



Declaration of Conformity



According to the In-vitro Diagnostic Medical Device Directive 98/79/EC

Manufacturer: Hangzhou Testsea Biotechnology Co.,Ltd

Address: Building 6 No. 8-2 Keji Road , Yuhang Street, Hangzhou -311121, China

Authorized Representative: Lotus NL B. V.

Address: T.a.v. de heer X. Wei Koningin Julianaplein 10 2595 AA's-Gravenhage

Product: COVID-19 Antigen Test

Model: TSCOVID-19AC

Classification: Other IVD

The manufacture, herewith, declares that the product as specified above meets the applicable provisions of the follow the Directive and standards and fulfil the obligations imposed by AnnexIII of Directive 98/79/EC. All supporting documentations is retained under the premise of authorized representative.

Directive:

In vitro Diagnostic Medical Device Directive: DIRECTIVE 98/79/EC OF THE EUROPEAN PARLLAMENT AND OF THE COUNCIL of October 1998 on invitro diagnostic medical device.

Standard:

All application harmonized standards(published in the Official Journal of the European Communities on 17th November 2017)

The above declaration of conformity is issued under the sole responsibility of the manufacture.

2020.09
杭州泰西生物技术有限公司
Signed for and on behalf of the manufacture
HANGZHOU TESTSEA BIOTECHNOLOGY CO., LTD

(Signature and Position)

