



Declaration of Conformity

Manufacturer: **Shenzhen Microprofit Biotech Co., Ltd.**
Rm. 405, 406, Zone B /4F, Rm. 205, 206-1, 207, West Side of Zone B/ 2F, Haowei Building, No. 8 Langshan 2nd Road, Songpingshan, Songpingshan Community, Xili Street, Nanshan District, Shenzhen, P.R. China

European Representative: CMC MEDICAL DEVICES & DRUGS, S.L.
C/ Horacio Lengo n18 · C.P 29006 · Málaga-Spain

Product Name: fluorecare SARS-CoV-2 & Influenza A/B & RSV Antigen Combo Test Kit

Common Name: SARS-CoV-2 & Influenza A/B & RSV Antigen Combo Test Kit (Colloidal Gold Chromatographic Immunoassay)

Brand: fluorecare®

Catalogue No.: MF-71-1, MF-71-2, MF-71-5

Classification: Self-testing Device of IVDD 98/79/EC

Conformity Assessment Route: Annex III of IVDD 98/79/EC

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|------------------------------|------------------------|---------------------|
| STANDARDS APPLIED | EN 13612:2002/AC: 2002 | EN ISO 13485:2016 |
| | EN ISO 14971:2012 | EN ISO 23640:2015 |
| | EN ISO 18113-1:2011 | EN ISO 18113-2:2011 |
| | EN ISO 15223-1:2016 | EN 13641:2002 |


We the manufacturer herewith declare on our solo responsibility that the above mentioned products meet the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer. The products comply with the essential requirements in accordance with Annex I of the IVDD 98/79/EC.

DIRECTIVES

General applicable directives:

In Vitro Diagnostic Medical Device Directive: COUNCIL DIRECTIVE 98/79/EC of 27 October 1998 concerning in vitro diagnostic medical devices (IVDD 98/79/EC).

Notified Body: CeCert

Identification number: 

(EC) Certificate(s): CeCert/092/W/E.2

Expire date of the Certificate: 2025.05.26

DATE OF ISSUE: 2022.05.18

SIGNATURE: