



## EC DECLARATION OF CONFORMITY

According to the *In Vitro* Diagnostic Medical Device Directive 98/79/EC

**Manufacturer:** Guangzhou Wondfo Biotech Co., Ltd.  
**Address:** No.8, Lizhishan Road, Science City, Luogang District,  
510663, Guangzhou, P.R. China

**EC Authorised Representative:** Qarad BV  
**Address:** Ciplastraat 3, 2440 Geel, Belgium

***In Vitro* Diagnostic Medical Device(s):**

**Product Name:** Finecare™ FIA Meter Plus  
**Model No.:** FS-113  
**IVDD Classification:** Other, for professional use

This declaration of conformity is issued under the sole responsibility of the manufacturer that the above product(s) meet(s) the provisions of the European Directive 98/79/EC for *In Vitro* Diagnostic Medical Devices, Directive 80/181/EEC and RoHS Directive 2011/65/EU.

The following (harmonized) standards have been applied:

|                     |                     |                  |
|---------------------|---------------------|------------------|
| EN ISO 13485:2016   | EN 61010-1: 2010    | EN 13612:2002    |
| EN ISO 14971:2019   | EN 61010-2-081:2015 | EN 62304:2006    |
| EN ISO 18113-1:2011 | EN 61010-2-101:2002 | EN 62366-1:2015  |
| EN ISO 18113-3:2011 | EN 61326-2-6: 2013  | EN 61326-1: 2013 |
| EN ISO 15223-1:2016 | EN 62321 Series     |                  |

The conformity with the requirements of the Directive has been assessed following the procedure(s) outlined in the following annexes of the Directive: Annex III, excluding 6

**Notified Body (if consulted):** Not applicable  
**Address:** /  
**EC Certificate(s):** /  
**Expiry date of the Certificate(s):** /

**Signature of manufacturer (Name and function):**

Vice-President of Regulatory Affairs

**Place and date of issue:** Guangzhou, P.R. China

December 10, 2021