## **SIEMENS**

## **EC Declaration of Conformity**



We hereby declare that the product described below conforms to all applicable requirements of Council Directive 98/79/EC for in vitro diagnostic medical devices.

**Legal Manufacturer:** Siemens Healthcare Diagnostics Inc.

511 Benedict Avenue

Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific

8365 Valley Pike

Middletown, VA, 22645-0307, USA

**EC Authorized Representative:** Siemens Healthcare Diagnostics Ltd.

Sir William Siemens Square

Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i PEROX SHEATH

Catalogue Number (REF): 03624240

Siemens Material Number (SMN): 10312275

Legacy Product Code: T01-3633-54

Classification: General IVD

Conformity Assessment Route: ANNEX III

**Document Control Number:** DoC\_ADVIA 120/2120/2120i PEROX SHEATH

Version: 1.0

Signature:

This declaration of conformity is issued under the sole responsibility of Siemens Healthcare Diagnostics Inc. This declaration supersedes any declaration issued previously for the same product.

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2015-11-12

Matthew Gee

Sr. Manager, Regulatory Affairs Siemens Healthcare Diagnostics Inc.

Tarrytown, NY, USA

Date [YYYY-MM-DD]