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Degree in Chemistry

10 years in Pharma companies, in QA, QC deparments

R&D Manager, Fedegari Autoclavi Spa, since 2010

The sentence which best describes me: «Stay hungry, stay foolish» S.Jobs

Specialties: Pharmaceutical industry - Commissioning - IQ/OQ/PQ - SOP -Sterile Process definition and Validation— Vaporized Phase Hydrogen Peroxide technology - Autoclaves - Washing machines and processes - Sterilization Oven – CIP/SIP skids — Aseptic isolators





Hydrogen peroxide decontamination

Maria Luisa Bernuzzi, Fedegari group







Hydrogen peroxide session: program of the day

AUDITORIUM:

OVERVIEW OF THE MAIN TOPICS TREATED

R&D DEPARTMENT:

PRACTICAL SESSION



HYDROGEN PEROXIDE MEASUREMENT





Hydrogen peroxide session: main topics

- Hydrogen peroxide definition
- Regulation
- Application fields
- Decontamination target
- Decontamination technologies:
 VPHP (dry or wet cycle), DRY FOG
- Sporicidal Concentration
- Materials
- Packaging Integrity verification

- Sensors
- Safety
- Catalyzer
- Example of dry cycle
- Hydrogen peroxide mapping
- Biological indicators and D-value
- SLR calculation





Hydrogen peroxide definition

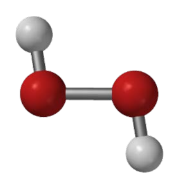


Hydrogen peroxide is a strong oxidizing agent used in aqueous solution as a ripening agent, bleach, and topical anti-infective. It is relatively unstable and solutions deteriorate over time unless stabilized by the addition of acetanilide or similar organic materials.





Hydrogen peroxide classification



USP 41, General Chapter (1072) - DISINFECTANTS AND ANTISEPTICS

Chemical Entity	Classification	Example
Hydrogen peroxide	Vapor phase sterilant, liquid sporicidal agent, antiseptic	4 μg per g H2O2 vapor, 10%– 25% solution, 3% solution





Hydrogen peroxide: European regulation

ANNEX 1 EU GMP, draft

5.34 Fumigation or vapour disinfection of clean areas such as Vapour Hydrogen Peroxide (VHP) may be useful for reducing microbiological contamination in inaccessible places.

5.19 For open, positive pressure isolators or closed isolators with decontamination by a sporicidal agent, the surrounding area should correspond to a minimum of grade D.





Hydrogen peroxide: US regulation

USP 41, General Chapter (1208) STERILITY TESTING; VALIDATION OF ISOLATOR SYSTEMS

Among the chemicals that have been used to treat isolators are peracetic acid, chlorine dioxide, ozone, and hydrogen peroxide; each has different requirements for exposure conditions and process control.







Decontamination: when?

✓ Heat sensitive materials (including electronic devices) that should be transferred between classified areas (class C,D → class A, B) in order to minimize the risk of contamination



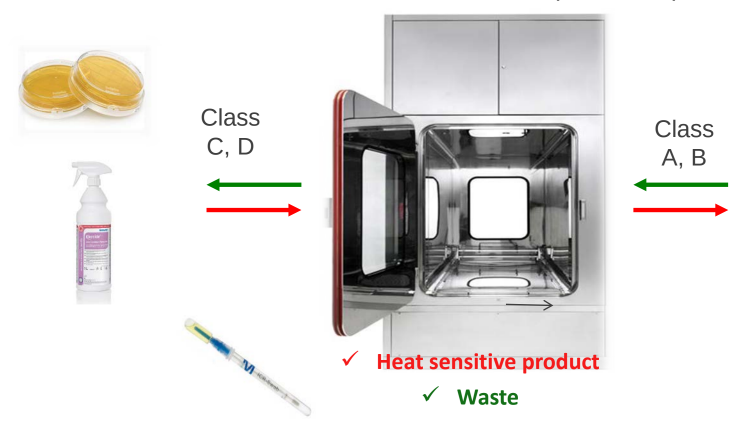
✓ Surface of aseptic processing rooms (ex. cleanroom) and of aseptic processing systems (ex. isolators)





Decontamination: when?

Decontamination unit (Pass Box)







Decontamination: when?

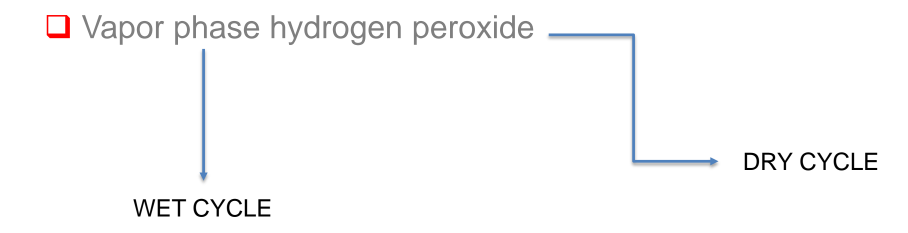






Decontamination technologies

The most widespread technologies



Dry fog





VAPOR PHASE HYDROGEN PEROXIDE

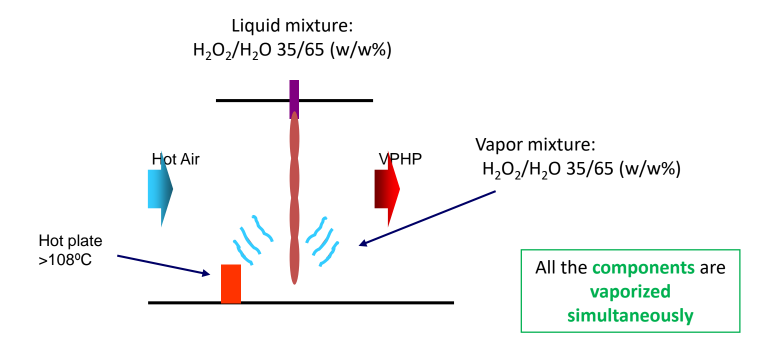
Vapor Phase Hydrogen Peroxide (VPHP): how is it produced?







VPHP Production: Flash Vaporization

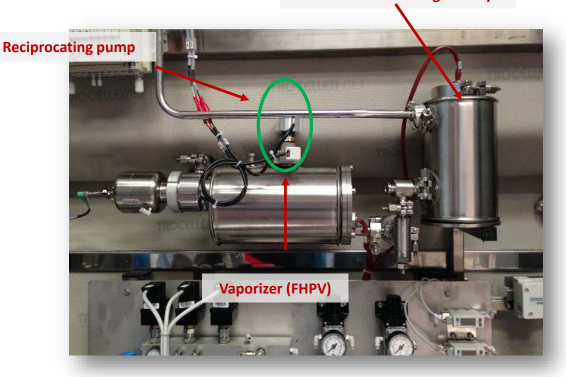






Fedegari Hydrogen Peroxide Vaporizer (FHPV)

Air electrical heating battery

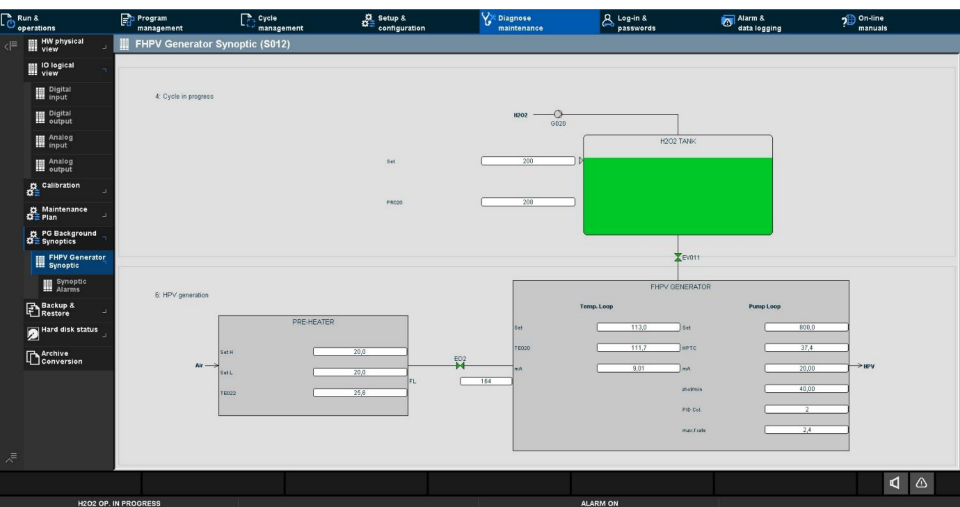


It produces vaporized hydrogen peroxide from the H₂O₂/H₂O liquid mixture



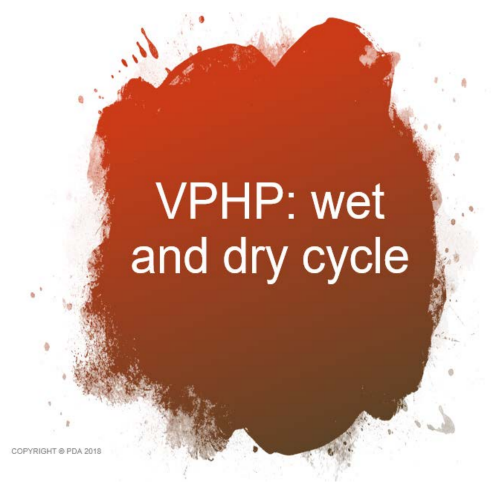


FHPV generator synoptic









Wet cycle

- effective in a short time lapse
- more penetrating
- wet load
- ☐ long cycle
- ☐ more agressive on materials
- concentration not well controlled

Dry cycle

- ☐ effective in a longer time lapse
- ☐ less penetrating
- ☐ dry load
- □ shorter cycle
- ☐ less agressive on materials
- □ concentration, well controlled





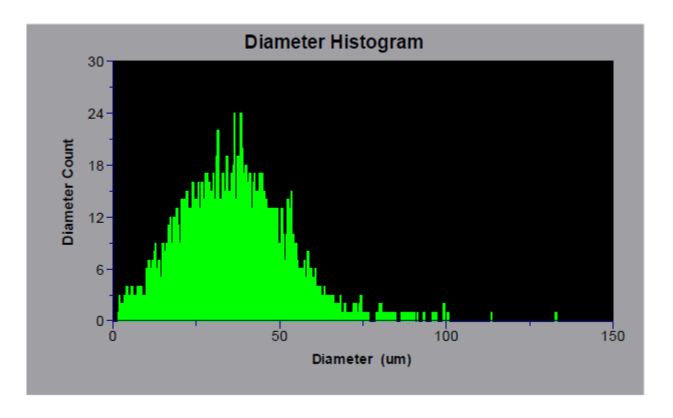
Dry fog

- Penetrating
- □ Control based on reading RH/ injecting grams
- □ No reliable concentration control





Dry fog



Droplets dimension distribution





Hydrogen peroxide concentration

3TANDARD» PERCENTAGE

SAFETY DATA SHEET
Hydrogen Peroxide 35% Durox® LRA

SDS #: 7722-84-1-35-27 Revision date: 2015-05-08

> Format: NA Version 1



1. PRODUCT AND COMPANY IDENTIFICATION

Product Identifier

Product Name Hydrogen Peroxide 35% Durox® LRA

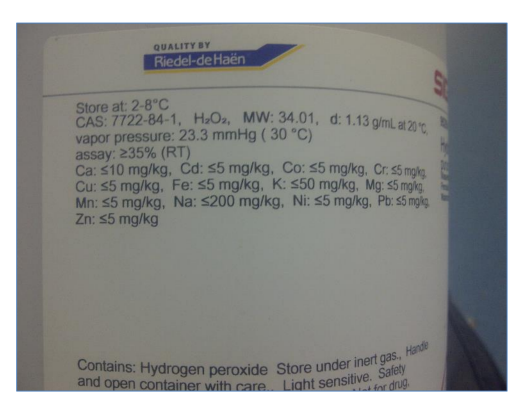
Other means of identification

CAS-No 7722-84-1





Hydrogen peroxide concentration









Hydrogen peroxide concentration

SUMMARY

On the test item "Metallic device in VHP system", analyses have been performed for the verification of the possible presence of residues. In particular, the presence of typical inorganic H_2O_2 stabilizers were investigated.

In fact the device underwent:

- Determination of silicon/silica (performed on a washing aqueous solution)
- Determination of phosphates, nitrates, sulphates (performed on a washing aqueous solution)

INTRODUCTION

On behalf of FEDEGARI AUTOCLAVI SpA has been performed a study for the verification of the possible presence of residues on the test item.





Hydrogen peroxide residues

4. SILICA

Silicon detected using ICP technique (see 3. Silicon paragraph) is silicon dissolved in the solution. Presumably all the silicon detected with this technique is related to the presence of dissolved silicates in the washing solution. Metallic silicon is not detectable not being dissolved.

The results obtained for silicon will then be processed so as to express the content of silicates in solution expressed as silica equivalent.

RESULTS

All the results are related to the analytes present in the washing solution (400ml).

1. NITRATES, SULPHATES and PHOSPHATES

Nitrates (mg/L)	Sulphates (mg/L)	Phosphates (mg/L)
5.47	21.34	74.82

2. SILICON and SILICA

Silicon (mg/L)	Silica (mg/L)
0.157	0.336

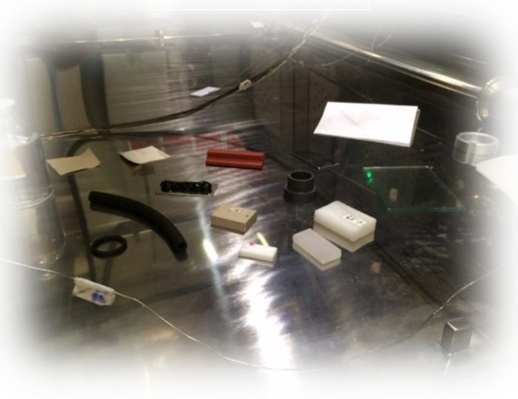




Materials compatibility

- Deterioration of elasticity or strenght or flexibility, visible damages
- Absorption Degassing time
- Penetration Product damage
- Microbiological effectiveness -Surface finish

MACHINE PARTS







Materials compatibility







EXAMPLES OF CRITICAL LOADS





Package integrity verification

USP 41, GENERAL CHAPTER (1208) STERILITY TESTING; VALIDATION OF ISOLATOR SYSTEMS: PACKAGE INTEGRITY VERIFICATION

Some materials are adversely affected which by decontaminating agents, can result in inhibition of microbial growth. Of concern is the penetration of decontaminating agents into product containers.





Material compatibility: SEM investigation A case study

Different materials were inoculated with 10⁶ Geobacillus Stearothermophilus spores and analyzed by scanning electron microscope (SEM)

Spore monolayers



Good substrate for decontamination

Spore clusters
Spores in grooves or cavities



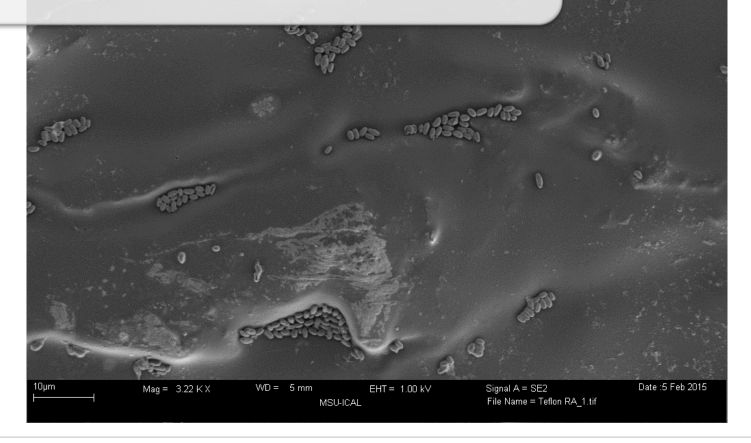
Bad substrate for decontamination





Teflon

The surface appears smooth, but spores can aggregate in some grooves

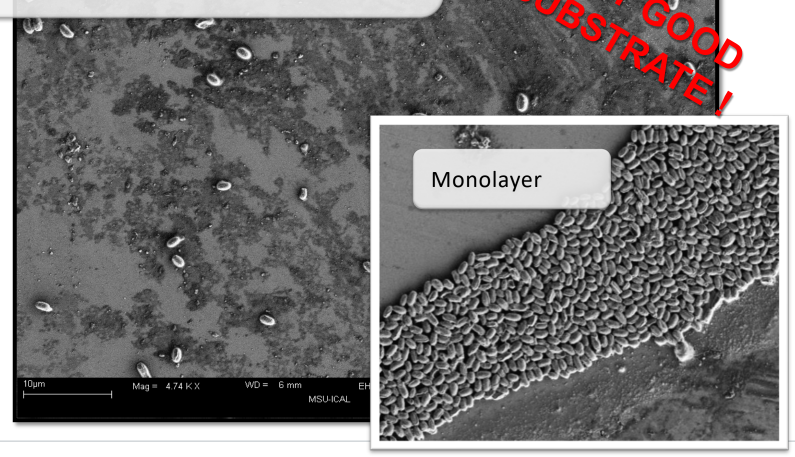






Glass

The spores are well dispersed on the very smooth surface







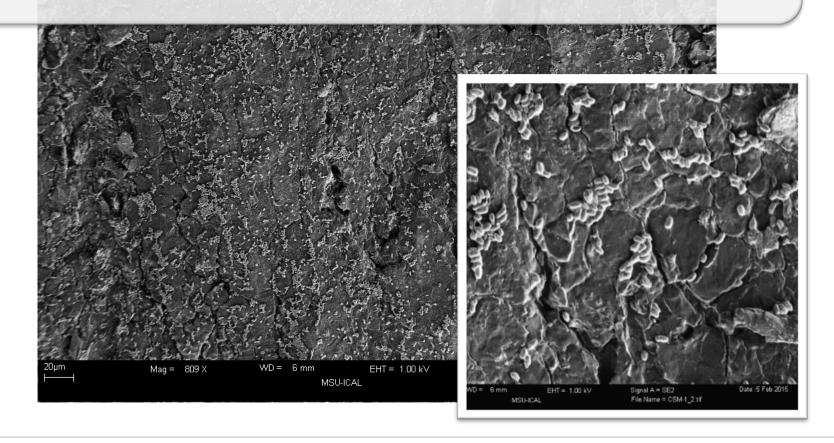






CSM (Hypalon™)

The spores are well dispersed, but they show slight clumping in some areas (relatively smooth surface)







Tyvek®

The spores are not visible, it is likely that they fell into cavities. However H_2O_2 vapors can easily penetrate through it.







Hydrogen peroxide detection







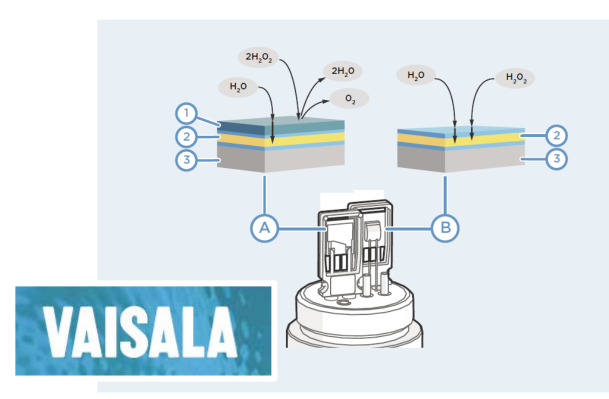
Electrochemical sensor

Measuring electrode: $H_2O_2 \longrightarrow O_2 + 2 H_+ + 2 e_-$ Counter electrode: $1/2O_2 + 2 H_+ + 2 e_- \longrightarrow H_2O$





Hydrogen peroxide detection



Operating principle of PEROXCAP measurement

- A HUMICAP sensor with a catalytic layer (under the probe filter). This sensor only senses water vapor.
- B HUMICAP sensor without a catalytic layer (under the probe filter). This sensor senses the air mixture with both hydrogen peroxide vapor and water vapor.
- 1 Catalytic protection layer over the thin-film polymer. This layer catalyzes hydrogen peroxide into water and oxygen and prevents it from entering the sensing polymer.
- 2 Thin-film polymer between two electrodes.
- 3 Alumina substrate.

The difference between the readings from these two sensors indicates the vapor concentration of H₂O₂.





Hydrogen peroxide detection



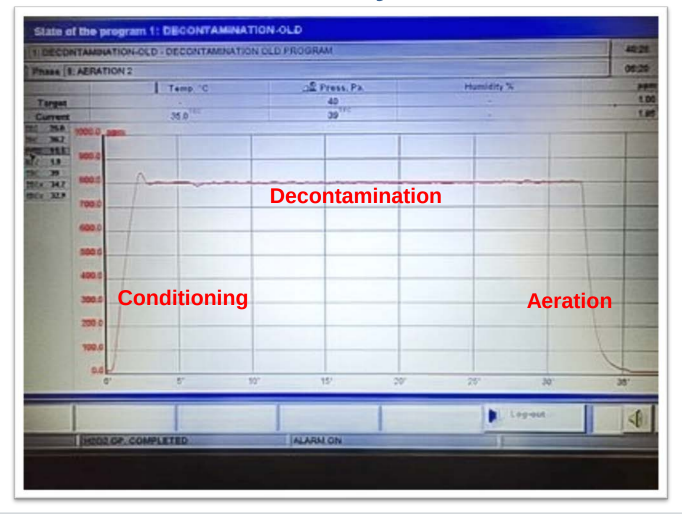
Cavity ring-down spectroscopy (CRDS)

Lower Detection Limit: < 3ppb





Decontamination cycle: its structure

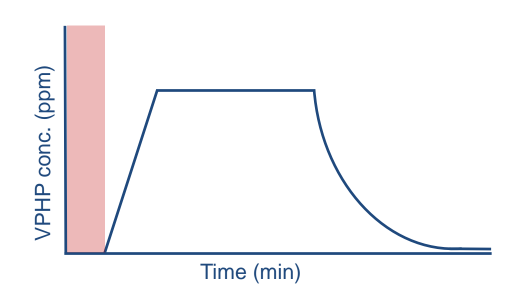






1. Preparation

✓ Achievement of the pre-defined temperature and relative humidity value (set point)

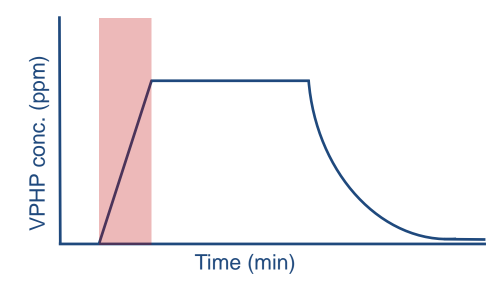






2. Conditioning

- ✓ VPHP injection at a high speed
- ✓ Achievement of the pre-defined VPHP concentration

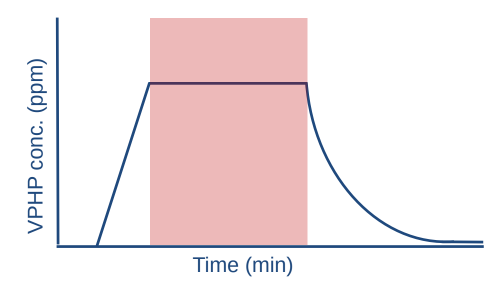






3. Decontamination (dwell time)

- ✓ VPHP injection at a reduced rate
- ✓ VPHP concentration is maintained constant for a pre-defined time

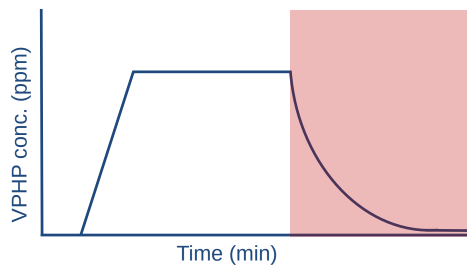






4. Aeration

- ✓ Air injection to replace (by dilution) H₂O₂
- ✓ H₂O₂< 1ppm (TLV/TWA, treshold limit value/time weighted average)
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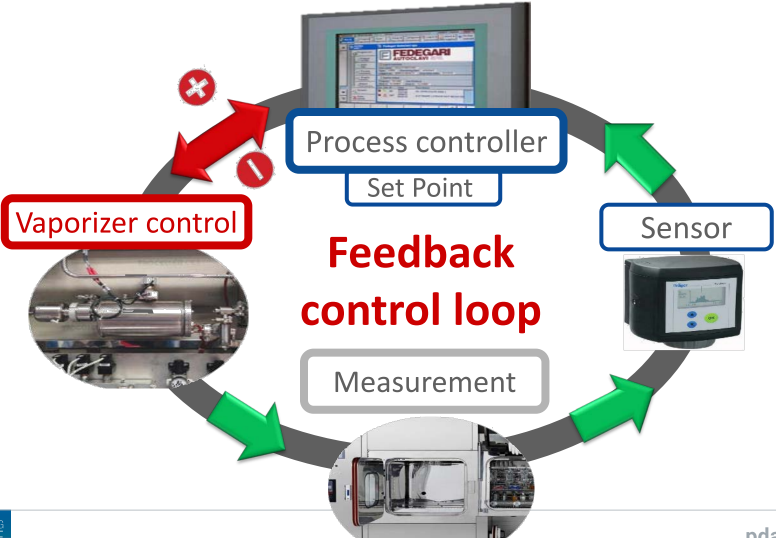


The time depends on both air exchange rate and H₂O₂ desorption from the decontaminated material (↑ temperature: ↑ v_{des})





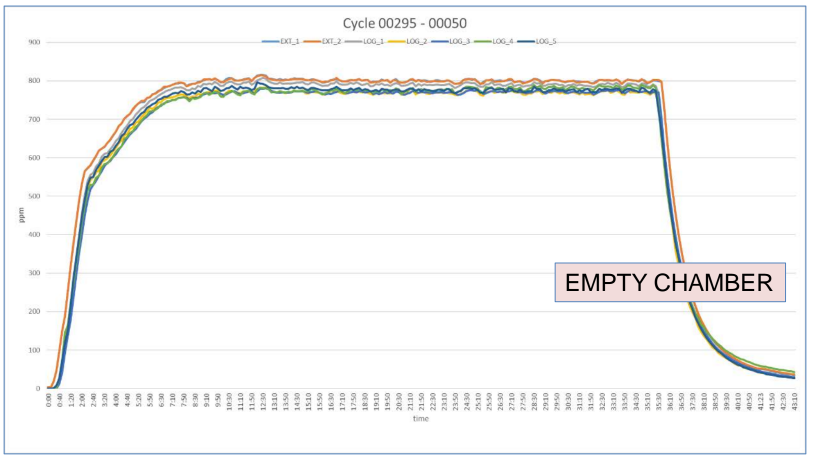
Biocide concentration: our approach







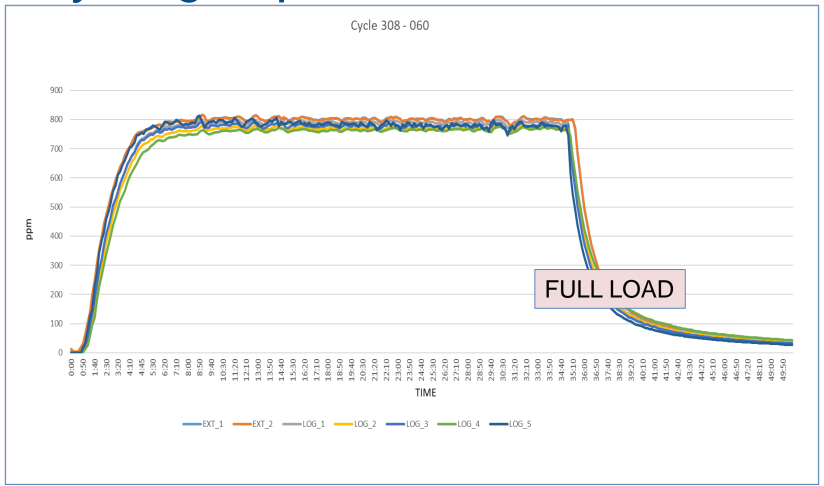
Hydrogen peroxide distribution







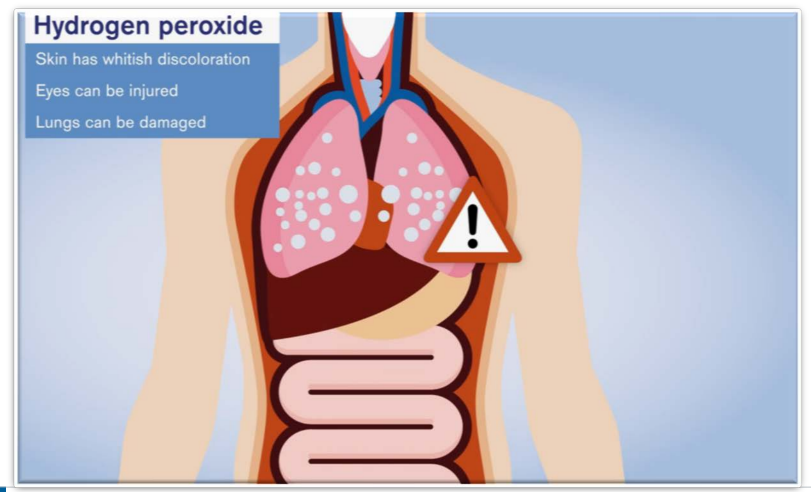
Hydrogen peroxide distribution







Health and Safety







Safety

1 ppm is the TLV, TWA declared by OSHA

TLVs® are not standards. They are guidelines designed for use by industrial hygienists in making decisions regarding safe levels of exposure to various chemical substances and physical agents found in the workplace.







Catalyzer







SLR achieved

Sporicidal process.

A gaseous, vapour or liquid treatment applied to surfaces, using an agent that is recognised as capable of killing bacterial and fungal spores. The process is normally validated using biological indicators containing bacterial spores. The number of spore log reductions is not specified in this definition, but a target of six log reductions is often applied. The process is applied to internal surfaces of the isolator and external surfaces of materials inside the isolator, when conventional sterilization methods are not required. The application of a sporicidal process to isolators is not considered to be a sterilization process in the same way as, for example, a sealed container subjected to a validated dry heat, moist heat or irradiation process.











SLR achieved



Cycle development starts with the definition of the required level of inactivation in terms of BIs. Sporicidal gassing cycles for critical areas used in aseptic processing are commonly validated to a minimum of 6-log reduction using biological indicators. Lower levels of log reduction may be acceptable in areas or on surfaces where risk of biocontamination transfer has been assessed as low.







Biological indicators

Process	Selected Organism	ATCC Derivation
Peracetic acid	Geobacillus stearothermophilus	7953 or 12980 (Ph. Eur.)
Hydrogen peroxide		
Ethylene oxide	Bacillus atrophaeus (formerly Bacillus subtilis var. niger)	9372 (Ph. Eur.)
Formaldehyde		
Peracetic acid		

For applications where the surface to be decontaminated is not in direct contact with the product, a BI with a population of $<10^6$ may be considered with a supporting rationale (40).

PDA, Technical report No.51



pda.org



Biological Indicators

Certificate of Analysis

Apex Biological Indicator (Reorder # HMV-091) for Gaseous Hydrogen Peroxide

Lot #: H0955

Manufacture: 2015 April 07 Expiration: 2016 January 31

Indicator: Geobacillus stearothermophilus 12980(1)

Mean population: 2.5 x 106 CFU per stainless steel carrier⁽²⁾

Storage conditions: 2 - 8°C; less than 50% RH; move to ambient conditions ≥ 1 hr

before use.

Shipping conditions: Ambient temperatures; cold pack and desiccant may be used to moderate conditions during shipping.

Resistance Characteristics:

D-value⁽³⁾: 1.0 minutes in 2mg/L gaseous H₂O₂

D-value is reproducible only when exposed and cultured under identical conditions used to obtain results reported here. MPN method used. Units are manufactured in compliance with Mesa Laboratory, Bozeman Manufacturing Facility's quality standards and ISO 11138-1 guidelines and all appropriate subsections.

Purity: No evidence of contaminants using standard plate count techniques.

Incubate at $55-60^{\circ}$ C for 7 days. The recommended growth medium is Soybean Casein Digest Medium (SCDM), Tryptic Soy Broth (TSB) or Mesa Releasat Medium (PM/100).







D-value determination

ISO 18742: Sterilization of health care products - Biological and chemical indicators - Test equipment

Resistometer

Test equipment designed to create defined combinations of the physical and/or chemical variables of a sterilization process.





D-value determination

USP 41, GENERAL CHAPTER <55> Bis - Resistance performance Tests

"...there is no standard process for the conduct of vapor phase hydrogen peroxide decontamination..."

"...there are no industry standard biological indicator evaluation methods for vapor hydrogen peroxide..."

... "it is more reasonable to consider resistance of biological indicators to be a relative or comparative measure from the manufacturer rather than a true D value."





Fedegari VPHP BIER









BI storage

- Refrigerate at 2÷8° C
- RH < 50% (insert a desiccant pouch inside the bag where they are kept)

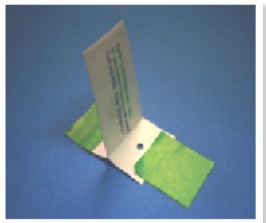


Move to ambient conditions ≥ 1h before use













Place the tape on the peel flaps, do not cover the spore location





- Do not use adhesive tapes or inks that absorb or catalyze hydrogen peroxide degradation
- Do not write on the spore location







Do not place the BI into or under a container







«Naked» BIs: spores are inoculated on a stainless steel ribbon not wrapped







Is the BI fault or our cycle is not a right one?







Re-run the cycle

Bls properly produced, stored and placed





 VPHP has a poor penetrating capability: it is a surface decontaminating agent



«...Quality control of Bls for sporicidal vapor-phase
 processes is imperative, since minor changes in the
 manufacture, storage, and presentation of the Bl may affect its
 sensitivity to the decontaminating agent...»

«PDA TR.51, «Biological Indicators for Gas and Vapor-Phase Decontamination Processes: Specification, Manufacture, Control and Use





RE-RUN A CYCLE WITH MULTIPLE BIS





Triplicate Bls at the «worst case»

locations allow to evaluate the situation with a statistical analysis





If we used **one Bl/location**, we might have:

BI (+)

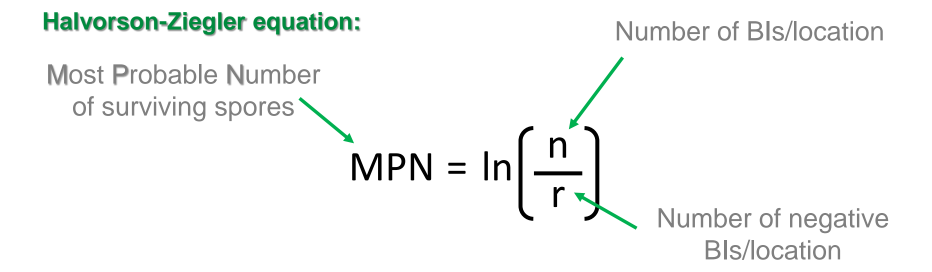
BI (-)

If we used **3 Bls/location**, we might have (+ + +) (- - -) (- - +) , (- + +)

Single BIs do not allow to perform a statistical analysis







- Applicable only with multiple Bls/location
- It allows to calculate the average number of surviving spores per BI





Example: after running a VPHP cycle we observed two positive and one negative BIs (++-) at a specific location

$$MPN = In (n/r)$$

n (number of Bls/location) = 3

r (negative BIs/location) = 1

MPN = In (3/1) = 1.099

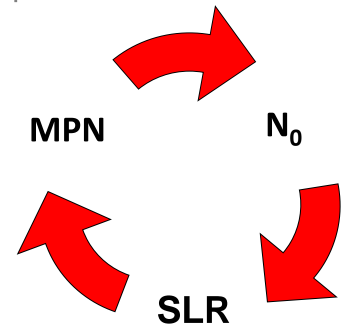


On average we have 1.099 survived spores per BIs





There is a link between MPN, the initial population of the used BIs (N_0) and the *Spore Log Reduction* (SLR) obtained at a specific location







Spore Log Reduction at the specific location where we observed Bls (++-):

$$SLR = Log_{10} N_0 - Log_{10} MPN$$

Example:

If spore population per BI = 2.8×10^6

$$Log_{10} 2.8 \times 10^6 = 6.447$$

If
$$(+ + -)$$
, MPN = 1.099

$$Log_{10} MPN = 0.041$$



SLR = 6.447 - 0.041 = 6.406





 Despite the growth of two BIs at that location, we can still prove that a 6 SLR was achieved at that specific test location

 This SLR value is what guidelines and/or rules require about decontamination

THE DECONTAMINATION CYCLE WAS
SUCCESSFUL!





 This calculation is ONLY possible when replicate Bls are used.



 If 100 Bls were placed at 100 different test locations, it would not be appropriate to perform this calculation as these
 100 individual Bls are not replicates of the others.

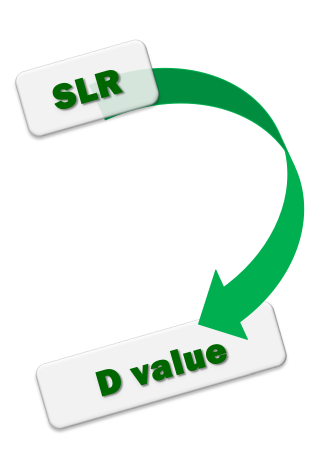




D value: time / SLR



Knowing D, how many SLR we have, we can add «x» minutes to reach a SAL 10⁻⁶







Before saying that your cycle has failed,

you should ask yourself...

- Was the BI correctly manipulated and stored?
- Is the BI not a good one («rogue» BI)?
- What is the microbiological result that I need (SLR)?
- Did we routinely observe multiple
 positive Bls at multiples locations?





Conclusions

- Hydrogen peroxide is a decontaminant, active on surfaces.
- Its process is a low temperature one, useful for heat sensitive loads.
- It migh provide a SAL 10⁻⁶ only on surfaces.
- Its validation includes considering:
- material compatibility,
- definition of the targets to achieve,
- assessing the homogeneity of distribution
- reaching the safety level required



Thank you for your attention

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