



## Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

| <b>MANUFACTURER</b>   |   |   |   |
|---|---|---|---|
| Name of Company   | Address   | SRN   |   |
| Bio/Data Corporation  | 155 Gibraltar Road,<br>Horsham, PA 19044 U.S.A.         | US-MF-000026991   |   |
| <b>AUTHORIZED REPRESENTATIVE</b>  |   |   |   |
| Name of Company   | Address   | SRN   | Telephone/email   |
| mdi Europa GmbH   | Langenhagener Str. 71<br>D-30855 Langenhagen<br>GERMANY | DE-AR-000006218   | +49-511-3908 9531 – phone<br><a href="mailto:info@mdi-europa.com">info@mdi-europa.com</a> |
| <b>PRODUCT IDENTIFICATION</b>   |   |   |   |
| Product / Trade Name  | Product Code / Catalog Number                           | Basic UDI-DI  |   |
| vW Factor Assay   | 103025  | ++G0561030253D  |   |
| Intended Purpose  |   | Photo   |   |
| <b>See Instructions for Use</b>   |   | <b>See website <a href="http://www.biodatacorp.com">www.biodatacorp.com</a></b> |   |
| vW Factor Assay® (Ristocetin CoFactor) Kit is a system of reagents and control plasmas that causes lyophilized platelets to agglutinate in the presence of patient Platelet Poor Plasma and Ristocetin for evaluation of von Willebrand Syndrome. |   |   |   |
| <b>IVDR RISK CLASS / COMMON SPECIFICATIONS</b>  |   |   |   |
| Device Classification   |   | Common Specifications   |   |
| Class   | A non-sterile   | No relevant common specifications have been published yet.                      |   |
| Rule  | 5a per Annex VIII of IVDR 2017/746                      |   |   |

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 An ISO 13485 Registered Company

**Bio/Data Corporation, the Manufacturer, declares that the above-mentioned product meets the provision of the following EU legislation:**

In Vitro Diagnostic Medical Device Regulation (IVDR) (EU) 2017/746

**Conformity Statement:**

Bio/Data Corporation confirms that the device covered by this declaration is in conformity with the (EU) IVDR 2017/746 Regulation and, if applicable, with any other relevant Union Legislation that provides for the issuing of an EU Declaration of Conformity.

**Notified Body (NB) Identification, conformity assessment procedure performed and identification if the certificate or certificates issued:**

Not applicable. The conformity assessment procedure for Class A Devices should be carried out, as a general rule, under the sole responsibility of manufacturers, since such devices pose a low risk to patients. (Self-Declaration)

**COMPANY REPRESENTATIVE: William M. Trolio****SIGNATURE:** **TITLE/FUNCTION:** Director of Quality Assurance & Regulatory Affairs for Bio/Data Corporation**PLACE:** Bio/Data Corporation, Horsham, PA. 19044 U.S.A.**DATE:** 1 April 2023