



EC Declaration of Conformity (Directive 93/42/EEC)

We,

Manufacturer's Name: Mortara Instrument, Inc.
Manufacturer's Address: 7865 North 86th Street
Milwaukee, WI 53224
USA

declare under our sole responsibility, that to the best of our knowledge, the product(s):

Product Name: ELI 150 and ELI 150Rx

Part Number: ELI150-XXX-XXXXX

(X designates alpha characters denoting system configuration management codes important for post distribution servicing)

Product Options (Configurations): Interpretive or Non-Interpretive, Standard or Expanded Memory, S/W (All Languages), Modem Module, LAN/ Wireless LAN; GSM/GPRS; optional Barcode Scanner.
ELI 150 Rx (Clinical Research version) provides management of multiple sites.

Lot and/or Serial Number(s): SN – "1YYWWXXXXXX"
(Where YY = 2 digit year;
WW = week code; and
XXXXXX = unique number starting at 0000001)

GMDN Code and Term: [11407] – Electrocardiograph, General Purpose

Class (according to the criteria of Annex IX, 93/42/EEC): IIa (Rule 10)

*are in conformity with the dispositions of the directive which are applicable to them.
This declaration is based on the following elements:*

Directive 93/42/EEC: The ISO 13485 Certificate N° 7473 rev.2 for approval of the quality system.
The EC Certificate ANNEX II N° 7472 rev.2 for approval of the quality system.
Technical file (ref. ELI 150, ANNEX VII) to demonstrate the conformity of the product to the essential requirements (ANNEX I).

Notified Body: LNE/G-MED (N° 0459)
1 rue Gaston Boissier
75724 Paris Cedex 15
France

European Union Representative: Mortara Rangoni Europe, Srl
(European Headquarters, Italy)
Via Cimarosa 103-105
40033 Casalecchio di Reno, BO

Identification of Individual Signing/Location: Charles Morreale
Director of Regulatory Affairs and Quality Assurance
Mortara Instrument, Inc.
7865 North 86TH Street
Milwaukee, WI 53224
USA

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