
**In vitro diagnostic medical devices —
Single-use containers for the
collection of specimens from humans
other than blood**

*Dispositifs médicaux de diagnostic in vitro — Récipients à usage
unique pour le prélèvement d'échantillons d'origine humaine autres
que le sang*

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Contents

Page

Foreword	iv
1 Scope	1
2 Normative references	1
3 Terms and definitions	1
4 Materials	3
5 Filling capacity/draw volume	4
6 Graduation lines	4
7 Design	4
8 Construction	5
9 Sterility and special microbiological states	5
10 Additives	5
11 Marking and labelling	6
Annex A (normative) Tests for filling capacity and/or graduation lines for non-evacuated specimen container	8
Annex B (normative) Draw volume test for evacuated containers	9
Annex C (normative) Test for leakage from the closure of a container	11
Annex D (normative) Test for the robustness of a container that is intended for centrifugation	13
Bibliography	14

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