

DECLARATION OF CONFORMITY CE

Calvenzano, December 20th 2010

LUROPAS del dottor Molinari & C. s.r.l., located in Calvenzano (BG), via Vivaldi 4, Italy,

DECLARES UNDER ITS OWN RESPONSIBILITY

that the following products that belong to the category of THERAPEUTIC HOSIERY K1, coded and named:

- 440 gambaletto classe 1 microfibre fine modello a punta aperta
- 443 gambaletto classe 1 microfibre fine modello a punta chiusa
- 441 calza autoreggente classe 1 microfibre fine modello a punta aperta
- 444 calza autoreggente classe 1 microfibre fine modello a punta chiusa
- 442 collant classe 1 microfibre fine modello a punta aperta
- 445 collant classe 1 microfibre fine modello a punta chiusa

are classified according to the 93/42/EEC Directive as **non active medical device class I**.

Besides, we declare that the above mentioned medical devices:

- are not instruments of measure
- are not used for clinics enquiries
- are commercialized in non sterile packaging
- satisfied the essential requirements of the Annex n. 1 and applicable instructions of the 93/42/EEC Directive.

LUROPAS, shall keep and have available to the Authority all the technical documentation specified in the Annex n. VII of the 93/42/EEC Directive for a period of five years from the last date of the making of the product.

President's signature


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