

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)



BSI Standards Publication

Prefilled syringes

Part 7: Packaging systems for sterilized subassembled syringes ready for filling

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

National foreword

This British Standard is the UK implementation of ISO 11040-7:2024. It supersedes BS ISO 11040-7:2015, which is withdrawn.

The UK participation in its preparation was entrusted to Technical Committee CH/212, IVDs.

A list of organizations represented on this committee can be obtained on request to its committee manager.

Contractual and legal considerations

This publication has been prepared in good faith, however no representation, warranty, assurance or undertaking (express or implied) is or will be made, and no responsibility or liability is or will be accepted by BSI in relation to the adequacy, accuracy, completeness or reasonableness of this publication. All and any such responsibility and liability is expressly disclaimed to the full extent permitted by the law.

This publication is provided as is, and is to be used at the recipient's own risk.

The recipient is advised to consider seeking professional guidance with respect to its use of this publication.

This publication is not intended to constitute a contract. Users are responsible for its correct application.

© The British Standards Institution 2024
Published by BSI Standards Limited 2024

ISBN 978 0 539 17730 5

ICS 11.040.25

Compliance with a British Standard cannot confer immunity from legal obligations.

This British Standard was published under the authority of the Standards Policy and Strategy Committee on 30 June 2024.

Amendments/corrigenda issued since publication

Date	Text affected
------	---------------



This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

ISO 11040-7

Prefilled syringes —

Part 7: Packaging systems for sterilized subassembled syringes ready for filling

Seringues préremplies —

*Partie 7: Systèmes d'emballage pour les seringues pré-assemblées
stérilisées préremplissables*

Second edition
2024-06

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)



COPYRIGHT PROTECTED DOCUMENT

© ISO 2024

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

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

Foreword	iv
Introduction	v
1 Scope	1
2 Normative references	1
3 Terms and definitions	2
4 Requirements for the packaging system	3
4.1 General.....	3
4.2 Nest and tub configuration.....	4
4.3 Nest.....	4
4.4 Tub.....	5
4.5 Insert liner.....	5
4.6 Sealing lid.....	5
4.7 Protective bag.....	5
4.8 Additional information to be provided by the manufacturer.....	6
5 Marking of the tub	6
6 Labelling	7
7 Packaging of tubs in trading units/bundles	7
Annex A (informative) Design of nests	8
Annex B (informative) Design of tubs	14
Annex C (informative) Schematic illustrations of examples of protective bag configurations	16
Annex D (informative) Design and dimensions of the protective bag for 3” and 4” tubs	18
Annex E (informative) Positioning of the tub and folding of inner protective bags for tubs packaged in a double bag configuration	21
Bibliography	25

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

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 document 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).

ISO draws attention to the possibility that the implementation of this document may involve the use of (a) patent(s). ISO takes no position concerning the evidence, validity or applicability of any claimed patent rights in respect thereof. As of the date of publication of this document, ISO had not received notice of (a) patent(s) which may be required to implement this document. However, implementers are cautioned that this may not represent the latest information, which may be obtained from the patent database available at www.iso.org/patents. ISO shall not be held responsible for identifying any or all such patent rights.

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*.

This second edition cancels and replaces the first edition (ISO 11040-7:2015), which has been technically revised.

The main changes are as follows:

- [Clause 3](#) was updated;
- former Annex B was removed because this specific method for determination of nest deflection was not commonly adopted by the industry;
- former Annex F was removed because the specific measurement method of determining the distance between the edge of the protective bag to rear end of the tub is not commonly adopted by the industry;
- a new [Annex E](#) was added to support automated processing;
- in [Annex A](#), [Annex B](#), the existing market ranges and tolerances have been revised and updated because fully automated and/or high-speed processes require smaller variations of certain tolerances;
- [Table D.1](#) for bag sizes was revised. Bags and header bags are combined, and two main groups of bag sizes have been defined based on market experience and future expectations. They were entered into [Table D.1](#) as column “recommended dimensions for 3” and 4” tub and “recommended dimensions for 4” tub”.

A list of all parts in the ISO 11040 series can be found on the ISO website.

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 www.iso.org/members.html.

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

At the start of prefilled syringe processing by the pharmaceutical industry, syringes made of tubing glass were delivered to the pharmaceutical companies in the form of a so called non-sterile “bulkware” only. The process steps such as washing, drying, inner lubrication, sealing the syringe with a closure system, sterilization, as well as filling, and closing were then performed by the pharmaceutical companies. Since their introduction to the market and with the emergence of specialized process equipment, sterilized subassembled syringes have more and more replaced the non-sterile “bulkware” to become the preferred approach for pre-filled syringe filling operations.

In the case of sterilized subassembled syringes ready for filling, responsibility for the aforementioned process steps relevant to the injectable product lies within the manufacturer of the subassembled syringes. Following the assembly of the needle shield on syringes with a staked needle or tip caps for the Luer cone version, the subassembled syringes are placed into so called nests. The nests, in turn, are placed into a plastic tub. The syringes in the nest are protected by means of an insert liner and the tub itself is sealed by a sealing lid (which is currently, and so far, primarily achieved using a porous material). Thus, the tub properly sealed with the sealing lid represents the “sterile barrier system”. The sealed tub is then wrapped into a sealable bag and, thus, ready for sterilization, which is currently, and so far, primarily performed using ethylene oxide.

In this form, the sterilized subassembled syringes ready for filling are delivered to the pharmaceutical companies in a sterile condition where they are processed on suitable machines.

The packaging design and material maintain sterility and should be compatible with the process of the customer. The packaging characteristics, material, thickness, shape, and resistance to deformations among others are such that they maintain, up to the point of use, the integrity of the product and a validated barrier against particulate and bacterial contamination.

The objective is to support the development of standardized equipment with automatic debagging process steps for aseptic processing.

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

This is a preview of BS ISO 11040-7:2024. [Click here to purchase the full version from the ANSI store.](#)

Prefilled syringes —

Part 7: Packaging systems for sterilized subassembled syringes ready for filling

1 Scope

This document specifies a packaging system that is used to deliver sterilized subassembled syringes ready for filling in tubs and nests.

Downstream processes (processes after filling such as in house/outside transport, reprocessing) can result in specific requirements on the packaging system used to deliver sterilized subassembled syringes ready for filling. However, these requirements are not within the scope of this document.

NOTE 1 Glass barrels and sterilized subassembled syringes ready for filling, plunger stoppers, and plastic barrels for injectables are specified in ISO 11040-4, ISO 11040-5 and ISO 11040-6.

NOTE 2 ISO 11607-2 addresses validation requirements of sealing and packaging processes for medical devices.

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 11040-1, *Prefilled syringes — Part 1: Glass cylinders for dental local anaesthetic cartridges*

ISO 11040-2, *Prefilled syringes — Part 2: Plunger stoppers for dental local anaesthetic cartridges*

ISO 11040-3, *Prefilled syringes — Part 3: Seals for dental local anaesthetic cartridges*

ISO 11040-4, *Prefilled syringes — Part 4: Glass barrels for injectables and sterilized subassembled syringes ready for filling*

ISO 11040-5, *Prefilled syringes — Part 5: Plunger stoppers for injectables*

ISO 11040-6, *Prefilled syringes — Part 6: Plastic barrels for injectables and sterilized subassembled syringes ready for filling*

ISO 11040-8, *Prefilled syringes — Part 8: Requirements and test methods for finished prefilled syringes*

ISO 11138-1, *Sterilization of health care products — Biological indicators — Part 1: General requirements*

ISO 11138-2, *Sterilization of health care products — Biological indicators — Part 2: Biological indicators for ethylene oxide sterilization processes*

ISO 11138-3, *Sterilization of health care products — Biological indicators — Part 3: Biological indicators for moist heat sterilization processes*

ISO 11138-4, *Sterilization of health care products — Biological indicators — Part 4: Biological indicators for dry heat sterilization processes*