



DECLARATION OF CONFORMITY

Manufacturer: Joinstar Biomedical Technology Co.,Ltd.

Address:10th Floor,Administration Building,NO.519,XingGuo RD.,Yuhang Economic and Technological Development Zone, Hangzhou, Zhejiang, China, 311188

EC Representative's Name: Lotus NL B.V.

EC Representative's Address: Koningin Julianaplein 10,1e Verd, 2595AA, The Hague, Netherlands.

Declares, that the product

Product Name and Model:

COVID-19 Antigen Rapid Test (Colloidal Gold)

1Test/Kit, 5Tests/Kit, 10Tests/Kit, 20Tests/Kit, 25Tests/Kit, 200Tests/Kit.

as described above are in conformity with the requirements as defined in IVDD98/79/EC Annex III.

Additional information:


Conformity assessment route: Directive 98/79/EC, Annex III

Classification: List Others

I, the undersigned, hereby declare that the medical devices specified above conform with the Directive 98/79/EC on in vitro diagnostic medical devices and pertinent essential requirements.

Date Signed:

2021.05.12



Zhong Wang

Management Representative

Joinstar Biomedical Technology Co.,Ltd.



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