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Needle-based injection systems for medical use - Requirements and test methods - Part 3: Finished containers (ISO 11608-3: 2012)

Contents	Page
Foreword	3
Introduction	4
1 Scope	5
2 Normative references	5
3 Terms and definitions	5
4 Requirements	7
4.1 General	7
4.2 Freedom from leakage	8
4.2.1 All containers	8
4.2.2 Cartridges	8
4.3 Plunger force	8
4.3.1 All containers	8
4.3.2 Cartridges	8
4.4 Dimensions	8
4.4.1 All containers	8
4.4.2 Cartridges	8
4.5 Eccentricity	9
4.5.1 All containers	9
4.5.2 Cartridges	9
4.6 Visibility of the medicinal product -- All containers	9
4.7 Meniscus -- Cartridge only	9
4.8 Resealability -- All containers	9
4.9 Coring (fragmentation)	10
4.10 Container materials	10
4.10.1 All containers	10
4.10.2 Cartridges	10
4.11 Cap -- Cartridge only	10
4.12 Plunger and disc -- Cartridge only	10
4.13 Particulates -- All containers	10
4.14 Dose accuracy -- All containers	10
4.15 Deliverable volume -- All containers	10
4.16 Freedom from damage -- All containers	10
5 Test methods	11
5.1 Test apparatus	11
5.2 Test conditions	11
5.3 Test fluid	11
5.4 Plunger force	11
5.5 Leakage (attribute)	11
5.6 Dimensions	12
5.6.1 Length reference, l3	12
5.6.2 Overall diameter, d6, and plunger insertion depth, h3	12
5.6.3 Eccentricity (attribute)	12
5.6.4 Meniscus (attribute)	12
5.6.5 Resealability (attribute)	12
5.6.6 Coring (variable)	12

6	Information supplied by the manufacturer	13
	Bibliography	14