



Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that: Corporation STERIS Canada

(also operating as STERIS Canada ULC)

490 boulevard Armand-Paris

Québec (Québec)

G1C 8A3 Canada

DUNS Number: 20-265-9140

Holds certificate No: MDSAP 687114

Statement of Conformity: The company listed on this certificate has been audited to and found to conform with the following criteria: ISO 13485:2016 and Australia - Therapeutic Goods (Medical Devices) Regulations, 2002, Schedule 3 Part 1 (excluding Part 1.6) - Full Quality Assurance Procedure [if design controls are part of the certification]; Brasil - 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; USA - 21 CFR 820, 21 CFR 803, 21 CFR 806, 21 CFR 807 – Subparts A to D

Design, manufacturing and technical support of washing disinfection and drying systems and associated accessories, and manufacture, distribution and service of sterile processing equipment for healthcare.

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For and on behalf of BSI:

arks Pitanga, Chief Operating Officer Assurance - Americas

Original Registration Date: 2018-10-23 Effective Date: 2018-10-23 Expiry date: 2021-10-22

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...making excellence a habit."



This certificate remains the property of BSI and shall be returned immediately upon request. To be read in conjunction with the scope above or the attached appendix.