

SLOVENSKI STANDARD SIST EN ISO 7199:2014

01-oktober-2014

Nadomešča:

SIST EN 12022:2000

Vsadki (implantati) za srce in ožilje ter umetni organi - Izmenjevalniki krvnih plinov (ISO 7199:2009 + Amd 1:2012)

Cardiovascular implants and artificial organs - Blood-gas exchangers (oxygenators) (ISO 7199:2009 + Amd 1:2012)

Kardiovaskuläre Implantate und künstliche Organe - Blutgasaustauscher (Oxygenatoren) (ISO 7199:2009 + Amd 1:2012) (standards.iteh.ai)

Implants cardiovasculaires et organes artificiels DÉchangeurs gaz/sang extracorporels (oxygénateurs) (ISO 2019:2009 Amd 2012) /sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014

Ta slovenski standard je istoveten z: EN ISO 7199:2014

ICS:

11.040.40 Implantanti za kirurgijo,

protetiko in ortetiko

Implants for surgery, prosthetics and orthotics

SIST EN ISO 7199:2014

en

SIST EN ISO 7199:2014

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7199:2014

 $https://standards.iteh.ai/catalog/standards/sist/b\overline{d4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014}$

EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM **EN ISO 7199**

August 2014

ICS 11.040.40

Supersedes EN 12022:1999

English Version

Cardiovascular implants and artificial organs - Blood-gas exchangers (oxygenators) (ISO 7199:2009 + Amd 1:2012)

Implants cardiovasculaires et organes artificiels -Échangeurs gaz/sang extracorporels (oxygénateurs) (ISO 7199:2009 + Amd 1:2012) Kardiovaskuläre Implantate und künstliche Organe -Blutgasaustauscher (Oxygenatoren) (ISO 7199:2009 + Amd 1:2012)

This European Standard was approved by CEN on 17 July 2014.

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-CENELEC 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-CENELEC Management Centre has the same status as the official versions.

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

SIST EN ISO 7199:2014

https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014



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

CEN-CENELEC Management Centre: Avenue Marnix 17, B-1000 Brussels

EN ISO 7199:2014 (E)

Contents	Page
Foreword	3
Annex ZA (informative) Relationship between this European Standard and the Essential	
Requirements of FII Directive 93/42/FFC on medical devices	4

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7199:2014 https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014

EN ISO 7199:2014 (E)

Foreword

The text of ISO 7199:2009 + Amd 1:2012 has been prepared by Technical Committee ISO/TC 150 "Implants for surgery" of the International Organization for Standardization (ISO) and has been taken over as EN ISO 7199:2014 by Technical Committee CEN/TC 205 "Non-active medical devices" 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 February 2015, and conflicting national standards shall be withdrawn at the latest by February 2015.

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 12022:1999.

This document has been prepared under a mandate given to CEN by the European Commission and the European Free Trade Association, and supports essential requirements of EU Directive(s).

For relationship with EU Directive(s), see informative Annex ZA, which is an integral part of this document.

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, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdom.

SISTENDORSement notice https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-

The text of ISO 7199:2009 + Amd 1:2012 has been approved by CEN as EN ISO 7199:2014 without any modification.

EN ISO 7199:2014 (E)

Annex ZA (informative)

Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EEC on medical devices

This European Standard has been prepared under a mandate given to CEN by the European Commission and the European Free Trade Association to provide a means of conforming to Essential Requirements of the New Approach Directive 93/42/EEC on medical devices.

Once this standard is cited in the Official Journal of the European Union under that Directive and has been implemented as a national standard in at least one Member State, compliance with the clauses of this standard given in table ZA.1 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding Essential Requirements of that Directive and associated EFTA regulations.

Table ZA.1 — Correspondence between this European Standard and Directive 93/42/EEC on medical devices

Clause(s)/sub-clause(s) of this EN	Essential Requirements (ERs) of Directive 93/42/EEC	Qualifying remarks/Notes
4.1.1; 4.1.2; 5.2; 6.2.1; 6.2.2; 6 3 eh	3 ² ΓANDARD PREV (standards.iteh.ai)	Some redundancies, but all these EN sections address performance and sterility issues.
4.3.3; 5.2; 5.4.3	7.3 <u>SIST EN ISO 7199:2014</u>	
4.2.1; 4.2.2; 4.2.4 https://standar	dszitsh.ai/catalog/standards/sist/bd4dd8c4-a94l 7902f0fd9622/sist-en-iso-7199-2014	o-4a72-83cd-
4.1.1; 6.2.1; 6.2.2; 6.3	8.1	
6.2.1; 6.2.2; 6.3	8.3	
6.2.1; 6.2.2; 6.3	8.4	
4.1.1; 5.2.1; 6.2.1; 6.2.2; 6.2.3	8.5	
6.2.1	8.6	
6.1; 6.2.1	8.7	
4.2.4	9.1	
6.1; 6.2; 6.3	13	

WARNING — Other requirements and other EU Directives may be applicable to the product(s) falling within the scope of this standard.

SIST EN ISO 7199:2014

INTERNATIONAL STANDARD

ISO 7199

Second edition 2009-04-15

Cardiovascular implants and artificial organs — Blood-gas exchangers (oxygenators)

Implants cardiovasculaires et organes artificiels — Échangeurs gaz/sang extracorporels (oxygénateurs)

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7199:2014 https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014



ISO 7199:2009(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 7199:2014 https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014



COPYRIGHT PROTECTED DOCUMENT

© ISO 2009

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

Forev	word	iv
Intro	duction	v
1	Scope	1
2	Normative references	1
3	Terms and definitions	2
4	Requirements	3
4.1	Biological characteristics	3
4.2	Physical characteristics	
4.3	Performance characteristics	
5	Tests and measurements to determine compliance with this International Standard	4
5.1	General	
5.2	Biological characteristics	
5.3	Physical characteristics	5
5.4	Performance characteristics	
6	Information supplied by the manufacturer	8
6.1	Information to be given on the oxygenator	8
6.2	Information to be given on the packaging	8
6.3	Information to be given in the accompanying documents	9
6.4	Information to be given in the accompanying documents in a prominent form	
7	SIST EN ISO 7199:2014 Packaging .https://standards.itch.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-	
-	\mathcal{E}	
Riblia	ography 7902f0fd9622/sist-en-iso-7199-2014	11

ISO 7199:2009(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 7199 was prepared by Technical Committee ISO/TC 150, *Implants for surgery*, Subcommittee SC 2, Cardiovascular implants and extracorporeal systems.

This second edition cancels and replaces the first edition (ISO 7199:1996), which has been technically revised. (standards.iteh.ai)

SIST EN ISO 7199:2014 https://standards.iteh.ai/catalog/standards/sist/bd4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014

ISO 7199:2009(E)

Introduction

This International Standard is intended to ensure that devices designed to affect the exchange of gases in support of, or as a substitution for, the normal respiratory function of the lungs have been adequately tested for both their safety and function, and that extracorporeal device characteristics are appropriately disclosed when labeling the device.

This International Standard therefore contains procedures to be used for evaluation of extracorporeal blood-gas exchangers (oxygenators). Type test procedures for determination of the gas transfer, blood cell damage and heat exchanger performance are described, although limits for these characteristics are not specified. Ready identification of the performance characteristics should, however, assist the user in the selection of an oxygenator that will suit the needs of the patient.

This International Standard also includes minimum reporting requirements, which will allow the user to compare performance characteristics of oxygenators of different designs in a standard way.

This International Standard makes reference to other International Standards in which methods for determination of characteristics common to medical devices can be found.

No provisions have been made for quantification of microbubble generation or for non-formed elements of bovine blood because there currently is no consensus regarding satisfactorily reproducible test methods.

Requirements for animal and clinical studies have not been included in this International Standard. Such studies may be parts of a manufacturer's quality system.

This International Standard contains only those requirements that are specific to oxygenators. Non-specific requirements are covered by references to other International Standards listed in the normative references section. Since non-toxicity is anticipated to be the subject of a future horizontal/level 1 standard, this International Standard does not cover non-toxicity.

SIST EN ISO 7199:2014

iTeh STANDARD PREVIEW (standards.iteh.ai)

SIST EN ISO 7199:2014

 $https://standards.iteh.ai/catalog/standards/sist/b\overline{d4dd8c4-a94b-4a72-83cd-7902f0fd9622/sist-en-iso-7199-2014}$