
**Sterilization of health care products —
General requirements for characterization
of a sterilizing agent and the
development, validation and routine
control of a sterilization process for
medical devices**

*Stérilisation des produits de santé — Exigences générales pour la
caractérisation d'un agent stérilisant et pour la mise au point, la
validation et la vérification de routine d'un processus de stérilisation
pour dispositifs médicaux*



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