

**DECLARATION OF CONFORMITY
TO COUNCIL DIRECTIVE 98/79/EC CONCERNING
In VITRO DIAGNOSTIC MEDICAL DEVICES**

Manufacturer: **BioCare Corporation**
4F, No.12, Ln. 5, Sec. 2, Nanshan Rd., Luzhu Dist., Taoyuan City
33852, Taiwan

European Representative: **MedNet GmbH**
Borkstrasse 10, 48163 Muenster, Germany

Product Name: **Blood Glucose Monitoring System**
Trademark / Model:

Blood Glucose Monitoring System : DIAVUE ToGo
Blood Glucose Meter : DIAVUE ToGo
Blood Glucose Test strip : DIAVUE ToGo
Control Solution : DIAVUE Y1 、 DIAVUE W2 、 DIAVUE B3

GMDN(UMDNS) Code: **16488**

Classification (IVDD, Annex II): **98/79/EC (IVDD) Annex II, List B (Self-Testing)**

Conformity Assessment Route: **98/79/EC (IVDD) Annex IV, Section 3**

We herewith declare that the above mentioned products meet the transposition into national law, the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer. Manufacture is exclusively responsible for DoC.

DIRECTIVES

General applicable directives:

Medical Device Directive: COUNCIL DIRECTIVE 98/79/EC concerning medical devices (IVDD 98/79/EC).

Notified Body: **TÜV SÜD Product Service GmbH, Ridlerstr. 65, 80339**
München, Germany

Identification number: **CE0123**

(EC) Certificate(s): **V1 077360 0016 Rev.01**

Expire date of the Certificate: **2022-01-22**

Start of CE Marking: **2013-01-01**

Place, Date of Issue: **Taiwan, April 29, 2019**

Signature: _____

Name: **Terry Liu**

Position: **Management Representative**

