

**FIND Evaluation of Mologic Ltd,**  
**COVID 19 RAPID ANTIGEN TEST**  
**External Report**  
*Version 1.2, 15 June 2021*

**Copyright and use of the report**

Copyright in this report is the property of FIND (or controlled by FIND). You are free to share, copy and redistribute the material in any medium or format provided that:

- (i) attribution: you must give appropriate credit to FIND and indicate if changes were made, you may do so in any reasonable manner, but not in any way that suggests that FIND endorses you or your use;
- (ii) non-commercial: you may not use the report for commercial purposes; and
- (iii) no derivatives: if you remix, transform, or build upon the materials or report, you may not distribute the modified materials or report unless with express authorization from FIND.

Presentation of data on our website does not impact any data ownership rights and FIND is not responsible for any use by any third party of these data. Data sources are provided.

**Evaluation process – private sector engagement**

FIND, the global alliance for diagnostics, seeks to ensure equitable access to reliable diagnosis around the world. It works closely with the private and public sectors and receives funding from donors and some of its industry partners. It has internal fire walls, policies and processes to protect it against any undue influence in its work or the publication of its findings.

More information on our policy and guidelines for working with private sector partners can be found here: <https://www.finddx.org/policies/>

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 history**

Document version	Date	Comment
1.0	23 April 2021	Initial release
1.1	08 June 2021	Updated shelf-life
1.2	15 June 2021	PCR result for one sample updated. Ag result for one sample updated.

## 1 Product Info:

Manufacturer name	Mologic Ltd.
Test name	COVID 19 RAPID ANTIGEN TEST
Product code(s)	11811125
Pack size(s)	25 tests per kit
Contents of kit	Lateral flow device, buffer capsule, sterile swab, swab extraction tube, instructions for use
Equipment and consumables required, but not provided	Tube stand, Stopwatch/timer
Product storage (temperature range)	2-30°C
Shelf-life (months)	18 months
Manufacturing site (country)	United Kingdom

## 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:	Verification of 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:	<p>Sensitivity was calculated as the proportion of true positive results detected by COVID 19 RAPID ANTIGEN TEST among all positives by the reference method and reported as a percentage.</p> <p>Specificity was calculated as the proportion of true negative specimens, identified as negative by COVID 19 RAPID ANTIGEN TEST among all negatives by the reference method, and reported as a percentage.</p>

	<p>The 95% confidence intervals were calculated to assess the level of uncertainty introduced by sample size, using the Wilson’s score method.</p> <p>Invalid Ag RDT results were not repeated.</p>
Ease of use	<p>A System usability survey and ease of use questionnaire assessing the quality of the test, test preparation, ease of test execution, procedure time, ease of result interpretation, storage conditions and perceived settings of use was completed by operators and a final score out of 100 was calculated.</p>

### 3 Evaluation details

<b>Country of collaborator</b>	<b>Germany</b>
Location of clinical site(s) (city, town)	<ol style="list-style-type: none"> <li>1. Heidelberg (HD)</li> <li>2. Berlin</li> </ol>
Health care level of site(s)	<ol style="list-style-type: none"> <li>1. Heidelberg: Drive-in testing Center</li> <li>2. Berlin: Ambulatory testing clinic of Charité – University Hospital</li> </ol>
Study period (date to date)	<ol style="list-style-type: none"> <li>1. HD: 11- 31 March 2021</li> <li>2. Berlin: 11 March – 15 April 2021</li> </ol>
Study cohort inclusion/exclusion	<p>Adults able to ambulate and meeting suspect definition of the Department of public health</p> <p>Provided informed consent</p>
Sample type, antigen test	Nasal (AN) (n=645) or NMT (n=20)
Reference PCR method	<ul style="list-style-type: none"> <li>• LightMix® Modular SARS-CoV (COVID19) E-gene (Tib Molbiol) <ul style="list-style-type: none"> <li>○ N = 323</li> </ul> </li> <li>• Cobas SARS-CoV-2 (Roche Diagnostics Inc) <ul style="list-style-type: none"> <li>○ N = 342</li> </ul> </li> </ul>
Sample type, PCR test	<ol style="list-style-type: none"> <li>1. HD: Nasopharyngeal swabs (oropharyngeal if NP contraindicated)</li> <li>2. Berlin: Combined nasopharyngeal/oropharyngeal swabs</li> </ol>

## 4 Results

### 4.1 Study cohort

<b>Country</b>	Germany
Total N (valid PCR results)	665
Age [mean (min-max), N]	38.65 (18-78), 665
Gender [%F, (n/N)]	50.2%, (333/664) <sup>1</sup>
Symptoms present [%Yes, (n/N)]	66.5%, (440/662)
Hospitalized (n, % Yes)	Not applicable
Days from symptom onset [median (Q1-Q3); N]	2 (1-4); 436 <sup>1</sup>
Days < 0-3 (n, %)	290, 67%
Days 4-7 (n, %)	121, 28%
Days 8+ (n, %)	25, 6%
Positivity [%, (n/N)]	29%, (195/665)
PCR Ct [median (Q1-Q3); N]	20.3 (17.83 – 22.95); 195
Ct > 33 (n, %)	5, 3%
Ct > 30 (n, %)	8, 4%
Ct > 25 (n, %)	28, 14%

<sup>1</sup> Age unknown for n=1; Date of symptom onset unknown for n=4

## 4.2 Estimation of clinical performance

Country	Germany
Clinical Sensitivity (95% CI), N	90.6% (85.6, 94), 192 <sup>§1</sup>
Sensitivity days ≤7, N	93.2% (88.3, 96.2), 162
Sensitivity Ct ≤ 33, N	92.5% (87.8, 95.5), 187
Sensitivity Ct ≤ 25, N	96.4% (92.3, 98.3), 166
Clinical Specificity (95% CI), N	100% (99.2, 100), 458 <sup>2</sup>
Invalid rate (% , n/N)	2.4% (16/665)

<sup>§</sup> 89/191 positives were tested using Roche Cobas, 103/191 using TibMolbiol.

<sup>1</sup> n=3 invalid for investigational RDT, excluded from the analysis

<sup>2</sup> n=13 invalid for investigational RDT, excluded from the analysis

### 4.2.1 Estimation of analytical performance

	Lowest dilution detected	Verified LOD concentration	Viral Copy equivalent	Supplier-reported LOD
<b>Analytical Sensitivity</b>	<b>2.5 x10<sup>2</sup> pfu/ml</b> ~ 3.52 x 10 <sup>2</sup> TCID <sub>50</sub> /ml	<b>2.5 x10<sup>2</sup> pfu/ml</b>	<b>5.9 x10<sup>5</sup> copies/ml</b> applied to test	<350 TCID <sub>50</sub> /ml ~ 245 <b>pfu/ml</b>

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

## 4.3 Ease of use

Mologic COVID 19 RAPID ANTIGEN TEST	57 out of 100	5 operators, Germany
-------------------------------------	---------------	----------------------