



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 Product Code	SI 1701.0105	MM YEAST BLOOD
Intended Purpose	<p>MM YEAST BLOOD is a qualitative in vitro diagnostic test, intended for professional clinical laboratory users, for the multiplex identification of nucleic acid sequences specific to yeasts starting from extracted DNA from a positive blood culture. The MM YEAST BLOOD cartridge contains all the necessary reagents to perform one single test through a multiplex Real Time PCR analysis in combination with MOLECULAR MOUSE SYSTEM.</p> <p>MM YEAST BLOOD analyzes the following targets: <i>Candida albicans</i>, <i>Candida glabrata</i>, <i>Candida krusei</i>, <i>Candida parapsilosis</i>, <i>Candida tropicalis</i>, <i>Candida auris</i>, <i>Candida lusitanae</i>, <i>Candida dubliniensis</i>, <i>Candida guilliermondii</i>.</p> <p>MM YEAST BLOOD provides a result to support the diagnosis of blood infection with yeast.</p> <p>The MM YEAST BLOOD test is intended for combined use with other clinical and analytical results within a diagnostic evaluation defined and regulated by each specific laboratory. MM YEAST BLOOD does not replace traditional methods based on culture testing.</p> <p>The result does not exclude the combined presence of targets other than those included in the list of identifiable targets for the identification of which other independent tests are necessary.</p>	
Risk Class	Other IVD Medical Device as it is neither in List A nor in List B of Annex II of the Directive 98/79/EC"	

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:


ALIFAX s.r.l.
Managing Director
Camillo Galiano