Quick reference guide
BD Veritor™ System for rapid COVID-19 (SARS-CoV-2) testing*

Analyze Now Mode

Read the complete test procedure, including recommended QC procedures before performing the test. Refer to the package insert for complete information about the test. Ensure ALL components are at room temperature (15–30 °C) when running the test.

Sample collection and preparation
1. Gather test materials and label test device with specimen ID.
2. Remove cap from extraction reagent tube.
3. Insert patient sample swab and vigorously plunge the swab up and down for 15 seconds.
4. Remove swab while squeezing tube to extract liquid. Properly dispose of swab.
5. Press dispensing tip on the tube firmly. Mix the sample by flicking swirling the bottom of the tube.

Running the assay on the BD Veritor™ Plus system
6. Add 3 drops of the processed sample to the test device sample well.
7. Allow test to develop for 15 minutes.*
8. When test is ready, power on instrument by pressing blue start button once. When prompted, insert test device to read.
9. Result will appear on screen. See interpretation chart below. Record result and remove test device. Properly dispose of test device.

Optional. If using the barcode scanning accessory, follow screen prompts to scan any required barcodes to start the test analysis.

Interpretation of results

<table>
<thead>
<tr>
<th>Display</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoV2: +</td>
<td>Positive Test for SARS-CoV-2 (antigen present)</td>
</tr>
<tr>
<td>CoV2: –</td>
<td>Presumptive Negative Test for SARS-CoV-2 (no antigen)</td>
</tr>
<tr>
<td>CONTROL</td>
<td>Test Invalid. Repeat the test.</td>
</tr>
<tr>
<td>INVALID</td>
<td></td>
</tr>
</tbody>
</table>

*CAUTION: Incorrect results may occur if development time is less than 15 minutes. Cover test device if working in a drafty environment.

*For use under Emergency Use Authorization (EUA) only
Quick reference guide

BD Veritor™ System for rapid COVID-19 (SARS-CoV-2) testing
For use under Emergency Use Authorization (EUA) only

**Interpretation of results**
Test results must **NOT** be read visually. The BD Veritor Plus System Analyzer (purchased separately) must be used for interpretation of all test results. Refer to table above.

**Positive Test Results**: SARS-CoV-2 antigen present; does not rule out coinfection with other pathogens.

**Negative Test Results**: Negative results are presumptive. Negative test results do not preclude infection and should not be used as the sole basis for treatment or other patient management decisions, including infection control decisions, particularly in the presence of clinical signs and symptoms consistent with COVID-19, or in those who have been in contact with the virus. It is recommended that these results be confirmed by a molecular testing method, if necessary for patient management.

**Invalid Test**: If the test is invalid the BD Veritor Plus System Analyzer will display a “CONTROL INVALID” result and the test or control must then be repeated.

<table>
<thead>
<tr>
<th>Display</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoV2: +</td>
<td>Positive Test for SARS-CoV-2 (antigen present)</td>
</tr>
<tr>
<td>CoV2: –</td>
<td>Presumptive Negative Test for SARS-CoV-2 (no antigen)</td>
</tr>
<tr>
<td>CONTROL INVALID</td>
<td>Test Invalid. Repeat the test.</td>
</tr>
</tbody>
</table>

**Warnings and precautions**
1. **For in vitro** Diagnostic use only.
2. All test results must be obtained using the BD Veritor Plus Analyzer.
3. **DO NOT** read the test results visually.
4. Handle all specimens and related materials as if capable of transmitting infectious agents.
5. Dispose of used materials as biohazardous waste in accordance with federal, state and local requirements.
6. Ensure all components are at room temperature (15–30 °C) when running the test.
7. Please refer to the package insert for detailed assay instructions, cautions, limitations and warnings.

**Specimen collection and handling**
Proper specimen collection and handling is required to ensure accurate results (see enclosed specimen collection guide). Additional training or guidance is recommended if operators are not experienced with specimen collection and handling procedures.

**External quality control procedure**
Swab controls are supplied with each kit. These swab controls should be used to ensure that the test reagents work properly and that the test procedure is performed correctly. For kit swab controls, insert the control swab into the extraction reagent tube and vigorously plunge the swab up and down for 15 seconds. Process according to the test procedures on the reverse side of this card beginning at step 4. BD recommends running controls for each new kit lot, each new operator, and each new shipment of test kits or at periodic intervals required by your facility. If the kit controls do not perform as expected, do not report patient results and contact BD Technical Support at 1.800.638.8663.

**Technical Information:**
In the United States contact BD Technical Service and Support at 1.800.638.8663 or bd.com

- 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.

BD Life Sciences
7 Loveton Circle, Sparks, MD, 21152, USA
800.638.8663

bd.com

BD, the BD Logo and Veritor are trademarks of Becton, Dickinson and Company or its affiliates. © 2020 BD.
All rights reserved. 574-US-0820 August 2020