

EC CERTIFICATE

FULL QUALITY ASSURANCE SYSTEM APPROVAL CERTIFICATE

(Annex II of the Directive 93/42/EEC on Medical Devices)

No. 41314523

We hereby declare that an examination of the under mentioned full quality assurance system has been carried out following the requirements of the national legislation LVFS 2001:6 to which the undersigned is subjected, transposing Annex II (with the exemption of section 4) of the Directive 93/42/EEC on medical devices. We certify that the full quality assurance system conforms with the relevant provisions of the aforementioned legislation.

Manufacturer: WR Medical Electronics Co.
123 North, Second Street
Stillwater, MN 55082-5047
USA

Product category: Paraffin Baths
- Therabath


Date of expiry: 29 December 2008

The Certificate is valid for the devices which are stated in the present MDD – Product list

Stockholm
29 December 2003

Intertek Semko AB
Notified Body MDD

The original certificate issued on
29 December 2003


Nils Bromander

Intertek Semko AB is a Notified Body according to the Council Directive 93/42/EEC concerning medical devices.
Identification number 0413.

Intertek ETL SEMKO