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Sterilization of health care products - Chemical indicators - Part 1: General requirements (ISO 11140-1:2014)

Contents	Page
Foreword	4
Introduction	5
1 Scope	6
2 Normative references	6
3 Terms and definitions	7
4 Categorization	9
4.1 General	9
4.2 Type 1: process indicators	9
4.3 Type 2: indicators for use in specific tests	10
4.4 Type 3: single critical process variable indicators	10
4.5 Type 4: multicritical process variable indicators	10
4.6 Type 5: integrating indicators	10
4.7 Type 6: emulating indicators	10
5 General requirements	10
6 Performance requirements	13
6.1 General	13
6.2 Type 1 indicators	14
6.3 Type 2 indicators	14
6.4 Types 3, 4, 5 and 6 indicators	14
7 Test methods	14
7.1 General	14
7.2 Off-set (transference)	14
7.3 Procedure -- Steam indicators	14
7.4 Procedure -- Dry heat indicators	15
7.5 Procedure -- EO indicators	15
7.6 Procedure -- Low temperature steam and formaldehyde indicators	16
7.7 Procedure -- Vaporized hydrogen peroxide indicators	16
8 Additional requirements for process (Type 1) indicators	17
8.1 Process indicators printed or applied on to packaging material	17
8.2 Process indicators for steam sterilization processes	17
8.3 Process indicators for dry heat sterilization processes	17
8.4 Process indicators for ethylene oxide sterilization processes	18
8.5 Process indicators for radiation sterilization processes	18
8.6 Process indicators for low temperature steam and formaldehyde sterilization processes	19
8.7 Process indicators for vaporized hydrogen peroxide sterilization processes	19
9 Additional requirements for single critical process variable (Type 3) indicators	20
10 Additional requirements for multicritical process variable (Type 4) indicators	20
11 Additional requirements for steam integrating (Type 5) indicators	21

12	Additional requirements for ethylene oxide integrating (Type 5) indicators	22
13	Additional requirements for emulating (Type 6) indicators	22
	Annex A (normative) Method for demonstrating shelf-life of the product	24
	Annex B (informative) Examples of testing indicators	25
	Annex C (informative) Rationale for the requirements for integrating indicators and the link to the requirements for biological indicators specified in ISO 11138 (all parts) and microbial inactivation	27
	formaldehyde indicators	
	Annex E (informative) Relationship of indicator and indicator system components	
35	Bibliography	
38	Annex ZA (informative) Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EC on medical devices	