

EU Declaration of Conformity

(N° dc90151ben)

WE THE MANUFACTURER

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Single Registration Number	FR-MF-000000320

TAKE SOLE RESPONSIBILITY FOR AND HEREBY DECLARE THAT THE PRODUCT(S)

Device category	Hematology Analyzer Accessory
Product name	Yumizen Compressor
Basic UDI-DI	361023ymz_compressorN9
Country of origin	FRANCE

Intended Use

The Yumizen Compressor is an accessory to the Yumizen H1500 or Yumizen H2500 analyzer used for *in vitro* diagnostic testing.
Clinical laboratories use.

MEET(S) THE PROVISIONS OF THE FOLLOWING DIRECTIVES, REGULATIONS, STANDARDS AND COMMON SPECIFICATIONS

Regulations	Regulation (EU) 2017/746 on <i>in vitro</i> diagnostic medical devices Risk Class: A ☒ B ☐ C ☐ D ☐
IVDR conformity assessment route	☒ ANNEX II + ANNEX III + ANNEX IV (<i>Class A devices excluding sterile devices</i>)
Directives	2011/65/EU - Amended by 2015/863/EU - ROHS Directive Category: 8- Medical Devices
Standards	EN IEC 63000: 2018
Common Specifications	Not applicable

Montpellier, France
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Claire MALLIÉ
Quality & Regulatory Affairs Junior
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