



## DECLARATION OF CONFORMITY

**According Directive 98/79/EC on In Vitro Diagnostic Medical Devices, Annex III.**

**Manufacturer:** Anbio (Xiamen) Biotechnology Co.,Ltd.

**Address:** No.2016, Wengjiao West Road, Xinyang Street, Haicang District, 361026 Xiamen, Fujian, China.

**European Representative:** Lotus NL B.V.

**Contact person:** Peter      **E-mail:** peter@lotusnl.com

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

**In Vitro Diagnostic Directive:**

- **Rapid COVID-19 Antigen Test (Colloidal Gold )**

**Category:** Others.

**Conformity assessment route:** Declaration of Conformity IVDD Annex III

**Applicable Standards:**

ISO 13485:2016

ISO 14971:2019

EN ISO 18113-1:2011

EN ISO 18113-2:2011

EN ISO 18113-3:2011

EN 13641:2002

ISO 15223-1:2016

EN 13612:2002

ISO 23640:2015

EN 62366-1:2015

We, the manufacturer, herewith declare with sole responsibility that our product/s mentioned above meet/s the provisions of the Directive 98/79/EC of the European Parliament and of the Council on In Vitro Diagnostic Medical Devices.

We agree to develop, implement and maintain a documented post-production monitoring process.

Signed on: 04/09/2020

Place: Xiamen, China

Name of authorized signatory: *Danny Wang*

Position held in the company: General Manager

Seal/Stamp:

Anbio (Xiamen) Biotechnology Co., Ltd.



**BUREAU VERITAS**  
Certification



## **Anbio (Xiamen) Biotechnology Co., Ltd**

No.2016, Wengjiao West Road, Xinyang Street, Haicang District, Xiamen City,  
P.R. China

### **Certified site:**

NO.2016, WENGJIAO WEST ROAD, XINYANG STREET, HAICANG DISTRICT, XIAMEN CITY,  
P.R. CHINA

*Bureau Veritas Italia S.p.A. certifies that the Management System of the  
above organisation has been audited and found to be in accordance  
with the requirements of the management system standards detailed below*

## **EN ISO 13485:2016**

*Scope of certification*

Design and manufacture of fluorescence-based immunoassay reagent and devices, as an aid in clinical assessment of cardiovascular, gastric, inflammation, diabetic and infectious diseases detection, as well as hormone, vitamin testing. Design and manufacture of IVD reagents as an aid in clinical assessment of blood type testing.


Certificate awarded in conformity with the requirements of ACCREDIA DT 02-DC Rev.00

Original cycle start date:	<b>22/06/2020</b>
Expiry date of previous cycle:	<b>n.a.</b>
Certification / Recertification Audit date:	<b>17/05/2020</b>
Certification / Recertification cycle start date:	<b>22/06/2020</b>

Subject to the continued satisfactory operation of the organization's  
Management System, this certificate expires on: **21/06/2023**

Certificate No. - Version: **IT298645-1**

Revision date: **22/06/2020**

  
**GIORGIO LANZAFAME - Local Technical Manager**

Certification body address:  
Bureau Veritas Italia S.p.A., Viale Monza, 347 - 20126 Milano, Italia



SGQ N° 009A  
Member of the Italian Association of Accredited Bodies (AIAA) and of the European Association of Accredited Bodies (EAAB)

Further clarifications regarding the scope of this certificate and the applicability of the  
management system requirements may be obtained by consulting the organisation.  
To check this certificate validity please refer to the website [www.bureauveritas.it](http://www.bureauveritas.it)