



Nabídka

Covid-19 testů ze slin

**Realy Tech Novel Coronavirus Antigen
Rapid Test Device (Saliva)**



Realy Tech (Saliva)

5 testů v krabičce

100 krabiček v krabici

Velikost krabičky: 12 x 7,1 x 7,1 cm

Velikost krabice: 50 x 37 x 39 cm

Objem: 0,0721 m³

Hmotnost krabička: 81 g

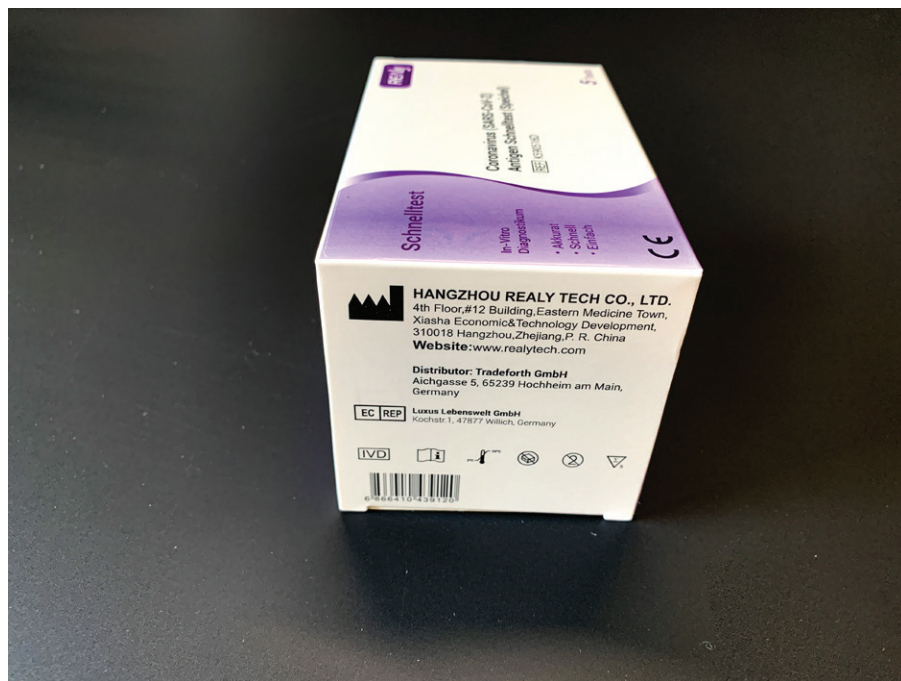
Hmotnost krabice: 8,1 kg

Délka testování: 10 minut

Citlivost: 93,94%

Specifita: >99%











Novel Coronavirus(SARS-COV-2) Antigen Rapid Test Device (Saliva)



20 Tests/kit

Features:

- Specimen type: Saliva
- Testing time: 10 minutes
- Sensitivity: 93.94%
- Specificity:>99%



By Saliva collect cup
or



By Saliva collector

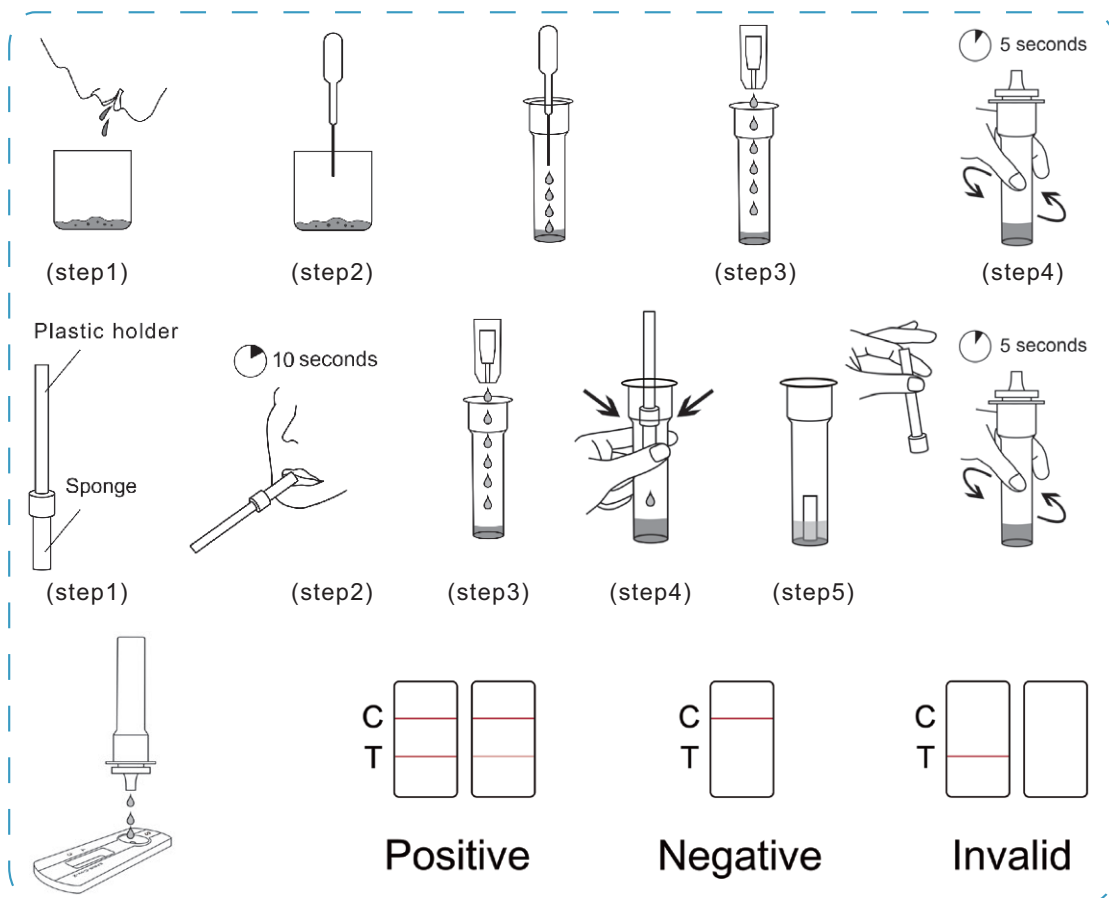
Product name	Ref.	Size	Storage Temp
Novel Coronavirus(SARS-COV-2) Antigen Rapid Test Device (Saliva)	K590516D	20 Test/kit	2-30 °C
Novel Coronavirus(SARS-COV-2) Antigen Rapid Test Device (Saliva)	K590516D	5 Test/kit	2-30 °C
Novel Coronavirus(SARS-COV-2) Antigen Rapid Test Device (Saliva)	K590516D	1 Test/kit	2-30 °C

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The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva) is an in vitro diagnostic test for the qualitative detection of novel coronavirus antigens in human saliva, using the rapid immunochromatographic method. The identification is based on the monoclonal antibodies specific for the novel coronavirus antigen. It will provide information for clinic doctors to prescribe correct medications.

DIRECTIONS FOR USE:



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HANGZHOU REALY TECH CO., LTD.

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Economic & Technology Development, 310018 Hangzhou,
Zhejiang, P. R. China
<http://www.realytech.com>
Email: info@realytech.com



Declaration of Conformity



in accordance with Directive 98/79/EC

Manufacturer:

Name: HANGZHOU REALY TECH CO., LTD.

Address: 4th Floor, #12 Building, Eastern Medicine Town, Xiasha Economic & Technology Development, 310018 Hangzhou, Zhejiang, P. R. China

Product/s	Catalogue number
Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Device (saliva)	K590516D

Category: Other Devices (All devices except Annex II and self-testing devices)

Conformity assessment route: Annex III, except Point 6, of Directive

Applicable Standards: EN ISO 13485: 2016; EN ISO 15223-1:2016; EN ISO 14971: 2012; EN ISO 13612:2002; EN ISO 17511:2003; EN ISO 18113-1:2011; EN ISO 18113-2:2011, EN ISO 23640:2015.

We, the Manufacturer, herewith declare with sole responsibility that our product/s mentioned above meet/s the provisions of the Directive 98/79/EC of the European Parliament and of the Council on In-Vitro Diagnostic Medical Devices.

We hereby explicitly appoint Luxus Lebenswelt GmbH, located at Kochstr.1,47877, Willich, Germany to act as our European Authorised Representative as defined in the aforementioned Directive.

HangZhou 2020-12-15

(Place and date of issue)



(Signature and position)

Signed for and on behalf of the manufacturer



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杭州睿丽科技有限公司
Hangzhou Realy Tech Co., Ltd.

**Clinical Validation report of Novel
Coronavirus (SARS-CoV-2) Antigen Rapid
Test device (saliva)**



Product name: Novel Coronavirus (SARS-CoV-2) Antigen Rapid
Test device (saliva)

Package Specification: 25 tests/kit

Manufacturer: Hangzhou Realy Tech Co., Ltd

I. Clinical validation time

This clinical evaluation was conducted from October 2020 to November, 2020.

II. Background information for clinical evaluation

Since December 2019, world has successively discovered multiple cases of patients with new-type coronavirus pneumonia. With the spread of the epidemic, China and abroad have also been found. As an acute respiratory infectious disease, the disease has been included in the Class B infectious diseases stipulated in the Law of the People's Republic of China on the Prevention and Control of Infectious Diseases, and is managed as a Class A infectious disease. Based on the current epidemiological investigation, the incubation period is 1-14 days, mostly 3-7 days.

The main manifestations are fever, dry cough, and fatigue. A few patients have symptoms such as nasal congestion, runny nose, sore throat, myalgia and diarrhea. Severe patients usually have dyspnea and / or hypoxemia one week after the onset of symptoms, and severe patients can quickly progress to acute respiratory distress syndrome, septic shock, difficult to correct metabolic acidosis, coagulation dysfunction and multiple organ Functional failure, etc. It is worth noting that in the course of severe and critically ill patients, there may be moderate to low fever, even without obvious fever.

Mild patients showed only low fever, mild fatigue, and no pneumonia. Judging from the current cases, most patients have a good prognosis, and a few patients are critically ill. The elderly and those with chronic underlying disease have a better prognosis. Symptoms in children are relatively mild.

The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva) developed by our company can help diagnose whether patients are infected with the Novel Coronavirus. It has further enriched the detection methods of Novel Coronavirus, expanded the supply of detection reagents, and fully served the needs of epidemic prevention and control.

III. Test purposes

The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva) produced by Hangzhou Realy Technology Co., Ltd. was used to verify the feasibility of clinical evaluation and the reliability of test results for Chinese subjects.

The purpose of research of the clinical test is to calculate the consistency percentage of negative/positive and the total consistency percentage and Kappa coefficient by statistically analyzing test results through comparative experimental research.

IV. Test design

1. Test plan selection and reasons

In vitro diagnostic reagents for testing and reference reagents were used to conduct comparative research tests on clinically suspected Novel Coronavirus saliva samples, and it was proved that the in vitro diagnostic reagents used in the test can achieve the expected assistance in infection of the Novel Coronavirus.

2. Sample volume required

The total number of clinical trials of this product is not less than 100 cases. The samples is classified into the positive group and the negative group as per the test results of the reference product. Meanwhile, the (unfrozen) nasopharyngeal swab samples shall be tested via the RT-PCR from the same patient at same time , then the saliva sample test results of the product tested and the

nasopharyngeal swab sample RT-PCR test results shall be compared, with statistical analysis being made.

3. Sample inclusion/exclusion certification.

The positive group and negative group in this experiment are applicable to the following inclusion/exclusion criteria.

Positive group inclusion:

PCR Test is positive;

symptoms are clinically positive;

Negative inclusion:

PCR test is negative;

Sample collection, processing

It is applicable to the diagnosis of the Novel coroinavirus from the samples of saliva. Use freshly collected samples for optimal test performance. Inadequate sample collection or improper sample handling may yield a false-negative result.

Sample collection procedure: The oral fluid specimen should be collected using the saliva collector provided with the kit. Follow the detailed Directions for Use refer to product IFU. No other collection cup should be used with this assay. Oral fluid collected at any time of the day may be used. **NOTE: Do not place anything in the mouth including food, drink, gum, tobacco, water and mouthwash products for at least 10 minutes prior to collection of oral fluid specimen.**

Specimen preparation:

Take out a sample extraction tube, remove the aluminum foil, Insert the sponge of sample collector with the saliva sample into the tube and twist close the whole cap of sample collector.

4. In vitro diagnostic reagents and reference products for testing

5.1 Test in vitro diagnostic reagents

Name: Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva)

Specification: 25 tests/kit

LOT: 202010001

Expiry: October,2022(Tentative)

Storage Conditions: Store in a dry place at 2-30°C, protected from light. After opening the inner package, the test card will become invalid due to moisture absorption. Please use it within 1 hour.

Source: Hangzhou Realy Tech Co.,Ltd

5.2 Reference products

Name: Novel Coronavirus(2019-nCoV) Nucleic Acid Diagnostic Kit (PCR-Fluorescence Probing)

Manufacturer: Sansure Biotech Inc.

Storage Conditions: Store in a dry place at 2-8°C, protected from light.

V. Experiment method

1. Open the package with the saliva collector, then remove the saliva collector from the sealed plastic bag.
2. Pre-process the saliva samples according to the instructions of the The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva) , and label the samples randomly.



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Hangzhou Realy Tech Co., Ltd.

- 2.1 Insert the sponge of saliva collector into the mouth, actively swab the inside of the mouth and tongue to collect oral fluid for approximately 10 second until the sponge becomes soft and fully saturated, The sponge will be free from hard spots when fully saturated.
- 2.2 . Take out a sample extraction tube, remove the aluminum foil. Remove the collector from the mouth and put the saturated oral fluid collector into the extraction tube.
- 2.3 Screw the cap into the extraction tube tightly so that saliva is squeezed out of the sponge into the extraction tube.
- 2.4 Gently shake the extraction tube vertically for about 5 seconds to allow saliva mix well with extraction buffer. Take out the saliva collector and discard it.
3. The operation steps of the in vitro diagnostic reagents for the test are as follows. For details, please refer to the product instruction manual:
 - 3.1 Remove the test device from the sealed foil pouch and use it as soon as possible. Best results will be obtained if the assay is performed immediately after opening the foil pouch. Put the test device on a clean and flat surface.
 - 3.2 twist open the small white cap from the extraction tube, transfer 3 drop of sample into the sample well of test device vertically.
 - 3.3 Read the result at 10~20 minutes. Don't interpret the result after 20 minutes.

Note: The detection steps need to be completed under protection against infection.

VI. Statistical methods of statistical analysis of clinical research data

A Methods evaluating clinical performance

Whether various indexes can reach the standards of clinical evaluation shall be judged by calculating the consistency percentage of negative/positive and the total consistency percentage in the test results of the product tested and the reference product, to validate the accuracy and applicability of the product in clinical applications. The product tested shall be subject to tests through the sample of different types, with statistics on the results. Meanwhile, different types of sample of the subjects shall be subject to determination by the product tested synchronously, and then the determination results of both shall be compared. The test results recorded shall be subject to statistical analysis upon completion of determination of all clinical samples, to calculate the consistency percentage of negative/positive and the total consistency percentage. Afterwards, equivalence of both shall be evaluated as per these statistical indexes

B Statistical method

The products launched on the market shall be subject to comparative study and evaluation. Kappa inspection: each sample shall be tested with the product tested and the reference product respectively, and then the consistency in statistical results of these two inspection methods shall be compared through Kappa inspection.

The data shall be subject to Kappa inspection and analysis and the Kappa coefficient shall be calculated. Favorable consistency can be proven if Kappa is >0.8 . The consistency in test results

of the product tested and the reference product is evaluated as per the evaluation standards.

VII Standards of clinical evaluation

The coincidence rate shall be calculated by comparing with the reference product whose marketing is approved. The product performance shall meet the following requirements.

1)Coincidence rate of negative: the sample whose test results are negative for both the product tested and the reference product and the proportion in the sample whose test results are negative for the reference product shall be more than 95%.

2)Coincidence rate of positive: the sample whose test results are positive for both the product tested and the reference product and the proportion in the sample whose test results are positive for the reference product shall be more than 85%.

3)Total coincidence rate: the sample whose test results are the same for the product tested and the reference product and its proportion in the total number of samples shall be more than 90%.

Method		2019-nCoV nucleic acid test kit (RT-PCR)		Total Results
The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva)	Result	positive	negative	
	positive	A	B	A+B
	negative	C	D	C+D
Total Results		A+C	B+D	A+B+C+D

Clinical sensitivity = $A/(A+C)*100\%$

Clinical specificity = $D/(B+D)*100\%$

Accuracy: $(A+D)/(A+B+C+D)*100\%$

If the coincidence rate of positive/negative can meet clinical requirements, two methods or Products are considered as equivalent; If the coincidence rate of positive/negative is greatly different, the clinical scheme should be re-designed.

4)Kappa consistency analysis shall be adopted for statistical analysis of reference reagents.

The results of the product tested are statistical materials and can be per the table below:

Method		2019-nCoV nucleic acid test kit (RT-PCR)		Total Results
The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva)	Result	positive	negative	
	positive	A	B	A+B
	negative	C	D	C+D
Total Results		A+C	B+D	A+B+C+D

$P_0=(A+D)/(A+B+C+D)*100\%$

$P_e=((A+B)(A+C)+(A+B)(B+D))/(A+B+C+D)^2$

Kappa: $(P_0 - P_e)/(1 - P_e)$

If conducting Kappa consistency analysis for the base data above, high consistency can be judged if the Kappa coefficient is >0.8, and both systems are considered as equivalent. Consistency is

considered if $0.4 < \text{Kappa coefficient} < 0.8$, and the coincidence rate of positive/negative shall be compared, with statistical analysis being made. Two such systems are considered as inconsistent and in-equivalent if the Kappa coefficient is < 0.4 .

VIII Provisions for amendments to clinical validation

In general, the clinical validation should not be changed. Any modification to the project during the test should be explained, and the time, reason, process of change, and whether there is a record of the change are explained in detail and its impact on the evaluation of the entire research result is explained.

IX. Results and Analysis of Clinical Tests

In total, 222 patients' samples are included for the unit, all the saliva samples and nasopharyngeal swab samples are tested. Statistics on rapid test results and those of the RT-PCR tested are as follows:

Method	2019-nCoV Nucleic Acid Test Kit (RT-PCR)		Total Results
	Results	Positive	
The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Cassette (Swab)	Positive	62	0
	Negative	4	156
Total Results		66	156
			222

Clinical sensitivity = $62/66 = 93.94\%$ (95%CI*84.99% to 98.06%)

Clinical specificity = $156/156 = 99.9\%$ (95%CI* 98.98% to 100%)

Accuracy: $(62+156)/(62+0+4+156) * 100\% = 98.20\%$ (95%CI* 95.29% to 99.46%)

$P_e = (62*66+62*156)/222*222 = 0.28$

Kappa: $(P_0 - P_e)/(1-P_e) = 0.97$

*: 95% confidence interval

According to the above table, 156 are proven negative of 156 negative specimens, 62 are proven positive of 66 positive specimens. The sensitivity and accuracy are more than 90%, indicating favorable consistency with the reference product. The Kappa=0.97>0.8, indicating favorable and high consistency of two methods and equivalence of two such systems.

X Analysis on Inconsistency in Test Results

NO.	Age	Gender	Rapid Test	RT-PCR	Clinical diagnostic
14	39	F	Negative	Positive (N gene)	Infection 25 days
25	43	F	Negative	Positive (N gene)	Infection 16 days
37	36	M	Negative	Positive (N gene)	Infection 28 days
49	30	M	Negative	Positive (RdRP gene)	Infection 22 days

XI Discussion and Conclusions

1. discussion

A Results of comparative analysis of the product tested and the reference product:

Test results of saliva specimen tested and the reference result: both the coincidence rate of negative/positive and the total coincidence rate are larger than 85%, indicating favorable consistency with the reference product. In the analysis results of Kappa inspection, Kappa was

proven >0.8, indicating favorable and high consistency of both methods. Both systems were proven equivalent.

2. Test conclusions

By analyzing the test results of the product tested and the reference product, the consistency percentage of negative/positive and the total consistency percentage are proven high. Moreover, according to the results of statistical analysis, there is no remarkable difference in test results of both, indicating favorable consistency in diagnosis and equivalence of two such systems and can be used for auxiliary diagnosis of those suffering from pneumonia triggered by COVID-19.

X. Quality control methods

On-site quality control

1) During the course of this study, clinical implementors appointed clinical inspectors to conduct regular on-site supervision visits to the research hospital. Through monitoring visits, it was found that all the contents of the research plan were strictly observed, and the correctness of the research data was also guaranteed. Participating researchers have undergone unified training, unified recording methods and judgment standards. The entire clinical trial process is conducted under strict operation, and the test content is complete and authentic. All observations and findings in the clinical trials have been verified and the data are reliable. The conclusions in the clinical trials are derived from the original data.

2) Quality control of clinical experiment process

During the evaluation, quality control was performed daily to ensure that the product was under control. Strict quality control is performed for each trial to ensure the quality of clinical trials.

XI. Prediction of adverse events

Because the Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test device (saliva) is an in vitro diagnostic reagent product, no direct contact with patients is required in clinical trials, no test report is provided to patients, and the test results are only used for comparative studies. It involves personal privacy, does not serve as a basis for auxiliary diagnosis, does not bring any risk to the subject, and does not cause adverse events.

References:

1. The "Technical Review Points for the Registration of New Coronavirus Antigen / Antibody Detection Reagents in 2019 (Trial)" issued by the State Drug Administration Medical Device Technical Evaluation Center on February 25, 2020;
2. "Pneumonitis Diagnosis and Treatment Program for New Coronavirus Infection (Trial Version 7)" issued by the National Health Committee on February 19, 2020.



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Annex II: Data of Clinical Tests

NO.	Age	Gender	Rapid Test	(RT-PCR)
1	49	F	Positive	Positive (RdRP and N gene)
2	32	F	Positive	Positive (RdRP and N gene)
3	31	F	Positive	Positive (RdRP and N gene)
4	32	F	Positive	Positive (RdRP and N gene)
5	21	F	Positive	Positive (RdRP and N gene)
6	51	M	Positive	Positive (RdRP and N gene)
7	22	F	Positive	Positive (RdRP and N gene)
8	46	F	Positive	Positive (RdRP and N gene)
9	23	F	Positive	Positive (RdRP and N gene)
10	14	M	positive	Positive (RdRP and N gene)
11	42	M	Positive	Positive (RdRP and N gene)
12	51	M	Positive	Positive (RdRP and N gene)
13	80	M	Positive	Positive (RdRP and N gene)
14	39	F	Negative	Positive (N gene)
15	67	M	Positive	Positive (RdRP and N gene)
16	44	M	positive	Positive (RdRP gene)
17	26	F	Positive	Positive (RdRP and N gene)
18	33	F	positive	Positive (N gene)
19	38	F	Positive	Positive (RdRP and N gene)
20	36	F	Positive	Positive (RdRP and N gene)
21	3	F	Positive	Positive (RdRP and N gene)
22	35	F	Positive	Positive (RdRP and N gene)

NO.	Age	Gender	Rapid Test	(RT-PCR)
23	23	F	Positive	Positive (RdRP and N gene)
24	43	M	Positive	Positive (RdRP and N gene)
25	43	F	Negative	Positive (N gene)
26	46	F	Positive	Positive (RdRP and N gene)
27	55	F	Positive	Positive (RdRP and N gene)
28	22	F	Positive	Positive (RdRP and N gene)
29	20	M	positive	Positive (N gene)
30	42	M	Positive	Positive (RdRP and N gene)
31	56	F	Positive	Positive (RdRP and N gene)
32	55	M	Positive	Positive (RdRP and N gene)
33	26	F	Positive	Positive (RdRP and N gene)
34	54	M	Positive	Positive (RdRP and N gene)
35	43	F	Positive	Positive (RdRP and N gene)
36	69	M	Positive	Positive (RdRP and N gene)
37	36	M	Negative	Positive (N gene)
38	37	F	Positive	Positive (RdRP and N gene)
39	44	F	Positive	Positive (RdRP and N gene)
40	43	F	Positive	Positive (RdRP and N gene)
41	67	F	Positive	Positive (RdRP and N gene)
42	51	F	Positive	Positive (RdRP and N gene)
43	75	F	Positive	Positive (RdRP and N gene)
44	60	F	Positive	Positive (RdRP and N gene)

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NO.	Age	Gender	Rapid Test	(RT-PCR)
45	25	M	Positive	Positive (RdRP and N gene)
46	75	F	Positive	Positive (RdRP and N gene)
47	43	F	Positive	Positive (RdRP and N gene)
48	30	F	Positive	Positive (RdRP and N gene)
49	30	M	Negative	Positive (RdRP gene)
50	26	F	Positive	Positive (RdRP and N gene)
51	32	F	Positive	Positive (RdRP and N gene)
52	73	M	Positive	Positive (RdRP and N gene)
53	58	F	Positive	Positive (RdRP and N gene)
54	66	F	Positive	Positive (RdRP and N gene)
55	29	F	Positive	Positive (RdRP and N gene)
56	56	M	Positive	Positive (RdRP and N gene)
57	24	M	Positive	Positive (N gene)
58	36	M	Positive	Positive (RdRP and N gene)
59	70	F	Positive	Positive (RdRP and N gene)
60	45	M	Positive	Positive (RdRP and N gene)
61	38	F	Positive	Positive (RdRP and N gene)
62	42	M	Positive	Positive (RdRP and N gene)
63	55	M	Positive	Positive (RdRP and N gene)
64	33	M	Positive	Positive (RdRP and N gene)
65	39	M	Positive	Positive (RdRP and N gene)
66	58	F	Positive	Positive (N gene)
67	77	F	Negative	Negative(Ct/Cq) >40

NO.	Age	Gender	Rapid Test	(RT-PCR)
68	62	F	Negative	Negative(Ct/Cq) >40
69	81	M	Negative	Negative(Ct/Cq) >40
70	18	F	Negative	Negative(Ct/Cq) >40
71	71	F	Negative	Negative(Ct/Cq) >40
72	37	M	Negative	Negative(Ct/Cq) >40
73	44	F	Negative	Negative(Ct/Cq) >40
74	79	M	Negative	Negative(Ct/Cq) >40
75	67	M	Negative	Negative(Ct/Cq) >40
76	61	F	Negative	Negative(Ct/Cq) >40
77	59	F	Negative	Negative(Ct/Cq) >40
78	28	F	Negative	Negative(Ct/Cq) >40
79	82	M	Negative	Negative(Ct/Cq) >40
80	63	F	Negative	Negative(Ct/Cq) >40
81	53	M	Negative	Negative(Ct/Cq) >40
82	43	M	Negative	Negative(Ct/Cq) >40
83	46	M	Negative	Negative(Ct/Cq) >40
84	46	F	Negative	Negative(Ct/Cq) >40
85	21	F	Negative	Negative(Ct/Cq) >40
86	46	F	Negative	Negative(Ct/Cq) >40
87	71	M	Negative	Negative(Ct/Cq) >40
88	60	F	Negative	Negative(Ct/Cq) >40
89	31	F	Negative	Negative(Ct/Cq) >40
90	72	M	Negative	Negative(Ct/Cq) >40

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NO.	Age	Gender	Rapid Test	(RT-PCR)
183	39	M	Negative	Negative(Ct/Cq) >40
184	61	F	Negative	Negative(Ct/Cq) >40
185	10	F	Negative	Negative(Ct/Cq) >40
186	37	M	Negative	Negative(Ct/Cq) >40
187	52	F	Negative	Negative(Ct/Cq) >40
188	41	M	Negative	Negative(Ct/Cq) >40
189	74	M	Negative	Negative(Ct/Cq) >40
190	51	F	Negative	Negative(Ct/Cq) >40
191	56	M	Negative	Negative(Ct/Cq) >40
192	62	F	Negative	Negative(Ct/Cq) >40
193	60	F	Negative	Negative(Ct/Cq) >40
194	54	F	Negative	Negative(Ct/Cq) >40
195	81	F	Negative	Negative(Ct/Cq) >40
196	79	F	Negative	Negative(Ct/Cq) >40
197	73	F	Negative	Negative(Ct/Cq) >40
198	35	F	Negative	Negative(Ct/Cq) >40
199	76	F	Negative	Negative(Ct/Cq) >40
200	23	M	Negative	Negative(Ct/Cq) >40
201	13	F	Negative	Negative(Ct/Cq) >40
202	14	M	Negative	Negative(Ct/Cq) >40

NO.	Age	Gender	Rapid Test	(RT-PCR)
203	43	M	Negative	Negative(Ct/Cq) >40
204	30	F	Negative	Negative(Ct/Cq) >40
205	57	M	Negative	Negative(Ct/Cq) >40
206	30	F	Negative	Negative(Ct/Cq) >40
207	65	M	Negative	Negative(Ct/Cq) >40
208	66	F	Negative	Negative(Ct/Cq) >40
209	38	F	Negative	Negative(Ct/Cq) >40
210	49	M	Negative	Negative(Ct/Cq) >40
211	23	F	Negative	Negative(Ct/Cq) >40
212	51	M	Negative	Negative(Ct/Cq) >40
213	64	F	Negative	Negative(Ct/Cq) >40
214	67	M	Negative	Negative(Ct/Cq) >40
215	34	M	Negative	Negative(Ct/Cq) >40
216	55	M	Negative	Negative(Ct/Cq) >40
217	58	M	Negative	Negative(Ct/Cq) >40
218	67	F	Negative	Negative(Ct/Cq) >40
219	20	F	Negative	Negative(Ct/Cq) >40
220	42	M	Negative	Negative(Ct/Cq) >40
221	59	M	Negative	Negative(Ct/Cq) >40
222	12	M	Negative	Negative(Ct/Cq) >40

Director: 
Date: 2020-12-10
Seal of company signature 

File No.:K590516D
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Effective date:2020-12-10



MINISTERSTVO ZDRAVOTNICTVÍ
Palackého náměstí 375/4, 128 01 Praha 2

Praha 17. března 2021

Č. j.: MZDR 10786/2021-2/OLZP



MZDRX01F22QC

ROZHODNUTÍ

Ministerstvo zdravotnictví (dále jen „Ministerstvo“) jako orgán příslušný k rozhodnutí podle ustanovení § 12 odst. 1 písm. h) zákona č. 22/1997 Sb., o technických požadavcích na výrobky a o změně a doplnění některých zákonů, ve znění pozdějších předpisů ve spojení s § 4 odst. 8 nařízení vlády č. 56/2015 Sb., o technických požadavcích na diagnostické zdravotnické prostředky in vitro (dále jen „nařízení vlády“), na základě žádosti společnosti

Capital Unicorns SE,

se sídlem Václavské náměstí 832/19, Nové Město, 110 00 Praha 1, IČO: 082 88 241
(dále jen „žadatel“)

rozhodlo v souladu s ustanovením § 67 a násł. zákona č. 500/2004 Sb., správní řád, ve znění pozdějších předpisů (dále jen „správní řád“) tak, že

povoluje

žadateli uvést na trh a do provozu diagnostický zdravotnický prostředek in vitro **Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Device (saliva)**, jehož výrobcem HANGZHOU REALY TECH CO., LTD., se sídlem 4th Floor, #12 Building, Eastern Medicine Town, Xiasha Economic & Technology Development, 310018 Hangzhou, Zhejiang, P.R. China, pro použití laickou osobou

a stanovuje

po dobu platnosti tohoto rozhodnutí žadateli následující povinnosti k zajištění ochrany veřejného zdraví:

- zajistit, aby konečný laický uživatel testu byl informován, že toto povolení se nevztahuje na variantu testu, která využívá nazofaryngeálního odběru vzorku
- informovat odběratele o povinnosti v rámci testování zajistit při pozitivitě antigenního testu provedeného laickou osobou bezprostřední informování poskytovatele zdravotních služeb za účelem provedení konfirmačního testu,
- v případě zájmu odběratele zajistit proškolení určené osoby,
- hlásit Státnímu ústavu pro kontrolu léčiv každou nepříznivou událost, ke které během používání výrobku dojde.

Platnost povolení: **do 30. 4. 2021.**

O důvodnění:

I.

Dne 11. 3. 2021 požádal žadatel o udelení výjimky podle § 4 odst. 8 nařízení pro diagnostický zdravotnický prostředek in vitro určený k sebetestování na onemocnění COVID-19 pod obchodním názvem Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Device (saliva), HANGZHOU REALY TECH CO., LTD., pro účely zavedení celoplošného testování v České republice, jakožto diagnostického zdravotnického prostředku in vitro, pro který nebyl proveden postup podle § 4 odst. 1 až 4 nařízení a jehož použití je v zájmu ochrany zdraví. Žádost zdůvodňuje potřebou pravidelně testovat populaci za účelem včasného odhalení výskytu nových případů onemocnění COVID-19 ještě před jejich rozšířením v kolektivu.

K žádosti přikládá následující dokumentaci:

- a) Declaration of Conformity
- b) Návod k použití v českém jazyce
- c) Fotodokumentace
- d) Clinical Report

II.

Ministerstvo posoudilo předmětný diagnostický zdravotnický prostředek in vitro na základě žadatelem předložených informací jako dostatečně funkčně způsobilý a pro uživatele bezpečný.

Ministerstvo se ztotožňuje s potřebou pravidelně testovat veřejnost rychlými antigenními testy za účelem včasného odhalení výskytu nových případů onemocnění COVID-19 ještě před jejich rozšířením v kolektivech, což při absenci antigenních testů určených pro sebetestování na celém trhu EU není možné řešit jinak, než s použitím vhodných antigenních testů určených pro profesionální použití, jež budou k tomuto účelu použity za účelem odhalení pozitivních osob ve společnosti. Povolení se vztahuje pouze na neinvazivní způsoby odběru vzorku.

Za účelem podpory opatření k ochraně veřejného zdraví je žadateli uložena povinnost informovat odběratele o povinnosti při zjištěné pozitivitě antigenního testu provedeného laickou osobou kontaktovat vzdáleným přístupem (telefonicky, e-mailem apod.) závodního lékaře (poskytovatele pracovně – lékařských služeb) nebo registrujícího praktického lékaře, který rozhodne o provedení konfirmačního testu a zajistí komunikaci v rámci systému ISIN. Za účelem minimalizace rizika chyb v provedení odběru a interpretaci výsledků testů je žadateli uložena povinnost v případě zájmu odběratele zajistit proškolení osoby určené odběratelem.

S ohledem na potřebu dalšího vyhodnocování z hlediska bezpečnosti a funkční způsobilosti testů je výjimka z procesu posouzení shody udělena do 30. 4. 2021.

S ohledem na výše uvedené rozhodlo Ministerstvo tak, jak je uvedeno ve výroku tohoto rozhodnutí.



Poučení:

Proti tomuto rozhodnutí je možné podat v souladu s § 152 odst. 1 správního řádu u Ministerstva rozklad, a to ve lhůtě 15 dnů ode dne doručení. O rozkladu rozhoduje ministr zdravotnictví.

doc. MUDr. Jan Blatný, Ph.D.
ministr zdravotnictví
podepsáno elektronicky

Str. 3 z 3

Elektronický podpis - 18.3.2021

Certifikát autora podpisu :

Jméno : doc. MUDr. Jan Blatný, Ph.D.

Vydal : PostSignature Qualified C...

Platnost do : 13.12.2021 17:26:01-000 +01:0



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