

Declaration of Conformity

according to Directive 98/79/EC, on in vitro diagnostic medical devices

Maker

(Name, Address)

Getein Biotech, Inc.

No. 9 Bofu Road, Luhe District, Nanjing, 211505, China

Authorized Representative

(Name, Address)

Lotus NL B.V.

Koningin Julianaplein 10, 1e Verd, 2595AA, The Hague, Netherlands.

Medical device

One Step Test for Novel Coronavirus(2019-nCoV) IgG antibody (Colloidal Gold)

One Step Test for Novel Coronavirus(2019-nCoV) IgM antibody (Colloidal Gold)

One Step Test for Novel Coronavirus(2019-nCoV) IgM/IgG antibody (Colloidal Gold)

Classification

Others

Applicable coordination standards

EN ISO 14971:2012
EN 13612:2002
EN 1041:2008

EN ISO 23640:2015
EN ISO15223-1:2012
EN ISO 18113-1:2011

EN ISO 13485:2016
EN ISO 18113-2:2011
EN ISO 18113-3:2011

Signatory representative declares herein the above mentioned device meets the basic requirements of the European Parliament and the Council's in vitro diagnostic medical devices directive: 98/79/EC Annex III. This declaration of conformity is based on European Parliament and the Council's 98/79/EC directive Annex III. The compiled technical file and quality system document according to 98/79/EC directive Annex III are testified and the quality system certificate has issued by TÜV Rheinland (Shanghai) Co., Ltd.

General Manager Enben Su

Nanjing, 4th Mar, 2020
(place and date of issue)

(name and signature or equivalent marking of authorized person)

