



## EU/UK/AU/CH Declaration of Conformity DC0006 Rev. 13

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

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| <b>Manufacturer Information:</b>                                          | TIDI Products, LLC<br>570 Enterprise Drive<br>Neenah, WI 54956<br>USA<br>SRN: US-MF-000012287                                                                                                                                                                                                                                                                                                                                         |
| <b>Person Responsible for Regulatory Compliance (PRRC):</b>               | Chris Rahn, VP Quality & Regulatory                                                                                                                                                                                                                                                                                                                                                                                                   |
| <b>European Union (EU) Authorized Representative Contact Information:</b> | MDSS GmbH<br>Schiffgraben 41<br>30175 Hannover<br>Germany<br>Phone: (+49) 511 6262 8630<br>SRN: DE-AR-000005430                                                                                                                                                                                                                                                                                                                       |
| <b>United Kingdom (UK) Authorized Rep Contact Information:</b>            | MDSS-UK RP Ltd.<br>Parkway House, Palatine Rd, Northenden, Wythenshawe,<br>Manchester M22 4DB<br>United Kingdom                                                                                                                                                                                                                                                                                                                       |
| <b>Swiss (CH) Authorized Representative Contact Information:</b>          | NA                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| <b>Product identification:</b>                                            | Zero-Gravity® Radiation Protection System                                                                                                                                                                                                                                                                                                                                                                                             |
| <b>Technical File No.:</b>                                                | TF-0023 Zero-Gravity® Radiation Protection System Personal Protective Equipment (PPE)                                                                                                                                                                                                                                                                                                                                                 |
| <b>Product Model Numbers:</b>                                             | See following page(s) for model numbers, and descriptions.                                                                                                                                                                                                                                                                                                                                                                            |
| <b>EU Legislation and Conformity Assessment Procedure:</b>                | <ol style="list-style-type: none"><li>1. EU Type Examination procedures under the requirements with Regulation (EU) 2016/425 relating to PPE Annex V (Module B) and meets the relevant health and safety requirements specified in Annex II.</li><li>2. Conformity to Type based on Quality Assurance of the Production Process under the requirements with Regulation (EU) 2016/425 relating to PPE Annex VIII (Module D).</li></ol> |
| <b>UK Legislation and Conformity Assessment Procedure:</b>                | UKCA Type-examination (module B) set out in Annex V, followed by conformity to type based on quality assurance of the production process (module D) set out in Annex VIII of Regulation 2016/425 on personal protective equipment as brought into UK law and amended.                                                                                                                                                                 |
| <b>Australia (AU) Legislation and Conformity Assessment Procedure:</b>    | Clause(s) 6.6 of Schedule 3 Part 6 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002.                                                                                                                                                                                                                                                                                                                            |

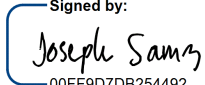


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| <b>CH Legislation and Conformity Assessment Procedure:</b> | 1. EU Type Examination procedures under the requirements with Regulation (EU) 2016/425 relating to PPE Annex V (Module B) and meets the relevant health and safety requirements specified in Annex II.<br>2. Conformity to Type based on Quality Assurance of the Production Process under the requirements with Regulation (EU) 2016/425 relating to PPE Annex VIII (Module D). |                                           |
| <b>Intended purpose:</b>                                   | Zero-Gravity Radiation Protection System: A protective shield for use during medical procedures requiring fluoroscopy, intended to protect users from radiation exposure and orthopedic strain.                                                                                                                                                                                  |                                           |
| <b>EMDN Code:</b>                                          | NA                                                                                                                                                                                                                                                                                                                                                                               |                                           |
| <b>Basic UDI-DI:</b>                                       | NA                                                                                                                                                                                                                                                                                                                                                                               |                                           |
| <b>PPE Category in EU/CH:</b>                              | Category III                                                                                                                                                                                                                                                                                                                                                                     |                                           |
| <b>PPE Category in UK:</b>                                 | Category III                                                                                                                                                                                                                                                                                                                                                                     |                                           |
| <b>Device Classification/ Rule in Australia (AU):</b>      | Class 1                                                                                                                                                                                                                                                                                                                                                                          | Rule 2.1                                  |
| <b>Australian Client ID No.</b>                            | TIDI's AU Client ID No.: 49283                                                                                                                                                                                                                                                                                                                                                   |                                           |
| <b>Reference to Common Specifications:</b>                 | N/A                                                                                                                                                                                                                                                                                                                                                                              |                                           |
| <b>EC Certificate:</b>                                     | Number:<br>CE 716486<br>CE 716567                                                                                                                                                                                                                                                                                                                                                | Issue Date:<br>15 Oct 2019<br>15 Oct 2019 |
| <b>UKCA Type -Certificate:</b>                             | Number:<br>750691<br>750690                                                                                                                                                                                                                                                                                                                                                      | Issue Date:<br>18 Nov 2021<br>18 Nov 2021 |
| <b>Quality Management Certificate - ISO 13485</b>          | Number:<br>FM 536366                                                                                                                                                                                                                                                                                                                                                             | Effective Date:<br>29 May 2023            |
| <b>MDSAP Certificate</b>                                   | Number:<br>MDSAP 703786                                                                                                                                                                                                                                                                                                                                                          | Effective Date:<br>29 May 2023            |
| <b>Notified Body for QMS:</b>                              | BSI Group The Netherlands B.V.<br>Say Building<br>John M. Keynesplein 9<br>1066 EP Amsterdam, The Netherlands<br>Notified Body Number: 2797                                                                                                                                                                                                                                      |                                           |
| <b>Notified Body for EU Conformity Assessment:</b>         | BSI Group The Netherlands B.V.<br>Say Building<br>John M. Keynesplein 9<br>1066 EP Amsterdam, The Netherlands<br>Notified Body Number: 2797                                                                                                                                                                                                                                      |                                           |



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| <b>Approved Body for UK<br/>Conformity Assessment:</b>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | BSI UK<br>Kitemark Court, Davy Avenue,<br>Knowlhill, Milton Keynes,<br>United Kingdom MK5 8PP<br>Approved Body Number: 0086 |                   |
| <b>For European Union (PPE products)</b><br><p>This declaration of conformity is issued under the sole responsibility of TIDI Products, LLC. The undersigned hereby declares, on behalf of TIDI Products, LLC, that the personal protective equipment complies with the applicable health and safety requirements to Annex II of the European Personal Protective Equipment Regulation 2016/425, and its relevant transposition into national laws of the member states into which the devices are placed and also self-declares compliance in part to the Machinery Directive 2006/42/EC as it applies to the overhead-body-shield-support functionality of the Zero-Gravity. We explicitly designate MDSS GmbH to act as our sole Authorized Representative in the European Union for the above indicated products.</p> |                                                                                                                             |                   |
| <b>For United Kingdom (PPE products):</b><br><p>This declaration of conformity is issued under the sole responsibility of TIDI Products, LLC. The undersigned hereby declares, on behalf of TIDI Products, LLC, that the personal protective equipment complies with Regulation 2016/425 on personal protective equipment as brought into UK law and amended also self-declares compliance in part to the Machinery Directive (2006/42/EEC) implemented in the UK by the Supply of Machinery (Safety) (Amendment) Regulations 2011 as it applies to the overhead-body-shield-support functionality of the Zero-Gravity Systems. We explicitly designate MDSS-UK RP Ltd. to act as our sole Authorized Representative in the UK under PPE Regulation 2016/425 for the above indicated products.</p>                        |                                                                                                                             |                   |
| <b>For Australia (Non-Sterile devices):</b><br><p>This declaration of conformity is issued under the sole responsibility of TIDI Products, LLC. The undersigned hereby declares, on behalf of TIDI Products, LLC, that each kind of medical device to which the technical documentation applies complies with the applicable provisions of the essential principles, and the classification rules before being supplied under clause(s) 6.6 of Schedule 3 Part 6 to the Australian Therapeutic Goods Administration Medical Device Regulation 2002.</p>                                                                                                                                                                                                                                                                   |                                                                                                                             |                   |
| <b>For Switzerland (PPE products):</b><br><p>This declaration of conformity is issued under the sole responsibility of TIDI Products, LLC. The undersigned hereby declares, on behalf of TIDI Products, LLC, that the personal protective equipment complies with the applicable health and safety requirements of the European Personal Protective Equipment Regulation 2016/425, and its relevant transposition into national laws of the member states into which the devices are placed and also self-declares compliance in part to the Machinery Directive 2006/42/EC as it applies to the overhead-body-shield-support functionality of the Zero-Gravity. We explicitly designate MDSS GmbH to act as our sole Authorized Representative in the European Union for the above indicated products.</p>               |                                                                                                                             |                   |
| <b>Signed for on behalf of TIDI Products LLC, in Neenah, WI 54956</b>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |                                                                                                                             |                   |
| <b>Name of TIDI<br/>Representative:</b><br>Joe Samz                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | <b>Title, Function</b>                                                                                                      | <b>Date</b>       |
| <b>Approval:</b><br><br>Signed by:<br>00FF9D7DB254492...                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | <b>Approval:</b><br>Sr. Compliance Manager                                                                                  | September 8, 2025 |



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| Product Model Number and Description to which this declaration applies. |                                                                                                                                     |
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| Model Numbers                                                           | Description                                                                                                                         |
| ZGCM-48                                                                 | Zero-Gravity™ Radiation Protection System Monorail 48                                                                               |
| ZGCM-66                                                                 | Zero-Gravity™ Radiation Protection System Monorail 66                                                                               |
| ZGHSA                                                                   | Zero-Gravity™ Radiation Protection System Hinged Swing Arm                                                                          |
| ZGCM-HSA                                                                | Zero-Gravity™ Radiation Protection System Monorail Hinged Swing Arm                                                                 |
| ZGM-6-5H                                                                | Zero-Gravity™ Radiation Protection System Floor Unit                                                                                |
| ZGCMRS                                                                  | Zero-Gravity™ Monorail Lead Acrylic Shield                                                                                          |
| ZG48                                                                    | Zero-Gravity™ Radiation Protection System - Body Shield with Extension Rail                                                         |
| ZGHH-CMHSA                                                              | ZGM-6-5H Upgrade to ZGHH-CMHSA Zero-Gravity™ Radiation Protection System Upgrade from Floor to Hybrid Monorail Design               |
| ZGHH-HSA                                                                | ZGM-6-5H Upgrade to ZGHH-HSA Zero-Gravity™ Radiation Protection System Upgrade from Floor to Hinged Swing Arm Design                |
| ZGHH-66-CMHSA                                                           | ZGCM-48/ZGCM-66 Upgrade to ZGHH-66-CMHSA Zero-Gravity™ Radiation Protection System Upgrade Monorail 48/66 to Hybrid Monorail Design |
| ZGHH-CM48                                                               | ZGM-6-5H Upgrade to ZGHH-CM48 Zero-Gravity™ Radiation Protection System Upgrade from Floor to 48" Hybrid Monorail Design            |
| ZGAV-XS                                                                 | Zero-Gravity™ Radiation Protection System Extra Small Vest                                                                          |
| ZGAV-S                                                                  | Zero-Gravity™ Radiation Protection System Small Vest                                                                                |
| ZGAV-M                                                                  | Zero-Gravity™ Radiation Protection System Medium Vest                                                                               |
| ZGAV-L                                                                  | Zero-Gravity™ Radiation Protection System Large Vest                                                                                |
| ZGAV-XL                                                                 | Zero-Gravity™ Radiation Protection System Extra-Large Vest                                                                          |
| ZGAV-2XL                                                                | Zero-Gravity™ Radiation Protection System Double Extra-Large Vest                                                                   |
| ZGAV-3XL                                                                | Zero-Gravity™ Radiation Protection System Triple Extra-Large Vest                                                                   |

| Glossary of Global Medical Device Nomenclature (GMDN) Terms (for AU only) |                                            |
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| GMDN                                                                      | Term                                       |
| 38373                                                                     | Radiation shielding panel, portable/mobile |

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The products to which this declaration relates are developed and manufactured in conformity with the following standard(s).

| Number                | Description                                                                                                                                                                                  | Year/Revision |
|-----------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------|
| <b>DIN EN 61331-1</b> | Protective devices against diagnostic medical X-radiation – Part 1: Determination of attenuation properties of materials (partial)                                                           | 2016          |
| <b>DIN EN 61331-3</b> | Protective devices against diagnostic medical X-radiation - Part 3: Protective clothing, eyewear and protective patient shields (partial)                                                    | 2016          |
| <b>EN 166</b>         | Personal Eye-Protection - Specifications (partial)                                                                                                                                           | 2001          |
| <b>ANSI Z87.1</b>     | Eye & Face Protection Standards (partial)                                                                                                                                                    | 2020          |
| <b>IEC 61331-1</b>    | Protective devices against diagnostic medical X-radiation – Part 1: Determination of attenuation properties of materials (partial)                                                           | 2014          |
| <b>IEC 61331-2</b>    | Protective devices against diagnostic medical X-radiation – Part 2: Translucent protective plates (partial)                                                                                  | 2014          |
| <b>IEC 61331-3</b>    | Protective devices against diagnostic medical X-radiation – Part 3: Protective clothing, eyewear and protective patient shields (partial)                                                    | 2014          |
| <b>IEC 60601-1</b>    | Medical electrical equipment – Part 1: General requirements for basic safety and essential performance (partial)                                                                             | 2020          |
| <b>IEC 60601-1-3</b>  | Medical electrical equipment – Part 1-3: General requirements for basic safety and essential performance – Collateral standard: Radiation protection in diagnostic X-ray equipment (partial) | 2021          |
| <b>IEC 62366-1</b>    | Medical devices – Part 1: Application of usability engineering to medical devices                                                                                                            | 2020          |
| <b>ISO 14971</b>      | Medical devices – Application of risk management to medical devices                                                                                                                          | 2019          |
| <b>ISO 13485</b>      | Medical devices – Quality management systems – Requirements for regulatory purposes                                                                                                          | 2016          |
| <b>ISO 780</b>        | Packaging – Distribution packaging – Graphical symbols for handling and storage of packages                                                                                                  | 2015          |
| <b>ISPM 15</b>        | International Standard for Phytosanitary Measures 15                                                                                                                                         | 2018          |
| <b>ASTM D5445</b>     | Standard practice for pictorial markings for handling of goods                                                                                                                               | 2021          |
| <b>EN 170</b>         | Personal Eye Protection - Ultraviolet Filters - Transmittance requirements and recommended use (partial)                                                                                     | 2002          |
| <b>DIN EN 14238</b>   | Cranes - Manually controlled load manipulating devices (partial)                                                                                                                             | 2010          |
| <b>EN ISO 12100</b>   | Safety Of Machinery - General principles for design - Risk assessment and risk reduction                                                                                                     | 2010          |
| <b>ISO 10993-1</b>    | Biological evaluation of medical devices. Evaluation and testing within a risk management process.                                                                                           | 2018          |