

## **Declaration of Conformity.**

## Class IIa Medical Device(s).

Manufacturer:

Viamed Limited.

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

Device(s):

Microstim DBS Mark 3

**Description of Device(s):** 

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 Device Directive) of 14<sup>th</sup> June 1993 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 the MDD (Annex II) retained by: Viamed Limited.
- EC Quality Assurance Certificate No. CE 01389, first issued by the British Standards Institute (CE0086) on the 23<sup>rd</sup> August 1996.
- Certificate of Registration No. FS28344 to BS EN ISO 9001:2000 of original registration date 15<sup>th</sup> June 1994, issued by the British Standards Institute (CE0086).

Signed:

Date: 01 101107

Position: Managing Director.