TECHNICAL SPECIFICATION

ISO/TS 13004

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Sterilization of health care products — Radiation — Substantiation of selected sterilization dose: Method VD_{max}SD

Stérilisation des produits de santé — Irradiation — Justification de la dose de stérilisation choisie: méthode VD_{max}^{SD}







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ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

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Foreword

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International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

In other circumstances, particularly when there is an urgent market requirement for such documents, a technical committee may decide to publish other types of normative document:

- an ISO Publicly Available Specification (ISO/PAS) represents an agreement between technical experts in an ISO working group and is accepted for publication if it is approved by more than 50 % of the members of the parent committee casting a vote;
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An ISO/PAS or ISO/TS is reviewed after three years in order to decide whether it will be confirmed for a further three years, revised to become an International Standard, or withdrawn. If the ISO/PAS or ISO/TS is confirmed, it is reviewed again after a further three years, at which time it must either be transformed into an International Standard or be withdrawn.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

ISO/TS 13004 was prepared by Technical Committee ISO/TC 198, Sterilization of health care products.

Introduction

This Technical Specification is intended to be used in conjunction with ISO 11137-1, *Sterilization of health care products – Radiation – Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices.* One of the activities encompassed within process definition in ISO 11137-1 is the option to select and substantiate a sterilization dose to be applied to health care products.

ISO 11137-2 includes Method VD_{max} for the substantiation of 25 kGy as a sterilization dose (termed Method VD_{max}^{25}) for product with an average bioburden less than or equal to 1 000 and Method VD_{max}^{15} for the substantiation of 15 kGy as a sterilization dose for product with an average bioburden less than or equal to 1,5.

This Technical Specification extends the methods of selection and substantiation of a sterilization dose specified in ISO 11137-2. It provides a methodology for the substantiation of selected sterilization doses of 17,5, 20, 22,5, 27,5, 30, 32,5 and 35 kGy, each of which is valid only for a specified upper limit of average bioburden.

NOTE Selected sterilization doses of 25 kGy and 15 kGy are not included in this Technical Specification. The seven methods in this Technical Specification follow the same technical steps as the methods given in ISO 11137-2 for selection and substantiation of sterilization doses of 25 kGy and 15 kGy. However, the descriptive text in this Technical Specification has been modified to better communicate the methods and hence the text occasionally differs from that in ISO 11137-2.

The method described in this Technical Specification is for substantiation of a selected sterilization dose to achieve a sterility assurance level (SAL) of 10^{-6} or less at that dose, (e.g. Method VD_{max}^{20} for a selected sterilization dose of 20 kGy). The application of the method is not limited by production batch size or production frequency, and the number of product items irradiated in the verification dose experiment remains constant. The method is founded on and embodies the following three principles:

- existence of a direct link between the outcome of the verification dose experiment and the attainment of an SAL of 10^{-6} at the selected sterilization dose;
- possession of a level of conservativeness at least equal to that of the standard distribution of resistances (SDR);
- for a given bioburden, use of a maximal verification dose (VD_{max}) corresponding to substantiation of a selected sterilization dose.

This approach to sterilization dose substantiation was first outlined by Kowalski and Tallentire [6] and, from subsequent evaluations involving computational techniques (Kowalski, Aoshuang and Tallentire [7]) and field evaluations (Kowalski et al [8]), it was concluded that the method is soundly based. An overview of the method and aspects of putting it into practice are provided in Kowalski and Tallentire. [9] [10] Application of the Method VD_{max} approach to doses other than 25 kGy is discussed in Kowalski and Tallentire. [11] [12]

The method described here and designated Method $VD_{max}^{\ SD}$ procedurally comprises elements that closely parallel those of dose setting Method 1 described in ISO 11137-2. One key area of difference is the number of product items used in the verification dose experiment. In the computer evaluations referred to above, changing the verification SAL value had little effect on the substantiation outcome and this finding led to a sample size of 10 product items being chosen for subsequent field evaluations and, ultimately, for inclusion in this document.

Manufacturers of health care products who intend to use this specification are reminded that the requirements contained in ISO 11137 apply to the manufacture and control of production batches destined for radiation sterilization. In particular, one requirement states that products have to be manufactured in circumstances such that the bioburden is controlled. Compliance with the requirements for controlling the quality of raw materials, the manufacturing environment, the health, hygiene and attire of personnel and for establishing the basic properties of packaging material is essential.

Sterilization of health care products — Radiation — Substantiation of selected sterilization dose: Method VD_{max}^{SD}

1 Scope

1.1 Inclusions

This Technical Specification describes a method for substantiating a selected sterilization dose of 17,5, 20, 22,5, 27,5, 30, 32,5 or 35 kGy that achieves a sterility assurance level (SAL) of 10^{-6} or less for radiation sterilization of health care products. This Technical Specification also specifies a method of sterilization dose audit used to demonstrate the continued effectiveness of the substantiated sterilization dose.

NOTE Selection and substantiation of the sterilization dose is used to meet the requirements for establishing the sterilization dose within process definition in ISO 11137-1.

1.2 Exclusions

This method is for the substantiation of a selected sterilization dose of 17,5, 20, 22,5, 27,5, 30, 32,5, or 35 kGy only and is not used to substantiate other sterilization doses. The method is not used for the substantiation of a selected sterilization dose if the average bioburden of the entire product item exceeds the limit specified for the selected sterilization dose (see <u>Table 3</u>).

NOTE The methods for substantiation of selected sterilization doses of 25 kGy and 15 kGy are not included in this Technical Specification; they are described in ISO 11137-2.

1.3 Application

If the decision is made to use this method of sterilization dose establishment, the method is to be followed according to the requirements (shall) and guidance (should) stipulated herein.

2 Normative references

The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

 $ISO\,11137-1:2006, Sterilization\,of\,health\,care\,products --Radiation\,--Part\,1: Requirements\,for\,development,\\ validation\,and\,routine\,control\,of\,a\,sterilization\,process\,for\,medical\,devices$

ISO 11737-1, Sterilization of medical devices — Microbiological methods — Part 1: Determination of a population of microorganisms on products

ISO 11737-2, Sterilization of medical devices — Microbiological methods — Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process

3 Terms and definitions

For the purposes of this document, the following abbreviations, terms and definitions apply.

3.1

batch

defined quantity of product, intended or purported to be uniform in character and quality, that has been produced during a defined cycle of manufacture

[SOURCE: ISO/TS 11139:2006, 2.1]

3.2

bioburden

population of viable microorganisms on or in product and/or sterile barrier system

[SOURCE: ISO/TS 11139:2006, 2.2]

3.3

correction

action to eliminate a detected nonconformity

Note 1 to entry: A correction can be made in conjunction with corrective action (3.4).

[SOURCE: ISO 9000:2005, 3.6.6, modified]

3.4

corrective action

action to eliminate the cause of a detected nonconformity or other undesirable situation

Note 1 to entry: There can be more than one cause for a nonconformity.

Note 2 to entry: Corrective action is taken to prevent recurrence whereas preventive action is taken to prevent occurrence.

Note 3 to entry: There is a distinction between correction and corrective action.

[SOURCE: ISO 9000:2005, 3.6.5]

3.5

dose

absorbed dose

quantity of ionizing radiation energy imparted per unit mass of specified material

Note 1 to entry: The unit of absorbed dose is the gray (Gy), where 1 Gy is equivalent to the absorption of 1 J/kg.

Note 2 to entry: For the purposes of this document, the term dose is used to mean absorbed dose.

[SOURCE: ISO 11137-1:2006, 3.1, modified]

3.6

dose mapping

measurement of dose distribution and variability in material irradiated under defined conditions

[SOURCE: ISO 11137-1:2006, 3.10]

3.7

false positive

test result interpreted as growth arising from product, or portion thereof, tested when either growth resulted from extraneous microbial contamination or turbidity occurred from interaction between the product, or portion thereof, and the test medium

[SOURCE: ISO 11137-2:2012, 3.1.3]

3.8

health care product(s)

 $medical\ device(s),\ including\ in\ vitro\ diagnostic\ medical\ device(s),\ or\ medicinal\ product(s),\ including\ biopharmaceutical(s)$

[SOURCE: ISO/TS 11139:2006, 2.20]

3.9

medical device

instrument, apparatus, implement, machine, appliance, implant, *in vitro* reagent or calibrator, software, material, or other related article intended by the manufacturer to be used, alone or in combination, for human beings for one or more of the specific purpose(s) of:

- diagnosis, prevention, monitoring, treatment, or alleviation of disease;
- diagnosis, monitoring, treatment, alleviation of or compensation for an injury;
- investigation, replacement, modification, or support of the anatomy or of a physiological process;
- supporting or sustaining life;
- control of conception;
- disinfection of medical devices;
- providing information for medical purposes by means of *in vitro* examination of specimens derived from the human body;

and which does not achieve its primary intended action in or on the human body by pharmacological, immunological or metabolic means, but which may be assisted in its function by such means

Note 1 to entry: This definition from ISO 13485 has been developed by the Global Harmonization Task Force (GHTF 2002)

[SOURCE: ISO 13485:2003, 3.7, modified]

3.10

Method VD_{max}

procedure for sterilization dose substantiation that uses the maximal verification dose for a given bioburden, consistent with the attainment of a SAL of 10^{-6} at a selected sterilization dose

Note 1 to entry: The substantiation method is generally referred to as Method $VD_{max}^{\quad SD}$, where SD takes the value of the selected sterilization dose.

3.11

microorganism

entity of microscopic size, encompassing bacteria, fungi, protozoa and viruses

Note 1 to entry: A specific standard might not require demonstration of the effectiveness of the sterilization process in inactivating all types of microorganisms, identified in the definition above, for validation and/or routine control of the sterilization process.

[SOURCE: ISO/TS 11139:2006, 2.26]

3.12

packaging system

combination of the sterile barrier system and protective packaging

[SOURCE: ISO/TS 11139:2006, 2.28]

3.13

positive test of sterility

test result for which there is detectable microbial growth from product, or portion thereof, subjected to a test of sterility

[SOURCE: ISO 11137-2:2012, 3.1.8]

3.14

product

result of a process

Note 1 to entry: For the purposes of sterilization standards, product is tangible and can be raw material(s), intermediate(s), sub-assembly(ies) or health care product(s)

[SOURCE: ISO 9000:2005, 3.4.2, modified]

3.15

sample item portion

SIP

defined portion of a health care product that is tested

[SOURCE: ISO 11137-2:2012, 3.1.9]

3.16

sterile barrier system

minimum package that prevents ingress of microorganisms and allows aseptic presentation of product at the point of use

[SOURCE: ISO/TS 11139:2006, 2.44]

3.17

sterility

state of being free from viable microorganisms

Note 1 to entry: In practice, no such absolute statement regarding the absence of microorganisms can be proven [see *sterilization* (3.19)].

[SOURCE: ISO/TS 11139:2006, 2.45]

3.18

sterility assurance level

SAL

probability of a single viable microorganism occurring on an item after sterilization

Note 1 to entry: The term SAL takes a quantitative value, generally 10^{-6} or 10^{-3} . When applying this quantitative value to assurance of sterility, an SAL of 10^{-6} has a lower value but provides a greater assurance of sterility than an SAL of 10^{-3} .

[SOURCE: ISO/TS 11139:2006, 2.46]

3.19

sterilization

validated process used to render product free from viable microorganisms

Note 1 to entry: In a sterilization process, the nature of microbial inactivation is exponential and thus the survival of a microorganism on an individual item can be expressed in terms of probability. While this probability can be reduced to a very low number it can never be reduced to zero [see *sterility assurance level* (3.18)].

[SOURCE: ISO/TS 11139:2006, 2.47]

3.20

sterilization dose

SD

dose selected to achieve the specified requirements for sterility

[SOURCE: ISO 11137-1:2006, 3.40, modified]

3.21

sterilization dose audit

exercise undertaken to confirm the appropriateness of an established sterilization dose

[SOURCE: ISO 11137-2:2012, 3.2.12]

3.22

test of sterility

technical operation performed as part of development, validation or requalification to determine the presence or absence of viable microorganisms on product or portion thereof

[SOURCE: ISO/TS 11139:2006, 2.54]

3.23

verification dose

dose predicted to give a predetermined SAL greater than or equal to 10^{-2} used in establishing the sterilization dose

Note 1 to entry: For the purpose of this Technical Specification, this predetermined SAL is 10^{-1} .

3.24

 VD_{max}^{SD}

 $maximal \, verification \, dose \, for \, a \, particular \, selected \, sterilization \, dose \, (SD) \, obtained \, in \, using \, Method \, \, VD_{max}^{SD}$

4 Definition and maintenance of product families for sterilization dose substantiation and sterilization dose auditing

4.1 General

The establishment of a sterilization dose, for which sterilization dose selection and substantiation can be undertaken, and the carrying out of sterilization dose audits are activities that are part of process definition and maintaining process effectiveness (see ISO 11137-1). For these activities, product may be grouped into families; definition of product families is based principally on the numbers and types of microorganisms on or in product (the bioburden), the type being indicative of the microorganism's resistance to radiation (see ISO 11737-1). Variables such as density and product configuration within its packaging system are not considered in the establishment of these product families because they are not factors that influence bioburden.

In using product families for establishing the sterilization dose and for carrying out sterilization dose audits, it is important to be aware of the reduction in the ability to detect an inadvertent change within the manufacturing process that influences the effectiveness of sterilization. Furthermore, with the use of a single product to represent the product family, changes that occur in other members of the product family might not be detected. The effect of a reduction on ability to detect changes in other members of the product family should be evaluated and a plan for maintaining product families developed and implemented before proceeding.

4.2 Defining product families

4.2.1 The criteria for defining a product family shall be documented. Product shall be assessed against these criteria and the similarities between potential product family members considered. Consideration shall include all product-related variables that affect bioburden, including, but not limited to:

5

- a) nature and sources of raw materials, including the effect, if any, of raw materials that might be sourced from more than one location;
- b) components;
- c) product design and size;
- d) manufacturing processes;
- e) manufacturing equipment;
- f) manufacturing environment;
- g) manufacturing location.

The outcome of the assessment and considerations shall be recorded (see 4.1.2 of ISO 11137-1:2006).

- **4.2.2** Product shall only be included in a product family if it is demonstrated that the product-related variables (see <u>4.2.1</u>) are similar and under control.
- **4.2.3** To include product within a product family, it shall be demonstrated that bioburden comprises similar numbers and types of microorganisms.
- **4.2.4** Inclusion of product from more than one manufacturing location in a product family shall be specifically justified and recorded (see 4.1.2 of ISO 11137-1:2006). Consideration shall be given to the effect on bioburden of:
- a) geographic and/or climatic differences between locations;
- b) any differences in the control of the manufacturing processes or environment;
- c) sources of raw materials and processing adjuvants (e.g. water).

4.3 Designation of product to represent a product family

4.3.1 Product to represent a product family

- **4.3.1.1** The number and types of microorganisms on or in product shall be used as the basis for selecting product to represent a product family.
- **4.3.1.2** A product family shall be represented by:
- a) a master product (see 4.3.2), or
- b) an equivalent product (see 4.3.3), or
- c) a simulated product (see 4.3.4).
- **4.3.1.3** A formal, documented assessment shall be undertaken to decide which of the three potential representative products in $\frac{4.3.1.2}{1.00}$ is appropriate. In this assessment, consideration shall be given to the following:
- a) number of microorganisms comprising the bioburden;
- b) types of microorganisms comprising the bioburden;
- c) environment in which the microorganisms occur;
- d) size of product;

- e) number of components;
- f) complexity of product;
- g) degree of automation during manufacture;
- h) manufacturing environment.

4.3.2 Master product

A member of a product family shall only be considered a master product if assessment (see 4.3.1.3) indicates that the member presents a challenge to the sterilization process that is greater than that of all other product family members. In some situations, there can be several products within the product family, each of which could be considered as the master product. In such circumstances, any one of these products may be selected as the master product to represent the family, either

- a) at random, or
- b) according to a documented procedure to include the different products each of which could be considered as the master product.

4.3.3 Equivalent product

A group of product shall only be considered equivalent if assessment (see 4.3.1.3) indicates that group members require the same sterilization dose. Selection of the equivalent product to represent the family shall be either a) at random, or b) according to a documented procedure to include different members of the product family. The manufacturing volume and availability of product should be considered in the selection of the equivalent product to represent the product family.

4.3.4 Simulated product

A simulated product shall only represent a product family if it constitutes an equivalent or greater challenge to the sterilization process than that provided by members of the product family. Simulated product shall be packaged in a manner and with materials used for the actual product.

NOTE A simulated product is not intended for clinical use; it is fabricated solely for the establishment or maintenance of the sterilization dose.

A simulated product may be:

- a) one that is similar to the actual product in terms of materials and size, and subjected to similar manufacturing processes; e.g. a piece of the material, used for implants, that goes through the entire manufacturing process, or
- b) a combination of components from product within the product family that would not typically be combined for use; e.g. a tubing set containing multiple filters, clamps and stopcocks that are components of other products within the product family.

4.4 Maintaining product families

4.4.1 Periodic review

Review shall be performed at a specified frequency to ensure that product families and product used to represent each product family remain valid. Responsibility for reviews of product and/or processes that might affect membership of product families shall be allocated to competent personnel. Such review shall be performed at least annually. The outcome of the review shall be recorded in accordance with 4.1.2 of ISO 11137-1:2006.

4.4.2 Modification to product and/or manufacturing process

Modifications to product, such as raw materials (nature and source), components or product design (including size), and/or modifications to the manufacturing process, such as equipment, environment or location, shall be assessed through a formal, documented change control system. Such modifications can alter the basis on which the product family was defined or the basis on which the selection of product to represent the product family was made. Significant changes can require definition of a new product family or the selection of a different representative product.

4.4.3 Records

Records of product families shall be retained (see 4.1.2 of ISO 11137-1:2006).

4.5 Consequence of failure of sterilization dose substantiation or sterilization dose audit

In the event of failure during substantiation of a selected sterilization dose or performance of the sterilization dose audit for a product family, all members of that family shall be considered to be affected. Subsequent actions shall apply to all members comprising the product family.

5 Selection and testing of product for substantiating and auditing a selected sterilization dose

5.1 Nature of product

5.1.1 Product for sterilization can consist of:

- a) an individual health care product in its packaging system;
- b) a set of components presented in a packaging system, which are assembled at the point of use to form the health care product, together with accessories required to use the assembled product;
- c) a number of identical health care products in their packaging system;
- d) a kit comprising a variety of procedure-related health care products.

Product items for sterilization dose substantiation and for sterilization dose auditing shall be taken in accordance with Table 1.

Table 1 — Nature of product items for sterilization dose substantiation and for sterilization dose auditing

Product type	Item for bioburden determination and verification dose experiment	Rationale	
Individual health care product in its packaging system	Individual health care product	Each health care product is used independently in clinical practice.	
Set of components in a packaging system	Combination of all components of the product	Components are assembled as a product and used together in clinical practice.	
Number of identical health care products in their packaging system	Single health care product taken from the packaging system	Each health care product is used independently in clinical practice; the SAL of an individual health care product within the packaging system meets the selected SAL, although the overall SAL associated with the packaging system might be higher.	
Kit of procedure-related health care products ^a	Each type of health care product comprising the kit	Each health care product is used independently in clinical practice.	
In dose establishment, the sterilization dose is chosen based on the health care product requiring the highest sterilization dose.			

^{5.1.2} If the product has a claim of sterility for part of the product, the sterilization dose may be established on the basis of that part only.

EXAMPLE If the product has a label claim of sterility for the fluid path only, the sterilization dose may be established based on bioburden determinations and outcomes of tests of sterility performed on the fluid path.

5.2 Sample item portion (SIP)

- **5.2.1** For product with an average bioburden greater than or equal to 1,0, whenever practicable, an entire product (SIP equal to 1,0) should be used for testing in accordance with <u>Table 1</u>. When the use of an entire product is not practicable, a selected portion of product (SIP) may be substituted. The SIP should be as large a portion of the item as practicable and should be of a size that can be handled during testing.
- **5.2.2** For a product with an average bioburden less than or equal to 0,9, an entire product (SIP equal to 1,0) shall be used for testing in accordance with <u>Table 1</u>.

NOTE When testing products with low average bioburden, it is possible that an SIP will not always be the portion of the product item possessing microorganisms. Therefore, the entire product (SIP = 1,0) is used for products with an average bioburden less than or equal to 0,9.

- **5.2.3** If the bioburden is evenly distributed on and/or in the item, the SIP may be selected from any portion of the item. If the bioburden is not evenly distributed, the SIP shall consist of either
- a) portions of product selected at random that proportionally represent each of the materials from which the product is made, or
- b) the portion of the product that is considered to be the most severe challenge to the sterilization process.

The value of SIP can be calculated on the basis of length, mass, volume or surface area (see <u>Table 2</u> for examples).

Basis for SIP	Product
I+1.	Tubing (consistent diameter)
Length	Rolls of bandage
M	Powders
Mass	Gowns
Volume	Liquids
C. C	Surgical drapes
Surface area	Tubing (variable diameter)

Table 2 — Examples for calculation of SIP

- **5.2.4** The preparation and packaging of an SIP shall be carried out under conditions that minimize alterations to bioburden. Environmentally-controlled conditions should be used for preparation of SIPs and, whenever possible, packaging materials should be equivalent to those used for the finished product.
- **5.2.5** The adequacy of a selected SIP shall be demonstrated. The bioburden of the SIP shall be such that either at least 17 of the 20 non-irradiated SIPs yield positive tests of sterility, or a bioburden of one or more is found on at least 85 % of 20 or more SIPs. If neither of these criteria is met, an SIP that is different than that examined originally and that meets one of the above criteria shall be used. If an entire product is tested (SIP equal to 1,0), the criteria specified above do not apply.
- **5.2.6** The same portion of product item (SIP) should be used in the performance of tests of sterility when carrying out the verification dose experiment as that used in the determination of bioburden when obtaining the verification dose.
- NOTE If the portion of product item (SIP) used in the performance of tests of sterility is different from that used in the determination of bioburden, caution should be exercised when selecting the sterilization dose and when calculating the value of SIP VD_{max}^{SD} . In carrying out these two activities, two separate determinations of bioburden are required: one for the SIP used to obtain the bioburden for the entire product item employed in the selection of the sterilization dose and the other for the SIP used to obtain the value of SIP VD_{max}^{SD} employed in the performance of the verification dose experiment.

5.3 Manner of sampling

- **5.3.1** Product for sterilization dose substantiation and for sterilization dose auditing shall be representative of that subjected to routine manufacturing procedures and conditions.
- **5.3.2** Each product item used in the determination of bioburden or in the performance of a test of sterility should be taken, where applicable (see <u>Table 1</u>), from a separate packaging system.
- **5.3.3** The period of time between the taking of product items from production and the determination of bioburden or the performance of the verification dose experiment should reflect the time period between completion of the last manufacturing step and sterilization of product.
- **5.3.4** If prolonged storage of product items is necessary prior to the determination of bioburden and/or in the performance of tests of sterility, product capable of supporting microbial growth shall be stored under conditions that inhibit such growth.
- **5.3.5** Product items may be selected from product rejected during the manufacturing process provided that they have been subjected to the same manufacturing procedures and conditions as the remainder of production, including packaging.

5.4 Microbiological testing

5.4.1 Bioburden determinations and performance of tests of sterility shall be conducted in accordance with ISO 11737-1 and ISO 11737-2, respectively.

NOTE Soybean Casein Digest Broth, with an incubation temperature of (30 ± 2) °C and an incubation period of 14 days, is generally recommended when a single medium is used for the performance of tests of sterility. If there is reason to suspect that this medium and temperature do not support the growth of microorganisms present, other appropriate media and incubation conditions should be used. See, e.g. Herring et al.[5]; Favero[4]; NHB 5340.1A.[3]

To reduce the possibility of false positives in carrying out tests of sterility, items may be disassembled and repackaged prior to irradiation. Manipulations prior to irradiation shall not change the magnitude of the bioburden or its response to radiation (i.e. manipulations that alter the chemical environment in the vicinity of the microorganisms, typically oxygen tension).

5.4.2 Bioburden determinations shall be carried out on product that has undergone the packaging process.

NOTE Generally, it is sufficient to perform a bioburden determination on a product item after removal from its packaging system and to omit the packaging system from the determination.

5.5 Irradiation

- **5.5.1** Irradiation of product in performing sterilization dose substantiation and sterilization dose auditing shall be conducted in an irradiator that has undergone installation qualification and operational qualification in accordance with ISO 11137-1.
- **5.5.2** Measurement of dose and the use of radiation sources shall be in accordance with ISO 11137-1.
- **5.5.3** For the performance of a verification dose experiment, sufficient performance qualification dose mapping shall be carried out to identify the highest and the lowest doses delivered to product.
- **5.5.4** Whenever practicable, for the performance of a verification dose experiment, product should be irradiated in its original form and in its packaging system.
- **5.5.5** Materials for repackaging product items for irradiation, if applicable (see <u>5.4.1</u>), shall be capable of withstanding the doses delivered and subsequent handling, thereby minimizing the likelihood of contamination.

NOTE See ISO 11137-3 for guidance on dosimetric aspects of radiation sterilization.

6 Method VD_{max}^{SD} — Substantiation of a selected sterilization dose of 17,5, 20, 22,5, 27,5, 30, 32,5, or 35 kGy

6.1 Rationale

Operationally, the method of substantiation for a selected sterilization dose is similar to dose setting Method 1 of ISO 11137-2; it too requires a determination of bioburden and the performance of a verification dose experiment.

In carrying out substantiation, the method verifies that bioburden present on product prior to sterilization is less resistant to radiation than a microbial population of maximal resistance consistent with the attainment of an SAL of 10^{-6} at the selected sterilization dose; verification is conducted through performance of a verification dose experiment at an SAL of 10^{-1} using 10 product items. The dose at an SAL of 10^{-1} for a population having this resistance (maximal verification dose, VD_{max}) is characteristic of the bioburden level, the sterilization dose and the associated maximal resistance. In

establishing the maximal resistance for a particular bioburden level and sterilization dose, due account has been taken of the various resistance components of the Standard Distribution of Resistances (SDR), the latter being the basis of Method 1. Components of the SDR of high resistance that have significant effect on the attainment of an SAL of 10^{-6} have been used to define the maximal resistances on which this substantiation method is based. In this way, the level of conservativeness of the SDR, and thus of Method 1, is preserved. See Kowalski and Tallentire [\mathfrak{I}]; Kowalski, Aoshuang, and Tallentire [\mathfrak{I}]; Kowalski and Tallentire.

In practice, a determination is made of the average bioburden. Based on this average value, a sterilization dose is selected from a table listing the upper limits of average bioburden that apply to specified selected sterilization doses. These upper limits are the numbers of microorganisms possessing a given maximal resistance commensurate with the attainment of a SAL of 10^{-6} at the selected sterilization dose. The VD_{max}^{SD} dose corresponding to the selected sterilization dose and the average bioburden is read from a second table; it is the dose at which the verification dose experiment is carried out. Ten product items, or portions thereof (if applicable, see 5.2), are exposed to the VD_{max}^{SD} dose and each item is subjected individually to a test of sterility. If there is no more than one positive test of sterility in the 10 tests, the pre-selected sterilization dose is substantiated.

The VD_{max} methods given in this Technical Specification are for selected sterilization doses of 17,5, 20, 22,5, 27,5, 30, 32,5 and 35 kGy. To distinguish these applications of Method VD_{max} and their associated sets of values of verification dose, a superscript has been added to the term VD_{max} where appropriate, viz. VD_{max}^{SD} , where SD is the selected sterilization dose.

NOTE Inspection of the values of the VD_{max} for the various levels of average bioburden given in each of the tables in Clause 8 reveals a change in the relationship between the bioburden level and the value of VD_{max}^{SD} . With increasing bioburden up to a certain level, values progressively increase, as might be expected. However, at a particular bioburden level, the VD_{max}^{SD} takes a maximum, and for higher bioburden levels, the corresponding VD_{max}^{SD} values progressively decrease. For example, for Method $VD_{max}^{17,5}$, $VD_{max}^{17,5}$ values progressively increase up to a bioburden level of 2,5. However, at a bioburden of 2,5, the value of $VD_{max}^{17,5}$ takes a maximum, and for higher bioburden levels, the corresponding $VD_{max}^{17,5}$ values decrease. A similar increase, followed by a decrease, is seen with the other VD_{max}^{SD} methods. This behaviour is not the result of an error in either the tables or the calculation of the VD_{max}^{SD} values. It is an inevitable outcome of building into Method VD_{max}^{SD} the same degree of conservativeness as that in Method 1 (see Kowalski and Tallentire[10]).

6.2 Procedure for Method VD_{max}SD for multiple production batches

6.2.1 General

6.2.1.1 In applying Method VD_{max}^{SD} for product with an average bioburden less than or equal to 0,9, the entire product item shall be used, whereas for product with an average bioburden greater than 0,9, an SIP may be used (see 5.2.5).

6.2.1.2 In applying Method VD_{max} SD, the six stages below shall be followed.

NOTE For worked examples, see 9.1 and 9.2.

6.2.2 Stage 1: Obtain samples of product

Select 10 product items from each of three independent production batches, in accordance with 5.1, 5.2 (if applicable) and <u>5.3</u>.

NOTE Product will also be needed to perform the verification dose experiment (see $\underline{6.2.6.1}$) and additional product might be needed to validate the adequacy of an SIP less than one (see $\underline{5.2.5}$) or to perform a confirmatory verification dose experiment (see $\underline{6.2.7.2}$ and $\underline{6.2.8}$).

6.2.3 Stage 2: Determine average bioburden

- **6.2.3.1** Apply the correction factor in the determination of bioburden (see ISO 11737-1).
- **6.2.3.2** Determine the bioburden of each of the selected product items and calculate:
- a) the average bioburden per item for each of the three batches of product items (batch average bioburden);
- b) the average bioburden per item for all selected product items (overall average bioburden).

NOTE 1 Bioburden is generally determined on individual product items, but when the bioburden is low (e.g. less than 10), it is permissible to pool the 10 product items for the determination of average bioburden. This guidance does not apply to SIP; SIPs should not be pooled, rather a larger SIP should be chosen (see $\underline{5.2.5}$).

NOTE 2 An observation of no colonies in the determination of bioburden is sometimes expressed as less than the limit of detection. Use of the limit of detection as a bioburden value in calculating average bioburden could lead to an overestimation of average bioburden. Overestimation could lead to selection of too high a sterilization dose and, in consequence, a high value for VD_{max}^{SD} , thereby affecting the validity of the verification dose experiment. The use of an approach for bioburden determination having a low limit of detection can reduce such overestimation.

For an SIP less than 1,0, calculate the average bioburden for the entire product item (SIP equal to 1,0) by dividing each of the three SIP batch average bioburdens and the overall SIP average bioburden by the SIP value.

6.2.4 Stage 3: Obtain the selected sterilization dose

- **6.2.4.1** In obtaining the selected sterilization dose, values of average bioburden for the entire product item (SIP equal to 1,0) shall be used (see 6.2.3.2).
- **6.2.4.2** From $\underline{\text{Table 3}}$, obtain the selected sterilization dose. In obtaining this dose, all three batch average bioburdens (SIP equal to 1,0) determined in Stage 2 shall be below or equal to the associated upper limit of the average bioburden given in $\underline{\text{Table 3}}$.
- **6.2.4.3** A sterilization dose greater than the lowest dose consistent with meeting the requirement in 6.2.4.2 may be selected. A rationale for selecting a greater dose might be based on factors such as:
- a) the difference between the average bioburden and the upper limit associated with the selected sterilization dose;
- b) available data on the variation in the numbers and types of microorganisms that comprise the bioburden;
- c) available data on the microbiological quality of similar products including the results of sterilization dose audits;
- d) the materials comprising the product and the control of the microbiological quality of materials;
- e) the manufacturing process and associated control and monitoring procedures, particularly steps that affect bioburden or its resistance; and
- the manufacturing environment, particularly the extent of microbiological control and monitoring, and available data on the stability of the manufacturing environment over time.

Table 3 — Upper limit of average bioburden for selection of a given sterilization dose

Upper limit for average bioburden (SIP equal to 1,0)	Selected sterilization dose (kGy)
9,0	17,5
45	20
220	22,5
5 000	27,5
23 000	30
100 000	32,5
440 000	35

6.2.5 Stage 4: Obtain VD_{max}^{SD}

6.2.5.1 From Table 4, identify the table in Clause 8 that gives values of SIP equal to 1,0 VD_{max}^{SD} , SIP dose reduction factor and dose augmentation value, corresponding to different values of average bioburden, for the selected sterilization dose.

Table 4 — Table in Clause 8 corresponding to the selected sterilization dose

Selected sterilization dose (kGy)	Corresponding table in <u>Clause 8</u>
17,5	Table 5
20	Table 6
22,5	Table 7
27,5	Table 8
30	Table 9
32,5	Table 10
35	Table 11

- **6.2.5.2** Compare the three batch average bioburdens to the overall average bioburden found in Stage 2 and determine whether any one of the batch average bioburdens is two or more times greater than the overall average bioburden.
- **6.2.5.3** From the identified table in <u>Clause 8</u>, obtain the value of SIP equal to 1,0 VD_{max}^{SD} using one of the following as the average bioburden:
- a) if a batch average bioburden is two or more times greater than the overall average bioburden, use the highest batch average bioburden, or
- b) if each of the batch average bioburdens is less than two times the overall average bioburden, use the overall average bioburden.

For an SIP equal to 1,0, if the average bioburden is not given in the identified table in <u>Clause 8</u>, use the closest tabulated value greater than the average bioburden to locate the value of SIP equal to 1,0 VD_{max}^{SD} .

For an SIP less than 1,0, use the average bioburden for the entire product item (SIP equal to 1,0), calculated in Stage 2 (6.2.3.2), to enter the identified table in Clause 8. If the calculated average bioburden is not given in the identified table in Clause 8, use the closest tabulated value greater than the average

bioburden to locate the value of SIP equal to 1,0 VD_{max}^{SD} and the corresponding SIP dose reduction factor. Use Equation (1) to calculate the SIP VD_{max}^{SD} (see Kowalski and Tallentire[10]).

SIP
$$VD_{max}^{SD} = (SIP \text{ equal to 1,0 } VD_{max}^{SD}) + (SIP \text{ dose reduction factor } \times \log SIP)$$
 (1)

NOTE Use of an SIP less than 1,0 is not permitted for product with an average bioburden less than or equal to 0.9 (see 6.2.1.1).

6.2.6 Stage 5: Perform verification dose experiment

- **6.2.6.1** Select 10 product items from a single batch of product. The 10 product items for the performance of Stage 5 may be selected from one of the batches on which a bioburden determination was carried out in Stage 2, or from a fourth batch manufactured under conditions that are representative of normal production (see <u>5.3</u>).
- **6.2.6.2** Irradiate these product items at VD_{max}^{SD} obtained from the identified table in <u>Clause 8</u> or calculated using Equation (1), whichever is appropriate.

The highest dose to product items shall not exceed $VD_{max}^{\ \ SD}$ by more than 10 % or 0,1 kGy, whichever is greater.

NOTE A tolerance of 0,1 kGy is allowed in order to accommodate the ability and practicality of irradiation facilities to deliver and measure $VD_{max}^{\ \ SD}$ doses below 1,0 kGy.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of $VD_{max}^{\ \ SD}$.

Determine the dose delivered (see 5.5).

If the highest dose to product items exceeds VD_{max}^{SD} by more than 10 % or 0,1 kGy, whichever is greater, the verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the verification dose experiment may be repeated. If this mean dose is less than 90 % of VD_{max}^{SD} and, on performance of tests of sterility, acceptable results are observed (see 6.2.7), the verification dose experiment need not be repeated.

6.2.6.3 Subject each irradiated product item individually to a test of sterility (see 5.4.1) and record the number of positive tests of sterility.

6.2.7 Stage 6: Interpretation of results

- **6.2.7.1** If no more than one positive test of sterility is obtained from the 10 tests carried out, accept verification and thereby substantiate the selected sterilization dose.
- **6.2.7.2** If two positive tests of sterility are obtained, perform a confirmatory verification dose experiment (see 6.2.8).
- **6.2.7.3** If three or more positive tests of sterility are obtained, do not accept verification as the selected sterilization dose might be inadequate.

If the occurrence of these positive tests of sterility can be ascribed to incorrect performance of the determination of bioburden, incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} or a specific bioburden-related cause, implement corrective action and repeat the verification dose

experiment using a further 10 product items from a batch manufactured under conditions that are representative of normal production. If, as a result of corrective action, the estimate of average bioburden changes, for the repeat verification dose experiment use the VD_{max}^{SD} (6.2.5) that corresponds to the changed average bioburden. If the estimate of average bioburden is unchanged, use the same VD_{max}^{SD} as that used in the verification dose experiment that was not accepted. Interpret the results of the repeat verification dose experiment in accordance with 6.2.7.

If the occurrence of these positive tests of sterility cannot be ascribed to incorrect performance of the determination of bioburden, incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} or a specific bioburden-related cause, the selected sterilization dose is not substantiated and another approach for establishing a sterilization dose shall be used. Other approaches are:

- a) selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 3 (6.2.4);
- b) Method 1;
- c) Method 2; and
- d) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

6.2.8 Confirmatory verification dose experiment

6.2.8.1 General

If a confirmatory verification dose experiment is to be carried out (see <u>6.2.7.2</u>), the three stages below shall be followed.

6.2.8.2 Stage 1: Obtain samples of product

Select 10 product items from a single batch of product. The 10 product items for the performance of the confirmatory verification dose experiment may be selected from one of the batches on which a bioburden determination was carried out in Stage 2 (see <u>6.2.3</u>), from the fourth batch used in Stage 5 (see <u>6.2.6.1</u>) or from a batch manufactured under conditions that are representative of normal production (see <u>5.3</u>).

6.2.8.3 Stage 2: Perform confirmatory verification dose experiment

6.2.8.3.1 Irradiate these product items at VD_{max} SD obtained in <u>6.2.5</u>.

The highest dose to product items shall not exceed $VD_{max}^{\ \ SD}$ by more than 10 % or 0,1 kGy, whichever is greater.

NOTE A tolerance of 0,1 kGy is allowed in order to accommodate the ability and practicality of irradiation facilities to deliver and measure VD_{max}^{SD} doses below 1,0 kGy.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of $VD_{max}^{\ \ SD}$.

Determine the dose delivered (see 5.5).

If the highest dose to product items exceeds $VD_{max}^{~~SD}$ by more than 10 % or 0,1 kGy, whichever is greater, the verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the confirmatory verification dose experiment may be repeated. If this mean dose is less than 90 % of

 VD_{max}^{SD} and, on performance of tests of sterility, acceptable results are observed (see <u>6.2.8.4</u>), the confirmatory verification dose experiment need not be repeated.

6.2.8.3.2 Subject each irradiated product item individually to a test of sterility (see <u>5.4.1</u>) and record the number of positive tests of sterility.

6.2.8.4 Stage 3: Interpretation of results

- **6.2.8.4.1** If there are no positive tests of sterility from the 10 tests carried out, accept confirmatory verification and thereby substantiate the selected sterilization dose.
- **6.2.8.4.2** If any positive tests of sterility are obtained, do not accept confirmatory verification as the selected sterilization dose might be inadequate.

If the occurrence of these positive tests of sterility can be ascribed to incorrect performance of tests of sterility or incorrect delivery of $VD_{max}^{\ SD}$, or a specific bioburden-related cause, implement corrective action and repeat the confirmatory verification dose experiment using a further 10 product items from a batch manufactured under conditions that are representative of normal production and the same $VD_{max}^{\ SD}$ as that used originally. Interpret the results of the repeat confirmatory verification dose experiment in accordance with 6.2.8.4.

If the occurrence of these positive tests of sterility cannot be ascribed to incorrect performance of tests of sterility or incorrect delivery of $VD_{max}^{\ \ SD}$, or a specific bioburden-related cause, the selected sterilization dose is not substantiated and another approach for establishing a sterilization dose shall be used. Other approaches are:

- a) selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 3 (6.2.4);
- b) Method 1;
- c) Method 2; and
- d) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

6.3 Procedure for Method VD_{max}SD for a single production batch

6.3.1 Rationale

This method is an adaptation of Method $VD_{max}^{\ \ SD}$ and is intended to be used only for the substantiation of a selected sterilization dose for a single production batch.

6.3.2 General

- **6.3.2.1** In applying Method VD_{max}^{SD} for product with an average bioburden less than or equal to 0,9, the entire product item shall be used, whereas for product with an average bioburden greater than 0,9, an SIP may be used (see <u>5.2.5</u>).
- **6.3.2.2** Product capable of supporting microbial growth should be stored under conditions that inhibit such growth for the time between manufacture of the single production batch and sterilization.
- **6.3.2.3** In applying Method VD_{max}^{SD} , the six stages below shall be followed.

6.3.3 Stage 1: Obtain samples of product

Select 10 product items from the single production batch, in accordance with 5.1, 5.2 (if applicable) and 5.3.

NOTE Product will also be needed to perform the verification dose experiment (see 6.3.7) and additional product might be needed to validate the adequacy of an SIP less than one (see 5.2.5) or to perform a confirmatory verification dose experiment (see 6.3.9).

6.3.4 Stage 2: Determine average bioburden

- **6.3.4.1** Apply the correction factor in the determination of bioburden (see ISO 11737-1).
- **6.3.4.2** Determine the bioburden of each of the selected product items and calculate the average bioburden.

NOTE 1 Bioburden is generally determined on individual product items, but when the bioburden is low (e.g. less than 10), it is permissible to pool the 10 product items for the determination of average bioburden. This guidance does not apply to SIP; SIPs should not be pooled, rather a larger SIP should be chosen (see <u>5.2.5</u>).

NOTE 2 An observation of no colonies in the determination of bioburden is sometimes expressed as less than the limit of detection. Use of the limit of detection as a bioburden value in calculating average bioburden could lead to an overestimation of average bioburden. Overestimation could lead to selection of too high a sterilization dose and, in consequence, a high value of VD_{max}^{SD} , thereby affecting the validity of the verification dose experiment. The use of an approach for bioburden determination having a low limit of detection can reduce such overestimation.

For an SIP less than 1,0, calculate the average bioburden for the entire product item (SIP equal to 1,0) by dividing the SIP batch average bioburden by the SIP value.

6.3.5 Stage 3: Obtain the selected sterilization dose

- **6.3.5.1** In obtaining the selected sterilization dose, the value of average bioburden for the entire product item (SIP equal to 1,0) shall be used (see 6.3.4.2).
- **6.3.5.2** From <u>Table 3</u>, obtain the selected sterilization dose. In obtaining this dose, the batch average bioburden (SIP equal to 1,0) determined in Stage 2 shall be less than or equal to the associated upper limit of the average bioburden given in <u>Table 3</u>.
- **6.3.5.3** A sterilization dose greater than the lowest dose consistent with meeting the requirement in <u>6.3.5.2</u> may be selected. A rationale for selecting a greater dose might be based on factors such as:
- a) the difference between the average bioburden and the upper limit associated with the selected sterilization dose;
- $b) \quad available \, data \, on \, the \, variation \, in \, the \, numbers \, and \, types \, of \, microorganisms \, that \, comprise \, the \, bioburden;$
- c) available data on the microbiological quality of similar products including the results of sterilization dose audits;
- d) the materials comprising the product and the control of the microbiological quality of materials;
- e) the manufacturing process and associated control and monitoring procedures, particularly steps that affect bioburden or its resistance; and
- f) the manufacturing environment, particularly the extent of microbiological control and monitoring, and available data on the stability of the manufacturing environment over time.

6.3.6 Stage 4: Obtain VD_{max}SD

- **6.3.6.1** From Table 4, identify the table in Clause 8 that gives the values of SIP equal to 1,0 VD_{max}^{SD} , SIP dose reduction factor and dose augmentation value, corresponding to different values of average bioburden, for the selected sterilization dose.
- **6.3.6.2** From the identified table, obtain the value of SIP equal to 1,0 VD_{max}^{SD} using the average bioburden for the entire product item (SIP equal to 1,0).

For an SIP equal to 1,0, if the average bioburden is not given in the identified table in <u>Clause 8</u>, use the closest tabulated value greater than the average bioburden to locate the value of SIP equal to 1,0 VD_{max}^{SD} .

For an SIP less than 1,0, use the average bioburden for the entire product item (SIP equal to1,0), calculated in Stage 2 (6.3.4.2), to enter the identified table in Clause 8. If the calculated average bioburden is not given in the table, use the closest tabulated value greater than the average value to locate the value of SIP equal to 1,0 VD_{max}^{SD} and corresponding SIP dose reduction factor. Use Equation (1) (see 6.2.5.3) to calculate the SIP VD_{max}^{SD} (see Kowalski and Tallentire[10]).

NOTE Use of an SIP less than 1,0 is not permitted for product with an average bioburden less than or equal to 0,9 (see 6.3.2.1).

6.3.7 Stage 5: Perform verification dose experiment

- **6.3.7.1** Select 10 product items from the single batch of product.
- **6.3.7.2** Irradiate these product items at VD_{max}^{SD} obtained from the identified table in <u>Clause 8</u> or derived using Equation (1), whichever is appropriate.

The highest dose to product items shall not exceed $VD_{max}^{\ \ SD}$ by more than 10 % or 0,1 kGy, whichever is greater.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of $VD_{max}^{\ \ SD}$.

Determine the dose delivered (see 5.5).

If the highest dose to product items exceeds VD_{max}^{SD} by more than 10 % or 0,1 kGy, whichever is greater, the verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the verification dose experiment may be repeated. If this mean dose is less than 90 % of VD_{max}^{SD} and, on performance of tests of sterility, acceptable results are observed (see <u>6.3.8</u>), the verification dose experiment need not be repeated.

6.3.7.3 Subject each irradiated product item individually to a test of sterility (see 5.4.1) and record the number of positive tests of sterility.

6.3.8 Stage 6: Interpretation of results

6.3.8.1 If no more than one positive test of sterility is obtained from the 10 tests carried out, accept verification and thereby substantiate the selected sterilization dose.

6.3.8.3 If three or more positive tests of sterility are obtained, do not accept verification as the selected sterilization dose might be inadequate.

If the occurrence of these positive tests of sterility can be ascribed to incorrect performance of the determination of bioburden, incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} or a specific bioburden-related cause, implement corrective action and repeat the verification dose experiment using a further 10 product items. If, as a result of corrective action, the estimate of average bioburden changes, use the VD_{max}^{SD} (6.3.6) that corresponds to the changed average bioburden. If the estimate of average bioburden is unchanged, use the same VD_{max}^{SD} as that used in the verification dose experiment that was not accepted. Interpret the results of the repeat verification dose experiment in accordance with 6.3.8.

If the occurrence of these positive tests of sterility cannot be ascribed to incorrect performance of the determination of bioburden, incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} or a specific bioburden-related cause, the selected sterilization dose is not substantiated and another approach for establishing a sterilization dose shall be used. Other approaches are:

- a) selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 3 (6.3.5);
- b) Method 1;
- c) Method 2; and
- d) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

6.3.9 Confirmatory verification dose experiment

6.3.9.1 General

If a confirmatory verification dose experiment is to be carried out (see <u>6.3.8.2</u>), the three stages below shall be followed.

6.3.9.2 Stage 1: Obtain samples of product

Select 10 product items from the single production batch of product.

6.3.9.3 Stage 2: Perform confirmatory verification dose experiment

6.3.9.3.1 Irradiate these product items at VD_{max}^{SD} obtained in <u>6.3.6</u>.

The highest dose to product items shall not exceed $VD_{max}^{SD\ D}$ by more than 10 % or 0,1 kGy, whichever is greater.

NOTE A tolerance of 0,1 kGy is allowed in order to accommodate the ability and practicality of irradiation facilities to deliver and measure VD_{max}^{SD} doses below 1,0 kGy.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of VD_{max}^{SD} .

Determine the dose delivered (see 5.5).

If the highest dose to product items exceeds VD_{max}^{SD} by more than 10 % or 0,1 kGy, whichever is greater, the confirmatory verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the confirmatory verification dose experiment may be repeated. If this mean dose is less than 90 % of VD_{max}^{SD} and, on performance of tests of sterility, acceptable results are observed (see <u>6.3.9.4</u>), the confirmatory verification dose experiment need not be repeated.

6.3.9.3.2 Subject each irradiated product item individually to a test of sterility (see <u>5.4.1</u>) and record the number of positive tests of sterility.

6.3.9.4 Stage 3: Interpretation of results

6.3.9.4.1 If there are no positive tests of sterility from the 10 tests carried out, accept confirmatory verification and thereby substantiate the selected sterilization dose.

6.3.9.4.2 If any positive tests of sterility are obtained, do not accept verification as the selected sterilization dose might be inadequate.

If the occurrence of these positive tests of sterility can be ascribed to incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} , or a specific bioburden-related cause, implement corrective action and repeat the confirmatory verification dose experiment using a further 10 product items and the same VD_{max}^{SD} as that used originally. Interpret the results of the repeat confirmatory verification dose experiment in accordance with 6.3.9.4.

If the occurrence of these positive tests of sterility cannot be ascribed to incorrect performance of tests of sterility or incorrect delivery of $VD_{max}^{\ \ SD}$, or a specific bioburden-related cause, the selected sterilization dose is not substantiated and another approach for establishing a sterilization dose shall be used. Other approaches are:

- a) selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 3 (6.3.5);
- b) Method 1;
- c) Method 2; and
- d) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

7 Maintaining process effectiveness

7.1 General

For product produced as multiple production batches, periodic determinations of bioburden and sterilization dose audits are carried out to confirm the continued effectiveness of the established sterilization dose. Requirements for maintaining process effectiveness given in Clause 12 of ISO 11137-1 apply to a selected sterilization dose substantiated using Method VD_{max}^{SD} together with those specific to Method VD_{max}^{SD} given below (see 7.2.2). None of these requirements apply to product produced as a single production batch.

7.2 Determination of bioburden

7.2.1 Background to frequency of determination

ISO 11137-1:2006 requires:

- a) for product with a bioburden greater than or equal to 1,5, determination of bioburden at a maximal time interval of three months (ISO 11137-1:2006, 12.1.2.1),
- b) for product with a bioburden less than 1,5 and for which the sterilization dose has been established using either Method 2 or Method VD_{max}^{25} , determination of bioburden at a maximal time interval of three months (ISO 11137-1:2006, 12.1.2.2), and
- c) for product with bioburden less than 1,5 and for which the sterilization dose has been established using either Method 1 or Method VD_{max}^{15} , the maximal time interval of one month (ISO 11137-1:2006, 12.1.2.3).

The requirement for the increased frequency of determination of bioburden given in ISO 11137-1:2006, 12.1.2.3, is to take account of wide variation in bioburden on product items that occurs, on occasions, with product of low bioburden. An increased frequency was specified for bioburden values below 1,5 in the light of this value being the upper bioburden limit of Method VD_{max}^{15} . An increased frequency of bioburden determination is also specified to Method $VD_{max}^{17,5}$ in this Technical Specification.

7.2.2 Frequency of determination specific to Method VD_{max}^{SD}

- **7.2.2.1** For product of average bioburden greater than or equal to 10 and for which the selected sterilization dose has been substantiated using Method VD_{max}^{SD} , the maximal interval of time between determinations of bioburden shall be three months.
- **7.2.2.2** For product of average bioburden less than 10 and for which the selected sterilization dose has been substantiated using Method $VD_{max}^{17,5}$, the maximal interval of time between determinations of bioburden shall be one month.

NOTE In these circumstances, this Technical Specification stipulates a shorter maximal interval of time than stipulated in ISO 11137-1 for an average bioburden greater than 1,5 but less than 10.

7.3 Sterilization dose audit

7.3.1 Frequency

The frequency at which sterilization dose audits are carried out shall be in accordance with 12.1 of ISO 11137-1:2006. Sterilization dose audits are not required during periods in which product is not produced. A review of environmental and manufacturing controls, together with determinations of bioburden, should be conducted in conjunction with sterilization dose audits. If the review indicates lack of control, appropriate action shall be taken.

7.3.2 Outcome

Actions resulting from the outcome of a sterilization dose audit shall apply to all product comprising the product family (see <u>Clause 4</u>).

7.3.3 Procedure for auditing a sterilization dose substantiated using Method VD_{max}^{SD}

7.3.3.1 **General**

- **7.3.3.1.1** For the performance of a sterilization dose audit for a selected sterilization dose, substantiated using Method VD_{max}^{SD} , use an SIP equivalent to that used originally in substantiating the sterilization dose.
- **7.3.3.1.2** In applying the sterilization dose audit, the four stages below shall be followed.

NOTE For a worked example, see 9.3.

7.3.3.2 Stage 1: Obtain samples of product

Select 20 product items from a single batch of product, in accordance with 5.1, 5.2 (if applicable) and 5.3.

7.3.3.3 Stage 2: Determine average bioburden

- **7.3.3.3.1** Apply the correction factor found from the most recent validation of the method of bioburden determination.
- **7.3.3.3.2** Determine the bioburden of each of 10 product items and calculate the average bioburden.
- NOTE 1 Bioburden is generally determined on individual product items, but when the bioburden is low (e.g. less than 10), it is permissible to pool the 10 product items for the determination of average bioburden. This guidance does not apply to SIP; SIPs should not be pooled, rather a larger SIP should be chosen (see $\underline{5.2.5}$).
- NOTE 2 An observation of no colonies in the determination of bioburden is sometimes expressed as less than the limit of detection. Use of the limit of detection as a bioburden value in calculating average bioburden could lead to overestimation of the average. The use of an approach for bioburden determination having a low limit of detection can reduce such overestimation.
- NOTE 3 These bioburden data are not intended to be used in obtaining the value of SIP equal to 1,0 VD_{max}^{SD} for the sterilization dose audit. They are used for process monitoring and control (e.g. trend analysis), investigation of sterilization dose audit failure and obtaining the dose augmentation value.

7.3.3.4 Stage 3: Perform verification dose experiment

7.3.3.4.1 Irradiate 10 product items at VD_{max}^{SD} used in the most recent successful substantiation of the selected sterilization dose.

The highest dose to product items shall not exceed $VD_{max}^{\ \ SD}$ by more than 10 % or 0,1 kGy, whichever is greater.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of VD_{max}^{SD} .

Determine the dose delivered (see <u>5.5</u>).

If the highest dose to product items exceeds VD_{max}^{SD} by more than 10 % or 0,1 kGy, whichever is greater, the verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the verification dose experiment may be repeated. If this mean dose is less than 90 % of VD_{max}^{SD} and, on

performance of tests of sterility, acceptable results are observed (see <u>7.3.3.5</u>), the verification dose experiment need not be repeated.

7.3.3.4.2 Subject each irradiated product item individually to a test of sterility (see <u>5.4.1</u>) using the media and incubation conditions used in the original dose substantiation. Record the number of positive tests of sterility.

7.3.3.5 Stage 4: Interpretation of results

- **7.3.3.5.1** If no more than one positive test of sterility is obtained from the 10 tests carried out, accept the sterilization dose audit.
- **7.3.3.5.2** If two positive tests of sterility are obtained, perform a confirmatory sterilization dose audit (see <u>7.3.3.6</u>).
- **7.3.3.5.3** If three or more positive tests of sterility are obtained, do not accept the sterilization dose audit as the selected sterilization dose might be inadequate.

If the occurrence of these positive tests of sterility can be ascribed to incorrect performance of tests of sterility or incorrect delivery of $VD_{max}^{\ \ SD}$, or a specific bioburden-related cause, implement corrective action and repeat the verification dose experiment as soon as practicable using a further 10 product items from a batch manufactured under conditions that are representative of normal production and the same $VD_{max}^{\ \ SD}$ as that used in the sterilization dose audit that was not accepted. Interpret the results of the repeat verification dose experiment in accordance with 7.3.3.5.

If the occurrence of these positive tests of sterility cannot be ascribed to incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} , or a specific bioburden-related cause, the following shall apply.

a) If three to six positive tests of sterility are obtained, augment the selected sterilization dose immediately (see 7.3.3.7) and re-establish the sterilization dose as soon as practicable using another approach. Other approaches are:

selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 1 (6.2.2 or 6.3.3, as appropriate);

- 1) Method 1;
- 2) Method 2; and
- 3) a method providing assurance, in regard to achieving maximally an SAL of 10⁻⁶, equivalent to that of other methods of dose establishment. Continue to use the augmented sterilization dose until re-establishment of the sterilization dose is completed.
- b) If seven or more positive tests of sterility are obtained, discontinue sterilization at the selected sterilization dose. Do not augment the selected sterilization dose and do not resume sterilization until the sterilization dose is re-established using another approach. Other approaches are:

selection and substantiation of a higher sterilization dose than that for which verification was not accepted using Method VD_{max}^{SD} , starting at Stage 1 (6.2.2 or 6.3.3, as appropriate);

- 1) Method 1;
- 2) Method 2; and
- 3) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

7.3.3.6 Confirmatory sterilization dose audit

7.3.3.6.1 General

7.3.3.6.1.1 For the performance of a confirmatory sterilization dose audit for a selected sterilization dose, substantiated using Method VD_{max}^{SD} , use an SIP equivalent to that used originally in substantiating the sterilization dose.

7.3.3.6.1.2 In applying the confirmatory sterilization dose audit, the three stages below shall be followed.

7.3.3.6.2 Stage 1: Obtain samples of product

Select 10 product items from a single batch of product, in accordance with 5.1, 5.2 (if applicable) and $\underline{5.3}$. The 10 product items for the performance of confirmatory sterilization dose audit may be selected from either the batch used for the verification dose experiment carried out in the original sterilization dose audit (see $\underline{7.3.3}$) or a second batch manufactured under conditions that are representative of normal production (see $\underline{5.3}$).

7.3.3.6.3 Stage 2: Perform confirmatory verification dose experiment

7.3.3.6.3.1 Irradiate 10 product items at VD_{max}^{SD} used in the most recent successful substantiation of the selected sterilization dose.

The highest dose to product items shall not exceed $VD_{max}^{\ \ SD}$ by more than 10 % or 0,1 kGy, whichever is greater.

The arithmetic mean of the highest and lowest doses to product items should not be less than 90 % of $VD_{max}^{\ \ SD}$.

Determine the dose delivered (see <u>5.5</u>).

If the highest dose to product items exceeds VD_{max}^{SD} by more than 10 % or 0,1 kGy, whichever is greater, the confirmatory verification dose experiment shall be repeated.

If the arithmetic mean of the highest and lowest doses to product items is less than 90 % of VD_{max}^{SD} , the confirmatory verification dose experiment may be repeated. If this mean dose is less than 90 % of VD_{max}^{SD} and, on performance of tests of sterility, acceptable results are observed (see 7.3.3.6.4), the confirmatory verification dose experiment need not be repeated.

7.3.3.6.3.2 Subject each irradiated product item individually to a test of sterility (see 5.4.1) using the media and incubation conditions used originally in the substantiation of the selected sterilization dose and record the number of positive tests of sterility.

7.3.3.6.4 Stage 3: Interpretation of results

Interpret the results of the confirmatory verification dose audit performed in accordance with 7.3.3.6, as follows:

a) If there are no positive tests of sterility from the 10 tests carried out, accept the sterilization dose audit.

- b) If one to four positive tests of sterility are obtained, do not accept the sterilization dose audit. Augment the selected sterilization dose immediately (see <u>7.3.3.7</u>) and re-establish the sterilization dose as soon as practicable using another approach. Other approaches are:
 - 1) selection and substantiation of a higher sterilization dose than that for which the sterilization dose audit was not accepted, starting at Stage 1 (6.2.2 or 6.3.3, as appropriate);
 - 2) Method 1;
 - 3) Method 2; and
 - 4) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

 $Continue\ to\ use\ the\ augmented\ sterilization\ dose\ until\ re-establishment\ of\ the\ sterilization\ dose\ is\ completed.$

- c) If five or more positive tests of sterility are obtained, do not accept the sterilization dose audit. Discontinue sterilization at the selected sterilization dose. Do not augment the selected sterilization dose and do not resume sterilization until the sterilization dose is re-established using another approach. Other approaches are:
 - 1) selection and substantiation of a higher sterilization dose than that for which the sterilization dose audit was not accepted, starting at Stage 1 (6.2.2 or 6.3.3, as appropriate);
 - 2) Method 1;
 - 3) Method 2; and
 - 4) a method providing assurance, in regard to achieving maximally an SAL of 10^{-6} , equivalent to that of other methods of dose establishment.

If the occurrence of one or more positive tests of sterility can be ascribed to incorrect performance of tests of sterility or incorrect delivery of VD_{max}^{SD} , or a specific bioburden-related cause, implement corrective action and repeat the confirmatory sterilization dose audit (see 7.3.3.6) using a further 10 product items from a batch manufactured under conditions that are representative of normal production and the same VD_{max}^{SD} as that used in the original sterilization dose substantiation. Interpret the results, in accordance with a) to c) above.

7.3.3.7 Augmentation of a sterilization dose substantiated using Method VD_{max}^{SD}

From the previously identified table in <u>Clause 8</u> for the selected sterilization dose, obtain the dose augmentation value corresponding to the average bioburden (SIP equal to 1,0) as determined according to <u>7.3.3.3</u>. If the average bioburden is not given in the table, use the closest tabulated value greater than the average bioburden to obtain the dose augmentation value. Use this dose augmentation value in Equation (2) to calculate the augmented sterilization dose.

augmented sterilization dose = selected sterilization dose + dose augmentation value (2)

7.3.4 Failure of a sterilization dose audit

Following failure of a sterilization dose audit requiring the re-establishment of the sterilization dose, the cause of failure shall be investigated and correction and/or corrective action taken (see 4.4 of ISO 11137-1:2006). As part of the investigation, the effect of irradiating product at the sterilization dose that has failed sterilization dose audit on the achievement of the specified SAL for previously irradiated batches of product shall be considered and a risk assessment undertaken on their suitability for use. The investigation and subsequent actions shall be recorded (see 4.1.2 of ISO 11137-1:2006).

NOTE It might not be possible to determine the effect of irradiation at the sterilization dose that has failed the sterilization dose audit on achievement of this SAL until the sterilization dose has been re-established.

Table 5-17,5 kGy selected sterilization dose for which the upper limit of average bioburden is 9,0

SD = 17,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} 17,5 (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
Less than or equal to 0,1	0,0	NAa	3,5	
0,15	0,6	NAa	3,4	
0,20	1,0	NAa	3,3	
0,25	1,3	NAa	3,2	
0,30	1,5	NAa	3,2	
0,35	1,7	NAa	3,2	
0,40	1,9	NAa	3,1	
0,45	2,0	NAa	3,1	
0,50	2,1	NAa	3,1	
0,60	2,4	NAa	3,0	
0,70	2,5	NAa	3,0	
0,80	2,7	NAa	3,0	
0,90	2,8	NAa	2,9	
1,0	2,9	2,92	2,9	
1,5	3,3	2,83	2,8	
2,0	3,6	2,78	2,8	
2,5	3,8	2,74	2,7	
3,0	3,7	2,50	2,8	
3,5	3,6	2,32	2,8	
4,0	3,5	2,17	2,8	
4,5	3,4	2,05	2,8	
5,0	3,3	1,95	2,8	
5,5	3,3	1,87	2,9	
6,0	3,2	1,79	2,9	
6,5	3,1	1,73	2,9	
7,0	3,1	1,67	2,9	
7,5	3,0	1,62	2,9	
8,0	3,0	1,57	2,9	
8,5	3,0	1,53	2,9	
9,0	2,9	1,49	2,9	

Table 6-20 kGy selected sterilization dose for which the upper limit of average bioburden is 45

SD = 20 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} 20 (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
less than or equal to 0,1	0,0	NAa	4,0	
0,15	0,7	NAa	3,9	
0,20	1,1	NAa	3,8	
0,25	1,5	NAa	3,7	
0,30	1,7	NAa	3,7	
0,35	2,0	NAa	3,6	
0,40	2,1	NAa	3,6	
0,45	2,3	NAa	3,5	
0,50	2,5	NAa	3,5	
0,60	2,7	NAa	3,5	
0,70	2,9	NAa	3,4	
0,80	3,1	NAa	3,4	
0,90	3,2	NAa	3,4	
1,0	3,3	3,33	3,3	
1,5	3,8	3,24	3,2	
2,0	4,1	3,17	3,2	
2,5	4,4	3,13	3,1	
3,0	4,6	3,09	3,1	
3,5	4,7	3,06	3,1	
4,0	4,9	3,03	3,0	
4,5	5,0	3,01	3,0	
5,0	5,1	2,99	3,0	
5,5	5,2	2,97	3,0	
6,0	5,2	2,95	3,0	
6,5	5,3	2,94	2,9	
7,0	5,4	2,92	2,9	
7,5	5,5	2,91	2,9	
8,0	5,5	2,88	2,9	
8,5	5,5	2,82	2,9	
9,0	5,4	2,77	2,9	
9,5	5,4	2,72	2,9	
10	5,3	2,67	2,9	
11	5,3	2,58	2,9	
12	5,2	2,51	3,0	
13	5,2	2,44	3,0	
14	5,1	2,39	3,0	
15	5,1	2,33	3,0	

Table 6 (continued)

SD = 20 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ²⁰ (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)	
16	5,0	2,28	3,0	
17	5,0	2,24	3,0	
18	5,0	2,20	3,0	
19	4,9	2,16	3,0	
20	4,9	2,13	3,0	
21	4,9	2,10	3,0	
22	4,8	2,07	3,0	
23	4,8	2,04	3,0	
24	4,8	2,01	3,0	
25	4,8	1,99	3,0	
26	4,8	1,97	3,1	
27	4,7	1,94	3,1	
28	4,7	1,92	3,1	
29	4,7	1,90	3,1	
30	4,7	1,89	3,1	
31	4,7	1,87	3,1	
32	4,6	1,85	3,1	
33	4,6	1,83	3,1	
34	4,6	1,81	3,1	
35	4,6	1,80	3,1	
36	4,6	1,78	3,1	
37	4,6	1,77	3,1	
38	4,5	1,76	3,1	
39	4,5	1,74	3,1	
40	4,5	1,73	3,1	
41	4,5	1,72	3,1	
42	4,5	1,71	3,1	
43	4,5	1,70	3,1	
44	4,5	1,68	3,1	
45	4,4	1,67	3,1	

Table 7-22,5 kGy selected sterilization dose for which the upper limit of average bioburden is 220

SD = 22.5 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ^{22,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
less than or equal to 0,1	0,0	NAa	4,5
a Not applicable: SIP equal to 1,0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u> .			

Table 7 (continued)

SD = 22,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{22,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
0,15	0,8	NAa	4,3	
0,20	1,3	NAa	4,2	
0,25	1,7	NAa	4,2	
0,30	2,0	NAa	4,1	
0,35	2,2	NAa	4,1	
0,40	2,4	NAa	4,0	
0,45	2,6	NAa	4,0	
0,50	2,8	NAa	3,9	
0,60	3,0	NAa	3,9	
0,70	3,3	NAa	3,8	
0,80	3,4	NAa	3,8	
0,90	3,6	NAa	3,8	
1,0	3,8	3,75	3,8	
1,5	4,3	3,64	3,6	
2,0	4,6	3,57	3,6	
2,5	4,9	3,52	3,5	
3,0	5,1	3,47	3,5	
3,5	5,3	3,44	3,4	
4,0	5,5	3,41	3,4	
4,5	5,6	3,38	3,4	
5,0	5,7	3,36	3,4	
5,5	5,8	3,34	3,3	
6,0	5,9	3,32	3,3	
6,5	6,0	3,30	3,3	
7,0	6,1	3,29	3,3	
7,5	6,1	3,27	3,3	
8,0	6,2	3,26	3,3	
8,5	6,3	3,25	3,2	
9,0	6,3	3,24	3,2	
9,5	6,4	3,22	3,2	
10	6,4	3,21	3,2	
11	6,5	3,20	3,2	
12	6,6	3,18	3,2	
13	6,7	3,16	3,2	
14	6,8	3,15	3,1	
15	6,8	3,14	3,1	
16	6,9	3,12	3,1	

Table 7 (continued)

SD = 22,5 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ^{22,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
17	6,9	3,11	3,1
18	7,0	3,10	3,1
19	7,0	3,09	3,1
20	7,1	3,08	3,1
21	7,1	3,07	3,1
22	7,2	3,06	3,1
23	7,2	3,06	3,1
24	7,3	3,05	3,0
25	7,3	3,03	3,0
26	7,3	3,00	3,1
27	7,2	2,97	3,1
28	7,2	2,94	3,1
29	7,2	2,92	3,1
30	7,2	2,89	3,1
31	7,2	2,87	3,1
32	7,1	2,85	3,1
33	7,1	2,82	3,1
34	7,1	2,80	3,1
35	7,1	2,78	3,1
36	7,1	2,74	3,1
37	7,1	2,74	3,1
38	7,0	2,73	3,1
39	7,0	2,71	3,1
40	7,0	2,69	3,1
41	7,0	2,68	3,1
42	7,0	2,66	3,1
43	7,0	2,65	3,1
44	7,0	2,63	3,1
45	6,9	2,62	3,1
46	6,9	2,60	3,1
47	6,9	2,59	3,1
48	6,9	2,58	3,1
49	6,9	2,56	3,1
50	6,9	2,55	3,1
55	6,8	2,50	3,1
60	6,8	2,44	3,1
65	6,8	2,40	3,2

 Table 7 (continued)

	SD = 22,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{22,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)		
70	6,7	2,36	3,2		
75	6,7	2,32	3,2		
80	6,7	2,29	3,2		
85	6,6	2,26	3,2		
90	6,6	2,23	3,2		
95	6,6	2,21	3,2		
100	6,5	2,18	3,2		
110	6,5	2,14	3,2		
120	6,5	2,09	3,2		
130	6,4	2,06	3,2		
140	6,4	2,03	3,2		
150	6,4	2,00	3,2		
160	6,3	1,98	3,2		
170	6,3	1,95	3,2		
180	6,3	1,93	3,2		
190	6,2	1,90	3,3		
200	6,2	1,88	3,3		
220	6,2	1,85	3,3		
^a Not applicable: SIP equal to	1,0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u>				

Table 8 - 27,5 kGy selected sterilization dose for which the upper limit of average bioburden is 5 000 $\,$

	SD = 27.5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{27,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)		
less than or equal to 0,10	0,0	NAa	5,5		
0,15	0,9	NAa	5,3		
0,20	1,6	NAa	5,2		
0,25	2,0	NAa	5,1		
0,30	2,4	NAa	5,0		
0,35	2,7	NAa	5,0		
0,40	3,0	NAa	4,9		
0,45	3,2	NAa	4,9		
0,50	3,4	NAa	4,8		
0,60	3,7	NAa	4,8		
0,70	4,0	NAa	4,7		
0,80	4,2	NAa	4,7		
^a Not applicable: SIP equal to 1,0 re	quired; see <u>6.2.1.1</u> and <u>6.3.2.1</u> .				

Table 8 (continued)

SD = 27,5 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ^{27,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
0,90	4,4	NAa	4,6
1,0	4,6	4,58	4,6
2,0	5,7	4,36	4,4
3,0	6,3	4,25	4,2
4,0	6,7	4,17	4,2
5,0	7,0	4,11	4,1
6,0	7,2	4,06	4,1
7,0	7,4	4,02	4,0
8,0	7,6	3,98	4,0
9,0	7,7	3,95	4,0
10	7,9	3,93	3,9
11	8,0	3,91	3,9
12	8,1	3,88	3,9
13	8,2	3,87	3,9
14	8,3	3,85	3,8
15	8,3	3,83	3,8
16	8,4	3,82	3,8
17	8,5	3,80	3,8
18	8,5	3,79	3,8
19	8,6	3,78	3,8
20	8,7	3,77	3,8
21	8,7	3,76	3,8
22	8,8	3,75	3,7
23	8,8	3,74	3,7
24	8,9	3,73	3,7
25	8,9	3,72	3,7
26	9,0	3,71	3,7
27	9,0	3,70	3,7
28	9,0	3,69	3,7
29	9,1	3,69	3,7
30	9,1	3,68	3,7
31	9,1	3,67	3,7
32	9,2	3,66	3,7
33	9,2	3,66	3,7
34	9,2	3,65	3,7
35	9,3	3,65	3,6
36	9,3	3,64	3,6

Table 8 (continued)

Average bioburden	SIP equal to 1,0 VD _{max} ^{27,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
37	9,3	3,63	3,6
38	9,4	3,63	3,6
39	9,4	3,62	3,6
40	9,4	3,62	3,6
41	9,4	3,61	3,6
42	9,5	3,61	3,6
43	9,5	3,60	3,6
44	9,5	3,60	3,6
45	9,5	3,59	3,6
46	9,5		
47	9,6	3,59	3,6
48	9,6	3,58	3,6
49		3,58	3,6
	9,6	3,58	3,6
50	9,6	3,57	3,6
55	9,7	3,55	3,6
60	9,8	3,54	3,5
65	9,9	3,52	3,5
70	10,0	3,51	3,5
75	10,0	3,49	3,5
80	10,1	3,48	3,5
85	10,2	3,47	3,5
90	10,2	3,46	3,5
95	10,3	3,45	3,4
100	10,3	3,44	3,4
110	10,4	3,42	3,4
120	10,5	3,40	3,4
130	10,6	3,39	3,4
140	10,6	3,38	3,4
150	10,7	3,36	3,4
160	10,7	3,35	3,4
170	10,8	3,34	3,3
180	10,8	3,33	3,3
190	10,9	3,32	3,3
200	10,9	3,31	3,3
220	11,0	3,30	3,3
240	11,1	3,28	3,3
260	11,1	3,26	3,3

SD = 27,5 kGy Dose			
Average bioburden	SIP equal to 1,0 VD _{max} ^{27,5} (kGy)	SIP dose reduction factor	augmentation valu
280	11,1	3,21	3,3
300	11,1	3,18	3,3
325	11,0	3,14	3,3
350	11,0	3,10	3,3
375	11,0	3,07	3,3
400	10,9	3,04	3,3
425	10,9	3,01	3,3
450	10,9	2,98	3,3
475	10,9	2,96	3,3
500	10,9	2,93	3,3
525	10,8	2,91	3,3
550	10,8	2,89	3,3
575	10,8	2,87	3,3
600	10,8	2,85	3,3
650	10,8	2,82	3,4
700	10,7	2,79	3,4
750	10,7	2,76	3,4
800	10,7	2,73	3,4
850	10,7	2,71	3,4
900	10,6	2,69	3,4
950	10,6	2,67	3,4
1 000	10,6	2,65	3,4
1 050	10,6	2,63	3,4
1 100	10,6	2,61	3,4
1 150	10,6	2,60	3,4
1 200	10,5	2,58	3,4
1 250	10,5	2,57	3,4
1 300	10,5	2,55	3,4
1 350	10,5	2,54	3,4
1 400	10,5	2,53	3,4
1 450	10,5	2,51	3,4
1 500	10,5	2,50	3,4
1 550	10,4	2,49	3,4
1 600	10,4	2,48	3,4
1 650	10,4	2,47	3,4
1 700	10,4	2,46	3,4
1 750	10,4	2,45	3,4

 Table 8 (continued)

SD = 27.5 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ^{27,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
1 800	10,4	2,44	3,4
1 850	10,4	2,43	3,4
1 900	10,4	2,42	3,4
1 950	10,4	2,41	3,4
2 000	10,4	2,41	3,4
2 100	10,3	2,39	3,4
2 200	10,3	2,37	3,4
2 300	10,3	2,36	3,4
2 400	10,3	2,35	3,4
2 500	10,3	2,34	3,4
2 600	10,3	2,32	3,4
2 700	10,3	2,31	3,5
2 800	10,2	2,30	3,5
2 900	10,2	2,29	3,5
3 000	10,2	2,28	3,5
3 200	10,2	2,26	3,5
3 400	10,2	2,24	3,5
3 600	10,2	2,23	3,5
3 800	10,1	2,21	3,5
4 000	10,1	2,20	3,5
4 200	10,1	2,19	3,5
4 400	10,1	2,17	3,5
4 600	10,1	2,16	3,5
4 800	10,1	2,15	3,5
5 000	10,1	2,14	3,5

Table 9 — 30 kGy selected sterilization dose for which the upper limit of average bioburden is $23\,000$

SD = 30 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
less than or equal to 0,10	0,0	NAa	6,0
0,15	1,0	NAa	5,8
0,20	1,7	NAa	5,7
0,25	2,2	NAa	5,6
0,30	2,6	NAa	5,5
a Not applicable: SIP equal to 1,0	O required; see <u>6.2.1.1</u> and <u>6.3.2.1</u>		

Table 9 (continued)

SD = 30 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
0,35	2,9	NAa	5,4
0,40	3,2	NAa	5,4
0,45	3,5	NAa	5,3
0,50	3,7	NAa	5,3
0,60	4,0	NAa	5,2
0,70	4,3	NAa	5,1
0,80	4,6	NAa	5,1
0,90	4,8	NAa	5,0
1,0	5,0	5,00	5,0
2,0	6,2	4,76	4,8
3,0	6,8	4,63	4,6
4,0	7,3	4,54	4,5
5,0	7,6	4,48	4,5
6,0	7,9	4,43	4,4
7,0	8,1	4,38	4,4
8,0	8,3	4,35	4,3
9,0	8,4	4,31	4,3
10	8,6	4,29	4,3
11	8,7	4,26	4,3
12	8,8	4,24	4,2
13	8,9	4,22	4,2
14	9,0	4,20	4,2
15	9,1	4,18	4,2
16	9,2	4,16	4,2
17	9,3	4,15	4,1
18	9,3	4,13	4,1
19	9,4	4,12	4,1
20	9,5	4,11	4,1
21	9,5	4,10	4,1
22	9,6	4,09	4,1
23	9,6	4,08	4,1
24	9,7	4,06	4,1
25	9,7	4,06	4,1
26	9,8	4,05	4,0
27	9,8	4,04	4,0
28	9,9	4,03	4,0
29	9,9	4,02	4,0

Table 9 (continued)

Table 9 (continued) SD = 30 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
30	9,9	4,01	4,0
31	10,0	4,00	4,0
32	10,0	4,00	4,0
33	10,0	3,99	4,0
34	10,1	3,98	4,0
35	10,1	3,98	4,0
36	10,1	3,97	4,0
37	10,2	3,96	4,0
38	10,2	3,96	4,0
39	10,2	3,95	4,0
40	10,3	3,95	3,9
41	10,3	3,94	3,9
42	10,3	3,94	3,9
43	10,3	3,93	3,9
44	10,4	3,92	3,9
45	10,4	3,92	3,9
46	10,4	3,92	3,9
47	10,4	3,91	3,9
48	10,5	3,91	3,9
49	10,5	3,90	3,9
50	10,5	3,90	3,9
55	10,6	3,88	3,9
60	10,7	3,86	3,9
65	10,8	3,84	3,8
70	10,9	3,82	3,8
75	11,0	3,81	3,8
80	11,0	3,80	3,8
85	11,1	3,78	3,8
90	11,1	3,77	3,8
95	11,2	3,76	3,8
100	11,3	3,75	3,8
110	11,3	3,73	3,7
120	11,4	3,71	3,7
130	11,5	3,70	3,7
140	11,6	3,68	3,7
150	11,7	3,67	3,7
160	11,7	3,66	3,7
Not applicable: SIP equal to 1	.,0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u>		

Table 9 (continued)

Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
170	11,8	3,65	3,6
180	11,8	3,63	3,6
190	11,9	3,62	3,6
200	11,9	3,61	3,6
220	12,0	3,60	3,6
240	12,1	3,58	3,6
260	12,2	3,57	3,6
280	12,2	3,55	3,6
300	12,3	3,54	3,5
325	12,4	3,52	3,5
350	12,4	3,51	3,5
375	12,5	3,50	3,5
400	12,6	3,49	3,5
425	12,6	3,48	3,5
450	12,7	3,47	3,5
475	12,7	3,46	3,5
500	12,8	3,45	3,4
525	12,8	3,44	3,4
550	12,8	3,43	3,4
575	12,9	3,42	3,4
600	12,9	3,42	3,4
650	13,0	3,40	3,4
700	13,0	3,39	3,4
750	13,1	3,38	3,4
800	13,2	3,37	3,4
850	13,2	3,35	3,4
900	13,1	3,32	3,4
950	13,1	3,30	3,4
1 000	13,1	3,27	3,4
1 050	13,1	3,25	3,4
1 100	13,1	3,23	3,4
1 150	13,0	3,21	3,4
1 200	13,0	3,19	3,4
1 250	13,0	3,18	3,4
1 300	13,0	3,16	3,4
1 350	13,0	3,14	3,4
1 400	13,0	3,13	3,4

Table 9 (continued)

SD = 30 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)
1 450	13,0	3,11	3,4
1,500	13,0	3,10	3,4
1 550	12,9	3,09	3,4
1 600	12,9	3,07	3,4
1 650	12,9	3,06	3,4
1 700	12,9	3,05	3,4
1 750	12,9	3,04	3,4
1 800	12,9	3,03	3,4
1 850	12,9	3,02	3,4
1 900	12,9	3,01	3,4
1 950	12,9	3,00	3,4
2 000	12,9	2,99	3,4
2 100	12,8	2,97	3,4
2 200	12,8	2,95	3,4
2 300	12,8	2,93	3,4
2 400	12,8	2,92	3,4
2 500	12,8	2,90	3,4
2 600	12,8	2,89	3,4
2 700	12,8	2,88	3,5
2 800	12,7	2,86	3,5
2 900	12,7	2,85	3,5
3 000	12,7	2,84	3,5
3 200	12,7	2,82	3,5
3 400	12,7	2,80	3,5
3 600	12,7	2,78	3,5
3 800	12,6	2,76	3,5
4 000	12,6	2,74	3,5
4 200	12,6	2,73	3,5
4 400	12,6	2,71	3,5
4 600	12,6	2,70	3,5
4 800	12,6	2,69	3,5
5 000	12,6	2,67	3,5
5 300	12,5	2,65	3,5
5 600	12,5	2,63	3,5
5 900	12,5	2,62	3,5
6 200	12,5	2,61	3,5
6 500	12,5	2,59	3,5

Table 9 (continued)

SD = 30 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ³⁰ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
6 800	12,5	2,58	3,5	
7 100	12,4	2,56	3,5	
7 400	12,4	2,55	3,5	
7 700	12,4	2,54	3,5	
8 000	12,4	2,53	3,5	
8 500	12,4	2,52	3,5	
9 000	12,4	2,50	3,5	
9 500	12,4	2,48	3,5	
10 000	12,4	2,47	3,5	
10 500	12,3	2,46	3,5	
11 000	12,3	2,44	3,5	
11 500	12,3	2,43	3,5	
12 000	12,3	2,42	3,5	
13 000	12,3	2,40	3,5	
14 000	12,3	2,38	3,6	
15 000	12,2	2,36	3,6	
16 000	12,2	2,35	3,6	
17 000	12,2	2,33	3,6	
18 000	12,2	2,32	3,6	
19 000	12,2	2,31	3,6	
20 000	12,2	2,29	3,6	
21 000	12,1	2,28	3,6	
22 000	12,1	2,27	3,6	
23 000	12,1	2,26	3,6	
ot applicable: SIP equal to 1	1,0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u>			

Table 10 - 32,5 kGy selected sterilization dose for which the upper limit of average bioburden is $100\,000$

SD = 32,5 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} 32,5 (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
less than or equal to 0,10	0,0	NAa	6,5
0,15	1,1	NAa	6,3
0,20	1,8	NAa	6,1
0,25	2,4	NAa	6,0
0,30	2,8	NAa	5,9
0,35	3,2	NAa	5,9
^a Not applicable: SIP equal to 1,	0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u> .		

Table 10 (continued)

SD = 32,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)	
0,40	3,5	NAa	5,8	
0,45	3,8	NAa	5,7	
0,50	4,0	NAa	5,7	
0,60	4,4	NAa	5,6	
0,70	4,7	NAa	5,6	
0,80	5,0	NAa	5,5	
0,90	5,2	NAa	5,5	
1,0	5,4	5,42	5,4	
2,0	6,7	5,16	5,2	
3,0	7,4	5,02	5,0	
4,0	7,9	4,92	4,9	
5,0	8,2	4,85	4,9	
6,0	8,5	4,79	4,8	
7,0	8,8	4,75	4,7	
8,0	9,0	4,71	4,7	
9,0	9,1	4,67	4,7	
10	9,3	4,64	4,6	
11	9,4	4,62	4,6	
12	9,5	4,59	4,6	
13	9,7	4,57	4,6	
14	9,8	4,55	4,5	
15	9,9	4,53	4,5	
16	9,9	4,51	4,5	
17	10,0	4,49	4,5	
18	10,1	4,48	4,5	
19	10,2	4,47	4,5	
20	10,2	4,45	4,5	
21	10,3	4,44	4,4	
22	10,4	4,43	4,4	
23	10,4	4,41	4,4	
24	10,5	4,40	4,4	
25	10,5	4,39	4,4	
26	10,6	4,38	4,4	
27	10,6	4,37	4,4	
28	10,7	4,36	4,4	
29	10,7	4,36	4,4	
30	10,8	4,35	4,3	

 Table 10 (continued)

Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
31	10,8	4,34	4,3
32	10,8	4,33	4,3
33	10,9	4,32	4,3
34	10,9	4,32	4,3
35	11,0	4,31	4,3
36	11,0	4,30	4,3
37	11,0	4,29	4,3
38	11,1	4,29	4,3
39	11,1	4,28	4,3
40	11,1	4,28	4,3
41	11,2	4,27	4,3
42	11,2	4,26	4,3
43	11,2	4,26	4,3
44	11,2	4,25	4,3
45	11,3	4,25	4,2
46	11,3	4,24	4,2
47	11,3	4,24	4,2
48	11,3	4,23	4,2
49	11,4	4,23	4,2
50	11,4	4,22	4,2
55	11,5	4,20	4,2
60	11,6	4,18	4,2
65	11,7	4,16	4,2
70	11,8	4,14	4,1
75	11,9	4,13	4,1
80	11,9	4,11	4,1
85	12,0	4,10	4,1
90	12,1	4,09	4,1
95	12,1	4,07	4,1
100	12,2	4,06	4,1
110	12,3	4,04	4,0
120	12,4	4,02	4,0
130	12,5	4,01	4,0
140	12,6	3,99	4,0
150	12,6	3,98	4,0
160	12,7	3,96	4,0
170	12,8	3,95	3,9

Table 10 (continued)

SD = 32,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
180	12,8	3,94	3,9	
190	12,9	3,93	3,9	
200	12,9	3,92	3,9	
220	13,0	3,90	3,9	
240	13,1	3,88	3,9	
260	13,2	3,86	3,9	
280	13,3	3,85	3,8	
300	13,3	3,83	3,8	
325	13,4	3,82	3,8	
350	13,5	3,80	3,8	
375	13,5	3,79	3,8	
400	13,6	3,78	3,8	
425	13,7	3,77	3,8	
450	13,7	3,76	3,8	
475	13,8	3,75	3,7	
500	13,8	3,74	3,7	
525	13,9	3,73	3,7	
550	13,9	3,72	3,7	
575	13,9	3,71	3,7	
600	14,0	3,70	3,7	
650	14,1	3,69	3,7	
700	14,1	3,67	3,7	
750	14,2	3,66	3,7	
800	14,2	3,65	3,7	
850	14,3	3,64	3,6	
900	14,4	3,63	3,6	
950	14,4	3,62	3,6	
1 000	14,4	3,61	3,6	
1 050	14,5	3,60	3,6	
1 100	14,5	3,59	3,6	
1 150	14,6	3,59	3,6	
1 200	14,6	3,58	3,6	
1 250	14,6	3,57	3,6	
1 300	14,7	3,57	3,6	
1 350	14,7	3,56	3,6	
1 400	14,7	3,55	3,6	
1 450	14,8	3,55	3,5	

 Table 10 (continued)

SD = 32,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
1 500	14,8	3,54	3,5	
1 550	14,8	3,54	3,5	
1 600	14,8	3,53	3,5	
1 650	14,9	3,53	3,5	
1 700	14,9	3,52	3,5	
1 750	14,9	3,52	3,5	
1 800	14,9	3,51	3,5	
1 850	15,0	3,51	3,5	
1 900	15,0	3,50	3,5	
1 950	15,0	3,50	3,5	
2 000	15,0	3,49	3,5	
2 100	15,1	3,49	3,5	
2 200	15,1	3,48	3,5	
2 300	15,1	3,47	3,5	
2 400	15,2	3,46	3,5	
2 500	15,2	3,46	3,5	
2 600	15,2	3,45	3,5	
2 700	15,3	3,44	3,5	
2 800	15,2	3,43	3,5	
2 900	15,2	3,41	3,5	
3 000	15,2	3,40	3,5	
3 200	15,2	3,37	3,5	
3 400	15,2	3,35	3,5	
3 600	15,2	3,33	3,5	
3 800	15,1	3,31	3,5	
4 000	15,1	3,29	3,5	
4 200	15,1	3,27	3,5	
4 400	15,1	3,25	3,5	
4 600	15,1	3,23	3,5	
4 800	15,1	3,22	3,5	
5 000	15,1	3,20	3,5	
5 300	15,0	3,18	3,5	
5 600	15,0	3,16	3,5	
5 900	15,0	3,14	3,5	
6 200	15,0	3,13	3,5	
6 500	15,0	3,11	3,5	
6 800	15,0	3,09	3,5	

 Table 10 (continued)

SD = 32,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)	
7 100	14,9	3,08	3,5	
7 400	14,9	3,07	3,5	
7 700	14,9	3,05	3,5	
8 000	14,9	3,04	3,5	
8 500	14,9	3,02	3,5	
9 000	14,9	3,00	3,5	
9 500	14,9	2,99	3,5	
10 000	14,9	2,97	3,5	
10 500	14,8	2,95	3,5	
11 000	14,8	2,94	3,5	
11 500	14,8	2,93	3,5	
12 000	14,8	2,91	3,5	
13 000	14,8	2,89	3,5	
14 000	14,8	2,87	3,6	
15 000	14,7	2,85	3,6	
16 000	14,7	2,83	3,6	
17 000	14,7	2,81	3,6	
18 000	14,7	2,79	3,6	
19 000	14,7	2,78	3,6	
20 000	14,7	2,76	3,6	
21 000	14,6	2,75	3,6	
22 000	14,6	2,74	3,6	
23 000	14,6	2,73	3,6	
24 000	14,6	2,72	3,6	
25 000	14,6	2,70	3,6	
26 000	14,6	2,69	3,6	
27 000	14,6	2,68	3,6	
28 000	14,6	2,67	3,6	
29 000	14,6	2,66	3,6	
30 000	14,6	2,66	3,6	
32 000	14,5	2,64	3,6	
34 000	14,5	2,62	3,6	
36 000	14,5	2,61	3,6	
38 000	14,5	2,60	3,6	
40 000	14,5	2,58	3,6	
42 000	14,5	2,57	3,6	
44 000	14,5	2,56	3,6	

 Table 10 (continued)

	SD = 32,5 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ^{32,5} (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)		
46 000	14,4	2,55	3,6		
48 000	14,4	2,54	3,6		
50 000	14,4	2,53	3,6		
54 000	14,4	2,51	3,6		
58 000	14,4	2,50	3,6		
62 000	14,4	2,48	3,6		
66 000	14,4	2,47	3,6		
70 000	14,3	2,45	3,6		
75 000	14,3	2,44	3,6		
80 000	14,3	2,42	3,6		
85 000	14,3	2,41	3,6		
90 000	14,3	2,40	3,6		
95 000	14,3	2,39	3,6		
100 000	14,3	2,38	3,6		
^a Not applicable: SIP equal to 1	1,0 required; see <u>6.2.1.1</u> and <u>6.3.2.1</u> .				

Table 11 - 35 kGy selected sterilization dose for which the upper limit of average bioburden is 440 000 $\,$

SD = 35 kGy				
Average bioburden	SIP equal to 1,0 VD _{max} ³⁵ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)	
less than or equal to 0,10	0,0	NAa	7,0	
0,15	1,2	NAa	6,8	
0,20	2,0	NAa	6,6	
0,25	2,6	NAa	6,5	
0,30	3,0	NAa	6,4	
0,35	3,4	NAa	6,3	
0,40	3,8	NAa	6,2	
0,45	4,0	NAa	6,2	
0,50	4,3	NAa	6,1	
0,60	4,7	NAa	6,1	
0,70	5,1	NAa	6,0	
0,80	5,4	NAa	5,9	
0,90	5,6	NAa	5,9	
1,0	5,8	5,83	5,8	
2,0	7,2	5,55	5,6	
3,0	8,0	5,40	5,4	

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} 35 (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)
4,0	8,5	5,30	5,3
5,0	8,9	5,22	5,2
6,0	9,2	5,16	5,2
7,0	9,4	5,11	5,1
8,0	9,6	5,07	5,1
9,0	9,8	5,03	5,0
10	10,0	5,00	5,0
11	10,1	4,97	5,0
12	10,3	4,94	4,9
13	10,4	4,92	4,9
14	10,5	4,90	4,9
15	10,6	4,88	4,9
16	10,7	4,86	4,9
17	10,8	4,84	4,8
18	10,9	4,82	4,8
19	11,0	4,81	4,8
20	11,0	4,79	4,8
21	11,1	4,78	4,8
22	11,2	4,77	4,8
23	11,2	4,75	4,8
24	11,3	4,74	4,7
25	11,3	4,73	4,7
26	11,4	4,72	4,7
27	11,5	4,71	4,7
28	11,5	4,70	4,7
29	11,5	4,69	4,7
30	11,6	4,68	4,7
31	11,6	4,67	4,7
32	11,7	4,66	4,7
33	11,7	4,66	4,7
34	11,8	4,65	4,6
35	11,8	4,64	4,6
36	11,8	4,63	4,6
37	11,9	4,62	4,6
38	11,9	4,62	4,6
39	11,9	4,61	4,6
40	12,0	4,60	4,6

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁵ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
41	12,0	4,60	4,6
42	12,0	4,59	4,6
43	12,1	4,59	4,6
44	12,1	4,58	4,6
45	12,1	4,57	4,6
46	12,2	4,57	4,6
47	12,2	4,56	4,6
48	12,2	4,56	4,6
49	12,2	4,55	4,6
50	12,3	4,55	4,5
55	12,4	4,52	4,5
60	12,5	4,50	4,5
65	12,6	4,48	4,5
70	12,7	4,46	4,5
75	12,8	4,44	4,4
80	12,9	4,43	4,4
85	12,9	4,41	4,4
90	13,0	4,40	4,4
95	13,1	4,39	4,4
100	13,1	4,38	4,4
110	13,2	4,35	4,4
120	13,3	4,33	4,3
130	13,4	4,31	4,3
140	13,5	4,30	4,3
150	13,6	4,28	4,3
160	13,7	4,27	4,3
170	13,7	4,25	4,3
180	13,8	4,24	4,2
190	13,9	4,23	4,2
200	13,9	4,22	4,2
220	14,0	4,20	4,2
240	14,1	4,18	4,2
260	14,2	4,16	4,2
280	14,3	4,14	4,1
300	14,4	4,13	4,1
325	14,4	4,11	4,1
350	14,5	4,10	4,1

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} 35 (kGy)	SIP dose reduction factor	Dose augmentation valu (kGy)
375	14,6	4,08	4,1
400	14,7	4,07	4,1
425	14,7	4,06	4,1
450	14,8	4,04	4,0
475	14,8	4,03	4,0
500	14,9	4,02	4,0
525	14,9	4,01	4,0
550	15,0	4,00	4,0
575	15,0	4,00	4,0
600	15,1	3,99	4,0
650	15,1	3,97	4,0
700	15,2	3,96	4,0
750	15,3	3,94	3,9
800	15,3	3,93	3,9
850	15,4	3,92	3,9
900	15,5	3,91	3,9
950	15,5	3,90	3,9
1 000	15,6	3,89	3,9
1 050	15,6	3,88	3,9
1 100	15,6	3,87	3,9
1 150	15,7	3,86	3,9
1 200	15,7	3,85	3,9
1 250	15,8	3,85	3,8
1 300	15,8	3,84	3,8
1 350	15,8	3,83	3,8
1 400	15,9	3,83	3,8
1 450	15,9	3,82	3,8
1 500	15,9	3,81	3,8
1 550	16,0	3,81	3,8
1 600	16,0	3,80	3,8
1 650	16,0	3,80	3,8
1 700	16,0	3,79	3,8
1 750	16,1	3,79	3,8
1 800	16,1	3,78	3,8
1 850	16,1	3,78	3,8
1 900	16,1	3,77	3,8
1 950	16,2	3,77	3,8

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} 35 (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
2 000	16,2	3,76	3,8
2 100	16,2	3,75	3,8
2 200	16,3	3,75	3,7
2 300	16,3	3,74	3,7
2 400	16,3	3,73	3,7
2 500	16,4	3,72	3,7
2 600	16,4	3,72	3,7
2 700	16,4	3,71	3,7
2 800	16,5	3,70	3,7
2 900	16,5	3,70	3,7
3 000	16,5	3,69	3,7
3 200	16,6	3,68	3,7
3 400	16,6	3,67	3,7
3 600	16,7	3,66	3,7
3 800	16,7	3,65	3,7
4 000	16,8	3,65	3,6
4 200	16,8	3,64	3,6
4 400	16,9	3,63	3,6
4 600	16,9	3,62	3,6
4 800	16,9	3,62	3,6
5 000	17,0	3,61	3,6
5 300	17,0	3,60	3,6
5 600	17,0	3,59	3,6
5 900	17,1	3,58	3,6
6 200	17,1	3,57	3,6
6 500	17,2	3,57	3,6
6 800	17,2	3,56	3,6
7 100	17,2	3,55	3,6
7 400	17,3	3,55	3,5
7 700	17,3	3,54	3,5
8 000	17,3	3,53	3,5
8 500	17,4	3,52	3,5
9 000	17,4	3,51	3,5
9 500	17,4	3,49	3,5
10 000	17,4	3,47	3,5
10 500	17,3	3,45	3,5
11 000	17,3	3,44	3,5

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} ³⁵ (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
11 500	17,3	3,42	3,5
12 000	17,3	3,41	3,5
13 000	17,3	3,38	3,5
14 000	17,3	3,35	3,6
15 000	17,2	3,33	3,6
16 000	17,2	3,31	3,6
17 000	17,2	3,29	3,6
18 000	17,2	3,27	3,6
19 000	17,2	3,25	3,6
20 000	17,2	3,24	3,6
21 000	17,1	3,22	3,6
22 000	17,1	3,20	3,6
23 000	17,1	3,19	3,6
24 000	17,1	3,18	3,6
25 000	17,1	3,17	3,6
26 000	17,1	3,16	3,6
27 000	17,1	3,14	3,6
28 000	17,1	3,13	3,6
29 000	17,1	3,12	3,6
30 000	17,1	3,11	3,6
32 000	17,0	3,09	3,6
34 000	17,0	3,08	3,6
36 000	17,0	3,06	3,6
38 000	17,0	3,04	3,6
40 000	17,0	3,03	3,6
42 000	17,0	3,02	3,6
44 000	17,0	3,00	3,6
46 000	16,9	2,99	3,6
48 000	16,9	2,98	3,6
50 000	16,9	2,97	3,6
54 000	16,9	2,95	3,6
58 000	16,9	2,93	3,6
62 000	16,9	2,91	3,6
66 000	16,9	2,90	3,6
70 000	16,8	2,88	3,6
75 000	16,8	2,86	3,6
80 000	16,8	2,85	3,6

Table 11 (continued)

SD = 35 kGy			
Average bioburden	SIP equal to 1,0 VD _{max} 35 (kGy)	SIP dose reduction factor	Dose augmentation value (kGy)
85 000	16,8	2,83	3,6
90 000	16,8	2,82	3,6
95 000	16,8	2,80	3,6
100 000	16,8	2,79	3,6
110 000	16,7	2,77	3,7
120 000	16,7	2,75	3,7
130 000	16,7	2,73	3,7
140 000	16,7	2,71	3,7
150 000	16,7	2,70	3,7
160 000	16,7	2,68	3,7
170 000	16,6	2,67	3,7
180 000	16,6	2,66	3,7
190 000	16,6	2,65	3,7
200 000	16,6	2,63	3,7
220 000	16,6	2,61	3,7
240 000	16,6	2,60	3,7
260 000	16,5	2,58	3,7
280 000	16,5	2,56	3,7
300 000	16,5	2,55	3,7
320 000	16,5	2,54	3,7
340 000	16,5	2,52	3,7
360 000	16,5	2,51	3,7
380 000	16,5	2,50	3,7
400 000	16,5	2,49	3,7
420 000	16,5	2,48	3,7
440 000	16,4	2,47	3,7

9 Worked examples

9.1 Substantiation of a selected sterilization dose of 17,5 kGy (SIP less than 1,0)

Table 12 shows a worked example for Method $VD_{max}^{17,5}$. The example is for a product with multiple production batches and that was too large for testing so a defined portion of the product item (SIP less than 1,0) was used.

Table 12 — Method VD_{max}^{17,5} substantiation (SIP less than 1,0)

Term	Value	Comment
Stage 1		

Table 12 (continued)

Term	Value	Comment
SAL	10-6	This method substantiates 17,5 kGy as a sterilization dose to achieve maximally an SAL of 10^{-6} .
SIP	0,5	The product was too large for the ready performance of tests of sterility; a 1/2 portion was selected for testing.
Number of product items	40	Ten from each of three independent batches for bioburden determination plus 10 for the verification dose experiment.
Stage 2		
Average bioburden	6	The SIP bioburden results from the three batches gave averages of 2, 3 and 4.
		The average bioburden for the entire product item for each of the batches was calculated:
		2/0,5 = 4
		3/0,5 = 6
		4/0,5 = 8
		The overall average bioburden was 6.
		The highest batch average bioburden was 8.
		The highest batch average bioburden of 8 is less than two times the overall average bioburden of 6. Therefore, 6 was used to obtain the SIP equal to $1.0~\rm VD_{max}^{17.5}$.
Stage 3		
Selection of sterilization dose	17.5 kGy	All three batch average bioburden values are less than or equal to 9,0, as is the overall average bioburden.
Stage 4		
SIP verification dose	2,7 kGy	Table 5 was used to obtain values of SIP equal to 1,0 $VD_{max}^{17,5}$ and SIP dose reduction factor at an average bioburden of 6. The value of SIP $VD_{max}^{17,5}$ dose for an SIP of 0,5 was calculated using Equation (1):
		$SIP VD_{max}^{17,5} =$
		(SIP equal to 1,0 $VD_{max}^{17,5}$) + (SIP dose reduction factor x log SIP)
		[Equation (1)]
		For the example:
		SIP $VD_{max}^{17,5} = 3.2 \text{ kGy} + (1.79 \text{ kGy} \times \log 0.5) = 2.7 \text{ kGy}$
Stage 5		
Verification dose experiment	2,3 to 2,9 kGy	The dose to SIPs in the verification dose experiment ranged from 2,3 to 2,9 kGy. The highest dose to product items is less than the calculated upper limit (3,0 kGy) and the arithmetic mean of the highest and lowest doses, 2,6 kGy, is not less than 90 % of the verification dose (90 % of 2,7 kGy is 2,4 kGy).
Stage 6		
Interpretation of results	0 positives	Doses are within the calculated limits and the results of tests of sterility are acceptable (i.e. less than or equal to one positive); therefore, verification was accepted and 17,5 kGy was substantiated as the sterilization dose.

9.2 Substantiation of a selected sterilization dose of 30 kGy (SIP equal to 1,0)

Table 13 shows a worked example for Method VD_{max}^{30} . The example is for a product with multiple production batches and for which the entire product item (SIP equal to 1,0) was used for testing.

Table 13 — Method VD_{max}^{30} substantiation (SIP equal to 1,0)

Term	Value	Comment
Stage 1	•	
SAL	10-6	This method substantiates 30 kGy as a sterilization dose to achieve maximally an SAL of 10^{-6} .
SIP	1,0	The entire product was used for testing.
Number of product items	40	Ten from each of three independent batches for bioburden determination plus 10 for the verification dose experiment.
Stage 2		
Average bioburden	5643	The bioburden results from the three batches gave averages of 5030, 5700, and 6200.
		The highest batch average bioburden is 6200.
		The highest batch average bioburden of 6200 was less than two times the overall average bioburden of 5643. Therefore, the overall average bioburden of 5643 was used to obtain the SIP equal to 1,0 VD_{max}^{30} .
Stage 3		
Selection of sterili-	30 kGy	All three batch average bioburden values are less than or equal to
zation dose		23 000, as is the overall average bioburden.
Stage 4		
Verification dose	12,5 kGy	An average bioburden of 5643 is not listed in <u>Table 9</u> so the next greater bioburden of 5900 was used to obtain SIP equal to 1,0 VD_{max}^{30} .
Stage 5		
Verification dose experiment	11,4 to 13,0 kGy	The dose to product items in the verification dose experiment ranged from 11,4 to 13,0 kGy. The highest dose to product items is less than the calculated upper limit (13,8 kGy) and the arithmetic mean of the highest and lowest doses, 12,2 kGy, is not less than 90 % of the verification dose (90 % of 12,5 kGy is 11,3 kGy).
Stage 6		
Interpretation of results	0 positives	Doses are within the calculated limits and the results of tests of sterility are acceptable (i.e. less than or equal to one positive); therefore, verification was accepted and 30 kGy was substantiated as the sterilization dose.

9.3~ Sterilization dose audit for a sterilization dose substantiated using Method $VD_{max}{}^{22,5}$, the findings from which necessitated augmentation of the sterilization dose

Table 14 shows the findings from a sterilization dose audit performed after a selected sterilization dose of 22,5 kGy had been substantiated.

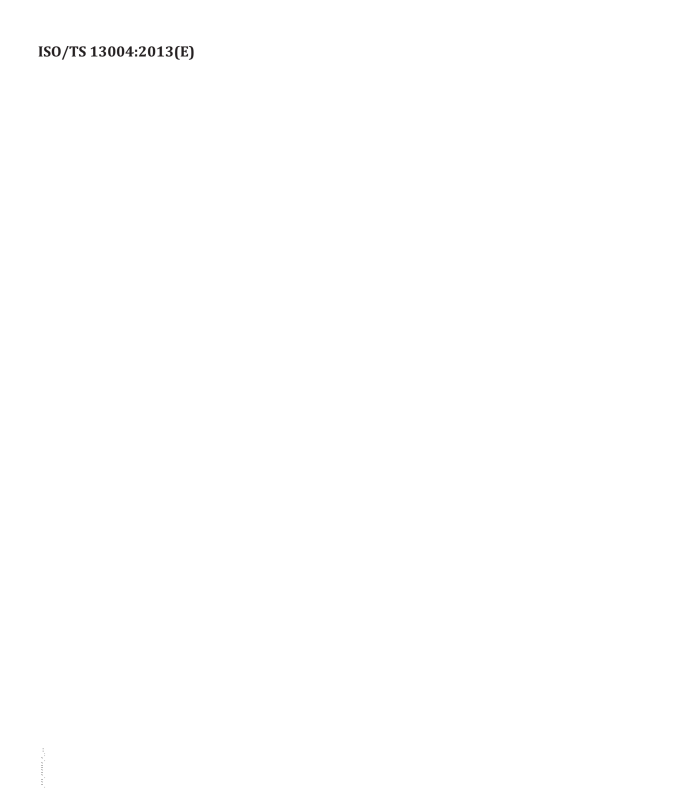
 $Table~14-Sterilization~dose~audit~following~which~augmentation~of~the~sterilization~dose~was~required~(selected~sterilization~dose~substantiated~using~Method~VD_{max}^{~22,5})$

Term	Value	Comment			
Sterilization dos	e audit				
Stage 1					
Number of prod- uct items	20	20 product items were obtained from a single production batch.			
Stage 2					
SIP	0,5	The original substantiation of 22,5 kGy was conducted using an SIP of 0,5.			
SIP average bioburden	99	The average bioburden for the 10 SIPs tested was 99.			
Average biobur- den	198	The average bioburden for the entire product was calculated as follows: 99/0,5 = 198			
Stage 3					
Verification dose experiment	5,2 to 6,0 kGy	The original substantiation of 22,5 kGy was conducted at a verification dose of 5,6 kGy. Ten SIPs were irradiated at this dose. The dose to SIPs in the verification dose experiment ranged from 5,2 to 6,0 kGy. The highest dose to SIPs is less than the calculated upper limit $(6,2\ kGy)$ and the arithmetic mean of the highest and lowest doses, 5,6 kGy, is not less than 90 % of the verification dose (90 % of 5,6 kGy is 5,0 kGy).			
Stage 4					
Interpretation of results	2 positives	Doses are within the calculated limits but the occurrence of two positive tests of sterility requires that a confirmatory sterilization dose audit be conducted.			
Confirmatory ste	erilization d	ose audit			
Stage 1					
Number of product items	10	Ten product items were obtained from a single production batch.			
Stage 2					
Confirmatory verification dose experiment	5,1 kGy to 5,9 kGy	The verification dose for the confirmatory sterilization dose audit is the same as the original verification dose. Ten SIPs were irradiated at this dose. The dose to SIPs in the confirmatory verification dose experiment ranged from 5,1 kGy to 5,9 kGy. The highest dose to SIPs is less than the calculated upper limit (6,2 kGy) and the arithmetic mean of the highest and lowest doses, 5,5 kGy, is not less than 90 % of the verification dose (90 % of 5,6 kGy is 5,0 kGy).			
Stage 3	Stage 3				
Interpretation of results	1 positive	Doses are within the calculated limits but the occurrence of one positive test of sterility requires that the 22,5 kGy sterilization dose be augmented immediately and that the sterilization dose be re-established using another approach.			
Sterilization dos	Sterilization dose augmentation				
Average biobur- den	198	The average bioburden for the entire product that was determined in performing the audit.			
Dose augmenta- tion value	3,3 kGy	The average bioburden and Table 7 is used to obtain the dose augmentation value. A bioburden of 198 was not listed in the table, so the next greater tabulated average bioburden of 200 is used.			
Augmented steri- lization dose	25,8 kGy	augmented sterilization dose (kGy) = 22,5 kGy + dose augmentation value (kGy) [Equation (2)]			
		For the example:			
		augmented sterilization dose (kGy) = 22,5 kGy + 3,3 kGy = 25,8 kGy			

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