Biological Indicators and Validation

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Agenda

Biological qualification concepts and normative requirements

Biological indicators description and practical case studies









Introduction to Steam Sterilization & Validation

All sterilization processes should be validated. Particular attention should be given when the adopted sterilization method is not described in the current edition of the European Pharmacopoeia, or when it is used for a product which is not a simple aqueous solution.

Where possible, heat sterilization is the method of choice.

[Annex 1 EU GMP, Clause 8.35/8.36]

"To validate a sterilization process, **physical conditions** are chosen that are expected to sterilise the items in the load to achieve a **Sterility Assurance Level (SAL) equal to or less than 10**-6".

[Eu Ph. 11.0 - Chapter 5.1.2]

The objective of a validation study is to demonstrate that the sterilization effectiveness anticipated from the physical process parameters is equivalent to the biological sterilization effectiveness.

[Eu Ph. 11.0 - Chapter 5.1.2]





Physical VS microbiological results

Validation of sterilization processes links **physical measurements** with **biological indicator performance** to establish method lethality.

[USP 43, Chapter 1229]

Consistency between **physical** and **microbiological** result is central to **sterilization validation**.

Likewise, the destruction of a BI without consideration of the physical parameters needed to kill the BI does not provide sufficient evidence of the **suitability** and **repeatability** of the cycle.







Spores don't lie and probes don't always know (or tell) the truth

The **probes** in your sterilization equipment are state-of-the-art, accurately calibrated, highly sensitive instruments and without them, controlling one's cycle would not be possible. But each one measures a limited environment and is only capable of sensing and measuring one parameter in a cycle.

Only the **Biological Indicator spores** can **accurately monitor the delivery of lethality** at various locations throughout the load.

[MesaLabs Spore News Vol 2 Issue 4 " spores don't lie and probes don't always know (or tell) the truth]





Biological Indicators for the sterilization process

In addition to the physical sterilization parameter **the effectiveness** of a sterilization process is dependent of a large numbers of variables, which may include, but is not necessarily restricted to:

- 1. The number and the resistance of the contaminant microorganisms;
- 2. Penetration of the sterilant;
- 3. Time;
- 4. Temperature;





Biological Indicators for the sterilization process

In addition to the physical sterilization parameter **the effectiveness** of a sterilization process is dependent of a large numbers of variables, which may include, but is not necessarily restricted to:

- 5. Concentration;
- 6. pH;
- 7. Moisture content;
- 8. Chemical composition of the product or item.

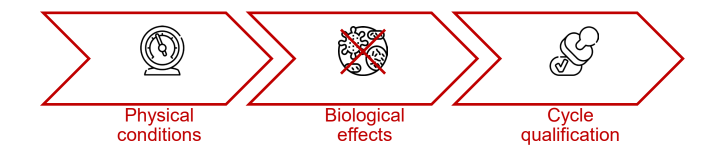






Biological qualification

The objective of the biological component of cycle qualification is to obtain biological data confirming that the developed cycle achieves the actual biological lethality requirements established during cycle design.









Biological qualification



Biological qualification using microbiological challenges follows a straightforward sequence:

- An appropriate microbial challenge system is devised based on the desired lethality (F-value) determined during the design of the process;
- The load is exposed to minimal or subminimal **sterilization conditions**;
- After completion of the cycle, the microbiological challenges are retrieved;





Biological qualification



Biological qualification using microbiological challenges follows a straightforward sequence:

- Each microbiological challenge is **individually incubated** in appropriate media and conditions for growth of survivors;
- The results are evaluated to ensure that the **spore log reductions** achieved for the microbiological challenges **meet predetermined acceptance criteria**;
- Growth of the microbiological challenge organism is required in positive controls.





Biological indicators at glance

What is a Biological Indicator?

Test systems that provide a defined challenge to verify required effectiveness of a specified sterilisation process.



Which type of microorganisms?

Microorganisms widely recognized as suitable for biological indicators are spore-forming bacteria because more resistant than normal microflora.

[USP 43, Chapter 1229.5]

Why to use Biological Indicators?

Bls are used to demonstrate the effectiveness of processes that render a product sterile.

CONNECTING PEOPLE SCIENCE M REGULATION® [USP 43, Chapter 1229.5]

IEu Ph. 11.0 5.1.2.1







Biological indicators are designed to show by the survival of test microorganisms whether specified sterilization conditions have been attained.

The **absence of growth** of a test microorganism after exposure to a sterilization process demonstrates that a specified level of microbiological **inactivation** has been delivered.





Survival of a test microorganism subjected to a sterilization process indicates that the process has **failed**.





Biological Indicators selection

The **selection of the types of biological indicator** used will depend on:

- The **nature of the sterilizing agent** (*Moist heat G. stearothermophilus, Dry Heat B. atrophaeus, Ethylene Oxide B. atrophaeus, H2O2 G. stearothermophilus*);
- The expected effectiveness of the treatment (e.g. the F_{Phys} calculated from the process parameters);
- The characteristics of the pharmaceutical product or item to be sterilized. (*e.g. final container, packaging material, utilities such us tubes or pumps*).

[Eu Ph. 11.0, Chapter 5.1.2]





Biological Indicators selection

The selection of a BI requires knowledge of the **resistance of the BI** system to the specific sterilization process.

It must be established that the **BI system provides a challenge** to the sterilization process greater than the resistance of the native bioburden.

[USP 43, Chapter 1229.5]





Use and Placement of **Biological Indicators**

For cycles designed using the overkill design approach, the challenge system is typically spores of G. stearothermophilus.

Lower levels of thermal input may be delivered for the product**specific design approach**. Consequently, the challenge organism used in qualification is often less resistant than spores of G. stearothermophilus.

Typical microorganisms used for qualification of cycles using the product-specific design approach include: *Clostridium sporogenes*, Bacillus smithii (formerly Bacillus coagulans) and Bacillus subtilis 5230. [PDA TR No.1 par. 3.2.1]





Biological indicators: Microorganisms & Regulatory References

		ISO 11138-3	Eu. Ph. XI ed.	USP 43
Strain		G. stearothermophilus	G. stearothermophilus (ATCC 7953, 12980, NCTC 10007, CIP 52.81, NCIMB 8157)	G. stearothermophilus (ATCC 12980, ATCC 7953)
				C. sporogenes (ATCC 7955)
				B. atropheus (ATCC 9372)
				B. subtilis (ATCC 5230)
Population		≥1,0*10 ⁵		
D value at 121°C		≥1,5 min	1,5 min to 4,5 min	
z value	Ŀ	≥6°C		







Biological Indicators: Types



Spore suspension





Inoculated carriers



Custom-made







Inoculated carriers

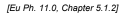


Consist of a **defined population** of bacterial spores inoculated into or onto a **suitable carrier** and in most cases in a protective envelope.

The **type of carrier** (and the envelope if used) may **influence the resistance** of the bacterial spores and must be compliant with the chosen sterilization process.

- No chemical/microbial contamination;
- No degraded by the sterilization process;
- They should minimize the loss of the original inoculum during transport, handling and shelf-life storage;
- Must not retain residual sterilizing agent such that it could hinder outgrowth of low numbers of surviving spores.

After the exposure the carrier is aseptically handled according to the manufacture's instruction, transferred to a suitable culture medium and incubated for a sufficient period of time at appropriate temperature.

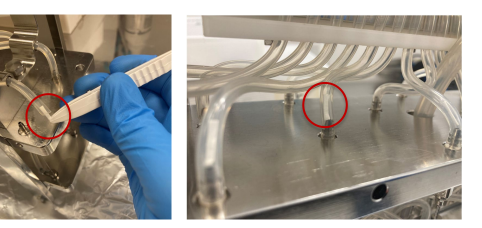






Load: Dosing system BIs: Inoculated carriers Cycle: Steam sterilization by direct contact









Load: Syringes Bls: Inoculated carriers Cycle: Air over steam sterilization













Load: Stoppers in double bags Bls: Inoculated carriers Cycle: Steam sterilization by direct contact







Self-container BIs

System consisting of an **inoculated carrier and a container** (e.g. ampoule) with a nutrient medium suitable for the test micro-organism used.

The sterilizing agent comes in contact with the inoculated carrier while the growth promoting properties of the nutrient medium are not adversely affected by the sterilization process. After the exposure the carrier is brought into contact with the nutrient medium **by simple manipulation**.

[Eu Ph. 11.0, Chapter 5.1.2]



TIPS: Using an SCBI provides an alternative option for customers who do not have the capability to perform aseptic culturing as the system is 'self-contained' and this eliminates the chance of post-process contamination.

[MesaLabs spore news 'biological indicator selection' vol 12 issue 5]





Load: Metal items in Tyvek bag Bls: Self contained Cycle: Steam sterilization by direct contact









Self-container BIs



Container (e.g. ampoule) of a population of the test microorganism **in an appropriate nutrient medium.**

After the exposure the container is incubated without any further manipulation.

This type of biological indicator is sensitive only to an exposure time and temperature and may be used primarily to monitor **sterilization of aqueous fluids**.

[Eu Ph. 11.0, Chapter 5.1.2]





Practical example Load: Liquid loads



BIs: Self contained ampoules **Cycle**: Steam sterilization by direct contact









Load: 250 ml plastic bottles (PP) filled with water Bls: Self contained ampoules Cycle: Super-heated water sterilization









Tech Tips

In cycle validation, the relevant **positions** in the load that are the most difficult to sterilise are determined and adequate **biological effectiveness is verified** by biological indicators in these positions or products, whichever is relevant.

[Eu Ph. 11.0, Chapter 5.1.1]

Monitor the process leaving biological indicators in the same position considered for thermal qualification: worst case locations and cold spots should be monitored.

The identification of the worst-case position of each standard load is part of the validation exercise of the sterilization process.







Spore suspension



Consist of a **defined population of bacterial spores**, prepared from clearly characterized ad suitable maintained strain of a spore forming bacterial species in a **stable suspension**.

[Eu Ph. 11.0, Chapter 5.1.2]







Custom made BIs

Test items (e.g. rubber stoppers), or products, inoculated with a suitable test micro-organism, usually from a characterized spore suspension.



The spore suspensions can be prepared from isolated environmental monitoring or other microbiological testing using a well-defined procedure designed to give satisfactory sporulation. The D- value and, when appropriate the z-value of the spore suspension must be determined.

The D-value and the z-value (if appropriate) of the spores of the inoculated items/products must be determined as this may be different from the spores in suspension.

After the exposure to the sterilization cycle, the custom-made biological indicator is enumerated or tested for the presence/absence of surviving test-microorganism using a validated, appropriate microbiological technique.



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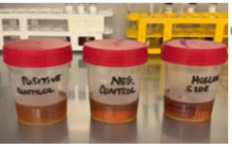
Load: Stoppers Bls: Spore suspension Cycle: Steam sterilization by direct contact



Steam sterilization











Spores do not have an intrinsic D-value

A major misunderstanding in the industry is that spores have an intrinsic D-value. This is not true if for no other reason than it is impossible to test the resistance of individual spores suspended in space. The spores are placed onto a surface or suspended in liquid and this system becomes the biological indicator (BI).

By definition, the D-value for a BI is the time (or dose) required at a specified set of exposure conditions to reduce the viable spore population by one log or 90%.

Don't forget that the D-value is a measure of resistance performance that refers to the entire BI package and is not a value for the spore itself.



[MesaLabs Spore News, Vol. 5 N. 1 - January 2008]

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Spores do not have an intrinsic D-value

Solid Surfaces

Not all surfaces are equal. **Spores will interact differently with a surface** depending on many factors:

- Surface topography
- Surface hydrophilicity
- Different elastomers type
- Etc...

[MesaLabs Spore News, Vol. 5 N. 1- January 2008]



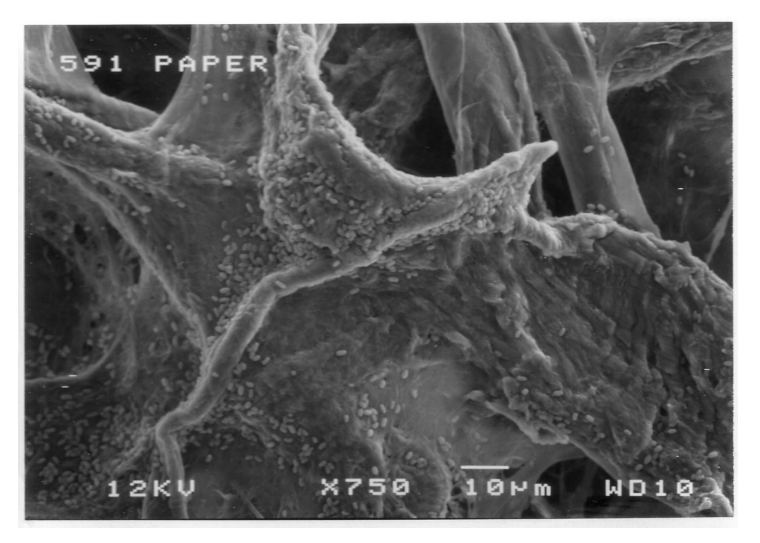


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PAPER









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[MesaLabs Spore News, Vol. 5 N. 1- January 2008]



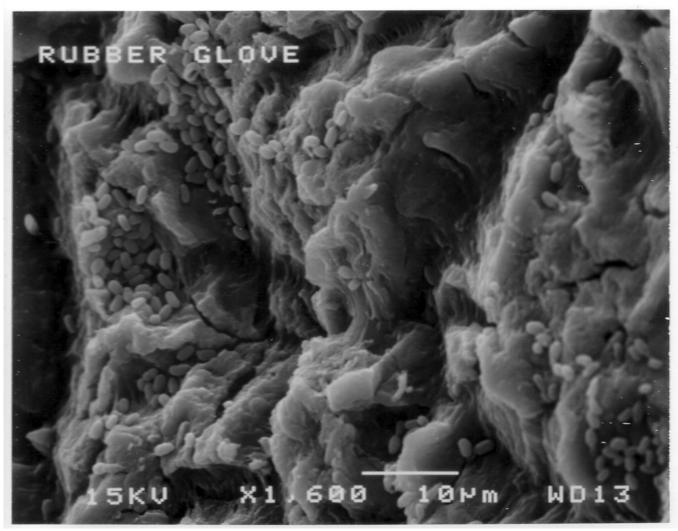


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RUBBER GLOVE









Spores do not have an intrinsic D-value

Solid Surfaces

Not all surfaces are equal. **Spores will interact differently with a surface** depending on many factors:

- Surface topography
- Surface hydrophilicity
- Different elastomers type
- Etc...

[MesaLabs Spore News, Vol. 5 N. 1- January 2008]





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STAINLESS STEEL









Biological indicators: Responsibility

Manufacturer



Determining the performance characteristics of each BI lot;



Provide information concerning the **microbial population and resistance** (D and z values, respectively, where appropriate);



The **resistance** of the BI should be determined by the manufacturer under defined conditions;

[USP 43, Chapter 1229.5]





Biological indicators: Responsibility

Manufacturer



Storage and **expiry** information;

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Should provide **directions for use**, including the medium and conditions used for the recovery of microorganisms after exposure to the sterilization process;

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Disposal instructions also should be provided by the manufacturer of the BI;



The manufacturer should provide, with each lot of BIs a **certificate of analysis** that attests to the validity of BI performance claims.







) Packaging and storage

Store under the conditions recommended on the label or under validated conditions, and protect from light, toxic substances, excessive heat, and moisture;



Expiration date

Use within the BI's labeled or determined expiration date;

Spore Suspension CERTIFICATE OF ANALYSIS

SSSE/6 Reorder No.: 7953(1) Geobacillus stearothermophilus Volume: 10 mL (40% Ethanol Suspension). In freezer (between -25° and -10°C). Storage: Incinerate or autoclave at 121°C for not less than 30 minutes Disnosal: No evidence of contaminants using standard plate count techniques Purity: Lot No. : SSS-753 Manufacture Date: 2017-11-13 Expiration: 2019-11-13 Heat Shocked Population: 1.8 x 10⁶ Spores / 0.1 mL Assayed Resistance: D-Value⁽²⁾ Steam 121°C 2.3 Z-value: 8.8°C Units are manufactured in compliance with Mesa Laboratories, Bozeman Manufacturing Facility's quality standards USP, and ISO 11138 guidelines and all appropriate subsections.

¹⁰Cahure is traceable to a recognized enhure collection identified in USP and ISO 11138.
¹⁰Resistance was determined in an AAM BER vesced using a paper carrier backaged in galaxies and exhaulted using the Fraction Negative method. The D-value is reproducible only when exposed and cultured under the exact conditions used to obtain results reported lore.

[USP 43 , Chapter 1229.5]







Identification

Where identification of the BI species is deemed necessary, as in the course of an investigation into unusual results, use either a phenotypic or genotypic identification method;

Spore Suspension Certificate of Analysis

SSSE/6 Reorder No.: Geobacillus stearothermophilus 7953(1) 10 mL (40% Ethanol Suspension). Volume: In freezer (between -25° and -10°C). Storage: Incinerate or autoclave at 121°C for not less than 30 minutes. Disposal: No evidence of contaminants using standard plate count techniques Purity: Lot No. : SSS-753 Manufacture Date: 2017-11-13 Expiration: 2019-11-13 Heat Shocked Population: 1.8 x 10⁶ Spores / 0.1 mL Assayed Resistance: D-Valuc(2) 2.3 Steam 121°C Z-value: 8.8°C

Units are manufactured in compliance with Mesa Laboratories, Bozeman Manufacturing Facility's quality standards USP, and ISO 11138 guidelines and all appropriate subsections.

mC_charms is strengther to recognized exhance collection identified in USP and ISO 11133.
W Restances was the determined in an AAAB URBE vessed ioning a payre carrier protacipation in glustance and exhaulted using the Fraction Negative method. The D-value is reproducible only when exposed and cultured under the exact conditions used to obtain retails reported hore.

[USP 43, Chapter 1229.5]





Purity

By examination of the colonies derived from the spores on a suitable plate culture medium, determine that there is no evidence of contamination with other microorganisms;

Disposal

Prior to discarding used spores, sterilize using a method recommended by the BI manufacturer or other equivalent means.

Spore Suspension

CERTIFICATE OF AUGINE				
Reorder No.: SSSE/6				
Geobacillus stearothermophilus 7953 ⁽¹⁾				
Volume: 10 mL (40% Ethanol Suspension).				
Storage: In freezer (between -25° and -10°C).				
Disposal: Incinerate or autoclave at 121°C for not less than 30 minutes.				
Purity: No evidence of contaminants using standard plate count techniques.				
Lot No. : SSS-753 Manufacture Date: 2017-11-13				
Expiration: 2019-11-13				
Heat Shocked Population: 1.8 x 106 Spores / 0.1 mL				
Assayed Resistance:				
D-Valuc ⁽²⁾				
Steam 121°C 2.3 min				
Z-value: 8.8°C				

Units are manufactured in compliance with Mesa Laboratories, Bozeman Manufacturing Facility's quality standards, USP, and ISO 11138 guidelines and all appropriate subsections.

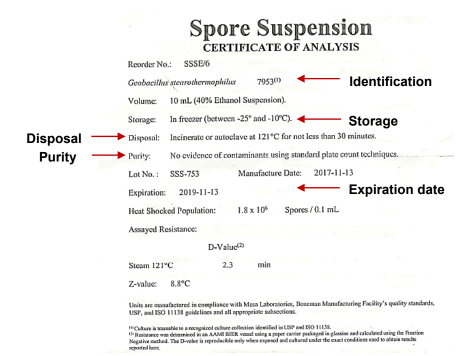
¹¹ Culture is tractable to a recognized onlune collection identified in USP and ISO 11138.
¹² Resistance was determined in an AAMI BIER vessel using a paper curiter packaged in glazzine and eakolated using the Franciso Negative neution. The D-value is reproducible only when exposed and cultured usder the easet conditions used to obtain readly reported by:

[USP 43, Chapter 1229.5]















Biological indicators: Responsibility

User



When BIs are purchased, their **suitability for use** in a specific sterilization process must be established.



The BI user should obtain a **certificate of analysis** for each lot of BIs and verify the manufacturer's label claims for spore population



When BIs are **used in accordance** with the BI manufacturer's directions, the **resistance of the BI need not be reconfirmed**.

[USP 43, Chapter 1229.5]







Biological indicators: Responsibility



User prepared biological indicators

A user of BIs may elect to propagate spore crops of a single species for use as a suspension. Alternatively, these spore suspensions may be purchased from a BI manufacturer. When liquid suspensions are applied to a substrate, it is the user's responsibility to determine the population and resistance of the microorganism used. The resistance determined for liquid suspensions relates only to other lots of the same suspension and is not representative of how that microorganism will perform on a substrate or in a different suspending medium. In these circumstances, the BI resistance and population should be reestablished.

[USP 43, Chapter 1229.5]





Tech Tips



The carrier and primary packaging **should not be damaged** or degraded by the specific sterilization process;



The **time** between completion of sterilization process and incubation should be within the manufacturer's stated time or should be justified;

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Transfer of microorganisms exposed to the sterilization process to the appropriate recovery medium should be done using aseptic technique.





Performance Qualification

PQ is the most demanding part of validation exercise, because it refers to **actual loads** and **operating conditions**. Basically, it includes:

Physical qualification → by **measuring physical parameters**;

Biological qualification \rightarrow by measuring the **actual kill of microorganism in a biological indicator.**

Both **physical and biological qualification** are concerned with definition of number of TE, TC, BIs and their placement in test with standard loads.







PQ on an actual load

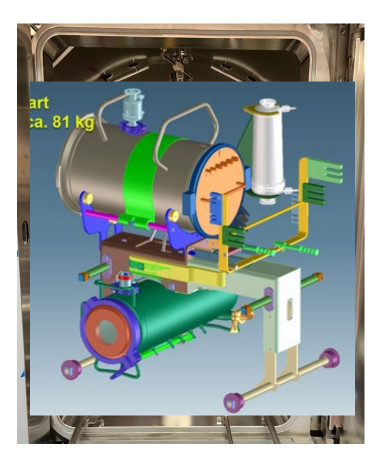
Dosing canister for vials filling process



Use of Biological Indicators



Use of Thermocouples









Biological Indicators used



Inoculated carrier with envelope

Hanged on the surfaces of the trolley and the chamber and inserted in the pipes with larger diameter





Inoculated carrier without envelope Inserted in the pipes with small diameter







Aim of the PQ

Physical qualification

Temperature monitoring in different points of the load and the chamber.

Biological qualification

Measure of the killing effect on the BIs – investigation in the same locations of the TC probes.

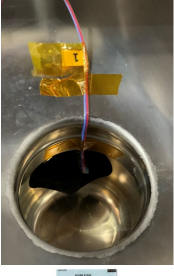
Dryness and Integrity of the load

To match the chosen process with the efficiency in terms of thermal and biological overcomes.





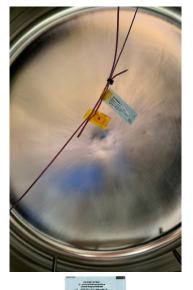
TC and BI 1 Drain of the autoclave



APORT ATTAIN Construction of the second seco TC and BI 2 and 3 TE4 and TE2 plant probe



TC and BI 4 In the centre of the autoclave chamber



TC and BI 5 Pall filter – Internal part of the cartridge







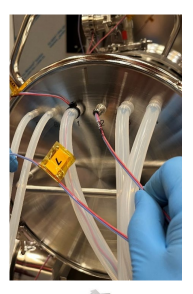




TC and BI 6 Inside plastic pipe



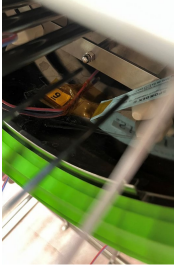
TC and BI 7 Inside needle No.2



TC and BI 8 Inside disposable filter -External part of the cartridge



Arque arter Arque arter Chantraperfision Lar obbets Exp persons arter arte TC and BI 9 Inside upper vessel (at the bottom)

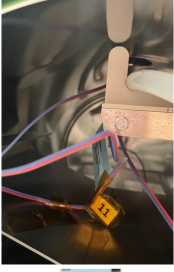








TC and BI 11 Inside upper vessel (at the top)





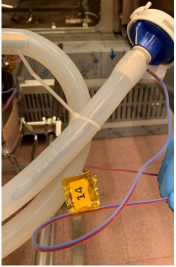
TC and BI 12 Inside Y connection



TC and BI 13 Inside plastic pipe with sterile connection (Y pipe)



TC and BI 14 Inside plastic pipe with sterile connection (Y pipe)









TC and BI 15 Pall filter – External part of the cartridge





TC and BI 17 Inside disposable filter – internal part of the cartridge



APORE STRIP G. starrothermophiles ChemVaporitiesm Lat 035-011 Exp 2011-06-13





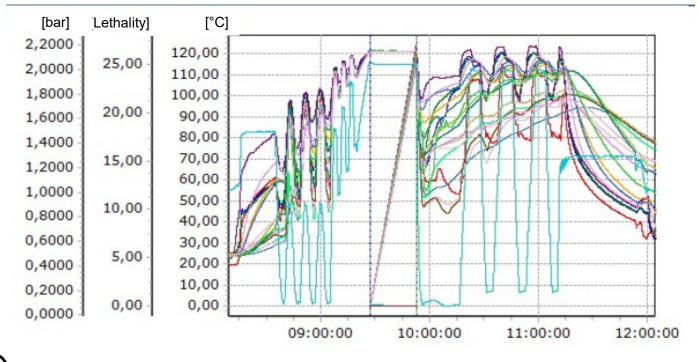
TC and BI 20 Inside plastic pipe







Temperature mapping



Good thermometric results: 6 seconds of equilibration time → every TC was within the sterilization range (120°C – 122°C).







Outcomes after BIs incubation





No Growth in any of the investigated positions







Outcomes after BIs incubation





No Growth in any of the investigated positions







Aim of the PQ VS Final results

Physical qualification

Sterilization temperature reached in every investigated point.

Biological qualification

No growth in any investigated position.

Dryness and Integrity of the load

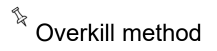
Dry load, filters not damaged.



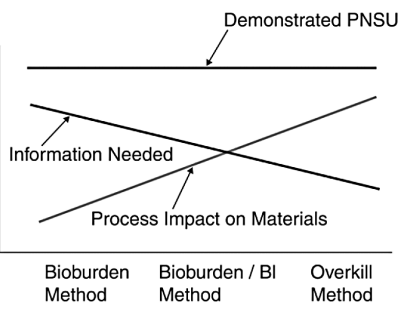
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Validation Methodologies at glance



- ^b Bioburden/BI method
- ${}^{\mbox{\sc b}}$ Bioburden based method



[USP 43, Chapter 1229]



The different approaches were developed because of the differences in the heat resistance of the materials to be sterilized.

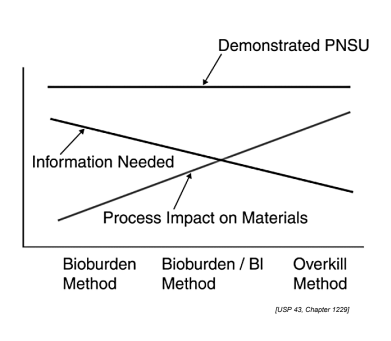




Overkill approach

Overkill sterilization is a method in which destruction of high the а concentration of resistant а microorganism is correlated with the destruction of reasonably anticipated bioburden during routine present processing.

When the load items can withstand substantial **heat without adverse consequence**, overkill is the **method of choice** for steam sterilization because of its ease of execution, reduced considerations for bioburden control, and overall simplicity.







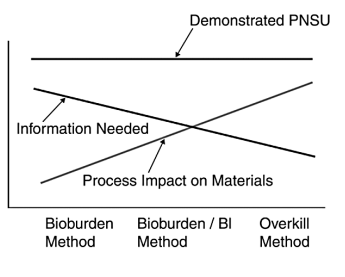
Overkill approach

Microbiological target: PNSU of ≤10⁻⁶;

Product: items that are unaffected by the process exposure (inert items);

Bioburden knowledge: some bioburden knowledge, not so extensive as bioburden data required for bioburden process or BI/BB process;

Use of BIs: G. stearothermophilus (ATCC 12980 or ATCC 7953).



[USP 43, Chapter 1229]

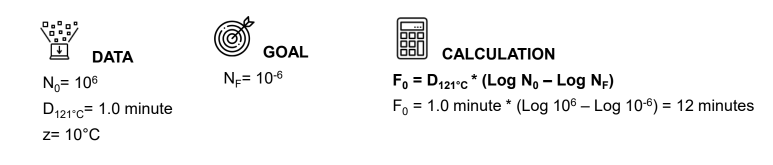




Overkill design approach

"...a cycle design with the overkill design approach can be defined as a sterilization cycle that is demonstrated to deliver an F_{PHY} and F_{BIO} of at least 12 minutes to the items being sterilize"

[PDA TR No 1, Paragraph 4.1.1.1]



[Overkill design approach, PDA TR No 1]

"The objective of the overkill design approach is to assure a level of sterility assurance **regardless of the number and heat resistance of the actual bioburden in the load**."

[PDA TR No 1 rev. 2007, Clause 4.1.1.1]

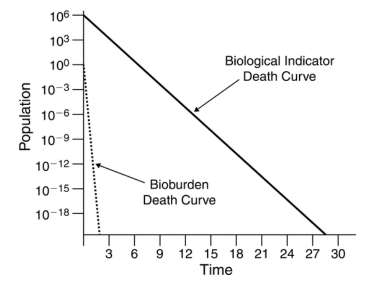




Product specific approach: BB/BI

Bioburden/Biological Indicator based sterilization is an approach in which the incomplete destruction (or destruction of a modest population) of a resistant biological indicator can be used to demonstrate the capability of the method to reliably destroy the bioburden present.

[UPS 43, Chapter 1229]



[Relative resistance and population of typical biological indicator and bioburden microorganisms. USP 43, Chapter 1229- Figure 4]



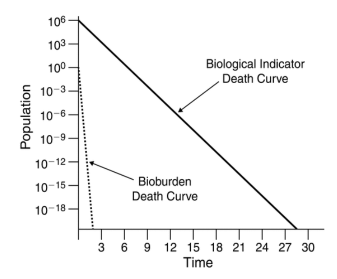
Product specific approach: BB/BI

Microbiological target: PNSU of $\leq 10^{-6}$;

Product: heat-sensitive one;

Bioburden knowledge: it requires detailed knowledge of the bioburden and biological indicator populations and their relative resistance. The relative resistance of the selected biological indicator to that of the bioburden must be established on or in the product;

Use of Bls: The conventional Bls are Clostridium sporogenes ATCC 7955 and Bacillus subtilis ATCC 5230, although other strains can be used.



[[]Relative resistance and population of typical biological indicator and bioburden microorganisms. USP 43, Chapter 1229- Figure 4]

[USP 43, Chapter 1229.2]





Product specific approach: Bioburden

BB method is similar to the BB/BI method. The difference lies in the isolation and characterization of the most resistant bioburden microorganism.

[USP 43, Chapter 1229.2]

The bioburden-based method is used when **material stability limited** or when there are **no suitable biological indicator microorganisms available** to use with the sterilizing process.





Product specific approach: Bioburden

Microbiological target: PNSU of $\leq 10^{-6}$ for the bioburden;

Product: heat-sensitive one;

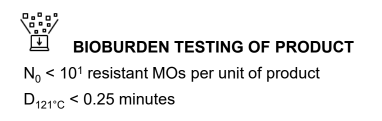
Bioburden knowledge: requires extensive knowledge of product bioburden, **routine monitoring** of the bioburden population and its resistance to the sterilization process is mandatory. The bioburdenbased method requires the user to develop suitable critical control points within the process to control the bioburden titer;

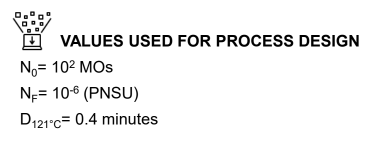
Use of BIs: none.





Product design approach: Example (I)





CALCULATED MINIMUM LETHALITY TO ACHIEVE A PNSU OF LESS THAN 10⁻⁶

 $F_{121^{\circ}C} = (Log N_0 - Log N_F) * D_T = (Log 10^2 - Log 10^{-6}) * 0.4 min = 3.2 min$

[Product- specific design approach , PDA TR No 1]



Since the design value for resistance (0.4 minutes) is only **slightly higher** than the heat resistance of the Mos present in the product, it is necessary to **conduct continuous monitoring of the bacterial load population** to ensure that no drift occurs in the product size of the population or resistance over time.





Product design approach: Example (II)



 $N_0 < 10^1$ resistant MOs per unit of product $D_{121^\circ C} < 0.25$ minutes

VALUES USED FOR PROCESS DESIGN N_0 = 10² MOs N_F = 10⁻⁶ (PNSU) $D_{121^{\circ}C}$ = 1 minutes

$ec{I}$ CALCULATED MINIMUM LETHALITY TO ACHIEVE A PNSU OF LESS THAN 10-6

 $F_{121^{\circ}C} = (Log N_0 - Log N_F) * D_T = (Log 10^2 - Log 10^{-6}) * 1 min = 8 min$

[Product- specific design approach , PDA TR No 1]



Since the design value selected far heat resistance is very conservative (1.0 minute), the need for ongoing product bioburden heat resistance testing is significantly reduced, but should still be monitored periodically.





Steam Sterilization & Validation Methodologies



Terminal sterilization processes require greater consideration of the **effects** of the treatment on **material properties**.



The preferred method for steam sterilization is the **overkill method**.



When the processed materials are susceptible to **damage** by moist heat at the overkill conditions, the **BB/BI** method is better suited because it results in reduced heat input while affording the same degree of process efficacy but with different controls.







European Medicines Agency – Guideline on the sterilization of the medicinal product, active substance, excipient and primary container.

Cycle	Type of process	Information in dossier*	Bioburden level before steam sterilisation or terminal heat treatment	Bioburden Characterised	Process hold temperature
Ph. Eur. 5.1.1 Reference Cycle	Sterilisation	1, 6	100 CFU/100ml (non-routine)	No	≥ 121 °C for ≥15 minutes
Overkill cycle F₀ >12 min	Sterilisation	1, 2, 3, 4, 7	100 CFU/100ml (non-routine)	No	≥ 121 °C
F₀ > 8 min	Sterilisation	1, 2, 3, 4, 7	100 CFU/100ml (routine)	No	> 115 °C
F₀ > 8 min	Sterilisation	1, 2, 3, 5, 7, 8	100 CFU/100ml (routine)	Yes**	> 115 °C
F₀ > 8 min	Sterilisation	1, 2, 3, 4, 7	100 CFU/100ml (routine)	Yes	> 110 °C
F₀ > 8 min	Sterilisation	1, 2, 3, 5, 7, 8	100 CFU/100ml (routine)	Yes**	> 110 °C
F₀ <8 min	Post-aseptic processing terminal heat treatment	1, 2, 3, 4, 7, 8	0 CFU/100ml, aseptic filtration and processing prior to terminal heat treatment (routine)	Yes***	> 110 °C****
F₀ <8 min	Post-aseptic processing terminal heat treatment	1 2, 3, 5, 7, 8	0 CFU/100ml, aseptic filtration and processing prior to terminal heat treatment (routine)	Yes***	> 110 °C****

* For clarification of the code numbers, see below

** In-process control demonstrating acceptable heat resistance of bioburden

*** The bioburden prior to the sterilisation step (i.e. filtration) should be characterised for heat resistance

**** Temperatures below 110 °C may be used if justified. The requirement for additional documentation for such cycles is evaluated on a case by case basis

Clarification of the information to be presented in the quality dossier

1: Sterilisation time, temperature profile

2: Sterilisation method (for instance saturated steam cycle, air/steam-overpressure cycle, vacuum phase) description including SAL

3: Validation of F_{0Phys} and F_{0Bio}

4: Biological indicator with a D₁₂₁ ≥ 1.5 minutes used in the validation

5: Biological indicator with a D₁₂₁ < 1.5 minutes used in the validation

6: No validation data requested in the dossier, only a confirmation that validation has been performed

7: Validation data to be provided in the dossier is presented below

8: Additional validation data to be provided in the dossier is presented below

Table 1: Cycles for steam sterilisation and post-aseptic processing terminal heat treatment and corresponding data required in the quality dossier



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Biological indicators for moist heat sterilisation in Ph. Eur. 5.1.2, 3-2-1

- **1. MOs**: *G. stearothermophilus* is the most widely accepted biological indicator micro-organism for moist heat sterilisation processes;
- **2. Heat resistance**: Reported $D_{121^{\circ}C}$ -values for its spores are in the range of 1.5 min to about 4.5 min;

- EMA guideline refers to these concepts for the selection of suitable BIs;

- EN ISO Standard 11138-3:2017, "Biological indicators for moist heat sterilization processes", prescribes as test organism "G. stearothermophilus or other strains of microorganisms of demonstrated equivalent performance". To comply with this standard the minimum $D_{121 \, ^{\circ}C}$ -value shall be 1.5 min and the minimum viable count 1 x 10⁵.





Biological indicators for moist heat sterilisation in Ph. Eur. 5.1.2, 3-2-1

- **3. Population**: It is recognised that a 10^5 or 10^6 population of G. stearothermophilus may not be suitable for sterilisation processes delivering an F₀ between 8 and 15, therefore a lower spore number (i.e. 10^3 or 10^4) or a different test micro-organism may be used;
- **4. Choice**: The charactertistics of the sterilisartion process [T, t, SAL, F₀] are the basis for the choice of the biological indicator [Type, M.O., Population].

⁻ EN ISO Standard 11138-3:2017, "Biological indicators for moist heat sterilization processes", prescribes as test organism "G. stearothermophilus or other strains of microorganisms of demonstrated equivalent performance". To comply with this standard the minimum $D_{121} \circ_{\rm C}$ -value shall be 1.5 min and the minimum viable count 1 x 10⁵.



⁻ EMA guideline refers to these concepts for the selection of suitable BIs;



Conclusions

Yelidation of sterilization processes requires knowledge of sterilization technology and use of the appropriate instrumentation and equipment to control and verify critical sterilization process parameters.



S Validation of sterilization processes links **physical measurements** with **biological indicator** performance to establish method lethality.



Product-oriented validation "Terminal sterilization processes require greater consideration of the effects of the treatment on material properties".



Validation methods: overkill, BB/BI and BB.



Validation lifecycle should be documented.





Thank you!



