

EU DECLARATION OF CONFORMITY

Manufacturer's Information

Manufacturer's Name:	Norma Instruments Zrt.	
Manufacturer's SRN:	HU-MF-000014244	
Manufacturer's Address:	Papírgyár u. 58-59., 1038 Budapest, Hungary	

Product Information

Product Name:	Norma Icon-5 OP Automated 5-part Hematology Analyzer
Product Code:	
Basic UDI-DI:	59991062Icon5OPU7
Intended Purpose:	Icon-5 OP is a 57 tests/hour laser and impedance based hematology analyzer for laboratory use, using specific reagents, performing 5-part differential analysis of 27 parameters of anti-coagulated human whole blood samples introduced in open vials.
Risk Classification:	Class A
EMDN code:	W0202010101

Applicable Regulations, Directives and Standards

Regulations:	2017/746/EU on in vitro diagnostic medical devices (IVDR)
Directives:	2011/65/EU on restriction of hazardous substances (ROHS) Classification: Category 8 Medical Device 2015/863/EU amending Annex II to Directive 2011/65/EU of the European Parliament and of the Council as regards the list of restricted substances 2014/30/EC on electromagnetic compatibility (EMC) 2014/35/EC on low voltage (LVD)
Standards:	EN 61010-1: 2010 IEC 61010-2-101: 2017 EN 60825-1: 2014 EN 61326-2-6: 2013

We, **Norma Instruments Zrt.**, as a manufacturer of the above-mentioned CE marked in-vitro medical devices, do hereby declare that this EU Declaration of Conformity is issued under our sole responsibility.

Devices covered by the present EU Declaration of Conformity is in conformity with **Regulation (EU)** 2017/746 on in vitro diagnostic medical devices (IVDR) and other directives and standards specified above. This declaration is supported by the harmonized quality management system approval of EN ISO 13485:2016 issued by SGS Hungary Kft. (HU15/7647). All supporting documentation is retained at the premise of the manufacturer.

Budapest, 2024/04/11

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