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 (Nasal/Saliva)

classified as Others according to the Annex II 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 (excluding Section 6) of the Directive.

Authorized Representative: MedNet GmbH Borkstrasse 10 48163 Muenster, Germany

This	Declaration	of Conformity	is valid	until 25	May, 2022.	

Signed this <u>5</u> day of <u>3</u> , <u>シン</u> in Hangzhou, China

Junny 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 Tel: +86-571-87963569 Fax: +86-571-87963570 E-mail: css@aconlab.com.cn