EU-Declaration of Conformity

Manufacturers Name: Ki Mobility 5201 Woodward Dr. Manufacturers Address: Stevens Point, WI 54481 SRN (Single Registration Number): US-MF-000012633 Authorized Representative Name (if **Etac Supply Center AB** applicable): Authorized Representative Address Langgatan 12 (if applicable): 33233 Anderstorp, Sweden Authorized Representative SRN: SE-AR-000001601 Basic UDI-DI: 0850013379SEATINGAG UDI-DI: 00850013379330 Name of the Device(s): Axiom PXP GMDN product code: 11100 Device Classification: Class I, Rule 1 Intended Purpose: A wheelchair component is a device intended for medical purposes that is generally sold as an integral part of a wheelchair but may also be sold separately as a replacement part. Notified Body name: Not applicable Notified Body Address: Not applicable Notified Body Identification Not applicable number: Conformity assessment route: Ki Mobility uses the following procedures for the CE-labeling of their products according to the Regulation MDR 2017/745: Class 1: EU conformity declaration according to annex VIII This declaration of conformity is issued under the sole responsibility of Ki Mobility. We hereby declare that the medical device(s) specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices. This declaration is supported by the Quality System in conformance to ISO 13485:2016 and on assessment of technical documentation. All supporting documentation is retained at the premises of Ki Mobility. Name: Douglas Munsey Title: President Signature:

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Date (YYYY-MM-DD) of issue: