

# MANAGEMENT SYSTEMS ACCREDITATION PROGRAM (MSAP)

### Scope of Accreditation

Accredited Legal Entity:	National Standards Authority of Ireland (NSAI)
Contact Name:	Mr. Eoin Banville Medical Device Operations Manager
LOCATION A	
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SCC File Number:	08028
Accreditation Standards: ISO/IEC 17021-1:2015 IAF MD1, MD2, MD4, MD5, MD11 (where applicable)	
Initial Accreditation Date:	2003-12-31
Reaccreditation Date:	2016-01-11
Accreditation Expiry Date:	2019-12-30

### Additional Critical/Key Locations:

Location	Country	Address	City
В	USA	NSAI Inc. 20 Trafalgar Square, Suite 603 Nashua NH 03063 USA	Nashua





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#### I: Medical Devices Quality Management Systems Program

Base program:	Medical Devices Quality Management Systems		
Additional accreditatio n standards	GD207; GD210; GD211; Q90R0 IAF MD9		
Certification standards:	ISO 13485:2003 under CMDCAS and ISO 13485:2016 under CMDCAS and ISO 13485:2003 and ISO 13485:2016		
Locations:	A, B		
Certification Body's main technical areas: (IAF MD 8 & MD 9)	Non-active Medical Devices	<ul> <li>General non-active, non-implantable medical devices</li> <li>Non-active implants</li> <li>Devices for wound care</li> <li>Non-active dental devices and accessories</li> <li>Non-active medical devices other than specified above</li> </ul>	
	Active Medical Devices (Non-Implantable)	<ul> <li>General active medical devices</li> <li>Devices for imaging</li> <li>Monitoring devices</li> <li>Devices for radiation therapy and thermo therapy</li> <li>Active (non-implantable) medical devices other than specified above</li> </ul>	
	Active Implantable Medical Devices	<ul> <li>General active implantable medical devices</li> <li>implantable medical devices other than specified above</li> </ul>	
	In Vitro Diagnostic Medical Devices (IVD)	<ul> <li>Reagents and reagent products, calibrators and control materials for: <ul> <li>Clinical Chemistry</li> <li>Immunochemistry (Immunology)</li> <li>Haematology/Haemostasis/Immunohematology</li> <li>Microbiology</li> <li>Infectious Immunology</li> <li>Histology/Cytology</li> <li>Genetic Testing</li> <li>In Vitro Diagnostic Instruments and software</li> <li>IVD medical devices other than specified above</li> </ul> </li> </ul>	
	Sterilization Method for Medical Devices	<ul> <li>Ethylene oxide gas sterilization (EOG)</li> <li>Moist heat</li> <li>Aseptic processing</li> <li>Radiation sterilization (e.g. gamma, x-ray, electron beam)</li> </ul>	





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		Sterilization method other than specified above
in sr	Devices Incorporating/utilizing pecific ubstances/technologies	<ul> <li>Medical devices incorporating medicinal substances</li> <li>Medical devices utilizing tissues of animal origin</li> <li>Medical devices incorporating derivatives of human blood</li> <li>Medical devices utilizing micromechanics</li> <li>Medical devices utilizing biological active coatings and/or materials or being wholly or mainly absorbed</li> </ul>

This document forms part of the Certificate of Accreditation issued by the Standards Council of Canada (SCC) to National Standards Authority of Ireland (NSAI). The original version is available in the Directory of Accredited Management Systems Certification Bodies on the SCC website at <u>www.scc.ca</u>.

Chantal Guay, ing., P.Eng. Vice President, Accreditation Services Date: 2017-05-16

