



EU DECLARATION OF CONFORMITY

Manufacturer Name:	POD Active Pty Ltd
Manufacturer Address:	Level 3, 115 Myers Street, Geelong, Victoria 3220 AUSTRALIA
Authorised Representative Name:	WellKang Ltd.
Authorised Representative Address:	Enterprise Hub, NW Business Complex 1 Beraghmore Rd., Derry, BT48 8SE NORTHERN IRELAND
Manufacturer's SRN (Single Registration Number):	AU-MF-000003989
Basic UDI-DI:	K8 3.0 (Singles) B-93440060K83SHX K8 3.0 (Pairs) B-93440060K83PHR
Name of the Device(s):	POD K8 3.0 Knee Brace
Product Codes:	POD K8 3.0 Knee Brace: K83004* (LEFT) K83005* (RIGHT) K83006* (PAIR)
Intended Purpose:	The device is intended for external support, stabilization and protection of the knee.
Classification:	Class 1 (non-sterile; no measuring function).
Conformity Assessment Route:	POD Active Pty Ltd uses the following procedure for the CE labelling of their products according to the Regulation MDR 2017 / 745: Class 1: EC conformity declaration according to Annex IV.
Validity:	This declaration of conformity is valid for one (1) year from the date of issue or when the technical documentation is revised.

This declaration of conformity is issued under the sole responsibility of POD Active Pty Ltd.

We hereby declare that the medical devices specified above meet the provisions of the Regulation (EU) MDR 2017 / 745 for medical devices.

All supporting documentation is retained at the premises of the Manufacturer.

Declared on Friday, 3 April 2026 by:

Signature:

Geoff Maloney
Director, POD Active Pty Ltd

Place and date of issue (DD.MM.YYYY):

Level 3, 115 Myers Street
Geelong, Victoria 3220
AUSTRALIA

Friday, 3 April 2026



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