

Welch Allyn[®] DECLARATION OF CONFORMITY

(in accordance with ISO/IEC 17050-1)

SAP DIR No.: 80016526 Version: G

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
U.S.A.



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

Product Name: Otoscope



901079 OTOSCOPE, STANDARD
901080 OTOSCOPE, POCKET



20000, 20000-L, 20001, 20097, 20098, 20200, 20201, 20203, 20250, 20251, 20270, 20282, 20283, 20284, 20285, 21110, 21111, 21140, 21141, 21307, 21308, 21504, 21700, 21701, 21770, 21782, 21783, 21784, 21785, 22009, 22091, 22100, 22800, 22811, 22820, 22821, 22822, 22831, 22840, 22841, 22860, 22861, 22870-BLK, 22870-BLU, 22870-PUR, 22870-WHT, 22880-BLK, 22880-BLU, 22880-PUR, 22880-WHT, 23510, 23510-L, 23520, 23520-L, 23540, 23557, 23804, 23810, 23810-L, 23811, 23811-L, 23814, 23820, 23820-L, 23821, 23821-L, 23824, 24222, 24224, 24330, 24610, 24612, 25020, 25020-L, 25021, 25035, 25070, 25082, 25270, 25282, 25283, 25284, 25285, 25582, 25583, 25584, 25585, 26538, 52133, 52134, 52135, 52700, 20201F, 21111F, 21601F, 21701F, 21783-C, 22820-CLX, 22821-LILLY, 22840S, 25272-MS, 25272-MSL, 25274-MS, 25282-B, 25282-BC, 25282-C, 52423-U & 97206-MVPS.

Annex: VII

Classification: I

Classification Rules: 5

GMDN Code and Term: 12849 – Otoscope, direct

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UMDNS Code and Term 12849 – Oscopes

Standards Applied:	EN 50581	Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances
	EN 1041	Information supplied by the manufacturer of medical devices
	EN/IEC 60601-1	Medical Electrical Equipment – General Guidelines for Safety
	EN/IEC 60601-1-2	Medical electrical equipment -- Part 1-2: General requirements for safety - Collateral standard: Electromagnetic compatibility - Requirements and tests
	EN/IEC 60601-1-6	Medical electrical equipment -- Part 1-6: General requirements for safety - Collateral standard: Usability
	EN/IEC 62366	Medical Devices – Application of Usability Engineering to Medical Devices
	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}

2014-11-04

Date

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Place of Issue