

EC Certificate No. 1434-IVDD-467/2021

EC Design-examination
Directive 98/79/EC concerning
in vitro diagnostic medical devices

Polish Centre for Testing and Certification certifies that manufactured by:

Xiamen AmonMed Biotechnology Co., Ltd Unit 503, 120 Xinyuan Road, Haicang District, Xiamen, Fujian, China

in vitro diagnostic medical devices for self-testing

COVID-19 Antigen Rapid Test Kit (Colloidal Gold) Saliva specimen CG01Ag-01S-ST, CG01Ag-05S-ST, CG01Ag-25S-ST

in terms of design documentation, comply with requirements of Annex III (Section 6) to Directive 98/79/EC (as amended) implemented into Polish law, as evidenced by the audit conducted by the PCBC

Validity of the Certificate: from 15.10.2021 to 27.05.2024

The date of issue of the Certificate: 15.10.2021

The date of the first issue of the Certificate: 15.10.2021



Issued under the Contract No. MD-128/2021 Application No: 233/2021 Certificate bears the qualified signature. Warsaw, 15/10/2021 Module A1 FBM-30-E 10

Vice-President