

FIND Evaluation of Core Technology Co., Ltd

Core tests COVID-19 Ag Test

External Report

Version 2.0, [19 July2022]

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Evaluation process - private sector engagement

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For the COVID-19 response, FIND has commissioned independent evaluations of in vitro diagnostics following an Expression of Interest (EOI) process available on FIND's website by which all test submissions were scored according to their regulatory status and time to market; the manufacturing and distribution capacity of the supplier; and the supplier-reported clinical and analytical performance.

| Document version | Date | Comment | |
|------------------|------------------|--|--|
| 1.0 | 14 December 2021 | First release | |
| 2.0 | 19 July 2022 | Analytical data from several variants added Clinical data from Uganda added | |

Document history



1 Product Info:

| Manufacturer name | Core Technology Co., Ltd. | |
|--|--|--|
| Test name | Coretests COVID-19 Ag Test | |
| Product code(s) | B291-20A | |
| Pack size(s) | 25 tests per kit | |
| Contents of kit | COVID-19 Ag Test Cassette, Instructions for use, Sample collection tube containing processing solution, nasal swab | |
| Equipment and consumables required, but not provided | PPE, timer | |
| Product storage (temperature range) | 2-30°C | |
| Shelf-life (months) | 24 months | |
| Manufacturing site (country) | China | |

2 Study details:

| Study design: | Prospective diagnostic evaluation studies across multiple, independent sites to determine the accuracy of COVID-19 antigen RDTs, using consecutive enrolment. Interim analyses are performed at 25% and 50% enrolment, and the evaluation is stopped if tests do not meet 95% specificity. Presence of symptoms, date of symptom onset and hospitalization status is collected for all enrolled participants. | |
|--------------------------|---|--|
| Index assays: | Novel lateral flow format tests that detect recombinant SARS-CoV-2 antigens. | |
| Reference method: | Results of the index test are compared to the routine, diagnostic RT-PCR result, which is used for clinical management | |
| Limit of detection: | Analytical sensitivity, i.e. Limit of detection, was performed at the Liverpool School of Tropical Medicine in which standardized serial dilutions of cultured viral isolate were prepared. Proprietary swab provided in the kit was soaked in viral dilution series. Dilutions were tested in triplicate and the LOD was defined as the last dilution where all repeats were interpreted as positive. | |
| Clinical performance: | Sensitivity was calculated as the proportion of true positive results detected by Core tests COVID-19 Test among all positives by the reference method, and reported as a percentage | |



| Specificity was calculated as the proportion of true negative specimens, identified as negative by Core tests COVID-19 Test among all negatives by the reference method and reported as a percentage. |
|---|
| The 95% confidence intervals were calculated to assess the level of uncertainty introduced by sample size, using the Wilson's score method. |

3 Evaluation details:

| Country of collaborator | Switzerland | Uganda |
|---|--|---|
| Location of clinical site(s) (city, town) | University Hospital of Geneva | Mulago National Referral Hospital Kiruddu National Referral Hospital Mbarara Regional Referral Hospital Masaka Regional Referral Hospital |
| Health care level of site(s) | Community Testing Clinic | National or regional referral hospitals |
| Study period (date to date) | 19 November – 3 December 2021 | 2- 22 June 2022 |
| Study cohort inclusion/exclusion | Adults in community meeting Department of Public Health definition of a suspected COVID- 19 case and being tested for SARS-CoV-2 part of routine medical care. Provided informed consent | Adults with acute respiratory illness who meets WHO case definition of a COVID-19 suspected case presenting to temporary testing locations around clinical sites. Provided informed consent |
| Sample type, antigen test | Nasal swab | Nasopharyngeal swab |
| Reference PCR method | Cobas SARS-CoV-2 (Roche Diagnostics) n=227; Xpert Xpress SARS-CoV-2 (Cepheid) n=5 | cobas® SARS-CoV-2 Test (Roche Diagnostics) |
| Sample type, PCR test | Nasopharyngeal | Nasopharyngeal swab |



4 Results:

4.1 Study cohort

| Country | Switzerland | Uganda |
|--|----------------------------|---------------------|
| Total N (valid PCR results) | 230 | 437 |
| Age [mean (min-max), N] | 38.7 (16-77), 230 | 35.2 (18-86), 437 |
| Gender [%F, (n/N)] | 62.7%, (142/230) | 44.9%, (196/437) |
| Symptoms present [%Yes, (n/N)] | 100%, (58/58) ¹ | 100%, (437/437) |
| Hospitalized (n, % Yes) | Not applicable | Not applicable |
| Days from symptom onset [median (Q1-Q3); N] | 2 (1-3), 55 ² | 4 (2-5), 437 |
| Days < 0-3 (n, %) | 44, 80% | 214, 49% |
| Days 4-7 (n, %) | 10, 18% | 183, 42% |
| Days 8+ (n, %) | 1, 2% | 40, 9% |
| Positivity [%, (n/N)] | 23%, (54/230) | 26%, (115/437) |
| PCR Ct [median (Q1-Q3); N] | 21.7 (19.3-25), 54 | 26.4 (22.6-29), 115 |
| Ct > 33 (n, %) | 4,7% | 6, 5% |
| Ct > 30 (n, %) | 5, 9% | 23. 20% |
| Ct > 25 (n, %) | 13, 24% | 60, 52% |

¹Symptom data only available for PCR and/or RDT positive samples

²Symptom onset data missing for n=3

4.2 Estimation of Clinical Performance

| Country | Switzerland | Uganda |
|----------------------------------|-------------------------|-------------------------|
| Clinical Sensitivity (95% CI), N | 87% (75.6, 93.6), 54 | 86.8% (79.4, 91.9), 115 |
| Sensitivity days ≤7, N | 88% (76.2, 94.4), 50 | 80.4% (71.2, 87.3), 92 |
| Sensitivity Ct ≤ 33, N | 92% (81.2, 96.8), 50 | 88.1% (80.7, 92.9), 109 |
| Sensitivity Ct ≤ 25, N | 97.6 (87.4, 99.6), 41 | 96.4% (87.7, 99), 55 |
| Clinical Specificity (95% CI), N | 98.3% (95.1, 99.4), 176 | 98.5% (96.4, 99.3), 322 |



| Invalid rate (%, n/N) | 0% (0/230) | 0% (0/437) |
|-----------------------|------------|------------|
| | | |

4.3 Estimation of analytical performance

Supplier-reported LOD = 2.25 x 10¹ TCID50/ml ~ 3.17 x 10¹ pfu/ml (isolate Gamma (P1))

• Verified LOD

| Variant | Lowest dilution detected | Verified LOD | Viral Copy equivalent |
|-----------------|---|-----------------------------|---------------------------------------|
| (lineage) | | concentration | |
| UK wild type | 5.0 x10³ pfu/ml ~ 7.05 x | 5.0 x10³ pfu/ml | 9.8 x10 ⁶ genome copies/ml |
| (B1) | 10 ³ TCID ₅₀ /ml | | applied to test |
| Alpha (B.1.1.7) | 2.5 x10² pfu/ml ~ 3.525 x | 2.5 x10² pfu/ml | 9.3 x10 ³ genome copies/ml |
| | 10 ² TCID ₅₀ /ml | | applied to test |
| Gamma (P1) | 1.0 x10¹ pfu/ml ~ 1.41 x | 1.0 x10 ¹ pfu/ml | 3.1 x10 ³ genome copies/ml |
| | 10¹ TCID₅₀/ml | | applied to test |
| Delta | 2.5 x10² pfu/ml ~ 3.525 x | 2.5 x10² pfu/ml | 9.9 x10⁵ genome copies/ml |
| (B.1617.2) | 10 ² TCID ₅₀ /ml | | applied to test |
| Omicron | 1.0 x10³ pfu/ml ~ 1.41 x | 1.0 x10³ pfu/ml | 1.8 x10 ⁵ genome copies/ml |
| (BA.1) | 10 ³ TCID₅₀/ml | | applied to test |
| | | | |

Note: viral dilution was applied directly to the test cassette, not to the provided swab