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Sterilization of health care products - E thylene oxide - Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices (ISO 11135-1:2007)

Contents

Page

Foreword.....	3
Introduction	4
1 Scope	6
2 Normative references	7
3 Terms and definitions	7
4 Quality management systems	14
4.1 Documentation	14
4.2 Management responsibility	14
4.3 Product realization.....	14
4.4 Measurement, analysis and improvement — Control of nonconforming product	15
5 Sterilizing agent characterization	15
5.1 Sterilizing agent	15
5.2 Microbicidal effectiveness	15
5.3 Material effects	15
5.4 Environmental considerations	15
6 Process and equipment characterization	15
6.1 Process characterization	15
6.2 Equipment characterization.....	16
7 Product definition	17
7.1 General.....	17
7.2 Product safety and performance.....	17
7.3 Microbiological quality	17
7.4 Documentation.....	18
8 Process definition.....	18
9 Validation.....	19
9.1 Installation qualification.....	19
9.2 Operational qualification.....	19
9.3 Performance qualification.....	19
9.4 Varying load configurations	21
9.5 Review and approval of validation.....	21
10 Routine monitoring and control	23
11 Product release from sterilization.....	24
12 Maintaining process effectiveness	24
12.1 General.....	24
12.2 Maintenance of equipment	24
12.3 Requalification	24
12.4 Assessment of change.....	25
Annex A (normative) Determination of lethal rate of the sterilization process — Biological indicator/bioburden approach.....	26
Annex B (normative) Conservative determination of lethal rate of the sterilization process — Overkill approach.....	28
Annex C (informative) General guidance	30
Bibliography	46
Annex ZA (informative) Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EEC Medical devices	47

Annex A (normative) Determination of lethal rate of the sterilization process -- Biological indicator/bioburden approach	26
Annex B (normative) Conservative determination of lethal rate of the sterilization process -- Overkill approach	28
Annex C (informative) General guidance	30
Bibliography	46
Annex ZA (informative) Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EEC Medical devices	47
Normative references	7