



# EU Declaration of Conformity

In accordance with EU Regulation 2017/746 of the European Parliament and of the Council of 5 April 2017 on *in vitro* diagnostic medical devices.

Manufacturer: **Roche Molecular Systems, Inc.**  
**1080 US Highway 202 South**  
**Branchburg, NJ 08876**  
**USA**

Single Registration Number (SRN) **US-MF-000018066**  
 Manufacturer:

Authorized Representative: **Roche Diagnostics GmbH**  
**Sandhofer Strasse 116**  
**68305 Mannheim**  
**Germany**

Single Registration Number (SRN) **DE-AR-000006262**  
 Authorized Representative:

This declaration is issued under the sole responsibility of Roche Molecular Systems, Inc.

## Product Information

Part Number:	Product Name:	Basic UDI-DI:
07958021190	<b>cobas</b> <sup>®</sup> PCR Media Dual Swab Sample Kit	761333601941B7

**Intended Purpose:** The **cobas**<sup>®</sup> PCR Media Dual Swab Sample Kit is used to collect and transport human specimens. The **cobas**<sup>®</sup> PCR Media serves as a nucleic acid stabilizing transport and storage medium for human specimens.

The complete Intended Use is contained in the **cobas**<sup>®</sup> PCR Media Dual Swab Sample Kit Package Insert.

**Risk Class and Classification Rule:** Class A, as per EU Regulation 2017/746, Annex VIII, Rule 5 (a)

**Common Specifications:** Not applicable as no Common Specifications exist for the concerned device.

Conformity of the product with EU Regulation 2017/746 and other applicable EU legislation has been established.



On behalf of Roche Molecular Systems, Inc.

Place: Tucson, AZ

Date: 21-Dec-2021

DocuSigned by:  
*Jeff Boone*

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**Jeff Boone**

Vice President, Quality Management

Place: Santa Clara, CA

Date: 20-Dec-2021

DocuSigned by:  
*Carolyn Glickman*

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**Carolyn Glickman**

Director, Regulatory Affairs