

Health study finds self-collected saliva and deep nasal swabs are equally effective for the diagnosis of COVID-19

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Self-collected saliva and deep nasal swabs collected by healthcare providers are equally effective for detecting SARS-CoV-2, the virus that



causes COVID-19, according to a new study conducted by ARUP Laboratories and University of Utah (U of U) Health.

The study, published in the *Journal of Clinical Microbiology*, represents one of the largest prospective specimen type comparisons to date, said Julio Delgado, MD, MS, ARUP chief medical officer. Other studies, including one from the Yale School of Public Health, have reached similar conclusions but with markedly fewer patients and specimens.

Researchers also found that specimens self-collected from the front of the nose are less effective than deep nasal swabs for virus detection. This finding prompted a subsequent study that has not yet been published in which researchers learned they could improve the sensitivity of anterior nasal <u>swab</u> testing to 98% by combining an anterior nasal swab with a swab collected from the back of the throat.

The results have important implications for patients and providers. The collection process for saliva and anterior nasal specimens is less invasive than the deep nasal, or nasopharyngeal, swab. In addition, both specimen types can be self-collected, reducing the risk of exposure for healthcare workers who collect nasopharyngeal specimens, said Kimberly Hanson, MD, MPH, section chief of clinical microbiology at ARUP and the primary author of the study.

"Saliva and nasal swab self-collection can resolve many of the resource and safety issues involved in SARS-CoV-2 diagnostic testing," Delgado said.

ARUP and U of U Health anticipate being able to start offering testing on saliva in some U of U Health clinical settings in early September. They already are using anterior nasal swabs in combination with throat swabs to test some asymptomatic individuals.



COVID-19 testing on these alternatives to nasopharyngeal swabs will increase with time, Delgado said. "From the start of the COVID-19 pandemic, ARUP has worked to build capacity for high-quality COVID-19 testing," he said. "Our goal is to make this testing available to hospitals and healthcare systems nationwide."

Hanson and her colleagues analyzed more than 1,100 specimens from 368 volunteers at the U of U Health Redwood Health Center drivethrough testing site from late May through June. Volunteers selfcollected saliva that they spit into a tube and swabbed from the front of both nostrils to produce specimens for testing. The researchers compared test results from these specimen types with test results from nasopharyngeal swabs <u>healthcare providers</u> collected from the volunteers. Discrepant results across specimens collected from the same patient triggered repeat testing using a second polymerase chain reaction (PCR)-based platform.

The study showed that SARS-CoV-2 was detected in at least two specimen types in 90% of the patients who tested positive for the virus.

As a standalone alternative specimen to nasopharyngeal swabs, saliva proved to be an excellent option, Hanson said. Positivity rates for <u>saliva</u> specimens were nearly the same as those for nasopharyngeal specimens.

The research showed that self-collected nasal swabs, when used alone, can miss nearly 15% of infections, which prompted researchers' further study combining them with oropharyngeal, or throat swabs.

More information: K. E. Hanson et al. Self-Collected Anterior Nasal and Saliva Specimens versus Healthcare Worker-Collected Nasopharyngeal Swabs for the Molecular Detection of SARS-CoV-2, *Journal of Clinical Microbiology* (2020). DOI: 10.1128/JCM.01824-20



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