







EU Quality Management System Certificate (IVDR)

Pursuant to Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices, Annex IX Chapters I and III (Class C and B Devices excluding self-/near-patient-testing and Companion Diagnostics)

No. V12 114429 0002 Rev. 00

Manufacturer:

Generi Biotech s.r.o.

Machkova 587/42 Třebeš 500 11 Hradec Králové 11 CZECH REPUBLIC

SRN Manufacturer - CZ-MF-000018857

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has established, documented and implemented a quality management system as described in Article 10 (8) of the Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices. Details on devices covered by the quality management system are described on the following page(s). The Report referenced below summarizes the result of the assessment and includes reference to relevant CS, harmonized standards, audit and test reports. The conformity assessment has been carried out according to Annex IX Chapter I and III of this regulation with a positive result. The quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis.

The certified quality management system is subject to periodical surveillance by TÜV SÜD Product Service GmbH. The surveillance assessment includes an assessment of the technical documentation for the device or devices concerned on the basis of further representative samples. For details and certificate validity see: www.tuvsud.com/ps-cert?g=cert:V12 114429 0002 Rev. 00

Report No.:	713230259
Valid from:	2023-10-18
Valid until:	2028-10-17

2023-10-18

Issue date:

Mortecondel

Marta Carnielli Head of Certification IVD

Page 1 of 2 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123 TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany







EU Quality Management System Certificate (IVDR)

Pursuant to Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices, Annex IX Chapters I and III (Class C and B Devices excluding self-/near-patient-testing and Companion Diagnostics)

No. V12 114429 0002 Rev. 00

Classification: Device Group: IVP Code: Intended Purpose:	Class C W0106 - GENETIC TESTING IVP 3011 - In vitro diagnostic devices which require knowledge regarding molecular biological testing including nucleic acid assays and next generation sequencing (NGS) IVR 0402 - Devices intended to be used to predict genetic disease/disorder risk and prognosis
Classification: Device Group: IVP Code: Intended Purpose:	Class C W0106 - GENETIC TESTING IVP 3011 - In vitro diagnostic devices which require knowledge regarding molecular biological testing including nucleic acid assays and next generation sequencing (NGS) IVR 0403 - Other devices intended to be used for human genetic testing

The validity of this certificate -nonedepends on conditions and/or is limited to the following:

Revision History:

 Rev.
 Dated
 Report

 00
 2023-10-18
 713230259

Description Initial issuance