



SLOVENSKI STANDARD

oSIST prEN ISO 11607-3:2025

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Embalaža za končno sterilizirane medicinske pripomočke - 3. del: Zahteve za razvoj procesa oblikovanja, tesnjenja in sestavljanja (ISO/DIS 11607-3:2025)

Packaging for terminally sterilized medical devices - Part 3: Requirements for process development for forming, sealing and assembly (ISO/DIS 11607-3:2025)

Verpackungen für in der Endverpackung zu sterilisierende Medizinprodukte - Teil 3: Anforderungen an die Prozessentwicklung der Formgebung, Siegelung und des Zusammenstellens (ISO/DIS 11607-3:2025)

Emballages des dispositifs médicaux stérilisés au stade terminal - Partie 3: Exigences relatives à la mise au point des procédés de formage, scellage et assemblage (ISO/DIS 11607-3:2025)

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11.080.30	Sterilizirana embalaža	Sterilized packaging
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en,fr,de



DRAFT International Standard

ISO/DIS 11607-3

Packaging for terminally sterilized medical devices —

Part 3: Requirements for process development for forming, sealing and assembly

*Emballages des dispositifs médicaux stérilisés au stade
terminal —*

*Partie 3: Exigences relatives au développement des procédés de
formage, scellage et assemblage*

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