EU Certificate

for the assessment of the quality management system



according to Medical Device Regulation (EU) 2017/745 Annex IX Chapter I+III

As a Notified Body of the European Union DEKRA Certification GmbH certifies, that the manufacturer

Stihler Electronic GmbH

Single Registration Number (SRN): DE-MF-000006188 Gaussstrasse 4, 70771 Leinfelden-Echterdingen, Germany

applies a quality management system according to Annex IX Chapter I+III of the Medical Device Regulation (EU) 2017/745 for the medical devices listed in the annex. This certificate is based on the assessments listed in CNo50192-00 and is only valid in conjunction with the successful completion of the annual surveillance audits.

EU Certificate no.: 50192-60-00-00 Certificate valid from: 2024-08-12 Certificate valid to: 2028-07-23



DEKRA Certification GmbH, Stuttgart Notified Body ID number: 0124

Annex to the EU Certificate no. 50192-60-00-00

Following devices/device categories are included in this certificate:

Class IIa

ASTOTUBE®

Article number: M77432002

M77449004 M77449003 M77455502 M77460002

Basic-UDI-DI: 426011414TUBE5P

EMDN Code: A030201MDN Code 0102

Change(s) to previous certificate: n.a.