

Certificate of Registration of Quality Management System to ISO 13485:2016

Australia - Therapeutic Goods (Medical Devices) Regulations, 2002,
 Schedule 3 Part 1 (excluding Part 1.6) - Full Quality Assurance Procedure.
 Brazil - RDC ANVISA n. 16/2013 RDC ANVISA n. 23/2012 RDC ANVISA n. 67/2009
 Canada - Medical Devices Regulations - Part 1- SOR 98/282
 Japan - MHLW Ministerial Ordinance 169, Article 4 to Article 68, PMD Act (,as applicable)
 United States - 21 CFR 803, 21 CFR 806, 21 CFR 807 - Subparts A to D,
 ∑ 21 CFR 820 - Quality System Regulation

The National Standards Authority of Ireland is an MDSAP Recognized Auditing Organization and certifies that:

Alcor Scientific Inc. 20 Thurber Boulevard Smithfield, RI 02917 USA

D-U-N-S: 884153748

has been assessed and deemed to comply with the requirements of the above standard and regulations in respect of the scope of operations given below:

The Design, Manufacture, and Distribution of Enteral Nutrition Feeding Pumps, Enteral Feeding Sets, and IVD Systems for Haematology.

Approved by: Geraldine Larkin Chief Executive Officer Approved by:
Caroline Dore Geraghty
Director of Medical Devices /
Head of Notified Body

Certificate Number: MP19.4997 / Rev 1 Certification Granted: 2020/02/25

Effective Date: 2020/12/02 Expiry Date: 2023/02/24



