The Airborne Isolation Hood

Instructions for Health Care Workers

Product Name: The Airborne Isolation Hood

The U.S. Food and Drug Administration has issued an Emergency Use Authorization (EUA) for Airborne Isolation Hood, for use by healthcare provider (HCP) as an additional layer of barrier protection in addition to personal protective equipment (PPE) to prevent HCP exposure to pathogenic biological airborne particulates by providing temporary isolation of hospitalized patients with suspected or confirmed diagnosis of COVID-19, at the time of definitive airway management, when performing airway-related medical procedures, or during certain transport of such patients during the COVID-19 pandemic.

Authorized non-transport use of Airborne Isolation Hood is only for definitive airway management (e.g., intubation, extubating and suctioning airways), or when performing any airway-related medical procedures (e.g., high flow nasal cannula oxygen treatments, nebulizer treatments, manipulation of oxygen mask or continuous positive airway pressure/bi-level positive airway pressure [CPAP/BiPAP] mask use, airway suctioning, percussion and postural drainage). Authorized use of Airborne Isolation Hood during patient transport is only within a hospital setting for temporary transfer with direct admission within the hospital in the presence of a registered nurse or physician. Maintenance of negative pressure with adequate air flow must be ensured. The patient should have constant monitoring of vital signs, electrocardiogram (EKG), SpO2%, and EtCO2, if available, throughout transport. If end-tidal CO2 monitoring is not available, then the use of the Airborne Isolation Hood Device must be limited to no longer than 30 minutes with inline blower fan on and under direct observation. For all authorized uses, the patient should always have supplemental oxygen during use of the Airborne Isolation Hood Device. The Airborne Isolation Hood Device is for use in addition to PPE for HCP during the COVID-19 pandemic and does not replace the need for PPE.

Airborne Isolation Hood has not been FDA-approved or cleared for this use; Airborne Isolation Hood has been authorized for emergency use by FDA under an EUA. Airborne Isolation Hood has been authorized only for the duration of the COVID-19 public health emergency declaration that circumstances exist justifying the authorization of the emergency use of medical devices under section 564(b)(1) of the Federal Food, Drug and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.

HCP should follow these instructions, as well as procedures at their healthcare facility to use the Airborne Isolation Hood.

The instructions below are to assist in using Airborne Isolation Hood. Airborne Isolation Hood is an adjunctive protective barrier designed to mitigate risk to HCP. Airborne Isolation Hood is not meant to be a stand-alone unit of PPE. Airborne Isolation Hood should always be used with appropriate PPE and pursuant to the guidance of your institution.

All connections should be tightly secured and checked frequently. Any time anyone is within Airborne Isolation Hood, direct observation is required. Inspect Airborne Isolation Hood prior to use. Any wear/tear of the chamber or other signs of degradation on Airborne Isolation Hood must promptly be reported to Airborne Isolation Hood Medical Solutions Inc. The healthcare facility must not use on
patients and must dispose of such a Airborne Isolation Hood. Rx Only

WARNINGS:

• Flammability of Airborne Isolation Hood has not been tested. No interventions that could create a spark or be a flammable source should be used within Airborne Isolation Hood.
• Remove Airborne Isolation Hood and use standard of care if there is difficulty visualizing or identifying anatomic landmarks or inability to intubate after the first try.
• Prolonged use of Airborne Isolation Hood may induce hypercarbia in a spontaneously breathing patient. Airborne Isolation Hood should be used with medical air flow and suction both on and working, under direct observation, and with EtCO₂ monitoring if available.
• Patient transport must only occur within a hospital setting for temporary transfer with direct admission within the hospital in the presence of a registered nurse or physician. Maintenance of negative pressure with adequate air flow must be assured. All patients should be on supplemental oxygen. Patients must have continuous monitoring of Sp-O₂, vital signs, EKG, and EtCO₂ if available during transport.
• Use caution prior to use on non-sedated or lightly sedated patients with severe claustrophobia and/or confined space anxiety.
• When using Airborne Isolation Hood, patients should always be receiving supplemental oxygen.
• Patients with diminished hearing may have difficulty understanding the provider while inside Airborne Isolation Hood.
• Airborne Isolation Hood is a single-use device and should be disposed of following the disposal instructions after use.
• Delayed emergency removal of the device may result in patient injury

CONTRAINdications:

Airborne Isolation Hood is not authorized:

• On patients needing emergent endotracheal intubation with severe hypoxemia • On patients with anticipated or known history of difficult airway
• On patients with other anatomical abnormalities that might interfere with clinical care including decreased neck mobility from arthritis or other causes
• On patients with anticipated or known history of claustrophobia and/or confined space anxiety
• On patients with communication disorders that might interfere with clinical care
• On patients under the age of 18
• On bariatric patients
The Airborne Isolation Hood Instructions for Use

Device Description:

The Airborne Isolation Hood provides respiratory isolation of patients to decrease risk of spread of COVID-19 to healthcare personnel (HCP) and visiting family.

The Airborne Isolation Hood is a clear, rigid chamber made of transparent plastic that covers a patient’s head and upper body and uses negative pressure airflow to filter respiratory particles from the chamber, through a high-efficiency particulate air (HEPA) filter, to isolate those outside the hood from the exhalation and respiratory secretions of the patient inside the hood. The hood is a protective barrier enclosure, with negative pressure and sealed ports, utilizing silicon gaskets, through which HCP hands are passed to perform medical procedures, for isolated access to the patient. There is also a small cutout at the bottom of the hood with a silicone barrier that acts as a cord feedthrough so that the hood can be removed without tangling cords attached to the patient. The hood has four tie-down points mounted to the polycarbonate panels and is attached to the bed with two black rubber elastic cords with metallic “S” hooks on each end. All components are disposable. During transport, the hood must be used with a battery pack adequate to power the blower.

System Components

- Respiratory Isolation Hood
- Exhaust hose-Clear Blo-R-Vac Duct Hose 4” ID; aluminum coated flexible hosing
- Blower Fan, 8-speed setting blower (AC Infinity CLOUDLINE S8)
- Levoit Air Purifier Filter; LV-H132XR
- Electrical Outlet/Power Source
- Cleaning supplies (Isopropyl Alcohol or similar) per hospital protocol
- Transport after use on a hospital owned cart and securement devices (e.g., bungee cords or nylon rope)
- Optional plastic drape for covering for duct

Note: The AC Infinity CLOUDLINE S8 Blower has an 8-speed setting, and the speed can be controlled currently by a connected remote. Increasing the speed would increase the suctioning, which would control the negative pressure setting.
Placement Instructions

To use this portable Isolation hood, have the patient positioned appropriately and place all supplies on the bed next to the patient within the area of the enclosure. Initial installation of the system above a patient should be done with adequate personal protective equipment (PPE).

1. Place the enclosure on the bed over the patient.
2. Ensure all lines pass through cord feedthroughs or the open face of the box (towards patient’s feet).
3. **Note:** Please do not insert wires/tubing through the iris ports to avoid tethering (to allow easy emergency removal of the protective hood if required)
4. Ensure in-line blower and exhaust hose are connected to device.
5. Plug into electrical outlet and turn on blower system. (Fan power setting of 4 for efficacy)
6. Apply respiratory modalities/care to the patient.
7. Use two bed attachments and connect one end to a tie-down point on the side of the Aerosol Hood, pass the attachment through the side of hospital bed rail, wrap it under the corner of the bed, pass up through the metal mattress support, and connect it to the adjacent tie-down point on the back side of the Aerosol Hood. Repeat this process for the other corner of the Aerosol Hood. Tie-downs can be disconnected easily in the event of a situation where the box needs to be moved quickly. Ensure bed attachments are in place if there is a need to elevate the head of the bed.

Placement Instructions for Intubation

The Airborne Isolation Hood provides effective respiratory isolation; thus, face mask ventilation is acceptable unless contraindicated for full stomach reasons; RSI may also be done when indicated

1. Patient supine on bed (aerosol unit is on)
   a. Have required materials ready and placed on the head of the bed next to the patient (e.g.; CMAC, ET tube with cuff syringe, eye tape, oral airway etc.). Introduce all lines through the cord feedthroughs or the open face of the box.

b. Clinician may consider mild sedation for patient comfort

2. Place device on bed over the patient:
   a. Provider positions patient and places box in preparation for intubation, begins passive pre-oxygenation with bag-mask ventilation with or without assistance

3. Administer anesthetics of your choice
   a. Utilize video laryngoscopy (preferred) or direct laryngoscopy for intubation
   b. Secure endotracheal tube after confirming appropriate placement with ETCO2 sensor and bilateral breath sounds
   c. Attach high-efficiency particulate air (HEPA) filter to endotracheal tube as required, connect to ventilatory circuit.

Removal Instructions

1. While wearing adequate PPE, wait 30 seconds after treatment before removing the hood. Aerosol will be cleared at a 99.9% reduction at 27.19 seconds when blower is at level 4 and 12.11 seconds at when blower is at level 8.
2. Turn off the blower system, unplug the device from the electrical outlet.
3. Detach the bed attachments from the bed.
4. Ensure all wires/tubing are free of the unit
5. Lift and raise the unit straight up off the bed
6. Place on transport cart for in-room cleaning and then transport per local policies for prescribed sanitation and reuse

**Replacement of Filter and Duct**

- It is recommended the filter and exhaust hose (duct) be replaced every 3 months.
- The replacement for the HEPA filter is a Levoit Air Purifier Replacement Filter for LV-H132XR. The filter can be removed from the filter housing by pulling on the side tabs of the filter, which are visible upon opening the housing thumb screws and opening the housing. Be sure to orient the replacement filter in the proper direction. The filter can be directly installed without modification needed.
- The replacement for the exhaust hose is a 6-foot length of Clear Blo-R-Vac Duct Hose 4" ID.
- Replacement components are commercially available, and can be sourced directly from Aspire Air, Inc, or from a commercial supplier.

**Emergency Removal Instructions**

1. Emergency removal of the hood should only be performed by healthcare workers who are wearing adequate PPE for COVID-19 patient care.
2. If the patient is actively coughing, and clinically appropriate, the patient should wear a mask before removal.
3. The particles under the hood will be a mixture of entrained air from the room and those generated by the patient.
4. Ensure all medical wires and tubing are not intertwined with the device
5. Ensure the attachments to secure the device to the patient bed are disconnected
6. Lift and raise the unit straight up off the bed
7. The system is not required to be turned off during the physical removal process, as the HEPA filtration system can continue to filter room air. The exception to this is if it is required to disconnect the outflow tubing from the unit in which case turning off the blower will reduce the possibility of entraining unfiltered air and spreading unfiltered particles within the room.
8. Once all attachments are disconnected, the healthcare worker does not need to wait any additional time before removing the hood.

**Sanitization and Reuse Instructions**

1. For cleaning/disinfection, the sponsor referred to the CDC’s recommendation for hospitals/facilities - [https://www.cdc.gov/coronavirus/2019-ncov/community/disinfecting-building-facility.html](https://www.cdc.gov/coronavirus/2019-ncov/community/disinfecting-building-facility.html). The disinfectant used should be from the CDC’s list of disinfectants for COVID19 found here: [https://www.epa.gov/pesticide-registration/list-n-disinfectants-coronavirus-covid-19](https://www.epa.gov/pesticide-registration/list-n-disinfectants-coronavirus-covid-19)
2. For cleaning, after each patient use, recommend spraying all inner and outer surfaces with approved disinfectant for surfaces per local guidelines (e.g. isopropyl alcohol-based disinfectant, such as Cavicide or Peroxide Multi Surface Cleaner and Disinfectant). Follow instructions per hospital requirements on disinfectant specific contact times.
3. Alternatively, after each patient use, wear gloves and wipe down the plastic outside surfaces, inside surfaces, and iris ports with disinfecting wipes or isopropyl alcohol such as Cavicide as used by the institution. All materials are isopropyl alcohol and disinfecting wipe compatible.

4. If possible, soil remains, repeat the procedure. Let all parts sit visibly wet for at least 2 minutes. Allow all parts to air dry. Remove and discard gloves.

5. For disinfection, spray all disassembled parts with a hospital approved EPA-registered isopropyl alcohol-based disinfectant, such as Cavicide or Peroxide Multi Surface Cleaner and Disinfectant. Let all parts sit visibly wet for at least the contact time indicated in the labeling (2 minutes for Cavicide) before allowing all parts to air dry. See List N: Disinfectants for Use Against COVID-19: https://www.epa.gov/pesticide-registration/list-n-advanced-search-page-disinfectants-coronavirus-covid-19.

6. Do not use harsh chemicals or abrasives to clean. Do not apply heat. Do not use high concentrations of cleaning agents than recommended as this may degrade components.

7. Store device at room temperature and a relative humidity between 40-60%. Store in a container, if possible, to avoid dust and grease build up on the device.
   - The blower fan, exhaust hose (duct) and HEPA filter may remain connected to the Airborne Isolation Hood when it is not in use. These can be placed on a shelf on the cart below the hood, or inside the hood, during transport to storage, and while being stored.

8. The exposed surfaces on the blower may be wiped down (same wipes) after each use. There is no need to take anything apart in the blower.

9. It is recommended that the connections remain intact unless changing out filter or tubing. Do not put hands in the interior space of the tubing or open the filter housing, to reduce the risk of contaminating this space.

10. It is recommended the filter and exhaust hose (duct) be replaced every 3 months.