## **Declaration of Conformity**

ACON Biotech (Hangzhou) Co., Ltd. No.210 Zhenzhong Road, West Lake District, Hangzhou, P.R. China, 310030

We declare under our sole responsibility that the in vitro diagnostic device:

Flowflex SARS-CoV-2 Antigen Rapid Test (Self-Testing) REF No.: L031-118M5 EAN Code: 6921756492427

classified as self-testing of the directive 98/79/EC, meets all the provisions of the directive 98/79/EC on *in vitro* diagnostic medical devices which apply to it.

This declaration is according to Annex III.6 of the Directive and thus is based on approval by the notified body TÜV SÜD Product Service GmbH, Ridlerstraße 65 80339 MÜNCHEN, Germany, notified under No. 0123 to the EC Commission.

> Authorized Representative: MedNet GmbH Borkstrasse 10 48163 Muenster, Germany

This declaration is valid until expiration of EC certificate No. V9 042074 0032 Rev.00 Expiration Date: 2024-05-26

2021

Signed this <u>|</u> day of \_ in Hangzhou, Qhina

کلاnny You International Regulatory Affairs Senior Director ACON Biotech (Hangzhou) Co., Ltd.



ACON BIOTECH (HANGZHOU) CO., LTD. No.210 Zhenzhong Road, West Lake District, Hangzhou, P.R. China, 310030