

EC Certificate Full Quality Assurance System: Certificate GB201965310

SGS

The management system of

Residual Barrier Technology Limited

The Die-Pat Centre, Broad March, Daventry, NN11 4HE, UK

has been assessed and certified as meeting the requirements of

Directive 93/42/EEC on medical devices, Annex II (excluding Section 4)

For the following products

P247 Ultimate Disinfectant used in the cleaning and disinfection
medical devices and equipment. P247 Viridis disinfectant used
in the cleaning and disinfection of medical devices and equipment.

Where the above scope includes class II medical device(s), a valid EC Design Examination
Certificate according to Annex II (Section 4) is a mandatory requirement for each device in
addition to this certificate to place that device on the market.

This certificate is valid from 27 March 2020 until 03 July 2022
and remains valid subject to satisfactory surveillance audits.
Issue 2. Certified since 18 October 2017
and first certified by SGS Belgium NV since 28 February 2020.

Certification is based on reports numbered GBPC 240660

Authorised by



SGS Belgium NV, Notified Body 1639

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UND6007 - Certificate CE1639 Annex II-4, EN rev. 02

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