

EC Certificate

FULL QUALITY ASSURANCE SYSTEM Directive 93/42/EEC on Medical Devices, Annex II (3)

Certificate Number 41313831

Initial Certification Date March 22, 2001

Certificate Valid from July 7, 2011

Certificate Expiry Date March 22, 2016

The certification is subject to the organization maintaining their system in compliance with the regulations stated in this certificate, allowing regular assessments and following the contracted requirements of the Notified Body.

Intertek Semko AB is a Notified Body according to Directive 93/42/EEC on medical devices, with identification number 0413.

Intertek Semko AB Box 1103, SE-164 22 Kista, Sweden Telephone +46 8 750 00 00 medtechsweden@intertek.com We hereby declare that an examination of the under mentioned full quality assurance system has been carried out following the requirements of the Swedish national legislation 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 directive, and the result entitles the organization to use the CE 0413 marking on those products listed below.

Organization:

TERATECH Corporation

77-79 Terrace Hall Avenue, Burlington, MA 01803, USA

Product Category:

- Portable Medical Ultrasound Imaging Equipment

For further identification of the products covered, see the MDD product list/product schedule.

July 7, 2011 Signed date

Marie Olsson, Certification Manager MDD Intertek Semko AB, Kista, Sweden