

SLOVENSKI STANDARD

SIST EN 868-6:2025

01-junij-2025

Nadomešča:
SIST EN 868-6:2017

Embalaža za končno sterilizirane medicinske pripomočke - 6. del: Papir za sterilizacijske procese z nizko temperaturo - Zahteve in preskusne metode

Packaging for terminally sterilized medical devices - Part 6: Paper for low temperature sterilization processes - Requirements and test methods

Verpackungen für in der Endverpackung zu sterilisierende Medizinprodukte - Teil 6: Papier zur Sterilisation mit Sterilisationsverfahren mit niedriger Temperatur - Anforderungen und Prüfverfahren

Emballages des dispositifs médicaux stérilisés au stade terminal - Partie 6 : Papier pour la fabrication d'emballages à usage médical pour stérilisation par l'oxyde d'éthylène ou par irradiation - Exigences et méthodes d'essai

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Ta slovenski standard je istoveten z: EN 868-6:2025

ICS:

11.080.30	Sterilizirana embalaža	Sterilized packaging
55.040	Materiali in pripomočki za pakiranje	Packaging materials and accessories

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en,fr,de

EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN 868-6

April 2025

ICS 11.080.30

Supersedes EN 868-6:2017

English Version

**Packaging for terminally sterilized medical devices - Part
6: Paper for low temperature sterilization processes -
Requirements and test methods**

Emballages des dispositifs médicaux stérilisés au stade
terminal - Partie 6 : Papier pour des procédés de
stérilisation à basse température - Exigences et
méthodes d'essai

Verpackungen für in der Endverpackung zu
sterilisierende Medizinprodukte - Teil 6: Papier für
Niedertemperatur-Sterilisationsverfahren -
Anforderungen und Prüfverfahren

This European Standard was approved by CEN on 14 March 2025.

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. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

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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

Contents	Page
European foreword	3
Introduction	5
1 Scope.....	6
2 Normative references.....	6
3 Terms and definitions	7
4 General requirements	7
5 Performance requirements and test methods	7
6 Sterilization compatibility.....	8
7 Labelling.....	9
7.1 Transport packaging	9
7.2 Labelling of individual units	9
8 Environmental declarations	9
Annex A (normative) Method for the determination of water repellency	10
Annex B (normative) Method for the determination of pore size	13
Annex C (informative) Repeatability and reproducibility of test methods	18
Annex D (informative) Environmental aspects.....	20
Bibliography	23

SIST EN 868-6:2025

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