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Sterilization of health care products - Biological indicators - Part 1: General requirements (ISO 11138-1:2017)

Contents	Page
European foreword	3
Foreword	5
Introduction	6
1 Scope	7
2 Normative references	7
3 Terms and definitions	7
4 General manufacturing requirements	10
4.1 Manufacturing controls	10
4.1.1 Quality management systems	10
4.1.2 Traceability	10
4.1.3 Finished product requirements	10
4.1.4 Personnel	10
4.2 Test organism	10
4.2.1 Strain	10
4.2.2 Originating inoculum for suspension	11
4.2.3 Test organism count	11
4.3 Information to be provided by the manufacturer (labelling)	11
4.4 Storage and transport	12
5 Specific manufacturing requirements	13
5.1 Suspensions	13
5.2 Carrier, primary and secondary packaging	13
5.3 Inoculated carrier	13
5.4 Biological indicators	14
5.5 Self-contained biological indicators	14
6 Determination of population and resistance	14
6.1 General resistance requirements	14
6.2 Test organism	14
6.3 Population of test organisms	14
6.4 Resistance characteristics	15
6.5 Test conditions	15
7 Culture conditions	16
7.1 Incubator	16
7.2 Growth medium	16
7.3 Incubation	16
7.4 Software validation	17
7.5 Incubation time using detection system	17
Annex A (normative) Determination of viable count	18
Annex B (normative) Determination of growth inhibition by carriers and primary packaging materials exposed to sterilization processes	20
Annex C (normative) D value determination by survivor curve method	23
Annex D (normative) D value determination by fraction negative method	27
Annex E (normative) Survival-kill response characteristics	43
Annex F (informative) Relationship between components of biological indicators	45
Bibliography	46