

EC Declaration of Conformity

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|---|---|
| Manufacturer Name: | RefLab Aps |
| Manufacturer Address: | Ole Maaløes Vej 3, COBIS 2200 Copenhagen N DK-Denmark |
| SRN (Single Registration Number): | DK-MF-000021844 |
| Basic UDI-DI: | 574400328STOPFW |
| Name of the Device(s): | Stop reagent (HClO ₄) 100 mL, Stop reagent (HClO ₄) 1 L |
| Catalogue number(s): | RLR011, RLA703 |
| Product code: | W0201020285 |
| Classification: | IVDR, Class A |
| Notified Body name: | N/A |
| Notified Body Address: | N/A |
| Notified Body Identification number: | N/A |
| Conformity assessment route: | 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, 20.05.2022

RefLab ApS