
**Containers and accessories for
pharmaceutical preparations —**

**Part 3:
Screw-neck glass bottles (veral) for
solid and liquid dosage forms**

AMENDMENT 1

Réipients et accessoires pour préparations pharmaceutiques —

*Partie 3: Flacons en verre à bouchon à vis (veral) pour formes sèches
et liquides*

AMENDEMENT 1

ISO 11418-3:2016/Amd 1:2017

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This document was prepared by Technical Committee ISO/TC 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*.

A list of all parts in the ISO 11418 series can be found on the ISO website.

[ISO 11418-3:2016/Amd.1:2017](https://standards.iteh.ai/ISO/11418-3:2016/Amd.1:2017)

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