



Declaration of Conformity



According to Art. 17 of Regulation 2017/746 (EU) on in vitro diagnostic medical devices

Manufacturer: E-LAB Biological Science & Technology Co., Ltd. (CN:MF-000020119)

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Bsaic UDI: 697517840003M3

Product: Auto Hematology Analyzer

Model: EC-20, EC-21, EC-28, EC-30, EC-31, EC-36, EC-38A, EC-38B, EC-38

Classification: Class A, according to Rule 5 of IVDR Annex VIII

We, manufacturer, herewith declare under our sole responsibility that the above-mentioned product meets the provisions of the following EC Council Directives and Standards. All supporting documentation is retained under the premises of the manufacturer.

List of Directive and Standard Applied:

EN ISO 14971:2019

EN ISO 13485:2016

ISO 15223-1:2021

EN ISO 18113-1:2011

EN ISO 18113-2:2011

IEC 61010-1:2017

IEC61010-2-101:2018

IEC 61326-1:2020

IEC61326-2-6:2020

IEC 62366-1:2015

Nanjing March 21, 2022

(Place and Date of Issue)

General Manager: 

(Signature and Position)

Signed for and on behalf of the Manufacturer