

## 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
<b>EC REP</b>	Regulatory Affairs Representative Welch Allyn Limited Navan Business Park Dublin Road Navan, County Meath Republic of Ireland
Product Name <sup>1,3</sup> :	Ophthalmoscopes
<b>REF</b> <sup>1,3</sup>	901081, OPHTHALMOSCOPE, STANDARD 901082, OPHTHALMOSCOPE, POCKET
<b>#</b> <sup>1,3</sup>	11470F, 11710, 11710F, 11720, 11720F, 11720-L, 11720-LF, 11720R, 11721, 11721F, 11722, 11723, 11724, 11730, 11730F, 11730-R, 11731, 11732, 11735, 11735F, 11736, 11736F, 11750, 11750-VBI, 11770, 11770-BI, 11772-BI, 11772-VC, 11772-VCI, 11772-VCL, 11772-VSM, 11782-VSM, 11790, 11792-SC, 11796-SC, 12800, 12811, 12812, 12820, 12821, 12831, 12850, 12851, 12860, 12861, 12870-BLK, 12870-BLU, 12870-PUR, 12870-WHT, 12880-BLK, 12880-BLU, 12880-PUR, 12880-WHT, 13000, 13010, 19090, 19091, 19092, 19093, 19190
Radio equipment <sup>2</sup> :	Not applicable, no radio
Object of the declaration <sup>2</sup> :	Not applicable, no radio
Accessories and components <sup>2</sup> :	Not applicable, no radio
Medical Device Conformity Assessment Route Annex <sup>1</sup> :	VII
Medical Device Classification <sup>1</sup> :	I

<sup>1</sup> applicable to the medical devices directive, 93/42/EEC

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

<sup>3</sup> applicable to the RoHS directive, 2011/65/EU

Medical Device Classification Rules <sup>1</sup> :	12	
GMDN Code and Term <sup>1</sup> :	46786 Direct ophthalmoscope, battery-powered	
UMDNS Code and Term <sup>1</sup> :	12817 – Ophthalmoscope, direct	
Standards Applied (Standards are applicable to the medical device directive, unless otherwise indicated):	Number	Title
	EN 50581 <sup>3</sup>	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 10942	Ophthalmic Instruments - Direct Ophthalmoscopes
	EN/IEC 62471	Photobiological Safety of Lamps and Lamp Systems
EN ISO 10993-1	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process	

Authorised Signatory:

*Fiona Butler*  
Fiona Butler, Manager Regulatory Affairs  
{EU Authorised Representative}

2018-04-30  
Date

Navan  
Place of Issue

<sup>1</sup> applicable to the medical devices directive, 93/42/EEC  
<sup>2</sup> applicable to the radio equipment directive, 2014/53/EU  
<sup>3</sup> applicable to the RoHS directive, 2011/65/EU