



## Declaration of Conformity

Product Designation: Support, Arm  
UMDNS- Code: 13-862

Issue Date: Oct. 12, 2018

Valid Date: Oct. 12, 2022

Model No's: Elbow - 1080, 1085, 1086, 1187, 1287, 1486, 1489, 2080, 2085, 2185, 2280,  
2285, 2480, 2580, 2587, 2589, 2685, 2686, 2986, 2987, 2988, 3086,  
4080, 4486, 2385  
Wrist - 1081, 1083, 1084, 1181, 1188, 1281, 2081, 2083, 2084, 2181, 2184,  
2281, 2481, 2583, 2586, 2681, 2980, 2383, 2384

We herewith declare that the products listed above are in compliance with the requirements set out in the Council Directive on the harmonization of the Laws of the Member States concerning Medical Device Directive 93/42/EEC.

Conformity Assessment Procedure: MDD 93/42/EEC Annex VII

Classification of the Product: Class I Rule: MDD 93/42/EEC Annex IX Rule 1

Applied Harmonized Standards: EN ISO9001:2015, EN ISO13485:2016, EN1041:2008,  
EN ISO14971:2012, EN ISO 15223-1:2016,  
EN ISO10993-5:2009, EN ISO 22523:2006

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

Oct. 12<sup>th</sup>, 2018

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Date

  
.....  
Mr. Jackson Chiang  
President

EU Authorized Representative: **MT Promedt Consulting GmbH**  
**Altenhofstrasse 80**  
**D-66386 St. Ingbert**  
**Germany**

MT Promedt Consulting GmbH (MTPC) has taken over the function and responsibilities of an European Authorized Representative of OPPO Medical Inc. in accordance to the requirements of the MDD 93/42/EEC.

The manufacturer has provided MTPC with the Technical Documentation.

The necessary notification of the above mentioned products has been performed. The products are registered under the Reg.-Nr.: DE/CA70/40838/0048123