

Declaration of Conformity

Manufacturer Name SD Biosensor, Inc.

Manufacturer Address Head Office

C-4th&5th, 16, Deogyeong-daero, 1556beon-gil, Yeongtong-gu,

Suwon-si, Gyeonggi-do 16690, KOREA

Manufacturing Site

74, Osongsaengmyeong 4-ro, Osong-eup, Heungdeok-gu, Cheongju-si, Chungcheongbuk-do 28161, KOREA

MT Promedt Consulting GmbH EC Representative Name

EC Representative Address Altenhofstrasse 80 66386 St. Ingbert Germany

Rapid Test Kit Common Name

Product Name STANDARDTM Q COVID-19 Ag Test

*Please refer to "Annex I. Product List" on page 2 in more detail.

Q-NCOV-01G Reference Number

Classification Others not covered by Annex II and self-testing according to

Directive 98/79/EC

Conformity Assessment Route Annex III of Directive 98/79/EC (EC Declaration of Conformity)

EN ISO 13485:2016 EN ISO 18113-1:2011 **Applied Standards**

> EN ISO 14971:2012 EN ISO 18113-2:2011 EN ISO 23640:2015 EN ISO 15223-1:2016

EN ISO 17511:2003 EN 62366:2008

EN 13612:2002

We herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for in vitro diagnostic medical devices. All supporting documentations are retained under the premises of the manufacturer. SD Biosensor, Inc. is exclusively responsible for the declaration of conformity.

Place: Suwon-si, Republic of Korea

Valid from: March 03, 2020

Signature

Hyo-Keun, Lee CEO / President



Annex I. Product List

<u>Q-NCOV-01G</u>	EDMA Code	Description of EMDA code
STANDARD TM Q COVID-19 Ag Test	15 70 90 90 00	Other Other Virology Rapid Tests
- Test Device (individually in a foil pouch with desiccant)		
- extraction buffer tube		
- Filter cap		
- Sterile swab A		
- Sterile swab B (option)		