Sterilization of health care products — Biological indicators — Guidance for the selection, use and interpretation of results

Stérilisation des produits de santé — Indicateurs biologiques — Directives générales pour la sélection, l’utilisation et l’interprétation des résultats

ISO 14161:2009
https://standards.iteh.ai/catalog/standards/sist/b0b91e0f-d372-4226-a95f-7e959a5f6d6/iso-14161-2009
## Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foreword</td>
<td>v</td>
</tr>
<tr>
<td>Introduction</td>
<td>vi</td>
</tr>
<tr>
<td>1. Scope</td>
<td>1</td>
</tr>
<tr>
<td>2. Normative references</td>
<td>1</td>
</tr>
<tr>
<td>3. Terms and definitions</td>
<td>2</td>
</tr>
<tr>
<td>4. General</td>
<td>5</td>
</tr>
<tr>
<td>5. Characteristics of biological indicators</td>
<td>7</td>
</tr>
<tr>
<td>5.1. General</td>
<td>7</td>
</tr>
<tr>
<td>5.2. Test organism suspension for direct inoculation of products</td>
<td>7</td>
</tr>
<tr>
<td>5.3. Inoculated carriers</td>
<td>8</td>
</tr>
<tr>
<td>5.4. Self-contained biological indicators</td>
<td>8</td>
</tr>
<tr>
<td>5.5. Other biological indicators</td>
<td>9</td>
</tr>
<tr>
<td>6. Selection of supplier</td>
<td>9</td>
</tr>
<tr>
<td>6.1. General</td>
<td>9</td>
</tr>
<tr>
<td>6.2. Documentation</td>
<td>10</td>
</tr>
<tr>
<td>7. Biological indicators in process development</td>
<td>11</td>
</tr>
<tr>
<td>7.1. General</td>
<td>11</td>
</tr>
<tr>
<td>7.2. Overkill approach</td>
<td>11</td>
</tr>
<tr>
<td>7.3. Combined biological indicator and bioburden method</td>
<td>12</td>
</tr>
<tr>
<td>7.4. Bioburden method</td>
<td>13</td>
</tr>
<tr>
<td>8. Biological indicators in sterilization validation</td>
<td>14</td>
</tr>
<tr>
<td>8.1. General</td>
<td>14</td>
</tr>
<tr>
<td>8.2. Placement and handling of biological indicators</td>
<td>14</td>
</tr>
<tr>
<td>8.3. Sterilizer qualification</td>
<td>14</td>
</tr>
<tr>
<td>8.4. Performance qualification</td>
<td>14</td>
</tr>
<tr>
<td>8.5. Review and approval of validation</td>
<td>15</td>
</tr>
<tr>
<td>8.6. Requalification</td>
<td>15</td>
</tr>
<tr>
<td>9. Biological indicators in routine monitoring</td>
<td>15</td>
</tr>
<tr>
<td>9.1. General</td>
<td>15</td>
</tr>
<tr>
<td>9.2. Placement and handling of biological indicators</td>
<td>15</td>
</tr>
<tr>
<td>9.3. Process challenge device (PCD)</td>
<td>16</td>
</tr>
<tr>
<td>10. General</td>
<td>16</td>
</tr>
<tr>
<td>10.1. Results</td>
<td>16</td>
</tr>
<tr>
<td>10.2. Interpretation of results</td>
<td>16</td>
</tr>
<tr>
<td>11. Application of biological indicator standards</td>
<td>17</td>
</tr>
<tr>
<td>11.1. General assessment of biological indicator performance by the user</td>
<td>17</td>
</tr>
<tr>
<td>11.2. Nominal population of test organism</td>
<td>17</td>
</tr>
<tr>
<td>11.3. Resistance determination</td>
<td>18</td>
</tr>
<tr>
<td>11.4. ( \varepsilon ) value determination</td>
<td>20</td>
</tr>
<tr>
<td>11.5. ( F_{T, 2} ) equivalent sterilization value determination</td>
<td>22</td>
</tr>
<tr>
<td>11.6. Establishing spore-log-reduction (SLR)</td>
<td>23</td>
</tr>
<tr>
<td>11.7. Sterility assurance level (SAL) calculation</td>
<td>23</td>
</tr>
<tr>
<td>11.8. Test equipment</td>
<td>24</td>
</tr>
<tr>
<td>12. Culture conditions</td>
<td>24</td>
</tr>
<tr>
<td>12.1. General</td>
<td>24</td>
</tr>
<tr>
<td>12.2. Incubation temperature</td>
<td>24</td>
</tr>
</tbody>
</table>
12.3 Incubation period

12.4 Choice of growth medium

13 Third-party requirements

13.1 General

13.2 Minimum requirements for replicates and total number of biological indicators

13.3 Test equipment

14 Personnel training

15 Storage and handling

16 Disposal of biological indicators

Annex A (informative) Microbiological inactivation kinetics and enumeration techniques

Annex B (informative) Process challenge devices

Annex C (informative) Formulae for fraction negative methods for $D$ value calculations

Annex D (informative) Examples of documentation for biological indicators prepared by the user

Annex E (informative) Calculation of $z$ value

Annex F (informative) $D$ value determination by survivor curve method

Annex G (informative) Survival-kill response characteristics

Bibliography
Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

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.

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 14161 was prepared by Technical Committee ISO/TC 198, Sterilization of health care products.

This second edition cancels and replaces the first edition (ISO 14161:2000), which has been technically revised.
Introduction

This International Standard provides guidance regarding the selection, use and interpretation of results of biological indicators when used to develop, validate and monitor sterilization processes. The procedures described in this International Standard are of a general nature and do not, of themselves, constitute a comprehensive development, validation or monitoring programme with regard to the sterilization of health care products. The intent of this International Standard is not to mandate the use of biological indicators in a process but, if they are used, to provide guidance for their proper selection and use in order to obviate misleading results.

In this International Standard, users will find guidance on selection of the correct biological indicator for their particular sterilization process and critical parameters as well as guidance on its appropriate use.

The user should select a biological indicator that is appropriate for the particular process to be used. There is a wide variety of sterilization processes in common use, and biological indicator manufacturers are not able to foresee all possible uses of their product. Manufacturers, therefore, label biological indicators according to their intended use. It is the responsibility of the users of biological indicators to select, use, recover and interpret the results as appropriate for the particular sterilization process used.

The certified performance of a biological indicator can be adversely affected by the conditions of storage and transport prior to its use, by the use of the biological indicator or by the sterilizer process parameters. In addition, the incubation procedure used after exposure to the process, including outgrowth temperature and culture medium type, supplier and specific lot, can affect measured resistance as a function of recovery and growth. For these reasons, the recommendations of the biological indicator manufacturer for storage and use should be followed. After exposure, biological indicators should be aseptically transferred (if applicable) and incubated as specified by the biological indicator manufacturer.

It should be noted that biological indicators are not intended to indicate that the products in the load being sterilized are sterile. Biological indicators are utilized to test the effectiveness of a given sterilization process and the equipment used, by assessing microbial lethality according to the concept of sterility assurance level. Suitably trained personnel should conduct these studies.
Sterilization of health care products — Biological indicators — Guidance for the selection, use and interpretation of results

1 Scope

This International Standard provides guidance for the selection, use and interpretation of results from application of biological indicators when used in the development, validation and routine monitoring of sterilization processes. This International Standard applies to biological indicators for which International Standards exist.

NOTE 1 See, for example, the ISO 11138 series.

NOTE 2 The general information provided in this International Standard can have useful application for processes and biological indicators not currently addressed by existing International Standards, e.g., new and developing sterilization processes.

This International Standard does not consider those processes that rely solely on physical removal of microorganisms, e.g., filtration.

This International Standard is not intended to apply to combination processes using, for example, washer disinfectors or flushing and steaming of pipelines.

This International Standard is not intended to apply to liquid sterilization processes.

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 11135-1, Sterilization of health care products — Ethylene oxide — Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices

ISO 11138-1:2006, Sterilization of health care products — Biological indicators — Part 1: General requirements

ISO 11138-2, Sterilization of health care products — Biological indicators — Part 2: Biological indicators for ethylene oxide sterilization processes

ISO 11138-3, Sterilization of health care products — Biological indicators — Part 3: Biological indicators for moist heat sterilization processes

ISO 11138-4, Sterilization of health care products — Biological indicators — Part 4: Biological indicators for dry heat sterilization processes

ISO 11138-5, Sterilization of health care products — Biological indicators — Part 5: Biological indicators for low-temperature steam and formaldehyde sterilization processes

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

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

3.1 accreditation
procedure by which an authoritative body gives formal recognition that a body or person is competent to carry out specific tasks

NOTE 1 See ISO/IEC 17011[3].

NOTE 2 Accreditation does not itself qualify the laboratory to approve any particular product. However, accreditation can be relevant to approval and certification authorities when they decide whether or not to accept data produced by a given laboratory in connection with their own activities.

3.2 aseptic technique
conditions and procedures used to exclude the introduction of microbial contamination

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

[ISO/TS 11139, definition 2.2]

3.4 biological indicator
Bi test system containing viable microorganisms providing a defined resistance to a specified sterilization process

[ISO/TS 11139, definition 2.3]

3.5 D value
$D_{10}$ value
time or dose required to achieve inactivation of 90 % of a population of the test microorganism under stated conditions

[ISO/TS 11139, definition 2.11]

3.6 holding time
period for which the sterilization variable within the sterilizer and at all points within the load are continuously within the limits specified for the sterilization stage

3.7 inoculated carrier
supporting material on or in which a defined number of viable test organisms have been deposited

NOTE 1 See ISO 11138-1.

NOTE 2 The test organism is a microorganism used for the manufacture of inoculated carriers.
3.8 installation qualification
IQ
process of obtaining and documenting evidence that equipment has been provided and installed in accordance with its specification

[ISO/TS 11139, definition 2.22]

3.9 inoculation
addition of a defined amount of a characterized microbial entity into or on to an item

3.10 log reduction
LR
reduction in number of viable microorganisms, expressed in log units

3.11 operational qualification
OQ
process of obtaining and documenting evidence that installed equipment operates within predetermined limits when used in accordance with its operational procedures

[ISO/TS 11139, definition 2.27]

3.12 performance qualification
PQ
process of obtaining and documenting evidence that the equipment, as installed and operated in accordance with operational procedures, consistently performs in accordance with predetermined criteria and thereby yields product meeting its specification

[ISO/TS 11139, definition 2.30]

3.13 process challenge device
PCD
item designed to constitute a defined resistance to a sterilization process and used to assess performance of the process

[ISO/TS 11139, definition 2.33]

3.14 process challenge location
PCL
site that simulates “worst case” conditions as they are given for sterilizing agent(s) in the goods to be sterilized

3.15 process parameter
specified value for a process variable

NOTE The specification for a sterilization process includes the process parameters and their tolerances.

[ISO/TS 11139, definition 2.34]

3.16 process variable
condition within a sterilization process, changes in which alter microbicidal effectiveness

EXAMPLE Time, temperature, pressure, concentration, humidity, wavelength.

[ISO/TS 11139, definition 2.35]
3.17 reference microorganism
microbial strain obtained from a recognized culture collection

[ISO/TS 11139, definition 2.39]

3.18 resistometer
test equipment designed to create defined reference combinations of the physical and/or chemical variables of a sterilization process

NOTE 1 Adapted from ISO 11138-1, definition 3.15 and ISO 18472:2006, definition 3.11.
NOTE 2 Also referred to as Biological Indicator Evaluator Resistometer (BIER).

3.19 spore-log-reduction
SLR
log of initial spore population, $N_0$, minus the log of the final population, $N_F$

3.20 sterile
free from viable microorganisms

[ISO/TS 11139, definition 2.43]

3.21 sterility assurance level
SAL
probability of a single viable microorganism occurring on an item after sterilization

NOTE 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 $10^{-3}$.

[ISO/TS 11139, definition 2.46]

3.22 sterilization
validated process used to render product free from viable microorganisms

NOTE 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.

[ISO/TS 11139, definition 2.47]

3.23 sterilization cycle development
procedure for determination of the appropriate processing parameters that are consistent with attaining the desired specifications and label claims for a given product or group of products

3.24 sterilization cycle validation
documented procedure for obtaining, recording and interpreting the results required to establish that a process will consistently yield product complying with predetermined specifications

3.25 supplier
organization or person that provides a product

EXAMPLE Producer, distributor, retailer or vendor of a product, or provider of a service or information.
NOTE 1  A supplier can be internal or external to the organization.

NOTE 2  In a contractual situation, a supplier is sometimes called “contractor.”

[ISO 9000, definition 3.3.6]

3.26  survival-kill window
extent of exposure to a sterilization process under defined conditions where there is a transition from all biological indicators showing growth (survival time) to all biological indicators showing no growth (kill time)

[ISO 11138-1, definition 3.18]

3.27  third party
person or body that is recognised as being independent of the parties involved, as concerns the issue in question

NOTE 1  See ISO/IEC Guide 2[1].

NOTE 2  Parties involved are usually supplier (“first party”) and purchaser (“second party”) interests.

3.28  user
person or body employing biological indicators for a given purpose

NOTE 1  See ISO 9000[4].

NOTE 2  The user is the customer who is the recipient of a product provided by the supplier. In a contractual situation, the user is called “purchaser.” The user could be the customer, beneficiary or purchaser. The user can be either external or internal to the organization and represents the “second party.”

3.29  z value
change in exposure temperature of a thermal sterilization process, which corresponds to a tenfold change in $D$ value

NOTE  See ISO 11138-3 and ISO 11138-4.

4  General

4.1  This International Standard provides guidance on biological indicators that can apply generally for any sterilization process, including new sterilization processes not yet covered by International Standards.

4.2  The use of biological indicators is normally documented in procedures and/or instructions.

NOTE  Employing quality management systems such as ISO 13485[7] usually satisfies this provision.

4.3  Biological indicators should always be used in combination with physical and/or chemical measurements in demonstrating the efficacy of a sterilizing process. When a physical and/or chemical variable of a sterilization process is outside its specified limits, the reason for the sterilizer's inability to achieve its process parameters should be evaluated and the problem corrected. Systems and/or procedures should be established to evaluate any deviations from the cycle process limits, and reasons for accepting any deviation should be fully documented.

4.4  A suitable biological indicator consists of carrier material and packaging and has a microbiological component that is known to be suitable for handling without special containment facilities. The growth conditions should be well documented, and the use of the indicator should be as simple and well described as possible to avoid misinterpretation by the user.
4.5 No formal approval system exists, internationally, for biological indicators that are marketed and used for stated purposes or under stated conditions. Some national regulatory authorities, however, have particular requirements for biological indicators and for the choice and use of biological indicators for the validation and control of products marketed as sterile or sterilized.

4.6 A biological indicator represents a microbiological challenge to a sterilization process and is used to verify that a sterilization process has the ability to inactivate microorganisms that have a known resistance to a referenced sterilization process. Test organisms employed in biological indicators typically have resistance to sterilization which exceeds that of common bioburden microorganisms, although some organisms can exhibit a resistance to sterilization in excess of that of the test organisms. The appropriate biological indicator provides a challenge to the sterilization process which exceeds that of the bioburden through a combination of population and resistance. If there is reason to believe that the goods to be processed could be contaminated with particularly resistant organisms, extended sterilization processing, based on the bioburden, could be required.

4.7 Biological indicators are not intended for use in any process other than that specified by the manufacturer on the product labelling. Microbial species and strains are selected as biological indicator candidates based on their known resistance to the specific method of sterilization as certified by the manufacturer. The use of an inappropriate biological indicator can give misleading results. The user should ensure that the biological indicator has been qualified for use with the particular range of sterilization conditions that are used. This could require information in addition to that given in the labelling. When biological indicators are used outside reference conditions, the user can require information on the reaction expected from the indicator, e.g., the effect of sub-optimal moisture conditions on the biological indicators used in an ethylene oxide process. Users who employ biological indicators outside the manufacturer’s labelled recommendations should thoroughly characterize the resistance of the biological indicators to the particular sterilization process. The relationship of the response of the biological indicator to process parameters should be clearly demonstrated.

4.8 It is incumbent upon those responsible for the sterilization of product to ensure that the type of biological indicator employed to validate and/or routinely monitor a given sterilization process is appropriate for that use.

4.9 The manufacturer’s recommendations for the use and storage of the biological indicators should always be followed. Failure to do so can compromise the integrity of the biological indicator. If the user removes the inoculated carrier from the biological indicator’s primary packaging, changes in the resistance characteristics can occur. Guidance should be sought from the manufacturer on the extent of this change, or the user can evaluate changes in the resistance characteristics. The user should document that the performance characteristics of the inoculated carrier are appropriate for their use.

4.10 Biological indicators should not be used beyond the expiration date stated by the manufacturer.

4.11 Those who employ biological indicators for validation and/or routine monitoring of sterilization should be properly trained in their use. The time between completion of the sterilization process and the testing of the BI should be justified as described in 8.2.4. Transferral of microorganisms exposed to the sterilization process to the appropriate recovery medium should be done using aseptic technique.

4.12 The ISO 11138 series gives requirements for the information that the manufacturer should provide for biological indicators. The information may be provided on the label, as a packet insert or as a general specification accompanying the biological indicators. This series of International Standards also includes minimum requirements for resistance characteristics. Testing conditions and methods are given as reference methods.

4.13 Users of biological indicators come from a wide variety of industries, private enterprises and health care facilities. Users are not generally required to perform resistance assays on biological indicators but can have differing requirements for their quality assurance systems, which include audits of vendors and/or manufacturers (see 6.2.2).

4.14 The verification of resistance characteristics by the user is an alternative to and/or complementary to an audit, when necessary.
5 Characteristics of biological indicators

5.1 General

5.1.1 Biological indicators provide means to assess directly the microbial lethality of a sterilization process (see References [8] and [9]). When used in conjunction with physical and/or chemical process monitors, biological indicators can provide an indication of the effectiveness of a given sterilization process.

5.1.2 A sterilization process should be considered as satisfactory only when the desired physical and/or chemical parameters and microbiological results, as determined by an appropriate sterilization cycle development, validation and monitoring programme have been realized. Failure to achieve the desired physical and/or chemical parameters and/or microbiological challenge forms the basis for declaring the sterilization process as nonconforming (see ISO 13485[7] and ISO 9001[22]).

5.1.3 Biological indicators consist of a defined population of test organisms presented in such a manner as to allow their recovery following sterilization processing. For example, test organisms employed for ethylene oxide sterilization processes can be spores of a suitable strain of *Bacillus subtilis* or *Bacillus atrophaeus*, as noted in ISO 11138-2. For steam sterilization or moist heat sterilization, the test organisms employed can be spores of a suitable strain of *Geobacillus stearothermophilus*, as noted in ISO 11138-3. Test organisms other than bacterial spores can be used if they have been shown to provide appropriate resistance to the sterilization process.

5.1.4 The basis of all formulae used to determine biological indicator resistance characteristics such as $D$ values is that the inactivation reaction follows first-order kinetics, with the requirement that the value for the coefficient of determination, $r^2$, for the linearity of the survivor curve be not less than 0.8 (see Annex E and Annex F). The strain, the production method, the suspension fluid, the carrier and packaging materials all affect the resistance characteristics of the finished product (see ISO 11138-1).

5.1.5 The design and construction of a biological indicator can result in unique resistance characteristics and can vary depending on whether the biological indicator is intended for use in the development and validation of a sterilization process or for use in routine monitoring. If the design of the biological indicator for use in routine monitoring differs from that employed to validate the sterilization processes, the challenge to the process during validation should be correlated with the challenge to the process during routine monitoring.

5.1.6 The resistance characteristics of biological indicators vary according to the manufacturing methods and the testing conditions. Depending upon placement within the load and the specific lethal conditions at those discrete locations, biological indicators from the same lot may show different survival capabilities (see 7.2.3). Users of biological indicators should note that ten indicators spread throughout the load are not considered replicates due to the differences in lethality that exist throughout the chamber and load (see Note to 11.3.1).

5.2 Test organism suspension for direct inoculation of products

5.2.1 Direct inoculation of test organisms on or in product can be necessary in cycle development and other studies when the use of a biological indicator is not feasible. Direct inoculation can be appropriate for assessing factors such as product sterilizability, identification of the more difficult to sterilize locations within the device, and localized microbiological effects, e.g., moist heat versus dry heat environments.

The rationale for the selection of the “most difficult to sterilize” site(s) on a medical device or within a sterilization load should be documented based on experimental data or derived from prior knowledge of the particular sterilization methodology. In practice, the “most difficult to sterilize” site represents those locations that are most likely to provide high resistance to the sterilization process. One should refer to specific sterilization standards (e.g., ISO 17665-1 and ISO 11135-1) for guidance in determining and selecting difficult-to-sterilize locations.

5.2.2 To assess the efficacy of sterilization at a particular site or location on the product, the desired species and population of test organisms can be inoculated at those sites. The use of suspensions of test organisms to prepare inoculated carriers or inoculated products requires caution. The materials on to which test organisms are inoculated can alter the test organisms’ resistance characteristics. The resistance can be
higher or lower due to deposition as a monolayer or multilayer, coating effects, and/or bacteriostatic or bactericidal effects of the material. The methods employed to recover the test organisms following processing should be validated to ensure an adequate level of recovery from the product (see ISO 11737-1). Test organism recovery should be expressed in terms of percent recovery of the population of the original inoculum.

5.2.3 Direct inoculation of products or materials with a test organism suspension can cause prolonged or decreased survival of test organisms. This may affect the observed percent recovery of the original inoculum relative to what is expected under specified sterilization conditions. Inoculated products may be assayed with either survivor curve (enumeration/direct counting) or fraction-negative procedures (see Figure A.4). These assays require aseptic techniques.

5.2.4 The $D$ value and, when appropriate, the $z$ value, are constant values only under determined and defined conditions. The resistance characteristics of a spore suspension provided by a manufacturer or supplier of biological indicators might not correspond to the resistance characteristics for direct product inoculation studies. The resistance characteristics should be measured for the carrier employed (solid carrier material or fluid) as well as for the specific sterilization cycle employed.

5.3 Inoculated carriers

5.3.1 Inoculated carriers consist of a defined population of test organisms inoculated on or in a suitable carrier material (see ISO 11138-1:2006, Annex B). Caution should be exercised to ensure that the integrity of the carrier material selected is sufficient to withstand sterilization processing without degradation and to minimize the loss of the inoculated test organisms during transport and handling.

5.3.2 The resistance characteristics of a test organism in suspension can be considerably changed upon deposition on or in carriers. Several factors can influence the resistance characteristics, such as the surface on to which the suspension is inoculated (e.g., solid materials, viscous products or fluids), the way the spores are dispersed and otherwise treated, the methods of drying, etc.

5.3.3 If an inoculated carrier is removed from the biological indicator primary package for cycle development, cycle validation studies, or for process challenge devices used for routine process monitoring, then it is the responsibility of the user to provide a rationale for this application. It should be recognized that the resistance of the microorganism on the inoculated carrier could differ from the labelled resistance of the packaged biological indicator.

5.3.4 The resistance characteristics of an inoculated carrier provided by the manufacturer of biological indicators might not correspond to the resistance characteristics established in direct product inoculation studies.

5.3.5 The carrier material should be evaluated by the biological indicator manufacturer or the user to establish that the sterilizing agent for which the biological indicator is intended neither retains nor releases inhibitory substances (e.g., sterilizing agent residuals) to such an extent that the recovery of low numbers is inhibited (see ISO 11138-1:2006, 5.2).

5.4 Self-contained biological indicators

Self-contained biological indicators consist of either a) or b).

a) An ampoule containing growth medium and a carrier inoculated with test organisms contained within an outer vial so that the sterilizing agent obtains access to the inoculated carrier through a sterile barrier or a tortuous path.

After exposure to the sterilization process, the growth medium is brought into contact with the inoculated carrier by breaking the ampoule of growth medium, thereby eliminating the need to aseptically transfer the inoculated carrier to a separate vial of growth medium. The biological indicator manufacturers’ recommendations should be followed for incubation of self-contained biological indicators.

NOTE Due to the low volume and the possibility of evaporation of the growth medium, prolonged post-exposure incubation might not be possible.
Chemical residuals resulting from processes such as ethylene oxide or vapour hydrogen peroxide can inhibit growth of surviving organisms. The biological indicator manufacturer’s recommendations should be followed for proper handling (including aeration) of biological indicators prior to incubation (see 8.2.4).

b) A hermetically sealed ampoule containing a suspension of test organisms in growth medium.

These are referred to as sealed-ampoule biological indicators. After exposure to the process, the sealed ampoule is incubated intact, and no aseptic transfer is required.

NOTE This type of indicator is sensitive only to exposure time and temperature and is primarily used to monitor moist heat sterilization of aqueous fluids.

Self-contained biological indicators are generally larger than biological indicators that consist only of an inoculated carrier in a primary packaging, and may not fit into locations within the device that represent the process challenge locations. Unless a biological indicator can be placed into a load without deforming it or otherwise potentially compromising its primary packaging, the biological indicator should not be used. Also, the user should be aware that the claimed resistance characteristics can be dependent on the air-removal method employed in the sterilization cycle.

5.5 Other biological indicators

Biological indicators in their simplest form consist of an inoculated carrier in primary packaging. The inoculated carrier can take a variety of forms, including paper strips, threads, metal coupons or other carriers suitable for inoculation. The primary packaging is chosen to permit the sterilizing agent to penetrate to the inoculated carrier while maintaining a sterile barrier after processing.

6 Selection of supplier

6.1 General

6.1.1 The user of biological indicators should, whenever possible, purchase to standard specifications, e.g., biological indicators manufactured according to specifications given in the ISO 11138 series, pharmacopoeial monographs or other applicable standards. The user should consider the particular sterilization process as the basis for the choice of biological indicator.

6.1.2 When the user has a process that requires performance characteristics that differ from the label claim for the biological indicator, it is the responsibility of the user to verify that the biological indicator has the performance characteristics needed.

6.1.3 The user of biological indicators should have a system in place to provide assurance that the biological indicators obtained consistently meet the specified characteristics. Such assurance may be provided by one or more of the following:

a) information from the manufacturer covering the performance characteristics of the lot of biological indicators prepared;

NOTE Requirements for information supplied by manufacturers of biological indicators are provided by the ISO 11138 series.

b) a statement of conformity from the manufacturer that the biological indicators meet the agreed specifications;

c) if needed, various degrees of testing of each lot of biological indicators received by the user, to verify that the performance characteristics meet the agreed specifications.

6.1.4 When the user has established a high level of confidence in the supplier (see 6.1.3), the testing performed by the user can be minimal. At a minimum, the user should have a mechanism to ensure that a shipment of biological indicators contains all agreed-upon documentation, such as appropriate label