




## Declaration of Conformity

  <b>SRN:</b>	SunTech Medical, Inc. 5827 South Miami Blvd, Ste 100 Morrisville, NC 27560 suntechmed.com USA  US-MF-000002189	<div>EC REP</div>          <b>SRN:</b>	Emergo Europe Westervoortsedijk 60 6827 AT Arnhem The Netherlands  NL-AR-000000116
<b>Product Name:</b>  	Disposable Blood Pressure Cuff and Single Patient Use Kits  DC100 & DC200	<b>Basic</b> <div>UDI</div>  <div>REF</div>	084093510000000000DC100H9  98-040X-XX, 98-050X-XX and 98-0700-XX (where X and - XX indicates any alphanumeric character 0 to 9 or A-Z)
<b>Description:</b>  <b>Intended Purpose:</b>	Disposable Blood Pressure Cuff  The Disposable Blood Pressure cuff is intended to be used with non-invasive blood pressure measurement systems to determine blood pressure parameters on neonate, pediatric and adult patients.  Single Patient Use (SPU) Kits contain a Disposable Cuff with an adhesive pad.		
<b>Classification:</b>	Class I, Rule 1	<b>Assessment Procedure:</b>	Annex II and III
<b>Notified Body:</b>	N/A	<b>Product Marking:</b>	
<b>GMDN Code and Term</b>	34978 - Blood pressure cuff, reusable	<b>UMDNS Code and Term</b>	11703- Devices that have an inflatable bladder in an inelastic sleeve (cuff) with a mechanism for inflating and deflating the bladder. These devices are used in conjunction with another device to determine a patient's blood pressure.

This declaration of conformity is issued under the sole responsibility of SunTech Medical Inc. The above system complies with MDR 2017/745 requirements, in accordance with Annex I (General Safety and Performance Requirements), Annex II (Technical Documentation), Annex III (Post-Market Surveillance), and Annex IV (EC Declaration of Conformity), WEEE Directive 2012/19/EU, the RoHS Directive 2015/863/EU. This declaration is supported by the Quality System approval to ISO 13485 issued by Intertek. All supporting documentation is retained at the premises of the manufacturer.

I, the undersigned, declare on the basis of the above information that the system described above is in compliance with the requirements of MDR Directive 2017/745. This declaration hereby authorizes the CE Mark to be affixed to the above-mentioned product(s).

Signed by:

*Tonia Bryant*



Signer Name: Tonia Bryant

Signing Reason: I approve this document

Signing Time: 12/12/2024 | 8:46:22 AM PST 12/12/2024

Reviewed and Approved by: 74D91508594B47A18C3113C71002CECD  
Tonia E. Bryant, Director of Regulatory Affairs

Signed at SunTech Medical, Inc, Morrisville, NC 27560

## Attachment to Declaration of Conformity

### Device variants

**REF**

#### Description

98-040X-XX 98-050X-XX	Disposable Cuff, various sizes, with 1 or 2 tubes, with various connectors and package sizes. (where X and -XX indicates any alphanumeric character 0 to 9 or A-Z)
98-0700-XX	Single Patient Use Kits (where -XX indicates any alphanumeric character 0 to 9 or A-Z)

### Standards Applied:

Cleaning/Disinfection	ISO 17664:2017	Processing of health care products – Information to be provided by the medical device manufacturer for the processing of medical devices
Safety	IEC 80601-2-30:2018	Medical Electrical Equipment - Part 2-30: Particular Requirements For The Basic Safety And Essential Performance of sphygmomanometers
Biocompatibility	EN ISO 10993-1:2018	Biological Evaluation of Medical Devices-Part 1: Evaluation and testing within a risk management process
Symbols	ISO 15223-1:2021	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
Quality System	EN ISO 13485:2016	Medical Devices – Quality Management Systems – Requirements for Regulatory Purposes
Risk Management	EN ISO 14971:2019	Medical devices – Application of risk management to medical devices