## **Certificate of Designation**



## **Eudamed Mandate Summary**

Client Ref.	USA/2015/08/04	Date of Issue: 26 October 2023
Issued To:	Avery Biomedical Devices Inc 61 Mall Drive Commack NY 11725 USA	Legal Manufacturer [SRN: US-MF-000001250]
Issued By:	Advena Limited Tower Business Centre, 2nd Flr, Tower Street, Swatar, BKR 4013. Malta.	EC-REP [SRN: MIT-AR-00000234]
EU Competent Authority:	Malta Medicines Authority (MMA) Sir Temi Zammit Buildings, Malta Life Sciences Park, San Gwann SGN 3000 Malta. Tel: +356 2343 9000 Email: info.medicinesauthority@gov.mt	Eudamed Actor ID: MT-CA-019
is issued and confirm acts as EU Authorise This certificate alon market. The Legal	ns the period of representation. Furthermore, t ed Representative for the Legal Manufacturer. e does not provide confirmation that the devi	acturer and Advena Limited, this Certificate of Designation his certificate confirms the medical devices Advena Limited ces listed in Appendix A can be legitimately placed on the ctory regulatory evidence that the devices mentioned in n and have the applicable valid certifications.
as applicable to EU	legislation: Advena Ltd. Tower Business Centre, 2nd Flr., Tower Street, Swatar, BKR 4013 Malta	U Authorised Representative, and in the following format,
Anthony Kirby – Ma	ATE OF maging Director	DESIGI
AR Cover Begins: [AIMDD]	01 November 2023 AR Cover Ends: 31 Mandate Start: 12 October 2022 M	October 2024 andate End: N/A Mandated for Vigilance: No
נסטואוסטן		andate that with with a standated for vignatice. NO

This certificate is subject to the organisation maintaining their documentation in compliance with the EU legislation as indicated in this certificate.

This certificate is for the exclusive use of Advena Ltd's clients and is provided pursuant of the European Authorised Representative agreement (Mandate) between Advena Ltd and the client. Advena's responsibility and liability is limited to the terms and conditions of this agreement. Advena Ltd assumes no liability to any party for any loss, expense or damage occasioned by the use of this certificate and the European Authorised Representative agreement (Mandate). Only the client is authorised to copy or distribute this certificate. Any use of the Advena Ltd name by others who are not covered by the above agreement, or any similar contract, is prohibited. This certificate remains valid until the expiry date has been reached or has been terminated by Advena Limited.



## **Appendix A**

Product Details, Names or Trade Names	EU Legislation	Classification	Date of Declaration
Implanted Diaphragmatic/Phrenic Nerve Stimulator (breathing pacemaker)	AIMDD	Class III	25/05/2021

, ADVENA LIMITED

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Table 1: List of devices (generic device group(s)) covered within the executed Mandate (authorised representative list).