

WelchAllyn® DECLARATION OF CONFORMITY

(in accordance with ISO/IEC 17050-1)

SAP DIR No.: 80016309

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: Battery Handles and Chargers



901087, INSTRUMENT HANDLE
901001, ACCESSORY, EYE, EAR, NOSE & THROAT



70000, 70500, 70500F, 70700, 70710, 70715, 70720, 70720F, 70762, 70764, 71000, 71001-B, 71001-C, 71010, 71020, 71021-C, 71062, 71064, 71066, 71670, 71712, 71714, 71716, 71732, 71734, 71736, 71900, 71902, 71904, 71906, 71907, 71910, 71911, 71930, 72800, 72801, 72830, 70720, 71000-A, 71000-B, 71000-C, 71020-A, 71020-B, 71020-C, 71062-C, 71064-C
71140, 71142, 71144, 71146, 71942, 71943, 71960-POD

Medical Device Conformity Assessment Route Annex: VII

Medical Device Classification: I

Medical Device Classification Rules: 1

GMDN Code and Term: 34158 – Battery, Secondary
17115 – Battery Charger

UMDNS Code and Term: 18557 - Any source of electric power, such as a power line or a battery. Batteries, direct-current electronic regulated power supplies, line power supply systems (including isolated and uninterruptible systems), and transformers are the main devices used in the field of medicine to deliver electric power to electric

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and electronic medical devices in hospitals and homes; batteries are also used as power supplies for implantable medical devices.

Standards Applied:	EN 50581	Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances
	EN 60601-1	Medical Electrical equipment – Part 1: General requirements for Safety.
	EN 60601-1-2	Medical electrical equipment – Part 1-2: General requirements for safety – Collateral Standard Safety requirements for medical electrical systems.

Authorised Signatory:



Fiona Butler, Manager Regulatory Affairs
{EU Authorised Representative}



Date

Navan

Place of Issue