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Verification Report

according to Regulation (EU) 2017/746 on In vitro Diagnostic Medical Devices – Annex IX, sec. 4.12 or Annex XI sec. 5

No. BRM-802330 24 11 070271

Manufacturer: **Bio-Rad Medical Diagnostics GmbH**
Industriestraße 1
63303 Dreieich

Product: **Seraclone Anti-E (RH3)**

Testplan: **TP-BRM-802330**

Batch: **3439021-00**


Basic UDI-DI: **361052A003948M**

Expiry Date: **2025-12-21**

The above mentioned batch meets the batch release criteria established during technical documentation assessment and may be placed on the market. The EU Technical Documentation Assessment Certificate (IVDR) issued for this product is V70 118577 0003 Rev. 00.

This verification report is valid for REF 802330 (1 x 5ml) and REF 802331 (1 x 10ml).

Date, 2024-11-13


Ellen Preuß (13. November 2024 15:00 GMT+1)
Adobe Acrobat Sign-Transaktionsnummer: CBJCHBCAABAAXShTscub19IG8GemYg7ijbY6drki3ELm
pp Dr. Ellen Preuß
In-vitro Diagnostics

TÜV SÜD Product Service GmbH is Notified Body according to Council Vitro Diagnostic Regulation 2017/746 concerning In-vitro Diagnostic Medical Devices with Identification No. 0123.

To:

Bio-Rad Medical Diagnostics GmbH

cc:

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