



SLOVENSKI STANDARD

oSIST prEN 868-5:2026

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Embalaža za končno sterilizirane medicinske pripomočke - 5. del: Vrečke in zvitki papirja z možnostjo tesnjenja (samolepilni) iz poroznega materiala in s plastičnimi folijami - Zahteve in preskusne metode

Packaging for terminally sterilized medical devices - Part 5: Sealable pouches and reels constructed of porous materials and plastic film - Requirements and test methods

Verpackungen für in der Endverpackung zu sterilisierende Medizinprodukte - Teil 5: Siegelfähige Klarsichtbeutel und -schläuche aus porösem Material und Kunststoffolie - Anforderungen und Prüfverfahren

Emballages des dispositifs médicaux stérilisés au stade terminal - Partie 5 : Sachets et gaines scellables constitués de matériaux poreux et de film plastique - Exigences et méthodes d'essai

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Ta slovenski standard je istoveten z: prEN 868-5

ICS:

11.080.30 Sterilizirana embalaža Sterilized packaging

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en,fr,de

EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

DRAFT
prEN 868-5

January 2026

ICS 11.080.30

Will supersede EN 868-5:2018

English Version

**Packaging for terminally sterilized medical devices - Part
5: Sealable pouches and reels constructed of porous
materials and plastic film - Requirements and test
methods**

Emballages des dispositifs médicaux stérilisés au stade
terminal - Partie 5 : Sachets et gaines scellables
constitués de matériaux poreux et de film plastique -
Exigences et méthodes d'essai

Verpackungen für in der Endverpackung zu
sterilisierende Medizinprodukte - Teil 5: Siegelfähige
Klarsichtbeutel und -schläuche aus porösem Material
und Kunststofffolie - Anforderungen und Prüfverfahren

This draft European Standard is submitted to CEN members for enquiry. It has been drawn up by the Technical Committee CEN/TC 102.

If this draft becomes a European Standard, CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration.

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Recipients of this draft are invited to submit, with their comments, notification of any relevant patent rights of which they are aware and to provide supporting documentation.

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

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