



International Standard

ISO 15223-1

Medical devices — Symbols to be used with information to be supplied by the manufacturer —

Part 1:

General requirements

AMENDMENT 1: Addition of defined
term for authorized representative
and modified EC REP symbol to not be
country or region specific

*Dispositifs médicaux — Symboles à utiliser avec les informations
à fournir par le fabricant —*

Partie 1: Exigences générales

*AMENDEMENT 1: Ajout du terme défini représentant autorisé
(mandataire) et modification du symbole EC REP pour ne pas
être spécifique d'un pays ou d'une région*

Fourth edition
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AMENDMENT 1

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