



Harness the power of simplicity

Easy to use. Flexible workflow. Fast, traceable results.

A streamlined point-of-care diagnostic experience is at hand with the BD Veritor™ Plus system.



BD Veritor™ Plus System

Enhancing the point-of-care testing experience



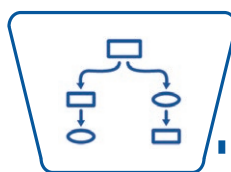
Time is limited. Staff are busy. Results are critical. Now is when your team needs unambiguous test results, streamlined workflows, simple training and clear documentation. The BD Veritor™ Plus system is designed to address these challenges with:

- Objective unambiguous results: “+” for positive, “-” for negative, give you confidence in testing results
- Flexible workflow options to adapt to your workplace needs
- Traceability and data storage help with impact analysis
- Timely results—15 minutes for SARS-CoV-2⁵ (COVID-19)*, less than 11 minutes for Flu A+B^{1,2} and RSV**, and less than 6 minutes for Group A Strep⁴—limiting patient wait time



Train your staff with ease

- Delivers easy-to-read results using a digital display
- Simplifies staff training with the **BD Veritor™ Plus eLearning platform**, featuring test training modules, how-to videos, progress, dashboards and more
- Minimizes training time and facilitates operation with intuitive sample processing
- CLIA-waived status allows for simple training for novice and nonlab personnel to confidently test for SARS-CoV-2 (COVID-19), Flu A+B, Group A Strep and RSV



Enable workflow flexibility

- Select the mode to fit your workflow:
 - **Analyze Now** mode displays results in seconds. Simply incubate specimen for time recommended for each assay, then insert test device into Analyzer
 - For multitasking, use **Walk Away** mode. Add specimen to test device and insert into Analyzer for automatic incubation and analysis

*Negative results do not rule out SARS-CoV-2 infection and should not be used as the sole basis for treatment or patient management decisions, including infection control decisions. Negative results should be considered in the context of a patient's recent exposures, history and the presence of clinical signs and symptoms consistent with COVID-19, and confirmed with a molecular assay, if necessary, for patient management. For additional information, please refer to the HCP fact sheet.

- 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
- 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.

**respiratory syncytial virus



Accelerate reliable results

- Proven performance for SARS-CoV-2 (COVID-19)*, Flu A+B detection vs PCR^{1,2}
- Analyzer technology helps improve specificity to reduce false-positive results⁴
- Timely results may improve patient experience
 - SARS-CoV-2⁵ (COVID-19)* in 15 minutes⁵
 - Flu A+B^{1,2} or RSV³ result in less than 11 minutes;
 - Group A Strep result in less than 6 minutes⁴
- Reduces the potential for manual errors by streamlining sample processing
 - Uses prefilled, color-coded tubes



Facilitate quality control

- BD Veritor™ InfoScan module provides reliable quality control and traceability through customizable documentation functionality (user ID, specimen ID and test kit/lot)
- May reduce risk of reporting errors by minimizing manual transcription and leveraging data storage/download of printable data via USB helps cut risk of manual transcription

Meet patient and staff needs with simple, timely point-of-care testing

- Easy-to-read results with 1-button functionality and minimal hands-on time to meet your staff's demands
- Flexible workflow options to adapt to your workplace needs
- Timely, reliable results (15 minutes for SARS-CoV-2⁵ (COVID-19), <11 minutes for Flu A+B¹ and RSV³; <6 minutes for Group A Strep⁴) for minimized wait times
- BD Veritor™ Plus eLearning platform simplifies staff training
- Quality control and traceability offered with the BD Veritor™ InfoScan module



Ordering information	Cat. no.	Qty.
BD Veritor™ System for Rapid Detection of SARS-CoV-2*	256082	30 tests
BD Veritor™ System RSV CLIA-waived kit	256038	30 tests
BD Veritor™ System RSV (Moderately complex)	256042	30 tests
BD Veritor™ System Group A Strep CLIA-waived kit	256040	30 tests
BD Veritor™ System Flu A+B CLIA-waived kit	256045	30 tests
BD Veritor™ System Flu A+B (Moderately complex)	256041	30 tests
BD Veritor™ Plus System Analyzer	256066	1
BD Veritor™ InfoScan module	256068	1
USB printer cable	443907	1

References: 1. BD Veritor System for Rapid Detection of Flu A+B, CLIA-waived kit [package insert], 8087667. Franklin Lakes, NJ: Becton, Dickinson and Company. 2. BD Veritor System for Rapid Detection of Flu A+B, laboratory kit [package insert], 8087666. Franklin Lakes, NJ: Becton, Dickinson and Company. 3. BD Veritor System for Rapid Detection of Respiratory Syncytial Virus (RSV), [package insert], 8086098. Franklin Lakes, NJ: Becton, Dickinson and Company. 4. BD Veritor System for Rapid Detection of Group A Strep, laboratory kit [package insert], 8087675. Franklin Lakes, NJ: Becton, Dickinson and Company. 5. BD Veritor System for Rapid Detection of SARS-CoV-2 [package insert], Franklin Lakes, NJ: Becton, Dickinson and Company.

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