Sterile hypodermic syringes for single use —

Part 1: Syringes for manual use

Seringues hypodermiques stériles, non réutilisables —
Partie 1: Seringues pour utilisation manuelle
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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 on 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 the following URL: www.iso.org/iso/foreword.html.

This document was prepared by Technical committee ISO/TC 84, Devices for administration of medicinal products and catheters.

This second edition cancels and replaces the first edition (ISO 7886-1:1993), which has been technically revised. It also incorporates the Technical corrigendum ISO 7886-1:1993/Cor.1:1995.

The main changes to the previous edition are the following:

a) clarified the Scope, e.g. excluding single-use syringes made of glass;
b) added new Normative references;
c) added new terms and definitions;
d) clarified the drawing to illustrate the component of the syringe;
e) included general requirements;
f) revised test methods for syringes;
g) revised the labelling requirement;
h) clarified the type of lubricant for the different types of syringes;
i) replaced Annex E (informative): Examples of test methods for incompatibility between syringes and injection fluids with Annex E (informative): Test method for the determination of forces required to operate the piston;
j) added Annex F (informative): Test method for the quantity of silicone;
k) informative annex on materials has been deleted.

A list of all parts in the ISO 7886 series can be found on the ISO website.
Introduction

The ISO 7886 series covers hypodermic syringes primarily intended for human use and provides performance and testing requirements. It permits broader variation in design so as not to limit innovation and methods of packaging. Its appearance and layout are consistent with other related standards which are designed to be more performance-based compared to design prescriptive.

General requirements as design guidelines for manufacturers are introduced in this document. Several limits for requirements which are historic based but confirmed in practice for many years have been kept.

Materials to be used for the construction and lubrication of sterile syringes for single use are not specified as their selection will depend to some extent upon the design, process of manufacture and sterilization method employed by individual manufacturers. The materials of the syringe should be compatible with injection fluids. If this is not the case, the attention of the user should be drawn to the exception by labelling on unit packaging. It is not practicable to specify a universally acceptable test method for incompatibility, as the only conclusive test is that an individual specific injection fluid is compatible with a specific syringe.

Manufacturers of pharmaceuticals use solvents in injectable preparations. Such solvents should be tested by the manufacturer of the injectable preparation for any possible incompatibility with the materials frequently used in syringe construction. If an incompatibility is identified, the injection fluid should be suitably labelled. The impossibility of testing any one injection fluid with all available syringes is recognized and it is strongly recommended that regulatory authorities and relevant trade associations should recognize the problem and take appropriate measures to assist manufacturers of injectable preparations.

Syringes should be manufactured and sterilized in accordance with recognized national or international codes of good manufacturing practice for medical devices.

The sampling plans for inspection selected for the ISO 7886 series are intended to verify the design at a high confidence level. The sampling plans for inspection do not replace the more general manufacturing quality systems requirements that appear in standards on quality systems, for example the ISO 9000 series and ISO 13485.

Manufacturers are expected to follow a risk-based approach and employ usability engineering during the design, development and manufacture of syringes.

Guidance on transition periods for implementing the requirements of ISO 7886 (all parts) is given in ISO/TR 19244.
Sterile hypodermic syringes for single use —

Part 1:
Syringes for manual use

1 Scope

This document specifies requirements and test methods for verifying the design of empty sterile single-use hypodermic syringes, with or without needle, made of plastic or other materials and intended for the aspiration and injection of fluids after filling by the end-users. This document does not provide requirements for lot release. The syringes are primarily for use in humans.

Sterile syringes specified in this document are intended for use immediately after filling and are not intended to contain the medicament for extended periods of time.

It excludes syringes for use with insulin (see ISO 8537), single-use syringes made of glass, syringes for use with power-driven syringe pumps, syringes pre-filled by the manufacturer, and syringes intended to be stored after filling (e.g. in a kit for filling by a pharmacist).

Hypodermic syringes without a needle specified in this document are intended for use with hypodermic needles specified in ISO 7864.

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 15223-1:2016, Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements

ISO 23908, Sharps injury protection — Requirements and test methods — Sharps protection features for single-use hypodermic needles, introducers for catheters and needles used for blood sampling

ISO 80369-7, Small-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for intravascular or hypodermic applications

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

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


— ISO Online browsing platform: available at http://www.iso.org/obp

3.1 nominal capacity
capacity of the syringe as designated by the manufacturer

EXAMPLE 1 ml, 5 ml, 50 ml
3.2 **graduated capacity**
volume of water, at 18 °C to 28 °C, expelled from the syringe when the fiducial line on the piston traverses a given scale interval or intervals

3.3 **total graduated capacity**
capacity of the syringe at the graduation line furthest from the zero graduation line

3.4 **maximum usable capacity**
capacity of the syringe when the piston is drawn back to its furthest functional position

3.5 **fiducial line**
leading edge on the plunger stopper that is in contact with and perpendicular to the syringe barrel and aligns with the zero marking on the syringe barrel when the piston is fully inserted

3.6 **unit packaging**
packaging which has direct contact with the device and maintains the sterility of the product

3.7 **user packaging**
packaging designed to contain one or more unit packages or self-contained syringe units

Note 1 to entry: Self-contained syringe units can be packed in multiple unit packs.

3.8 **two-piece syringe**
syringe assembly comprises the barrel and piston, whereas plunger and plunger stopper form one component made of the same material

3.9 **three-piece syringe**
syringe assembly includes the barrel and piston, whereas plunger and plunger stopper are two separate components of different materials

3.10 **nozzle cap**
sheath intended to physically protect the nozzle prior to use

3.11 **plunger stopper**
component connected to the leading end of the plunger and seals the open end of the syringe barrel

3.12 **self-contained syringe**
syringe with protective end caps [i.e. plunger cap, and nozzle cap or needle cap (3.17)] intended to maintain the sterility of the interior of the syringe

3.13 **dead space**
residual volume of fluid left in syringe when the plunger stopper (3.11) is fully depressed

3.14 **multiple unit pack**
multiple syringes packaged with a single seal that maintains the sterility of the product
3.15 piston
assembled component of plunger and \textit{plunger stopper} (3.11)

3.16 barrel flanges
flanges that protrude from the barrel (also referred to as finger grips) to provide the user an ergonomic means of gripping the syringe during injection

3.17 needle cap or shield
sheath intended to physically protect the needle prior to use

3.18 plunger
device component which advances the \textit{plunger stopper} (3.11) to deliver the medicinal product

4 Nomenclature

The nomenclature for the components of hypodermic syringes for single use is shown in \textit{Figure 1}. 
Key

1  needle cap or shield (if used)  10  plunger cap
2  nozzle cap  11  barrel flanges (finger grips)
3  nozzle lumen  12  fiducial line
4  nozzle  13  nominal capacity
5  barrel  14  graduation lines
6  plunger stopper (3-piece only)  15  zero line
7  seals  16  needle tube
8  plunger  17  hub
9  push-button

NOTE The figure is intended to be illustrative of the components of a syringe. The plunger stopper/plunger assembly can or cannot be of integral construction and can or cannot incorporate more than one seal.

Figure 1 — Schematic representation of hypodermic syringe for single use
5 General requirements

The general requirements are considered to be design inputs for manufacturers.

a) Syringes shall be free from defects affecting appearance, safety and performance for their intended use. Syringes with integrated or add-on sharps protection shall comply with ISO 23908.
   — The syringe’s barrel flanges shall be of adequate size, shape and strength for the intended purpose. The design specifications for the barrel flanges shall be determined through risk analysis and confirmed through usability validation testing.
   — The materials shall not cause the syringes to yield, under conditions of normal use, significant amounts of toxic substances and shall permit them to satisfy the appropriate national requirements or regulations for freedom from pyrogenic materials and abnormal toxicity.
   — Materials used in the construction of the wall of the syringe barrel shall have sufficient clarity to enable dosages to be read without difficulty.
   — The standard does not specify materials to be used for the construction and lubrication of sterile syringes with or without needles for single use, because their selection will depend, to some extent, upon the manufacturers specific syringe design, process of manufacture and sterilization method.

b) The design and validation of the packaging shall take into consideration the final use of the syringe and the storage and shipping conditions and the defined shelf life.

6 Extraneous matter

6.1 General

The surfaces of the syringe that come in contact with injection fluids during normal use shall be free from particles and extraneous matter.

NOTE Compliance with this requirement will be determined through inspection by an individual with normal vision (or corrected-to-normal vision), without magnification.

6.2 Limits for acidity or alkalinity

Exposure of distilled water to the finished syringe product shall not change its pH value by more than one unit.

Compliance with this requirement shall be demonstrated by preparing the solutions described in Annex A. The results shall show that the pH value of the syringe assessment fluid is within one pH unit of the pH value of the control fluid.

The pH value of both solutions may be determined with a laboratory potentiometric pH meter using a general purpose electrode.

6.3 Limits for extractable metals

Exposure of distilled water to the finished syringe product shall not change its content of metals by more than a combined total of 5 mg/kg of lead, tin, zinc and iron; the cadmium content shall be less than 0,1 mg/kg.

Compliance with this requirement shall be demonstrated by preparing the solutions described in Annex A and testing them using a recognized micro-analytical method, for example, by an atomic absorption method or by an inductively coupled plasma mass spectrometry method (ICP).