

EC Declaration of Conformity

This declaration of conformity is issued under the sole responsibility of the manufacturer in accordance with the Council Directive 93/42/EEC in its currently valid version (Annex I) concerning medical devices.

Product: **Reusable Pulse Oximetry Sensor
Fingerclip**

Typ:	F-2182-3 Minolta	F-3212-9 Nellcor	F-3226 EnviteC	F-3512-20 Datex
	F-2203-16 Nihon Kohden	F-3212-31 Nellcor	F-3226-20 EnviteC	F-3512-40 Datex
	F-2412 HP/Philips	F-3212-31 M Nellcor	F-3227 EnviteC	F-3512-9 Datex
	F-2414-15 HP/Philips	F-3213-30 Mindray	F-3412 Ohmeda	F-3513-30 GE Datex-O hmeda
	F-3003-9 FlexMon	F-3213-9 Mindray	F-3412-10 Ohmeda	
	F-3012 Nonin	F-3222-12 BCI	F-3412-9 Ohmeda	
	F-3012-30 Nonin	F-3222-24 BCI		
	F-3013-30 Nonin	F-3222-30 BCI		
	F-3015-30 Nonin			

Classification:
(RL 93/42/EEC,
Annex IX)

Class IIb

CE mark:



Notified Body: TÜV SÜD Product Service GmbH, Ridlerstr. 65, 80339 Munich, Germany

Conformity
Assessment Process:

Annex II, section 3 of the Directive 93/42/EEC

Particular product
standard applied:

ISO 80601-2-61

Issued by:

EnviteC-Wismar GmbH
Alter Holzhafen 18
D-23966 Wismar
Germany

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2015-07-01

Valid until:

2018-06-30

Place, Date:

Wismar, 2015-07-01

Authorized Signature:

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