COVID-19 Case Classification

Clinical Criteria:
At least two of the following symptoms:

- Fever (subjective or measured)
- Chills
- Rigors
- Myalgia
- Headache

OR

At least one of the following symptoms:

- Cough
- Shortness of breath
- Difficulty breathing
- New olfactory disorder
- New taste disorder

OR

Severe respiratory illness with at least one of the following:

- Clinical or radiologic evidence of pneumonia, or
- Acute respiratory distress syndrome (ARDS)

AND

No alternative more likely diagnosis
Laboratory Criteria:
Method used as laboratory evidence should be approved or authorized by the FDA*

Confirmatory laboratory evidence:
- Detection of SARS-CoV-2 RNA in a clinical or autopsy specimen using a molecular amplification test (various nomenclature: NAT, NAA, NAAT, PCR, RNA, RT-PCR, rRT-PCR, etc.)

Presumptive** laboratory evidence:
- Detection of SARS-CoV-2 by antigen test in a respiratory specimen

Supportive** laboratory evidence:
- Detection of specific antibody to SARS-CoV-2 (IgM, IgG, or total [IgM + IgG]) in serum, plasma, or whole blood; may be indicative of previous infection or vaccination
- Detection of specific SARS-CoV-2 antigen by immunocytochemistry in an autopsy specimen


**The terms confirmatory, presumptive, and supportive are categorical labels used here to standardize case classifications for public health surveillance. The terms should not be used to interpret the utility or validity of any laboratory test methodology.

Epidemiological Evidence:
One or more of the following exposures in the last 14 days before onset of symptoms or positive test:
- Close contact* with a confirmed or probable case of COVID-19 disease; or
- Member of a risk cohort** as defined by public health authorities during an outbreak

*Close contact is defined as being within 6 feet for at least 15 minutes, and close personal activities, such as hugging or kissing. In healthcare settings, this may be defined as any exposure of greater than a few minutes.

**Examples of a “risk cohort” might be health care workers treating patients with certain respiratory illnesses, staff and residents in long-term care facilities and other confined population environments (behavioral health hospitals, prisons, etc.) with confirmed cases, travel to a defined geographic location with known COVID-19 outbreaks/clusters, or exposure to defined outbreaks/clusters, including business, child care, church, sports team, and other settings.

Vital Records Criteria:
A death certificate that lists COVID-19 disease or SARS-CoV-2 as a cause of death or a significant condition contributing to death (Note: If the cause of death is clearly unrelated to COVID-19, this will not qualify despite laboratory evidence of SARS-CoV-2 infection at time of death).
**Case Classification:**

**Confirmed**
- Meets confirmatory laboratory evidence

**Probable**
- Meets clinical criteria AND epidemiologic evidence with no confirmatory laboratory testing performed for COVID-19 OR
- Meets presumptive laboratory evidence OR
- Meets vital records criteria with no confirmatory laboratory testing performed for COVID-19 unless additional epidemiologic or clinical evidence exists to refute the vital records criteria

**Suspect**
- Meets supportive laboratory evidence with no prior history of being a confirmed or probable case OR
- Meets clinical criteria and is epi-linked to a confirmed case, but with one negative confirmatory laboratory test result within two days prior to, or 7 days after symptom onset.*

* Cases with more than one negative confirmatory laboratory test result would not be considered a Suspect case

**Distinguishing a New Case from an Existing Case when there is a Repeat Positive Test:**
A repeat positive test for SARS-CoV-2 RNA using a molecular amplification detection test or antigen test within 3 months of the initial report should not be enumerated as a new case for surveillance purposes. However, any repeat positive test for SARS-CoV-2 RNA using molecular amplification detection or antigen testing greater than 90 days after an initial positive test or case identification should be enumerated as a new case. Timing is based upon specimen collection dates and would start from the date of collection of the initial positive test (or from the onset of symptoms if testing not performed) that identified the case. Detection of antibody (IgG) greater than 90 days after an initial positive test or case identification would be indicative of past infection and would NOT be reported as a new case.