E)

Note that 20 µg/d for 31 d would be an exposure of 620 µg, 10 µg/d for 365 d would be an exposure of 3 650 µg, and 1,5 µg/d for 3 650 d would be an exposure of 5 475 µg. Each of these theoretical extreme approaches would therefore be less conservative.

**iTeh STANDARD PREVIEW
(standards.iteh.ai)**

[SIST EN ISO 10993-18:2020/oprA1:2021
https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-
d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021](https://standards.iteh.ai/catalog/standards/sist/8bc2d68b-d5a7-4486-97b5-d04e5551559a/sist-en-iso-10993-18-2020-opra1-2021)