

## **Declaration of Conformity**

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

MANUFACTURER						
Name of Company		Address		SRN		
Bio/Data Corporation		155 Gibraltar Road,		US-MF-000026991		
			Horsham, PA 19044 U.S.A.			
AUTHORIZED REPRESENTATIVE						
Name of Company Addre		255	SRN	Telephone/email		
mdi Euro	pa GmbH	Langenhagener Str. 71		DE-AR-000006218	+49-511-3908 9531 – phone	
		D-308 GERM	55 Langenhagen IANY		info@mdi-europa.com	
PRODUCT IDENTIFICATION						
Product / Trade Name			Product Code / Catalog Number		Basic UDI-DI	
ADP / Adenosine-5'-		101312		++G05610131235		
diphosphate						
Intended Purpose					Photo	
See Instructions for Use					See website	www.biodatacorp.com
ADP Reagent (Adenosine-5'-Diphosphate) is for routine use in						
eliciting a concentration dependent activation or aggregation						
response in a Platelet Rich Plasma sample.						
IVDR RISK CLASS / COMMON SPECIFICATIONS						
Device Classification Comr			Common Specifica	ommon Specifications		
Class	A non-st	erile	No relevant common specifications have been published yet.			
Rule	5a per Ar VIII of IV 2017/7	/DR				

155 Gibraltar Road, Horsham, PA 19044 U.S.A. Worldwide: (215) 441-4000 U.S.A.: (800) 257-3282 Fax Worldwide: (215) 443-8820 www.biodatacorp.com e-mail: customer.service@biodatacorp.com An ISO 13485 Registered Company



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

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