EC Certificate Full Quality Assurance System: GB98/13044

The management system of

Research Instruments Ltd

Bickland Industrial Park, Falmouth, Cornwall, TR11 4TA, 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

The scope of registration appears on page 2 of this certificate.

This certificate is valid from 05 May 2016 until 05 August 2018 and remains valid subject to satisfactory surveillance audits.

Re certification audit due before 07 July 2016 Issue 15. Certified since 29 May 1998

Certification is based on reports numbered GB/PC 08973

This is a multi-site certification. Additional site details are listed on the subsequent page.

Authorised by



SGS United Kingdom Ltd, Notified Body 0120

SGS United Kingdom Ltd 202B Worle Parkway, Weston-super-Mare, BS22 6WA UK t +44 (0)1934 522917 f +44 (0)1934 522137 www.sgs.com

SGS CE 02 0315 M2

Page 1 of 2





This document is issued by the Company subject to its General Conditions of Certification Services accessible at www.sgs.com/terms_and_conditions.htm.
Attention is drawn to the limitations of liability, indemnification and jurisdictional issues established therein. The authenticity of this document may be verified at http://www.sgs.com/en/Our-Company/Certified-Client-Directories/Certified-Client-Directories.aspx. Any unauthorized alteration, forgery or falsification of the content or appearance of this document is unlawful and offenders may be prosecuted to the fullest extent of the law.



Research Instruments Ltd

Directive 93/42/EEC

on medical devices, Annex II (excluding section 4)

Issue 15

Detailed scope

Integra micromanipulation system, Saturn laser systems, electrically heated plates and temperature control units for use in RI Witness systems, sterile single-use glass RI pipettes, sterile single-use plastic pipettes (EZ Range) and sterile single-use migration sedimentation chambers (RI MSC) all for use in handling and treatment of human reproductive samples.

Where the above scope includes class III 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

Additional facilities

Site B, Bickland Industrial Park, Falmouth, Cornwall, TR11 4TA, UK Site C, Bickland Industrial Park, Falmouth, Cornwall, TR11 4TA, UK