# PRODUCTS AND APPLICATIONS

## General Requirements

Infection control and prevention requires a disruption of the transmission of a pathogen. Understanding the different routes of disease transmission is imperative for an informed, multi-layered approach to mitigate the health hazard. Transmission of pathogens can be by different routes: direct contact, fomite, aerosol (airborne), oral (ingestion), or vector-borne.

Understanding of the pathogen can help air carriers make an informed selection of disinfection products relevant to the pathogen and its transmission path while at the same time taking into consideration the unique environment of the aircraft. Just as important as selecting the correct product, the product or method chosen must not compromise, damage, or adversely affect the functionality of the aircraft structure, systems or components.

Appendix A lists the multiple documents used to develop this paper. Many of those documents used the terms “cleaning” and “disinfection” interchangeably. While this document’s focus is related to product selection and procedures relevant to disinfection, it is necessary to clarify the similarities and differences in the two important processes needed to break the pathogen transmission route.

### Pathogens

In the past, aviation has needed to respond to health events like Severe Acute Respiratory Syndrome (SARS), Swine Influenza A (H1N1), and Zika virus. The current threat pathogen, SARS-CoV-2, has been identified as either present on surfaces or in aerosol form with both chemical and non-chemical processes being utilized against the SARS-CoV-2 virus. Different agencies in the US and internationally have evaluated the toxicity and health implications of chemicals in other work places. Non-chemical approaches, including solutions currently in use and emerging solutions in development, vary in terms of description (equipment, processes, procedures) and documented efficacy.

Sections 3.2 and 3.3 provide more detailed information on both chemical and non-chemical approaches to combatting pathogens, including evaluating their effectiveness and use in the aircraft.

Specific pathogen characteristics are beyond the scope of this guidance and those data should be made available through agreed sources, such as the US CDC, WHO and other resources. Likewise, a specific disinfectant product or procedure related to the effectiveness against a pathogen, or the specific disinfection frequency is beyond the scope of this guidance. As noted above, an SMS process should be followed for evaluating both chemical and non-chemical processes and disinfection frequencies based on the information gathered from the multiple sources including the respective health organizations, aircraft manufacturers and others as appropriate.

### Cleaning

Cleaning is an important first step in the disinfection process. Cleaning as defined by IATA Ground Operation Manual (IGOM) and IATA notes that “Guidance for Aircraft cleaning and disinfection during and post pandemic", is the removal of visible dirt or particles through mechanical action. During a pandemic, general cleaning procedures are still a best practice, but may need to be conducted more frequently or the process adjusted. Cleaning and disinfection can be combined into one process if disinfectants are used during the cleaning. There are a number of considerations for an effective cleaning procedure:

* Cleaning personnel require proper training in correct cleaning procedures to safely clean without damaging the aircraft or harming themselves or others.
* Cleaning product selection should be subject to a safety risk assessment (SRA). Cleaning products and it should also be based on OEM recommendations, SAE standards and approved for the country of use (if applicable).
* Consideration should also be given to the methods appropriate for specific areas of the aircraft including the flight deck, galley, passenger cabin, lavatory, and cargo area. High-touch surfaces, such as door handles, armrests or interphones may require additional cleaning frequency or cleaning products as well as additional training.
* Appropriate personal protective equipment (PPE) for the environment and product used should be provided, and training on the proper use of PPE for cleaning staff should be provided.

### Disinfection

The purpose of disinfection is to eliminate, reduce or prevent the further spread of pathogens in the aircraft environment. The disinfection process builds upon the assumption of a previously cleaned surface. This document provides guidance on two methods of disinfection, specifically chemical and non-chemical.

Disinfection schedules, techniques, and products may be different at each aircraft operator considering the operational circumstances and the duration of the disinfecting effects of the substance used. The procedure should be updated when new information becomes available.

Routine disinfection procedures are in place to prevent the spread of diseases and are standard procedures which are routinely performed in addition to the cleaning process.

Aircraft operators should review and update their disinfection matrices based on specific configurations of their aircraft types, continue monitoring the high contact surface areas in aircraft, Based on the conducted risk assessment, each aircraft operator should implement policies and procedures for disinfection of aircraft.

Post-event (also known as event-driven) disinfection is performed after a specific event (e.g. after the transport of suspected or confirmed cases of communicable diseases onboard, spill of body fluids in the aircraft). This disinfection is not a frequent or standard practice and the requirements, methods, procedures as well as training for personnel will most likely differ. The World Health Organization (WHO) recommends that post-event disinfection procedures should meet the requirements under 3.2.4 and Annex F of Guide to Hygiene and Sanitation in Aviation, ICAO Annex 9 Chapter 2 (E) requirements. IATA Guidelines for aircraft cleaning and disinfection to manage affected aircraft carrying suspected communicable disease, also outlines general considerations. In case such event happens during the flight, inflight personnel may need to perform inflight disinfection.

It is important to ensure that the disinfection and other measures meet conditions required by applicable regulatory authorities to fulfil the following requirements:

* The disinfectants should be tested by a certified laboratory according to the specifications of the aircraft manufacturers for material compatibility tests, and not be corrosive or detrimental to aircraft components.
* The disinfectant should be applied according to the label instructions (e.g. concentration, method and contact time).
* Any contaminated items should be handled appropriately to mitigate the risk of transmission.

It is necessary to exercise caution in selecting disinfecting products or processes suitable for aircraft use. While being nationally approved for use, all disinfectants used should be aircraft-component compatible in that they must not have any negative effects on the individual parts or the structure of the aircraft. The products or processes should not have any negative effects on human health.

There are two types of disinfections:

* Chemical disinfection (e.g. wiping methods spraying, fogging…)
* Non-chemical disinfection (e.g. HEPA filters, UVC, ionization, ….)

Please see 3.2 for specific guidance for chemical products and 3.3 for non-chemical products.

#### Frequency of application of aircraft cleaning/disinfection substances or processes

The table in appendix B that follows was designed to assess the usefulness, effectiveness and practicality of a multitude of products and systems proposed to the airlines for reducing the spread of a pathogen causing viruses as transmitted by passengers and crew aboard commercial transport aircraft. One of the distinct challenges is assessment of the duration of effectiveness of the products or systems, as it drives the frequency of application. As an example, some liquid disinfection products are highly effective until the product dries and/or evaporates, after which the treated surface is susceptible to re-contamination when touched or otherwise brought in fresh contact with the virus. Similarly, it is known that ultraviolet light can be effective, but generally only where the light directly falls.

During an outbreak of disease (prior to effective vaccine or fully reliable pre-boarding screening), the criteria for selection of a product will be effectiveness of the product at the time of application, health risks associated with the substance or treatment, and what, if any, residual effectiveness remains following application. Re-contamination of treated surfaces can occur at any time, however some products have demonstrated a residual anti-microbial action.

Primary references on anti-viral effectiveness are guidelines and substance list established on the country level (e.g. the US Centers for Disease Control (CDC) guidelines and US Environmental Protection Agency substance lists). Regarding health risks of products, reference must be made to the manufacturers’ Safety Data Sheets (SDS), which are documents produced in alignment with the UN’s Globally Harmonized System (GHS) of Classification and labelling of chemicals.

Substance or process application could adversely affect or degrade any flight safety-related items or aircraft systems therefore it is imperative to identify any effect and mitigate and address them through maintenance tasks, processes, training and other mitigation actions. Substance or process application could adversely affect airworthiness of aircraft structure and systems. It is imperative to identify any adverse effects and adequately address them through the implementation of maintenance tasks, training and other mitigating actions, such as occasional deep cleaning to remove any residual product. Aircraft operators should employ Safety Management System (SMS) practices to assess the adverse effects on the airplane of their cleaning/disinfection programs

There is no one answer to cleaning/disinfection application intervals as that will completely depend upon the means employed and the expected level of interaction between the occupants and the aircraft cabin. Different cabin areas may require different disinfection frequencies, if exposed to different level of interaction with occupants, and depending on the level of separation or isolation with other cabin areas. For example, for aircraft where the flight crew compartment is not separated from the passenger cabin, the frequency of preventive disinfection of the flight crew compartment should be the same as for the passenger cabin. Aircraft Operators must assess the minimum application effectivity based on OEMs, disinfectant product and equipment, manufacturer’s data against more frequent application they may wish to publicize, designed to create good will and passenger confidence. There remains, however, a need to reliably evaluate application processes and products and mechanical systems using direct quality assessment. Measurement of effective application and confirmation of expected duration of effectiveness is not widely available and remains an outstanding challenge. Such means will be required to complete an SMS program.

Consideration for evaluation and frequency of application are listed in Appendix B.

The current threat pathogens have been identified as either present on surfaces or in aerosol form with aerosol believe to be the dominant path of transmission. Both chemical and non-chemical processes are being utilized against the virus causing COVID-19 and other pathogens. While chemical effectiveness has been evaluated, the non-chemical approaches listed herein, while in use, are still under varying degrees of evaluation and development, as some validations have been carried out and further evaluations are in process. Both chemical and non-chemical processes have new solutions expected that will need evaluation. The next two sections provide more detailed information on both chemical and non-chemical approaches to combatting pathogens, including evaluating their effectiveness and use on the aircraft.

The specific pathogen characteristics are beyond the scope of this guidance and those data should be made available through agreed sources, such as the CDC in USA, WHO and other resources as needed, available. As noted above, an SMS process should be followed for evaluating both chemical and non-chemical processes.

## Chemicals

### Selection and Approval Process

The selection of chemicals to be used to mitigate a pathogen must be found to be safe for the aircraft, crew, passengers and employees, including personnel performing the aircraft disinfection. The specific identification of the chemicals to be used is outside the scope of this document, however the best practice for the use and application is within scope. Section 3.2.2 through 3.2.10 will assist in the process of making the decision if the chemical choice will be safe for all parts and efficacious.

Air carriers should use their Safety Management System in the selection of their chemical disinfectant and application process. An integral part of that selection process is to utilize information from organizations like the US CDC, the WHO, and other public health organizations in a collaboarative approach to understanding the chemical's safety, effectiveness against a pathogen, and any health hazards of the chemical.

Certain agencies, like the US CDC or the European CDC (ECDC), have lists of disinfectant substances that can be efficient against certain pathogens. In the US, the Environmental Protection Agency (EPA) has published a list of products that are effective against COVID-19. Information from these organizations should be utilized in the selection process.

Given the nature of some chemicals, it is important to protect the health of the cleaning crew as well as other occupants with specific procedures for use of the chemicals. As example, ensuring that all occupants are off the aircraft before the chemical is used to ensure the other occupants are not incidentally exposed to the chemical.

During disinfection, any personnel not involved in conducting the actual disinfection should not enter the aircraft. If others need to enter the aircraft, they must also be protected to ensure they are not exposed to the disinfection chemicals. Sufficient time must be allowed for chemicals to dissipate prior to personnel, crew, or passengers entering the cabin. Ventilation of the aircraft may be necessary during the disinfection to reduce the build up of harmful aerosols in the aircraft, if recommended by the product manufacturer.

Coordination with the aircraft manufacturer or the component manufacturer is recommended when selecting a chemical and process for use in an aircraft to ensure the chemical does not have a negative effect upon the aircraft systems or structures, such as corrosion of metal components or degradation of any visual panels. Ensuring the safety of the aircraft components is critical to the selection process.

For purposes of this document, several distinct areas of concern were noted for their electronics and or furnishings. These are: passenger cabin, galleys, lavatories, crew rest areas, cargo compartments and the flight deck. Due to their unique features, an individual SRA for each section may be necessary.

Both Appendix B and C provide spreadsheets that can be used in the evaluation and selection process.

To ensure a continued effective response, after a chemical selection and application process has been selected it is imperative that there be a periodic SMS monitoring process, or feedback loop, to evaluate the chemical, the process, and any possible degradation of aircraft systems or components, or any health risks.

### Effects on Aircraft Interiors and Components

The US Environmental Protection Agency (EPA) List N contains hundreds of disinfectant products for use against SARS-CoV-2, additionally other national agencies have created lists of chemicals for use, however, it is imperative to recognize that not all of these products are appropriate for use on commercial airplanes. Commercial aircraft cabins, galleys, flight decks, and cargo compartments are made of parts with many different materials, some based on very specific safety requirements. Special attention should be made to consider the effects of the different chemical solutions for cleaning and disinfecting these areas of the aircraft on the different material parts. Given the many different materials, different chemical disinfectants may be needed for different parts of the aircraft.

Liquid chemical solutions can be of concern if not applied properly. Only trained personnel should be cleaning and disinfecting the airplane. Problems can arise if liquid chemicals are over applied, allowed to pool, seep into crevices, or soak into porous surfaces, such as airplane seat fabric. Some chemical disinfectants are only intended to be used on hard surfaces. Operators should carefully follow the chemical solution manufacturer’s recommendations for applications.

Frequent and repeated use of chemical solutions may cause undesired effects on some materials. These include material degradation, corrosion, discoloration, staining, or other visual defects on surfaces. Airlines performing multiple flights a day must be particularly attuned to long term impacts and accelerated deterioration of materials with frequent chemical disinfection.

Chemical disinfectants can also be highly flammable. Careful precaution should be taken to ensure these flammable disinfectants be kept away from heat, sparks, flames, and other sources of ignition and that proper ventilation of the aircraft is accomplished, if recommended by the product manufacturer. Operators should coordinate with the OEM for power on/off during the disinfection process, taking precautions to collar appropriate circuit breakers to avoid any unintentional re-application of power.

Operators should implement an inspection interval to ensure no detrimental effects have occurred from repeated use of chemical disinfectants. Depending on the aircraft part and use, operators should inspect for functionality, durability, material degradation, aesthetics, readability (for placards, etc.), and visibility (no scratches or crazing).

Specifically, for the flight deck, careful attention should be given to determine disinfectant compatibility with all parts and also the quantity of any liquid disinfectant used to disinfect. Given the critical flight controls located in the flight deck, operators need to avoid any over saturation, pooling, or moisture ingression on any surface. Also, ensure the flight deck is properly ventilated during the disinfection process.

### Chemical Makeup

In addition to the need to ensure any chemicals selected do not pose a safety hazard to the aircraft, a challenge all operators may face are the regulations around chemical disinfectants. Different countries, governing regulatory agencies, and local governments may each have separate requirements and restrictions for use of specific chemical disinfectants. Operators should understand and follow these requirements and restrictions for every destination. Some chemical disinfectants may not be able to be procured in certain parts of the world due to restrictions, high demand, etc.

Operators should follow the chemical manufacturer’s instructions for application of disinfectant product.

### Efficacy

The chemical disinfectant products used must be safe for the aircraft, crew, passengers, and employees while at the same time effective. Individual operators should use the Safety Management System (SMS) to determine safety The distinct areas of concern were noted earlier due to their electronics and furnishings. A safe and effective chemical disinfectant must be identified for each area. Each area will need a Safety Risk Assessment (SRA) to protect the aircraft, crew, and employees. In addition to the standard members of the SRA, Subject Matter Experts (SME) should be added to the SRA. SMEs will be the best source for determining which chemicals meet Original Equipment Manufacturer (OEM) recommendations, approved for the country of use, and most important, are safe and effective. Due to the dynamic nature of the environment and pathogens, the following SMEs should be added to the SRA.

* Original Equipment Manufacturer (OEM) to identify risks and threats (e.g. potential material or equipment damage) to the aircraft. In addition, make recommendations on products that are safe for the equipment.
* Industrial Hygienist to identify disinfectants that are approved and available in specific locales.
* A SME educated in microbiology or epidemiology for review of tests and efficacy.
* A dangerous goods SME related to any proposed shipping of the chemical disinfectants.

It is important to evaluate the disinfection process and safety of that process as part of the SRA feedback loop. The feedback should happen through testing and evaluation at set intervals as established by stakeholders and Subject Matter Experts (SME). The ongoing testing and evaluation of the four areas should be accomplished by the original SRA stakeholders and SMEs as noted above, that is the OEM, an industrial hygienist and a microbiology or epidemiologic SME.

For each of these areas the stakeholder and SMEs will evaluate the current chemical disinfectants used, the application process for those chemicals, and the effectiveness of those chemicals for the environment and current pathogens. The type and frequency of the testing should be determined and accomplished by the operator, regulator, third party, or a combination of those. This ongoing testing and evaluation process will monitor the effectivity of the disinfectant chemicals used in the four areas of the aircraft. The data from this testing and evaluation process will provide feedback for the SRA process.

### Application Locations, Methods, Phases of Flight

For purposes of this document, several distinct areas were noted with respect to their unique electronics or furnishings. These are: passenger cabin, galleys, lavatories, crew rest areas, cargo compartments and the flight deck. Given the uniqueness of each area, the air carrier will need to take into consideration the needs of each area in relation to the number of occupants, the use of the area, the ability to disinfect the area, and any electronics or furnishings that could be damaged by the chemical disinfection. Additional information is provided in section 3.2.2 related to possible adverse effects on aircraft interiors and components.

The passenger cabin is the largest area in relation to the disinfection process. This area can also present the biggest challenge in relation to disinfection because of the high-volume of passengers that could potentially infect another passenger or crewmembers during the boarding process, the inflight segment, and the deplaning segment. Passenger movement during the inflight segment can also add to the potential exposure to an infected person/surfaces. A best practice for efficient disinfection involves utilizing different cleaning tools for different areas of the aircraft cabin to limit cross-contamination. Likewise starting from the either the front or the back, top to bottom of the area can reduce cross-contamination.

Other cabin areas include the chemical disinfection of the lavatory, both inside and out as it can be a high-touch area. Other areas that air carriers should take into consideration are disinfecting any crew rest compartments, including any bedding and furnishings. Consideration related to health hazards such as crew allergic reactions to chemicals need to be tracked to ensure an additional health hazard is not induced.

Aircraft galleys can be considered high-touch areas, especially for crewmembers. A unique aspect of galleys is that the area is considered a food preparation area, so a chemical that may be acceptable for cleaning overhead bins may not be acceptable for cleaning a galley countertop surface. Any chemical utilized on that surface may need to be reviewed for appropriateness related to food service. Like other areas care should be taken to ensure any chemical chosen for the area takes into consideration the electronic items such as ovens coffee makers, and, the possibility of chemical spray entering any food service compartments especially if food is contained in the compartment. Concerns over possible food contamination may limit the chemical disinfection to pre- or post-flight application versus inflight application.

Cargo compartments, although not high touch environments, should be disinfected. Unique characteristics can include a limited size and limited ventilation, therefore proper application of chemical disinfectants should take into consideration those characteristics related to health risks. Use of flammable chemicals in poorly ventilated areas can also be a concern. Cargo compartments can contain unique safety design features such as smoke detectors, electronic door operation equipment and fire extinguisher discharge nozzles, so care should be taken to ensure that these systems are protected during the chemical application and ensure the safety benefits of these items is not compromised. As example, detectors resulting in spurious warnings because of mis-application. Cargo compartments are designed to only be utilized at the gate area during the boarding and deplaning process so any type of chemical used during flight is not applicable.

Flight decks, although separated from the passenger compartment, due to the frequency of crew transitions will require disinfection. This area of the aircraft is also the one that may require a broader list of chemical disinfectants because of the potential to damage the flight critical equipment. Careful evaluation of any chemical, including quantities, chemical types such as liquid (including spray) or wipes, need to be taken into consideration as they can each present a hazard if used incorrectly. The limited area of the flight deck can also present a hazard related to the time when disinfection occurs, specifically an inhalation hazard. Disinfection of the flight deck at the gate may permit adequate ventilation to minimize health risks. However, if disinfection of the flight deck is required during the flight, the disinfection process may need to be revised to take into consideration any inhalation concerns related to the selected chemical. The SRA process should be used to evaluate any additional hazards that may result from any proposed disinfection processes used during flight.

Air carriers should also take into consideration what type of personnel should be permitted to disinfect the flight deck to reduce the probability that aircraft switch settings are not disturbed during the disinfection process, including whether powering on/off aircraft systems is appropriate during the process. After the flight deck is disinfected the operator should verify the aircraft configuration to ensure all switches are in the correct positions, especially prior to turning back on if it was turned off. This is to avoid personal injury and damage to the aircraft.

### Effects on Humans

Two considerations should be made when discussing the effects of chemicals on humans: acute and chronic (repeated) exposure. The primary reference documentation for acute chemical exposure to humans is the Safety Data Sheet (SDS), which is sometimes referred to by the now obsolete term Material Safety Data Sheet (MSDS), as specified in the United Nations’ SDS standard called Globally Harmonized System of Classification and Labelling of Chemicals (GHS) revision 8 (2019). SDS provide critical information about the chemical's identity and ingredients, health and physical hazards, safe handling and storage procedures, emergency procedures and disposal considerations. Use of these documents is an essential starting point in the selection process especially for consideration of the chemical's effect on humans. When evaluating a chemical for possible use, the air carrier should consider any health hazards to the personnel actively applying the disinfectant, as well as any other persons that may be in the disinfected environment.

In the US, the two portions of the SDS paramount to the discussion on any effects on humans are Section 2, Hazard(s) identification, and Section 4, First Aid measures.

Section 2 Hazard Identification contains information on:

* + Acute toxicity
	+ Skin corrosion/irritation
	+ Serious eye damage/eye irritation
	+ Respiratory or skin sensitization
	+ Germ cell mutagenicity
	+ Carcinogenicity
	+ Reproductive toxicity
	+ Specific target organ toxicity-single exposure
	+ Specific target organ toxicity-repeated exposures
	+ Aspiration hazard

Section 4 First Aid contains:

* + Description of necessary measures, subdivided according to the different routes of exposure, i.e. inhalation, skin and eye contact and ingestion
	+ Most important symptoms/effects, acute and delayed
	+ Indication of immediate medical attention and special treatment needed, if necessary

The information located within the SDS is valuable for both emergency and chronic exposure details. Any employee or passenger should have access to the appropriate SDS of the chemical’s used within the airplane. Education and training will be required for the employees and crew on proper protocols and reporting of adverse reactions as defined by section 3.2.9 and 3.2.10.

While much is known about acute exposure to most disinfecting and cleaning material, little is known about prolonged chronic exposure experiences in humans. Although the GHS SDS does address specific target organ toxicity-repeated exposures, few large-scale studies have been done on repeated human exposures on most products. Likewise, as pointed out by the WHO:

*Safety of active ingredients for humans*: In spite of best practices in the decontamination of environmental surfaces, human exposure to microbiocidal chemicals cannot be prevented altogether; this is particularly the case in confined spaces such as aircraft cabins. Therefore, formulations with the safest possible ingredients must be selected for such use, including proper ventilation as recommended by the product manufacturer.2

Also, the WHO explains:

*Freedom from off-gassing and volatile organic chemicals (VOCs)*: Pungent odors are obviously undesirable, but addition of even strong scents/perfumes to disinfectants is now discouraged because of increasing numbers of individuals with multiple chemical allergies. Formulations that may release corrosive gases (e.g. chlorine) and VOCs must be avoided because of potential exposure of sensitive and vital components of the aircraft. Advice from the equipment manufacturer or aircraft operator’s engineering department should be followed. Appropriate ventilation during cleaning is also important2

Furthermore, EASA goes on to point out:

3.4 Any residual disinfection substances that may be harmful to humans should be removed from the seat covers or any other surfaces. This is essential especially when using cleaning and disinfection products which can cause skin irritation or harm.3

These identified risks can be minimized by providing a feedback loop into the operator's SMS system. Ongoing monitoring and reporting should feed back into the SMS feedback loop. This will provide protection for the employees, crew, and passengers. There should be a designated point of contact for the employees and crew in the event of exposure or adverse effects. These exposures must be tracked for the SMS process.

Lastly and best stated within the WHO International Health Regulations in Article 22, to the regulation authorities in each state, health measures “shall be carried out so as to avoid injury and as far as possible discomfort to persons, or damage to the environment in a way which impacts on public health, or damage to baggage, cargo, containers, conveyances, goods or postal parcels”1

Reference:

1. [WHO International Health Regulation](https://apps.who.int/iris/bitstream/handle/10665/43883/9789241580410_eng.pdf;jsessionid=B5C9DFA0BEF762DD189276D1267EB000?sequence=1)
2. [WHO Guide to Hygiene and Sanitation in Aviation](https://apps.who.int/iris/bitstream/handle/10665/44164/9789241547772_eng.pdf?sequence=1)
3. [EASA Guidance on aircraft cleaning and disinfection](https://www.easa.europa.eu/sites/default/files/dfu/EASA%20Guidance%20on%20aircraft%20cleaning%20and%20disinfection-issue%202.pdf)

### Frequency

The chemical disinfection frequency may vary from air carrier to air carrier based in part by their types of operation, their aircraft environments, and their destinations. Disinfection frequency is also dependent on the chemical chosen, the location of the disinfection, and operational circumstances. It is essential that air carriers conduct a safety risk assessment, coordinate with local health authorities and, possibly include the SMEs referenced in 3.2.4 to assess the chemical. It is important to note that any measures or frequencies utilized may need to be adjusted relative to any changes in the pathogen, or in relation to any regional conditions given the global nature of aviation. Frequency may also be determined by the location specifics, as example the passenger cabin, galleys, cargo compartments and the flight deck due to their unique electronics or furnishings in those various areas. Informed selections and correct use of products should support the determined frequency schedule.

### Relevant PPE use and limitations

Personal Protective Equipment (PPE) is worn to minimize exposure to workplace hazards. PPE may include, but is not limited to, items such as face coverings, gloves, safety glasses or shoes, respirators, coveralls or body suits. Chemicals in general can create a need for the use of PPE, making it more relevant to ensure protection of personnel assigned the task of apply chemical disinfectants in the aircraft environment. Air carriers or the contract service provider should ensure personnel who will be working with the selected chemicals, should be provided the appropriate PPE based on the chemical hazards. Although not recommended, if others need to enter the aircraft during the actual disinfection process, they must also be provided the appropriate PPE for the specific chemical hazard. Employees who shall be assigned roles for use of chemical must understand:

1. The purpose of each PPE item to be used
2. The use of correct PPE item(s) for each function specified
3. The proper selection, fit, and use, including donning and doffing as necessary, of each PPE item.

NOTE: it is recommended when appropriate;

1. Disposable gloves that are recommended by the manufacturer of the chemical should be worn.
2. Disposable gowns should be worn while disinfecting the cabin and lavatories.
3. If splashing is possible, eye protection, such as a face shield or goggles and facemask may be required according to the manufacturer’s label.
4. Other personal protective strategies e.g., bio-safety measures that must be applied during the application phase
5. Useful life of PPE, maintenance & care of PPE (if applicable) and appropriate disposal methods
6. Limitations of PPE
7. Risk, and hazards for not using PPE, or improper use, while working with specific chemicals
8. Process for reporting any breaches or concerns with selected PPE

### Training requirements – Chemical Specific

Air carriers should provide training to personnel related to the chemicals selected for use in the aircraft. In addition to the training for all personnel, those employees specifically tasked with performing chemical disinfection, either on the ground or during the flight, should be provided an additional specialized training either by the air carrier or the contract service provider on exposure control procedures in addition to being provided appropriate PPE. The correct application process, properties, risk exposures and hazards of the selected chemical should be part of the training. Inflight disinfection may need to take into consider additional aspects and provide more details on the specific tasks for inflight personnel. Employees who shall be assigned roles for use of chemical must understand:

1. Authorized chemical products to be used based on each jurisdiction or regulator
2. Recommendation of aircraft manufacturer regarding any new chemical composition introduced
3. Recommendation of chemical product manufacturer’s instructions to ensure that the proper application, ventilation and personal protection equipment is used
4. Chemical composition type and use of the chemical products
	* Chemical composition and effects of product used on aircraft
	* Instruction of use e.g.; Mixing ratios
	* Understanding of the Safety Data Sheets (SDS) for the products used
	* PPEs required for products used
5. Application methods and effective contact time in different areas of the aircraft e.g., Flight deck, cabin, cargo holds/compartments

Note: Use of chemicals in the flight deck must be approved by and supervised as per airline requirements. Effects on contact with aircraft electrical wiring should also be known and avoided where possible

1. Compatibility with different chemical or with non-chemical methods
2. Hazards and risks associated with the use of the products on personnel health and aircraft
3. Safe storage, hazards of storage, and disposal methods
4. Reporting channels in case of any incidents or accidents with use of the chemical

### SRA for Chemicals

The SMS process is a closed feedback loop that needs to be ongoing and reviewed. A minimum standard must be set that is scalable to address the emergent threat of pathogens. Furthermore, evaluation of the process must be completed periodically and increased when a known threat has been identified. Monitoring of the aircraft needs to be established for threats of degradation of systems and potential for safety risks. Employees and crews will require a reporting system to monitor their health and exposure complications.

It is recommended that the SRA committee be included in the monitoring process as the chemical application, process, and choice may need to be changed over time.

## Non-Chemical Disinfection Methods

Two general considerations are involved when evaluating non-chemical process; aviaiton airworthiness evaluation, approvals and the efficacy of the disinfection process for a particular pathogen. Generally, the non-chemical disinfection methods, and processes include portable devices and in-line installation devices which currently are part of aircraft type certification or are post-production installed devices requiring FAA and EASA approval. Non-chemical devices and processes include options in current use, in development, emerging technology, or an application of existing technology. There will likely be other processes and or solutions developed in the future.

One difference from chemical disinfectant solutions is that non-chemical processes may include installed devices requiring a different review and/or certification process where the focus is on safety, and airworthiness impacts. Generally, aviation safety regulators like the FAA or EASA approve the part or installation as a safe, or non-hazardous, component of the aircraft.

Note that aviation airworthiness authorities (regulator or regulators) are not the relevant agency for evaluating the efficacy of the disinfection process in relation to a pathogen.

Examples of non-chemical disinfection methods are:

* HEPA Air Filtration
* Ionization
* Ultraviolet
* Thermal

### Selection and Approval Process

* Applicable to all currently identified non-chemical processes in this guidance and listed in section 3.3.2, and other processes as may be agreed as non-chemical processes.
* Intended use, within the interior of an aircraft, of non-chemical processes to eliminate, neutralize surface and, or aerosol COVID-19 virus and other existing, known specified pathogens and scalable for pandemic responses.
* Non-chemical processes should be applicable to all aircraft pressurized spaces.
* For each device, process, if an aviation regulator approved device, OR NOT, an evaluation process following an agreed, nominal safety management systems (SMS), shall be employed, identifying the hazards, and analyzing, assessing, and controlling risk.
	+ A key characteristic differentiator is the specific process used in the presence of passengers, ground crew, flight crew or the process used when passenger, ground crew, flight crew are not present, and the SMS process shall include consideration, evaluation and accommodation for this differentiation.
* As a part of a suggested nominal SMS, the application of a Safety Risk Analysis (SRA) process should address intended use specifically regarding interaction with pathogens, including minimum performance expectations, scalable for pandemic responses.
	+ Recommended minimum SRA participants:
		- Applicable aircraft OEM and, or device OEM (preferably, both OEMs)
		- Aviation regulator, if applicable
		- Industrial hygienist, or equivalent
		- Subject matter experts, as needed
		- Operator (or representative), as applicable
	+ Review & recommendations regarding regulator documentation shall be undertaken.
		- If approval documents are in existence, provide and/or referenced.
		- Review device OEM supporting documentation, qualifications, testing.
	+ Review approval requirements with applicable aviation regulator to determine if regulatory approval required and process.
* Specifically, a nominal SMS process shall address the following issues, providing guidance to meet objectives as determined in the nominal SMS.
	+ Efficacy
		- Per the nominal SMS process, the non-chemical processes shall demonstrate that through the proper operation and application of the process, the process eliminates, or neutralizes the stated, intended targeted pathogens and per recommendation of the SRA.
			* Review and reference the testing documentation of independent testing and device OEM testing.
			* Determine process and periodic review feedback loop for continuous effectivity.
	+ Application locations / methods / phases of flight
	+ Effects on aircraft interiors / components
	+ Effects on humans
	+ Other Safety considerations
	+ Frequency
	+ Training requirements
	+ Use of non-chemical process, devices shall be same as determined by aviation regulator, and as required.
	+ Supplemental training, if required in consideration of intended use in pathogen elimination, neutralization, shall be determined through the agreed nominal SMS process and implemented by the device or aircraft OEM.

### Devices (Products, including air filtration)

* Non-chemical processes and associated devices identified in current use, under study, or in development.
	+ Aircraft Environmental Control System (ECS)/ HEPA Filters & Related
	+ Ionization
	+ Ultraviolet
	+ Thermal
	+ Other devices as may be identified
* Three of the four identified non-chemical processes, ECS, ionization and UV, are currently in use and as such have reference data to be used in the SMS process. Thermal process is a relatively new development when associated with civil aircraft. Here is brief discussion on each of the three in-use systems, with references to supporting data.

#### Aircraft Environmental Control System (ECS)/ HEPA Filters & Related

The ECS on civil aircraft is part of an aircraft type certification and as such has been approved by the regulator. This is airworthiness and does not necessarily mean the process is “approved” as effective for disinfection, neutralization and, or elimination of pathogens. The regulator approved documentation will or will not address the pathogen issues, such as efficacy, and may include effects on humans, for example. A best practice is to check with the aircraft or system OEM with regard to regulatory approval details, if there are any.

Most turbojet aircraft and some turboprop powered aircraft have some type of environmental control system that uses a mixture of engine bleed air and ambient air for airflow. HEPA filters are utilized on these aircraft and are capable of eliminating pathogens through the filtration process of airflow from the ECS,. These ECS are normally approved as part of the aircraft Type Certificate (TC) by FAA, EASA and other aviation regulatory organizations.

High-efficiency particulate air (HEPA), also known as high-efficiency particulate absorbing and high-efficiency particulate arrestance, is an efficiency standard of air filters. Filters meeting the HEPA standard must satisfy certain levels of efficiency. Common standards require that a HEPA air filter must remove—from the air that passes through—at least 99.95% (European Standard) or 99.97% (ASME, US DOE) of particles whose diameter is equal to 0.3 μm; with the filtration efficiency increasing for particle diameters both less than and greater than 0.3 μm.

The efficacy of HEPA filters is dependent on factors such as filter rating, airflow volume, airflow patterns, and how often the filters are changed.

Here is a listing of links to online information and reference documents regarding ECS and HEPA filtration that may be considered as a part of the SMS process for ECS:

* Federal Aviation Administration Regulations, 14 C.F.R. § 25.831(a); European Union Aviation Safety Agency, Certification Specifications, CS-25.831(a)2.
* ASHRAE 62.2-20163 (American Society of Heating, Refrigeration and Air Conditioning Engineers)
* American Society for Healthcare Engineering – Hospital HEPA usage in “protective environment" rooms https://www.ashe.org/compliance/ec\_02\_05\_01/01/airfiltration/
* The Airliner Cabin Environment and the Health of Passengers and Crew, National Research Council (US) Committee on Air Quality in Passenger Cabins of Commercial Aircraft. Washington (DC): National Academies Press (US); 2002

Computational Fluid Dynamics Modeling and the Transport of Cough Particles in an Aircraft Cabin - Atmur, Cummins, Olson, et al [White Paper], The Boeing Company https://www.boeing.com/confident-travel/downloads/Boeing-Computational-Fluid-Dynamics-Modeling-and-the-Transport-of-Cough-Particles-in-an-Aircraft-Cabin.pdf

#### Ionization

Ionization, on-aircraft installations currently available and in use are NOT part of aircraft type certifications and are installed under supplemental type certification (STC) process or an equivalent airworthiness approval process. Ionization installations are normally fitted in the aircraft airflow output ducting of an aircraft environmental system. herein s.Rsafe installationon aircraft, as approved These aviation safety regulators are not involved in determining disinfection efficacy of any process.

In general ionization can produce ozone, the concentration and amount being dependent on a number of variables. Current suppliers of non-portable ionization devices, installed on aircraft , claim their ionization devices do not produce ozone. As such, ozone presence testing should be a part of the SMS process.

Extensive on-aircraft installed ionization efficacy testing has, to date, been limited to large commercial air transport aircraft. The Boeing and Airbus general conclusion is that more extensive ionization efficacy testing is required and that, at this time, Boeing and Airbus do not recommend installation of ionization devices on aircraft. Boeing has published efficacy testing results to support their position not recommending ionization and recommending additional testing. To date, the Boeing white paper is the most extensive test and evaluaiton of ionization use on aircraft. The white paper is available through the following link.

Licht, Hehir, Trent, et al (2020).  Use of Bipolar Ionization for Disinfection within Airplanes [white paper].  The Boeing Company.

<https://www.boeing.com/confident-travel/research/use-of-bipolar-ionization-for-disinfection-within-airplanes.html>

Following is only intended as general information regarding ionization.

* Generally, the ionization process eliminates pathogens, in an airspace, such as an aircraft cabin, by electronically creating positive (H+) and negative (OH-) ions from hydrogen and oxygen atoms in the water vapor present in the air. Pathogens infect a host by binding to sites on the cell membrane. Viruses expelled from a person through mucus or saliva are airborne in aerosol form. Ionization works by leveraging an electronic charge to create a high concentration of positive and negative ions. These ions travel through the air binding to particles, which sets in motion a process of particle agglomeration. As these particles become larger, they are eliminated from the air.
* Additionally, positive and negative ions have microbicidal effects on all pathogens, rendering the COVID-19 virus non-infectious, while neutralizing and removing other viruses, allergens, contaminates and even mold spores. Here is a listing of reference documents that should be considered as a part of the SMS process involving ionization (NPBI) processes accessible at Aviation Clean Air online, [www.aviationcleanair.com](http://www.aviationcleanair.com) .

One supplier of aviation-specific ionization equipment, Aviation Clean Air, has available online (www.aviationcleanair.com) several test and evaluation reports that provide detailed information which should be useful in SMS process.

Tests for the Component's Ability to Neutralize Bacteria

* [SARS-CoV-2 Neutralization by Needlepoint Bipolar Ionization by Innovative Bioanalysis](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/phase_2_aca-iae_covid_test_official-2_final.pdf)
* [Efficacy of a Bipolar Ionization System - (C. difficile) by EMSL Analytical, Inc.](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/gps_cdiff_test_results.pdf)
* [Efficacy of a Bipolar Ionization System - (ECOLI) by EMSL Analytical, Inc](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/gps_ecoli_test_results.pdf)
* [Efficacy of a Bipolar Ionization System - (MRSA) by EMSL Analytical, Inc](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/gps_mrsa_test_results.pdf)
* [Efficacy of a Bipolar Ionization System - (TB) by EMSL Analytical, Inc](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/gps_tb_test_results.pdf)
* [Efficacy of a Bipolar Ionization System - (Reduction of L. Pneumophila) by EMSL Analytical, Inc](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/pneumonia_test_emsl_labs.docx)
* Ozone Emissions Testing by [Underwriter's Laboratories (UL)](https://www.aviationcleanair.com/uploads/1/3/3/2/133274601/gtr-aca-oz-0001_rev_nc-_aca_ionizer_ozone_emission_test_results_june_2019.pdf)

The efficacy of ionization is impacted by various factors in the dynamic environment of an operating aircraft. Current on-aircraft testing data, utilizing actual aircraft systems (i.e. Environmental Systems), in actual aircraft operations are not available and may not reflect real operational environments. Most testing has relied on relatively static simulations. Boeing, Airbus and the USA Center for Disease Control have indicated they feel more test and evaluation is needed, particularly when considering aerosol pathogens.

#### Ultraviolet

Ultraviolet (UV) treatment is an example of a process that is in current use by many airlines. A best practice guidance would be to refer to the aircraft and system OEM for detailed information and secure regulatory approval details, if available.

Ultraviolet light is the portion of the electromagnetic spectrum between 100 and 400 nm. It is divided into the vacuum-UV region below 200 nm, the UV-C region between 200 and 280 nm, and the UV-B and UV-A regions at higher wavelengths. The vacuum-UV region is so called because at these wavelengths oxygen from the air begins to absorb some of the light, limiting transmission and generating ozone. The UV-C region is most commonly used for germicidal treatment, since the peak wavelength for germicidal action is at 264 nm. Some germicidal activity remains in the UV-B and UV-A regions, and even with visible light, but much higher doses are required.



Figure 3‑1 The Spectrum of Light

##### Selection & approval process

Use of ultraviolet light for germicidal treatment, while widespread in hospitals and other settings, has only recently been considered for aircraft. Germicidal treatment of air using UV is well known for use in buildings[[1]](#footnote-2), and has now begun to be used in commercial aircraft. Treatment of surfaces using ultraviolet light is rapidly gaining acceptance and is useful in treating surfaces that are touched by passengers or crew in the cabin, lavatories, galley or flight deck. When UV treatment is completed while the aircraft is on the ground and no passengers are present, certification by airworthiness authorities may not be needed. Local and regional laws and regulations should be reviewed. In the United States, UV devices are regulated by the EPA[[2]](#footnote-3). These rules will include provisions to ensure electrical safety and the safety of personnel using the equipment.

##### Products

Ultraviolet germicidal products for aircraft use include both hand-held wands or lamps, and larger systems on wheels. Handheld systems are lightweight and can be used in tight spaces where the larger systems will not fit. Typically, the lamps are lower in intensity, so they must be held closer to the surfaces to be disinfected and/or used to irradiate the surfaces for a longer period of time. The angle between the surface to be treated and the lamp focus direction is also important. Operator training and diligence is very important with hand-held units, since holding the lamps too far from the surfaces or using them too quickly will result in inadequate treatment. As a result, the SMS approval for handheld devices to be used should specify the distance and exposure time for surfaces. UV systems mounted on wheels are larger and heavier than the hand-held units. Frequently equipped with a rechargeable battery, they can move down the aisle of an aircraft cabin, disinfecting all surfaces exposed to the UV light. Because of the higher light intensity, they can be used more rapidly than a hand-held system, and because of the fixed geometry, they expose surfaces in a reproducible way. The UV dose will be determined by the light intensity (a function of the distance between the lamp and the surface) and the speed with which the device is moved through the cabin. The UV system can be operated by a single individual, appropriately protected to avoid UV light exposure, or robotic systems which avoid the risk of human exposure. The SMS should specify the rate to be used, and further directions to ensure that all necessary touchpoints will receive an adequate dose.

UV devices can also vary in the wavelength of the light used. The most common is 254 nm light, which has commonly been used in hospital and other applications. These devices generally use mercury vapor lamps similar to fluorescent lamps. Pulsed xenon lamps provide a broader spectrum of UV light, including the UV-C region. UV-LED lights can also be used and are significantly smaller, with wavelengths from 280 nm to 400 nm. “Far-UV” lamps with a wavelength of 222 nm have recently been proposed and use krypton chloride excimer lamps. The efficacy of these different wavelengths will be discussed below. Some lamps emit multiple wavelengths of light . If emission occurs below 200 nm, there is potential for ozone to be generated. Filters or phosphors are sometimes used to prevent emission below 200 nm.

 Filters or phosphors are sometimes used to prevent emission below 200 nm.

##### Efficacy

Properly applied, Ultraviolet Germicidal Irradiation (UVGI) has been found to reduce pathogens - including tested bacteria and viruses - on multiple surfaces and in multiple environments.[[3]](#footnote-4),[[4]](#footnote-5),[[5]](#footnote-6) UV treatment can be performed on surfaces, in which case the applied intensity and treatment time are easily measured, or on a volume of air. In the latter case, the intensity will change with the distance from the lamps, and the residence time of the microorganisms in the irradiated zone will depend on the air velocity.

The reduction in live organisms on a surface or in the air is a function of the dose or fluence of ultraviolet light, and the micro-organism specific and wavelength specific rate constant. Dose (fluence) is defined as the intensity of UV light at the surface multiplied by the irradiation time. Typical units for intensity are mW/cm2 and typical units for dose (fluence) are mJ/cm2. The relationship between dose and fractional reduction is expressed[[6]](#footnote-7) in equation 1, where K is the rate constant, H0 is the dose and D is the fractional reduction. Increasing the intensity or increasing the exposure time increases the % reduction.

$$H\_{0}= \frac{-ln⁡(1-D)}{K} (1)$$

It follows from this equation that, to ensure adequate reduction of infectious organisms, the dose at the surface to be disinfected must be known or estimated, and compared with the dose required to achieve the desired reduction. For surfaces, the intensity of the UV source(s) must be known, the distance from these sources to the surface, and the exposure time. Frequently, manufacturers of UV devices will provide guidance for exposure time, and will provide estimates for the dose when the device is being used in accordance with their guidelines. It is also possible to measure the dose with a commercially available dosimeter. The dose can be compared to the dose required to treat various bacteria, viruses etc. using published compilations. For example, for UV radiation at 254nm, the International Ultraviolet Association (IUVA) has published a compendium of numerous studies with dose data[[7]](#footnote-9). A more limited list is also available for UV radiation at 222 nm[[8]](#footnote-10).

Disinfection for COVID-19 is of especial concern. Blatchley et al[[9]](#footnote-11) have published a collection of many recent studies to determine the dose required, and found that a dose of 5 mJ/cm2 (254 nm radiation) corresponds to 99.9% removal on surfaces. Information on the dose required to use 222 nm radiation against COVID has been published[[10]](#footnote-12),[[11]](#footnote-13), and dose information for 275 nm LEDs and for pulsed xenon radiation is also available[[12]](#footnote-14).

UVGI has been used to inactivate viruses in hospitals and other critical public and military environments for years.[[13]](#footnote-15) When disinfection is event-driven or concerns group 4 pathogens, UV treatment should be replaced or supplemented by chemical cleaning by companies certified for this task e.g. ebola, human, animal liquids on board, death on board, etc.

In addition to the wavelengths already discussed, other wavelengths have been used. The relative germicidal efficiency vs. wavelength is well known[[14]](#footnote-16), and reaches a peak at 264 nm. At higher wavelengths, higher doses are required. Scientific studies indicate that 405 nm, which is near the high wavelength limit for UV-A light, may be effective in reducing certain bacteria but are not conclusive with regard to its ability to inactivate certain viruses. . The referenced studies found no effectiveness against viruses in some cases, and potential effectiveness against viruses only when suspended in specific organic media in other cases. The authors conclude that further work should be carried out to establish the effects of 405nm light.[[15]](#footnote-17),[[16]](#footnote-18)

Safe limits for human exposure are discussed in section 3.3.2.3.6.

##### Application Locations/Methods/Phases of Flight

UV treatment can be completed while the aircraft is on the ground and does not contain passengers. In the future, UV treatment technologies for use during flight may be developed. Since UV leaves no residual disinfecting residue, it is only effective to disinfect surfaces at the time of disinfection exposure. It may be performed in between flights as part of a normal cleaning cycle, or more intense cleaning may be performed when more time is available or when there is concern that an infected person may have been present in the aircraft. A UV dose, as described above, should be selected with the assistance of the SMS, and should meet or exceed the dosage on surfaces including seats, seat belts, arm rests, trays, in-flight entertainment screens and other displays, overhead bins and passenger service units. The SMS in this case should include representation from the original equipment manufacturer, the UV equipment provider, and a hygiene expert cognizant of local and regional standards. With hand-held units, this requires that the operator hold the lamps in an appropriate distance from the surface for an appropriate length of time, according to the manufacturer’s directives. For larger units on wheels, the manufacturer’s recommendations for the rate of travel with the device should be reviewed. One product manufacturer has suggested 30 rows/minute pace down the aisle of a commercial aircraft for a rapid cleaning, and a 10 rows/minute pace for in depth treatment. In contrast to chemical disinfection, there is no safety-related need for UV treatment to ventilate the disinfected location prior to entry of personnel.

##### Effects on Aircraft Interiors/Components

In order for UV-C light to affect the properties of a material, two things must happen: (1) absorption of the light, and (2) chemical reaction[[17]](#footnote-22). Many materials that are transparent to visible light are opaque to UV light, limiting penetration of the light deep into the material. Other materials, like leather or other fabric materials commonly used for aircraft seating, are opaque because they scatter the light or because of materials blended into the polymers from which they are made. For these materials, the effect of UV-C light will be only on the exposed surface. Once light absorption has occurred, surfaces exposed to air can oxidize, with possible impact on their strength or other properties. In the absence of oxygen from the air, other chemical reactions can occur which may result in color changes to the material.

All UV-C effects on materials are dose-dependent. Dose can be generally measured by the light intensity and time, so the same effect can be observed with low intensity over a long period of time or high intensity for a short time. Since the intensity of the light on a surface will depend on the distance between the surface and the light source, and the angle between the plane of the surface and the incident light, it will be different for different devices or treatment techniques. The cumulative dose experienced by materials in the cabin can be estimated by multiplying the single treatment dose by the number of treatments per day, and the number of treatment days.

Cumulative dose = Treatment dose \* Treatments/day \* Treatment days

Aircraft materials in the cabin can include wool, polyester, leather or synthetic leather seat coverings, carpets, seat belts, tray tables, in-flight entertainment LCD screens, and other displays, transparent window coverings, decorative foil laminates and window shades. Flight deck materials include a similarly wide range of materials. Materials compatibility information is specific to each material and UV wavelength, and must cover the range of cumulative dose corresponding to realistic use. Materials property information that is useful to airlines includes the effect of UV light on flame retardancy, the effect on strength of the materials, and the effect on appearance. Recent studies using 254 nm UV-C radiation[[18]](#footnote-23),[[19]](#footnote-24) and 222 nm Far-UV-C radiation[[20]](#footnote-25) summarized measurements of the effects on these materials.

The results can be summarized as follows:

* None of the materials tested experienced any detectable change in flame retardancy. The maximum dose tested was 269 J/cm2 for 254 nm radiation and 100 J/cm2 for 222 nm radiation.
* None of the materials tested experienced any detectable changes in tensile strength. The maximum dose tested was 191 J/cm2 for 254 nm radiation and 100 J/cm2.

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* Color changes were observed for lightly colored materials after extensive UV exposure. Materials that were most affected included polyvinyl chloride/ polycarbonate thermoplastics (used in tray tables and seat assemblies, and the glues used to mount decorative laminates. Darkening was observed after a dose of 17-34 J/cm2 for 254 nm radiation and a similar dose for 222 nm radiation.

Using the equation a dose of 50 J/cm2 would correspond to 10,000 treatments if a single treatment was chosen to be 5 mJ/cm2.

##### Effects on humans

Exposure to an excessive dose of UV light can be harmful, especially to exposed skin or eyes. The limits for safe exposure have been established. Threshold limit values for UV exposure depend on wavelength and are expressed as cumulative dose over an 8 hour period. For the wavelengths in potential use for aircraft germicidal treatment, the limits are shown below from the ACGIH handbook[[21]](#footnote-27) and from the International Commission on Non-Ionizing Radiation Protection[[22]](#footnote-28). EU Directive 2006/25/EC also provides threshold limit values which are the same as those shown for ACGIH.[[23]](#footnote-29)

Table 3‑3 Threshold Limit Values for 8 hour exposure interval

|  |  |  |
| --- | --- | --- |
| Wavelength (nm) | ACGIH TLV (mJ/cm2) | ICNIRP (mJ/cm2) |
| 400 | 1 x 105 | Not stated |
| 254 | 6 | 6 |
| 220 | 25 | 25 |

The person operating UV equipment should ensure that other personnel are not in close proximity. The operator can be protected by a UV-opaque shield if possible and should wear clothing that cover the arms and legs. Shoes should completely enclose the foot. If no shield is used, gloves may be worn and a faceshield should be used. If a protective shield is damaged, compromised or is not correctly positioned to protect the operator, the equipment should not be used. Similarly, if autonomous or robotic equipment is used, personnel exposure to UV may be avoided.

##### Other Safety considerations

UV systems are electrically powered and frequently use batteries. The operator should read and be familiar with the manufacturer’s manual and safety warnings, and training should include safe operation. Like any electrical appliance, the system and any battery charger should be kept away from water, heat sources, flammable liquids or areas with flammable gases, vapors, or explosive/airborne dust. If the equipment has been dropped, damaged, left outdoors or dropped in water, it should be disconnected from electrical power, and not used until the equipment has been checked.

UV equipment may use bulbs, LEDs or other UV sources. UVC bulbs are very similar to common fluorescent bulbs and contain a very small amount of mercury. Care should be taken to avoid breaking them, and the equipment should be correctly designed to reduce the risk of lamp breakage. If a lamp does break, the clean-up procedure is similar to that for fluorescent lamps.

##### Training requirements

UV equipment should be operated only by trained personnel. Manufacturers will provide a manual for the safe operation of the equipment, and personnel should be trained in and familiar with the contents of this manual. The manual will also provide equipment-specific safety warnings and guidance, and this should also be included in training. Supervisors should oversee UV equipment operators to ensure that the equipment is being operated safely and effectively.

Ultraviolet light sources used for disinfection should not generate ozone. . It is important to check the specifications for any UV light source to check that it does not generate ozone, and that it is compliant with any local or regional regulations or standards. Do not use replacement lamps for any UV source from a source other than the original vendor without making sure that these lamps also comply with these regulations.

####  Thermal - are we deleting this section for the section that follows?

Thermal Heating or Thermal Disinfection is an example of an emerging technology to combat SARS-CoV-2. Viruses and bacteria can be inactivated (die off) when exposed to heat for a certain period of time. The specific thermal inactivation temperature with associated relative humidity, and length of heat exposure required to be effective for each virus and bacteria is different.

Efficacy of thermal disinfection is related to a combination of other environmental factors.

Specifically, it appears humidity plays a major role in the relationship between temperature, time, and kill rate for SARS-CoV-2. Currently, studies are showing at certain relative humidity ranges that SARS-CoV-2 thermal inactivation can be achieved from around 50 degrees C at 30 minutes exposure to around 70 degrees C at 10 minutes exposure.[[24]](#footnote-30) Airplane manufacturers and the U.S. military are studying thermal heating as a potential disinfection solution for the flight deck, cabin, and cargo compartment. More testing is needed before conclusions can be made about the efficacy and viability of thermal heating for an aircraft. Testing also needs to address safety of equipment and parts after repeated heating cycles, functional inspection after testing, and proper safety guidelines.

While testing may prove heat and humidity can eliminate biological contamination, for aircraft operations, the operational heat limits still need to be addressed to ensure safety. As the process evolves or the suspect pathogen changes, operators wishing to utilize the thermal heating should coordinate with knowledgeable entities such as safety regulators, aircraft and equipment OEMs to research the viability of the option, including conducting a SMS and SRA process.

#### Thermal

Thermal Heating or Thermal Disinfection is an example of an emerging technology to combat SARS-CoV-2. Thermal Disinfection involves heating surfaces for prolonged periods of time to deactivate viruses and bacteria. Thermal disinfection also prevents the risk of moisture ingress posed by liquid-based disinfectants and limits the potential for missed spots and ergonimic issues that may occur when maually disinfecting surfaces by hand. The specific thermal inactivation temperature with associated relative humidity, and length of heat exposure required to be effective for each virus and bacteria is different. Studies have been conducted to show the functional capability and efficacy for thermal heating against SARS-CoV-2 30 (superscript). Efficacy of thermal disinfection is related to a combination of other environmental factors.

Specifically, humidity plays a major role in the relationship between temperature, time, and kill rate for SARS-CoV-2. Studies have shown that thermal inactivation temperatures for SARS-CoV-2 can be achieved at 40 degrees C to 55 degrees C depending on the amount of time surfaces are exposed to these temperatures. Airplane manufacturers and the U.S. military are studying thermal heating as a potential disinfection solution for the flight deckOperators performing thermal disinfection also need to address safety of equipment and parts after repeated heating cycles, functional inspection after testing, and proper safety guidelines.

While the testing may prove heat and humidity can eliminate biological contamination, for aircraft operations, the operational heat limits still need to be addressed to ensure safety. Thermal disinfection should be performed using external heaters and in an airplane depowered state so as to avoid cooling air being introduced from the on board aircraft cooling systems which may prevent surface temperatures to rise to the thermal disinfection temperature. Electrical heater blankets are not recommended for thermal disinfection due to potential for overheat and damage.

As the process evolves or the suspect pathogen changes, operators wishing to utilize thermal heating should coordinate with knowledgeable entities such as safety regulators, aircraft, and equipment OEMs to research the viability of the option, including conducting a SMS and SRA process.

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