

Declaration of Conformity

Number: 60031771_D

Manufacturer: Welch Allyn, Inc.
Address: 4341 State Street Road,
 Skaneateles Falls, NY 13153-0220
 USA

Manufacturing site Welch Allyn de Mexico S. de R.L. de C.V.
 Calle Emilio Flores #271-A
 Colonia Cañon de Padre
 Tijuana, Baja California C.P. 22203
 Mexico

Representative: European Regulatory Representative,
 Welch Allyn Ltd.
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 Dublin Road,
 Navan,
 County Meath,
 Republic of Ireland
 Tel. +353 46 9067700
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We Welch Allyn declare under our sole responsibility that the Product

Name:	GS 600	
Model Number:	44600	GS600 General Exam light with Mobile Stand DOM
	44602	GS600 General Exam light with Mobile Stand EUR
	44604	GS600 General Exam light with Mobile Stand UK
	44606	GS600 General Exam light with Mobile Stand AU
	44607	GS600 General Exam light with Mobile Stand ZA
	44608	GS600 General Exam light with Mobile Stand CN
	44610	GS600 General Exam light with Table/Wall Mount DOM
	44612	GS600 General Exam light with Table/Wall Mount EUR
	44614	GS600 General Exam light with Table/Wall Mount UK
	44616	GS600 General Exam light with Table/Wall Mount AU
	44617	GS600 General Exam light with Table/Wall Mount ZA
	44618	GS600 General Exam light with Table/Wall Mount CN
Accessory	52630	Disposable Sheets for GS600/900 (100/box, 5 boxes/case)
	48955	Table/Wall Mount for GS Exam Light IV/GS 300/GS 600

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To which this declaration relates is in conformity with the following standards

EN 14971: 2007	Medical Devices- Application of Risk Assessment to medical devices
EN 60601-1: 1990 (incl. amendments)	Medical Electrical Equipment- part 1: General requirements for basic safety and essential requirements
EN 60601-1-1: 2000	Medical Electrical Equipment- part 1-1-General requirements for safety- Collateral Standard: Safety requirements for Medical Electrical Equipment
EN 60601-1-2: 2004	Medical Electrical Equipment- part 1-2- General requirements for safety- Collateral Standard: Electromagnetic Compability- Requirements and Test
EN 60601-1-4: 1997	Medical Electrical Equipment- part 1-4- General requirements for safety- Collateral Standard: General requirements for programmable electrical medical systems
EN 60601-1-6: 2004	Medical Electrical Equipment- part 1-6- General requirements for safety- Collateral Standard: Usability
EN 1041: 2008	Information supplied by the manufacturer with Medical devices
EN 980: 2003	Graphical symbols for use in the labeling of medical devices

The quality system that controls the manufacture of this product complies with the following standards, EN ISO 13485:2003.

Following the provisions of European Council Directive 93/42/EEC of 14 June 1993 concerning medical devices Annex VII. Device is Class I under rule(s) 12 of Annex IX of this directive and following the provisions of European Council Directive 2006/95/EC of 12 December 2006 concerning electrical equipment designed or used within certain voltage limits.

European Regulatory Representative

Signed:


Paul Reynolds

Place of issue, Welch Allyn Ltd. Ireland.

Date:

2010-04-12