





## CERTIFICATE

No. QS6 082277 0008 Rev. 02

Certificate Holder: BioPorto Diagnostics A/S

Tuborg Havnevej 15, st.

2900 Hellerup DENMARK

**Certification Mark:** 



Scope of Certificate: Design and Development, Manufacture and Distribution

of In-Vitro Diagnostic Medical Devices used in the Diagnosis of Immune Status and Renal Disorders

Standard(s): ISO 13485:2016

Regulatory Authority(ies): Health Canada, USA FDA. See attached for listing of

specific regulatory requirements.

The Certification Body of TÜV SÜD America Inc. certifies that the quality management system of the manufacturer listed above has been audited against the stated criteria and found to conform to those criteria for the scope of certification listed. Validity of this certificate can be obtained by visiting the website www.tuvsud.com/ps-cert

TÜV SÜD America Inc. is an MDSAP Recognized Auditing Organization.

REPs Facility ID: F002670

**Effective Date: 2023-01-16** 

**Expiry Date:** 2025-01-16

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Date of Issue: 2023-02-24

( Renee Walker )

Director, US Certification Body, MHS





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Regulatory Requirements: Audit/Certification Criteria

Canada

- Medical Device Regulations - Part 1- SOR 98/282

**United States** 

- 21 CFR Part 803 - 21 CFR Part 806

- 21 CFR Part 807 - Subparts A to D

- 21 CFR Part 820

Facility(ies): BioPorto Diagnostics A/S

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