

EC Declaration of Conformity

I, the undersigned, hereby declare that the products specified below conform to Annex III of Directive 98/79/EC of the European Parliament and Council of 27 October 1998, on in-vitro diagnostic medical devices (and its relevant transposition into the national laws of the Member States in which the device is intended to be placed on the market).

Manufacturer: Ventana Medical Systems, Inc.

Address: 1910 E. Innovation Park Drive
Tucson, AZ USA 85755

European Authorized Representative: ROCHE DIAGNOSTICS GmbH
Sandhofer Strasse 116
D-68305 Mannheim
Germany

Manufacturing Site: Tucson, AZ USA

Product name: PAS Staining Kit

Ventana P/N: 860-014
Roche P/N: 05279291001

Classification: General IVD
Technical Data File: TDF-0054
Date of First CE Marking: 11-Jul-2006

Name of Authorized Signatory: Peter Martin
Vice President, Regulatory Affairs

Signature:  Date: 22-Aug-2010