Declaration of Conformity

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SunTech Medical, Inc.

5827 South Miami Boulevard, Suite 100

Morrisville, NC 27650-8394

suntechmed.com

EU Rep:

EMERGO Europe Prinsessegracht 20 2514 AP The Hague The Netherlands

Product Name:

NBP One

Model Number:

250D

Description:

Non-Invasive Ambulatory Blood Pressure Monitor and AccuWin Pro V4

Software

Classification:

NBP One System: Class IIa, Rule 10 Orbit ABPM Cuff: Class 1, Rule 1

Assessment Procedure

NBP One System: Annex II Orbit ABPM Cuff: Annex VII

Notified Body

Intertek Medical Notified Body AB Torshamnsgatan 43, Box 1103

SE-162 22 Kista

Sweden

(€ 0413

Product Marking

The above NBP One ABPM system complies with Council Directive 93/42/EEC, with LVFS 2003:11 transposing European Medical Devices, amendments to Council Directive 93/42/EEC outlined in Directive 2007/47/ED, Annex I (Essential Requirements) and Annex II (EC Declaration of Conformity -Quality System Production), and with WEEE Directive 2002/96/EC, and with the European RoHS Directive 2011/65/EU.

I, the undersigned, declare on the basis of the above information that the system described above is in compliance with the requirements of Directive 93/42/EEC. This declaration hereby authorizes the CE Mark to be affixed to the above-mentioned product(s).

DocuSigned by:

Signer Name: Tonia Bryant Signing Reason: I approve this document Signing Time: 7/26/2022 | 10:28:51 AM PDT

7/26/2022 Date:

Reviewed and Approved by:

Tonia E. Bryant, Regulatory Affairs 400 msag 324 B47A18C3113C71002CECD

