

Document #:	FRM-000751
Revision #:	00
Issue Date:	See Agile

Page 1 of 5

### EC DECLARATION OF CONFORMITY

Legal Manufacturer Name and Address:	Teleflex Medical 3015 Carrington Mill Boulevard Morrisville, NC 27560 USA
Authorized Representative Name and Address:	TFX Medical Ltd. IDA Business and Technology Park Dublin Road, Athlone, Co. Westmeath Ireland
Notified Body Name and Address:	<ul> <li>☐ Class I: Not Applicable</li> <li>☑ Class Is, Im, Ila, IIb, III</li> <li>SGS Belgium NV</li> <li>SGS House Noorderlaan 87 2030</li> <li>Antwerp Belgium CE1639</li> </ul>

#### Class I

Teleflex Medical declares that the products herewith comply with the requirements of the Council Directive 93/42/EEC dated 14, June 1993 as amended by 2007/47/EC and is in accordance with Annex II (excluding Section 4) and EN ISO 13485:2016, as implemented by the European Union's Medical Devices Regulations.

#### 🛛 Class Is, Im, Ila, Ilb, III

Teleflex Medical declares that the products herewith comply with the requirements of the Council Directive 93/42/EEC dated 14, June 1993 as amended by 2007/47/EC and is in accordance with Annex II (excluding Section 4) and EN ISO 13485:2016, as implemented by the European Union's Medical Devices Regulations as verified by the Notified Body listed above:

Teleflex Medical confirms that no other application has been lodged with another Notified Body for the same devices related Quality Management System.

Teleflex Medical agrees to develop, implement, and maintain a formally-recognized Quality Management System to ensure continued adequacy and efficacy.

Teleflex Medical agrees to develop, implement and maintain a documented post-production experience monitoring process, including the notification of reportable events under the European Medical Device Vigilance System Guidelines.

Teleflex Medical confirms that no medicinal products/drugs are incorporated in any devices covered by the Device Schedule.

Teleflex Medical agrees to inform the appointed Notified Body of any planned or unplanned substantial change to the Quality Management System.

Teleflex Medical agrees to inform the appointed Notified Body of any planned or unplanned significant change to the Device Schedule, if applicable.

Product Name:	Non-Sterile Cannula and Supply Tubing
Classification:	Class Ila Rule 5 (Nasal Cannula)
	Class IIa Rule 2 (Oxygen Supply Tubing)
EC Certificates No.:	Canadian – ISO 13485:2016-US18/81827522
	European – ISO 13485:2016, EN ISO 13485:2016 –
	US19/819943645.00, Directive 93/42/EEC –US19/819943647.00

For Use by Affiliates of Teleflex



Document #:	FRM-000751
Revision #:	00
Issue Date:	See Agile

Page 2 of 5

Conformity Assessment Routes:

Annex II (excluding Section 4) of the MDD (93/42/EEC), full Quality Assurance System

Product Code	Product Description	CE Distribution Date	GMDN Code
Nasal Cannulas			
1103	Cannula, Over-the-Ear, with Tubing	September 6, 2005	35201
	Cannula, Over-the-Ear, with Flared Prongs,		35201
1104	Tubing	September 6, 2005	
	Cannula, Over-the-Ear, with STARLUMEN		35201
41110	Tubing, Standard Connectors	September 6, 2005	
	Cannula, Softech, Adult with STARLUMEN		35201
41820	Tubing, Standard Connectors	September 6, 2005	
	Cannula, Softech, Pediatric with STARLUMEN		35201
41826	Tubing and Universal Connectors	September 6, 2005	
	Cannula, Softech, Infant with STARLUMEN		35201
41828	Tubing	September 6, 2005	
1811	Cannula, 1103 w/25' Star-Lumen Tubing	October 22, 2014	35201
1813	Cannula, 1103 w/50' Star-Lumen Tubing	October 22, 2014	35201
1070	Softech Plus Adult Cannula,7' Green Star-		35201
1870	Lumen Tubing	January 24, 2013	
1871	Softech Plus Pediatric Cannula,7' Green Star-		35201
10/1	Lumen Tubing	January 24, 2013	
1070	Softech Plus Infant Cannula,7' Green Star-		35201
1872	Lumen Tubing	January 24, 2013	
1873	Softech Plus Neonatal Cannula,7' Green Star-		35201
10/3	Lumen Tubing	January 24, 2013	
1874	Softech Plus Adult, Cannula,7' Green Star-		35201
10/4	Lumen Tubing w/ Universal Connector	January 24, 2013	
1876	Softech Plus Pediatric, Cannula,7' Green Star-		35201
1070	Lumen Tubing w/ Universal Connector	January 24, 2013	
1852	Softech Plus Adult Cannula,14' Green Star-		35201
1052	Lumen Tubing w/ Universal Connector	January 24, 2013	
1853	Softech Plus Pediatric Cannula,14' Green Star-		35201
1000	Lumen Tubing w/ Universal Connector	January 24, 2013	
1877	Softech Plus Adult Cannula,14' Green Star-		35201
1077	Lumen Tubing	January 24, 2013	
1878	Softech Plus Pediatric Cannula, 14' Green Star-		35201
1070	Lumen Tubing	January 24, 2013	
Oxygen Supply			
1115	Tubing, STARLUMEN, 2.1 m	September 6, 2005	12875
1900	Tubing, Oxygen Supply, 50'	October 22, 2014	12875
1986	Tubing, Tinted Star-Lumen, 30'	October 22, 2014	12875
1987	Tubing, Tinted Star-Lumen, 40'	October 22, 2014	12875
1988	Tubing, Tinted Star-Lumen, 50'	October 22, 2014	12875
1989	Tubing, Oxygen, 25'	October 22, 2014	12875
1991	Tubing, Oxygen Supply, 2.1 m	March 24, 2014	12875
41113	Tubing, STARLUMEN, 2.1 m	September 6, 2005	12875
41118	Tubing, STARLUMEN, 4.2 m	September 6, 2005	12875
41119	Tubing, STARLUMEN, 7.6 m	September 6, 2005	12875
41120	Tubing, STARLUMEN, 15.2 m	September 6, 2005	12875
41925	Oxygen Tubing w/ Universal Connector 2.1 m	September 6, 2005	12875
1679	Oxygen Tubing Water Trap	September 6, 2005	41679

\* Indicates the item is within the scope of and in compliance with European Directive 2011/65/EU, The Restriction of the Use of Certain Hazardous Substances in Electrical and Electronic Equipment

For Use by Affiliates of Teleflex
This document contains proprietary information. It may not be reproduced without prior written approval.
>>THE USER OF THIS DOCUMENT IS RESPONSIBLE FOR CHECKING THE CURRENT ISSUE DATE BEFORE USING THIS DOCUMENT<<
Document #: FRM-000751 Revision#: 00 Issue Date: See Agile Parent Document: WI-003907



Document #:	FRM-000751
Revision #:	00
Issue Date:	See Agile

## Page 3 of 5

Name and Title of Approver:	Nicole Schaffer, Regulatory Affairs Manager, Respiratory
Signature of Approver:	Micole Athoffer
Date Approved:	07 June 2021
Site Where Approved:	Teleflex Medical, Inc. 3015 Carrington Mill Blvd. Morrisville, NC 27560 USA
Canadian Classification	

The following devices meet Canadian requirements as listed:

Product Code	Product Description	Canadian License #	Issue Date	Class of Product 2
1103	Cannula, Over-the-Ear, with Tubing	7582	07/09/99	
1104	Cannula, Over-the-Ear, with Flared Prongs, Tubing	7582	07/09/99	2
41110	Cannula, Over-the-Ear, with STARLUMEN Tubing, Standard		10/15/12	2
1115	Tubing, STARLUMEN, 2.1 m	7588	07/09/99	2
1991	Tubing, Oxygen Supply, 2.1 m	7588	3/21/2014	2
41113	Tubing, STARLUMEN, 2.1 m	7588	8/24/09	2
41118	Tubing, STARLUMEN, 4.2 m	7588	10/15/12	2
41119	Tubing, STARLUMEN, 7.6 m	7588	10/15/12	2
41120	Tubing, STARLUMEN, 15.2 m	7588	10/15/12	2
41925	Oxygen Tubing w/ Universal Connector 2.1 m	7588	10/15/12	2
1679			11/1/2002	2
1811	Cannula, 1103 w/25' Star-Lumen Tubing	7582	10/22/2014	2
1813	Cannula, 1103 w/50' Star-Lumen Tubing		10/22/2014	2
1900	Tubing, Oxygen Supply, 50'	7588	10/22/2014	2
1985	Tubing, Tinted Star-Lumen, 25'	7588	10/22/2014	2
1986	Tubing, Tinted Star-Lumen, 30'	7588	10/22/2014	2
1987	Tubing, Tinted Star-Lumen, 40'	7588	10/22/2014	2
1988	Tubing, Tinted Star-Lumen, 50'	7588	10/22/2014	2
1989	Tubing, Oxygen, 25'	7588	10/22/2014	2
41820	Cannula, Softech, Adult with STARLUMEN Tubing, Standard		8/12/2016	2
Cannula, Softech, Pediatric with STARLUMEN Tubing and Universal Connectors		94237	8/12/2016	2
41828	Cannula, Softech, Infant with STARLUMEN Tubing	94237	8/12/2016	2

# For Use by Affiliates of Teleflex This document contains proprietary information. It may not be reproduced without prior written approval. >>THE USER OF THIS DOCUMENT IS RESPONSIBLE FOR CHECKING THE CURRENT ISSUE DATE BEFORE USING THIS DOCUMENT<< Document #: FRM-000751 Revision#: 00 Issue Date: See Agile Parent Document: WI-003907



Document #:	FRM-000751
Revision #:	00
Issue Date:	See Agile

Page 4 of 5

Product Description	<ul> <li>while providing part of the patient's ins oxygen is provided from a nearby sour inhaled from the room environment.</li> <li>Oxygen Supply Tubing</li> <li>Oxygen supply tubing is an accessory oxygen supply and a device, such as a requires an oxygen supply. This prod as a component included with other pr soft, flexible PVC in a clear or green til end connectors or the slightly larger "u</li> </ul>	Nasal cannulas are designed to administer low-flow oxygen to the patient while providing part of the patient's inspiratory gas flow needs. Supplemental oxygen is provided from a nearby source or regulator and the remaining air is inhaled from the room environment.		
Indications for Use	the nasal passage. Oxygen Supply Tubing Oxygen supply tubing is used to delive	<ul> <li>Nasal cannulas are used to deliver a low flow of oxygen to the patient through the nasal passage.</li> <li>Oxygen Supply Tubing</li> <li>Oxygen supply tubing is used to deliver oxygen (or other medical gases) from a gas source to a patient, typically in conjunction with a mask, nasal cannula,</li> </ul>		
Intended Use	the nasal passage. Oxygen Supply Tubing Oxygen supply tubing is used to delive	Nasal cannulas are used to deliver a low flow of oxygen to the patient through the nasal passage. <b>Oxygen Supply Tubing</b> Oxygen supply tubing is used to deliver oxygen (or other medical gases) from a gas source to a patient, typically in conjunction with a mask, nasal cannula,		
Contraindications	These products do not list any contrair	These products do not list any contraindications on the labeling.		
Manufacturing Site(s)	Manufacturing Site Name           Manufacturing Facility for _Cat.No.           1103, 1104, 1104L, 1115, 1811,           1813, 1900, 1985, 1986, 1987,           1988, 1989, 1991, 41110, 41113,           41118, 41119, 41120, 41925, 1679:           Manufacturing Facility for _Cat.No.           1870, 1871, 1872, 1873, 1874,           1876, 1852, 1853, 1877, 1878           41820, 41826, 41828. These           products are Purchased Finished           Goods (PFG) from Soundway. The           Soundway DoC, CE Certificate and           contract may be found in Section 2.	Manufacturing Site AddressTeleflex MedicalAve. Industrias No 5954Parque Industrial FinsaNuevo Laredo, Tamaulipas, 88275MexicoNingbo Shengyurui MedicalAppliances Co., Ltd (akaSoundway)No. 138, Binhaisi Road,Hangzhou Bay New Zone,Cixi City, Zhejiang ProvinceChina		
Sterilizer	N/A: The product is sold non-sterile	<ul> <li>N/A: The product is sold non-sterile.</li> <li>☐ The product is sold sterile.</li> </ul>		

For Use by Affiliates of Teleflex This document contains proprietary information. It may not be reproduced without prior written approval. >>THE USER OF THIS DOCUMENT IS RESPONSIBLE FOR CHECKING THE CURRENT ISSUE DATE BEFORE USING THIS DOCUMENT<< Document #: FRM-000751 Revision#: 00 Issue Date: See Agile Parent Document: WI-003907



CMDR

EN 1041+A1:2013

21CFR Part 820

21 CFR Part 801

EU MEDDEV

2.7.1/rev 4

Document #:	FRM-000751
Revision #:	00
Issue Date:	See Agile

2013

2011

2011

2016

Page 5 of 5

Quality System Regulation

Canada Medical Device Regulations

Labeling

90/385/EEC

Sterilization Site Name         Sterilization           Standards			Sterilization Site Add	Iress
Standard Number	Standard Name			Issue Date
93/42/EEC MDD	Medical Device	Directive as amended by 2007/47/EC		2007
ISO 13485	Medical Devices	- Quality management systems - Req	uirements for	2016
	regulatory purposes			
EN ISO 13485	Medical Devices – Quality management systems – Requirements for			2016
	regulatory purpos	ses		
ISO 14971	Medical devices – Application of risk management to medical devices			2019
EN ISO 14971	Medical devices – Application of risk management to medical devices			2019
ISO 10993-1	Biological evaluation of medical devices Part 1: Evaluation and testing within a risk management process			2020
ISO 10993-5	Biological evaluation of medical devices Part 5: Tests for in vitro cytotoxicity			2009
ISO 10993-10	Biological evalua sensitization	ation of medical devices Part 10: Tests f	or irritation and skin	2013
ISO 15223-1		<ul> <li>Symbols to be used with medical de to be supplied — Part 1: General requir</li> </ul>		2016

Information supplied by the manufacturer of medical devices

Guidelines on Medical Devices Clinical Evaluation: A Guide for

Manufacturers and Notified Bodies under directives 93/42/EEC and

For Use by Affiliates of Teleflex