1. DISEASE REPORTING

1.1 Purpose of Reporting and Surveillance
To monitor severe illness and deaths associated with COVID-19 in Oregon, inform efforts to reduce transmission to others, promote health equity and better understand the epidemiology of the disease.

1.2 Physician Reporting Requirements
Healthcare providers are required to report, within 1 working day,
- All pediatric deaths, among persons with COVID-19.
- All cases of Multisystem Inflammatory Syndrome in Children (MIS-C; §3.5).
1.3 Local Public Health Authority Responsibilities

1. Educate and consult with local providers and facilities to promote compliance with outbreak reporting, isolation, and infection-control procedures.

2. Investigate outbreaks of COVID-19 associated with high-consequence settings (see Respiratory Disease Outbreak Investigative Guidelines).

3. Consult with the Oregon Health Authority (OHA) as needed about patient isolation and protection of contacts, including healthcare personnel, and about strategies for public health response, testing, and access to therapeutics.

1.4 State Public Health Division Responsibilities

1. Update LPHAs on changes to criteria for investigation (e.g., through HAN, multijurisdictional conference calls, etc.).

2. Support investigation and response to high-consequence outbreaks.

3. Advise LPHA, Tribal, and private-sector health professionals concerning:
   - Isolation of cases and symptomatic persons in high-consequence congregate settings;
   - Protection of healthcare personnel;
   - Diagnostic evaluation;
   - Required reporting and surveillance activities.

2. THE DISEASE AND ITS EPIDEMIOLOGY

2.1 Etiologic Agent

Coronaviruses are enveloped, single-stranded RNA viruses. With the notable exceptions of SARS-CoV and MERS-CoV, most human coronaviruses typically cause mild upper respiratory illness. SARS-CoV-2, the coronavirus causing COVID-19, was first identified in Wuhan, China in December 2019 among patients with severe respiratory illness and pneumonia and has spread around the globe through person-to-person transmission. Genomic sequencing of isolates demonstrates that SARS-CoV-2 is a betacoronavirus with roughly 80% genome identity with SARS-CoV and 50% with MERS-CoV. Variants with demonstrated or suspected characteristics of public health importance such as increased transmissibility, severity, vaccine resistance or diagnostic or therapeutic escape have been labeled ‘variants of concern’ or ‘variants of interest,’ respectively.

2.2 Description of Illness

Symptoms are non-specific and may include fever (temperature of ≥100.4°F or 38.0°C), chills, sore throat, cough, shortness of breath (dyspnea), myalgia, fatigue, loss of smell (anosmia) or taste (ageusia), headache, nasal congestion, runny nose (rhinorrhea), nausea, vomiting and diarrhea. A significant proportion of cases are asymptomatic. Pneumonia typically presents with patchy, multilobar infiltrates on chest X-ray. Reported complications have included but are not limited to acute respiratory distress syndrome, cardiac events, and death.

COVID-19-associated multisystem inflammatory syndrome in children (MIS-C) is defined by fever, multisystem involvement (cardiac, renal, respiratory, hematologic, gastrointestinal, dermatologic or neurologic), laboratory evidence of inflammation and recent SARS-CoV-2 infection.

COVID-19 associated multisystem inflammatory syndrome in adults (MIS-A) is defined by fever, multisystem involvement which must include severe cardiac illness or the combination of rash and conjunctivitis, laboratory evidence of inflammation and recent SARS-CoV-2 infection.
infection.

2.3 Reservoirs
Members of the coronavirus family are common in many different species of animals, including camels, cattle, cats, and bats. Rarely, animal coronaviruses can infect people and then spread from person to person, as occurred with MERS-CoV and SARS-CoV. The frequency with which the COVID-19 virus is transmitted from its original animal reservoir(s) to humans is unknown, but such events are probably rare. Since the start of the COVID-19 pandemic, SARS-CoV-2 infection has been confirmed in many animal species; whether any (other than humans) will become important reservoirs is unknown.

2.4 Sources and Routes of Transmission
This virus probably originated from an animal source and was followed by rapid person-to-person spread. Person-to-person transmission occurs primarily from respiratory droplets and aerosols produced when an infected person coughs, sneezes, breathes or speaks. Although coronaviruses may persist on surfaces for up to several days, surfaces and fomites are probably not significant routes of transmission. Virus is detectable in the urine and feces of infected persons, and replication-competent virus has been demonstrated. While no concrete evidence exists for the fecal-oral spread of SARS-CoV-2, one study has demonstrated probable evidence of fecal-aerosol transmission of SARS-CoV-2. Transmission from blood or other body fluids has not been identified.

2.5 Incubation Period
Typically, 3–6 (range, 2–14) days.

2.6 Period of Communicability
Most SARS-CoV-2 transmission occurs early in the course of illness, generally in the 2 days before symptom onset and during the 2–3 days thereafter. Various studies pre-dating the emergence of the Omicron variant indicated an infectious period ranging from 3–10 days after symptom onset. Patients with more severe illness—i.e., hospitalized or severely immunocompromised—may shed replication-competent virus for longer periods of time.

2.7 Treatment and Prevention

2.7.1 Vaccines against COVID-19
COVID-19 vaccination recommendations continue to change. The following vaccines have FDA Emergency Use Authorization or full approval for use in the United States:

- Moderna mRNA vaccine
- Pfizer-BioNTech mRNA vaccine
- Novavax adjuvanted protein subunit vaccine

See CDC's Overview of COVID-19 Vaccines and CDC's Use of COVID-19 Vaccines in the United States for up-to-date details regarding approved age ranges and recommended vaccination schedules.

Individuals are considered up to date with their COVID-19 vaccinations when they have received all recommended primary series and booster doses. Individuals who were vaccinated outside the United States and have completed the primary series (1 or 2 doses) of a vaccine accepted in the United States and are not yet eligible for a booster are considered up to date. Visit CDC's Stay up to Date with COVID-19 Vaccines webpage for more information.
Each of the COVID-19 vaccines is contraindicated in patients who have had a severe allergic reaction (e.g., anaphylaxis) to a previous dose of that vaccine or to any of its components.

### 2.7.2 Treatment
Please refer to the National Institutes of Health COVID-19 Treatment Guidelines for the most current information regarding COVID-19 therapeutics.

### 3.0 CASE AND CLINICAL DEFINITIONS

**Note:** Oregon no longer requires or recommends individual-level surveillance for SARS-CoV-2, a common and endemic pathogen. These case definitions inform historical classification.

#### 3.1 Close Contact

A close contact is a person with an epidemiologic exposure to a person with confirmed or presumptive COVID-19. The exposure may be close contact with a confirmed or presumptive case—being within 6 feet of a COVID-19 case for ≥15 minutes\(^1\)—or contact with their infectious secretions or clinical specimens.

Note: This definition only applies to persons who have close contact with a confirmed or presumptive case. Persons who have an epidemiologic exposure to a close contact do not meet this definition.

#### 3.2 Suspect Case

A suspect case is a person with:

- New onset of symptoms consistent with COVID-19, including fever or chills, cough, shortness of breath or difficulty breathing, fatigue, muscle or body aches, headache, new loss of taste or smell, sore throat, nasal congestion, rhinorrhea, nausea, vomiting, or diarrhea;

  AND

- No more likely alternative diagnosis

  Note: This includes people who had close contact with a presumptive\(^2\) case and have an acute illness featuring at least two of the following: shortness of breath, cough, fever, new loss of smell or taste, radiographic evidence of viral pneumonia.

  OR

- A test result that, in combination with their symptoms, does not meet the definition of a confirmed or presumptive case, including:
  - An indeterminate reverse transcriptase polymerase chain reaction (RT-PCR), other nucleic acid amplification test (NAAT)\(^3\) or antigen result;
  - A close contact who is getting tested

These criteria are for epidemiologic classification and are not meant to direct clinician testing or clinical care.

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\(^1\) This time is cumulative over a 24-hour period and does not have to be consecutive.

\(^2\) If a contact of a presumptive case has symptoms consistent with COVID-19 but neither the contact nor the case has tested positive, the contact remains a suspect case.

\(^3\) e.g., a polymerase chain reaction (PCR) test.
3.3 Confirmed Case
A confirmed case is someone who tests positive using a laboratory-based FDA Emergency Use Authorized (EUA) diagnostic test. Any positive result from a laboratory-based RT-PCR, other NAAT, or antigen platform developed under an FDA EUA, even if conducted as asymptomatic screening, is considered a positive result. A negative follow-up test does not negate the initial positive test.

3.4 Presumptive Case
A presumptive case is a person without a positive laboratory-based COVID-19 RT-PCR, NAAT, or antigen test result, with:
- An acute illness featuring at least two of the following: shortness of breath, cough, fever, new loss of smell or taste, radiographic evidence of viral pneumonia; AND
- No more likely alternative diagnosis; AND
- Within the 14 days before illness onset:
  - Had close contact with a confirmed case; OR
  - Lived in the same household or congregate setting as a confirmed case; OR
  - Is identified as having been exposed in an outbreak OR
- A COVID-19-specific ICD-10 code listed as a primary or contributing cause of death on a death certificate.
  OR
- A person with a positive test result from an at-home test kit

3.5 Multisystem Inflammatory Syndrome in Children (MIS-C)
- An individual aged <21 years presenting with fever, laboratory evidence of systemic inflammation, and evidence of clinically severe illness requiring hospitalization, with new onset manifestations in at least 2 of the following categories: cardiac involvement, mucocutaneous involvement, shock, gastrointestinal involvement, or hematologic involvement; AND
- No alternative more likely diagnosis; AND
- Evidence for current or recent SARS-CoV-2 infection by RT-PCR, NAAT, serology, or antigen testing; or COVID-19 exposure within the 60 days prior to the onset of symptoms

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4 Even with a negative test, a person with an identified epi-link, compatible symptoms, and no more likely diagnosis is still considered a presumptive case.
5 Fever can be objective (≥100.4°F) or subjective.
6 Fever may be objective (≥100.4°F) or subjective.
7 Indicated by C-reactive protein of ≥3.0 mg/dL (30mg/L).
If a person is diagnosed with MIS-C, create a suspect MIS-C case in Orpheus and route to Nasreen Abdullah for confirmation. Contact tracing is not indicated for MIS-C. **Consider MIS-C in any pediatric death with evidence of SARS-CoV-2 infection.**

### 3.6 Multisystem Inflammatory Syndrome in Adults (MIS-A)

- An individual ≥21 years with fever, laboratory evidence of inflammation (at least two of the following must be elevated: CRP, ferritin, IL-6, ESR, procalcitonin), and evidence of clinically severe illness requiring hospitalization with at least 3 of the following clinical criteria, one of which must be a primary clinical criterion
  - **Primary clinical criteria**
    - Severe cardiac illness (e.g., myocarditis, pericarditis, coronary artery dilatation, new ventricular dysfunction, 2nd- or 3rd-degree AV block, or ventricular tachycardia)
    - Rash or
    - Non-purulent conjunctivitis
  - **Secondary clinical criteria**
    - New-onset neurologic signs and symptoms (e.g., encephalopathy, seizures, meningeal signs, peripheral neuropathy)
    - Shock or hypotension not attributable to medical therapy
    - Abdominal pain, vomiting or diarrhea
    - Thrombocytopenia

**AND**

- No alternative more likely diagnosis;
**AND**

- Evidence for current or recent SARS-CoV-2 infection by RT-PCR, NAAT, serology, or antigen testing; or COVID-19 exposure within the 28 days prior to the onset of symptoms.

If a person is diagnosed with MIS-A, create a suspect MIS-A case in Orpheus and route to Nasreen Abdullah for confirmation. Contact tracing is not indicated for MIS-A.

### 4.0 LABORATORY TESTING

#### 4.1 Testing at the Oregon State Public Health Laboratory

Testing through the Oregon State Public Health Laboratory (OSPHL) must be approved by the Regional Epidemiologist supporting the outbreak. The [Criteria for COVID-19 Testing at OSPHL](https://www.oregon.gov/oha/PH/diseases/covid/pages/criteria-for-covid-testing-at-osphl.aspx) provide general information about testing policies and targeted populations tested at OSPHL. Generally, OSPHL testing is prioritized for high-risk individuals and in support of outbreak investigations of public health significance. COVID-19 is an extremely common infection and OSPHL is not able to support testing for all outbreaks. Current guidance for specimen collection, handling, and transport is posted on OSPHL’s [Lab Test Menu](https://www.oregon.gov/oha/PH/diseases/covid/what-testing-ltdl/intl_pages/lab-test-menu.aspx).

Whole genome sequencing for SARS-CoV-2 is available at OSPHL. Please review the [Criteria for Requesting COVID-19 Sequencing at OSPHL](https://www.oregon.gov/oha/PH/diseases/covid/what-testing-ltdl/intl_pages/sequencing-at-osphl/) for details on how to make a request, the approvals process, and required specimen types.

#### 4.2 Collecting Specimens

Specimens should be collected while using proper personal protective equipment (PPE). See CDC’s [healthcare infection control guidance](https://www.cdc.gov/).
For specimen collection that involves an aerosol-generating procedure (§10):
- Using an airborne infection isolation room (AIIR) is ideal, but if one is not available, use a private room and keep the door closed.
- Mask the patient with a surgical facemask during any movement within clinic or facility.
See CDC Guidelines for Collection and Handling Specimens for COVID-19 Testing.

Many common respiratory infections present with symptoms similar to those of COVID-19. If a person tests positive for a common respiratory pathogen, testing for COVID-19 remains indicated, as co-infections may occur.

4.3 Guidance Regarding Serologic Tests
Serologic test results do not currently alter case classifications and serology should not be used for diagnosis of COVID-19 infection.

Except where specifically identified, all references in this guide to a “test” or “testing” refer to RT-PCR, NAAT, or antigen tests and not to serology.

4.4 Guidance Regarding At-Home Test Kits and Point-of-Care Tests
At-home COVID-19 test kits are widely available. Patients with positive test results should be encouraged to follow-up with a medical provider if they have questions, require medical evaluation, or are at increased risk of severe disease and may be eligible for antiviral therapy.

5.0 PREVENTING TRANSMISSION OF COVID-19

CDC’s Respiratory Virus Guidance provides information on how to prevent transmission of respiratory viruses including COVID-19.

6.0 OUTBREAK RESPONSE

Please see the Respiratory Disease Outbreak Investigative Guidelines for additional information.

7.0. COVID-19 IN HIGH-RISK SETTINGS

7.1 Healthcare Settings
The following COVID-19-specific Oregon Administrative Rules remain in effect:
- **OAR 411-061** – COVID-19 vaccination reporting requirements for licensed assisted living, nursing, and residential care facilities
- **OAR 411-060** (under revision) – COVID-19 testing in licensed assisted living, nursing and residential care facilities

For additional healthcare specific guidance, please see:
- Infection Control: Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) | CDC
- Interim Guidance for Managing Healthcare Personnel with SARS-CoV-2 Infection or Exposure to SARS-CoV-2 | CDC
- Strategies to Mitigate Healthcare Personnel Staffing Shortages | CDC
- Best practices for COVID-19 related admissions from hospitals to long-term care facilities | OHA
7.1.1 Quarantine for healthcare personnel, patients, and residents in healthcare settings

In general, quarantine is not recommended for asymptomatic healthcare facility inpatients or residents following an exposure to someone with suspected or confirmed COVID-19, regardless of vaccination status. Additionally, asymptomatic healthcare personnel (HCP) no longer require quarantine following close contact with someone suspected or confirmed COVID-19, regardless of vaccination status.

HCP who have a higher-risk exposure should take additional steps to reduce the risk of spreading COVID-19. Generally, a higher-risk exposure is considered prolonged close contact with a patient, visitor, other HCP, or other members of the community outside of the workplace (e.g., household contacts) with confirmed SARS-CoV-2 and the HCP was not:

- Wearing a respirator (or if wearing a mask, the person with SARS-CoV-2 was not wearing a mask);
- Wearing eye protection if the person with SARS-CoV-2 was not wearing a mask; or
- Wearing all recommended PPE while in the room for an aerosol-generating procedure

Following any close contact with someone with SARS-CoV-2 for asymptomatic patients and residents or following a higher-risk exposure for asymptomatic HCP, the individual should:

- Complete a series of 3 viral tests (i.e., antigen or NAAT) for SARS-CoV-2 if they have not recovered from COVID-19 in the prior 30 days, typically at day 1 (where day of exposure is day 0), day 3 and day 5. That is at least 24 hours after the initial exposure, and if negative 48 hours later and again another 48 hours later if the second test is also negative;
- Wear well-fitting source control;
- Monitor themselves for fever or other symptoms of COVID-19; and
- Stay home when ill or if they test positive for SARS-CoV-2.

Note: Smaller congregate-care facilities (e.g., adult foster homes) may not have routine and consistent access to the testing and specimen-collection resources needed to conduct serial testing as recommended above. Though it is recommended that facilities that house high-risk or medically fragile residents test according to the testing schedule above, AFHs could otherwise follow testing strategies as outlined in CDC’s Additional Information for Community Congregate Living Settings.

HCP who are put under quarantine can return to work, and transmission-based precautions can be lifted for patients:

- After day 7 following the exposure if they remain asymptomatic and all viral testing conducted is negative; or
- After 10 days if testing is not performed and they remain asymptomatic.

Testing recommendations for patients or residents with exposure or during outbreak scenarios can be found in CDC Interim Infection Prevention and Control Recommendations.

Outpatients who have been exposed to COVID-19 should be cared for using appropriate Transmission-Based Precautions.

7.1.2 HCP Work Restriction
HCP with COVID-19 should be restricted from work following the criteria outlined below. After returning to work, HCP should continue to monitor for symptoms and seek re-evaluation if symptoms recur or worsen.

Table. Healthcare personnel restrictions

<table>
<thead>
<tr>
<th>Illness Severity* and Immune Status</th>
<th>Work Restriction Recommendations</th>
<th>Additional Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>No symptoms or mild or moderate illness AND Not immunocompromised</td>
<td>At least 7 days after symptoms began or first positive test if asymptomatic with a negative viral test within 48 hours of returning to work AND At least 24 hours fever free AND Other symptoms improving</td>
<td>10 days if no viral testing completed or test positive on day 5-7 If using antigen test, obtain a negative test on day 5 and day 7</td>
</tr>
<tr>
<td>Severe illness AND Not immunocompromised</td>
<td>At least 10 days and up to 20 days since symptoms began AND At least 24 hours fever free AND Other symptoms improving</td>
<td>Can consider using a test-based strategy in consultation with infectious disease expert to determine when the staff can return to work</td>
</tr>
<tr>
<td>Moderate or severely immunocompromised† AND Any level of symptom severity</td>
<td>Test-based strategy in consultation with infectious disease expert to determine when the staff can return to work</td>
<td>2 consecutive negative tests at least 24 hours apart If symptomatic, should be fever free for at least 24 hours and other symptoms improving to end isolation Retest for SARS-CoV-2 if symptoms return or worsen</td>
</tr>
</tbody>
</table>

† See §10

7.1.3 Contact tracing in healthcare settings
Contact tracing in healthcare settings, though not feasible in all scenarios, may be of benefit in high-risk settings to halt early outbreak spread or when being utilized to guide exposure testing. Formal contact tracing for exposures in healthcare settings may be infeasible and of limited benefit during periods of high transmission or when staffing is insufficient to maintain this work. In these scenarios, healthcare facilities should consider forgoing contact tracing for exposures in a healthcare setting in favor of broad infection control measures (e.g., well-fitting masks, ventilation, frequent symptom monitoring, and readily accessible testing). Additional infection prevention and control recommendations, including more details about universal source control in healthcare settings, are available from the CDC.

Where there is a known healthcare-associated exposure, healthcare facilities, including LTCFs, should conduct a risk assessment of HCW exposures and ensure exposure testing and/or work restrictions are implemented, if needed, as outlined in CDC’s Interim Guidance for...
Managing Personnel with SARS-CoV-2 Infection or Exposure.

In cases of HCP exposures of staff or patients in healthcare systems in which a designated individual or team, qualified by education, training, and experience or certification, is responsible for carrying out facility infection prevention protocols and is available to serve as primary point of contact for the facility regarding COVID-19 outbreaks, a risk assessment may be performed that takes into account presence of symptoms, proximity and duration of encounters, and the use of personal protective equipment. The healthcare facility or system will take the lead on contact tracing and patient exposure notifications and will consult their local public health authority as needed. Healthcare facilities or systems have some discretion in identifying exposures that are higher risk and warrant notification and quarantine. Risk stratification should be aligned with CDC guidance.

Features of higher-risk exposures:
- Longer duration of exposure
- Healthcare provider close contact with patient airway (e.g., intubation, pharyngeal examination, bronchoscopy, laryngoscopy)
- Patient unmasked

Features of lower-risk exposures:
- Shorter duration of exposure
- No close contact with airway or mucous membrane
- Patient masked

In addition to any determination made due to the above factors, healthcare systems should notify contacts of HCP with COVID-19 if the patient with known exposure resides in or will be transferred to a congregate care setting, in order to inform testing efforts.

7.1.4 Caring for hospitalized COVID-19 cases
HCP who enter the room of a patient with suspected or confirmed SARS-CoV-2 infection should adhere to standard precautions and use a fit-tested N95 or higher-level respirator (or a facemask if respirator supply is genuinely limited and measures to obtain N95-level or higher respiratory protection via local or state resource requests have been exhausted), gown, gloves, and eye protection. Performing or assisting with an aerosol-generating procedure warrants airborne precautions, including an N95 or higher-level respiratory protection. Any necessary aerosol-generating procedures (§10) should be undertaken in an airborne infection isolation room, if available. Additional PPE considerations are provided in CDC Interim Infection Prevention and Control Recommendations. Transmission-based precautions should continue to be followed until discontinuation of isolation criteria are met.

7.1.5 Responding to a case or outbreak in LTCFs (SNF, ALF, RCF, MC)
COVID-19 can present with a broad range of symptoms (see §3.2), making identification of outbreaks difficult. LPHAs should have a low threshold for investigation when there is a cluster of illnesses in a congregate residential setting. Because COVID-like illness (CLI) and influenza-like illness (ILI) are similar, it is a priority to investigate any CLI or ILI in LTCFs and other congregate settings because they may indicate an outbreak of either; see §8.2 for guidance specific to outbreaks in other congregate settings and correctional facilities. Respiratory specimens should be collected from all ill persons in such outbreaks to be tested for COVID-19; and, during influenza season, for influenza. Testing for other pathogens may be considered.
Please remember that while individual influenza cases are not reportable, ILI outbreaks are reportable. If an ILI outbreak is identified, create an outbreak in Opera Outbreaks or call the regular ACDP line (971-673-1111) to report the outbreak.

Following the identification of a single new case of COVID-19 in a resident or HCP, LTCFs should determine if others in the facility are ill or have been exposed. LTCFs have the option to conduct contact tracing if resources and experience allow. However, a broad-based testing approach (e.g., unit, floor, full facility) is preferred if resources and experience in conducting contact tracing are not available within the LTCF, all potential contacts cannot be identified, or if contact tracing fails to halt transmission.

Broad-based testing or testing of all close contacts (if contact tracing implemented) is recommended regardless of vaccination status, as identified by the LTCF. Viral testing is recommended at days 1, 3, and 5, with the exposure occurring on day 0. Quarantine is recommended as outlined in §8.1.1, however all individuals being tested for a potential exposure should wear source control. If no additional cases are identified during contact tracing or broad-based testing, no further testing is indicated.

In the event of ongoing transmission that is not controlled with initial interventions, a facility should consider implementing quarantine for residents and HCP with higher-risk exposures. If additional cases are identified, facilities should strongly consider moving to broad-based testing and implementing quarantine for residents in affected areas of the facility.

7.1.6 Testing guidance in LTCFs
ODHS is in the process of amending OAR 411-060 to align with the CDC testing recommendation. CDC recommendations include broad-based testing (“unit-wide”) or contact tracing options in response to identification of a staff or a resident COVID-19 case. ODHS is expected to adopt contact tracing or broad-based testing options, however, facilities should continue to utilize broad-based testing when the facility does not have the experience and resources to perform individual contact tracing or should contact tracing efforts fail to control ongoing transmission.

An outbreak investigation and initial testing would not be triggered when a resident with known COVID-19 is admitted directly into transmission-based precautions, or when a resident known to have close contact with someone with COVID-19 is admitted directly into transmission-based precautions and develops COVID-19 before precautions are discontinued.

7.2 Non-Healthcare Congregate and Carceral Settings (e.g., shelters, supported/supportive living, temporary/transitional housing, employer-provided congregate housing, prisons, jails, youth detention facilities)

CDC’s Respiratory Virus Guidance provides information on how to prevent transmission of respiratory viruses including COVID-19.

The guidance above does not apply to dedicated patient care areas within these settings. HCP or individuals who are routinely in the patient care areas should refer to the healthcare recommendations in §7.1.

7.2.1 Responding to outbreaks in non-healthcare congregate and carceral settings
Similar to healthcare settings, these settings benefit from early risk assessment and
When COVID-19 cases are identified in residents, staff or volunteers including providing guidance for testing, implementing infection control practices (e.g., cohorting, ventilation, PPE use), and access to vaccines and medical care. Enhanced COVID-19 prevention strategies include moving activities outdoors, requiring universal masking indoors, and increased physical distancing. Refer to CDC’s guidance for congregate residential settings and shelters and carceral settings for additional considerations.

Although not required, non-healthcare congregate and carceral settings should test residents and staff who have been exposed or who have symptoms compatible with COVID-19. If additional cases are identified, consult with OHA to determine if more widespread testing or screening testing would be appropriate. LPHAs are encouraged to offer infection control consultations in collaboration with the OHA COVID-19 Congregate Care Epidemiologist. If the LPHA experiences challenges with facilities, they should work with their COVID-19 Regional Epidemiologist to elevate concerns and connect with the appropriate licensing and regulatory body.

7.2.2 Managing outbreaks associated with the Oregon Department of Corrections
When there is an outbreak of COVID-19 in an Oregon Department of Corrections (ODOC) facility, ODOC will perform a contact investigation within the facility and notify OHA of identified cases. OHA staff serve as the leads on outbreaks associated with ODOC facilities and are responsible for data management associated with the outbreaks. If an outbreak is identified in a local carceral facility not under ODOC jurisdiction, the LPHA should work with Community Corrections to investigate the outbreak.

8.0 GLOSSARY OF TERMS

Aerosol-generating procedures:
Include, but are not limited to:
- Intubation, extubation, and related procedures such as manual ventilation and open suctioning
- Cardiopulmonary resuscitation
- Tracheotomy and tracheostomy procedures (insertion, open suctioning, removal)
- Bronchoscopy
- Surgery and post-mortem procedures involving high-speed devices
- Some dental procedures (such as high-speed drilling)
- Non-invasive ventilation (NIV) such as bi-level positive airway pressure (BiPAP) and continuous positive airway pressure ventilation (CPAP)
- High-frequency oscillating ventilation (HFOV)
- High-flow nasal oxygen (HFNO) [i.e., oxygen delivered through high-flow nasal cannula (HFNC) at ≥15L/min].
- Induction of sputum
- Medication administration via continuous nebulizer

COVID-19-related pediatric death: As of May 11, 2023, only COVID-19-related deaths in persons <18 years of age are reportable. A death is considered to be related to COVID-19 in any of the following circumstances:
- Death of a confirmed or probable COVID-19 case within 60 days of the earliest available date among exposure to a confirmed case, onset of symptoms, or date of specimen collection for the first positive test;
- Death from any cause in a hospitalized person during their hospital stay or in the
60 days following discharge and a COVID-19-positive laboratory diagnostic test at any time since 14 days prior to hospitalization; or

- Death of someone with a COVID-19-specific ICD-10 code listed as a primary or contributing cause of death on a death certificate, regardless of the dates of diagnosis or death.

**Healthcare personnel (HCP):** Any paid or unpaid person serving in a healthcare setting who has the potential for direct or indirect exposure to patients or infectious materials, including body substances (e.g., blood, tissue, and specific body fluids); contaminated medical supplies, devices, and equipment; or contaminated environmental surfaces. HCPs may include, but are not limited to, emergency medical service personnel, nurses, nursing assistants, physicians, technicians, therapists, personal support workers, home care workers, phlebotomists, pharmacists, students and trainees, veterinarians, dentists, contractual staff not employed by the health care facility, and persons (e.g., clerical, dietary, environmental services, laundry, security, maintenance, engineering and facilities management, administrative, billing, and volunteer personnel) not directly involved in patient care but potentially exposed to infectious agents that can be transmitted between HCPs and patients.

**Healthcare setting:** Means any place where health care, including physical, dental or behavioral health care is delivered and includes, but is not limited to any health care facility or agency licensed under ORS chapter 441 or 443, such as hospitals, ambulatory surgical centers, birthing centers, special inpatient care facilities, long-term acute care facilities, inpatient rehabilitation facilities, inpatient hospice facilities, nursing facilities, assisted living facilities, residential facilities, residential behavioral health facilities, adult foster homes, group homes, pharmacies, hospice, vehicles or temporary sites where health care is delivered or is related to the provision of health care (for example, mobile clinics, ambulances) outpatient facilities, such as dialysis centers, health care provider offices, dental offices, behavioral health care offices, urgent care centers, counseling offices, offices that provide complementary and alternative medicine such as acupuncture, homeopathy, naturopathy, chiropractic and osteopathic medicine, and other specialty centers.

**Moderate or severely immunocompromised person:** One who requires care in a protected environment, (e.g., bone marrow transplant recipients, individuals with severe combined immunodeficiency) and HIV+ persons with CD4+ percentages <15% or CD4+ T lymphocyte counts <200. Immunocompromised persons include but are not limited to those who:

- Are currently or have recently been receiving cancer treatment for tumors or cancers of the blood
- Received an organ transplant and are taking medicine to suppress the immune system
- Received chimeric antigen receptor (CAR)-T-cell therapy (a treatment to help your immune system attach to and kill cancer cells) or received a stem cell transplant (within the last 2 years)
- Have a moderate or severe primary immunodeficiency (e.g., DiGeorge syndrome or Wiskott-Aldrich syndrome)
- Have advanced or untreated HIV infection
- Are receiving treatment with high-dose corticosteroids or other drugs that may suppress the immune response

**Period of transmissibility:** The time when cases can transmit disease to others. For symptomatic COVID-19 cases, this begins 48 hours prior to symptom onset. For asymptomatic cases, this is presumed to have begun 48 hours prior to the collection of the first specimen that
tested positive. The period of transmissibility extends until the case has met criteria for discontinuation of isolation.

**Physical distancing:** Maintaining distance from others to the greatest extent possible. Physical distancing measures reduce opportunities for person-to-person virus transmission and can help slow the spread of the disease, as well as save lives.

**REFERENCES**


**UPDATE LOG**

March 11, 2024. Removed outdated sections and updated isolation guidance to align with CDC Respiratory Virus Guidance. (Amanda Faulkner, Melissa Sutton, Bailey Weissenfels)

May 22, 2023. Updated outbreak reporting threshold for congregate care settings; updates to section 2.7.1 to reflect updated vaccination recommendations; removed test reporting requirements; updated isolation recommendations for the general population; updates throughout to reflect these changes. (Becca Pierce, Lee Peters, Melissa Sutton, Paul Cieslak).


January 27, 2023. Updated outbreak monitoring period for all outbreaks; removed outbreak monitoring for outbreaks in K-12 and childcare settings; added clarification around contact investigations in §6.2.2. (Lee Peters, Melissa Sutton, Meagan McLafferty, Kristen Hollywood).

January 6, 2023. Updated MIS-C case definition in alignment with recent CDC changes; updated links to OSPHL documents and treatment options; removed outdated links. (Lee Peters, Melissa Sutton, Paul Cieslak, Sarah Humphrey King).

December 15, 2022. Streamlined section 2.7 to reference federal guidance; removed recommendation to prioritize collection of a line list for outbreaks in K-12 and early learning settings; combined sections 8.2 and 8.3 to updated guidance for non-healthcare congregate and carceral settings in alignment with CDC; removed section 9.2 REDCap Platform and references throughout including presumptive case definition criteria following the retirement of the survey on December 9, 2022; added additional information on data management in Opera Outbreaks; removed links to retired guidance documents; updates throughout (Lee Peters, Meagan McLafferty, Becca Pierce, Melissa Sutton, Paul Cieslak).

October 20, 2022. Updated section 8.1 healthcare settings to align with CDC guidance released September 23rd including adding section 8.1.2 isolation for healthcare workers, patients, and residents in healthcare settings; updated vaccine information and moved to table format; removed vaccine up to date table and sections 3.5 breakthrough case definition, 8.4 managing special situations in K-12 and ELD
childcare settings, 8.5 cases with recent or planned travel, and 9.4 managing close contacts; numerous updates throughout (Lee Peters, Melissa Sutton, Amanda Faulkner, Becca Pierce, Paul Cieslak).

May 27, 2022. Revised outbreak response section; updated treatment and vaccines section; removed historic K-12, ELD, and agriculture outbreak subsections; added new CDC guidance for outbreaks in carceral settings (Amanda Faulkner, Lee Peters, Paul Cieslak).

March 12, 2022. Removed requirement for quarantine among the general population; updated K-12 and childcare guidance sections; formatting changes in the healthcare and other congregate settings sections; updated vaccines section. (Amanda Faulkner, Lee Peters, Becca Pierce, Melissa Sutton, Paul Cieslak).


December 29, 2021. Adopted new CDC shortened quarantine and isolation guidance for the general population and HCW; updated vaccine and treatment section; removed guidance regarding active monitoring; removed 7-day shortened quarantine with test; removed outbreak guidance regarding general workplaces; added language and intention for prioritizing public health response for COVID-19. (Amanda Faulkner, Paul Cieslak, Becca Pierce, Tom Jeanne).

December 6, 2021. Defined extracurricular activities in the test to stay guidance; clarified that masked staff on school buses are also eligible for test to stay. (Amanda Faulkner, Melissa Sutton).

December 2, 2021. Added language regarding the risk of outdoor exposures and variables to consider when determining if quarantine is needed for contacts; added language for a modified quarantine option for exposures in K-12 settings where universal masking is in place, updated test interpretation table (Lee Peters, Tom Jeanne, Paul Cieslak, Melissa Sutton, Amanda Faulkner).

November 18, 2021. Updated presumptive case definition to specify symptoms for persons who test positive using an at-home test; removed recommendation for people who test positive at-home to follow-up with a confirmatory test; recommended use of 7-day quarantine with negative test option for close contacts who work in or attend K-12 schools (Amanda Faulkner, Tom Jeanne, Melissa Sutton).

October 19, 2021. Added language about close contact exceptions for outdoor K–12 settings; revised ideal post-exposure test window to 5–7 days (Meagan McLafferty, Amanda Faulkner).

September 24, 2021. Added language about new case investigation protocol; clarified
April 29, 2021. Updated duration of quarantine to 14 days for all unvaccinated close contacts; updated surge conditions guidance section. Added detail to vaccination/treatment section. (Amanda Faulkner).

March 22, 2021. Added clarification surrounding vaccine breakthrough case surveillance follow-up; clarified use of test-based discontinuation of isolation; provided language regarding upcoming OSPHL whole genome sequencing capacity; clarified at-home test kits. (Amanda Faulkner).

February 17, 2021. Added Surge Conditions Guidance section; refined new quarantine guidance for fully-immunized close contacts in health care settings to match CDC’s; updated infection control language to align with OHA Clinical and Infection Control Guidance; added breakthrough case surveillance project information (Amanda Faulkner, Rebecca Pierce).

January 20, 2021. Updated treatment, prevention and limitation of spread section; provided new quarantine guidance for fully-immunized close contacts; clarified timing of isolation period for asymptomatic cases who subsequently develop symptoms. (Amanda Faulkner).

December 9, 2020. Removed language regarding creation of suspect cases based on negative test results; added options for shorter quarantine, adopting CDC options in part (Amanda Faulkner, Melissa Sutton, Paul Cieslak).

November 25, 2020. Added clarification for assessment and notification regarding persons exposed to cases among healthcare personnel, removed test-based discontinuation of isolation, modified close contact definition to include ‘24-hour’ time frame in line with CDC, included direction for sharing case information with schools, directed LPHA to classify MIS-C cases as Suspect until chart review is complete. (Kristen Hollywood, Melissa Sutton, Amanda Faulkner).

September 18, 2020. Clarified the recommended isolation period for cases who live in congregate settings, updated language to reflect that all jurisdictions are on ARIAS, defined first- and second-generation in the context of linking cases to outbreaks, added required data elements for outbreak reporting, added the definition of COVID-19-related hospitalization, sundry edits (Steve Rekant).

July 23, 2020. Changed all mentions of Orpheus to Opera, updated discontinuation of isolation criteria for symptoms from 72 hours to 24 hours, deemphasized test-based
discontinuation of isolation and added the longer minimum period for specific groups, included new testing rules and guidance, added positive antigen tests to the confirmed case definition and added language about any test developed under an FDA EUA, added description of criteria for possible work exemptions for quarantine and isolation, sundry edits (Steve Rekant)

July 2, 2020. Clarified language around using test-based discontinuation of isolation in LTCFs, added requirement for LPHAs to share information with employers (Steve Rekant)

June 24, 2020. Added details about investigating outbreaks, added references to ARIAS, clarified definition of suspect and presumptive cases including information about antigen testing, added MIS-C, disentangled discontinuation of isolation and assessment of recovery, harmonized language across sections, sundry edits (Steve Rekant)

May 1, 2020. Added presumptive case definition and revised recommended follow-up with contacts, defined recovery and clarified release from isolation, defined COVID-19-related deaths, clarified language around testing, added required follow-up for close contacts. (Steve Rekant, Kelly Cogswell)

April 1, 2020. Added language for emergency rule regarding reporting deaths and hospitalizations; reduced expectations for follow-up of potentially exposed persons; clarified language regarding testing in clusters; removed negative influenza test as a requirement for automatic testing approval at OSPHL; modified exposure period per new CDC guidance; added revised flowcharts. (Steve Rekant, Madeline LeVasseur, Amanda Faulkner, Rebecca Pierce)

March 23, 2020. Changed requirements for LPHA follow-up and investigation of PUMs, suspect cases, and confirmed cases. Updated guidance on monitoring and restrictions of exposed persons. Updated criteria for testing at OSPHL and overall testing prioritization recommendations. Changed language from PUI to suspect case and changed suspect and confirmed case definitions (Madeline LeVasseur, Steve Rekant, Amanda Faulkner, Orion McCotter)

March 12, 2020. Added information about other laboratories. Sundry edits. (Steve Rekant)

March 8, 2020. Edited testing criteria, PUM, PUI definitions. Updated guidance for discontinuation of isolation. Sundry edits. (Kelly Cogswell, Alexia Zhang)


February 28, 2020. Updated PUI case definition and testing criteria. Updated testing availability at the OSPHL. Added current list of geographic areas with widespread or sustained community transmission. (Tasha Poissant, Madeline LeVasseur)

February 20, 2020. Provided guidance on discontinuation of isolation for PUIs or COVID-19 cases and pregnant persons, and revised figures. (Alexia Zhang, Madeline LeVasseur, Steve Rekant)

February 12, 2020. Clarified expectations of LPHAs regarding contacting PUMs, provided guidance on interpreting testing, and revised figures. (Amanda Faulkner, Steve Rekant, Alexia Zhang)

February 7, 2020. Provided minor clarifications to date of PUM guidance implementation, DGMQ PUM forms, and Figures. (Amanda Faulkner, Steve Rekant)

January 2020. First draft. (Nicole West, Amanda Faulkner, Steve Rekant)