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## 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

Welch Allyn, Inc.

Name and

4341 State Street Road

Business Address:

Skaneateles Falls, NY 13153-0220

U.S.A

EC REP

Regulatory Affairs Representative

Welch Allyn Limited Navan Business Park

**Dublin Road** 

Navan, County Meath Republic of Ireland

Product Name<sup>1,3</sup>:

KleenSpec® Vaginal Specula

REF 1,3

901071 VAGINAL SPECULUM

# 1,3

58600, 59000, 59001, 59004, 59005, 59006, 58000S, 58001S, 58004S, 59000-B,

59001-B, 59004-B, 590XS, 590XS-B

Radio equipment<sup>2</sup>:

Not applicable

Object of the

Not applicable

declaration<sup>2</sup>:

Accessories and

Not applicable

components<sup>2</sup>:

Medical Device

VII

Conformity

Assessment Route

Annex1:

Medical Device

I

Classification<sup>1</sup>:

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Medical Device

5 and 12

Classification

Rules1:

GMDN Code and

37468 Vaginal Speculum, single use

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

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

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

## Welch Allyn<sup>o</sup>

Term<sup>1</sup>:

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UMDNS Code and Term <sup>1</sup> :	13666 Speculum, Vaginal	
Standards Applied (Standards are applicable to the medical device directive, unless otherwise indicated):	EN 50581	Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances
	IEC 60601-1	Medical Electrical Equipment – Part 1: General Requirements for Basic Safety and Essential Performance
	IEC 60601-1-2	Medical Electrical Equipment – Part 1-2: General Requirements for Safety – Collateral Standard:

IEC 62366 Medical devices - Application of usability engineering to medical devices

Electromagnetic Compatibility

IEC 60601-1-6 Medical Electrical Equipment - Part 1-6: General Requirements for Safety - Collateral Standard: Usability

EN ISO 10993-1 Biological Evaluation of Medical Devices - Part 1:

**Evaluation and Testing** 

Authorised Signatory:

Fiona Butler, Manager Regulatory Affairs

for Butter

{EU Authorised Representative}

Navan Place of Issue

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

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