Aseptic handling of the MOMA Mass Spectrometer after Dry Heat Microbial Reduction

Contamination, Coatings, Materials, & Planetary Protection Workshop
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ExoMars 2020

Mars Organic Molecule Analyzer (MOMA):

- ExoMars 2020 - an ESA lander and rover:
  - Scheduled Launch Date: July 2020
  - Life detection mission
  - Samples will be collected up to 2m below the surface by a drill
- Mars Organic Molecule Analyzer (MOMA) is an instrument suite on the rover:
  - **Mass Spectrometer (MS) – NASA/GSFC**
  - Sample Ovens – MPS
  - Gas Chromatograph (GC) – LISA and LATMOS
  - Laser Desorption (LD) - LZH
MOMA-MS Hardware Bioburden Requirements

• **Sample path:** <0.03 spores/m²
  - **Accessible areas:**
    - Base of MS
    - Internal surface of pseudo-Ultra Clean Zone (pUCZ)
  - **Inaccessible areas:**
    - Internal surfaces of Mass Spectrometer (MS)
    - Internal surfaces of Wide Range Pump (WRP)
    - Internal surfaces of Gas Processing System (GPS)
    - Achievable only by dry heat microbial reduction (DHMR)

• **Surfaces not in contact with sample path:** 300-1000 spores/m²
  - Exterior of MS, pUCZ, WRP, GPS,
  - Internal and external surfaces of electronics boxes
  - Achievable by good contamination control practices and biocidal cleaning solutions

Mass Spectrometer model

Base of Mass Spectrometer

16 cm
Verifying MOMA-MS Bioburden Requirements

- Sample path (Ultra Clean Zone): $<0.03$ spores/m$^2$
  - Accessible areas:
    - Bioassay to 300 spores/m$^2$ at final access before 4 order of magnitude bioburden reduction with Dry Heat Microbial Reduction (DHMR)
  - Inaccessible areas:
    - Bioassay surfaces with similar handling, calculate bioburden reduction credit from (DHMR)

- Surfaces not in contact with sample path: $300-1000$ spores/m$^2$
  - Internal volumes of electronics boxes: Inspect and bioassay before final assembly
  - Exterior surfaces: Inspect and bioassay before shipment and delivery to ESA.
Pre-DHMR Handling

- Clean all hardware and tools before entering the cleanroom:
  - Multi step solvent cleaning process (detergent, water, acetone, hexane or toluene, isopropyl alcohol)
  - Inspect to VCHS+UV
  - Bake for dryness (GSE) or contamination bake out (flight hardware or critical GSE).
  - Contamination bakeout can double as bioburden reduction bakeout at the right times and temperatures.
- Clean (laundered) non-sterile garments (change 2x/week), with surgical masks and non-sterile gloves
- Handle hardware in an ISO class 7 clean room, mostly in an ISO class 5 tent or ISO 5 flow bench
- Reclean/reinspect before and after any major move (between cleanrooms)
DHMR: Dry Heat Microbial Reduction

- Standard approved method of bioburden reduction on flight hardware
  - Exposing hardware to temperatures of at least 110°C with controlled humidity
  - 4 orders of magnitude decrease in viable bioburden
  - Higher temperatures = shorter bake, but many components are not compatible with high temperatures
  - Encapsulated or mated surfaces take 10x longer than surface bioburden reduction
- Viking: DHMR entire lander
- Today, subcomponents are usually treated
- Alternates to DHMR have to be analyzed, proven, and approved by PPO
• Only sample path is included in DHMR
  • Electronic components and Laser are not compatible with DHMR temperatures

• 4 orders of magnitude reduction of viable microorganisms for surface contamination only.
  • 60 hours at 110 °C
  • A longer DHMR bake would be required to microbial reduce mated surfaces or microorganisms encapsulated in bulk materials

• All post-DHMR sample path exposure must be conducted using aseptic operations
Post DHMR Overview

• Maintain hardware in biologically controlled (but not sterile) cleanroom
  • Increase cleaning frequency, add biocidal solvents, increase bioburden monitoring
• Non sample-path portions of the hardware do not need special handling post DHMR except for increased bioburden control in the cleanroom
• Minimize sample path exposure as much as possible
  • Implement aseptic handling when sample path must be exposed
  • Environmental monitoring for bioburden and particulate during sample path exposure
Activities that Require Aseptic Protocols

- Any exposure of sample path surfaces post-DHMR
- Sample path surfaces include:
  - Gas processing system
  - Mass Spectrometer interior
    - WRP
    - Laser window
  - Pseudo-Ultra Clean Zone (pUCZ)) with sample carousel
    - Aperture Valve interface with Mass Spectrometer (MS)

- Planned Aseptic Activities
  - Connections/disconnections at the gas processing system
  - Move MS from pUCZ to vibration plate
  - Move MS from vibration plate to pUCZ
  - Replace laser window
Aseptic Procedure Overview

• Prepare:
  • Prepare personnel through aseptic handling training
  • Prepare tools by sterilization or isolation of non-sterilizable components
  • Prepare the cleanroom by cleaning with biocidal solutions and bioburden assays

• Monitor during the activity
  • Monitor movement of tools, personnel, and parts with special attention to contact transfer and air flow direction
  • Monitor workspace for bioburden during the activity
  • Three days for bioburden results from activity monitoring (not the hardware itself)
  • If aseptic conditions (<1 CFU measured) are compromised then microbial reduction must be repeated
Prepare: Aseptic Handling Training

- Half day training in addition to full day PP training
- Review expected aseptic activities
- Sterile gowning overview with demo of sterile gloves and single use sterile garments
- Tool preparation: sterilization or isolation
- Tool handling: careful attention to sterile and non sterile surface, contact transfer awareness, multi person handling to ensure no contact between sterile and non sterile surfaces
- Sterile tools may only touch sterile surfaces (gloves, hardware, work surface)
- If sterility is compromised through contact transfer, change out whatever was contacted (gloves, garment, tools, sterile work field, etc.)
Kimtech Pure® A5 Sterile Cleanroom Apparel Gowning Procedure

Before Gowning

Step 1
(Pre-Entry) Don Hair Net and Shoe Covers after removing all jewelry and cosmetics.

Step 2
(Gowning) Wash hands and gown first pair of sterile gloves. Sanitize gloves after gowning each article thereafter.

Step 3
Apply mask and hood ensuring snug fit.

Step 4
Open vacuum-packed apparel. Tear at notched edge.

Step 5
Grasp the blue line. Located on the inside middle back.

Step 6
Gently unfold coverall utilizing blue indicator line. Arms and legs are pre-drawn and snapped in place. Garment is already folded inside-out and unzipped.

Begin Gowning

Step 7
Hold garment at waist.

Step 8
Put one leg in and pull over through opening until snap releases.

Step 9
Do the same with the other leg.

Step 10
Insert one arm and extend until snap releases.

Step 11
Do the same with the other arm.

Step 12
Slip thumb through thumb loops.

Final Step
Cross legs and snap coverall.

After A5 Sterile Cleanroom Apparel is gowned
• Add boot covers.
• Complete gowning by adding goggles and a second pair of sterile gloves.

For more information, contact your local Kimberly-Clark Professional® Sales Representative at 1-800-255-6401 or visit www.kimtech.com.
Kimtech Pure® G3
Sterile Sterling® Nitrile Gloves Donning Procedure

Before starting the donning procedure, wash hands thoroughly and dry.

**Step 1**
Peel open sterile pouch and unfold glove wallet (DO NOT touch the exterior surface of gloves). Pinch the sides of wallet to open.

**Step 2**
Apply first glove to hand by sliding palm up into glove (thumb facing outward). Bend thumb toward center of palm and slide into glove while pulling up on the cuff. Leave the cuff rolled up.

**Step 3**
Apply second glove to hand by sliding the four gloved fingers into cuff of the second glove. Slide ungloved palm (thumb facing outward) into glove. Bend thumb toward center of palm and slide into glove while pulling up with fingers of gloved hand.

**Step 4**
Complete donning the gloves by pulling up the cuff of the first glove with the fingers of the second hand.
Prepare: Sterile Tools

- Tools compatible with sterilization:
  - DHMR: 60 min, 165 °C
  - Autoclave sterilization: 20 min 121°C, 100 kPa (15psi)
- Tools not compatible with sterilization
  - Not be used in direct contact with sample path surfaces post DHMR
  - May be used during aseptic process if non sterile components are wrapped in sterile foil or handled by non-sterile operator
- Prepare extra tools in case sterility is compromised
Preparing an aseptic work space

• Four days prior to aseptic procedure:
  • Mop with 7% hydrogen peroxide, then 70% IPA
  • Wipe horizontal and vertical surfaces in work area with sterile 70% IPA
  • Sample work surfaces and air for bioburden using ATP (rapid test for risk reduction) and CFU (3 day test, gold standard)

• Day of aseptic procedure
  • Confirm low bioburden from earlier surface samples
  • Clean again as above
  • Check work surfaces for bioburden using ATP
  • Ensure sterile garmenting/gloves are stocked
  • Prepare aseptic monitoring kit
  • Establish a sterile work field with sterile foil as a place to set tools or parts
Selected for biocidal action without leaving a residue

Use multiple cleaning solutions with different biocidal mechanisms to prevent selecting for resistant organisms

70% Isopropyl Alcohol (IPA) / 30% deionized water
  - Denatures proteins and damages cell membrane
  - 70% IPA is a more effective biocide than 100% IPA
  - Effective against vegetative microorganisms, but not spores. Some effectiveness against spores in mechanical removal.

7.5% $\text{H}_2\text{O}_2$ in deionized water
  - Disinfests by oxygen radical damage to DNA and proteins
  - Effective against spores at long exposure times
ATP rapid Bioburden Assessment

- Pre-wet swab is used to sample a surface, swished in the reactant buffer
  - ATP is the energy carrying molecule in all cell types
  - ATP in the sample will react with the luciferase and luciferin in the buffer and produce light
  - Less than 5 minutes to sample
- Pre-wet swab contains *Chlorhexidine digluconate*
  - Used to sample environmental surfaces or GSE
  - Not to be used on sensitive hardware
  - Residue removed by 70% IPA wiping
Most cleanroom and hardware samples do not have any CFU
- 99% of environmental microorganisms do not grow in a laboratory setting
- <15% of cleanroom samples had CFU within 72h
- RLU and CFU does not directly correlate in environmental samples

### Laboratory Experiment

<table>
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<tr>
<th>RLU Range</th>
<th># Samples</th>
<th># with CFU</th>
<th>% Positive</th>
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<tr>
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Processed according to ESA protocol ECSS-Q-ST-70-55C

- Spore specific or general viable microbe screen (with or without heat shock)
- Swab a 25cm² area on work surface with a damp swab
- Plated onto multiple pre-poured agar plates
- 72 hour incubation to count colonies (reported CFU/m²)

Bacterial colonies growing on a plate

ECSS-Q-ST-70-55C, D.2.1

dry swab → 1 ml H₂O → 25 cm² surface → 2.5 ml PBS + Tween 80

moistening → swabbing → resuspension

transport + storage → heat shock → 80°C 15'

at 4-8°C ≤ 6h

spread-plating → incubation at 32°C

0.5 ml → 0.5 ml → 0.5 ml

R2A

incubation after 24h, 48h and 72h

colony counting

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Ultraviolet Light to Sanitize work surfaces

- Ultraviolet-C (UV-C 100-290nm), 250-260nm is germicidal
  - Kills by crosslinking DNA, which prevents the organisms from faithfully replicating its DNA
  - Line of sight limited
- 22,000 $\mu$Ws/cm$^2$ is a sufficient energy dose to kill 99% of most common bacteria and bacterial spores on an exposed surface
- UV-C lamps (253nm) to be implemented in clean tent if bioburden levels are high
- UV-C intensity at work surface to be measured to determine appropriate exposure time

(NASA/David Herring/JPL)
Monitoring aseptic processing

- Dedicated Contamination Control personell to monitor:
  - Sterile gowning and gowning
  - Any contact between sterile gloves and non sterile surface (change gloves!)
  - Handling and passage of sterile tools
  - Tracking sterile and non sterile GSE/tool components
- Establish sterile work field with sterile foil
- Active and passive bioburden air sampling during the duration of sample path exposure
- Active air sampling with particle counter to ensure ISO class 5 is maintained
Aseptic Process Sterile Tool Handling

- Only be exposed to aseptic ISO Class 5 or cleaner conditions
- Must be handled wearing sterile gloves and garments
- Sterile pouches or foil opened by an assistant who is not handling sterile items
- Sheets of foil will be sterilized to establish sterile working surfaces
  - single use and only for the continuous working session
- Tools only be set on sterile surfaces
Airborne microbial monitoring

- **Active monitoring:** pulling air through a filter which is later transferred to a plate
  - Almost no growth seen in weekly cleanroom samples (20 min, 1 m³)
  - Most aseptic activities should be <20 min exposure time

- **Passive monitoring:** Allowing airborne microbes to settle onto a plate surface
  - Standard: Use agar plate to capture settling microorganisms. Not used in GSFC cleanrooms because of high volatile content of plates
  - Alternate: Use dry gelatin filters as fallout witnesses to transfer to an agar plate.
Summary

• As part of a life detection Mars Rover mission, MOMA-MS planetary protection requirements necessitate DHMR to reach 0.03 spores/m²
• After DHMR, any sample path exposure must be handled in an aseptic ISO class 5 work spaces
  • Sample path exposure is kept to a minimum, but is unavoidable
  • Aseptic ISO class 5 work space will be established by cleaning with multiple biocidal solutions confirmed by bioburden monitoring
• All personnel in the clean room during sample path exposure activities must be trained in sterile gowning practices and aseptic handling
• During sample path exposure, sterile gowning, sterile gloves, sterile tools, and aseptic handling will be implemented
  • Contamination Control personnel will monitor for non-sterile contacts
  • Active monitoring for bioburden during sample path exposure
  • Compromise in aseptic handling can result in high cost to schedule and risk to hardware.

Credit: NASA
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