
**Plastics collapsible containers
for human blood and blood
components —**

**Part 1:
Conventional containers**

*Poches en plastique souple pour le sang et les composants du sang —
Partie 1: Poches conventionnelles*





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Foreword

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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*.

This third edition cancels and replaces the second edition (ISO 3826-1:2013), which has been technically revised.

The main changes compared to the previous edition are as follows:

- in [Clause 3](#) 'Terms and definitions' four new entries have been added;
- in [Clause 4](#), the designation example has been removed;
- [Clause 5](#) 'Design' has been revised, especially regarding the pilot samples, collection and transfer tube(s), blood-taking needle and outlet port(s);
- the physical requirements in [6.2](#) have been slightly amended;
- [Clause 8](#) 'Labelling' has been reviewed and amended with barcoding information;
- the normative references in [Clause 2](#) and the Bibliography have been updated.

A list of all parts in the ISO 3826 series can be found on the ISO website.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.