

# EC Declaration of Conformity

<b>Manufacturer Name:</b>	RefLab Aps
<b>Manufacturer Address:</b>	Ole Maaløes Vej 3, COBIS 2200 Copenhagen N DK-Denmark
<b>SRN (Single Registration Number):</b>	DK-MF-000021844
<b>Basic UDI-DI:</b>	5744003280STRIP6M
<b>Name of the Device(s):</b>	Stripping buffer, 100 mL, Stripping buffer 1 L
<b>Catalogue number(s):</b>	RLA701, RLR016
<b>Product code:</b>	W0201020285
<b>Classification:</b>	IVDR, Class A
<b>Notified Body name:</b>	N/A
<b>Notified Body Address:</b>	N/A
<b>Notified Body Identification number:</b>	N/A
<b>Conformity assessment route:</b>	RefLab Aps uses the following procedures for the CE-labeling of their products according to the Regulation IVDR 2017/746:  <u>Class A:</u> EC conformity declaration according to Annex IV.

This declaration of conformity is issued under the sole responsibility of RefLab Aps. We hereby declare that the in-vitro diagnostic medical device(s) specified above meet the provision of the Regulation (EU) IVDR 2017/746. This declaration is supported by the ISO 13485 Quality Management System approval issued by DQS. All supporting documentation is retained at the premises of the manufacturer.

**Signature:**



Lise Bo  
Quality Manager

**Place and date (dd.mm.yyyy) of issue:**

Copenhagen, 24.05.2022

**RefLab ApS**