

EU DECLARATION OF CONFORMITY

Code: MOL_DECO_SI17010105_MMYEASTBLOOD-1-0bis_EN

Date:

2023-10-13

The Manufacturer:

Manufacturer Name/Registered trade Name	ALIFAX S.R.L.
Address	Via Merano 30
	33045 NIMIS (UD)
	ITALY

declares on his own responsibility that the following device:

Product Name and	SI 1701.0105	MM YEAST BLOOD
Product Code		
Intended Purpose	clinical laboratory users, for the multiply specific to yeasts starting from extracted YEAST BLOOD cartridge contains all the test through a multiplex Real Time PCI MOUSE SYSTEM. MM YEAST BLOOD analyzes the foliglabrata, Candida, krusei, Candida pa Candida lusitaniae, Candida dubliniens MM YEAST BLOOD provides a result to yeast. The MM YEAST BLOOD test is intend analytical results within a diagnostic specific laboratory. MM YEAST BLOOD on culture testing. The result does not exclude the combined in the specific laboratory and years.	itro diagnostic test, intended for professional plex identification of nucleic acid sequences ed DNA from a positive blood culture. The MM he necessary reagents to perform one single R analysis in combination with MOLECULAR lowing targets: Candida albicans, Candida rapsilosis, Candida tropicalis, Candida auris, sis, Candida guilliermondii. support the diagnosis of blood infection with ed for combined use with other clinical and evaluation defined and regulated by each does not replace traditional methods based bined presence of targets other than those regets for the identification of which other
Risk Class	Other IVD Medical Device as it is neithed Directive 98/79/EC"	er in List A nor in List B of Annex II of the

complies with the Essential Requirements of the Directive 98/79/EC related to the In Vitro Diagnostic Medical Devices (IVDD).

We declare also that the device has been designed and manufactured in conformity to the below:

Harmonised Standards	EN ISO 13485:2016/A11:2021: Medical devices - Quality management systems - Requirements for regulatory purposes EN ISO 14971:2019: 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 EN ISO 18113-1:2011: In vitro diagnostic medical devices - Information supplied by the manufacturer -Part1: Terms, definitions and general requirements EN ISO 18113-2:2011: In vitro diagnostic medical devices - Information supplied by the manufacturer - Part 2: In vitro diagnostic reagents for professional use EN 13612:2002/AC:2002: Performance evaluation of in vitro diagnostic medical devices EN ISO 23640:2015 In vitro diagnostic medical devices - Evaluation of stability of in vitro diagnostic reagents
Technical Standards	IEC 62366-1:2015: Application of usability engineering to medical devices



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	IEC TR 62366-2:2016: Guidance on the application of usability engineering to medical device
Other	Directive 2015/863/EU (RoHS3)

The device has been CE Marked as IVD Medical Device/Devices according to Annex III of the Directive 98/79/EC as they are neither in List A nor in List B of Annex II.

Any change made to the devices without the prior consent of the Manufacturer will automatically void the present Declaration of Conformity.

Place and date of issue: NIMIS (UD), 24th May 2022

Name and function: Camillo Galiano, Managing Director

Signature: