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Declaration of Conformity

MANUFACTURER'S DECLARATION OF CONFORMITY

AUSTRALIAN THERAPEUTIC GOODS (MEDICAL DEVICES) REGULATIONS 2002

FULL QUALITY ASSURANCE PROCEDURES

This is a declaration of conformity made under clause 1.8 of Schedule 3 to the Therapeutic Goods (Medical Devices) Regulations 2002.

| Manufacturer's name: | SunTech Medical, Inc. | |
|-----------------------|--|--|
| Business address: | 5827 South Miami Blvd, Suite 100, Morrisville, NC 27560 | |
| Medical device(s): | Non-Invasive Ambulatory Blood Pressure device, Oscar 2, Model 250D (Bravo Mini) | |
| Classification: | Па | |
| GMDN code and term: | 36888 – Blood pressure ambulatory recorder | |
| Scope of application: | 99-0233-XX (where -XX indicates any number 00 to 99); NJXXX (where -XXX indicates any number 000 to 999) | |
| Basic UDI: | 08409351000000000250D92 | |

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

| Full quality assurance procedures certificate: | ISO 13485:2016 / MDSAP Certificate 006 | 58171-05 | | |
|--|--|----------|-----------|--|
| Design examination certificate (if applicable): | None | | | |
| Standards applied: | See attached Schedule A | | | |
| Authorised signatory: | | | | |
| Signed by: | | | | |
| Tonia Bryant | | | | |
| Signer Name: Tonia Bryant Signing Reason: Lapprove this document Signing Time: 3/24/2025 11:53:55 AM PDT 3/24/2025 | | | 3/24/2025 | |
| 74D91508594B47A18C3113C71002CECD | | | | |
| Tonia E. Bryant, Regulatory Affairs Director | or Da | ate | | |
| Declaration of Conformity – Schedule A | | | | |

Standards Applied:

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| Safety | | Medical electrical equipment – Part 1: General requirements for basic safety and essential performance | |
|---------------------------|-------------------------------|---|--|
| Performance/Safety | | Medical electrical equipment – Part 2-30: Particular requirements for the basic safety and essential performance of automated non-invasive sphygmomanometers | |
| | | Non-invasive sphygmomanometers — Part 1: Requirements and test methods for non-automated measurement | |
| | | Medical electrical equipment - Part 1-11: General requirements for basic safety and essential performance — Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment | |
| EMC/EMI/ ESD | | Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance – Collateral Standard: Electromagnetic disturbances – Requirements and tests | |
| Software | IEC 62304: Ed. 1.1 (2015) | Medical device software – Software life cycle processes | |
| Usability | | Medical electrical equipment – Part 1-6: General requirements for basic safety and essential performance – Collateral standard: Usability | |
| | EN 62366-1: 2015+AMD1:2020 | Medical devices – Part 1: Application of usability engineering to medical devices | |
| Clinical | | Non-Invasive sphygmomanometers - Part 2 Clinical investigation of intermittent automated measurement type | |
| Biocompatibility (System) | | Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk managemen process | |
| Risk Management | ISO 14971:2019 | Medical devices — Application of risk management to medical devices | |
| Quality System | ISO 13485:2016 | Medical devices – Quality management systems – Requirements for regulatory purposes | |
| Symbols | | Medical Devices – Symbols to be used with information to be supplied by the manufacturer – Part 1 General requirements | |
| Information | ISO 20417:2021 | Medical devices - Information to be supplied by the manufacturer | |