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Needle-based injection systems for medical use - Requirements and test methods - Part 1: Needle-based injection systems (ISO 11608-1:2022)

Contents	Page
European foreword	4
Foreword	5
Introduction	7
1 Scope	10
2 Normative references	10
3 Terms and definitions	11
4 Symbols	14
5 Requirements	15
5.1 General	15
5.2 System designations	16
5.3 Risk approach	17
5.4 Usability engineering	17
5.5 Uncertainty of measurement and conformance with specifications	17
5.6 General design requirements	18
5.7 Design verification	20
5.7.1 General	20
5.7.2 Primary functions requirements	20
6 Reagent and apparatus	21
6.1 General	21
6.2 Test liquid	22
6.3 Test surface for free-fall testing	22
7 Assessment of dose accuracy and other primary functions	22
7.1 General	22
7.2 Dose accuracy	22
7.2.1 General	22
7.2.2 Dosing regions	23
7.2.3 Dose settings	23
7.3 Sampling for other primary functions	25
7.4 Assessment	25
7.4.1 General	25
7.4.2 Determination of dose accuracy limits	25
7.4.3 Determination of last-dose error and last-dose accuracy limits (system designations A and C)	27
7.4.4 Calculation of dose delivery efficiency (system designations B1 and D1, user-filled)	28
7.4.5 Acceptance criteria	28
8 Preparation and operation of NISs	29
9 Test case matrix	29

10	Testing procedures	31
10.1	General	31
10.2	Normal/anticipated condition test cases	31
10.2.1	Cool, standard and warm atmosphere in-use testing	31
10.2.2	Last-dose accuracy testing (system designations A and C only)	32
10.2.3	Life-cycle testing (systems designations A and B only) — Preconditioning	32
10.3	Stressed/challenge condition test cases	32
10.3.1	Free fall testing – Preconditioning	32
10.3.2	Dry-heat storage – Preconditioning	34
10.3.3	Cold-storage - Preconditioning	34
10.3.4	Damp-heat testing (system designations A and B only) — Preconditioning	35
10.3.5	Cyclical testing (system designations A and B only) — Preconditioning	35
10.3.6	Vibration testing — Preconditioning	35
10.3.7	Transport – Preconditioning	36
10.3.8	Functional stability – Preconditioning	36
10.3.9	Fluid leakage (system designations A and B only) - Preconditioning	36
11	Inspection	37
11.1	General	37
11.2	Legibility of markings	37
11.3	Freedom from defects	37
12	Information supplied with the NIS	38
12.1	General	38
	Annex A (informative) Testing rationale	39
	Annex B (normative) One- and two-sided tolerance limit factors, k (for normally distributed data)	44
	Annex C (informative) Biological evaluation according to ISO 10993-1	55
	Annex D (informative) Functional stability	57
	Annex E (normative) Instructions for use, marking and age warning	59
	Annex F (informative) Rationale for recommended sample sizes	61
	Annex G (informative) Considerations for assessing impact on primary functions due to exposure to or contact with medicinal product	66
	Annex H (informative) Introduction of primary functions	68
	Bibliography	77