

## DECLARATION OF CONFORMITY

### Medical Device(s)

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

Device(s): Supramaximal Nerve Stimulator

Class: IIa

Part number: 2510000

Description: Supramaximal Nerve Stimulator

It is hereby declared that the medical device(s) specified above is (are) in compliance with EC Council Directive 93/42/Eec (Medical Devices Directive) of the 14<sup>th</sup> June 1993 and considering the amendments by directive 2007/47/EC - as transposed into National Law in the countries where the product is intended to be marketed.

This declaration is supported by:

Technical documentation required by MDD ( Annex II ) retained by: Viamed Ltd.

EC Quality Assurance Certificate No. CE 01389, first issued by the British Standards Institute (CE0086) on the 23rd August 1996.

Certificate of Registration No. MD 78787 to ISO 13485:2003 of original registration date 27th January 2004, issued by the British Standards Institute (CE0086)

Certificate of Registration No. FM 540797 to ISO 13485:2003 (CMDCAS recognized) of original registration date 27th January 2004, issued by the British Standards Institute (CE0086)

Certificate of Registration No. FS 28344 to ISO 9001:2008 of original registration date 15th June 1994, issued by the British Standards Institute (CE0086)

In accordance with 2011/65/EU RoHS Directive  
For and on behalf of Viamed Ltd. \_\_\_\_\_

Signature:



Name: **Derek Lamb - Managing Director**

Date: 26 / 06 / 2014