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Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

| MANUFACTURER | | |
|-----------------------------|--|-----------------|
| Name of Company | Address | SRN |
| Biodex Medical Systems, Inc | 49 Natcon Dr Shirley, NY 11967, USA | US-MF-000028834 |

| AUTHORIZED REPRESENTATIVE | | | |
|---------------------------|---|-----------------|--|
| Name of Company | Address | SRN | Phone/email |
| Emergo Europe | Westervoortsedijk 60 6827 AT Arnhem The Netherlands | NL-AR-000000116 | +31.70.345.8570 EmergoEurope@ul.com |

| PRODUCT IDENTIFICATION | |
|--|-----------------------|
| Product Name | Code / Catalog Number |
| System 4 MVP | 852-000 |
| Contains MODEL NUMBER(s) <u>852-000 System 4 MVP includes:</u> 850-000-E300 Dell CPU 840-109-J800 Software C07-015 Biodex Advantage Software Manual C08-051 Biodex Multi Joint System Installation Instructions C08-246 System 4 Multi Joint System Poster C14113 HP Office Jet printer C12940 Touch Screen Monitor with Stand 900-860 Power head / Gimbal 830-000-K904 Limb Support Kit: 830-154 Arm / Leg Support 830-155 Foot Rest Tube 820-153 Small Tee Cap Screws (14 pieces) 945-300-M322 Levelers (6 pieces) 840-000-K900 Quick Set Attachments Kit: 830-157 Elbow / Shoulder Attachment 830-158 Wrist Attachment 830-174 Knee Attachment, Left 830-175 Knee Attachment, Right 830-321 Shoulder Attachment 830-332 Ankle Attachment 830-350 Calibration Weight 830-550 Hamstring 830-269 Work Sim Tools 830-260 Anti-Shear, Left 830-271 Anti-Shear, Right 830-240 Attachment Rack 832-135-A000 MVP Single Chair Assembly 852-145-A000 Tee Base | |

Biodex Medical Systems, Inc.

49 Natcon Drive, Shirley, New York, 11967-4704, Tel:800-224-6339 (Int'l 631-924-9000), Fax:631-924-9241, Email: info@biodexrehab.com, www.biodex.com

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|--|---------------------|
| 835-210-A000 CDS Cart | |
| Intended Purpose | Basic UDI-DI |
| System 4 is intended to identify, treat, and document the physical impairments that cause functional limitations typical of sports injuries, orthopedics, pediatric medicine, and neurorehabilitation. | 07181752014S |

| RISK CLASS FOR DEVICES | | |
|-------------------------------|-----|--|
| Device Classification | | Common Specifications / Standards |
| Class: | Ila | <ul style="list-style-type: none"> • ISO 13485:2016 Medical devices -- Quality management systems -- Requirements for regulatory purposes • ISO 14971: 2019 Medical devices -- Application of risk management to medical devices • ISO 15223-1:2021 Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements • ISO 7000:2019 Graphical symbols for use on equipment -- Registered symbols • ISO 10993-1:2018 Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process • IEC 62366-1:2015+AMD1:2020 CSV Consolidated version Medical devices - Part 1: Application of usability engineering to medical devices • IEC 62304:2006+AMD1:2015 CSV Consolidated version Medical device software - Software life cycle processes • IEC 60601-1 Ed. 3.2 en:2020 Medical Electrical Equipment - Part 1: General Requirements For Basic Safety And Essential Performance • ANSI AAMI ES60601-1:2005/(R)2012 & A1:2012, C1:2009/(R)2012 & A2:2010/(R)2012 (Cons. Text) [Incl. AMD2:2021] Medical electrical equipment – Part 1: General requirements for basic safety and essential performance (IEC 60601-1:2005, MOD) [Including Amendment 2 (2021)] • CAN/CSA-C22.2 No. 60601-1:2014 • IEC 60601-1-2:2014+AMD1:2020 CSV Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests |
| Rule: | 9 | |

| NOTIFIED BODY | | | |
|------------------------|------------------|--|---------------------------------|
| Name of Company | ID Number | Conformity Assessment Procedure | Certificate Reference(s) |
| Intertek Semko AB | 0413 | Annex II MDD 93/42/EEC Council Directive (excluding Section 4) | 41313009-03 |

QUALITY SYSTEM REGISTRAR

Biodex Medical Systems, Inc.

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| Name of Company | Certified Quality Management System | Certificate Reference(s) |
|-------------------|-------------------------------------|--------------------------|
| Intertek Services | ISO 13485:2016 | 0084059 |

Biodex Medical Systems declares that the above-mentioned products meet the provision of the following EU legislation:

- Medical Devices Regulation (EU) 2017/745
- Directive 2011/65/EU (RoHS 2)
- Directive 2015/863/EU (RoHS 3)
- Directive 2017/2102/EU
- Directive 2006/42/EC (Machinery Directive)

COMPANY REPRESENTATIVE: Amaris Ajamil, PhD, RAC

TITLE: Vice President, Quality and Regulatory Affairs, Salona Global

SIGNATURE: *Amaris Ajamil*
Amaris Ajamil (Oct 9, 2023 10:27 EDT)

PLACE: Shirley, New York, USA
No. 111C Rev. Q

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





S4 MVP 111C REV. Q

Final Audit Report

2023-10-09

| | |
|-----------------|--|
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