

EC – Declaration of Conformity

Manufacturers Name:	Orfit Industries N.V.	· · · · · · ·
SRN (Single Registration	BE-MF-000007872	· · · • • • •
Number):	BE-IVIF-000007872	· · · • • • •
Number):		0 0
Manufacturers Address:	Vosveld 9a, 2110 Wijnegem, Belgium	· · • • •
Manufacturers Address.	vosvelu 5a, 2110 Wijnegelli, belgiulli	• • •
Basic UDI-DI:	5420028700244Y	
Basic ODI-DI.	54200287002441	
Norma of the Davias(s):		
Name of the Device(s):	HP PRO® SOLUTION	
Intended use:	Patient Positioning and Immobilization for Radiation Oncology	
		· · · • • • •
Product code(s):	25000/1, 25000/22, 25000/25, 25000/20, 25000/30, 25000/18, 25000/20	5 25000/1/MR
1100000000000	25000/27, 25000/16, 25000/36, 25000/35, 25000/28, 25000/29, 25000/33	
	25000/27, 25000/10, 25000/30, 25000/33, 25000/28, 25000/29, 25000/33 25000/6, 25000/58/20, 25000/39/20, 25000/17/A, 25000/17/B, 25070	
	25100/16MI+N, 25100/16Mi+N/NH, 25113/24MI+N, 25114/24Mi+N,	
		15/24MI+N/NH,
	25116/24MI+N, 25103/16MI+N/NH, 25101/16MI+N, 25101/16MI+N/NH, 25110/24MI+N,	
	25107/2MI+N/NH, 25106/2MI+N, 25106/2MI+N/NH, 25112/24MI+N, 45	
	25000/75, 25000/27/MR, 25000/76, 25000/62, 25000/67, 25000/69, 25000/69/A, 25000/69/B, 25000/7, 25000/7/20, 25000/74, 25000/85, 33129, 32138, 25100/2MI+/NH,	
	25100/2MI+, 25105/2MI+, 25105/2MI+/NH, 25000/26, 25000/70, 25000	/71, 25000/68,
	25000/61, 25000/83, 25000/84, 25000/88, 25000/81, 25000/89, 25000	/23, 25000/50,
	25000/82, 25000/87, 25000/58, 25000/34, 25000/38, 25000/53, 25000	
	2500/7/20, 25000/77, 25000/78, 25000/79, 25000/80, 25000/8	· · · · · •
	2000, 7, 20, 2000, 7, 7, 2000, 70, 2000, 70, 2000, 00, 2000, 0	
Classification:	Class I, according the rules of Annex VIII	· · · • • • •
Classification.	class I, according the rules of Annex vill	
C	Oufit is durative NLV were the fallowing groups during fauths OF labelling of th	· · · •
Conformity assessment route:	Orfit Industries N.V. uses the following procedures for the CE-labelling of th	ieir products
	according the Regulation MDR 2017/745:	0 0
		· · • • •
	Class I: EC conformity declaration according to Annex IV.	· · · • • • •
Applied norms:	ISO 13485:2016	· · • • •
	ISO 14971:2019	· · · • • • • •
	ISO 15223-1:2021	
	ISO 10993-5:2009	· · · • • • • •
	ISO 10993-10:2010	· · · · · · · · · · · ·

This declaration of conformity is issued under the sole responsibility of Orfit Industries N.V. We hereby declare that the medical device(s) specified above meet the provisions of the Regulation (EU) MDR 2017/745 for medical devices. This declaration is supported by the Quality System approval ISO 13485:2016 issued by LRQA. All supporting documentation is retained at the premises of the manufacturer.

Eddy Marivoet, Wijnegem, 29 May 2024

Quality Assurance & Regulatory Affairs Manager



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