



## Ce Cert.

## **CERTIFICATE**

DIRECTIVE 98/79/EC
EC DESIGN-EXAMINATION

CeCert Sp. z o.o. hereby confirms that manufactured by

Safecare Biotech (Hangzhou) Co., Ltd.

Building 2/203, No. 18 Haishu Rd., Cangqian Sub-district, Yuhang District, Hangzhou, 311121, Zhejiang, P.R. China

in vitro diagnostic medical device for self-testing

## COVID-19 & Influenza A+B Antigen Combo Rapid Test

catalogue number: FCO-6032H

in term of the design conforms to the requirements of Annex III section 6 to Directive 98/79/EC (as amended) implemented into Polish Law, as evidenced by the assessment conducted by CeCert Sp. z o.o.

 $\epsilon$ 

2934

Validity date: 29.04.2022 - 26.05.2025 Issue date: 29.04.2022

Check it



CeCert Sp. z o.o. ul. Żurawia 32/34 00-515 Warszawa Green volu

Kamil Szczurowski Director of in Vitro Diagnostic Medical Device Certification Department

www.cecert.pl e-mail: biuro@cecert.pl

Certificate no: CeCert/063/W/E.1



