

## **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:
05279771001 (alternative P/N: 950-102)	EZ Prep Concentrate (10X)	761333601901AT

Intended Purpose: Ventana Medical Systems' (Ventana) EZ Prep Concentrate (10X) solution

(EZ Prep) is used for paraffin removal from tissue samples during

immunohistochemistry and in situ hybridization reactions, and to dilute 2X SSC during stringency washes during in situ hybridization reactions carried

out on Ventana automated slide stainers.

This product is designed for use on BenchMark Series automated slide

stainers.

This product is intended for in vitro diagnostic (IVD) use.

Risk Class: Class A

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

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	
29-Mar-2022 Date:		
Jeff Boone	Ben Curson	
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Site Head of Quality Function Site Head of Regulatory Affairs Function