

Simplify COVID-19 Testing



The portable, easy-to-use BD Veritor™ Plus System provides reliable COVID-19 (SARS-CoV-2) results in 15 minutes.

Positive % Agreement (PPA)

BD Veritor System for Rapid Detection of SARS-CoV-2 with **84% PPA¹**

Negative % Agreement (NPA)

BD Veritor System for Rapid Detection of SARS-CoV-2 with **100% NPA¹**

Results in 15 minutes

“How much confidence can I have in this result?”

Negative Result
-

98.2%¹

Probability that the negative result is **correct****

Positive Result
+

>99%¹

Probability that the positive result is **correct*****

1.8% Probability that the negative result is incorrect

<1% Probability that the positive result is incorrect

1,000

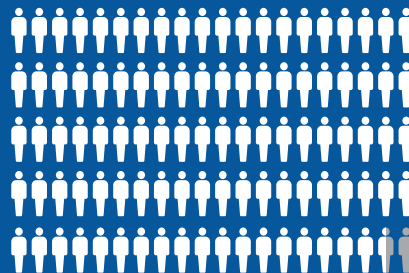
People tested (suspected of infection)[†]

100

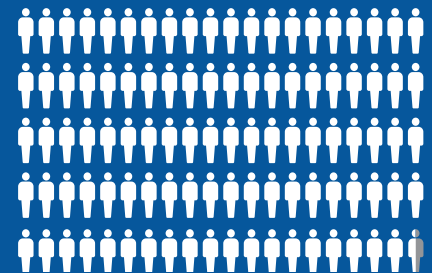
People confirmed infected with SARS-CoV-2

10%

Positivity rate*



Fewer than 2% of the 1,000 people tested might receive a **false negative** result.



Fewer than 1% of the 1,000 people tested might receive a **false positive** result.

[†] The intended use of the BD Veritor System for Rapid Detection of SARS-CoV-2 assay only includes those who are suspected of COVID-19 by their health care provider within the first five days of the onset of symptoms.

- This test has not been FDA cleared or approved;
- This test has been authorized by FDA under an EUA for use by authorized laboratories;
- This test has been authorized only for the detection of proteins from SARS-CoV-2, not for any other viruses or pathogens; and,
- This test is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.

References: 1. BD Veritor System for Rapid Detection of SARS-CoV-2 package insert. Franklin Lakes, NJ: Becton, Dickinson and Company. *CDC, accessed July 21, 2020 @<https://www.cdc.gov/coronavirus/2019-ncov/covid-data/covidview/index.html>. **At a prevalence of 10%; the negative predictive value is the probability that persons with a negative test result truly do not have the disease. ***At a prevalence of 10%; the positive predictive value is the probability that persons with a positive test result truly have the disease.

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