



SLOVENSKI STANDARD

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Sterilizatorji za uporabo v medicini - Mali parni sterilizatorji - Zahteve in preskušanje

Sterilizers for medical purposes - Small steam sterilizers - Requirements and testing

Sterilisatoren für medizinische Zwecke - Dampf-Klein-Sterilisatoren - Anforderungen und Prüfung

Stériliseurs à usage médical - Petits stériliseurs à la vapeur d'eau - Exigences et essais

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ICS:

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**Sterilizers for medical purposes - Small steam sterilizers -
Requirements and testing**

Stérilisateurs à usage médical - Petits stérilisateurs à la
vapeur d'eau - Exigences et essais

Sterilisatoren für medizinische Zwecke - Dampf-Klein-
Sterilisatoren - Anforderungen und Prüfung

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