EC DECLARATION OF CONFORMITY

We, as the Manufacturer, certifies that the following medical device:

LATEX TOURNIQUET, LATEX FREE TOURNIQUET, LATEX FINGER COT Classification: Classified as class I according to Annex IX, rule 1 of the Directive 93/42/EEC

meets all applicable requirements of the Medical Devices Directives

Directive Name / Number

93 / 42 / EEC and 47/2007/EC

The declaration is sole responsibility of the manufacturer

Name of manufacturer Jiangsu High Hope International Group Sunshine I/E Corp. No. 50,Zhonghua Road,Nanjing,China

The Authorized Representative within EU who has been empowered to enter into commitments on our behalf:

Name of Representative in EU:

MedNet GmbH Borkstrasse 10,48163 Muenster,Germany

Date: July 15,2018

Signature: Zhu Baoyu General Manager

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