Interim Guidance: Use of Personal Protective Equipment by Healthcare Personnel in Resource-Constrained Settings

Background

The Oregon Health Authority (OHA) is following federal guidance on the use of personal protective equipment (PPE) by healthcare personnel (HCP). At the same time, we recognize that more granular guidance, based on the best evidence we have to this point, will be useful to HCP and health systems. A common set of strategies listed in order of efficacy for infection prevention is important for our shared goal of protecting HCP. To this end, OHA has worked with a technical advisory group of infectious disease physicians and infection preventionists from around the state to develop this guidance. In most cases, it represents expert opinion, rather than stronger levels of evidence, which are currently lacking. Nonetheless, we have heard from multiple partners that a uniform “playbook” for response to this challenging situation would be valuable. We hope you will find it useful.

Definitions

- **Administrative controls**: work practices that prevent or reduce hazardous exposures.

- **Aerosol-generating procedures (AGPs)**: procedures that generate small droplet nuclei in high concentration, presenting a risk for airborne transmission of pathogens not otherwise able to spread by the airborne route (e.g., coronavirus, influenza).

- **Emergency Use Authorization (EUA)**: permission granted under section 564 of the Federal Food, Drug, and Cosmetic Act (FD&C Act) by the FDA Commissioner to allow unapproved medical products or unapproved uses of approved medical products for use in an emergency to diagnose, treat, or prevent serious or life-threatening diseases or conditions caused by chemical, biological, radiological, nuclear, and high yield explosives (CBRN) threat agents when there are no adequate, approved, and available alternatives.

- **Engineering controls**: measures that reduce or eliminate worker exposures to hazards, such as through exhaust ventilation or by placing a barrier between the worker and the hazard.
- **Expedient patient isolation room approach**: the use of portable of high-efficiency particulate air (HEPA) filtration systems to establish a high-ventilation-rate, negative pressure, inner isolation zone that sits within a “clean” larger ventilated zone.

- **Expired PPE use**: use of PPE beyond the manufacturer-designated shelf life during patient care.

- **Extended use**: the practice of wearing the same PPE for repeated encounters with several patients, without removing the device between patient encounters.

- **Face mask**: mask covering nose and mouth to protect the wearer or the environment from respiratory droplets; includes procedure mask or surgical mask.

- **Industrial-grade PPE with EUA**: PPE designed for use in other industries determined by federal regulatory agencies to provide adequate protection in a healthcare setting.

- **NIOSH-approved particulate filtering facepiece respirator**: particulate respirators that meet or exceed standards for design suitability, quality assurance, and laboratory performance, which includes filtration efficiency and ability to maintain a seal on the user’s skin. NIOSH, under authorization of the Occupational Safety and Health Act of 1970, provides a testing, approval, and certification program assuring respirators used in the workplace meet the standards of 42 CFR Part 84. Since 1994, NIOSH has maintained a searchable, online version of the Certified Equipment List.

- **Respirators with exhalation valves**: Respirators with a valve allowing unfiltered breath from the wearer to escape into the environment. Such respirators should not be used for surgical and other procedures requiring a sterile field, are not appropriate for source control, and are not approved for decontamination under FDA-issued EUAs.

- **Re-use**: use of the same PPE by a single provider for care of several patients, with removal between patients. It also includes extended use of several pieces of equipment in rotation on successive days.

- **Re-use after decontamination**: the practice of using PPE that is permitted by the manufacturer to be decontaminated following an FDA authorized method so that it retains regulatory agency (e.g., NIOSH, FDA) approvals and authorizations. Such PPE can be returned to the general PPE supply or to the original user following decontamination.

- **Single use**: the practice of using the PPE for one encounter with one patient, then discarding it.

- **Source control**: use of protective equipment or other measures to prevent the spread of illness from an infectious person to others, for example, use of a mask by an infectious person to limit the spread of a respiratory illness.
Scope

This guidance is specific to PPE in the context of a COVID-19 pandemic in healthcare settings across the continuum of care. It outlines strategies for optimizing the supply of PPE in healthcare settings when PPE resources are constrained. Additional guidance for dental practices is available on the OHA website. Compliance with Oregon OSHA standards remains a requirement. A few standards that apply include, but are not limited to, Bloodborne Pathogens (29 CFR 1910.1030), Respiratory Protection (29 CFR 1910.134) and other PPE (OAR 437-002-0134).

Oregon OSHA is addressing supply chain considerations, including respirator shortages, through enforcement flexibilities, as discussed in the federal OSHA Enforcement Memoranda section of the Standards page.

See information on PPE flexibilities and prioritization in the Personal Protective Equipment Flexibilities section within the Interim Guidance for U.S. Workers and Employers of Workers with Potential Occupational Exposures to SARS-CoV-2.

Framework

With the understanding that we may face sustained PPE shortages during the COVID-19 response, the guidance comprises three components while applying the hierarchy of hazard controls system:

1. Implement engineering and administrative control measures to optimize PPE use
   - These measures must fully be implemented when possible before other strategies for optimizing PPE supply are considered and implemented.

2. Use a tiered approach to optimize PPE supply
   - Strategies for optimizing the supply of PPE are arranged in four tiers, with each successive tier representing increasingly resource-constrained situations. Keep in mind, employers must provide effective protections to healthcare providers during resource-constrained situations.
   - Within each tier and for each PPE type, the strategies are further organized by three elements that can be considered: the PPE type, the duration of its use, and any additional engineering or administrative controls. (See Table 1.) Table 1 also summarizes the CDC’s conventional capacity strategies by PPE type.
   - Generally, Tier 1 and 2 strategies align with CDC’s contingency capacity strategies, while Tier 3 and 4 strategies correspond to CDC’s crisis capacity strategies.
3. Monitor PPE availability to guide optimal PPE use

- Facilities and providers understand their PPE inventory, supply chain, and utilization rate.
- Facilities and providers are in communication with local healthcare coalitions, as well as local and state public health partners regarding identification of additional PPE supplies.
- Facilities and providers must regularly assess the need for all optimization strategies by PPE type, and move to a lower-number tier when the supply of a specific PPE type allows, with the goal of eventually following conventional capacity strategies for all PPE types. The regularity of these assessments should increase as higher-numbered tier strategies are implemented.

General Required Measures to Minimize PPE Use

**Engineering controls**

- Promptly isolate symptomatic patients
  - Move from common waiting area to private room with door closed
  - Selective use of airborne infection isolation rooms (AIIRs), e.g., when performing AGPs.
- Use of physical barriers
  - Glass or plastic windows in reception and patient areas (e.g., intake desk at emergency department, triage station, information booth, or pharmacy service counter)
  - Closed suctioning systems for airway suctioning for intubated patients
- Properly maintained HVAC systems
  - Investigate increasing percentage of outdoor air in the HVAC air supply
  - Increase air filtration to highest level (MERV 13 or 14) compatible with HVAC
  - Provide air movement from clean areas to contaminated areas.
  - Provide 6 room air changes per hour (ACH) in older buildings and 12 ACH in newer buildings (appropriate for high-risk areas).
- Consider use of portable HEPA air filtration units for source control (especially in higher-risk areas).
- Consider use of ultraviolet germicidal irradiation (UVGI) as a supplement to help inactivate viruses.
**Administrative controls**

- Limit the number of patients going to hospital or outpatient settings.
  - Implement a patient screening system and instruct those with mild to moderate symptoms who do not require a higher level of care to remain at home.
  - Use telemedicine when appropriate.
- Limit face-to-face HCP encounters with suspected or confirmed COVID-19 patients
  - Limit patient room and care area entry to medically essential HCP as much as possible (e.g., exclude dietary and housekeeping staff; medically essential personnel would assume these responsibilities).
  - As much as possible, while ensuring patient safety, reduce room entries by HCP providing direct patient care.
    » Bundle activities while in the room with patients.
    » Use out-of-room monitoring, phone and intercom communication, and self-administration of oral medications.
  - Decrease the length of hospital stays for medically stable patients with COVID-19.
- Follow OHA’s COVID-19 [Guidance on Screening and Visitation at Acute Health Care Facilities](#).
- During large, sustained surge in healthcare need or severe shortage of PPE, deferral of elective surgeries and procedures could be considered
- **Source control**
  - Use of face masks for suspected COVID-19 patients before they are placed in an AIIR or private room.
  - Consider universal masking for all staff, patients, and visitors to healthcare facilities.
- **Cohorting patients**
  - Cohort patients confirmed to have the same infection that requires transmission-based precaution (e.g., COVID-19) to facilitate extended PPE use.
  - Patients co-infected with other pathogens that require transmission-based precautions (e.g., influenza, RSV) should not be placed in the same room with patients only infected with SARS-CoV-2 (i.e., COVID-19).
- **Cohorting HCP**
  - Assign designated teams to provide care for all patients with suspected or confirmed COVID-19 to facilitate extended PPE use.
- **Enhance training on indications for use of each PPE type. Training must be provided to healthcare providers on how to properly use and maintain PPE according to the [Oregon OSHA PPE standard](#).**
Tiered approach to optimizing PPE

Tier 1 strategies
Consider implementing these strategies if PPE supply and rate of use make conventional capacity strategies (Table 1) unsustainable.

- Use of unexpired, medical-grade PPE that follows FDA regulations obtained through usual vendors, other healthcare venues, the Strategic National Stockpile, or through State purchase and allocation.

- Use of protective equipment designed for use in other industries and determined by federal regulatory agencies to provide adequate protection in a healthcare setting. For example, FDA issued Emergency Use Authorizations (EUAs) on March 2 and March 27 allowing such use of filtering facepiece respirators (including NIOSH-approved N95s) and of powered air-purifying respirators (PAPRs). [www.fda.gov/media/135763/download](http://www.fda.gov/media/135763/download)

- Use of PPE designed for reprocessing and re-use is an acceptable and, in many cases, preferred strategy, and can preserve PPE supplies. This includes use of goggles, re-processable face shields, launderable gowns, and re-processable alternatives to N95 respirators (e.g., elastomeric or powered air-purifying respirators [PAPRs]) after appropriate reprocessing.

- PPE should be changed with every patient encounter, with following exceptions:
  - Reuse of re-processable PPE devices with appropriate reprocessing between patient use.
  - Extended use of eye and respiratory protection among cohorted COVID-19 patients; however, gloves should be always be changed with every patient encounter (i.e., single use) even in this context. Note: Under Tier 1, a cohort includes only patients with laboratory-confirmed COVID-19. It would not include patients with suspected, non-lab-confirmed illness.
  - Extended use of medical grade face masks by HCP for source control when a facility-wide universal masking policy is in place; mask should be changed when doffed and prior to any care for a patient in droplet precautions.

- Implementation of additional administrative controls applicable to the PPE type to conserve supply (e.g., prioritizing N95 respirators for AGPs or use of face shield over masks or respirators, with reprocessing of face shield between patients).

- Given the potential for future PPE supply shortage, consider collection and storage of doffed single-use respirators for future decontamination when their scarcity could make Tier 1 strategies unsustainable. This pre-emptive measure decreases the risk of having to implement Tier 3 or 4 strategies should severe, sustained supply chain disruptions occur in the future.

- Refer to Table 1 for more details.
**Tier 2 strategies**

*Consider implementing these strategies if PPE supply and rate of use make Tier 1 strategies unsustainable.*

- Use of the same medical grade PPE that follows FDA regulations through the regular authorization process or approved under EUAs as specified in Tier 1.

- Extended use of PPE may be implemented for eye and respiratory protection when providing care exclusively to patients without suspected or confirmed communicable respiratory infection or infection spread through contact transmission.

- Extended use of surgical masks, NIOSH-approved N95 FFRs, and other single-use respirators may be implemented if worn with an impermeable face shield completely covering them, so long as care does not involve an AGP for a patient with suspected or confirmed communicable respiratory illness. They should be removed and discarded if they become visibly soiled or damaged, or after use during an AGP for a patient with suspected or confirmed COVID or other serious infectious respiratory disease. Otherwise, they can be doffed and decontaminated as mentioned below.

- Use of decontamination systems approved by FDA under EUAs to decontaminate compatible NIOSH-approved N95 or N95-equivalent respirators for subsequent limited re-use. (See Re-use of PPE Implementation Guidance for more details). At this time, several decontamination systems have received EUAs. NIOSH-approved respirators only retain their NIOSH approval status after decontamination if the respirator manufacturer permits decontamination with the specific system and cycle parameters used. For up-to-date information on FDA-approved decontamination systems, refer to FDA’s EUA information on PPE and related devices.

- Implementation of additional engineering or administrative controls applicable to the PPE type (e.g., extended use of secondary coverage of N95 respirator, preferably with a face shield, to prevent droplet spray contamination of the respirator).

- Refer to Table 1 for more details.

**Tier 3 strategies**

*Consider implementing these strategies if PPE supply and rate of use make Tier 2 strategies unsustainable.*

- Use of PPE beyond the manufacturer-designated shelf life. (See Expired PPE implementation guidance below for more details.) CDC recommends against use of N95s beyond the manufacturer-designated shelf life in surgical settings.

- Use of respirators and other PPE that meet standards in other countries and have been approved for use in the healthcare setting under an FDA-issued emergency use authorization. NIOSH has raised concerns that it is difficult to pass a fit test and perform successful seal checks with N95 and other single-use respirators that have
ear loops rather than over-the-head straps. A number of these products are on the FDA-approved list. We provide NIOSH’s concern for your consideration in deciding whether or not to obtain and use this type of respirator.

- Extended use of gowns can be considered when caring for a cohort of patients with the same, laboratory-confirmed pathogen. However, they must be doffed if used during care of a patient with known or suspected MDRO infection or colonization.

- Re-use of disposable face shields with appropriate reprocessing between patient use, limited re-use of face masks, and limited re-use of N95 or N95-equivalent respirators designed for single use without decontamination or with decontamination using a decontamination system without an FDA EUA. (See Extended use of PPE implementation guidance and Re-use of PPE implementation guidance below for more details.) Refer to CDC’s Decontamination and Re-use of Filtering Facepiece Respirators for summary of decontamination methods, effect on device performance, and use recommendations after decontamination. Per CDC, ethylene oxide (EtO) is not recommended as a decontamination method, as it may be harmful to the wearer.

- Implementation of additional administrative controls applicable to the PPE type to conserve supply (e.g., prioritize non-sterile gloves for activities resulting in contact with hazardous substances, including blood and bodily fluids, such as wound care and AGPs).

- Refer to Table 1 for more details.

**Note:** Gloves must still be changed with every patient encounter.

**Tier 4 strategies**

*Consider implementing these strategies if PPE supply and rate of use make Tier 3 strategies unsustainable.*

Tier 4 strategies should only be implemented when all local and regional resources have been exhausted. Note: If healthcare facilities anticipate the need for or emergently begin to use Tier 4 strategies, the local public health authority should be notified.

- If no medical grade PPE are available, see:
  - CDC’s [Considerations for Selecting Protective Clothing used in Healthcare for Protection against Microorganisms in Blood and Body Fluids](https://www.cdc.gov/coronavirus/2019-ncov/hcp/protective-clothing.html) for gowns and coveralls.
  - [OSHA](https://www.osha.gov) standards and recommendations for protective clothing based on protective properties.

- Use of re-purposed items as PPE. Examples include safety glasses that cover side of eyes, laboratory coats, patient gowns, disposable, impermeable aprons, combination of clothing.
Use of self-made or locally produced PPE with characteristics that suggest efficacy in providing necessary level of infection prevention. Examples include:

- **Eye protection**: impermeable, transparent shield with sufficient coverage of the face and side of eyes, fashioned from plastic beverage bottles or other clear plastic
- **Masks**: multi-ply, tightly woven material that is reasonably moisture resistant, such as GORE-TEX, sterilization wrap (typically used to wrap surgical instruments to maintain sterility), or material from furnace filters or vacuum cleaner bags might be used to produce masks that completely cover the nose, mouth, and chin
- **Gowns**: use multi-ply or moisture-resistant material designed to cover all clothing

Various durations of use can be implemented, depending on PPE type and device. (See Table 1.)

Implementation of additional engineering and administrative controls applicable to the PPE type (e.g., use of expedient patient isolation room approach, designate convalescent HCP for provision of care to known or suspected COVID-19 patients). The latter example should be evaluated as an option based on the most up-to-date science regarding the duration and degree of protection conferred by past infection.

Refer to Table 1 for more details.
## Table 1. Personal Protective Equipment Usage Guideline

<table>
<thead>
<tr>
<th>Conventional Capacity Strategies</th>
<th>Equipment Specifications</th>
<th>Face Masks</th>
<th>N95 Respirators or Approved Equivalent*</th>
<th>Isolation Gowns</th>
<th>Gloves</th>
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<tbody>
<tr>
<td><strong>Baseline (Pre-Pandemic)</strong></td>
<td><strong>Equipment Specifications</strong></td>
<td>Medical grade, FDA-cleared</td>
<td>Medical grade; must be FDA-cleared for activities where splashes and sprays are anticipated</td>
<td>Medical grade, NIOSH-approved, fit-tested†</td>
<td>Medical grade, conforms to U.S. or international standards, or NIOSH specifications</td>
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<tr>
<td>Duration</td>
<td>Single use</td>
<td>Single use</td>
<td>Single use</td>
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<tr>
<td><strong>General Considerations</strong></td>
<td>● Prioritize FDA-cleared surgical masks for activities with anticipated splashes and sprays (e.g., surgical procedures)</td>
<td>● Implement respiratory protection program that complies with Oregon OSHA’s respiratory protection standard [29 CFR 1910.134]†</td>
<td>Prioritize surgical gowns for surgical or other sterile procedures</td>
<td>● Double gloving not recommended for care of suspected or confirmed COVID-19 patients</td>
<td>● See CDC guidance on conventional capacity practices for more details, including prioritization of surgical N95 respirators, and use of alternatives to N95 respirators (e.g., elastomeric half-mask and full facepiece air purifying respirators, PAPRs)</td>
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<td>Contingency Capacity Strategies</td>
<td>Tier 1</td>
<td>Equipment Specifications</td>
<td>Face Masks</td>
<td>N95 Respirators or Approved Equivalent*</td>
<td>Isolation Gowns</td>
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| **Eye Protection**              |        | ● Medical grade, authorized by FDA under [EUA](https://www.fda.gov)  
● Reprocessable face shields or goggles preferred  
● Consider preferential use of PAPRs or full-face elastomeric respirators, which have built-in eye protection  | Medical grade; must be FDA-cleared for activities where splashes and sprays are anticipated  | ● Medical grade, NIOSH-approved, fit-tested†  
● Industrial grade, approved by FDA under [EUA](https://www.fda.gov), fit-tested  
● [Alternatives to N95 respirators, NIOSH-approved*](https://www.cdc.gov), fit-tested  | Medical grade, conforms to U.S. or [international standards](https://www.cdc.gov), or [NIOSH specifications](https://www.niosh.gov) (laundry gowns or coveralls preferred)  | Medical grade, FDA-cleared or conforming to other U.S. and [international standards](https://www.cdc.gov)  |
| **Duration**                    | Single use  
Exceptions: extended use in care of cohorted COVID-19 patients; re-use of goggles and reprocessable face shields with appropriate reprocessing between patient use.  | Single use  
Exceptions: extended use in care of cohorted COVID-19 patients; extended use by HCP for source control in the context of a universal masking policy; mask should be changed when doffed and prior to any care for a patient in droplet precautions.  | Single use  
Exceptions: extended use in care of cohorted patients with the same infection requiring airborne precautions; re-use of reprocessable alternatives to N95 respirators such as elastomeric or powered air purifying respirators with appropriate cartridges.  | Single use | Single use |
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<tr>
<th>Contingency Capacity Strategies</th>
<th>Tier 1</th>
<th>Eye Protection</th>
<th>Face Masks</th>
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<th>Isolation Gowns</th>
<th>Gloves</th>
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<tr>
<td>Additional Engineering or</td>
<td></td>
<td>● Control access by visitors (e.g., remove from public areas, but made available for symptomatic patients)</td>
<td>● Control access by visitors (e.g., remove from public areas, but made available for symptomatic patients)</td>
<td>● Prioritize for AGPs</td>
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<td>● Use expired (when indicated by manufacturer) gloves for training</td>
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<td>Administrative Controls</td>
<td></td>
<td>● Preserve medical grade face masks by supplying visitors and asymptomatic patients with cloth face coverings for source control</td>
<td>● Preserve medical grade face masks by supplying visitors and asymptomatic patients with cloth face coverings for source control</td>
<td>● Plan for and implement just-in-time fit testing in context of a full respiratory protection program†</td>
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<td></td>
<td>● Preserve medical grade face masks by supplying visitors and asymptomatic patients with cloth face coverings for source control</td>
<td>● Consider changing from quantitative to qualitative fit testing</td>
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<td>● <strong>Temporarily suspend annual fit testing</strong> (initial fit testing as well as additional actions are required)†</td>
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<td></td>
<td>● Retain and reserve expired N95 respirators for training and fit testing</td>
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<td>Tier 2</td>
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<td><strong>Duration</strong></td>
<td>Extended use also an option when providing care exclusively to patients without suspected or confirmed communicable respiratory infection or infection spread through contact transmission.</td>
<td>Extended use also an option when providing care exclusively to patients without suspected or confirmed communicable respiratory infection or infection spread through contact transmission. Use with an impermeable face shield completely covering the face mask is strongly recommended.</td>
<td>● Extended use (exception: single use only for AGP in care of patient with suspected/confirmed communicable respiratory infection) ● <strong>Limited</strong> re-use of single-use respirators after reprocessing by a compatible decontamination system <a href="https://www.fda.gov">approved by FDA under an EUA</a>, NIOSH-approved respirators only retain their NIOSH- approved status after decontamination if manufacturer permits decontamination with the specific system and cycle parameters used; may only be used after</td>
<td>Same as Tier 1</td>
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<td>Tier 2</td>
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<td>AGPs if manufacturer or third-party guidance or procedures available</td>
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<td>• <strong>Limited</strong> Re-use of single-use respirator during the same shift (e.g., after lunch break) if respirator is completely covered by impermeable face shield, hands are cleaned before and after donning and doffing, and respirator is not damaged or visibly soiled.</td>
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<td>• Rotating re-use of multiple respirators assigned to a given HCP with at least a 5-day interval between uses of a given respirator</td>
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<td>Tier 2 Additional Engineering or Administrative Controls</td>
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<td>• Same as Tier 1, and</td>
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<td>● Restrict use to HCP only (homemade masks, tissues, or other barriers can be used by patients for source control)</td>
<td>● Same as Tier 1, and</td>
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<td>● Use secondary barrier (cleanable face shield preferred) and masking patients to prevent droplet contamination of N95 respirators</td>
<td>Same as Tier 1</td>
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<td>Crisis Capacity Strategies</td>
<td>Tier 3</td>
<td>Equipment Specifications</td>
<td>Eye Protection</td>
<td>Face Masks</td>
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|                            |        | Same as Tiers 1 and 2, but expired product can also be used. | Same as Tiers 1 and 2, but expired product can also be used. (must be FDA-cleared for activities where splashes and sprays are anticipated) | Same as Tiers 1 and 2, but expired product can also be used. (must be FDA-cleared for activities where splashes and sprays are anticipated) | ● Medical grade, NIOSH-approved, expired but evaluated by NIOSH, fit-tested; CDC recommends against use of N95s beyond manufacturer-designated shelf life in surgical settings.  
● Medical grade, approved under standards used in other countries similar to NIOSH (see CDC guidance regarding use limitations during AGPs)  
● Medical grade, NIOSH approved, expired but not evaluated by NIOSH, fit-tested | Same as Tiers 1 and 2, but expired product can also be used; extended use for cohorted patients with lab-confirmed COVID-19 | Same as Tiers 1 and 2, but expired (when indicated by manufacturer) – expired sterile gloves should not be used for surgical or other sterile procedures |
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| Tier 3                    | ● Same as Tier 2 and  
● Re-use of disposable face shields after appropriate reprocessing | ● Same as Tier 2 and  
● Limited re-use | Limited re-use, other than as described in Tier 2 above, without decontamination or decontamination using a system without an FDA EUAΔ | Extended use when caring for cohort with lab-confirmed COVID or other shared pathogen; gown must be doffed if used during care of patient with known or suspected MDRO infection/colonization | Same as Tiers 1 and 2 |
| Duration                  |                |            |                                        |                  |        |
| Additional Engineering or Administrative Controls | ● Prioritize for selected activities (e.g., splashes and sprays anticipated as in AGPs, prolonged face-to-face contact with potentially infectious patient) | ● Same as Tier 2, and  
● Prioritize for HCP in procedures with high risk of transmission (e.g., emergent surgeries or procedures, activities with anticipated splashes and sprays, activities requiring prolonged face-to-face or close contact with potentially infectious patient, AGPs if respirators no longer available) | ● Prioritize by activity type (e.g., AGPs) and based on distance from patient and use of source control  
● If no respirators are available, whenever possible, AGPs should be deferred or patients should be transferred to facilities where adequate respiratory protection is available | ● Prioritize for selected activities (e.g., splashes and sprays anticipated as in AGPs, high-contact patient care activities including bathing or showering, changing linens, wound care) | ● Same as Tiers 1 and 2, and  
● Prioritize non-sterile gloves for activities resulting in contact with hazardous substances, including blood and bodily fluids (e.g., wound care, AGPs) |
<table>
<thead>
<tr>
<th>Crisis Capacity Strategies</th>
<th>Equipment Specifications</th>
<th>Eye Protection</th>
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<tbody>
<tr>
<td>Tier 4 (medical grade PPE scarce)</td>
<td><strong>Equipment Specifications</strong></td>
<td>Consider safety glasses (e.g., trauma glasses), swim masks, or other protection that extends to cover side of eyes</td>
<td>● Face shield covering chin and sides of face or in combination with homemade mask</td>
<td>● AGPs if respirators not available; transfer to another facility with necessary PPE isn’t feasible</td>
<td>● Launderable gown (see duration below)</td>
<td>● Same as Tier 1, 2, or 3</td>
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<tr>
<td><strong>Duration</strong></td>
<td>● Extended use</td>
<td>● Extended use</td>
<td>● Re-use after appropriate reprocessing</td>
<td>● Re-use of launderable isolation gowns without laundering in between</td>
<td>● Re-use of gown alternatives that have not been evaluated as effective (e.g., lab coats, patient gowns, disposable aprons, combinations of clothing)</td>
<td>● Non-healthcare glove alternatives (e.g., food service or industrial chemical resistance gloves) only for situations with no exposure to pathogens</td>
</tr>
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<td><strong>Duration</strong></td>
<td>● Extended use</td>
<td>● Extended use</td>
<td>● Re-use after appropriate reprocessing (face shields) and laundering (homemade masks)</td>
<td>● Re-use of launderable isolation gowns without laundering in between</td>
<td>● Single use or re-use of gown alternatives</td>
<td>● Extended use of disposable medical grade gloves (exception: Dispose of gloves after direct care of patients with MDRO infection (e.g., MRSA, VRE, ESBL-producing organisms); ● Single (not extended) use of non-healthcare gloves</td>
</tr>
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<td>● Use <a href="#"><em>expedient patient isolation room</em></a> approach</td>
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<td>● Use <a href="#"><em>expedient patient isolation room</em></a> approach</td>
<td>● Use <a href="#"><em>ventilated headboards</em></a> in combination with HEPA filter units</td>
</tr>
</tbody>
</table>

* Approved equivalents for an N95 include NIOSH-approved P95 and R95 FFRs and others approved for medical use through an FDA EUA, such as respirators providing greater respiratory protection including elastomeric half- and full-face respirators or PAPRs with appropriate cartridges. However, use of respirators with an exhalation valve is not recommended for use during surgical procedures due to concerns about potential contamination to the procedural field.

† In any medical, dental, or other facility in which respirators are used for a procedure, the facility must establish and maintain a [respiratory protection program](#) for its employees.

∆ Decontaminating with ethylene oxide is discouraged at this time due to potential harmful effects to the wearer.
Extended Use of PPE Implementation Guidance

Extended use is preferred over re-use on the assumption that it is safer for HCP to leave their respiratory protection and eye protection in place, to reduce the risk of self-contamination through frequent donning and doffing.

1. Eye protection
   - Should not be touched or adjusted; hand hygiene to be performed immediately if touched or adjusted by HCP
   - Should be removed and reprocessed when visibly soiled or difficult to see through
   - Should be discarded if damaged
   - Should be removed when leaving patient care area
     » If a disposable face shield is reprocessed, it should be dedicated to one HCP and reprocessed whenever it is visibly soiled or removed (e.g., when leaving the isolation area) prior to putting it back on. See protocol for removing and reprocessing eye protection.

2. Gowns
   - Should only be used in Tier 3 for the serial care of cohorts of patients with laboratory-confirmed COVID-19 another shared pathogen
     » Should only be considered if there is no infection or colonization among patients with another pathogen transmitted by contact (e.g., *Clostridioides difficile*, multidrug-resistant organism)
     » Should be removed and discarded or laundered as per usual practices when visibly soiled

3. Face masks
   - Should be removed and discarded if soiled, damaged, or hard to breathe through
   - Should not be touched or adjusted during care; hand hygiene to be performed immediately if touched or adjusted by HCP
   - HCP should leave patient care area if they need to remove the face mask
   - Should not be worn for multiple shifts
   - Should be doffed before meals and restroom breaks. Re-use during the same shift is acceptable (e.g., after lunch break) if hands are cleaned before and after donning and doffing, and face mask is not damaged or visibly soiled

4. N95 filtering facepiece respirators (NIOSH-approved)
   - Well suited to situations wherein multiple patients with the same infectious disease diagnosis requiring airborne precautions are cohorted, e.g., tuberculosis, varicella, measles, or other infectious diseases where use of an N95 respirator or higher is recommended
   - Maximum recommended extended use period is 12 contiguous hours
   - Re-use during the same shift is acceptable (e.g., after lunch break) if the respirator is completely covered by an impermeable face shield, hands are
cleaned before and after donning and doffing, the respirator is not damaged or visibly soiled, and it continues to pass a seal check.

- If touching respirator is necessary for comfort, to maintain fit, or for doffing before meals or restroom breaks, hand hygiene should be performed before and after touching or adjusting respirator. HCP should leave the patient care area if they need to adjust/remove the respirator.

- Should be discarded:
  - Following use during AGP for a patient with suspected or confirmed communicable respiratory infection
  - If visibly contaminated with blood, respiratory or nasal secretions, or other bodily fluids from patients
  - If obviously damaged or the respirator becomes hard to breathe through
  - Following close contact with, or exit from, the care area of any patient co-infected with an infectious disease requiring contact precautions

- Consider collection for future decontamination and re-use, as appropriate, if no criteria requiring disposal are met

- Can be combined with re-use using either of the following methods:
  - Decontamination using an FDA-approved process that is compatible with the model of respirator, according to the manufacturer.
  - Rotating use of multiple respirators assigned to a given HCP with at least a 5-day interval between uses of a given respirator. As noted above, respirator should be discarded after use during an AGP for a patient with suspected or confirmed communicable respiratory infection

5. Gloves

- To be used only in a Tier 4/emergent scenario

- Applicable only to disposable medical grade gloves, and not to non-healthcare glove alternatives

- Most easily implemented when patients are cohort based on the same confirmed infectious disease diagnosis (e.g., confirmed COVID-19) in a shared or adjacent location

- Must be sanitized between patients within cohort to prevent cross transmission of any other pathogens from patient to patient, and at other intervals where gloves would normally be changed (e.g., when moving from “dirty” to “clean” task); refer to CDC guidance regarding methods for performing hand hygiene of gloved hands

- Should always be discarded after:
  - Visible soiling or contamination with blood, respiratory or nasal secretions, or other bodily fluids
  - Any signs of damage or degradation
  - Maximum of four hours of continuous use
  - Doffing (i.e., re-use should never occur)

- Hand hygiene should be performed after removing gloves for any reason
Re-use of PPE Implementation Guidance

1. Eye protection
   - If a disposable face shield is reprocessed, it should be dedicated to one HCP and reprocessed whenever it is visibly soiled or removed (e.g., when leaving the isolation area) prior to putting it back on. See protocol for removing and reprocessing eye protection.

2. Gowns
   - In Tier 4, consider limiting re-use without laundering in between where the gown is used as part of standard precautions to protect HCP from a splash, as the risk may be lower. The risk of re-use without laundering when caring for suspected or confirmed COVID-19 is unclear. Note that this strategy aims at minimizing exposure to HCP and does not necessarily prevent transmission of pathogens between patients.
   - Should be removed and laundered if becomes visibly soiled

3. Face masks
   - Ideally reserved for activities with low transmission risk such as dispensing medications or other activities that do not involve close, direct contact with patients in droplet precautions.
   - Not all face masks can be re-used: single-use masks with elastic ear hooks may be more suitable for re-use than those fastened via ties.
   - Should not be touched or adjusted during care; hand hygiene to be performed immediately if touched or adjusted by HCP
   - Removal and replacement of mask should be performed in a careful and deliberate manner
   - Should be removed and discarded if soiled, damaged, or hard to breathe through
   - HCP should leave patient care area if they need to remove the face mask. If possible, face masks should be carefully folded with outer surface held inward and against itself to reduce contact with outer surface during storage. Masks can be stored between uses in a clean sealable paper bag or breathable container.
   - Re-use should be limited to one shift.

4. N95 filtering facepiece respirators (NIOSH-approved)
   - Prioritize re-use for pathogens for which contact transmission is not a concern (e.g., tuberculosis)
   - The maximum number of re-uses (i.e., doffs) should be in accordance with manufacturer guidance or, if not available, be no more than five uses per device.
   - Should not be shared by multiple HCP unless decontaminated in between using a system with an FDA-approved EUA that is compatible with manufacturer recommendations.
   - Place used respirators in a designated storage area, or keep them in a clean, breathable container such as a paper bag between uses. To minimize potential cross-contamination, store respirators so that they do not touch each other, with
the person using the respirator clearly identified by labeling storage containers or respirators themselves (e.g., on the straps). Storage containers should be disposed of or cleaned regularly.

- If touching respirator is necessary for comfort or to maintain fit, hand hygiene should be performed before and after touching or adjusting respirator.
- Avoid touching inside of respirator. If inadvertent contact is made with the inside of the respirator, discard the respirator and perform hand hygiene as described above.
- Use a pair of clean (non-sterile) gloves when donning a used N95 respirator and performing a user seal check, as re-use may change the shape of a disposable respirator and affect fit. Discard gloves after the N95 respirator is donned and any adjustments are made to ensure the respirator is sitting comfortably with a good seal.
- Should be discarded:
  - Following use during AGP for a patient with suspected or confirmed communicable respiratory infection
  - If contaminated with blood, respiratory or nasal secretions, or other bodily fluids from patients
  - Following close contact with, or exit from, the care area of any patient co-infected with an infectious disease requiring contact precautions
  - If obviously damaged, seal check is unsuccessful, or if hard to breathe through
  - If respirator has been used for five shifts or five separate procedures
- Consider collection for compatible decontamination and re-use, as appropriate, if no criteria requiring disposal are met
- Additional guidance on potential methods can be found here.

5. Gloves

- Disposable, medical grade gloves should never be re-used.

**Expired PPE Use Implementation Guidance**

If no date is indicated on the device or packaging, facilities should contact the manufacturer.

Reserve for settings where there is a lower risk of transmission (e.g., non-surgical); prioritize the use of unexpired FDA-cleared surgical masks and NIOSH-approved respirators, as appropriate, for healthcare providers in procedures where there is high risk of transmission to the healthcare provider or the patient due to exposure to blood, respiratory secretions, or other body fluids, including AGPs.

The user should visually inspect the device prior to use and, if there are concerns (e.g., degraded materials or visible tears), discard the product.
**Acronyms:**

- AGP: aerosol generating procedure
- EUA: Emergency Use Authorization
- HCP: healthcare personnel
- MDRO: multidrug-resistant organism
- NIOSH: National Institute for Occupational Safety and Health
- OSHA: Occupational Safety and Health Administration
- PAPR: powered air purifying respirator

**References**


**Document accessibility:** For individuals with disabilities or individuals who speak a language other than English, OHA can provide information in alternate formats such as translations, large print, or braille. Contact Mavel Morales at 1-844-882-7889, 711 TTY or OHA.ADAModifications@dhsoha.state.or.us.