EU Declaration of Conformity

Name and address of the manufacturer: Shandong Lianfa Medical Plastic Products Co., Ltd.

No.1 Shuangshan Sanjian Road, 250200, Zhangqiu City, Jinan,

Shandong, PEOPLE' S REPUBLIC OF CHINA

EC Authorized Representative: Linkfar Healthcare GmbH

Niederrheinstraße 71, 40474 Düsseldorf, Germany

SRN CODE DE-AR-000005107

We declare under our sole responsibility that:

The medical device: Lancing Device

UMDNS code: 18866 Basic UDI-DI 694951701VA

Trade name / Classification |

According to annex VIII(Rule 1) of Regulation EU 2017/745(MDR)

Intended use Assisting the lancet for peripheral blood sampling form the fingertip or the

paw for blood Glucose testing or other testing ulilising a small amount of blood, it is the non-invasive device and does not come into contact with blood, it only makes a brief contact with the skin surface for a few seconds.

Model: A,B,C,D,E,F,G

We,the manufacture herewith declare that the above-mentioned products meet the provisions of the Regulation EU 2017/745(MDR) . The products meet prospective uses and all supporting documentation is retained under the premise of manufacturer.

Shandong Lianfa Medical Plastic Products Co., Ltd is solely responsibility for the Declaration of Conformity.

Conformity assessment procedure: Annex II & Annex III of Regulation EU 2017/745(MDR)

Declaration of Conformity is valid until: 2024-12-29

Signature:_

Name: Yang Lianying Position: President

Place, Date: Jinan, 2023-12

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