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### The EU Directives covered by this Declaration

93/42/EEC MEDICAL DEVICE DIRECTIVE

### The Products Covered by this Declaration

Autoclave tape

**EC – Representative : Medica & Co Srl (Mr Letterio OLIVA)**  
**Viale della liberta isol 521, N395 Messina 98121**  
**ITALY**

### The Basis on which Conformity is being Declared

The product identified above complies with the requirements of the Medical Device Directory and its annexes above by meeting the following standard: *EN 867*

*The product is defined as Clas 1 product and Declaration of Conformity is issued as per Annex VII of 93/42/EEC MDD.*

The technical documentation required to demonstrate that the product meets the requirements of the Medical Device Directive has been compiled by the signatory below and is available for inspection by the relevant enforcement authorities. The CE mark was first applied in May 2000

The products described above comply with the essential requirements of the directives specified.

**Authority : Hizir Aykin**  
**Date : January 2010**

### **ATTENTION!**

The attention of the specifier, purchaser, installer, or user is drawn to special measures and limitations to use which must be observed when the product is taken into service to maintain compliance with the above directives. Details of these special methods and limitations to use are available on request.

**CE DECLARATION OF CONFORMITY**