

## New COVID-19 technology to deliver faster, more accurate test results

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Process flow from saliva collection to data reporting. (A) Schematic diagram of the main steps of the process flow. (B) Sentinel robot for LAMP reaction setup, incubation, and detection. Individual process steps: 1. Saliva collection via microswabs into test tubes containing inactivation solution. 2. Sample tubes with saliva dilutions were heat-inactivated for 10 min at 95 °C. 3. Automated uncapping of a complete sample tube rack. 4. Sample tube barcode reading during Sentinel robot loading. 5. The automated pipetting system adds a selected sample volume to the RT-LAMP master mix in clear reagent plates. 6. Plate sealing by automated heat sealer. 7. Plate crane transport of the reaction plate onto a 65 °C incubation field. 8. LAMP reactions were incubated at 65 °C for 25 min; digital, parallel scanning of multiple 96-well microplates. 9. Real-time, secure reporting of deidentified data and analysis. 10. Disposal of scanned microplates. Credit: *Scientific Reports* (2022). DOI: 10.1038/s41598-022-08263-4

The latest research from Murdoch University and the Perron Institute is paving the way for the development of a new technological screening platform that will rapidly and accurately detect COVID-19, without compromising quality.

The Avicena Sentinel saliva sampling approach involves sensitive molecular processes and ultra-high throughput technology for screening potentially infectious, asymptomatic carriers.

It can run up to four thousand samples per hour, with results reported within 25 minutes.

Latest data published in the *Nature* journal *Scientific Reports* shows 98% accuracy in identifying the SARS-CoV-2 virus in samples.



Senior author, Professor Sulev Koks, Head of Genetic Epidemiology Research at Murdoch University and at Western Australia's Perron Institute, is among the scientists and others contributing.

He is collaborating with local specialists from Avicena Systems, Telethon Kids Institute, The University Western Australia and further afield in the United Kingdom and Spain. The key overseas organizations evaluating the technology are the Lancs LAMP Laboratory in Preston, UK and Biodonostia Health Research Institute in San Sebastian, Spain.

Professor Koks said although present testing methods are accurate, they are not capable of providing a result to people with COVID-19 that have no symptoms. Additionally, PCR testing puts a strain on the health system as health staff need to undertake rigorous training to obtain results.

"The rapid antigen tests (RAT) are easier to use but not sufficiently sensitive in detecting the virus in infected people who are contagious but have no symptoms. The <u>viral load</u> samples from presymptomatic or asymptomatic people can take several days to reach sufficient levels for detecting infection.

"PCR results take hours and throughput is, at most, a few thousand samples processed per instrument per day. Also, nasopharyngeal PCR sampling requires trained personnel and has decreased participant acceptance if multiple testings are required within a short period.

"Effective viral surveillance to contain outbreaks requires frequent testing with more sensitive assays, particularly as new variants arise."

Professor Koks explained the new screening technology, that will identify infectious patients sooner and eliminate the threat of unwanted community transmission, will help hasten the pandemic's closure.



"The proposed large-scale and convenient saliva testing platform combines accuracy and scalability to mitigate the risk of viral transmission as restrictions are lifted and to guard against future threats.

"It would make frequent testing feasible in complementing vaccines to contain the spread of highly infectious pandemic agents such as COVID-19."

Further validation of Avicena's rapid, ultra-sensitive Sentinel screening system is continuing.

First authors of the recent publication are Robert Dewhurst and Tatjana Heinrich (Perron Institute and Avicena Systems), and the other authors are Paul Watt and Paul Ostergaard (Avicena Systems), Jose Maria Marimon (Biodonostia Institute), Mariana Moreira and Philip Houldsworth (Lancs LAMP Laboratory), Jack Rudrum (Perron Institute) and David Wood (UWA).

**More information:** Robert E. Dewhurst et al, Validation of a rapid, saliva-based, and ultra-sensitive SARS-CoV-2 screening system for pandemic-scale infection surveillance, *Scientific Reports* (2022). DOI: 10.1038/s41598-022-08263-4

Provided by Murdoch University

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