

Third edition
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Containers and accessories for pharmaceutical preparations —

Part 1: Drop-dispensing glass bottles

*Réipients et accessoires pour préparations pharmaceutiques —
Partie 1: Flacons compte-gouttes en verre*



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Foreword

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The committee responsible for this document is ISO/TC 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*.

This third edition cancels and replaces the second edition (ISO 11418-1:2005), which has been technically revised by

- updating [Figure 1](#) on typical drop-dispensing glass bottle and [Table 1](#) on nominal volume, brimful capacity and dimensions of drop-dispensing glass bottles,
- deleting the sizes 25 ml and 75 ml in [Table 1](#), and
- editorially revising this document.

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