

## **EU Declaration of Conformity**

Manufacturers Name:	Boule Medical AB
Manufacturers Address:	Domnarvsgatan 4 SE-16353 Spånga, Sweden
SRN (Single Registration Number):	SE-MF-000001911

## **Product information**

Basic UDI-DI:	735004079A001Y6
EMDN code	W020201

UDI-DI	Product code	Product name
7350040796153	1420021	Medonic M-series M32B
7350040796160	1420022	Medonic M-series M32M
7350040796184	1420024	Medonic M-series M32C AR
7350040796207	1420026	Medonic M-series M32S BD AR

Table 1 – Products included in this Declaration of Conformity

Risk Classification	Class B	According to Rule 6
Intended purpose	The Medonic M-series M32 is a quantitative, multi-parameter automated hematology analyzer intended for blood cell counting and sizing for in vitro diagnostic use in screening of human capillary or venous whole blood samples collected in K2EDTA or K3EDTA blood collection tubes to collect data reflecting the patient's hematological status. The analyzer must be used together with Medonic M-series Diluent and M-series Lyse.	
	The device is intend	ded for professional use.
	the absolute numbe lymphocytes (LYM hemoglobin (HGB) cell hemoglobin (M distribution relative	ries M32 is used for enumeration of white blood cells (WBC); r and percentage concentration for granulocytes (GRA), (I), mid-sized white cells (MID); red blood cells (RBC); ; mean cell volume of red cells (MCV); hematocrit (HCT); mean (ICH); mean cell hemoglobin concentration (MCHC); red cell width (RDW%); platelets (PLT); mean platelet volume (MPV).
	the age of 1 month.	1 1
IVDR assessment	Assessment for CE	marking following Annex IX (Chapter I and III), Conformity
route	Assessment Based	on a Quality Management System and on Assessment of
	Technical Document	ntation

## Applicable European directives and regulations

Regulation/Dir	rective
EU 2017/746	In Vitro Medical Devices Regulation
2011/65/EU	Restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS) Directive
2006/42/EC	Machinery Directive

Table 2 – Applicable regulations/directives



Notified body	Identification number	
BSI Group The Netherlands B.V.	NB 2797	
Say Building, John M. Keynesplein 9, 1066 EP		
Amsterdam, Netherlands		

Certificate number	Date of issue
IVDR 745210	2022-11-01

Boule Medical AB hereby declares that the medical device(s) listed in *Table 1* are in conformity with applicable requirements of the Directive/Regulation/s listed in *Table 2*.

This declaration of conformity is issued under the sole responsibility of and signed on behalf of Boule Medical AB

Spånga, 2023-02-14

Place and date of issue

Jesper Söderqvist, CEO

Spånga, 2023-02-14

Place and date of issue

Terese Wasell, PRRC