EC DECLARATION OF CONFORMITY

Name and address of the

manufacturer:

Shenzhen Viatom Technology Co., Ltd. 4E,Building 3, Tingwei Industrial Park, No.6 Liufang Road, Block 67, Xin'an Street, Baoan District, 518101 Shenzhen, P.R.China

Name and address of Authorized

Representative:

MedNet EC-REP GmbH

Borkstrasse 10, 48163 Muenster, Germany

Telefon: +49 251 32266-61 Telefax: +49 251 32266-22

We declare under our sole responsibility that

the medical device: Pulse Oximeter

Model: O2 RING, Wear O2

UMDNS 17148

of class: Class IIa

according to annex IX of directive 93/42/EEC meeting all the provisions of the Directive 93/42/EEC, as amended by Directive 2007/47/EEC, which apply to it.

Conformity assessment procedure: MDD 93/42/EEC Annex II excluding (4)

Registration No.: HD 601373560001

Notified Body: TÜV Rheinland LGA Products GmbH

Tillystraße 2 90431 Nürnberg Deutschland CE 0197

Shenzhen, 2021/03/22

Place, date

General Manager Zhou Saixin

Name and function