# SIEMENS

## **Declaration of Conformity**

**LEGAL MANUFACTURER:** 

Siemens Healthcare Diagnostics Inc.

**511 Benedict Avenue** 

Tarrytown, New York 10591-5097

**USA** 

PLACE OF MANUFACTURE:

Siemens Healthcare Diagnostics Products GmbH

Emil-von-Behring-Str. 76

35041 Marburg Germany

**EU AUTHORIZED REPRESENTATIVE** 

Siemens Healthcare Diagnostics Ltd.

Sir William Siemens SQ.

Frimley, Camberley, UK GU16 8QD

PRODUCT:

Xprecia Systems™ PT Controls

PRODUCT CATEGORY:

See Attachment 1

**CLASSIFICATION:** 

**Self-Declaration** 

**CONFORMITY ASSESSMENT ROUTE:** 

**Annex III Applied** 

STANDARDS APPLIED:

<u>EN ISO 13485:2003</u> – Medical devices – Quality Management Systems – Requirements for Regulatory Purposes

<u>EN ISO 14971:2012</u> – Medical devices – Application of risk management to medical devices

**EN ISO 18113-1:2011** – In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 1: Terms, definitions and general requirements

EN ISO 18113-2:2011 – In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 2: In vitro diagnostic reagents for professional use

**EN ISO 18113-3:2011** – In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 3: In vitro diagnostic instruments for professional use

BS EN 980:2008 – Symbols for use in the labelling of medical devices

<u>EN 13612:2002</u> – Performance evaluation of in vitro diagnostic medical devices

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Date

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#### STANDARDS APPLIED:

<u>EN 13640:2002</u> – Stability testing of in vitro diagnostic reagents

<u>ISO 15198:2004</u> – Clinical laboratory medicine — In vitro diagnostic medical devices — Validation of user quality control procedures by the manufacturer

<u>ISO 17511:2003</u> – In Vitro Diagnostic Medical Devices-Measurement of Quantities in Biological Samples-Metrological Traceability of Values assigned to Calibrators and Control Materials.

<u>ISO 17593:2007</u> – Clinical laboratory testing and in vitro medical devices – Requirements for in vitro monitoring systems for self-testing of oral anticoagulant therapy (Chapter 6 to be used for verification testing)

<u>WHO Technical Report Series 889 - Annex 3</u> – Guidelines for Thromboplastins and Plasmas used to control Oral anticoagulant therapy

<u>ISO 5725-2:1994</u> – Accuracy (trueness and precision) of measurement methods and results -- Part 2: Basic method for the determination of repeatability and reproducibility of a standard measurement method

Reach Article 33 & 67 – The system shall not contain Substances of Very High Concern (SVHC) in excess of limits set by Reach Articles 33 & 67

<u>IEC 62321, Ed.1:2008</u> – Procedures for the determination of levels of six regulated substances (Lead, Mercury, Cadmium, Hexavalent Chromium, Polybrominated Biphenyls, Polybrominated Diphenyl Ethers) in electrotechnical products

(EC) 1907/2006 – Regulation (EC) 1907/2006 of the European Parliament and the Council concerning the Registration, Evaluation, Authorization and Restriction of Chemicals (REACH)

**EN 61010-2-101:2002** – Safety requirements for electrical equipment for measurement, control, and laboratory use – Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

<u>CAN/CSA C22.2 No. 61010-1:2009</u> – Safety Requirements for Electrical Equipment for Measurement, Control, and Laboratory Use – Part 1: General Requirements.

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CAN/CSA C22.2 No. 61010-2-101:2009 - Safety requirements for electrical equipment for measurement. control, and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

**UL 61010-1-2008** – Safety Requirements for Electrical Equipment for Measurement, Control, and Laboratory Use - Part 1: General Requirements.

EN 61326-2-6:2006 - Electrical equipment for measurement, control and laboratory use. EMC requirements. Particular requirements. In vitro diagnostic (IVD) medical equipment

IEC/EN 61010-1:2001 - 2nd Edition - Safety requirements for electrical equipment for measurement, control, and laboratory use. General requirements

IEC/EN 61010-1:2010 - 3rd Edition - Safety requirements for electrical equipment for measurement, control, and laboratory use. General requirements

IEC 61010-2-101 Ed. 1 - Safety requirements for electrical equipment for measurement, control and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

EN 62304:2006 - Medical device software - Software life-cycle processes

EN 62366:2008 - Medical devices - Application of usability engineering to medical devices

ISTA Procedure 3A - Packaged-Products for Parcel Delivery System Shipments 70kg (150 lb) or Less (standard, small, flat or elongated)

2002/96/EC - Council Directive relating to the waste of electrical and electronic equipment (WEEE)

AS 60417.1:2004 - Graphical symbols for use on equipment

ASTM D3363-05 - Standard Test Method for Film Hardness by Pencil Test

IEC 60068-2-64:1993 - Environmental testing - Part 2: Test methods – Test Fh: Vibration, broadband random (digital control) and guidance

**IEC 60529:2001** – Degrees of protection by enclosures (IP code)

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We herewith declare that the below-mentioned product(s) meet the provisions of the Council Directive 98/79/EC for in vitro diagnostic medical devices and therefore has fulfilled all requirements for applying the CE mark to the medical device(s). The Manufacturer retains all supporting documentation.

### Attachment 1

SMN	REF	Product Code	Description
10873436	10873436	10873436	Xprecia™ Systems PT Controls
10873633	10873633	10873633	Xprecia™ System PT Controls
End of List			

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