

DECLARATION OF CONFORMITY (DOC-031)

Nea International bv declares that this EU declaration of conformity is issued under the sole responsibility of the manufacturer.

The devices covered by this declaration are in accordance with the requirements of Regulation (EU) 2017/745 of the European parliament and of the Council of 5 April 2017.

In accordance with the rules in Annex VIII chapter III, the devices are classified in MDR class 1.

Push® CARE		Push® MED	
Art.no.	Description	Art.no.	Description
1.10.1	Wrist Brace	2.10.1	Wrist Brace
1.20.1	Ankle Brace	2.10.2	Wrist Brace Splint
1.30.2	Knee Brace	2.20.1	Ankle Brace
1.40.1	Back Brace	2.20.2	Ankle Brace Aequi® Flex
1.60.1	Neck Brace 8 cm	2.20.3	Ankle Brace Aequi® Flex
1.60.2	Neck Brace 10 cm	2.30.1	Knee Brace
		2.30.2	Patella Brace
		2.40.2	Back Brace
		2.50.2	Shoulder Brace Plus
		2.50.3	Shoulder Brace
		2.60.1	Neck Brace 8 cm
		2.60.2	Neck Brace 10 cm
		2.70.1	Elbow Brace Epi
		2.70.2	Elbow Brace
Push® ORTHO		Push® Sports	
Art.no.	Description	Art.no.	Description
3.10.1	Thumb Brace CMC	4.10.1	Wrist Brace
3.10.2	Thumb Brace CMC/ MetaGrip®	4.10.2	Wrist Support
3.20.1	Ankle Brace Aequi®	4.10.3	Thumb Brace
3.20.2	Ankle Brace Aequi® Junior	4.20.1	Ankle Brace Kicx®
3.20.3	Ankle Foot Orthosis AFO	4.20.2	Ankle Brace 8
		4.30.1	Knee Brace
		4.30.2	Patella Brace
		4.70.1	Elbow Brace

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