



# Declaration of Conformity



according to the In vitro diagnostic Medical Devices Directive 98/79/EC

**Manufacturer:** Changsha Sinocare Inc.

No.265 Guyuan Road, Hi-tech Zone, Changsha, Hunan Province,  
410205, People's Republic of China.

**Address:**

**We declare under our sole responsibility that**

**the Medical  
Device**

Product Name : Blood Glucose monitoring system

Type/model,  
batch/serial number, Safe AQ Angel Blood Glucose Meter  
possibly sources and : Safe AQ Angel Blood Glucose Strip  
number of items Blood Glucose Control Solution  
(Where applicable)

**of class**

according to directive : The product classification is ListB.  
98/79/EC

meets all the provisions of the directive 98/79/EC which apply to it.

Applied  
harmonised  
standards, national  
standards or other  
normative  
documents

## EN ISO 15197:2015

In vitro diagnostic test systems  
requirements for blood-  
Glucose monitoring systems for  
Self-testing in managing  
diabetes mellitus(ISO  
15197:2013)

## EN23640:2013

In vitro diagnostic medical  
devices-evaluation of stability of  
in vitro diagnostic reagents (ISO  
23640:2011)

## IEC 61326-1: 2012

Electrical equipment for  
measurement, control and  
laboratory use - EMC  
requirements - Part 1: General  
requirements

## IEC-61326-2-6: 2012

Electrical requirements for  
measurement, control and  
laboratory use -EMC  
requirements- Part 2-6:  
Particular requirements -In vitro  
diagnostic (IVD) medical  
equipment

## EN ISO14971:2012

Medical devices - Application of  
risk management to medical  
devices (ISO 14971:2007,  
Corrected version 2007-10-01)

## IEC 61010-1: 2010

Safety requirements for electrical  
equipment for measurement,  
control, and laboratory use - Part  
1: General requirements

## IEC 61010-2-101: 2015

Safety requirements for electrical  
equipment for measurement,  
control, and laboratory use - Part  
2-101: Particular requirements for  
in vitro diagnostic (IVD) medical  
equipment

## EN ISO 18113-1:2011

In vitro diagnostic medical  
devices - Information supplied by  
the manufacturer (labelling) - Part  
1: Terms, definitions and general  
requirements (ISO 18113-1:2009)

**EN ISO 18113-4:2011**

In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 4 : In vitro diagnostic reagents for self-testing (ISO18113-4:2009)

**IEC 62366:2007(First Edition)+A1:2014**

Medical devices-Application of usability engineering to medical devices

**EN ISO 18113-5: 2011**

In vitro diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 5: In vitro diagnostic instruments for self-testing (ISO 18113-5:2009)

**IEC 62304 : :2015**

Medical device software- Software life cycle processes

Conformity assessment procedure Council directive 98/79 /ECAnnex IV except setion 4、 6

Notified Body (if consulted) TÜV SÜD Product Service GmbH  
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Li Shaobo  
General Manager

2/28/2017  
(place and date)

Li Shaobo.  
(name and signature, function)

