



DECLARATION OF CONFORMITY

(in accordance with ISO/IEC 17050-1)

SAP DIR No.: 80016303

Version: F

We declare, under our sole responsibility, that the product listed below conforms to the provisions of:

- the European Council Directive 93/42/EEC of 14 June 1993 concerning medical devices
- the Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS)

Manufacturer's Name and Business Address: Welch Allyn, Inc.
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Skaneateles Falls, NY 13153, USA



Regulatory Affairs Representative
Welch Allyn Limited
Navan Business Park
Dublin Road
Navan, County Meath
Republic of Ireland

Product Name: Fiber Optic Laryngoscopes



901038, LARYNGOSCOPE
901087, INSTRUMENT HANDLE



Blades:
68060, 68061, 68062, 68063, 68064, 68065 (Miller Fiber-Optic Blades)
69061, 69062, 69063, 69064 (MacIntosh Fiber-Optic Blades)
69211, 69212, 69213, 69214 (English MacIntosh Fiber-Optic Blades)

Handles:
60813, 60813-LED, 60814, 60814-LED, 60815, 60815-LED, 60713, 60835 (Fiber Optic Battery Handles)

Kits:
65101, 65102, 65103, 65104, 65121, 65122, 65123, 65124, 65125, 65126, 68696, 68696-LED, 69696, 69696-LED 69697 & 69697-LED

Medical Device Conformity Assessment Route Annex: VII

Medical Device Classification: I

Medical Device Classification Rules: 5

GMDN Code and Term: 15076 – Laryngoscope, intubation

UMDNS Code and Term: 15076 - Laryngoscopes designed with a non-flexible (i.e., rigid) structure that can only follow a straight path through the airway. They are constructed of metal and contain



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straight or curved blades for manipulation of the tongue and pharynx during the procedures. Rigid endoscopes are frequently used for insertion of endotracheal tubes.

Standards Applied:	EN 50581	Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances
	EN / IEC 60601-1	Medical Electrical Equipment, Part 1: General Requirements for Safety
	ISO 7376-3	Laryngoscope fittings - Part 3: Fibre-illuminated re-usable rigid laryngoscopes
	EN / ISO 7376	Anaesthetic and respiratory equipment -- Laryngoscopes for tracheal intubation NOTE: Compliance report applicable to blades used with -LED handles.
	EN / ISO 10993-1	Biological evaluation of medical devices -- Part 1: Evaluation and testing within a risk management process

Authorised Signatory:

Fiona Butler, Manager Regulatory Affairs
{EU Authorised Representative}

Date

Place of Issue