

Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

MANUFACTURER		
Name of Company	Address	SRN
Orthomerica Products Inc.	6333 North Orange Blossom Trail Orlando, FL 32810 USA	US-MF-000009882

AUTHORIZED REPRESENTATIVE			
Name of Company	Address	SRN	Phone/email
Emergo Europe	Prinsessegracht 20 2514 AP The Hague The Netherlands	NL-AR-000000116	+31.70.345.8570 EmergoEurope@ul.com

PRODUCT IDENTIFICATION	
Product Name	Code / Catalog Numbers
Newport III Orthosis	3640, 3641, 6342, 6343, 3645, 3646, 3647, 3648, 3640.40, 3641.41, 3642.42, 3643.43
Intended Purpose	Basic UDI-DI
Post-operative hip revision patients Primary arthroplasty patients at risk to dislocate Patients needing stability after dislocation Inoperable patients requiring hip stabilization Patients who can benefit from a hip orthosis to reinforce hip precautions	Being Assigned UDI 00195003004350 - 00195003004381 00195003004411 - 00195003004442 00195003004473 - 00195003004480 00195003004497 - 00195003004503 00195003004367 - 00195003004398 00195003004428 - 00195003004459

RISK CLASS FOR DEVICES		
Device Classification		Common Specifications / Standards
Class:	1	EN ISO 13485:2016 EN ISO 15223-1
Rule:	1	

Orthomerica declares that the above-mentioned products meet the provision of the following EU legislation:

- Medical Devices Regulation (EU) 2017/745

COMPANY REPRESENTATIVE: Najiba Katir

TITLE: Regulatory Compliance

SIGNATURE: *Najiba Katir*

PLACE: Orlando

DATE: 14/07/2021