

Welch Allyn, Inc. is a subsidiary of Hill-Rom Holdings, Inc.

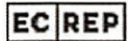
We declare, under our sole responsibility, that the product listed below conforms to the provisions of:

- European Council Directive 93/42/EEC of 14 June 1993 concerning medical devices, and
- Directive 2014/53/EU of the European Parliament and of the Council of 16 April 2014 on the harmonization of the laws of the Member States relating to the making available on the market of radio equipment, and
- 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 as amended by Commission Delegated Directive (EU) 2015/863 of 31 March 2015 (RoHS3).

Document Number 80016302      Version K

Product Name      ProBP 3400 Series

 Manufacturer's Name and Business Address      Welch Allyn, Inc.      SRN: US-MF-000013394  
 4341 State Street Road  
 Skaneateles Falls, NY 13153  
 USA

 EC Certificates Declaration of Conformity Validity      EC Certificate 314505 MR2  
 Expiry Date: 2024-05-26

 Welch Allyn Limited      SRN: IE-AR-000000768  
 Navan Business Park, Dublin Road  
 Navan Co. Meath  
 C15 AW22 Ireland


34XFHT-B	34XXHT-B	34XXHT-B	34XFST-B	34XXST-B
34XFWT-B	34XXWT-B	34XXHT-2	34XFWT-2	34XXST-2
34XFHT-2	34BXWT-2	34XXWT-2	34XFST-2	34XFHT-4
34XXHT-4	34XFST-4	34XFWT-4	34XXST-4	34XXWT-4

901055, DIGITAL BLOOD PRESSURE DEVICE

Radio equipment      Laird Tech, Model BTM411

Object of the declaration



Medical Device Conformity Assessment Route Annex      Annex II



## DECLARATION OF CONFORMITY

(in accordance with ISO/IEC 17050-1)

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Medical Device Classification	Ila
Medical Device Classification Rule	10
Standards	See Appendix A
GMDN Code and Term	45617 – Automatic-inflation electronic sphygmomanometer, portable, arm/wrist
UMDNS Code and Term	16173 - Sphygmomanometers, Electronic, Automatic
Notified Body	DQS Medizinprodukte GmbH, August-Schanz-Str.21, 60433 Frankfurt am Main Notified Body Number: 0297



**DECLARATION OF CONFORMITY**

(in accordance with ISO/IEC 17050-1)

Welch Allyn, Inc. is a subsidiary of Hill-Rom Holdings, Inc.

**Authorised Signatory**

A handwritten signature in black ink, appearing to read 'Joshua Kim', written over a horizontal line.

Joshua Kim  
Senior Manager  
Global Regulatory Affairs

2022.04.20  
Date

Skaneateles Falls NY, USA  
Place of Issue

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**Appendix A: Standards and Common Specifications**

Standards Applied	Number	Version/Date of Issue	Title
Directive 93/42/EEC	EN 60601-1	2013	Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
	EN 60601-1-2	2015	Medical Electrical Equipment, Part 2: Collateral Standard: Electromagnetic Compatibility: Requirements and Test
	EN ISO 81060-1	2012	Non-Invasive Sphygmomanometers - Part 1: Requirements and Test Methods for Non-Automated Measurement Type
	EN ISO 81060-2	2014	Non-Invasive Sphygmomanometers - Part 2: Clinical Validation of Automated Measurement Type
	EN 80601-2-30	2018	Medical Electrical Equipment - Part 2-30: Particular Requirements for The Basic Safety and Essential Performance of Automated Non-Invasive Sphygmomanometers
	EN 62366-1	2015	Medical devices -- Part 1: Application of usability engineering to medical devices
	EN ISO 10993-1	2010	Biological evaluation of medical devices - Part 1: Evaluation and testing
	EN 62304	2006 check	Medical Device Software - Software Life Cycle Processes
	EN 62353	2008	Medical Electrical Equipment - Recurrent Test and Test After Repair of Medical Electrical Equipment
Directive 2014/53/EU	EN 62311	2008 Check	Assessment of electronic and electrical equipment related to human exposure restrictions for electromagnetic fields (0 Hz – 300 GHz)
	EN 301 489-1	2019-03 check	ElectroMagnetic Compatibility (EMC) standard for radio equipment and services; Part 1: Common technical requirements; Harmonised Standard covering the essential requirements of article 3.1(b) of Directive 2014/53/EU and the essential requirements of article 6 of Directive 2014/30/EU

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Standards Applied	Number	Version/Date of Issue	Title
	EN 301 489-17	2017-03 check	ElectroMagnetic Compatibility (EMC) standard for radio equipment and services; Part 17: Specific conditions for Broadband Data Transmission Systems; Harmonised Standard covering the essential requirements of article 3.1(b) of Directive 2014/53/EU
	EN 300 328	2016-11 check	Electromagnetic compatibility and Radio spectrum Matters (ERM); Wideband transmission systems; Data transmission equipment operating in the 2,4 GHz ISM band and using wide band modulation techniques; Harmonized EN covering the essential requirements of article 3.2 of Directive 2014/53/EU
	EN 301 893	2017-03 Check	5 GHz RLAN; Harmonised Standard covering the essential requirements of article 3.2 of Directive 2014/53/EU
Directive 2011/65/EU + (EU) 2015/863	EN IEC 63000	2018	Technical documentation for the assessment of electrical and electronic products with respect to the restriction of hazardous substances