

Certificate of Registration

In accordance with European Medical Device Regulation MDR (EU) 2017/745, we hereby declare that:

- An examination has been made of this organization's Declaration of Conformity(s) and where appropriate Notified Body certification(s) exist.
- The EU Authorized Representative contract has been fulfilled.
- Device registrations for the medical devices mentioned within this certificate have duly been completed with an EU Competent Authority.

Therefore, these devices have met the requirements of the MDR (EU) 2017/745 and the CE mark may be applied to the products listed below.

Certificate:	Issue Date: 21st December 2021
Legal Manufacturer	EU Authorized Representative (EC REP)
PROTEOR USA, LLC	PROTEOR SAS
3 Morgan	6 rue de la Redoute
Irvine, CA 92618	21850 Saint APOLLINAIRE
USA	France

Product Details, Names or Trade Names
Artificial Limbs & Prosthetic Devices

Competent Authority

ANSM - Site de Saint Denis

143/147, boulevard Anatole France

93285 SAINT-DENIS CEDEX

FRANCE

This certificate is issued by:	Authorized Signature:
PROTEOR SAS	
6 rue de la Redoute	
21850 Saint APOLLINAIRE	
France	

This certificate is subject to the organization maintaining their documentation in compliance with the directive stated in this certificate.

This certificate is for the exclusive use of PROTEOR USA, LLC and is provided pursuant of the European Authorized Representative agreement (Mandate) between PROTEOR SAS and PROTEOR USA, LLC. PROTEOR SAS responsibility and liability is limited to the terms and conditions of the European Medical Device Authorized Representative Mandate signed between both parties. Only PROTEOR USA, LLC and PROTEOR SAS are authorized to copy or distribute this certificate. This certificate remains valid until the expiry date has been reached or has been terminated by PROTEOR SAS.

PROTEOR USA, LLC

EU Declaration of Conformity

Version: 5.0

Date: 12/21/2021

Declaration of Conformity

for Cosmetic Foot Shell

European Medical Device Regulation MDR (EU) 2017/745

The undersigned declares that the products described in this document meet the MDR (EU) 2017/745 provisions that apply to them and the CE Mark may be affixed.

General Product Name:	See Appendix II Description/Name list	
Legal Manufacturer: (Name on Label)	PROTEOR USA, LLC 3 Morgan, Irvine, CA 92618	
Variants:	As per Appendix II (This document) - Product Listing/Schedule	
Intended Use:	Lower Limb Prosthetic Device	
MDR Classification:	Class I, in accordance with the rules set out in Annex VIII	
Notified Body:	Not Applicable for Class I	
EU Authorized	PROTEOR SAS, 6 rue de la Redoute, 21850 Saint APOLLINAIRE,	
Representative:	FRANCE	
	Self-certification by Medical Device Directive Annex IV	
MDR Assessment Route:	Article 19: EU Declaration of Conformity	
	Article 15: Person responsible for regulatory compliance	

Valery BARBOUR.

VP of Quality and Regulatory Affairs

Person responsible for Regulatory compliance

Who is the natural and legal person with responsibility for the design, manufacture, packaging and labelling before the device is placed on the market under this manufacturer's name regardless of whether these operations are carried out by the manufacturer or on his behalf by a third party.

December 21st, 2021



PROTEOR USA, LLC

EU Declaration of Conformity

Version: 5.0

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Appendix I – Applicable Standards

This present declaration is also in conformity with the following European standards and Common Specifications:

Standard/Document Name	Description	
MDR (EU) 2017/745	Regulation (EU) 2017/745 of the European Parliament and of the	
WIDK (EG) 2017/743	Council on Medical Device	
EN ISO 13485:2016	Medical Devices – Quality Management Systems – Requirements for	
EN 130 13483.2010	Regulatory Purposes	

Appendix II – Product Listing/Schedule

Part/Catalogue Number	Description/Name	Basic UDI
FTC-3M-1	Generation 3 (No Cap)	0888349FTC-3M-1V9
FTC-3M-0	Generation 3 with CAP	0888349FTC-3M-0V7
FTC-2M-1XXXX-SX	Generation 2 Sandal Toe (No Cap)	0888349FTC-2M-1XXXX-SXGP
FTC-2M-1	Generation 2 Mid Profile (No Cap)	0888349FTC-2M-1V2

Version History

Version	Compiled by	Date	Description
1.0	Jean Chen	04/10/2017	First issue
2.0	Jean Chen	04/23/2019	2 nd issue
3.0	Valery BARBOUR	12/14/2020	Change of ownership to PROTEOR USA, LLC Change of EU Rep to PROTEOR SAS Removal of: - Generation 1 Mid Profile (No Cap)
4.0	Valery BARBOUR	05/25/2021	Compliance to EU MDR 2017/745
5.0	Valery BARBOUR	12/21/2021	Removal of: - Generation 2 High Profile (No Cap)