

# EC Declaration of Conformity

We hereby declare under our sole responsibility  
that the product

## VM-2160 with accessories

**Handheld Pulse Oximeter for continuous or spot check monitoring of functional arterial  
oxygen saturation (SpO<sub>2</sub>) and pulse rate (PR)**

REF	Type	Product
12020112001E	SMARTsat® Technology	VM2160 - 0012165 complete device - Central Europe
12020112001N	SMARTsat® Technology	VM2160 - 0012166 complete device - Scandinavia
12020112001S	SMARTsat® Technology	VM2160 - 0012167 complete device - Special Europe

Complies with the essential requirements of Annex I and Annex II of the Council Directive 93/42/EEC as amended by 2007/47/EC concerning medical devices as well as the requirements of Regulation (EU) 2017/745, Article 120, Chapter (3).

In accordance with Annex IX of the Council Directive 93/42/EEC the product has been  
classified as Class IIb.

Application of the CE-marking:



MEDCERT GmbH, Pilatuspool 2, 20355 Hamburg

**Validity:**

**Date of expiry: 02 November 2023**

**Issuer:**

bluepoint medical GmbH & Co. KG  
An der Trave 15  
23923 Selmsdorf  
Germany

**Place, Date:**

Selmsdorf, 25 May 2021

**Legally binding signature:**

  
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Bernd Lindner  
General Manager