



International Standard

ISO 15378

Primary packaging materials for medicinal products — Particular requirements for the application of ISO 9001:2015, with reference to good manufacturing practice (GMP)

AMENDMENT 1: Climate action changes

*Articles d'emballage primaire pour médicaments — Exigences
particulières pour l'application de l'ISO 9001:2015 prenant en
considération les Bonnes Pratiques de Fabrication (BPF)*

AMENDEMENT 1: Actions relatives aux changements climatiques

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