
Packaging for terminally sterilized medical devices —

Part 1: Requirements for materials, sterile barrier systems and packaging systems

AMENDMENT 1: Application of risk management

Emballages des dispositifs médicaux stérilisés au stade terminal —

*Partie 1: Exigences relatives aux matériaux, aux systèmes de barrière
stérile et aux systèmes d'emballage*

AMENDEMENT 1: Application de la gestion des risques





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