TECHNICAL SPECIFICATION

First edition 2019-12

Medical devices — Transfusion set and blood bag compatibility test method

Dispositifs médicaux — Méthode d'essai de compatibilité entre les appareils de transfusion et les poches de sang

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>ISO/TS 23128:2019</u> https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-73f4c4ce8651/iso-ts-23128-2019



Reference number ISO/TS 23128:2019(E)

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>ISO/TS 23128:2019</u> https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-73f4c4ce8651/iso-ts-23128-2019



COPYRIGHT PROTECTED DOCUMENT

© ISO 2019

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

Page

Contents

Forew	vord	iv
Introd	duction	v
1	Scope	
2	Normative references	
3	Terms and definitions	
4	Materials and equipment	1
5	Labelling	2
6	Preparation6.1General6.2Insertion force equipment6.3Transfusion sets6.4Blood bags6.5Test worksheet	2 2 2 2 2 2 2 2 2 2 2
7	Test method	3
8	Analysing results	4
9	Guidance on interpretation of results	4
Annex	xe A (normative) Insertion force test equipment measures	5
Annex	xe B (informative) Example of a suitable report format	8
Biblio	ography	9

<u>ISO/TS 23128:2019</u> https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-73f4c4ce8651/iso-ts-23128-2019

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 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*. https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-

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 <u>www.iso.org/members.html</u>.

Introduction

The connection between a blood bag (as specified in ISO 3826-1, ISO 3826-3 and ISO 3826-4) and a transfusion set (ISO 1135-4 and ISO 1135-5) is provided by the bag port and transfusion set closure piercing device referred to in this document by the abbreviation 'spike'. The spike is a rigid structure with tightly defined dimensions whereas the blood bag port is of flexible material in order to accommodate the spike. Transfusion sets are compatible with a range of commercially available blood bags and vice versa. It is vitally important in setting up a blood transfusion that the force required to insert the spike into the port is not excessive. This can lead to difficulties in piercing the port septum, damage to the blood bag, leakage of its contents and injuries to bedside staff.

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>ISO/TS 23128:2019</u> https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-73f4c4ce8651/iso-ts-23128-2019

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>ISO/TS 23128:2019</u> https://standards.iteh.ai/catalog/standards/sist/bc3109c3-ea91-4fc9-b352-73f4c4ce8651/iso-ts-23128-2019

Medical devices — Transfusion set and blood bag compatibility test method

1 Scope

This document details suitable equipment, a test method, acceptance criteria and advisable limits to help to ensure that there is compatibility (by measuring the insertion force) between a transfusion set closure piercing device (referred to in this document by the abbreviation 'spike') and a blood bag outlet port.

The test procedure in its entirety is complex and beyond the scope of each of the relevant transfusion set and blood bag standards. This document was therefore developed to support the implementation of the existing standards for blood bags and transfusion sets.

The procedure described in this document can be used by manufacturers of blood bags to test the compatibility with transfusion set spikes available on the market or by manufacturers of the transfusion set spikes to test the compatibility with blood bags available on the market.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their 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 1135-4, Transfusion equipment for medical use — Part 4: Transfusion sets for single use, gravity feed ISO/TS 23128:2019

ISO 1135-5, Transfulsion/equipment for medical-usels/sisPart 59 Transfulsion sets for single use with pressure infusion apparatus 73f4c4ce8651/iso-ts-23128-2019

ISO 3826-1, Plastics collapsible containers for human blood and blood components — Part 1: Conventional containers

ISO 3826-3, Plastics collapsible containers for human blood and blood components — Part 3: Blood bag systems with integrated features

ISO 3826-4, Plastics collapsible containers for human blood and blood components — Part 4: Aphaeresis blood bag systems with integrated features

3 Terms and definitions

No terms and definitions are listed in this document.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <u>https://www.iso.org/obp</u>
- IEC Electropedia: available at <u>http://www.electropedia.org/</u>

4 Materials and equipment

The following materials and equipment shall be used for the test set-up.

- Transfusion set spike in conformance with ISO 1135-4 or ISO 1135-5.
- Blood bags conforming to ISO 3826-1, ISO 3826-3 or ISO 3826-4.

— Spike insertion test equipment including standard masses.

5 Labelling

Remove each transfusion set and blood bag system from its outer packaging and mark each device with the date, the initials of tester, the Lab Test Request (LT) number and a reference number that provides traceability to the manufacturer's product code and lot number.

6 Preparation

6.1 General

The test shall be conducted at a temperature between (23 ± 2) °C. All materials and equipment shall be stored at a temperature between (23 ± 2) °C for at least 12 hours.

6.2 Insertion force equipment

An example of suitable test equipment is shown in <u>Figure 2</u> with dimensions given in <u>Annex A</u>. Set up the equipment on a rigid, level surface. Ensure that the shaft/chuck assembly and standard masses have been calibrated. When no additional mass is used, the weight of the shaft, chuck and nuts is equal to 2 000 g, thus pushing the tested spike against the outlet port with a force of 20 N.

6.3 Transfusion sets **iTeh STANDARD PREVIEW**

If not already isolated, remove (e.g. cut off) the spikes from 3 separate unused transfusion sets to be used in the test so that at least 5 mm of the portion above the spike, see Figure 1, is available to be gripped within the chuck of the test equipment. When the spike is attached directly to the drip chamber it is alternatively possible to grip a concentric portion of the isolated drip chamber/spike assembly leaving the spike fully exposed for the insertion test. Jose 1, iso-ts-23128-2019



Figure 1 — Closure piercing device

NOTE Figure 1 is reproduced from ISO 1135-4:2015, without dimensions.

6.4 Blood bags

Select and isolate, by heat sealing, 3 fluid filled (either anticoagulant or additive solution) blood bags.

NOTE It is important that the test is carried out on plastic in contact with aqueous fluid to simulate the condition of a bag in contact with blood or blood components. Water is an acceptable alternative to anticoagulant and additive solution.

6.5 Test worksheet

Prepare the worksheet shown in <u>Annex B</u> in readiness for the commencement of testing.

7 Test method

7.1 Select the equipment in Figure A.1 and set up as in <u>6.2</u>.



iTeh Figure 2 Example of test equipment

(standards.iteh.ai)

7.2 Position the shaft (3) so that the chuck (4) is 5 cm to 10 cm above the orifice (5) and rotate the lower (collar) nut (2) to lock the shaft in place in this position.

NOTE Numbers in brackets refer to keys in <u>Annex A</u>, Figure A.1.

7.3 Open the chuck to accommodate a spike prepared in accordance with <u>6.3</u>.

7.4 Clamp the spike firmly within the chuck (see <u>6.3</u>).

WARNING — Beware of the sharp spike.

7.5 Remove the protective cover from a blood bag port prepared as in 6.4 according to the manufacturer's instructions for use.

7.6 Rotate the lower (collar) nut (2) and slowly, carefully allow the spike to be introduced into the equipment orifice (5) until the chuck prevents it from entering further.

7.7 With one hand, raise the unloaded shaft (3) 15 cm to 20 cm (or greater if the spike is attached to part of the drip chamber) above the orifice (5) and with the other hand simultaneously introduce the blood bag port vertically from beneath the orifice until it can enter no further.

7.8 Holding the port securely in place within the orifice (by gripping the blood bag film adjacent to the port and without compressing the port sleeve), lower the shaft so that the spike is introduced into the port.

7.9 With the spike in the port and whilst continuing to hold the port as in <u>7.8</u>, twist the shaft 5 times through 180°.

7.10 Assess if the spike has penetrated the port septum. If the septum has been completely penetrated record this on the worksheet (Annex B) at an insertion force of 20 N. If not, proceed to 7.11.