

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