

SLOVENSKI STANDARD SIST EN ISO 7405:2009

01-marec-2009

Nadomešča:

SIST EN ISO 7405:2000

Zobozdravstvo - Ovrednotenje biokompatibilnosti medicinskih pripomočkov v zobozdravstvu (ISO 7405:2008)

Dentistry - Evaluation of biocompatibility of medical devices used in dentistry (ISO 7405:2008)

Zahnheilkunde - Beurteilung der Biokompatibilität von in der Zahnheilkunde verwendeten Medizinprodukten (ISO 7405:2008) standards.iteh.ai)

Art dentaire - Évaluation de la biocompatibilité des dispositifs médicaux utilisés en art dentaire (ISO 7405:2008) andards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

Ta slovenski standard je istoveten z: EN ISO 7405:2008

ICS:

11.060.01 Zobozdravstvo na splošno Dentistry in general
 11.100.20 Biološko ovrednotenje Biological evaluation of medicinskih pripomočkov medical devices

SIST EN ISO 7405:2009 en

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009

https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

EUROPEAN STANDARD

EN ISO 7405

NORME EUROPÉENNE EUROPÄISCHE NORM

December 2008

ICS 11.060.10: 11.100

Supersedes EN ISO 7405:1997

English Version

Dentistry - Evaluation of biocompatibility of medical devices used in dentistry (ISO 7405:2008)

Art dentaire - Évaluation de la biocompatibilité des dispositifs médicaux utilisés en art dentaire (ISO 7405:2008)

Zahnheilkunde - Beurteilung der Biokompatibilität von in der Zahnheilkunde verwendeten Medizinprodukten (ISO 7405:2008)

This European Standard was approved by CEN on 5 December 2008.

CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN Management Centre has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Bulgaria, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland and United Kingdom.

SIST EN ISO 7405:2009

https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009



EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

Management Centre: rue de Stassart, 36 B-1050 Brussels

EN ISO 7405:2008 (E)

Contents	Pag
Foreword	

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009

EN ISO 7405:2008 (E)

Foreword

This document (EN ISO 7405:2008) has been prepared by Technical Committee ISO/TC 106 "Dentistry" in collaboration with Technical Committee CEN/TC 55 "Dentistry" the secretariat of which is held by DIN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by June 2009, and conflicting national standards shall be withdrawn at the latest by June 2009.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN [and/or CENELEC] shall not be held responsible for identifying any or all such patent rights.

This document supersedes EN ISO 7405:1997.

According to the CEN/CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland and the United Kingdom.

iTeh STANDARD PREVIEW Endorsement notice

The text of ISO 7405:2008 has been approved by CEN as a EN ISO 7405:2008 without any modification.

SIST EN ISO 7405:2009 https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009

https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

INTERNATIONAL STANDARD

ISO 7405

Second edition 2008-12-15

Dentistry — Evaluation of biocompatibility of medical devices used in dentistry

Art dentaire — Évaluation de la biocompatibilité des dispositifs médicaux utilisés en art dentaire

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009 https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009



ISO 7405:2008(E)

PDF disclaimer

This PDF file may contain embedded typefaces. In accordance with Adobe's licensing policy, this file may be printed or viewed but shall not be edited unless the typefaces which are embedded are licensed to and installed on the computer performing the editing. In downloading this file, parties accept therein the responsibility of not infringing Adobe's licensing policy. The ISO Central Secretariat accepts no liability in this area.

Adobe is a trademark of Adobe Systems Incorporated.

Details of the software products used to create this PDF file can be found in the General Info relative to the file; the PDF-creation parameters were optimized for printing. Every care has been taken to ensure that the file is suitable for use by ISO member bodies. In the unlikely event that a problem relating to it is found, please inform the Central Secretariat at the address given below.

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009

https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009



COPYRIGHT PROTECTED DOCUMENT

© ISO 2008

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

Published in Switzerland

Contents Page

Forew	ord	iv
Introdu	uction	v
1	Scope	1
2	Normative references	
3	Terms and definitions	2
4 4.1 4.2	Categorization of medical devices Categorization by nature of contact Categorization by duration of contact	3
5 5.1 5.2 5.3 5.4 5.5	Biological evaluation process General Selection of tests and overall assessment Selection of test methods Types of test Re-evaluation of biocompatibility	4 4 5
6 6.1 6.2 6.3 6.4 6.5	Test procedures specific to dental materials	8 10
Annex	63f9bc02b0b5/sist-en-iso-7405-2009 A (informative) Types of test to be considered for evaluation of biocompatibility of medical devices used in dentistry	23
Annex	B (informative) Dentine barrier cytotoxicity test	25
Annex	C (informative) Acute toxicity testing	32
Bibliog	graphy	33

ISO 7405:2008(E)

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 7405 was prepared by Technical Committee ISO/TC 106, Dentistry.

This second edition cancels and replaces the first edition (ISO 7405:1997) which has been technically revised. The following changes have been made:

- a) addition of dentine barrier cytotoxicity test to Annex B;
- b) improved description of test methods; SIST EN ISO 7405;2009 https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-
- c) updated cross-references to ISO 10993 series. b0b5/sist-en-iso-7405-2009

ISO 7405:2008(E)

Introduction

This International Standard concerns the evaluation of the biocompatibility of medical devices used in dentistry. It is to be used in conjunction with the ISO 10993 series of standards. This International Standard contains special tests, for which ample experience exists in dentistry and which acknowledge the special needs of dentistry.

Only test methods for which the members of the committee considered there was sufficient published data have been included. In recommending test methods, the need to minimize the use of animals was given a high priority. It is essential that the decision to undertake tests involving animals be reached only after a full and careful review of the evidence indicating that a similar outcome cannot be achieved by other types of test. In order to keep the number of animals required for tests to an absolute minimum, consistent with achieving the objective indicated, it can be appropriate to conduct more than one type of test on the same animal at the same time, e.g. pulp and dentin usage test and pulp capping test. However, in accordance with ISO 10993-2 these tests are performed both in an efficient and humane way. On all occasions when animal testing is undertaken, such tests are conducted empathetically and according to standardized procedures as described for each test.

This International Standard does not explicitly describe test methods for occupationally related risks.

Annexes B and C are included to encourage the development of *in vitro* and *ex vivo* test methods which will further reduce the use of animals in the evaluation of the biocompatibility of medical devices used in dentistry.

(standards.iteh.ai)

SIST EN ISO 7405:2009 https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7405:2009

https://standards.iteh.ai/catalog/standards/sist/03eadc18-b53c-4c81-8ecc-63f9bc02b0b5/sist-en-iso-7405-2009

Dentistry — Evaluation of biocompatibility of medical devices used in dentistry

1 Scope

This International Standard specifies test methods for the evaluation of biological effects of medical devices used in dentistry. It includes testing of pharmacological agents that are an integral part of the device under test.

This International Standard does not cover testing of materials and devices that do not come into direct or indirect contact with the patient's body.

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.

(Standards.iteh.ai)

ISO 1942, Dentistry — Vocabulary

ISO 6344-1, Coated abrasives — Grain size analysis — Part 1: Grain size distribution test

ISO 10993-1, Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process

ISO 10993-2, Biological evaluation of medical devices — Part 2: Animal welfare requirements

ISO 10993-3, Biological evaluation of medical devices — Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity

ISO 10993-5, Biological evaluation of medical devices — Part 5: Tests for in vitro cytotoxicity

ISO 10993-6, Biological evaluation of medical devices — Part 6: Tests for local effects after implantation

ISO 10993-10¹⁾, Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization

ISO 10993-11, Biological evaluation of medical devices — Part 11: Tests for systemic toxicity

ISO 10993-12:2007, Biological evaluation of medical devices — Part 12: Sample preparation and reference materials

ISO 14971, Medical devices — Application of risk management to medical devices

_

¹⁾ To be published.