

STN	Injekčné systémy na báze ihiel na zdravotníckej použítie Požiadavky a skúšobné metódy Časť 3: Zásobníky a integrované dráhy tekutín (ISO 11608-3: 2022/Amd 1: 2026) Zmena A1	STN EN ISO 11608-3/A1 85 5930
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Needle-based injection systems for medical use - Requirements and test methods - Part 3: Containers and integrated fluid paths - Amendment 1 (ISO 11608-3: 2022/Amd 1: 2026)

Táto norma obsahuje anglickú verziu európskej normy.

This standard includes the English version of the European Standard.

Táto norma bola oznámená vo Vestníku ÚNMS SR č. 07/26

Obsahuje: EN ISO 11608-3:2022/A1:2026, ISO 11608-3:2022/Amd 1:2026

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EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN ISO 11608-3:2022/A1

April 2026

ICS 11.040.25

English Version

**Needle-based injection systems for medical use -
Requirements and test methods - Part 3: Containers and
integrated fluid paths - Amendment 1 (ISO 11608-
3:2022/Amd 1:2026)**

Systèmes d'injection à aiguille pour usage médical -
Exigences et méthodes d'essai - Partie 3: Conteneurs et
chemins de fluide intégrés - Amendement 1 (ISO
11608-3:2022/Amd 1:2026)

Kanülenbasierte Injektionssysteme zur medizinischen
Verwendung - Anforderungen und Prüfverfahren - Teil
3: Behälter und integrierte Flüssigkeitsbahnen -
Änderung 1 (ISO 11608-3:2022/Amd 1:2026)

This amendment A1 modifies the European Standard EN ISO 11608-3:2022; it was approved by CEN on 27 April 2026.

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EUROPEAN COMMITTEE FOR STANDARDIZATION
COMITÉ EUROPÉEN DE NORMALISATION
EUROPÄISCHES KOMITEE FÜR NORMUNG

CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

EN ISO 11608-3:2022/A1:2026 (E)

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European foreword

This document (EN ISO 11608-3:2022/A1:2026) has been prepared by Technical Committee ISO/TC 84 "Devices for administration of medicinal products and catheters" in collaboration with Technical Committee CEN/TC 205 "Non-active medical devices" the secretariat of which is held by DIN.

This Amendment to the European Standard EN ISO 11608-3:2022 shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by October 2026, and conflicting national standards shall be withdrawn at the latest by October 2026.

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Endorsement notice

The text of ISO 11608-3:2022/Amd 1:2026 has been approved by CEN as EN ISO 11608-3:2022/A1:2026 without any modification.



International Standard

ISO 11608-3

Needle-based injection systems for medical use — Requirements and test methods —

Part 3: Containers and integrated fluid paths

AMENDMENT 1

*Systèmes d'injection à aiguille pour usage médical — Exigences
et méthodes d'essai —*

Partie 3: Conteneurs et chemins de fluide intégrés

AMENDEMENT 1

**Third edition
2022-04**

**AMENDMENT 1
2026-04**

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ISO 11608-3:2022/Amd.1:2026(en)**Foreword**

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This document was prepared by Technical Committee ISO/TC 84, *Devices for administration of medicinal products and catheters*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 205, *Non-active medical devices*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

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Needle-based injection systems for medical use — Requirements and test methods —

Part 3: Containers and integrated fluid paths

AMENDMENT 1

Clause 2 Normative references

Replace “ISO 10555-1:2013” with “ISO 10555-1:2023”.

koniec náhľadu – text ďalej pokračuje v platenej verzii STN