

## **EU Declaration of Conformity**

as per Annex IV of the Regulation EU 2017/746 on in-vitro diagnostic medical devices

**Manufacturer:** Roche Diagnostics GmbH  
**Address:** Sandhofer Strasse 116  
 68305 Mannheim  
 Germany

**Single Registration Number:** DE-MF-000006260

*Roche Diagnostics GmbH declares, under the sole responsibility, that the product/the product line*

<b>Product Name</b>	<b>Cat. No.</b>	<b>Basic UDI-DI</b>
CalSet Vials	11776576322	761333601249AL

### ***Intended Use:***

The CalSet Vials are empty SnapCap bottles intended to be used as an IVD accessory to aliquot the reconstituted or pre-filled Assay CalSets into smaller portions for storing. The CalSet vials are placed in a 5 position rack for further processing.

***Risk Class:***  A  B  C  D

***Conformity Route:***  *Self-Declaration of Conformity (Class A)*  
 *Self-Declaration of Conformity after Notified Body involvement for sterile manufacturing conditions acc. Art. 48 (10) (Class A sterile)*  
 *Technical Documentation Assessment Class B/C – Annex IX*  
 *Technical Documentation Assessment Class D – Annex IX*  
 *Technical Documentation Assessment Class B/C/D for Self-Testing – Annex IX*  
 *Technical Documentation Assessment Class B/C/D for Near-Patient Testing – Annex IX*  
 *Technical Documentation Assessment Class C/D for Companion Diagnostics – Annex IX*

***Certificates:***  *EU QM Certificate No.:*  
 *EU Technical Documentation Assessment Certificate No. (Class D, Near-Patient Testing, Self-Testing and Companion Diagnostics):*

***Other:***  *Common Specifications:*

***Notified Body (NB) Name:*** N/A  
***NB Address:***

***NB Ident. No.:*** N/A

*to which this declaration relates fulfils the requirements of Regulation EU 2017/746 on in-vitro diagnostic medical devices.*

Mannheim, 19 June 2023

Roche Diagnostics GmbH

*i.V./on behalf of the company*

*ppa./on behalf of the company*

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