

EU DECLARATION OF CONFORMITY

RR Mechatronics Manufacturing B.V.

P.O.Box 225

This declaration of conformity is issued under the sole responsibility

1620 AE HOORN

of the manufacturer:

The Netherlands

De Corantijn 13

RR Mechatronics Manufacturing B.V.

1689 AN ZWAAG

De Corantijn 13, 1689 AN Zwaag, The Netherlands

The Netherlands

SRN: NL-MF-000023105

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www.rrmechatronics.com

We declare that the:

The Automatic Erythrocyte Sedimentation Rate analyzer series:

Trade name/ Model:

Starrsed TL

Variants:

Variant:	Product-ID (REF):	UDI-DI
Starrsed TL 120V	A0026352	08719189137750
Starrsed TL 230V	TLAX109100	08719189137330

Basic UDI-DI (BUDI-DI): 8719189137A385ZC87

EMDN-code:

W02029001 Erythrocyte sedimentation rate devices

Classification IVDR:

Class A

Intended purpose:

Automated analyzer for in vitro determination of the Erythrocyte Sedimentation Rate

(ESR) of human blood samples in conformity with the Westergren standard.

For professional use in medical laboratories only. To be used for all patient populations,

not restricted by age or any other anatomical or physiological particulars.

is in conformity with the requirements of the following EU legislation(s):

Regulation (EU) 2017/746

In vitro diagnostic medical devices

(conformity assessment according Article 48, §10 of this regulation)

DOC-259 F4401 DoC Starrsed TL.v1

Conditions: By the ORGALIME GENERAL CONDITIONS S2012 of March 2012. Any other conditions are herewith explicitly rejected by us. VAT number: NL 8557 65 392 B01 CoC Number 64657353



Directive 2011/65/EU

Restriction of the use of certain hazardous substances

Including the amendment of Annex II; 2015/863

(conformity assessment according Article 7 of this directive)

The following harmonized standards have been applied:

EN ISO 13485:2016

Medical devices – Quality management systems – Requirements for regulatory purposes

EN ISO 14971:2012

Medical devices – Application of risk management to medical devices

EN-ISO 15223-1:2021

Medical devices – Symbols to be used with medical device labels, labelling and information to

be supplied – Part 1: General requirements

EN ISO 18113-3:2011

In vitro medical devices – Information supplied by the manufacturer (labelling) – Part 1: Terms,

definitions and general requirements

EN IEC 61010-2-101:2002 Safety requirements for electrical equipment for measurement, control, and laboratory use –

Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment

EN IEC 61326-2-6:2006

Electrical equipment for measurement, control and laboratory use – EMC requirements – Part

2-6: Particular requirements – In vitro diagnostic (IVD) medical equipment

EN IEC 62366-1:2008

Medical devices – Application of usability engineering to medical devices

The CE mark was applied for the first time on this type of IVD device in 2015.

Place:

Zwaag, The Netherlands

Date:

May 14, 2024

Signature:

Stamp:

Name:

.S van der Meer

Function title:

CEO

RR Mechatronics Manufacturing B.V.

De Corantiin 13 1689 AN Zwaag The Netherlands Walter Park - V.8 2 May

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