

SO YOU KNOW. ON THE GO. ➡

QuickFinder™ COVID-19 Antigen Self Test

Convenient, Reliable and Easy-To-Use

For *in vitro* diagnostic use only
For use under FDA EUA only
For use with anterior nasal swab specimens



*See QuickFinder COVID-19 Antigen Self Test instructions for use



- Large sample loading port
- Large results window
- Thick results lines

A low-cost and reliable solution for COVID-19 self testing



Rapid self testing
anytime, anywhere

25M+



Quality manufacturing
at significant scale

100+

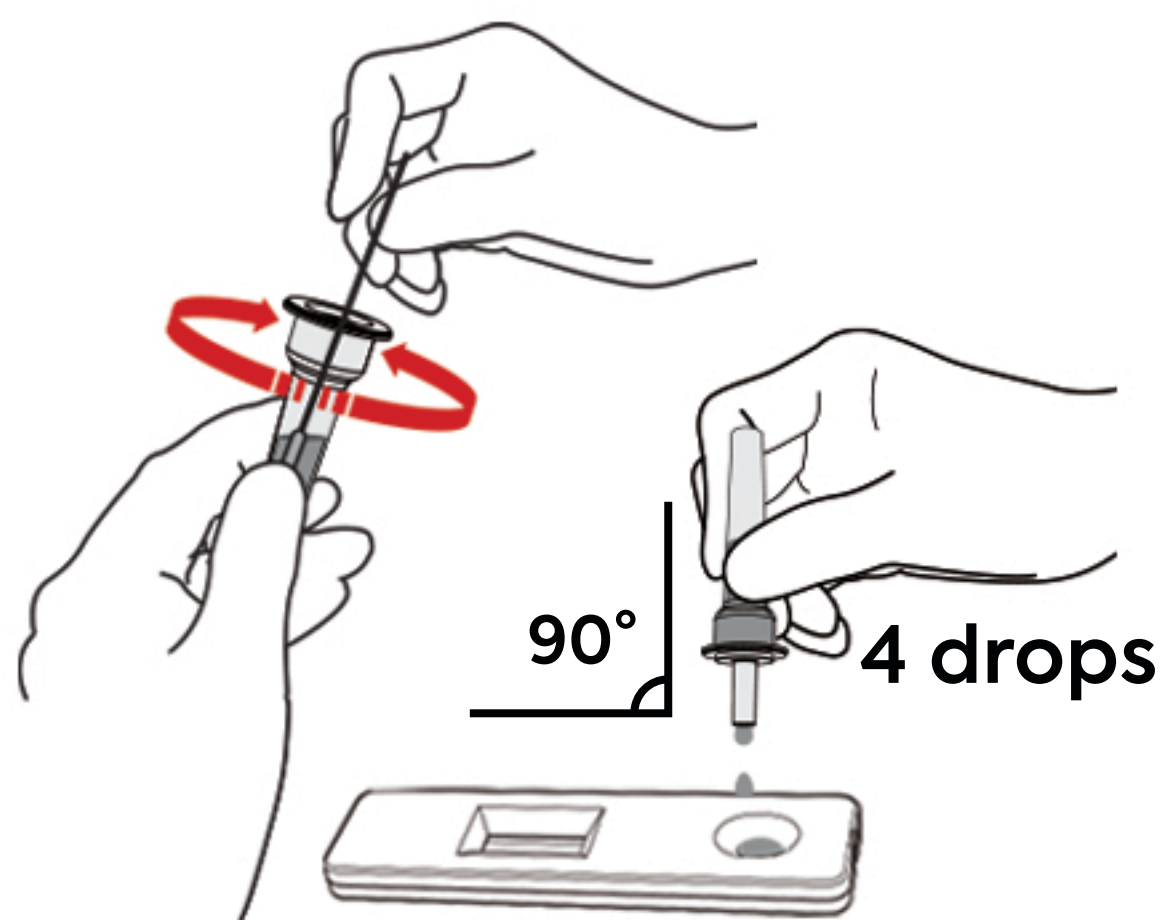


OHC products trusted in
markets across the globe

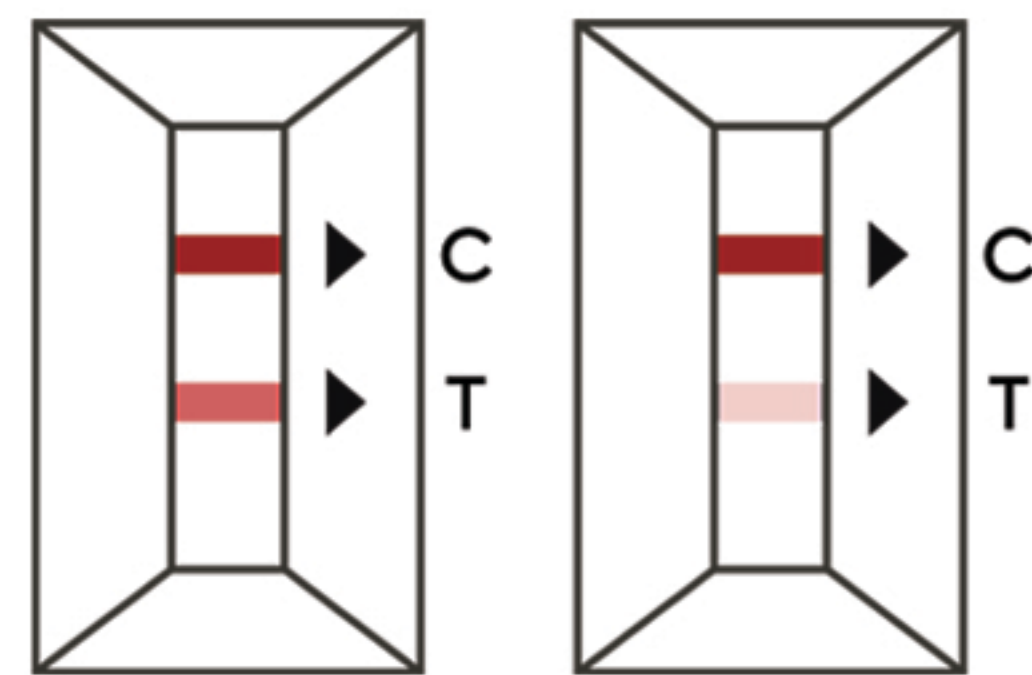
3 simple steps for on the go results



Swab



Transfer



Read



Trusted and Tested

FDA Approved
NIH RADx graduate
NIH ITAP graduate



Comprehensive

Detects known SARS-CoV-2
clinical variants



Integrate into Daily Life

ConnectedHealth
Capability – in process



Flexible

3PL and reformatting available
POC and Self Test labeling



Performance



91% relative sensitivity*



99% relative specificity*

*The QuickFinder COVID-19 Self Test has not been FDA cleared or approved; but has been authorized by FDA under an Emergency Use Authorization.

*Clinical Data shows, QuickFinder™ COVID-19 Antigen Self Test showed a relative sensitivity of 91% (95% CI:82.8 to 95.6%) at high viral loads (less than a cycle threshold (Ct)-value of 30) and relative specificity of 99% (95% CI: 95.2 to 99.6%)

Detecting All Mutation

*The QuickFinder™ Ag Rapid Test Diagnosis Test's anti-body design was validated by the United States NIH (National Institution of Health) and the ITAP (Independent Test Assessment Program) with outstanding recognition rates for all mutations.

* Through the United States NIH (National Institution of Health) and the ITAP (Independent Test Assessment Program) we were able to test and confirm the detection of the SARS-CoV-2 despite any previous or current MOH VOC which includes Omicron BA.4 & BA.5.

**Learn about purchase or distribution of the QuickFinder™ COVID-19 Self Test
at www.osangllc.com or email us at sales@osangllc.com**

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