

EC Declaration of Conformity

We hereby declare under sole responsibility that the product

VM-2103

Finger Oximeter for monitoring of functional
arterial oxygen saturation (SpO₂) and pulse rate,

Product No.
0012103

conforms with the essential requirements of Annex II of the Council Directive 93/42/EEC
of 14 June 1993 considering the amendments by the directive 2007/47/EC as
transposed into national law in the countries where the medical product is intended to be
marketed.

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

Application of the CE-marking:

CE 0086

Issuer:

Viamed Ltd.
15 Station Road
Cross Hills
Keighley
West Yorkshire, BD20 7DT
United Kingdom

Place, Date:

Keighley, 26 April 2012

Legally binding signature:


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Derek Lamb (Managing Director)