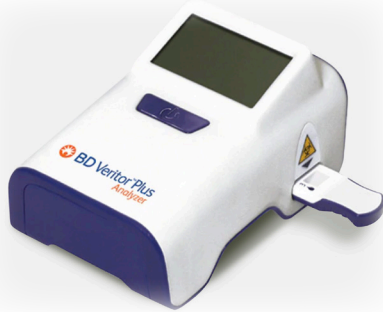


# COVID TESTING!

## Bring Diagnostic Testing to Your Office Starting with COVID Testing! Announcing Henry Schein's Dental Diagnostic Testing Program

Keep your staff and patients safe with in-office COVID results in 10–20 minutes.



### Choose from 4 COVID testing platforms\*:

- Cue Health molecular
- BD Veritor antigen
- Abbott BinaxNOW antigen
- Premier Biotech antibody

Henry Schein Dental has developed a comprehensive program to guide you through setting up and conducting testing in your practice.



- Laboratory Licensure
- Setting Up Testing Environment
- Ordering Supplies
- Administering Tests
- Reporting Results
- Reimbursement and/or Billing the Patient

We are here to support you through the process.

To learn more about how to get started, reach out to Henry Schein today at  
[DentalDX@henryschein.com](mailto:DentalDX@henryschein.com).

For technical questions, feel free to contact  
Diagnostic Consultant Deborah Forren, MT (ASCP) at  
[deborah.forren@henryschein.com](mailto:deborah.forren@henryschein.com)

\*These tests have not been cleared or approved by FDA, but they have been authorized by FDA under Emergency Use Authorizations for use in Authorized Laboratories (e.g., point-of-care settings with CLIA Certificates of Waiver). The Cue Health molecular test is authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens; the BD Veritor antigen test and the Access Bio antigen test are authorized only for the detection of proteins from SARS-CoV-2, not for any other viruses or pathogens; the Premier Biotech antibody test has been authorized only for detecting the presence of IgM and IgG antibodies to SARS-CoV-2, not for any other viruses or pathogens. The emergency use of all of these tests 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 Federal Food, Drug and Cosmetic Act, 21 U.S.C. § 360bbb3(b)(1), unless the declaration is terminated or authorization is revoked sooner.

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