

A close-up, angled view of a metal incubator tray with a grid of circular wells. The tray is shown in a perspective view, with the top edge of the tray and the grid of wells visible. The background is a solid green color.

CELL CULTURE INCUBATORS CLEANING AND DECONTAMINATION METHODS

Basic care for optimal performance and care in case of contamination. Includes tips to minimize contamination, proper laboratory techniques and information about active vs. passive decontamination.

Models:

MCO-233AICUVHX-PE
MCO-233AIC-series
MCO-173AICUVHX-PE
MCO-173AIC-series
MCO-171AICD-series
MCO-170AC-series
MCO-170M-series
MCO-50AIC-series
MCO-50M-series



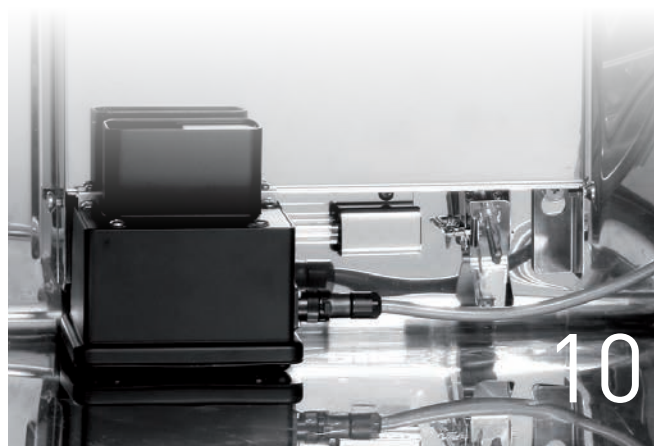
Cell culture incubators Cleaning and Decontamination Methods

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Introduction to Cell Culture Incubators



Cleaning and Decontamination Methods



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Introduction to cell culture incubators

A cell culture incubator is designed to artificially replicate in vitro conditions essential to in vivo physiology typical of human and animal models. Cell growth outside of a natural environment presents a multitude of challenges associated with exposure to microorganisms that are not present in the in vivo state. Depending on the type of cell cultures being managed, several operating parameters must be carefully controlled with accuracy, repeatability and flexibility in setpoint choices. These include temperature and gas control.

- Cell culture incubators are designed to establish and maintain a controlled, stable environment by regulating temperature at a typical setpoint of 37°C or over a range from ambient to points above 37°C.
- Incubator gases typically include CO₂ and/or O₂.

CO₂ is controlled at a precise setpoint to maintain desired pH in the cell culture media, whether liquid or gel. The CO₂ concentration in the incubator functions as a critical pH buffer.

Some biological materials may require different pH levels.

Desired CO₂ setpoint concentrations may differ. Most media contain an indicator which helps detect the change in pH.

- Optimal cell culture environments must include humidification to prevent desiccation of cell culture media.

While some incubators have internal humidification systems with heated water reservoirs, most incubators include simplified, removable humidifying trays designed to hold sterile distilled water which evaporates to naturally increase the relative humidity within the chamber.

The MCO-233AICUVHX and MCO-173AICUVHX feature an integrated active humidity system that comprises a humidifying unit capable of regulating the chamber humidity to 80–90% by means of a heating and evaporation system, and a humidity sensor. Combining this with a conventional system using natural evaporation of humidifying water further improves the stability of the culture environment.

If the chamber humidity decreases, humidifying water is atomized by heating and evaporation at 140°C in the humidifying unit to rapidly restore the humidity to 90%.

TYPES OF CELL CULTURE CONTAMINATION

Contamination of a cell culture in vitro is usually caused by the inadvertent introduction of one or more organisms that can damage or destroy the cell culture in progress. These organisms include:

- Bacteria (including Thermophilic Bacteria) and Mycoplasma
- Molds and Yeasts
- Viruses

Other contaminants include dust, volatile organic compounds (VOC's) from adjacent instrumentation or processes, cross contaminants from other cultures in a shared incubator environment and particulates found in the natural environment. Regardless of the contaminant or its source, prudent laboratory techniques can help avoid the recurrence of contamination.



THE INCUBATOR BUBBLE

Unlike closed systems, such as hollow-fiber substrates, stirred tank or airlift bioreactors, the typical cell culture incubator is a conditioned chamber with a door that closes against a soft gasket. When the door is closed, the incubator creates an ideal environment for the cell culture process based on user defined setpoint parameters for temperature, CO₂ and O₂. Humidification is naturally evaporative from the humidifying tray, and an elevated relative humidity is sufficient to eliminate desiccation, especially in microplates with small media volumes. Some larger cell culture incubators use immersion heaters to supplement the natural humidification process.

When the incubator door is opened, however, the conditioned bubble is lost. Accessing cell culture labware for transport to a biological safety cabinet (BSC) or other processes is a normal part of laboratory workflow. Opening the door exposes the incubator interior walls, shelves, humidifying water tray and culture vessels to ambient conditions that carry the potential for contamination from molds, yeasts, fungi or other microorganisms such as mycoplasma and viruses. In a practical sense, unless the incubator is installed in a clean room, this exposure cannot be avoided. Proper technique can reduce the potential. The first consideration is to understand basic incubator systems and how they can harbor contamination.

INCUBATOR DESIGN PREREQUISITES

The first step in managing cell culture contamination is to consider the incubator design, the interior in particular. Components exposed to the interior's high humidity atmosphere must be manufactured using materials with superior environmental performance such as high quality stainless steel. Components must also be easily removable (preferably using no tools) for autoclaving or manual cleaning. Components include: shelves, shelf brackets, plenums, floors, humidifying trays, blower wheels, sensor housings, innerdoor gaskets, humidifying tray covers, fan covers, duct covers, inner doors' inner sides and any other

item in the chamber during cell culture. Control probes are often protected by stainless steel sheath housings. These must be cleaned according to manufacturer instructions.

Components manufactured from copper enriched or copper supplemented stainless steel contain an inherent germicidal property that resists airborne organisms introduced to the chamber during door openings. Such materials are considered "passive" contamination control insofar as the inability of organisms to sustain growth on these surfaces.



Important points for installation

There are many factors to consider when determining the permanent location of the cell culture incubator. It is desirable to locate the unit where there is minimal foot traffic and where air disturbance is of little consequence. This reduces the volatility of outside air entering the incubator during a door opening. Avoid installing the incubator near windows, air conditioners, ceiling or floor HVAC air diffusers and return air intakes, all of which are sources of airborne contamination.

INSTALLATION, LOCATION AND CLEARANCES

It is important to consider the function of the biological safety cabinet when planning for incubator contamination mitigation.

If feasible, locate the incubator as close to the biological safety cabinet (BSC) as possible. This limits the exposure when removing or replacing cell cultures for processing.

Improper use of the BSC, wrong sash window height, blockage of downflow slots and use of instrumentation or equipment on the BSC work surface can create pathways for contaminants to attach to the cell culture labware when working in the hood. These contaminants are then returned to the incubator where they can migrate to other cultures via cross contamination or to interior surfaces exposed to a conditioned atmosphere ideal for cell growth.

Although the cell culture lab may normally be under positive pressure, this can change to neutral or even negative pressure when a BSC is operating, especially when the BSC has an exhaust transition connected to or over the exhaust filter.

Other laboratory equipment such as centrifuges, stirrers, shakers and robotic plate readers can aggravate an otherwise calm air environment to create aerosols that are easily airborne.

It is important to establish clearances adjacent to and behind the incubator because this space is required to provide easy access to gas supply tubing, tubing filters, gas input ports, pass-thru ports and blanking plugs and any interior components such as blower motors, fans or sensors that must be removed for maintenance.

Most CO₂ cylinders contain a CO₂ supply in liquid form wherein the CO₂ gas evaporates and moves through the two-stage pressure regulator as a gas. It exits the regulator at a pressure of approximately 4.4 PSIG - 14.5 PSIG (CO₂) / 7.3 PSIG - 14.5 PSIG (N₂/O₂), sufficient to prevent the introduction of contaminants into the gas system. The CO₂ itself, however, often contains microscopic particles that may provide surfaces for contaminants.



Locations where there is a lot of foot traffic are not suitable for the unit.

- Locate the incubator in a clean room or location where few people enter
- Choose a clean room that is safe or a place where there are as few people as possible.

Set up the unit as high off the floor as possible

- Since there are fewer airborne bacteria in the upper part of a room, the incubator should be placed on a laboratory table or a special stand.
- If stacking two or three units on top of each other, use a special roller base for that purpose.

Install the unit away from products that cause vibration and away from heat sources

- Do not install the incubator near products that cause vibration. Vibration may adversely affect cultures.
- Do not install the incubator near significant heat sources, such as heaters, boilers, ovens, or autoclaves. Heat will adversely affect the performance of the incubator.

Place in a location that is not directly affected by outside air

- Avoid putting the unit in a location that will be directly affected by air from a window, door, or air conditioning/heating vent.





Cleaning and decontamination methods

Most incubator manufacturers recommend a solution of ethanol and manual cleaning prior to initial start-up and regularly thereafter. The ethanol solution is intentionally diluted to give it time to kill the contaminant before the ethanol evaporates.

Why is an ethanol solution better than 100% ethanol in bacterial inhibition? 100% ethanol coagulates and dehydrates proteins so quickly that a layer of relatively impermeable denatured protein forms in the exterior parts of the bacterial cell (in and under the cell wall), and this prevents further diffusion of the alcohol into the cell. This protects the core of the cell from denaturation.

With an ethanol solution, the process is slower and the alcohol manages to diffuse throughout the cell denature proteins. In addition to conventional manual wipe down using ethanol, the incubator may be equipped with a sterilisation cycle such as a dry heat (180°C) system or a hydrogen peroxide vapor H₂O₂ system. The cycle should be performed prior to first use.

If commissioning and Current Good Manufacturing Practices (cGMP) criteria are in place, all contamination control efforts must be in compliance with previously approved best practices and facility protocol.

INSTRUMENTATION AND EQUIPMENT INTERFACE

Shakers, cell bottle rollers, magnetic stirrers and other devices are commonly used in the cell culture incubators. These must be free of contaminants before they are placed in the incubator.

Cell culture vessels usually include flasks with and without vent caps, petri dishes, roller bottles and multi-well plates. These are usually prepackaged and sterilised by gamma radiation prior to shipment. They should be opened only in a biosafety cabinet to preserve the integrity of the sterilisation.

Other labware returned from a central sterilisation room must be considered a source of contamination if exposed to ambient air during cart transit and shelf storage.

SUMMARY CONTAMINATION SOURCES

The following contamination points must be included in a regular schedule for cleaning in situ or removal and cleaning manually or by autoclaving.

INSIDE THE INCUBATOR

- Walls
- Ceiling
- Floor, shelves
- Chamber corners
- Ductwork, duct cover and plenums
- Humidifying tray
- UV light housing (if equipped) or UV-LED (for MCO-171AICUVD, MCO-173AICUV and MCO-233AICUV series)
- Temperature control probe and probe housing
- Probe wire to control panel
- Circulation fan and fan shaft
- Fan cover
- Humidifying tray cover
- Access port silicon cap
- Water-proof cap of the H₂O₂ generator connector

INCUBATOR CABINET

- Inner door gasket and feather surfaces
- Inner door latch
- Inner door glass
- Inner door hinges and fasteners
- Cool spots where condensation may accumulate due to insufficient cabinet insulation

GAS SYSTEM

- CO₂ or O₂ sensor
- Sensor housing and connectors
- Injection tubing from control solenoid(s)
- Air pump
- Filters and housings
- Fan, shaft and seal

MCO-233AICUVHX AND MCO-173AICUVHX

- Humidifying water tank and humidifying water supply tube

Basic care of cell culture incubators

Always puts on rubber gloves before cleaning the unit.

As a basic rule, do not clean the incubator with bare hands.
Be sure to use rubber gloves.

Necessary materials

- Rubber gloves
- Ethanol
- Gauze, non-woven cloth, or sterile paper

STEP 1

Turn off the power



STEP 2

Remove the interior components.

Remove the components in the correct order.

1. Remove the shelves.
2. Take out the humidifying tray and remove the humidifying tray cover.
3. Remove the connector cap*¹, duct cover, fan cover and back duct.
4. Remove the fan.
5. Remove the stopper for the access port.

For MCO-233AICUVHX and MCO-173AICUVHX remove the humidifying water tank from the holder. [See page 13 for more details]

*1) In the case of MCO-233AICUVHX, MCO-173AICUVHX, MCO-233AICUVH and MCO-173AICUVH with the optional H₂O₂ decon kit MCO-HK, or MCO-233AIC/MCO-233AICUV/MCO-173AIC/MCO-173AICUV with the optional H₂O₂ decon kit MCO-HK and electric lock MCO-170EL.



STEP 3

Clean the interior components

Use the correct cleaning procedure.

1. Wash with a neutral detergent (soap).
2. Rinse well with distilled water.
3. Wipe with gauze, non-woven cloth, or sterile paper.

For MCO-233AICUVHX and MCO-173AICUVHX

Clean the inside of the humidifying water tank using a brush. Wipe the tank caps, luer cap, tank tube, and the exterior of the tank using gauze moistened with a proper amount* of ethanol for disinfection.



*1) The amount that does not form droplets on the surface.

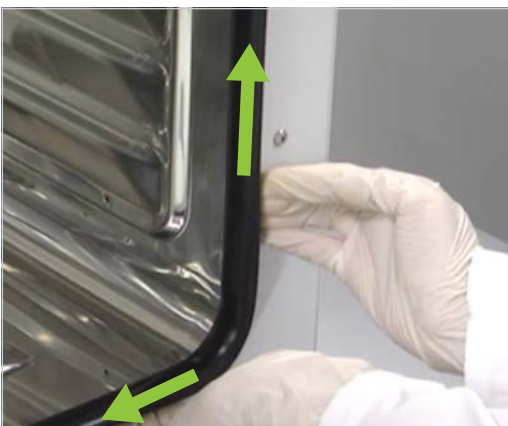
STEP 4

Spray disinfecting alcohol inside the unit and wipe (ethanol).

Do not spray ethanol into the CO₂ density measurement port directly! Just wipe off with gauze, non-woven cloth, or sterile paper which is sprayed with ethanol. Thoroughly wipe clean the inner walls of the chamber, the removed inner attachments, the temperature sensor, the humidity control bar, the UV-LED (Only for MCO-171AICUVD/ MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series) and the humidity sensor (for MCO-233AICUVHX and MCO-173AICUVHX) using gauze moistened with a proper amount of ethanol for disinfection. Make sure to take particular care in cleaning corners and joints of the product.



Picture 1



Picture 2

STEP 5

Disinfect all interior surfaces, internal components, shelves and water tray with ethanol.

STEP 6

Thoroughly wipe clean the inner door, inner door gasket, handle, and inner door glass. Take particular care to spread the ethanol for disinfection to all corners of the inner door gasket and sufficiently remove stains while wiping it clean. (picture 1)

If it is operated with the inner door gasket not in place, the humidified air will leak and will cause condensation between the unit and outer door. After wiping, confirm that the inner door gasket is securely in place and doesn't have any creases.

Adjust shape of inner door gasket after wiping. (picture 2)

Adjust the shape of inner door gasket by sliding fingers from each corner in direction of arrows. Specifically, insert fingers behind fin of inner door gasket and slide.

The inner door gasket has an important role to maintain the chamber humidity. If it is operated with the inner door gasket not in place, the humidified air will leak and will cause condensation between the unit and outer door. After wiping, confirm that the inner door gasket is securely in place and doesn't have any creases. If the inner door gasket is not in place, please refer to the back side and adjust the shape of the inner door gasket.



Basic care of cell culture incubators

STEP 7

Replace the interior components.

Replace the components in the reverse order of [STEP 2], and put sterilised distilled water in the humidifying tray.

1. For MCO-233AICUVHX and MCO-173AICUVHX, replace the humidifying water tank. [See page 13 for more details]
2. Replace the stopper for the access port.
3. Replace the fan, check to see if the fan spins smoothly by turning it with your hand.
4. Replace the back duct, duct cover and fan cover.
5. Put back the connector cap and the humidifying tray cover and replace the humidifying tray.
6. Replace the shelves.



STEP 8

Let it dry with the door ajar.

Before turning the power back on (restarting) let the inside dry out, check that there is no alcohol smell remaining.

If you turn the power on while it is still damp inside, the O_2 and CO_2 sensors may be damaged.



If using a unit that is equipped with heat sterilisation function or H_2O_2 (hydrogen peroxide) decontamination function, performing sterilisation/decontamination before use will make prevention of contamination (bacterial contamination) more effective.

inCu-saFe® CO_2 Incubator with heat sterilisation:
MCO-170AICD-series

inCu-saFe® CO_2 / Multi-Gas Incubators with H_2O_2 decontamination:

MCO-233AICUVHX, MCO-233AIC series (optional)*¹,
MCO-173AICUVHX, MCO-173AIC series (optional)*¹,
MCO-50AIC series (optional), MCO-50M (optional),
MCO-170M series (optional)



*1) Standard for MCO-233AICUVH and MCO-173AICUVH.

REMOVING THE HUMIDIFYING WATER TANK FOR MCO-233AICUVHX AND MCO-173AICUVHX

1. On the control panel, long tap the tank holder unlock button on the right side of the Home screen.
2. Fully pull the humidifying tank holder toward you.
3. Remove the humidifying water supply tube connected to the top part of the tank cap on the front side by rotating the luer fitting counterclockwise.
4. Hold the top part of the water tank and tilt it a little, and then remove the water tank from the tank holder.
5. Close the humidifying tank holder.



PLACING THE HUMIDIFYING WATER TANK FOR MCO-233AICUVHX AND MCO-173AICUVHX

1. On the control panel, long tap the tank holder unlock button on the right side of the Home screen.
2. Fully pull the humidifying tank holder toward you.
3. Replace the components in the reverse order of "Removing the humidifying water tank".
4. Connect the humidifying water supply tube to the tank cap.
5. Close the humidifying tank holder.

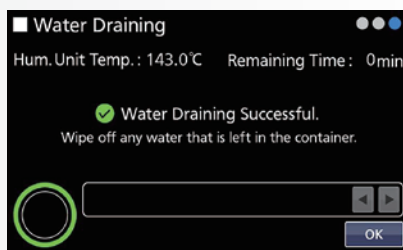


DRAINING THE WATER IN THE HUMIDIFYING WATER SUPPLY TUBE FOR MCO-233AICUVHX AND MCO-173AICUVHX

If the incubator will not be used for an extended period, please discard the humidifying water in the humidifying tray and tank and drain the water from the humidifying water supply tube in the humidifying unit. Then remove all moisture from inside the chamber, and when the inside is completely dry, close the door. Leaving the incubator unused for an extended period with high humidity inside the chamber can lead to malfunction.

1. On the Home screen, tap the Cleaning button.
2. Tap the Water Draining button.
3. Open the humidifying tank holder by long tapping the Tank Open button, remove the humidifying water tank from the holder, and close the holder.

Then, place the empty humidifying pan (referred to as "drainage tray" in the on-screen instructions) under the steam injection port, and close the outer door. Once the outer door is closed, tap the Start button. Be sure to close the humidifying tank holder and outer door after preparation for draining. If left open, the draining procedure will not start.



4. Wipe off any water left in the chamber and tap the OK button.

Basic care of cell culture incubators

CLEANING THE HUMIDIFYING UNIT WITH ETHANOL FOR DISINFECTION FOR MCO-233AICUVX/ MCO-233AICUVHX AND MCO-173AICUVX/MCO-173AICUVHX

Before cleaning the humidifying unit, prepare a 10 ml size (recommended) luer-lock tip type syringe and about 2 ml of ethanol for disinfection.

STEP 1 Tap the Cleaning button in the menu bar.
Tap the Hum. System Cleaning button.

STEP 2 Remove any contents from the chamber and install the empty humidifying pan (referred to as "drainage tray" in the on-screen instructions) under the steam injection port.

STEP 3 Long tap the Tank Open button to open the humidifying tank holder. After pouring sterile distilled water or pure water into the humidifying water tank, connect the humidifying water supply tube to the tank cap. Then, close the humidifying tank holder.



STEP 4 Tap the Start button.
The humidifying unit is automatically cooled down.

STEP 5 Long tap the Tank Open button to open the humidifying tank holder. Then, detach the humidifying water supply tube from the humidifying water tank and remove the tank from the holder. After that, fill the prepared syringe with ethanol for disinfection and connect it to the tube detached from the humidifying water tank. Then, tap the Start button.

Use a syringe with luer lock tip and a 10 ml capacity (recommended). Fill the syringe with about 2 ml of ethanol for disinfection.



Connect the syringe containing ethanol to the detached tube.

STEP 6 Tap the Yes button after checking the connection.

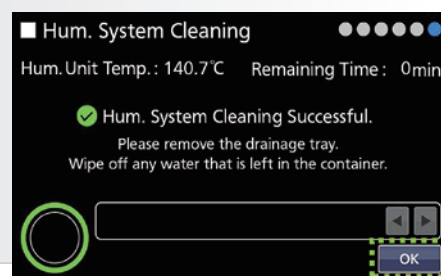
The ethanol for disinfection in the syringe is automatically moved to the humidifying water supply tube and humidifying block.

STEP 7 Remove the syringe from the humidifying water supply tube and place the humidifying water tank with sterile distilled water or pure water into the humidifying tank holder. Then, connect the tube to the humidifying water tank and tap the Start button.

STEP 8 Check that the humidifying water supply tube is connected to the tank and close the humidifying tank holder. Then, tap the Yes button.

The sterile distilled water or pure water in the humidifying water tank is supplied to the humidifying water supply tube and humidifying block to drain the ethanol for disinfection in them and clean them.

STEP 9 Remove the humidifying pan and wipe off any water left in the chamber. Then, tap the OK button.



CLEANING PRECAUTION

Get into the habit of wiping carefully

- Be sure to wear rubber gloves to avoid cutting your hands on the interior component
- Do not use acid, alkali, or chlorine-based cleansers, disinfectants, or sanitizers

IMPORTANT POINT

Do not wipe with the same portion of the sterile non-woven cloth more than once.

If you wipe another area with same portion of the cloth, you will be spreading bacteria around. Don't forget to wipe the gasket and the inside of the door.



CLEANING THE HUMIDIFYING WATER

When replacing the water, also clean the humidifying tray. Do this at least once every week.

- Lift the humidifying tray cover and pull out the humidifying tray.
- Wash the humidifying tray with neutral detergent, rinse it thoroughly with distilled water and wipe it off with gauze, non-woven cloth, or sterile paper.
- Spray ethanol to the humidifying tray and wipe off thoroughly.
- Wipe any moisture on the bottom of the chamber.
- Place the humidifying tray under the humidifying tray cover and pour sterile distilled water (preferably pre-heated to 37°C) into it.

IMPORTANT POINT

Do not use ultrapure water, tap water or reverse osmosis water as these are not suitable for incubators.

Please avoid adding any chemicals to the humidifying tray.



Tips for the incubator which is prone to be contaminated easily

TIPS TO MINIMIZE CONTAMINATION

- Increase the frequency of cleaning and humidifying water replacement.
- Spray Biocidal ZF to the inside of chamber once every week.

Replacement of humidifying water

Recommended frequency: Once every week (depending on frequency/ environment of use)

1. Lift the humidifying tray cover and pull out the humidifying tray.
2. Wash the humidifying tray with neutral detergent, rinse it thoroughly with distilled water and wipe it off with gauze, non-woven cloth, or sterile paper.
3. Wipe any moisture on the bottom of the chamber.
4. Spray ethanol to the humidifying tray and wipe off thoroughly.
5. Place the humidifying tray under the humidifying tray cover and pour sterile distilled water (preferably pre-heated to 37°C) into it.

IMPORTANT POINT

Do not replenish the humidifying water. The humidifying tray area is a major air way which is prone to collect dust and/or dirt that UV sterilisation can't remove.

The use of biodegradable Biocidal ZF spray-dis-infectant in incubators will help protect cultures against bacteria, fungi and enveloped viruses.

Non-volatile:

The active microbiocidal ingredients of Biocidal ZF are non-volatile. These protect cell cultures from microbial contamination and do not invade cell cultures via air. Thus cell cultures are protected against contamination and the disinfectant itself.



TIPS TO MINIMIZE THE RISK OF CONTAMINATION

- Locate the incubator in a clean room or a site where there are few people.
- Install the incubator some distance above floor level (The higher you go, the less floating bacteria are found). Use a roller base to facilitate cleaning around and under the incubators.
- Install the incubator in an area away from draughts and easy air intrusion when opening and closing the incubator doors. Beware of air dust and the air flow direction of any air conditioning.
- Ensure that there is no condensation inside chamber.
- Always keep the interior of an incubator clean and free from culture medium and/or water and fingerprints. They must be wiped off immediately if ever spilled or smeared (When there is a film or foreign matter foamed or placed on the surface of the copper alloy, sterilisation effect will be lost).
- Always maintain and handle culture vessels under the maximum aseptic conditions possible. It is recommended to wipe off the bottom and periphery of the culture vessels with ethanol for sterilisation when taking them into or out of an incubator.
- Minimize the door opening and closing frequency.
- Use sterile distilled water for the humidifying pan. Do not use ultrapure water, which may cause red rust-like particles in the humidifying tray. Change the water in the humidifying tray once a week.
- Take care when putting items that emit heat into the chamber. Putting items such as shakers which emit heat into the chamber may cause condensation and lead to possible contamination.

Care in the event of contamination

Get into the habit of wiping carefully

- Be sure to wear rubber gloves to avoid cutting your hands on the interior component
- Do not use acid, alkali, or chlorine-based cleansers, disinfectants, or sanitizers

Necessary materials

- Rubber gloves
- Ethanol
- Gauze, non-woven cloth, or sterile paper

24 HOUR UV DECONTAMINATION FOR MCO-170M SERIES (OPTIONAL), MCO-170AC SERIES (OPTIONAL), MCO-50AIC SERIES (OPTIONAL) AND MCO-50M (OPTIONAL)



STEP 1

Turn off the power.

STEP 4

Spray ethanol inside the unit and wipe.

STEP 2

Remove the interior components.

Remove the components in the correct order.

1. Remove the shelves and the fan cover
2. Take out the humidifying tray and remove the humidifying tray cover.
3. Remove the back duct
4. Remove the fan
5. Remove the stopper for the access port
6. Remove the gas injection nozzle and the gas injection nozzle tube (for multi-gas incubators).

STEP 5

Disinfect all interior surfaces, internal components, shelves and water tray with ethanol.



STEP 3

Clean the interior components.

Use the correct cleaning procedure.

1. Wash with a neutral detergent (soap)
2. Rinse well with distilled water
3. Wipe with a gauze, non-woven cloth, or sterile paper

SEE PAGE 10/11 FOR MORE DETAILS ABOUT STEP 1 TO 5

Care in the event of contamination

STEP 6

Turn on the power and (activate the UV lamp for 24 hours)*

Once the interior components and UV lamp cover have been removed, carry out UV sterilisation for 24 hours. It is not necessary to wipe the interior with ethanol afterwards.

STEP 8

Let it dry with the door ajar.

Before turning the power back on (restarting) let the inside dry out, check that there is no alcohol smell remaining.

If you turn the power on while it is still damp inside, the O₂ and CO₂ sensors may be damaged.

STEP 7

Replace the interior components.

Replace the components in the reverse order of [STEP 2], and put sterilised distilled water in the humidifying tray.

Before replacing all the components, check to see if the fan spins smoothly by turning it with your hand.

*** Models equipped with the 24 hour UV decontamination system:**

PHCbi Multigas Incubator with optional SafeCell UV™ Lamp:

MCO-170M series, MCO-170AC series, MCO-50AIC series and MCO-50M



NB: MCO-171AICUVD, MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series do not include the 24 hour UV decontamination system.

ACTIVATION UV LAMP 24H FOR MCO-170M (OPTIONAL)



Lighting the UV lamp for 24 hours

If the chamber has been contaminated by dirt or by spilling the medium, use the following procedure to decontaminate it by lighting the UV lamp for 24 hours.

1. Remove all attachments from the chamber, including the trays, the fan cover, the duct, the fan, the humidifying tray, and the humidifying tray cover. Disinfect all the attachments in an autoclave or with ethanol.
2. Clean and wipe down the inside of the chamber with ethanol.
3. Set the CO₂ density to 0% and the O₂ density to 20%. Go to the Stand-by-Setting screen (Menu > Set), enter 0% at CO₂ density and 20% at the O₂ density. Press "Apply" to save the entered values.

4. Go to the Tools #1 screen (Menu > Tools#1). Press "UV Setting" to display the UV Setting screen.

5. Turn the UV 24h Mode to ON and press "Apply".

6. The UV lamp will now light continuously for 24 hours. "UV 24h Mode ON" is displayed on the UV lamp condition display.

Notes:

- The UV 24-hour mode may activate the automatic set temperature alarm because of rising chamber temperature.
- If the outer door is opened when the UV lamp is lit, the UV lamp is turned OFF and UV 24-hour mode is cancelled. Repeat procedures 4 to 6 to restart the UV 24-hour mode.

7. After 24 hours, the UV lamp turns OFF automatically. Reinstall all the attachments removed in procedure 1.



Care in the event of contamination

ACTIVATION UV LAMP 24H FOR MCO-170AC, MCO-50AIC AND MCO-50M SERIES (OPTIONAL)

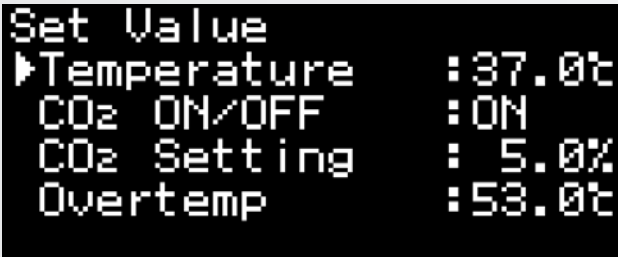
Lighting the UV lamp for 24 hours

If the chamber has been contaminated by dirt or by spilling the medium, use the following procedure to decontaminate it by lighting the UV lamp for 24 hours.

1. Remove all internal items from the chamber (racks, fan cover, duct, fan, humidifying tray and humidifying tray cover).
Disinfect all the attachments in an autoclave or with ethanol.

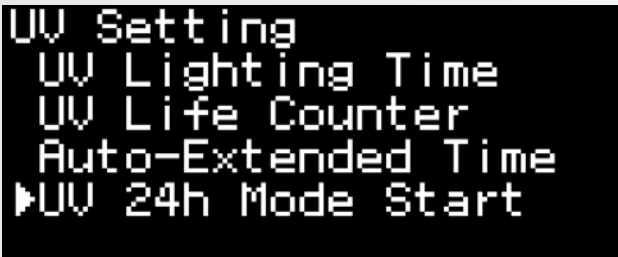
2. Clean and wipe down the inside of the chamber with ethanol.

3. Set the CO₂ density to 0 %.



```
Set Value
▶ Temperature      : 37.0°C
CO2 ON/OFF        : ON
CO2 Setting       : 5.0%
Overtemp          : 53.0°C
```

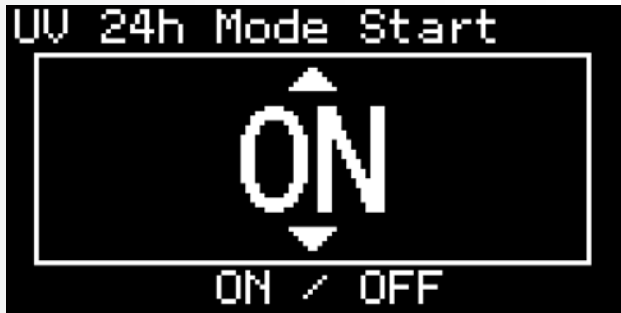
4. On the home screen press MENU. The left side of the display will change to Menu screen.
 - Move the cursor to Alarms & Controls by using the up/down keys and press ENTER.
 - Move the cursor to UV Setting by using the up/down keys and press ENTER.



```
UV Setting
UV Lighting Time
UV Life Counter
Auto-Extended Time
▶ UV 24h Mode Start
```

5. Move the cursor to UV 24h Mode Start by using the up/down keys and press ENTER.
The right side of the display will change to the UV 24h Mode Start setting screen, the current setting (OFF) will be displayed.

6. Use the up/down keys to change the UV 24h lighting mode setting to ON and press ENTER.



```
UV 24h Mode Start
  ON
ON / OFF
```

7. Press the MENU key to display the home screen.
The UV lamp will now light continuously for 24 hours.

Notes:

- The UV 24-hour mode may activate the automatic set temperature alarm because of rising chamber temperature.
- If the outer door is opened when the UV lamp is lit, the UV lamp is turned OFF and UV 24-hour mode is cancelled. Redo procedures from 4 to restart the UV 24-hour mode.

8. After 24 hours, the lamp UV turns OFF automatically.
Reinstall all the attachments removed in procedure 1.



H₂O₂ DECONTAMINATION FOR MCO-233AICUVX*¹/MCO-233AICUVHX, MCO-233AIC SERIES*², MCO-173AICUVX*¹/ MCO-173AICUVHX, MCO-173AIC SERIES*² AND MCO-170M SERIES*³



No need to remove the UV lamp/UV-LED and inner parts. It doesn't use a heater, so it conserves energy.

STEP 1

Preparation Time:
10 - 15 minutes



1. Remove all interior components
2. Wipe down the inside of the incubator
3. Reposition interior components to specified locations for in situ decontamination
4. For MCO-170M series: Install the H₂O₂ generator (MCO-HP).^{*4}
For MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series: Install the H₂O₂ generator (MCO-50HP).^{*4}



MCO-173AICUVHX decontamination layout

**Placement
H₂O₂ generator**

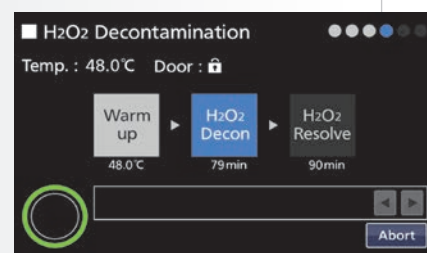
STEP 2

Decontamination Time:
Approx. 135 minutes



Internal decontamination can be completed by simple operation of the control panel

1. For MCO-170M series: On the Top screen, press the "H₂O₂" button for 3 seconds to open the Attachment Position Confirmation window. After confirming, press "Next."
For MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series: Tap the "Cleaning" button on the menu bar, then tap the "H₂O₂ Decontamination" button to open the Attachment Position Confirmation window. After confirming, press "Next."
2. The system check starts automatically, and if the system is normal the "OK" button is displayed. Press to start decontamination.
3. To improve the effects of decontamination, the chamber temperature increases to 45°C (48°C for the MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series).
4. H₂O₂ vapour generation starts
5. Interior fan circulates vapour
6. The H₂O₂ is reduced to water and oxygen.



STEP 3

Finish Time:
Approx. 10 minutes



1. Open the outer and inner doors. Disconnect the connector, cable, and H₂O₂ generator, and remove the H₂O₂ generator and cable from the chamber.
Note: Put on protective glasses and rubber gloves to do this.
2. Wipe off remaining liquid with sterile non-woven cloth
3. Reposition interior components to normal positions. Cover the connector on the H₂O₂ generator with the connector cap.

*1) MCO-233AICUVX and MCO-173AICUVX require H₂O₂ decon kit MCO-HKX-PW for H₂O₂ decontamination.

*2) Standard for MCO-233AICUVH and MCO-173AICUVH. MCO-233AIC, MCO-233AICUV, MCO-173AIC and MCO-173AICUV require H₂O₂ decon kit MCO-HK-PW and electric lock MCO-170EL for H₂O₂ decontamination.

*3) MCO-170M/170ML require H₂O₂ decontamination control board MCO-170HB UV system set MCO-170UVS and electric lock MCO-170EL for H₂O₂ decontamination.

*4) Optional extra. This process requires a dedicated H₂O₂ decontamination reagent (sold separately).

Care in the event of contamination

H₂O₂ DECONTAMINATION MCO-50AIC SERIES AND MCO-50M SERIES (OPTIONAL)

FOLLOW STEPS 1 TO 3 FROM PAGE 20

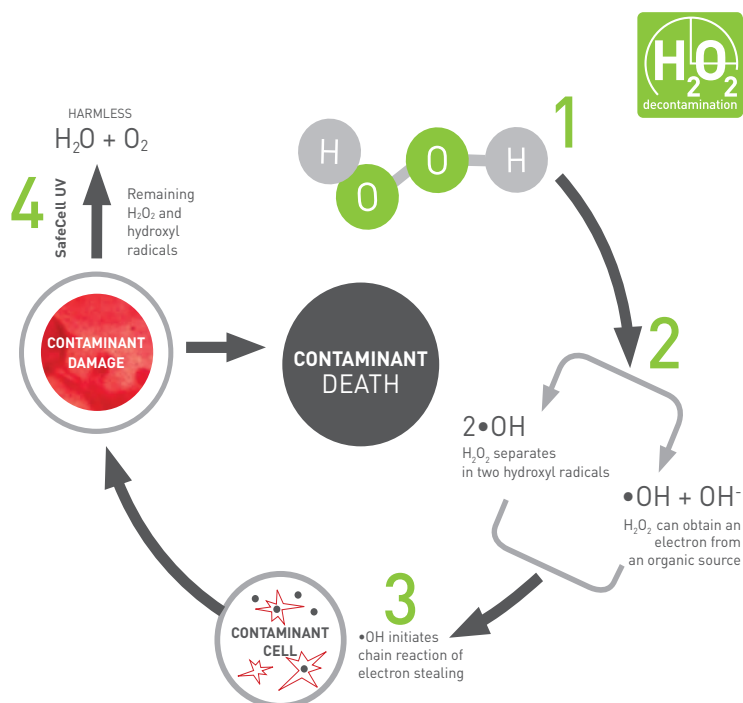
(Screen appearance may slightly vary depending on the model. Please refer to the user manual)



HOW DOES IT WORK?

1. Hydrogen peroxide (aqueous) is converted to vapour using high frequency ultrasonics. During this process, the fan motor remains active, ensuring H₂O₂ vapour accesses every point of the chamber and the tubing to and from, and the inside of the CO₂ sensor.
2. The H₂O₂ vapour breaks down into hydroxyl radicals naturally.
3. The hydroxyl radicals initiate a chain reaction of electron stealing.
4. This unstable internal environment leads to death of contaminants. Remaining hydroxyl radicals and H₂O₂ are resolved to H₂O (aqueous) & O₂ (gas).

PHCbi's H₂O₂ decontamination achieves at least a 6 log reduction of major contaminants. The full decontamination process takes less than three hours.



DNA is very susceptible to oxidative damage. Since most bacteria have a single chromosome controlling all their life functions, this kind of effect can be detrimental to their normal function. Prokaryotic organisms often lack repair mechanisms to limit such damage, making them more prone to change.

HEAT STERILISATION FOR MCO-171AICD SERIES CONVENIENT APPROX. 11 HRS. / 180°C HEAT STERILISATION



STEP 1

Preparation Time: 10 - 15 minutes

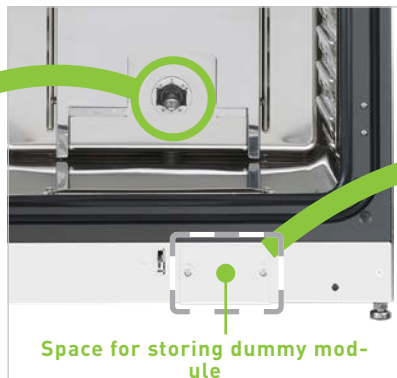


1. Take out the trays, humidifying tray, humidifying tray cover, fan cover, duct, duct cover, and fan from the chamber.
2. Dispose of the water in the humidifying pan, and wipe the inner attachments removed from the chamber with a gauze containing ethanol.
3. Wipe the inner walls of the chamber with a gauze containing ethanol.
4. Attach the dummy module (for the models with UV-LED).

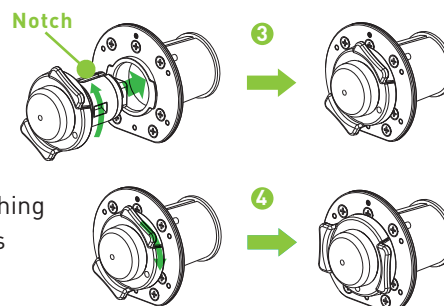
Attaching the dummy module

When performing dry heat sterilisation, the UV-LED module in the chamber needs to be exchanged with the dummy module to protect it from the heat.

- 1 Remove the UV-LED module in the chamber by holding the handles of the coupling ring, turning it counterclockwise 45°, and then pulling it hard.
- 2 Remove the dummy module stored in the space for storing dummy module at the bottom front of the incubator by pulling it out. Then, insert the UV-LED module (removed in step 1) into the space and store it.



- 3 Turn the coupling ring of the dummy module counterclockwise 45° so that the **notch** of it faces upward. Then, insert the dummy module into the place where the UV-LED module was placed so that the notch on the dummy module fits the projected part in the sleeve.
- 4 Turn the coupling ring of the dummy module clockwise 45° by pushing it to fix the dummy module. Then, check that the dummy module is securely fixed by pulling the coupling ring.



5. Attach the fan, duct, duct cover, and fan cover.
6. Set the trays on the 3rd, 5th, 6th, and 7th tray runners from the top.

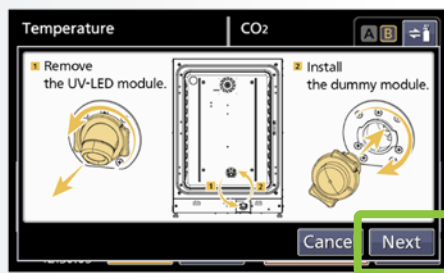
Care in the event of contamination

STEP 2

Sterilisation time: approx. 11 hours



1. On the Top screen, long tap the Sterilisation button.
2. Check that the dummy module is correctly inserted and fixed in the position shown in the screen and tap the Next button.
3. Check that the inner attachments are properly set by looking at the description on the screen. Then, tap the Next button.
4. In Sterilisation STEP 1, a system check is performed automatically.
5. In Sterilisation STEP 2, slide the slide button to the right, select "Yes" and tap the OK button.
6. In STEP 3 the chamber is warmed up to 180°C.
7. STEP 4 The chamber is kept at 180°C or higher for 60 minutes to sterilise the chamber.
8. STEP 5: The chamber is cooled down to 45°C.
9. STEP 6: Tap the OK button.



Both chambers can be in use at the same time even when double stacked.

One unit in heat sterilisation mode the other in incubation mode.

Heat leak reduction



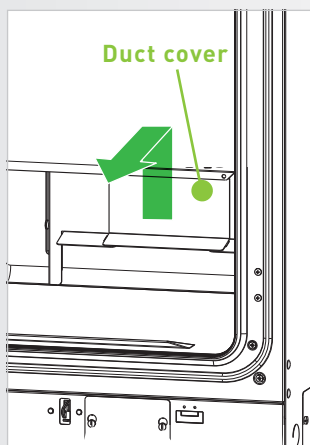
To prevent burning during the heat sterilisation cycle, the outer door is electronically locked. The top surface temperature of the MCO-171AICUVD/ MCO-171AICD during heat sterilisation is within the tolerance described in the International Safety Standard IEC61010 10.1 Surface temperature limits for the burn prevention.

STEP 3

Finish time: Approx. 10 minutes



1. Outer door is unlocked upon completion.
2. Open the outer and inner door
3. Take out the duct cover from the chamber by lifting it to disengage the protruded parts from the duct.
4. Then, put the UV-LED module (for the models with UV-LED) and dummy module in the original place. Attach the dummy module." after wiping the UV-LED module with a gauze containing ethanol.
5. Return the duct cover
6. Reposition interior components to normal positions.



UV-LED DISINFECTION FOR MCO-171AICUVD, MCO-233AICUVHX, MCO-233AIC SERIES*1, MCO-173AICUVHX AND MCO-173AIC SERIES*1



With the MCO-171AICUVD, MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series, the UV-LED can be set to turn on for sterilisation of the water in the humidifying tray after the outer door of the unit is closed. To ensure minimal effect on the temperature inside the chamber, the UV-LED cycles on and off automatically during the period when "UV Light Mode" is displayed. The length of time the UV-LED is on is calculated automatically from the chamber temperature setting, the UV Light Time selection (either Normal or Extended), and the UV Timer Ext. setting. The UV Setting allows the user to select UV-LED ON or OFF, and the light time duration when ON is selected.

MCO-171AICUVD:

1. On the Top screen, tap the "Menu" button.
2. On the "Menu" screen, tap the "Tools #1" button.
3. On the "Tools #1" screen, tap the "Settings" button.

MCO-233AICUVHX, MCO-233AIC series, MCO-173AICUVHX and MCO-173AIC series:

1. Tap the "Settings" button on the Menu Bar.
2. Tap the ► button or the center of the page indicator radio button.
3. Tap the "UV" button.

UV-LED ON/OFF Setting (UV Light Mode)

This sets whether or not the UV-LED comes on when the outer door is closed. When it is set to ON, the duration of the UV-LED light time is displayed in the box on the right. The default UV-LED ON time depends on the setting. Since the light time that is displayed is calculated

automatically, it cannot be set by the user. The UV-LED is set to ON or OFF by moving the slide button right or left.

4. Enter parameters in the UV Setting screen

Normal: For normal use.

Extended: For when higher sterilisation capability is required. This is selected by tapping the radio button. The duration of the UV-LED ON period is selected. The ON period is automatically calculated depending on the mode that has been selected and is displayed in the box next to "UV Light Mode".

5. Tap the "Apply" button.



CLEANING THE EXTERIOR, INTERIOR, AND INNER ATTACHMENTS

Clean the exterior and interior of the incubator and inner attachments once a month. Regular cleaning keeps the incubator in good condition. When the chamber is contaminated or when cleaning the chamber prior to starting a culture, you can perform dry heat sterilisation.

1. Take the humidifying pan out of the incubator, dispose of the water in it, and turn off the power to the incubator.
2. Take the inner attachments out of the chamber.
3. Clean the exterior, inner attachments, and chamber.

If the outside panels are dirty, clean them by wiping them with a gauze moistened with a diluted neutral detergent (using an undiluted solution of detergent may cause the unit's plastic areas to crack. Follow

the directions on the detergent for details of dilution). After that, be absolutely sure to wipe the surfaces using a cloth moistened with clean water to remove traces of the detergent and wipe the surfaces with a dry cloth. After that, wipe the surfaces with a cloth moistened with ethanol.

4. Put back the inner attachments to the original positions.
5. Open the doors for a while to dry the chamber.
6. Wipe the control panel with a dry cloth.

If the stain will not come out, wipe it using a cloth moistened with a proper amount of water (the amount that cannot form droplets) and then, wipe thoroughly with a dry cloth. If the water enters the control panel, it may cause the panel to malfunction. (Do not apply spray directly, it may cause damage.)

*1) MCO-233AIC and MCO-173AIC require UV system set (MCO-LUVS) for UV-LED disinfection. This item come standard on MCO-233AICUV/MCO-233AICUVH and MCO-173AICUV/MCO-173AICUVH.

Care in the event of rust

Always put on gloves before cleaning the unit.

As a basic rule, do not clean the incubator with bare hands.

Be sure to use rubber gloves.

Be careful as hands may get cut on the interior components.

Necessary materials

- Rubber gloves
- Ethanol
- Gauze, non-woven cloth, or sterile paper

FIRST FOLLOW STEPS 1 TO 5 FROM PAGE 9/10

STEP 6

Remove the rust with a cream cleanser.

Use a suitable amount of fine-grained cream cleanser, and carefully remove the rust.

STEP 7

Wipe with ethanol.



STEP 8

Activate the UV lamp for 24 hours.

*If it is a UV-lamp models

Once the interior component and UV lamp cover have been removed, carry out UV sterilisation for 24 hours. It is not necessary to wipe the interior with alcohol afterwards.

STEP 9

Replace the interior components.

Replace the components in the reverse order of [STEP 2], and put sterile distilled water in the humidifying tray.

Before replacing all the components, check to see if the fan spins smoothly by turning it with your hand.

KEEP IN MIND

These are the conditions that promote rust

- Insufficient wiping after the use of acid, alkali, or chlorine-based cleansers, disinfectants, or sanitizers
- Scratches on the surface of the unit interior or interior components
- The incubator is used with foreign matter left stuck to the inside of the unit or interior components
- Sodium dodecyl sulfate (SDS) is added to the humidifying water
- Using ultrapure water, deionized water or reverse osmosis water in the humidifying tray
- Adding chemicals to the humidifying water
- If an autoclave is used
If articles are stored wet in an enclosed place after coming out of an autoclave, rust can easily occur.
- If a heat steriliser is used
After coming out of a heat steriliser, once oxide scale appears on the surface that turns yellow or black, rust can easily occur.

Best practice and good laboratory technique

The most obvious approach to contamination-free incubator operation is to keep the incubator clean. A combination of manual cleaning and automatic (if equipped) decontamination processes managed on a regular schedule help protect cultures in situ and minimize loss of work due to contamination and downtime. Predictive maintenance is analogous to preventive maintenance, whereby cleaning processes can be documented for standardization and compliance, scheduled in advance and assigned to laboratory staff as required. There is no substitute for aseptic technique when handling cell cultures. Both personal and laboratory hygiene are essential to a holistic contamination management program.

ACTIVE VS. PASSIVE DECONTAMINATION

Active decontamination, whether by manual wipe down, dry heat sterilisation, H_2O_2 vapor or other method, must be initiated by the user. Design attributes inherent to a properly engineered cell culture incubator offer an additional layer of protection by working in the background to inhibit and destroy contaminants as they occur.

Active Decontamination



Dry Heat. A dry heat process utilizes time and temperature, typically 160°C to 170°C for a two-hour period, for a proven method of decontamination. The PHCbi brand, new Copper Enriched Stainless Steel (marketed as inCu-saFe® under the PHCbi brand) is a stainless steel and copper composite alloy that

forms a germicidal barrier to prevent thermal decontamination system operates at a higher temperature. It is the fastest and most effective active method of decontamination in a cell culture incubator reaching 180°C for a two-hour dwell before returning to ambient temperature. To minimize downtime, total cycle time is less than 12 hours. This energy-efficient process does not require the removal of the CO_2 sensor in the PHCbi brand incubator.



Hydrogen Peroxide (H_2O_2) Vapor. PHCbi brand incubators permit the use of active hydrogen peroxide (H_2O_2) vapor decontamination with complete safety and zero impact on the surrounding environment. Hydrogen peroxide starts in aqueous form and is converted to vapor using a nebulizer; this exposes all interior surfaces to the H_2O_2 vapor which ultimately resolves to water and oxygen at less than 1 ppm when catalyzed by a UV lamp.



Humidity regulation function by heat evaporation.

If the chamber humidity decreases due to door opening/closing or a restart, humidifying water is atomized by heating and evaporation at 140°C in the humidifying unit, which regulates the humidity to 80–90%, for as long as the humidity is reduced. By increasing the speed of restoration of the chamber humidity,*1 fluctuations in the culture environment are minimized and the culture is protected from drying out. Having an integrated humidity sensor unit reduces the risk of condensation formation and contamination from placing measurement or analytical equipment in the incubator. Regulation of the chamber humidity to 80–90% by the humidity sensor controls humidity increases and condensation formation from heat sources placed inside the incubator if the heater stops.

Passive Decontamination



Copper Enriched Stainless Steel (marketed as inCu-saFe® under the PHCbi brand) is a stainless steel and copper composite alloy that forms a germicidal barrier to prevent growth of organisms on surfaces. All interior surfaces, shelves and brackets are comprised of the inCu-saFe® composite. This material is a hybrid of Type 304 stainless steel. It is 100% corrosion-proof and will not corrode or discolor like conventional C100 copper surfaces.



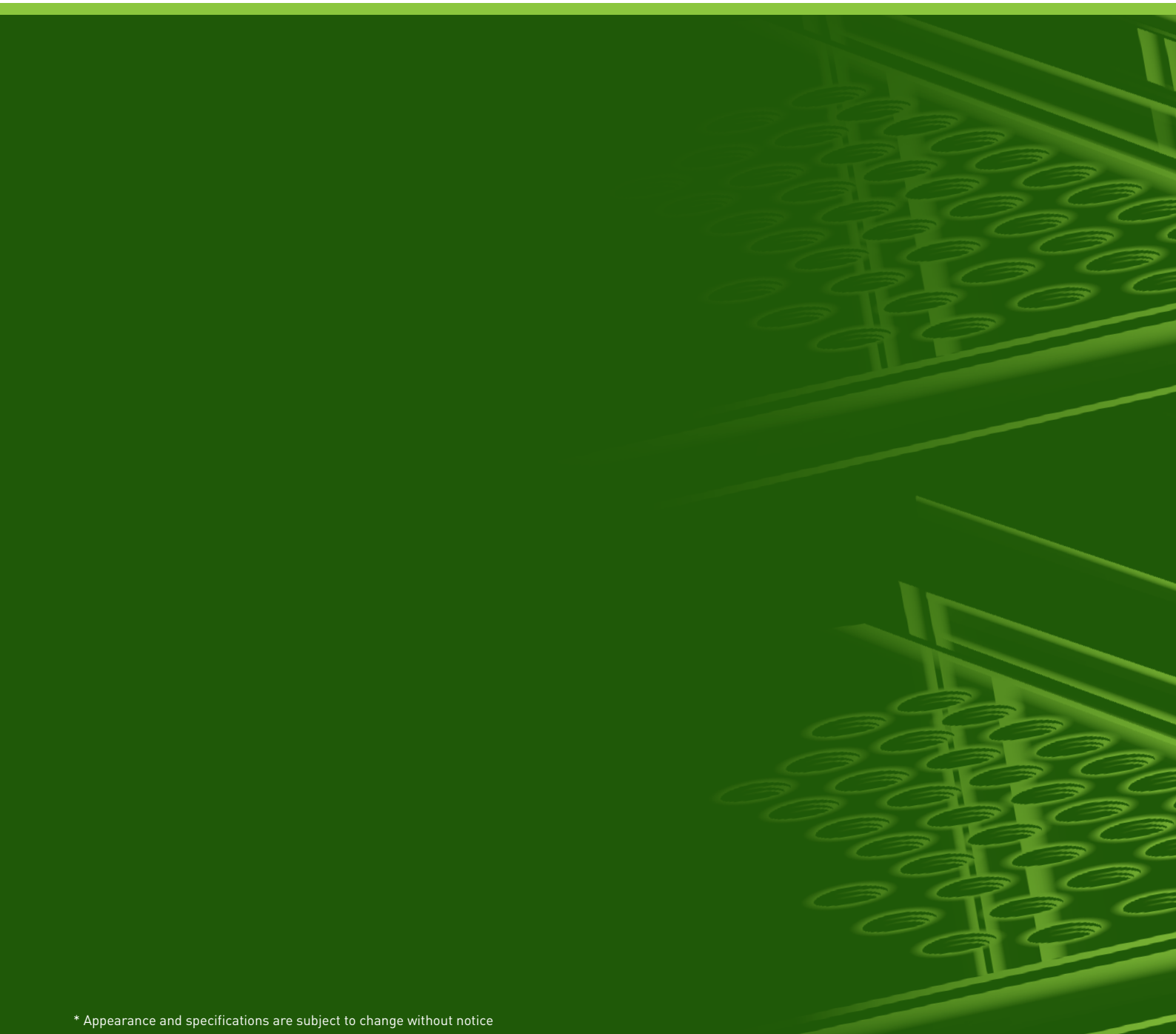
Ultraviolet Light (marketed as SafeCell UV™ under the PHCbi brand) is provided in either of two configurations: One is a UV lamp concealed behind the duct that creates a serial exposure of 257.3 nm wavelength. The other is a UV LED*, that shines UV light directly on the humidifying tray. The UV sterilisation system employs the UV-C (wavelengths between 100–280 nm) that provide, even for UV, very high effectiveness for destroying the DNA of any

organism passing through the airflow system as well as surface water contaminants in the removable humidity tray. The UV lamp/UV LED*2, automatically operates with a door opening/closing event. SafeCell UV™ inhibits the growth of mycoplasma, bacteria, molds, spores, viruses, yeasts and fungi without costly HEPA filter air scrubbers that accumulate contaminants in filter media. The UV lamp can also be programmed for a timed 100% ON cycle*3 for supplemental chamber decontamination.

*1) Restoration of humidity after the door is opened and closed is 50% faster than a conventional device (after the door is opened for 60 seconds and then closed, humidity is restored in 15 minutes.)

*2) For MCO-233AICUVHX, MCO-233AICUVH, MCO-233AICUV, MCO-233AIC (optional), MCO-173AICUVHX, MCO-173AICUVH, MCO-173AICUV, MCO-173AIC (optional), MCO-171AICD (optional), MCO-171AICUV

*3) For the MCO-50AIC and MCO-50M series.



* Appearance and specifications are subject to change without notice



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