## Understanding Sterilization

- Sterilization basics
- Radiation Technology & Ethylene Oxide



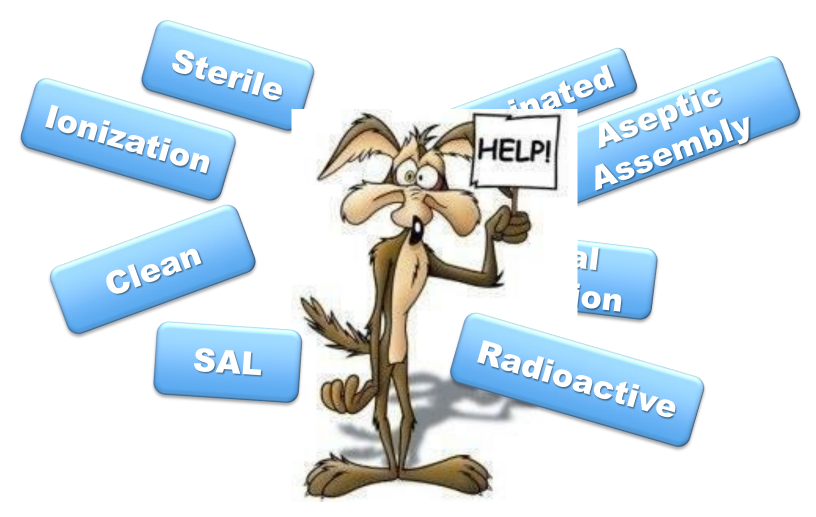
ANNICK GILLET













#### Content



#### Basics of sterilization

- Distinguish disinfection, sterilization and decontamination
- Definition
- Selection of sterilization method
- Difference between Aseptic Assembly and Terminal Sterilization
- Sterilization using Irradiation
  - o **Gamma**
  - o E-Beam
- Sterilization by Ethylene Oxide
- Comparison between technologies





## **Sterilization Basics**

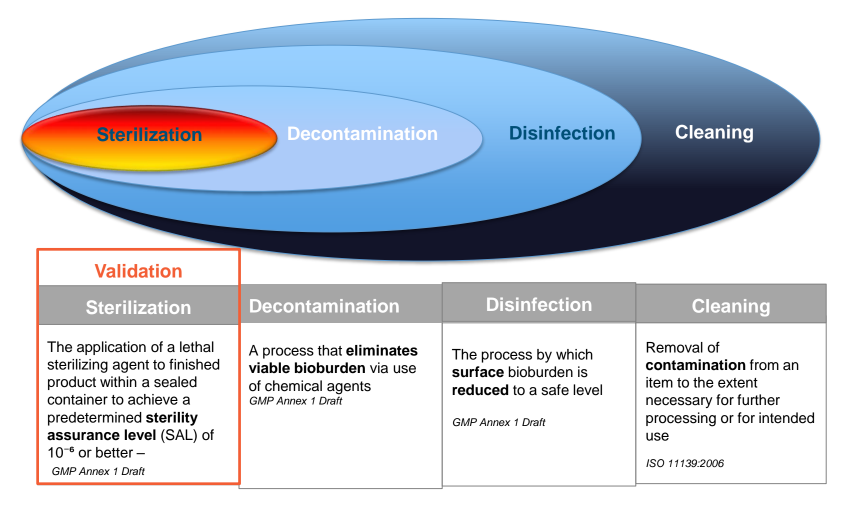
- Decontamination Vs Sterilization
- Terminal Sterilization Vs Aseptic Assembly
- Method selection







## Decontamination Vs Sterilization









#### A sterile product is one that is free of viable microorganisms



#### Absolute sterility can never be guaranteed!

- 100% control of the batch is not possible
- No assurance that any microorganism can be detected during Sterility Test





## Sterility Assurance Level (SAL) = The probability

of a single item in a batch being non-sterile after being subjected to a sterilization process.

## Sterile: SAL ≤ 10<sup>-6</sup>

## SAL likelihood of surviving organisms

 $10^{-1} = 1:10$ 

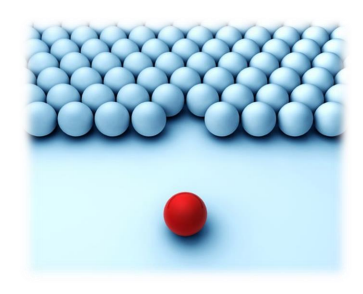
 $10^{-2} = 1:100$ 

 $10^{-3} = 1:1,000$ 

 $10^{-4} = 1:10,000$ 

 $10^{-5} = 1:100,000$ 

 $10^{-6} = 1:1,000,000$ 







## Sterility is much more than just a process!

### **Initial** contamination level

- Microbiological status raw material and components
- Cleaning and disinfection procedures
- **Environment control**
- Personnel Hygiene

#### **Equipment**

- Control
- Maintenance
- Calibration

## **Product** preservation

- **Packaging**
- Storage









## Pharmaceutical Product Life Cycle



Think about sterilisation as soon as possible during product development





### Sterile means: Safe Product & Functional product



Selection of the right sterilization method is critical!





# There are two (2) methods to produce a sterile drug product:

Aseptic Assembly

## Terminal Sterilisation

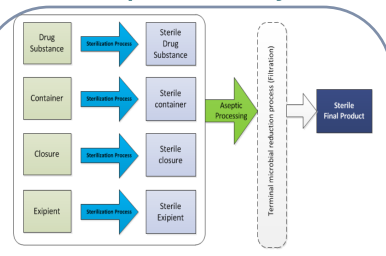
- Heat
- Irradiation
- Gas (EO, NO<sub>2</sub> ...)



#### **Sterilization – Basics**



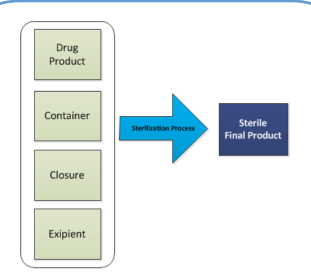
#### **Aseptic Assembly**



Maintain sterility of a product that is assembled from components, each of which has been previously sterilized

## **Sterile**

#### **Terminal Sterilization**



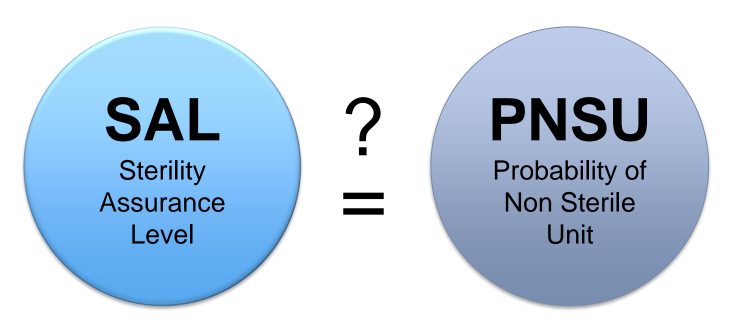
**Exposure** to a physical or chemical sterilizing agent for a predetermined extent of treatment

## **Sterilized**





## Is the effectiveness of a sterilization process assessed the same way for AA or TS products?



Reference: ISO TS 19930:2017





**PNSU** 

## Probability of Non Sterile Unit (PNSU):

The probability of one or more microorganism being present on a product item in a population of items.

- No SAL can be calculated for aseptically assembled products (no killing kinetic)
- Rely on multiple factors
- Estimated by Media Fill Test (process simulation)
- Bracketing approach (MFT before and after process)







#### Selection of the Sterilization Method:



"Wherever possible, a process in which the product is sterilized in its final container (terminal sterilization) is chosen"

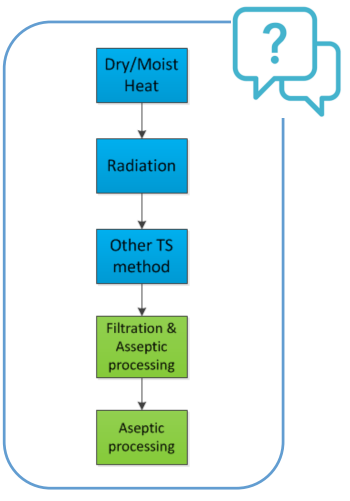
**European Pharmacopoeia 9.7** 

Per PDA 2017 Survey – 30% of Aseptically assembled product could be Terminally sterilized!



#### **Sterilization – Basics**





## Selection of the Sterilization Method:

Use a **structured approach** to select the most appropriate sterilisation method

Based on EMA - CPMP/QWP/054/98 Decision Tree for the selection of sterilisation methods



#### Sterilization - Basics



## Prior to making your choice, consider mitigation options:

 Can your formula be adapted (limit degradation and impurities)?

- Can the **container** be adapted?
- Can you select compatible component with selected sterilization process?
- Can the process can be optimized (limit impact)?

CPMP/QWP/054/98 Decision Tree for the selection of sterilisation methods









#### **Ethylene Oxide**

• (EO) gas



#### Irradiation

- Gamma ray
- Accelerated electrons (E-beam)
- X-rays



for terminal sterilization of single use medical devices



#### Other

- Moist heat
- Dry heat
- Vaporized hydrogen peroxide (VHP)
- Gas plasma
- Low temp Steam Formaldehyde (LTSF)
- Nitrogen Dioxide (NO<sub>2</sub>)





## Radiation Technology

- General principles
- Gamma
- E-Beam
- Sterilization validation



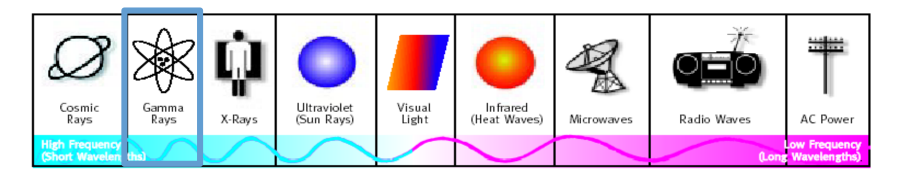


#### **Sterilization by Irradiation – General principles**

### **General Terminology**

#### Radioactivity:

Electromagnetic radiation (photons) produced by radioactive decay.



Ionising

Non-Ionising

**E-beam** = Electrons (with a mass)





#### **Sterilization by Irradiation : General principles**

## **General Terminology**

#### **Radiation**

Energy in the form of waves or moving subatomic particles

#### Radioactive

Substance emitting radiation



Exposure to radiation

≠ Making something radioactive





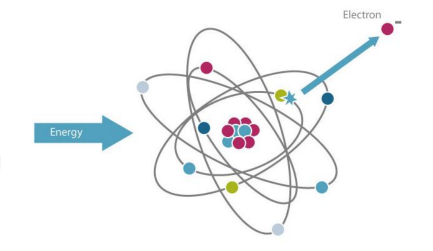


#### **Sterilization by Irradiation : General principles**

## **General Terminology**

#### **Ionising Radiation**

Radiation capable of knocking electrons out of their thermal orbits in atoms or molecules



#### (Absorbed) Dose

Measure of the amount of energy that is absorbed by the material while exposed to a radiation source.

Unit: Gray 1 Gy = 1 Joule per Kg material





#### **Sterilization by Irradiation: Monitoring**

## Irradiation process monitoring:

#### **Dosimeter**

Device having a reproducible, measurable response to radiation, which can be used to measure the obsorbed dose in a given system.



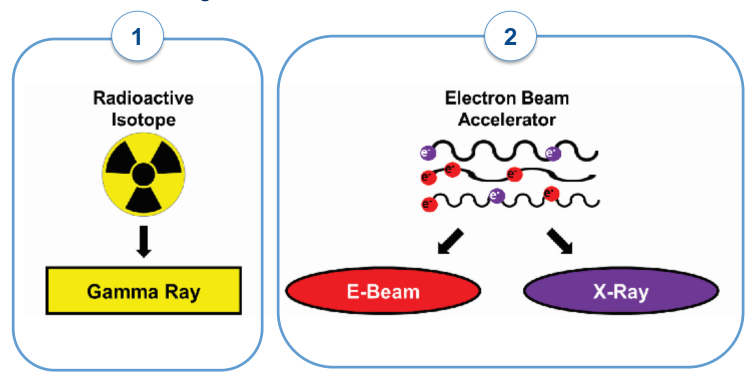
0 kGy 12 kGy 25 kGy 50 kGy 0kGy





#### **Sterilization by Irradiation – General principles**

#### Two methods to generate irradiation:









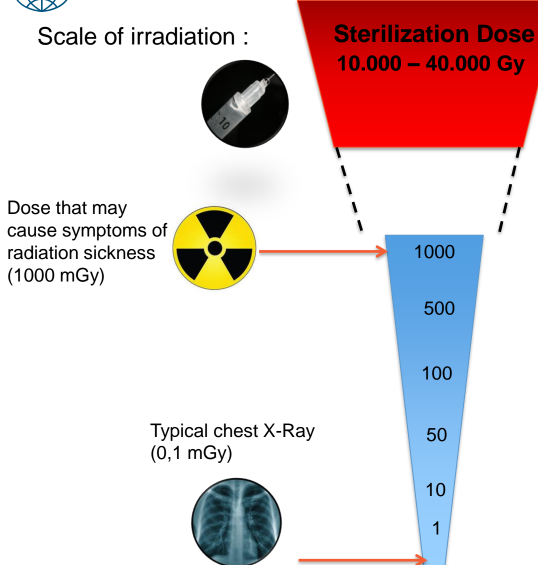
### **Gamma Irradiation**

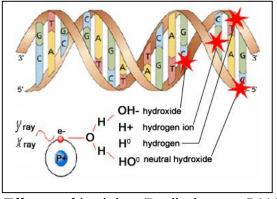






#### **Sterilization by Irradiation: Gamma**





Effects of ionizing Radiation on DNA



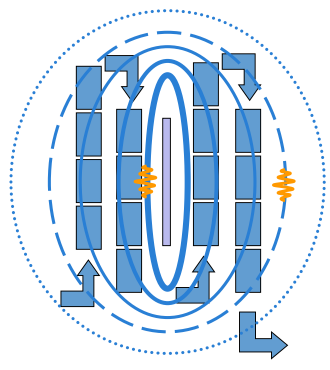


#### **Sterilization by Irradiation: Gamma**

**Source:** 60Co (mostly)

**Decay rate:** 12% per year (Half life 5,3 years)

Source Activity: Several Million Ci



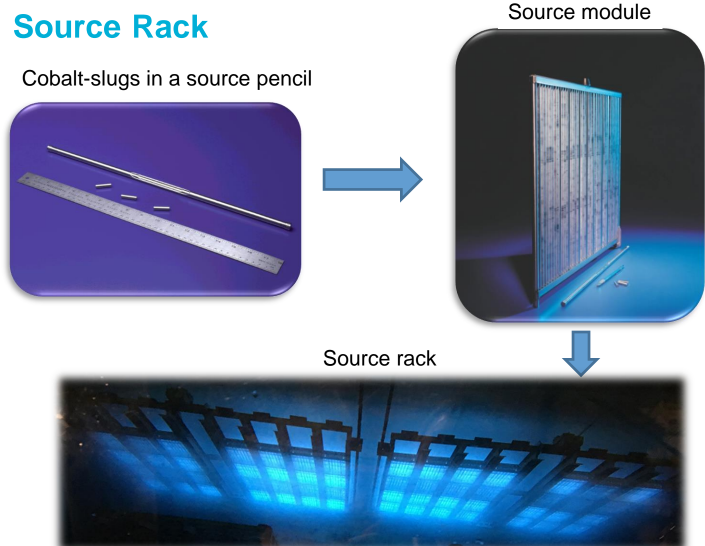








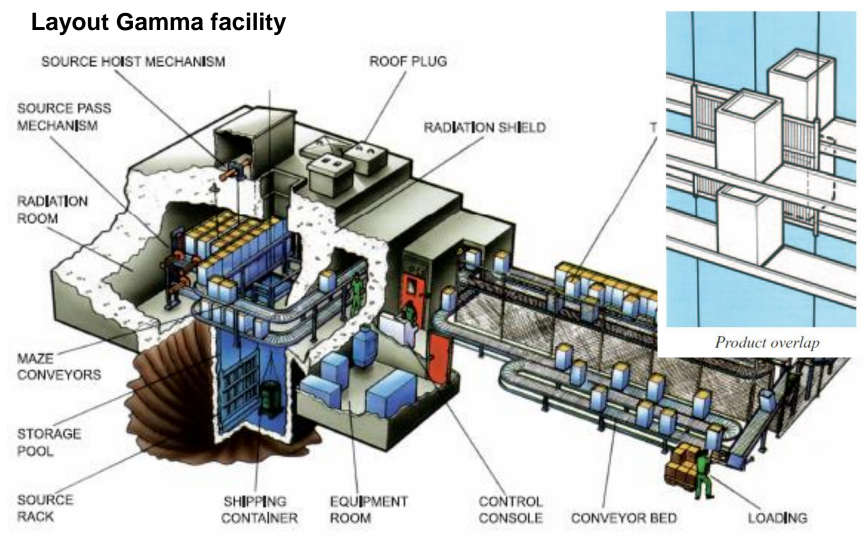
### **Sterilization by Irradiation : Gamma**







#### **Sterilization by Irradiation: Gamma**

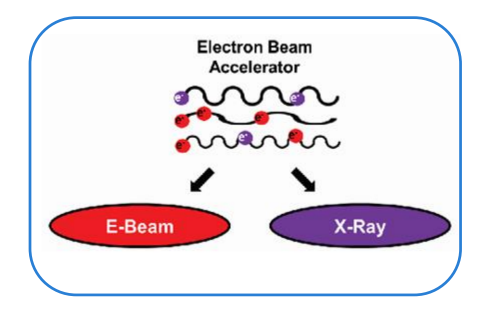








#### **E- Beam irradiation**







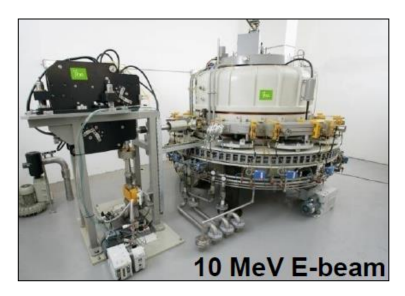
#### **Sterilization by Irradiation : E-Beam**

#### **Electron Beam**

Directed stream of electrons (B radiation) produced by a particle accelerator

#### Beam energy

Speed of the electrons. Parameter related to depth of penetration Limited to 10 MeV for medical device sterilisation (ISO 11137-1) to avoid radioactivity induced in product



**IBA Rhodotron** 





### **Sterilization by Irradiation : E-Beam**

#### **Layout E-Beam facility**

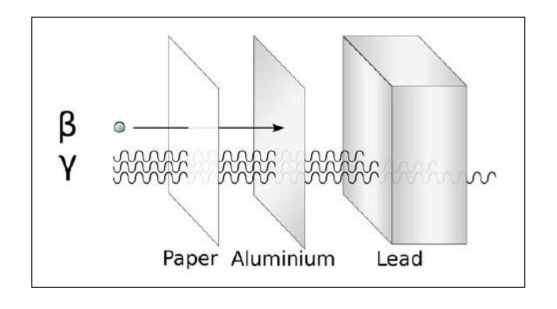








### **Electron Beam & Gamma, Penetration**







### **Sterilization by Irradiation: comparison**

Parameter	Gamma	E-Beam
Irradiation parameter	Cycle Time Density	Conveyor speed Density Scan width Beam energy
<b>Radiation Field</b>	Isotroptic	Highly directional
Geometry of material and heterogeneity of Product	Important to consider	Critical





## **Sterilization by Irradiation: comparison**

Parameter	Gamma	E-Beam
Product Treatment	Pallet/Tote	Boxes
Dose Rate (Dmin 25KGy)	Hours	Seconds
Dose uniformity ration (DUR)	Low sensitivity to product thickness	sensitivite to product thickness
On/Off Technology	No	Yes
Flexible Target Dose	No	Yes
Process validation	Straightforward	Potentially complicated





#### **Sterilization by Irradiation : validation principles**

#### **Relevant Standards:**

#### ISO 11137-1:2015

Sterilization of health care products – Radiation – Part 1: Requirements for development, validation, and routine control of a sterilization process for medical devices

#### ISO 11137-2: 2015

Sterilization of health care products – Radiation – Part 2: Establishing the sterilization dose

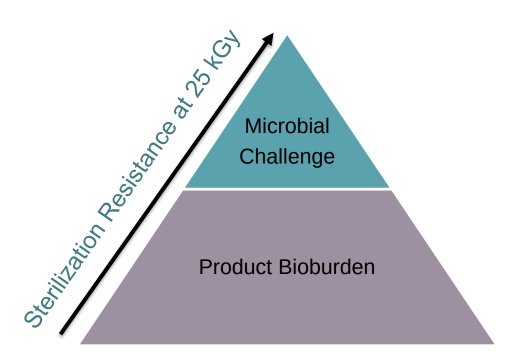
#### **GMP – Annex 12**

Use of ionising radiation in the manufacture of medicinal products

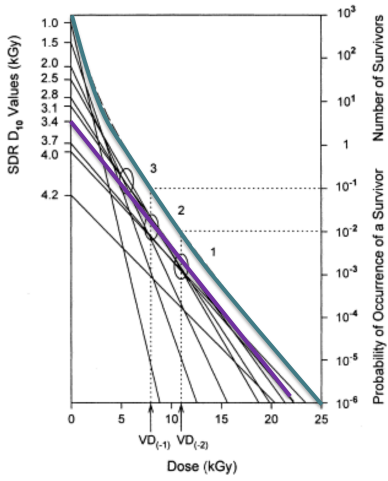






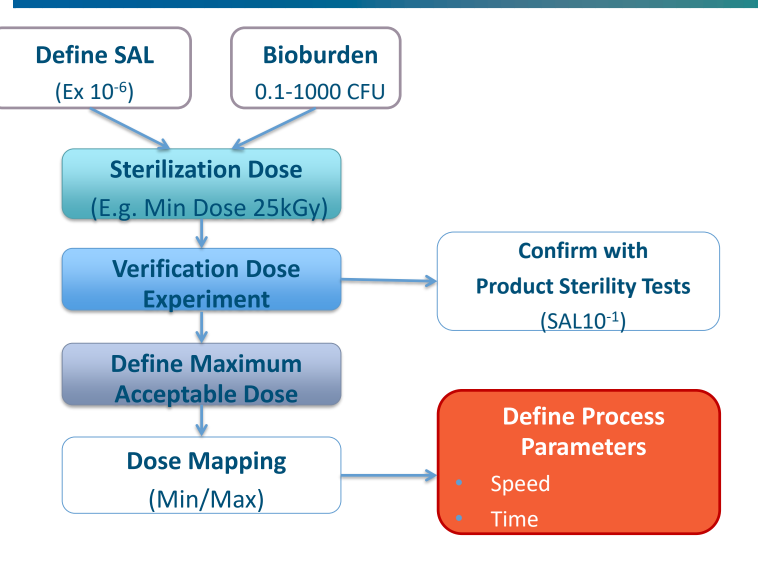


#### Standard Distribution of resistances (SDR)













#### Bioburden is critical parameter in Irradiation technology

Sample Item Portion (SIP) is frequently used for bioburden evaluation . Basis for SIP can be:

#### Length

Consistent diameter tubing



#### Mass

- Powders
- Gowns
- Absorbable implants



#### Volume

•



#### **Surface Area**

- Non-absorbable implants
- Variable diameter tubing







## Select Sterilization Dose

 $Method\ VD_{max}$ 

Example minimum

Dose to apply related to bioburden

Bioburden Range	Dose (kGy)
≤ 0.1 to 1.5	15.0
≤ 0.1 to 9.0	17.5
≤ 0.1 to 45	20.0
≤ 0.1 to 220	22.5
≤ 0.1 to 1000	25.0
≤ 1.0 to 5000	27.5
$\leq$ 1.0 to 23,000	30.0
$\leq$ 1.0 to 100,000	32.5
≤ 1.0 to 440,000	35.0





Select Verification Dose: VD<sub>max</sub><sup>25</sup>

Bioburden	Verification Dose (kGy	
40	8.6	
45	8.7	
50	8.8	
55	8.9	

Verification is conducted at an SAL of 10-1 with 10 product items irradiated.

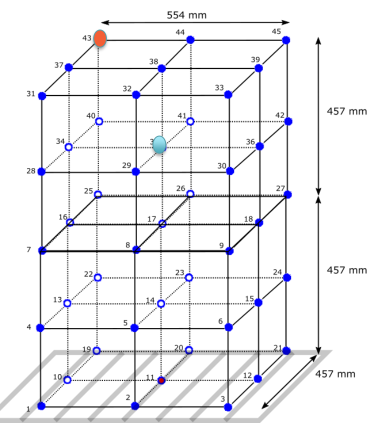




## **Dose Mapping**

Establish the distribution of absorbed dose within the irradiation container when packed with product in a defined configuration

- Min and Max limits of absorbed Dose
- Define cycle time
- Establish monitoring points
  - Min Dose = 28KGy
  - Max Dose = 37KGy







#### Quarterly Dose Audit (QDA)

#### Check bioburden

Can vary due to

- Season
- Environment ...

#### **Verification Dose**

(Often In a research irradiator)

Ex: 8,7KGy

## **Sterility Test**

SAL10<sup>-1</sup>



Confirm Product
SAL 10<sup>-6</sup>

With Routine Dose

Ex:25KGy





## **Sterilization by Irradiation: examples**

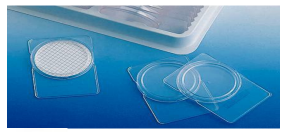


























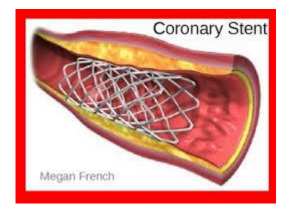




## **Sterilization by Irradiation: examples**

## ... But also







Grafts





**API** 







## Summary

Minium & Maximum dose to product shall be defined

Methods 1, 2, VDmax, "equivalent method"

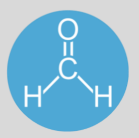
Based on natural product bioburden

Routine process monitored with dosimeters

**Quarterly Dose Audit (QDA)** required





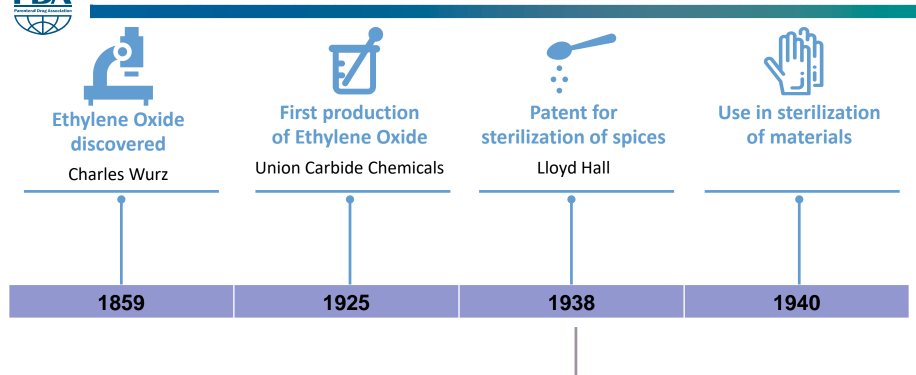


## Ethylene Oxide Sterilization

Introduction



### **Sterilization by Ethylene Oxide: History**





Dr. Lloyd Augustus Hall, a food scientist, while working for Griffith Laboratories, devised a process known as the Ethylene Oxide Vacugas treatment to control the growth of molds and bacteria. Griffith and Hall received US Patent 2,189,949 in 1940.

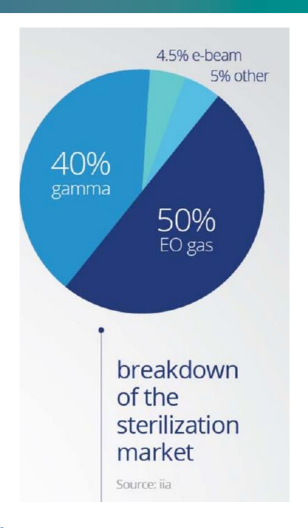






## **Properties**

- Toxic gas
- "Sweet smell" from ca. 500 ppm concentration
- Forms with air explosive mixtures (2.6 %)
- Oncogenic by inhalation
- Irritating for skin and respiratory system
- Mutagenic for animals and very likely for humans



Last choice but sometimes the only one!

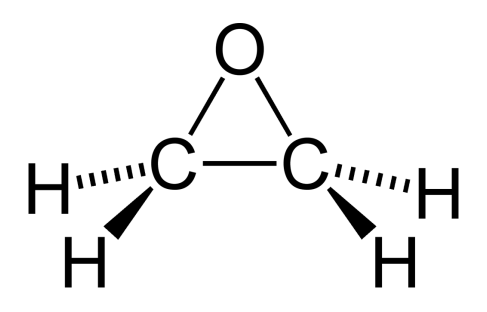






## Mode of Action

- Extremely reactive
- Irreversible reaction with DNA and proteins (alkylation)
  - The molecule is loses function
  - Replication stops
  - The cell dies







#### Mainly used to sterilize:

- Heat-sensitive material
- Material sensitive to ionizing radiation
- High Volumes
- Packs with multiple components







# Device/packaging must be permeable to the gas

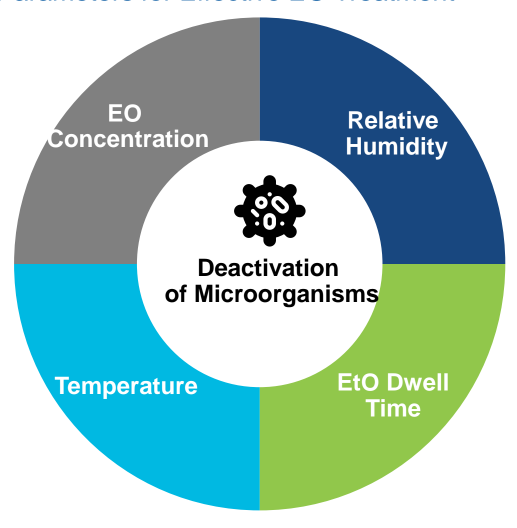
- No aqueous substances
- No protein-type materials
- Powders, batteries, electronic circuits have to be assessed (risk of explosion)
- Vacuum/heat can have adverse impact on some packaging (bubble wrap packaging, polystyrene)







#### Critical Parameters for Effective EO Treatment







## Temperature (T)

## EtO kills microorganis

ms even at temperatures below 10°C (50oF)

## Industrial sterilization

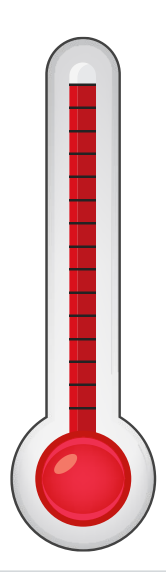
performed in 40-60 °C (104–140oF) temperature range

#### **Q10 Effect**

increase by 10°C (18°F) = 2x Deactivation Rate

## Temperature increase

may increase of permeability of gases through materials





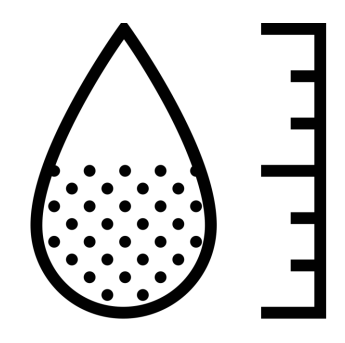


## Relative Humidity (RH)

Necessary for alkylation reaction

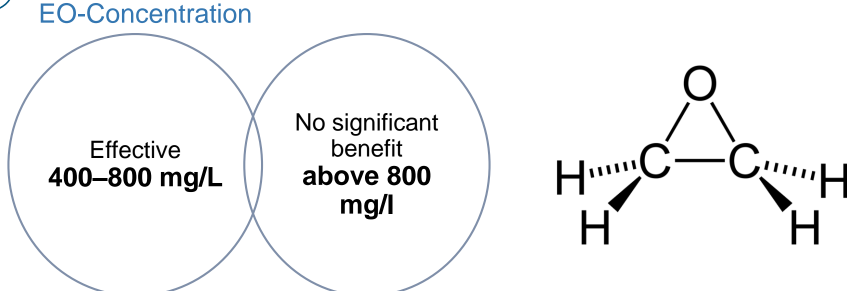
Relative
humidity may
assist
penetration of
EO through
materials

EO is most effective at RH > 30%









At constant T and RH – if EO concentration increases microbiological Deactivation is more effective - up to c. 800 mg/l

- ~ 500 mg/L @ 131°F
- ~ 800 mg/L @ 86°F





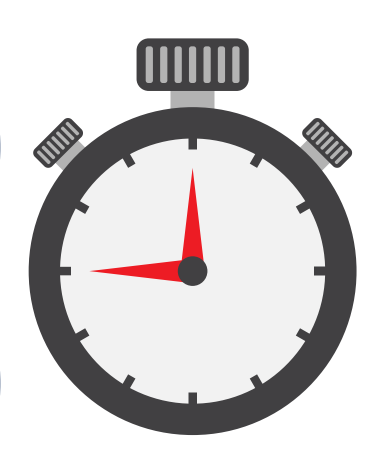
#### Time

## Microbiological deactivation

is more effective with longer gas dwell phase

## **Industry cycles**

2 to 10 hours gas dwell phase Typically 3-4 hours









## **Customer Needs To Define**

Product
Families/Processing
Categories

Finalize Packaging

**Load Configuration** 

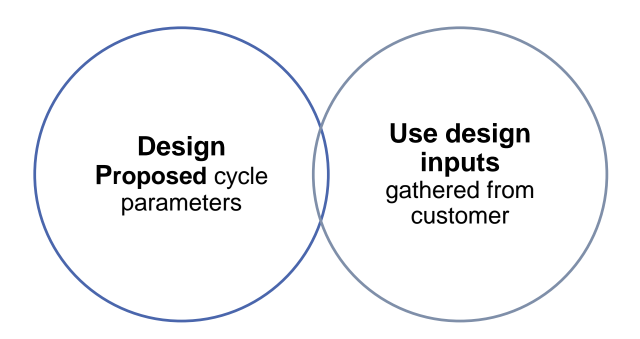
Bioburden

Internal PCD













## **Sterilization Process**



3 key phases



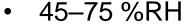


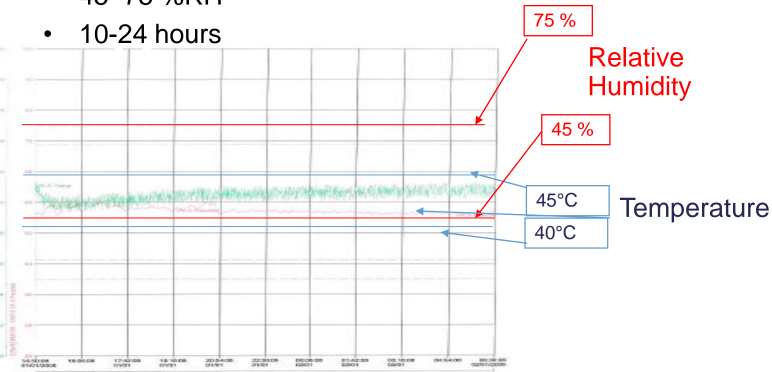
## Preconditioning (





40-45°C





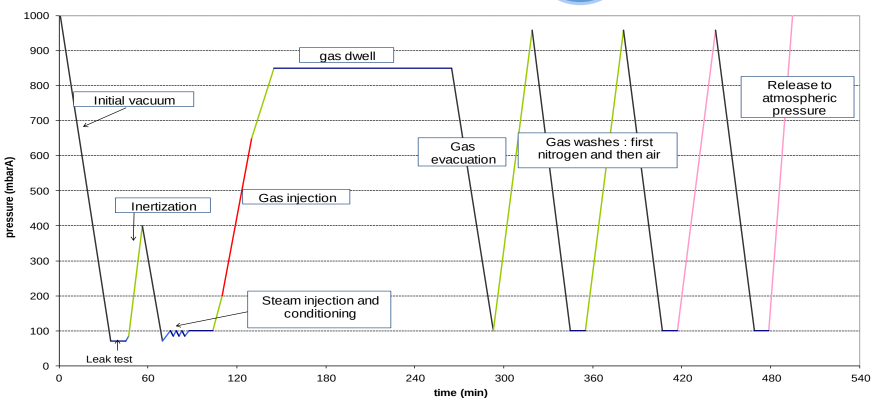




## Typical EO Cycle Design – Deep Vacuum



#### **GENERIC CYCLE**





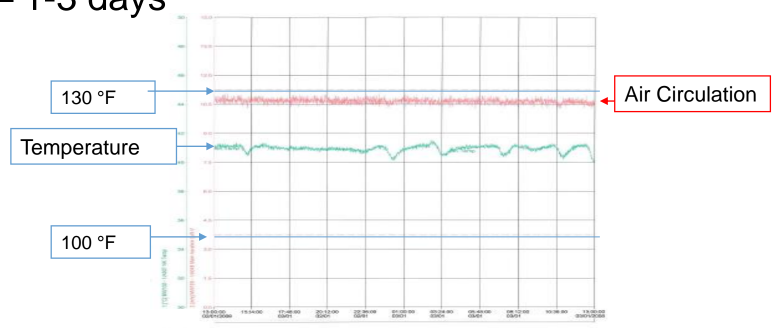






- -45°C
- Forced circulation

1-3 days

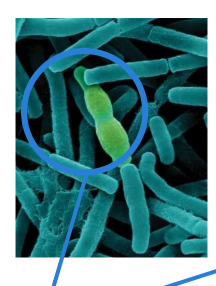


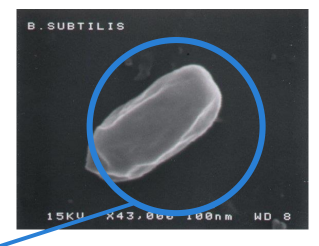




#### Monitoring EO Sterilization - Biological Indicators

- Usually, the BI contains at least a million spores (>10Exp6) of an organism that is highly-resistant to the EO process (Bacillus atrophaeus)
- Growth is very characteristic (orange ring)





**Spore** 



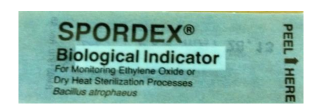




## Process Challenge Device (PCD)

Item designed to constitute a defined resistance to the sterilization process and used to assess performance of the process

- Internal PCD (IPCD)
- External PCD (EPCD)



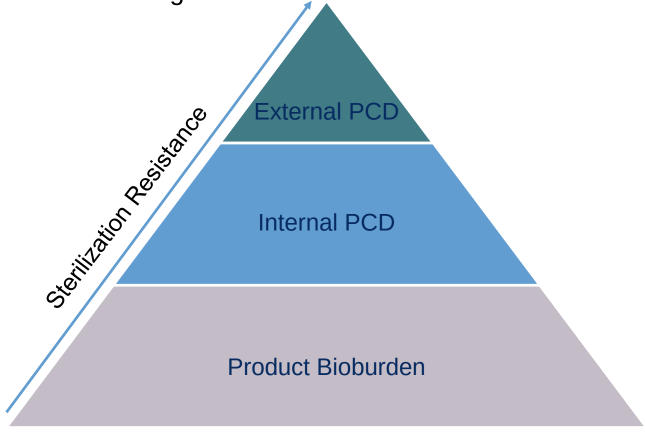






## Monitoring EO Sterilization – Biological Indicators

We design the validation to show that the **BI** is more difficult to kill than natural occurring bioburden



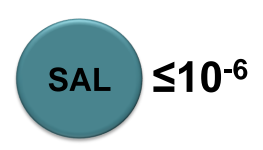




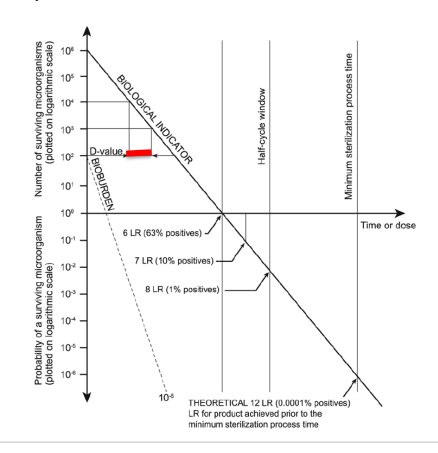


#### **D** Value

The Time needed to deactivate 90% of population of microorganisms (or 1 Log Reduction)



The sterilization cycle is validated to predict achievement of an SAL equal to or less than a specified value (≥12LR)







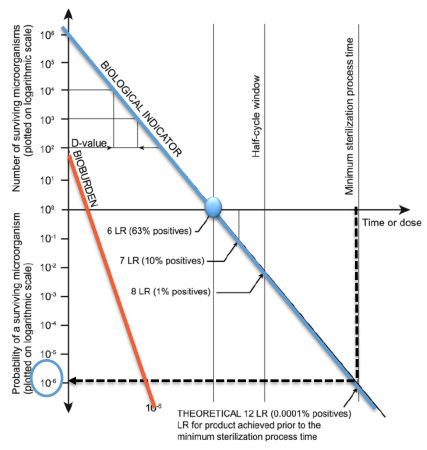
### Sterilization by Ethylene Oxide: Validation principle

## Level of Sterility Assurance

#### Example:

 $D_{value} IPCD = 15min = 1LR$ 

6 LR = 90 min (Half cycle) 12 LR =180 min (Full cycle)







### **Sterilization by ETO: validation principles**

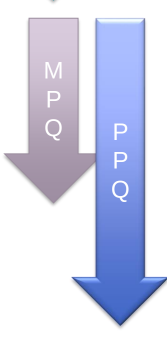


- Establish Product/IPCD D<sub>Value</sub>
- Product Natural bioburden killed
- Define Challenges (IPCD -EPCD)





- Confirm IPCD selection (SAL ≤10<sup>-1</sup>)
- Confirm External Challenge (EPCD)





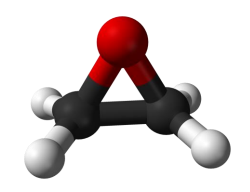
- SAL ≤10<sup>-6</sup>
- Aeration validation Residue Tests



#### Sterilization by Ethylene Oxide: Validation principle

## Compounds that remain on product after EO sterilization

- Ethylene Oxide (EO)
- Ethylene Chlorohydrin (ECH) = EO + HCL
- Ethylene Glycol (EG) = EO + H2O



ISO 10993-7:2008

"Biological Evaluation Of Medical Devices-

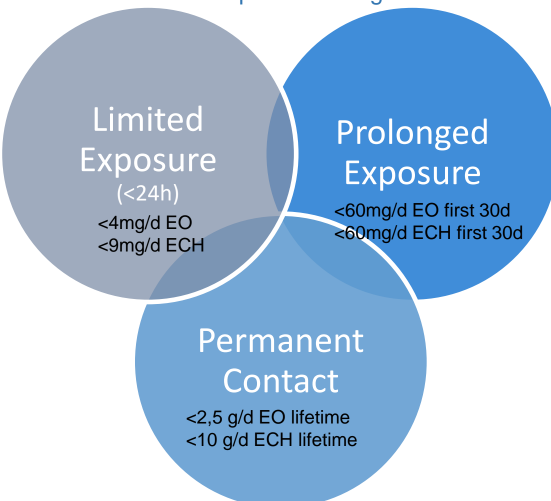
**Part 7: Ethylene Oxide Sterilization Residuals**"





#### **Sterilization by Ethylene Oxide: Validation principle**

There are Three Patient Exposure Categories







#### ISO 11135:2014

Sterilization of medical devices – Requirements for the development; validation and routine Control of a Sterilization Process for Medical Devices – Ethylene Oxide

#### **ETO Residuals**

ISO 10993-7:2008 (R) 2012

Biological evaluation of medical devices - Part 7: Ethylene oxide sterilization residuals

#### **GMP – Annex 1 (Draft)**

Manufacture of Sterile medicinal Products

#### Bioburden

ISO 11737-1:2018

Sterilization of medical devices (Microbiological methods) Part 1: Determination of a population of microorganisms on products





#### **Product Sterility**

• ISO 11737-2:2009 (R) 2014

Sterilization of medical devices (Microbiological methods) Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process

- United States Pharmacopeia (USP) Chapter <71> Sterility Tests
- European Pharmacopeia (EP) Chapter 2.6.1 Sterility
- Japanese Pharmacopeia (JP) Chapter 54. Sterility Test

#### **Biological Indicator Tests**

• ISO 11138-1:2017

Sterilization of health care products (Biological indicators) Part 1: General requirements

• ISO 11138-2:2017

Sterilization of health care products (Biological indicators)Part 2: Biological indicators for ethylene oxide sterilization processes

ISO 14161: 2009 (R) 2014

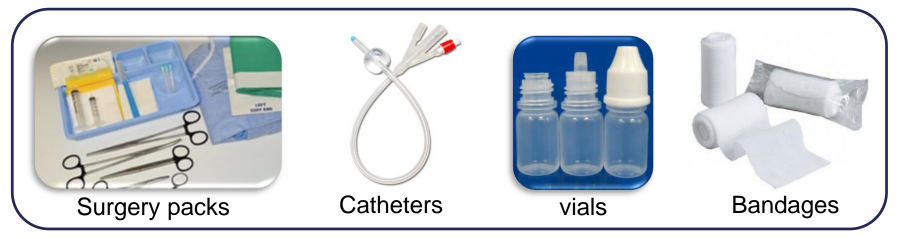
Biological indicators. Guidance for the selection, use and interpretation of results



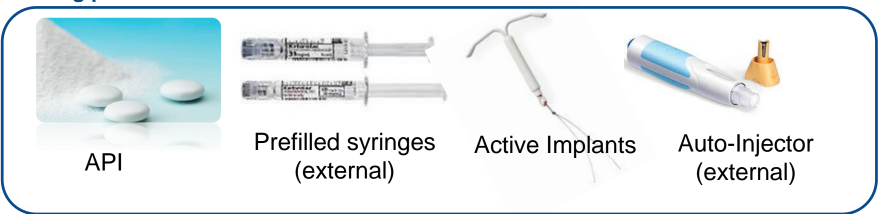


### **Sterilization by Ethylene Oxide : Product examples**

#### **Medical Devices**



#### **Drug products**







## **Sterilization: Comparison Radiation & Ethylene Oxide**

Parameter	Gamma	E-Beam	EO
Process	Individual product, box, tote, pallet	Boxes	Pallets
Material compatibility	Not compatible with some type of polymers (PTFE and PVC affected)	Wider polymer compatibility compared to Gamma	Very good No liquid/proteins Low Temperature (40-55°C)
Validation	Straightforward	Straightforward	Complicate
Validation principle	Based on bioburden	Based on bioburden	Based on Bio Indicators
Requalification	Every 3 months (QDA)	Every 3 months (QDA)	Every 2 years (1 cycle)
SAL	<10exp6	<10exp6	<10exp6
Residues	None	None	ETO,ECH,(EG)





## **Sterilization: Comparison Radiation & Ethylene Oxide**

Parameter	Gamma	E-Beam	EO
Process	Individual product, box, tote, pallet	Boxes	Pallets
Material compatibility	Not compatible with some type of polymers (PTFE and PVC affected)	Wider polymer compatibility compared to Gamma	Very good No liquid/proteins Low Temperature (40-55°C)
Tolerance for density variation	High	Low	Medium
Routine monitoring	<ul><li>Only a few parameters (Time, Size, density)</li><li>Dosimeter</li></ul>	<ul><li>Higher Nb of parameters</li><li>Dosimeter</li></ul>	<ul> <li>Multiple cycle parameters</li> <li>BI (unless parametric release)</li> </ul>
Volumes	High	Limited	High
Turn time	Fast (<24 hours)	Very Fast (<8 hours)	Long (1 week)







You should now feel better at ease with the following concepts:

- What is sterilization Vs decontamination
- Aseptic Assembly Vs Terminal sterilization
- Irradiation sterilization (Gamma and Ebeam)
- Ethylene Oxide sterilization





#### **Reference Slide**

- ISO 11135:2014 Sterilization of medical devices Requirements for the development; validation and routine Control of a Sterilization Process for Medical Devices – Ethylene Oxide
- ISO 10993-7:2008 (R) 2012 Biological evaluation of medical devices Part 7: Ethylene oxide sterilization residuals
- 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
- ISO 11137-2 Sterilization of health care products Radiation Part 2: Establishing the sterilization dose
- ISO 11737-1:2018 Sterilization of medical devices (Microbiological methods) Part 1: Determination of a population of microorganisms on products
- ISO 11737-2:2009 (R) 2014
- Sterilization of medical devices (Microbiological methods) Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process
- ISO 11138-1:2017
- · Sterilization of health care products (Biological indicators) Part 1: General requirements
- ISO 11138-2:2017
- Sterilization of health care products (Biological indicators)Part 2: Biological indicators for ethylene oxide sterilization processes
- ISO 14161: 2009 (R) 2014
- · Biological indicators. Guidance for the selection, use and interpretation of results





#### **Reference Slide**

- ISO 11737-2:2009 (R) 2014
   Sterilization of medical devices (Microbiological methods) Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process
- ISO TS 19930:2017 Guidance on aspects of a risk-based approach to assuring sterility of a terminally-sterilized, single use health care product unable to withstand processing to achieve maximally a sterility assurance level of 10-6
- AAMI TIR 33 Sterilization of health care products—Radiation—Substantiation of a selected sterilization dose Method Vdmax
- United States Pharmacopeia (USP) Chapter <71> Sterility Tests
- Eudralex Volume 4 GMP Annex 1
- Eudralex Volume 4 GMP Annex 12
- European Pharmacopeia (EP) Chapter 2.6.1 Sterility
- The Aseptic and Sterile Processing: Control, Compliance and Future Trends Edited by Tim Sandle, Edward Tidswell PDA – 2017
- PDA Survey: 2017 PDA Aseptic Processing
- A comparison of Gamma, E-beam, X-Ray and ETO technologies for the industrial Sterilization of MD and Health care products – GIPA, IIA – 31 Aug 2017







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