## **EC Declaration of Conformity**

Manufacturer/
Supplier Information:

BioFire Diagnostics, LLC

515 Colorow Drive
Salt Lake City, Utah 84108, USA
Phone: 1-801-736-6354
Regulatory@BioFireDX.com
http://www.BioFireDX.com

We, BioFire Diagnositics, LLC, declare under our sole responsibility, that the product

# FilmArray® Blood Culture Identification (BCID) Panel (RFIT-ASY-0126, RFIT-ASY-0127)

meets the provisions of the European Directive 98/79/EC for *In vitro* Diagnostic Medical Devices. The device is classified as a General In Vitro Diagnostic (IVD) Device.

BioFire Diagnostics' quality system is registered to EN ISO 13485:2016.

The following relevant standards have been met:

#### ISO 13485:2016/EN ISO 13485:2016

Medical devices - Quality Management System - Requirements for regulatory purposes

#### EN ISO 14971:2019

Medical devices – Application of risk management to medical devices

#### EN 13641:2002

Elimination or reduction of risk of infection related to in vitro diagnostic reagents

#### EN 62366:2008

Medical devices-Application of usability engineering to medical devices

#### EN 13612:2002

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

### EN ISO 15223-1:2016

Medical Devices – Symbols to be used with medical device labels, labeling and information to be supplied – Part 1: General requirements

#### EN ISO 18113-1:2011

In vitro diagnostic medical devices – Information supplied by the manufacturer (labeling) – Part 1: Terms, definition and general requirements

#### EN ISO 18113-2:2011

In vitro diagnostic medical devices – Information supplied by the manufacturer (labeling) – Part 2: In vitro diagnostic reagents for professional use

Technical documentation demonstrating compliance as described in Annex III of the European Directive 98/79/EC is kept by the manufacturer and can be made available by the authorized representative in Europe - QARAD EC-REP BV, Pas 257, B-2440 Geel, Belgium.

The notified body for this product is BSI (Notified body #2797; Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, Netherlands).

Salt Lake City, UT, USA	
(Place and date of issue)	Kevin Bourzac
	Vice President, Regulatory and Clinical Affairs



www.biofiredx.com