EC Certification



PRODUCTION QUALITY ASSURANCE Directive 93/42/EEC for Medical Devices, Annex V

We hereby declare that an examination of the under mentioned production quality assurance system has been carried out following the requirements of the UK national legislation to which the undersigned is subjected, transposing Annex V of the Directive 93/42/EEC on medical devices. We certify that the production quality system conforms with the relevant provisions of the aforementioned legislation, and the result entitles the organization to use the CE 0473 marking on those products listed below.

DTR MEDICAL LIMITED

17 Clarion Court, Clarion Close, Swansea Enterprise Park, Swansea SA6 8RF

Aspirating needles, aspirating cannula and flushing needles

Single-use surgical instruments (comprising forceps, scissors, biopsy punches, rongeurs, retractors, aspirating dissectors, curettes, hooks, probes and points)

Minor procedure packs (including myringotomy packs, ENT packs, suture packs, delivery packs and neuro-burr trays)

Tracheal dilators

Arterial cannula

Gigli saw blades

Suction handles, micro-suction handles, micro-suction devices, Zoellner fine ends, suction regulators, spigots and silicone suction tubing

Silicone bands and silicone slings Single-use vascular booties

Single-use insufflation cannula

Certificate Number:6Initial Certification Date:1Certificate Effective Date:1Certificate Expiry Date:1

601-01 CE 19 October 2005 19 October 2015 18 October 2020

Brian Johnson

AMTAC Certification Services Limited, Milton Keynes, UK This certificate is the property of AMTAC Certification Services Ltd

In the issuance of this certificate, Intertek assumes no liability to any party other than to the Client, and then only in accordance with the agreed upon Certification Agreement. This certificate's validity is subject to the organization maintaining their system in accordance with Intertek's requirements for systems certification. Validity may be confirmed via email at <u>certificate.validation@intertek.com</u> or by scanning the code to the right with a smartphone.

This Certificate is for the exclusive use of AMTAC's client and is provided pursuant to the agreement between AMTAC and its Client. AMTAC's responsibility and liability are limited to the terms and conditions of the agreement. AMTAC assumes no liability to any party, other than to the Client in accordance with the agreement, for any loss, expense or damage occasioned by the use of this Certificate. Only the Client is authorized to permit copying or distribution of this Certificate. Any use of the AMTAC name or one of its marks for the sale or advertisement of the tested material, product or service must first be approved in writing by AMTAC.

The certificate remains the property of Intertek, to whom it must be returned upon request.

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.

AMTAC Certification Services Limited is a Notified Body according to Directive 93/42/EEC for medical devices, with identification number 0473.

