

VIAMED



Declaration of Conformity Class IIa Medical Devices

Manufacturer's name:

Viamed Limited

Manufacturer's address: 15 Station Road, Cross Hills

Keighley, West Yorkshire BD20 7DT

United Kingdom

Device:

Microstim DB

Supramaximal Nerve Stimulator

It is hereby declared that the medical devices specified above conform with the essential requirements listed in Annex I of the EC Council Directive 93/42/EEC of 14 June 1993.

This declaration is supported by:

Technical documentation required by the MDD (Annex VII) retained by: Viamed Limited

EC Quality Assurance BS EN ISO 9001 FS28344 Certificate No. FS 28344

EN 46001

Certificate No. CE 01389

Issued by: British Standards Institute

Date: 8 February 1999 Amended 18th October 1999

Notified body number CE 0086

Signed:

Date: December 2, 1999

Position: Managing Director

