

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):	Rectal / Oesophageal Temperature Probe
Description of Device(s):	Rectal / Oesophageal probes for the continuous measurement of temperature of the patient
Council Directive 93/42/EEC	medical device(s) specified above is (are) in compliance with EC (Medical Device Directive) of 14 th June 1993 as transposed into where the product is intended to be marketed.
This declaration is supported b	y:
Technical documentation required by the MDD (Annex II) retained by: Viamed Limited.	
EC Quality Assurance Institute (CE0086) on to	e Certificate No. CE 01389, first issued by the British Standards the 23 rd August 1996.
	tion No. FS28344 to BS EN ISO 9001:2000 of original registration sued by the British Standards Institute (CE0086).
Signed:	Date:/
Position: Managing Director.	