



SLOVENSKI STANDARD

oSIST prEN ISO 7864:2026

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Sterilne podkožne igle za enkratno uporabo - Zahteve in preskusne metode (ISO/DIS 7864:2025)

Sterile hypodermic needles for single use - Requirements and test methods (ISO/DIS 7864:2025)

Sterile Injektionskanülen für den Einmalgebrauch - Anforderungen und Prüfverfahren
(ISO/DIS 7864:2025)

Aiguilles hypodermiques stériles à usage unique - Exigences et méthodes d'essai
(ISO/DIS 7864:2025)

Ta slovenski standard je istoveten z: prEN ISO 7864

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ICS:

11.040.25	Injekcijske brizge, igle in katetri	Syringes, needles and catheters
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DRAFT International Standard

ISO/DIS 7864

Sterile hypodermic needles for single use — Requirements and test methods

*Aiguilles hypodermiques stériles, non réutilisables — Exigences et
méthodes d'essai*

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