

EU DECLARATION OF CONFORMITY

RR Mechatronics Manufacturing B.V.

This declaration of conformity is issued under the sole responsibility of the manufacturer:

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The Netherlands

RR Mechatronics Manufacturing B.V.
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We declare that:

The whole blood diluent:

Trade name: **Starrsed Diluent**

Product models:

Model name:	Product-ID (REF):	Basic UDI-DI
Starrsed Diluent	QRR010931	08719189137194

EMDN-code: W0103010105 Cbc-reagents (cleaning-/diluting-/lysing-/sheat-fluids)
Classification IVDR: Class A

Intended purpose: Product for the automatic dilution of blood samples in Starrsed ESR analyzers

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)

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 18113-1:2011	In vitro medical devices – Information supplied by the manufacturer (labelling) – Part 1: Terms, definitions and general requirements
EN ISO 18113-2:2011	In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 2: In vitro diagnostic reagents for professional use
EN ISO 23640:2015	In vitro medical devices – Evaluation of stability of in vitro diagnostic reagents

The CE mark was applied for the first time on this type of IVD reagent in 1998.

Place: Zwaag, The Netherlands
Date: May 12, 2022

Signature:

Stamp:

Name: Jan Buis

Function title: CEO
RR Mechatronics Manufacturing B.V.

