

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 Diagnostics, LLC, declare under our sole responsibility, that the product

**FilmArray® Gastrointestinal (GI) Panel
(RFIT-ASY-0104, RFIT-ASY-0116)**

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:

EN ISO 13485:2016 Medical devices – Quality Management System – Requirements for regulatory purposes
EN ISO 14971:2012 Medical devices – Application of risk management to medical devices
EN 13641:2002 Elimination or reduction of risk of infection related to <i>in vitro</i> 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 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



BY BIOMÉRIEUX

515 Colorow Drive | Salt Lake City | Utah | 84108
p: 1-801-736-6354 | f: 1-801-588-0507
www.biofire.com