
Pen systems —

Part 2:

Plungers and discs for pen-injectors for
medical use

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Systèmes de stylos-injecteurs —
Partie 2: Bouchons-pistons et rondelles d'étanchéité pour stylos-injecteurs
à usage médical

ISO 13926-2:1999

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Printed in Switzerland

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 3.

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 13926-2 was prepared jointly by Technical Committees ISO/TC 76, *Transfusion, infusion and injection equipment for medical use* and ISO/TC 84, *Medical devices for injections*.

ISO 13926 consists of the following parts, under the general title *Pen systems* :

- *Part 1: Glass cylinders for pen-injectors for medical use*
- *Part 2: Plungers and discs for pen-injectors for medical use*

Annexes A, B and C form an integral part of this part of ISO 13926.

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Pen systems —

Part 2:

Plungers and discs for pen-injectors for medical use

1 Scope

This part of ISO 13926 specifies the design, dimensions, material performance requirements and marking of plungers and discs for medical pen systems. It is applicable to primary packs used in direct contact with drugs.

NOTE The potency, purity, stability and safety of a drug during its manufacture and storage can be strongly affected by the nature and performance of the primary pack.

This part of ISO 13926 does not apply to laminated or lacquered plungers.

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2 Normative references

The following normative documents contain provisions which, through reference in this text, constitute provisions of this part of ISO 13926. For dated references, subsequent amendments to, or revisions of, any of these publications do not apply. However, parties to agreements based on this part of ISO 13926 are encouraged to investigate the possibility of applying the most recent editions of the normative documents indicated below. For undated references, the latest edition of the normative document referred to applies. Members of ISO and IEC maintain registers of currently valid International Standards.

ISO 48:1994, *Rubber, vulcanized or thermoplastic – Determination of hardness (hardness between 10 IRHD and 100 IRHD)*.

ISO 3302:1990, *Rubber – Dimensional tolerances for use with products*.

ISO 7864:1993, *Sterile hypodermic needles for single use*.

ISO 8871:1990, *Elastomeric parts for aqueous parenteral preparations*.

ISO 11040-3:1993, *Prefilled syringes – Part 3: Aluminium caps for dental local anaesthetic cartridges*.

ISO 13926-1:1998, *Pen systems – Part 1: Glass cylinders for pen-injectors for medical use*.

3 Dimensions and designation

3.1 Dimensions

The plungers shall be type PSF, with dimensions as shown in Figure 1 and given in Table 1, and for discs as shown in Figure 2 and given in Table 2.

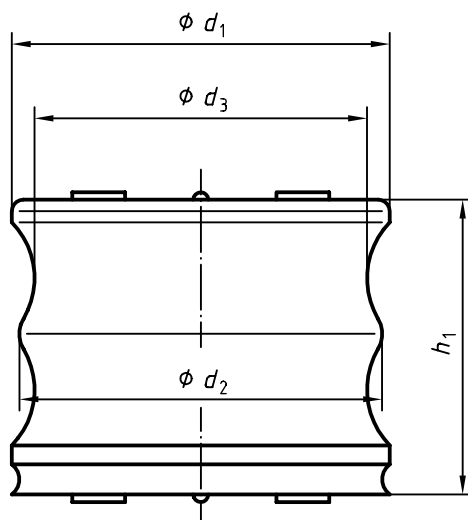


Figure 1 — Dimensions and configuration of plungers for medical pen-injectors

Table 1 — Dimensions of the plunger

Dimensions in millimetres

Nominal volume ml	Diameter (standards.iteh.ai)			Height h_1
	d_1 $\pm 0,1$	d_2 $\pm 0,1$	d_3 $\pm 0,15$	
1,5	7,2	6,9	6,4	5,5
2	9,1	8,8	8,3	8,1
2,5	9,6	9,3	8,8	8,7
3	10	9,7	9,2	11,0
4	12,5	12,1	11,7	11
5	12,35	11,95	11,55	11
6	16,6	16,15	15,7	13

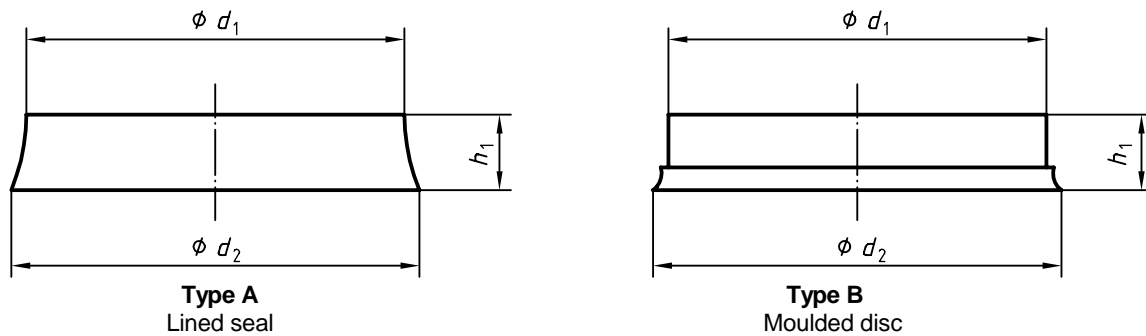


Figure 2 — Dimensions and configuration of discs for medical pen-injectors

Table 2 — Dimensions of the disc

Dimensions in millimetres

Nominal volume ml	Type	Diameter		Height
		d_1	d_2	h_1 $\pm 0,15$
1,5	A, B	7,1 min.	7,8 max.	1,5
3	A, B	$7,65 \pm 0,1$	7,85 max.	
4 to 6	B	$9,85 \pm 0,15$	10 max.	

3.2 Designation

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Plungers and discs for medical pen systems shall be designated with the appropriate block descriptor followed by a reference to this part of ISO 13926, followed by the type of disc (if applicable), followed by the nominal volume, expressed in millilitres, of the glass cylinder with which it is to be used.

EXAMPLE 1 Designation of a plunger for a glass cylinder with a nominal volume of 1,5 ml complying with the requirements in this part of ISO 13926:

Plunger ISO 13926-2 - 1,5

EXAMPLE 2 Designation of a disc type A for a glass cylinder with a nominal volume of 3 ml complying with the requirements in this part of ISO 13926:

Disc ISO 13926-2 - A - 3

4 Material

The type of elastomeric material used shall be chosen such that the plungers and discs meet the requirements specified in clause 5.

5 Requirements

5.1 Physical requirements

5.1.1 Dimensions

If not otherwise specified, general dimensional tolerances shall be in accordance with ISO 3302.

The trimmed part of the disc may be slightly conical and eccentric. The trimming edge shall not extend beyond diameter d_2 .

Discs may be knurled on one or both sides in order to avoid sticking together in a package.

In order to avoid adhesion of the plungers to each other, interrupted rings or bridges should be used as spacers in packaging. The height of the spacers shall not exceed 0,2 mm.

Sprues, if present on the surface of the plunger, shall not protrude beyond spacers.

5.1.2 Hardness

The hardness shall be agreed between manufacturer and user; hardness shall be determined in accordance with ISO 48.

5.1.3 Fragmentation (coring)

When testing discs for fragmentation in accordance with annex A, not more than three fragments per 50 piercings shall be observed.

5.1.4 Leakage

When testing discs or plungers according to annex B, no leakage of liquid from the glass cylinder shall be observed.

5.1.5 Sliding properties

When testing plungers according to annex C, the break-loose force and restarting force shall not exceed 30 N. The force to sustain continuous movement shall not exceed 15 N and there shall be no chattering¹⁾

5.2 Chemical requirements

The chemical properties of the material of the plungers and discs shall not exceed the limits specified in Table 3.

ISO 13926-2:1999
Table 3 — Chemical limits for plungers and discs
<https://standards.iteh.ai/catalog/standards/sis/007901eb-04ac-43c1-a867-3d72931b8b22/iso-13926-2-1999>

Test	Requirement	Test procedure as described in ISO 8871:1990, annex ^a
Reducing matter (oxidizables)	≤ 7 ml of $c(\text{KMnO}_4) = 2 \text{ mmol/l}$ per 20 ml	C
Heavy metals (calculated as Pb^{2+})	≤ 10 $\mu\text{g Pb}^{2+}/10 \text{ ml}$	D
Ammonium (calculated as NH_4^+)	≤ 20 $\mu\text{g NH}_4^+/10 \text{ ml}$	E
Acidity/alkalinity	≤ 1 ml of $c(\text{HCl})$ or $c(\text{NaOH}) = 5 \text{ mmol/l}$ per 20 ml	G
Residue on evaporation (total solids)	≤ 4 mg/100 ml	H
Volatile sulfides (at $\text{pH} \approx 2$)	coloration of lead acetate paper ≤ 50 $\mu\text{g Na}_2\text{S}/20 \text{ cm}^2$ rubber surface	J
Zinc (calculated as Zn^{2+})	$\text{Zn}^{2+} \leq 30 \mu\text{g}/10 \text{ ml}$	K
Conductivity	≤ 40 $\mu\text{S/cm}$	L
Turbidity	not exceeding opalescence suspension number 3	M

^a ISO 8871:1990 is under revision at present. Therefore, the requirements specified in this table could change.

1) Chatter (stick-slip) is the phenomenon of irregular motion of the plunger.

5.3 Biological requirements

The elastomeric plunger shall not release any substances which may adversely affect the therapeutic effectiveness of the injectable products or substances which may exhibit toxic, pyrogenic or haemolytic reactions.

NOTE Since biological tests are usually requested by most of the national Pharmacopoeias or related regulations of health authorities, they are mandatory for producers and users in countries where they exist. If this is not the case, reference should be made to biological tests, e.g. as described in the United States Pharmacopoeia, European Pharmacopoeia, national Pharmacopoeias or ISO 10993 series.

6 Marking

The package of plungers or discs may be marked with a designation in accordance with 3.2.

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