
Prefilled syringes —

Part 4:

**Glass barrels for injectables and
sterilized subassembled syringes
ready for filling**

AMENDMENT 1

Seringues préremplies —

*Partie 4: Cylindres en verre pour produits injectables et seringues pré-
assemblées stérilisées préremplissables*

AMENDEMENT 1

ISO 11040-4:2015/Amd 1:2020

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ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Fax: +41 22 749 09 47
Email: copyright@iso.org
Website: www.iso.org

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