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## Declaration of Conformity

(For Britain, Wales & Scotland)

	SunTech Medical, Inc.		International Associates Limited	
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	100	UK REP	Glasgow,	
	Morrisville, NC 27560		Lanarkshire, G1 3DX, UK	
	suntechmed.com		UKRP@ia-uk.com	
	USA			
SRN:				
	US-MF-000002189			
Product Name:	Oscar 2	UDI	08409351000000000002507E	
		Basic		
#	250		99-0133-XX, (where -XX indicates any	
		REF	number 00 to 99).	
Description:	Non-Invasive Ambulatory Bloo	d Pressure device		
Intended	The Operation 2 Medel 250 events	m ia o non-invoci	we esseillemetric embulatory blood	
Purpose:	The Oscar 2, Model 250 system is a non-invasive oscillometric ambulatory blood			
Fulpose.	pressure monitor that is intended to be used with AccuWin Prom, a PC-based computer			
	program for the recording and displaying of up to 250 measurements of systolic and			
	diastolic blood pressure and heart rate. It is intended for use as an aid or			
	adjunct to diagnosis and treatment when it is necessary to measure an adult and			
	pediatric (> 3yrs.) patient's systolic and diastolic blood pressures over an			
	extended period of time. The system is only for measurement, recording, and display. It makes no diagnoses.			
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	Optionally, the Model 250 will provide a derived ascending aortic blood pressure			
	waveform and a range of central arterial indices. These measurements are provided			
	non-invasively through the use of a brachial cuff. It is to be used on those			
	patients where information related to ascending aortic blood pressure is desired,			
	but the risks of cardiac catheterization procedure or other invasive monitoring may			
	outweigh the benefits (excludes pediatric subjects). Bluetooth wireless			
	connectivity may be offered	-	-	
Classification:	Class IIa, Rule 10	Assessment	Annex II (with the exception of	
		Procedure:	section 4)	
Notified Deduc	Intertek Medical Notified	Product		
Notified Body:			~ ~	
	Body AB	Marking:	CE	
	Torshamnsgatan 43, Box		0413	
	1103 SE-162 22 Kista			
	SE-162 22 Kista Sweden			
GMDN Code and		UMDNS Code and	18364 - Recorder, physiologic, blood	
	36888 - Blood pressure			
Term	ambulatory recorder	Term	pressure	

This declaration of conformity is issued under the sole responsibility of SunTech Medical Inc. The above system complies with the requirements set out in the Medical Devices Regulations 2002 (SI 2002 No 618, as amended) (UK MDR 2002), in accordance with Annex I (Essential Requirements), Annex II (EC Declaration of Conformity - Quality System Production), with 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 the Medical Device Directive 93/42/EEC. This declaration hereby authorizes the CE Mark to be affixed to the above-mentioned product(s).



Reviewed and Approved by: Tonia E. Bryant, Manager, Regultable 2015/08594847418C3113C71002CECD

Signed at SunTech Medical, Inc, Morrisville, NC 27560 Document Expiry Date: 20 June 2023 (maximum of 1 year upon release)

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### Attachment to Declaration of Conformity

Device variants

Description99-0133-00System, Oscar 2, Model 250, Standard99-0133-01System, Oscar 2, Model 250, Standard, Bluetooth99-0133-02System, Oscar 2, Model 250, Central BP99-0133-03System, Oscar 2, Model 250, Central BP, Bluetooth99-0133-10System, Oscar 2, Model 250, Standard, without software99-0133-11System, Oscar 2, Model 250, Standard, Bluetooth Without Software99-0133-12System, Oscar 2, Model 250, Central BP w/o Software99-0133-13System, Oscar 2, Model 250, Central BP w/o Software99-0133-13System, Oscar 2, Model 250, Central BP, Bluetooth W/O SW

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Standards Applied:			
Safety	IEC 60601-	Medical electrical equipment - Part 1: General requirements	
	1: Ed. 3.1	for basic safety and essential performance	
	(2012)		
Performance/S	IEC80601-2-	Medical electrical equipment - Part 2-30: Particular	
afety	30: Ed. 2.0	requirements for the basic safety and essential performance	
	(2018)	of automated non-invasive sphygmomanometers	
	ISO 80601-	Medical electrical equipment - Part 2-56: Particular	
	2-56:2017	requirements for basic safety and essential performance of	
		clinical thermometers for body temperature measurement	
	ISO 80601-	Medical electrical equipment - Part 2-61: Particular	
	2-61:	requirements for basic safety and essential performance of	
	Ed.2.0	pulse oximeter equipment	
	(2017)		
	ISO 81060-	Non-invasive sphygmomanometers - Part 1: Requirements and	
	1: 2012	test methods for non-automated	
		measurement	
EMC/EMI/	IEC 60601-	Medical electrical equipment - Part 1-2: General	
ESD	1-2: Ed.	requirements for basic safety and essential performance -	
	4.0 (2014)	Collateral Standard: Electromagnetic disturbances -	
		Requirements and tests	
Software	IEC 62304:	Medical device software - Software life cycle processes	
	Ed. 1.1		
	(2015)		
Usability	IEC 60601-	Medical electrical equipment - Part 1-6: General	
	1-6:2010	requirements for basic safety and essential performance -	
	+A1:2015	Collateral standard: Usability	
	EN 62366-1:	Medical devices - Part 1: Application of usability	
	2015	engineering to medical devices	
Clinical	IEC 81060-	Non-Invasive sphygmomanometers - Part 2 Clinical	
	2: 2018	investigation of intermittent automated measurement type	
Biocompatibil	ISO 10993-	Biological evaluation of medical devices - Part 1:	
ity (System)	1: 2018	Evaluation and testing within a risk management process	
	ISO 10993-	Biological evaluation of medical devices - Part 5: Tests for	
	5: 2009	in vitro cytotoxicity	
	ISO 10993-	Biological evaluation of medical devices - Part 10: Tests	
	10: 2010	for irritation and skin sensitization	
Risk	ISO ISO	Medical devices - Application of risk management to medical	
Management	14971:2019	devices Application of fisk management to medical	
Quality	ISO 13485:	Medical devices - Quality management systems - Requirements	
System	2016	for regulatory purposes	
Symbols	ISO 15223-	Medical Devices - Symbols to be used with information to be	
	1:2021	supplied by the manufacturer - Part 1: General requirements	
Information	ISO	Medical devices - Information to be supplied by the	
	20417:2021	manufacturer	