

EU Declaration of Conformity

In accordance with EU Regulation 2017/746 of the European Parliament and of the Council of 5 April 2017 on *in vitro* diagnostic medical devices.

Manufacturer: Ventana Medical Systems Inc.

1910 E Innovation Park Drive Tucson, AZ 85755, USA

Single Registration Number (SRN)

Manufacturer:

US-MF-000016993

Authorized Representative: Roche Diagnostics GmbH

Sandhofer Strasse 116

68305 Mannheim

Germany

Single Registration Number (SRN)

Authorized Representative:

DE-AR-000006262

This declaration is issued under the sole responsibility of Ventana Medical Systems Inc.

Product Information

Part Number:	Product Name:	Basic UDI-DI:
05929903001 (alternative P/N: 790-4451)	CONFIRM anti-CD5 (SP19) Rabbit	761333601857BH
	Monoclonal Primary Antibody	

Intended Purpose: CONFIRM anti-CD5 (SP19) Rabbit Monoclonal Primary Antibody is intended

for laboratory use in the qualitative immunohistochemical detection of CD5 by light microscopy in sections of formalin fixed, paraffin embedded tissue

stained on a BenchMark IHC/ISH instrument.

Risk Class C

Common Specifications: Not applicable as no Common Specifications exist for the concerned device.

Name, Address and Identification number of the Notified Body:

TÜV SÜD Product Service GmbH (No. 0123)

Ridlerstraße 65 80339 MÜNCHEN

Germany

Conformity Assessment was established through the procedure described in Annex IX of EU Regulation 2017/746, including an assessment of the Technical Documentation as described in Annex IX, Chapter II. This declaration is supported by the following certificate(s):

EU Quality Management System Certificate: V12 096981 0003



Conformity of the product with EU Regulation 2017/746 and other applicable EU legislation has been established.

On behalf of Ventana Medical Systems Inc.

Place: Tucson, AZ 85755, USA Place: Tucson, AZ 85755, USA

Date: 22-Jun-2022 Date: 21-Jun-2022

Jeff Boone Ben Curson

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Site Head of Quality Function Site Head of Regulatory Affairs Function