

## **SLOVENSKI STANDARD** SIST EN ISO 10555-5:2000

01-januar-2000

## Sterilni žilni katetri za enkratno uporabo - 5. del: Periferni katetri z notranjo iglo (ISO 10555-5:1996)

Sterile, single-use intravascular catheters - Part 5: Over-needle peripheral catheters (ISO 10555-5:1996)

Sterile intravaskuläre Katheter zur einmaligen Verwendung - Teil 5: Periphere Katheter mit innen liegender Kanüle (ISQ 10555-5:1996) D PREVIEW

Cathéters intravasculaires stériles, non réutilisables - Partie 5: Cathéters périphériques a aiguille interne (ISO 10555-5:1996)<sub>SIST EN ISO 10555-5:2000</sub>

https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588-

Ta slovenski standard je istoveten z: EN ISO 10555-5-2000

ICS:

11.040.25 Injekcijske brizge, igle in katetri

Syringes, needles an catheters

SIST EN ISO 10555-5:2000

en

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

SIST EN ISO 10555-5:2000 https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588ff6a8aa94abb/sist-en-iso-10555-5-2000

#### SIST EN ISO 10555-5:2000

#### EUROPEAN STANDARD

## EN ISO 10555-5

#### NORME EUROPÉENNE

## EUROPÄISCHE NORM

July 1997

ICS 11.040.20

a Ese

Descriptors: see ISO document

English version

#### Sterile, single-use intravascular catheters - Part 5: Over-needle peripheral catheters (ISO 10555-5:1996)



SIST EN ISO 10555-5:2000

https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588-

ff6a8aa94abb/sist-en-iso-10555-5-2000

This European Standard was approved by CEN on 1997-06-09. 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 Central Secretariat or to any CEN member.

The European Standards exist 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 Central Secretariat has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Czech Republic, Denmark, Finland, France, Germany, Greece, Iceland, Ireland, Italy, Luxembourg, Netherlands, Norway, Portugal, Spain, Sweden, Switzerland and United Kingdom.

## CEN

European Committee for Standardization Comité Européen de Normalisation Europäisches Komitee für Normung

Central Secretariat: rue de Stassart, 36 B-1050 Brussels

Ref. No. EN ISO 10555-5:1997 E

2 . .

Page 2 EN ISO 10555-5:1997

#### Foreword

The text of the International Standard from Technical Committee ISO/TC 84 "Medical devices for injections" of the International Organization for Standardization (ISO) has been taken over as a European Standard by Technical Committee CEN/TC 205 "Non-active medical devices", the secretariat of which is held by BSI.

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 January 1998, and conflicting national standards shall be withdrawn at the latest by January 1998.

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

**Endorsement notice** 

The text of the International Standard ISO 10555-5:1996 has been approved by CEN as a

NOTE: Normative references to international Standards are listed in annex ZA (normative).

ENTRA CITA ICOTTA OS TALES

3 3330

iTeh STANDARD PREV

European Standard without any modification S. Iten.al)

1990 - **1**9e



#### Annex ZA (normative) Normative references to international publications with their relevant European publications

This European Standard incorporates by dated or undated reference, provisions from other publications. These normative references are cited at the appropriate places in the text and the publications are listed hereafter. For dated references, subsequent amendments to or revisions of any of these publications apply to this European Standard only when incorporated in it by amendment or revision. For undated references the latest edition of the publication referred to applies.

Publication	Year	Title	EN	Year
ISO 594-1	1986	Conical fittings with à 6 % (Luer) taper for syringes, needles and certain other medical equipment - Part 1: General requirements	EN 20594-1	1993
ISO 9626	1991	Stainless steel needle tubing for the manufacture of medical devices	EN ISO 9626	1995
ISO 10555-1	1995	Sterile, singlesusenintravasculareh.ai) catheters - Part 1: General requirements <u>SIST EN ISO 10555-5:2000</u> https://standards.iteh.ai/catalog/standards/sist/dba10dcb-58 ff6a8aa94abb/sist-en-iso-10555-5-2000	EN ISO 10555-1 33-4bc6-b588-	1996

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

SIST EN ISO 10555-5:2000 https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588ff6a8aa94abb/sist-en-iso-10555-5-2000

# INTERNATIONAL STANDARD



First edition 1996-06-15

## Sterile, single-use intravascular catheters —

## Part 5:

Over-needle peripheral catheters

## (standards.iteh.ai)

Cathéters intravasculaires stériles, non réutilisables — Partie 5: Cathéters périphériques à aiguille interne https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588ff6a8aa94abb/sist-en-iso-10555-5-2000



Reference number ISO 10555-5:1996(E)

#### SIST EN ISO 10555-5:2000

## 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.

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.

International Standard ISO 10555-5 was prepared by Technical Committee IEW ISO/TC 84, *Medical devices for injections*, Subcommittee SC 1, *Syringes*, needles and intravascular catheters for single use **Carcs.iten.al** 

ISO 10555 consists of the following parts, under the general title *Sterile*, single-use intravascular catheters: https://standards.iteh.ai/catalog/standards/sist/dba10dcb-5833-4bc6-b588-

— Part 1: General requirements

ff6a8aa94abb/sist-en-iso-10555-5-2000

- Part 2: Angiographic catheters
- Part 3: Central venous catheters
- Part 4: Balloon dilatation catheters
- Part 5: Over-needle peripheral catheters

Annexes A and B form an integral part of this part of ISO 10555. Annexes C, D and E are for information only.

© ISO 1996

Case Postale 56 • CH-1211 Genève 20 • Switzerland

Printed in Switzerland

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 the publisher.

International Organization for Standardization

## Sterile, single-use intravascular catheters —

## Part 5:

Over-needle peripheral catheters

## 1 Scope

This part of ISO 10555 specifies requirements for over-the-needle peripheral intravascular catheters, intended for accessing the peripheral vascular system, supplied in the sterile condition and intended for single use. **3.1 peripheral intravascular catheter:** Catheter designed for the introduction or withdrawal of liquids or devices into or from the peripheral vascular system.

**3.2 needle:** Assembly comprising at least a needle tube attached to, and communicating with, a needle hub.

## NOTE 1 Attention is drawn to ISO 11070, which specifies See figure 1. TEW requirements for accessory devices for use with intra-

(standards.i33 needle tube: Rigid tube with one end sharpened to facilitate entry into body tissue.

## **2** Normative references

SIST EN ISO 10555-5:2000

The following standards contain provisions which,

through reference in this text, constitute provisions which, through reference in this text, constitute provisions of this part of ISO 10555. At the time of publication, the editions indicated were valid. All standards are subject to revision, and parties to agreements based on this part of ISO 10555 are encouraged to investigate the possibility of applying the most recent editions of the standards indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.

ISO 594-1:1986, Conical fittings with a 6 % (Luer) taper for syringes, needles and certain other medical equipment — Part 1: General requirements.

ISO 9626:1991, Stainless steel needle tubing for the manufacture of medical devices.

ISO 10555-1:1995, Sterile, single-use intravascular catheters — Part 1: General requirements.

## 3 Definitions

For the purposes of this part of ISO 10555, the definitions given in ISO 10555-1 and the following definitions apply. **3.5 vent fitting:** Fixed or removable fitting permitting venting of air while restricting or preferably preventing the escape of blood.

**3.6 catheter unit:** Assembly comprising the catheter tube, catheter hub and any integral fittings.

See figure 1.

3.7 flashback: Blood flow into the needle hub.

## **4** Requirements

#### 4.1 General

Unless otherwise specified in this part of ISO 10555, catheters shall comply with ISO 10555-1.

## 4.2 Radio-detectability

It is recommended that catheters be radio-opaque.