



CERTIFICATE



Management System as per EN ISO 13485 : 2016

In accordance with TÜV AUSTRIA procedures, it is hereby certified that

LERIVA PHARMA S.A.

HEAD OFFICE: 33, Pigis Av. & 6, Anapiron Polemou Str., GR-151 27 MELISSIA, GREECE

BRANCH 1: Lakkos Kamaterou, GR-193 00 ASPROPYRGOS, GREECE

BRANCH 2: 10th km Thermis – Thessalonikhs, GR-570 01 THERMI, GREECE

Applies a Quality Management System for Medical Devices in line with the above Standard for the following Scope

IMPORT, TRADING AND DISTRIBUTION OF:

- ACTIVE MEDICAL DEVICES AND ASSOCIATED CONSUMABLES IN THE FIELD OF RADIOLOGY AND CARDIOLOGY
- GENERAL, NON-ACTIVE, NON-STERILE SELF-CARE MEDICAL DEVICES

TECHNICAL SUPPORT OF ACTIVE MEDICAL DEVICES IN THE FIELD OF RADIOLOGY AND CARDIOLOGY

Certificate Registration No.: 20302190002308

Valid until: 2025-10-16

Initial certification: 2019-10-17

Maria Agapitou
Head of Management Systems & Products Certification Division

Certification Body
at TÜV AUSTRIA

Athens, 2024-10-10

This certification was conducted in accordance with TÜV AUSTRIA auditing and certification procedures and is subject to regular surveillance audits.

TÜV AUSTRIA HELLAS
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GEMI No: 1650201000



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Headquarters in Athens bear the responsibility of the Certification decision