Welch Allyn[•]

SAP DIR: 80019192 Version: F Page 1 of 2

DECLARATION OF CONFORMITY

(in accordance with ISO/IEC 17050-1)

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. 4341 State Street Road Skaneateles Falls, NY 13153, USA
ECREP	Regulatory Affairs Representative Welch Allyn Limited Navan Business Park Dublin Road Navan, County Meath Republic of Ireland
Product Name ^{1,3} :	PanOptic Ophthalmoscope
REF 1,3	901022 OPHTHALMOSCOPE, WIDEVIEW
# 1,3	11800DEM, 11801, 11810, 11810-CE, 11810DEM, 11811, 11812-V, 11812- VSM, 11816-V, 11816-VC, 11816-VSM, 11820, 11820-CE, 11820-CEL, 11820DEM, 11820-L, 11821, 11821-L, 11822-V, 11822-VSM, 11824-V, 11824- VC, 11824-VSM, 11826-V, 11826-VC, 11826-VSM, 11870, 11875
Radio equipment ² :	Not Applicable, No Radio
Object of the declaration ² :	Not Applicable, No Radio
Accessories and components ² :	Not Applicable, No Radio
Medical Device Conformity Assessment Route Annex ¹ :	VII
Medical Device Classification ¹ :	Ι
Medical Device Classification Rules ¹ :	12

applicable to the medical devices directive, 93/42/EEC

² applicable to the radio equipment directive, 2014/53/EU

³ applicable to the RoHS directive, 2011/65/EU

Template DIR 80019151 Ver. F

Welch Allyn[•]

SAP DIR: 80019192 Version: F Page 2 of 2

GMDN Code and 46788 Indirect more Term¹:

46788 Indirect monocular ophthalmoscope, battery-powered

12818 Ophthalmoscope, indirect

UMDNS Code and Term¹:

Standards Applied (Standards are applicable to the medical device directive, unless otherwise indicated):

Number	Title
EN 50581 ³	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 – 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 15004-1	Ophthalmic Instruments – Fundamental Requirements and Test Methods – Part 1: General Requirements Applicable to All Ophthalmic Instruments
EN/ISO 15004-2	Ophthalmic Instruments – Fundamental Requirements and Test Methods – Part 2: Light Hazard Protection
EN/ISO 10943	Ophthalmic Instruments - Indirect Ophthalmoscopes
EN ISO 10993-1	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process

Authorised Signatory:

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Fiona Butler, Manager Regulatory Affairs {EU Authorised Representative}

2019-03-19

Date

Navan Place of Issue

applicable to the medical devices directive, 93/42/EEC

² applicable to the radio equipment directive, 2014/53/EU

³ applicable to the RoHS directive, 2011/65/EU

Template DIR 80019151 Ver. F