## **EC Declaration of Conformity**

Manufacturer/ Supplier Information:	Maine Molecular Quality Controls, Inc.
	23 Mill Brook Road Saco, Maine 04072 USA
	Phone: 1-207-885-1072
	Email: info@mmqci.com
	Website: www.mmqci.com
European Representative	QARAD EC-REP BV
	Pas 257 2440 Geel, Belgium

We, Maine Molecular Quality Controls Inc., declare under our sole responsibility, that the product:

## BioFire® RP2.1/RP2.1 plus Control Panel M441

Catalog #: M441

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. The MMQCI quality system is registered to EN ISO 13485:2016. Conformity Assessment Method: General IVD Annex III.

The undersigned declares that the device listed above conforms to the essential requirements described in ANNEX I of the European Directive 98/79/EC. Technical documentation demonstrating compliance as described in Annex I is kept by the manufacturer and can be made available by the authorized representative in Europe - QARAD EC-REP BV, Pas 257, 2440 Geel, Belgium.

Saco, Maine USA / Offospore

(Place and date of Issue)

(Signature and Title)

