

Declaration of Conformity

EMDN CND Code: Y061209

Issue Date: May 26, 2022

EMDN CND Description: Knee orthoses

Valid Date: Dec. 31, 2025

This declaration of conformity is issued under the sole responsibility of the manufacturer:

OPPO Medical Inc., address at: Seattle City Center, 1420 Fifth Ave., Ste. 220080, Seattle, WA98101, USA

Single registration number (SRN): US-MF-000005202

Authorised Representative: MT Promedt Consulting GmbH, address at: Altenhofstrasse 80,

D-66386 St. Ingbert Germany Single registration number (SRN): DE-AR-000000085

The basic UDI-DI: 8402706CL113L2

We herewith declare that the products as referred to in Attachment I are in conformity with the requirements set out in the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL.

Conformity Assessment Procedure: MDR (EU) 2017/745 Art 52. (7), ANNEX II (Technical documentation) & ANNEX III (Technical documentation on post-market surveillance)

Risk Classification of the Product: Class I Rule: MDR (EU) 2017/745, Annex VIII, Section 4, 4.1, Rule 1"All non-invasive devices are classified as class I."

Applied Standards: EN ISO13485:2016, EN1041:2008, EN ISO14971:2012, EN ISO 15223-1:2016, EN ISO 10993-1:2009, EN ISO10993-5:2009, EN ISO 22523:2006, EN 62366:2008, ISO 10993-10:2021, ISO 9001:2015

Intended purpose: This is a brace for pain relief from osteoarthritis and degenerative meniscal tears through unicompartmental unloading of the knee.

May 26, 2022/ Seattle

Date/Place

Jackson Chiang President



Attachment 1

EMDN CND Code: Y061209

EMDN CND Description: Knee orthoses

Reference number	Product Trade name
2935	OppO genu X