



EU/UK/AU/CH Declaration of Conformity DC0041 Rev. 15

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

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| Manufacturer Information: | TIDI Products, LLC 570 Enterprise Drive Neenah, WI 54956 USA SRN: US-MF-000012287 |
| Person Responsible for Regulatory Compliance (PRRC): | Chris Rahn, VP Quality & Regulatory |
| European Union (EU) Authorized Representative Contact Information: | MDSS GmbH Schiffgraben 41 30175 Hannover Germany Phone: (+49) 511 6262 8630 SRN: DE-AR-000005430 |
| United Kingdom (UK) Responsible Person Contact Information: | Emergo Consulting (UK) Limited c/o Cr360 UL International Compass House, Vision Park Histon Cambridge CB24 9BZ, United Kingdom Phone: +44(0) 1223 772 671 |
| Swiss (CH) Authorized Representative Contact Information: | MDSS CH GmbH Laurenzenvorstadt 61 5000 Aarau Switzerland CHRN: CHRN-AR-20001035. |
| Product identification: | Non-Sterile Equipment Covers & Non-Sterile Urology Drain Bags |
| Technical File No.: | TF-0022: TIDI Products Medical Barriers Family |
| Product Model Numbers: | See following page(s) for model numbers, GMDNs, descriptions and photos, where appropriate |
| EU Legislation and Conformity Assessment Procedure: | Annex II & Annex III: Technical documentation including PMS of Regulation (EU) 2017/ 745 of the European Parliament and the Council of the European Union. |
| UK Legislation and Conformity Assessment Procedure: | UK Medical Devices Regulation 2002 (SI 618) as subsequently amended by the EU Exit Regulations of 2019 (SI 791) and 2020 (SI 1478). Conformity to Part II of the UK MDR 2002, Annex VII (as modified by Part II of Schedule 2A to the UK MDR 2002). |
| Australia (AU) Legislation and Conformity Assessment Procedure: | Clause(s) 6.6 of Schedule 3 Part 6 to the Australian Therapeutic Goods (Medical Devices) Regulations 2002 |
| CH Legislation and Conformity Assessment Procedure: | Annex II & Annex III: Technical documentation including PMS of Regulation (EU) 2017/ 745 of the European Parliament and the Council of the European Union. |
| Intended purpose: | The medical barriers are intended to protect the equipment it covers. |



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| | The Urology drain bags are intended for use as accessories to an urological table for fluid containment. | |
| EMDN Code: | EMDN code for medical barriers and urology drain bags: <ul style="list-style-type: none"> • T030102 – Cover Sheaths, Instruments and Equipment • A060303 – Urine collection systems and bags, single use | |
| Basic UDI-DI: | Basic UDI-DI for medical barriers and urology drain bags: <ul style="list-style-type: none"> • General - non-sterile 0618125TF-0022-BWY • Urology Drain Bags 0618125TF-0022-FX8 | |
| Device Classification/ Rule in EU/CH: | Risk Class I | Rule 1 |
| Device Classification/ Rule in UK: | Risk Class I | Rule 1 |
| Device Classification/ Rule in Australia (AU): | Risk Class I | Rule 2.1 |
| Australian Client ID No. | TIDI's AU Client ID No.: 49283 | |
| Reference to Common Specifications: | NA | |
| EC Certificate: | EC Number: N/A Self-Declared | Issue Date: N/A Self-Declared |
| UKCA Certificate: | EC Number: N/A Self-Declared | Issue Date: N/A Self-Declared |
| Quality Management Certificate - ISO 13485 | Number: FM 536366 | Effective Date: 29 May 2023 |
| MDSAP Certificate | Number: MDSAP 703786 | Effective Date: 29 May 2023 |
| Notified Body for QMS: | BSI Group The Netherlands B.V. Say Building John M. Keynesplein 9 1066 EP Amsterdam, The Netherlands NB No. 2797 | |
| Notified Body for EU Conformity Assessment: (if applicable) | NA | |
| Approved Body for UK Conformity Assessment: (if applicable) | NA | |
| <p>:For European Union (Non-sterile products)</p> <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 medical devices referenced in this declaration comply with the European Medical Devices Regulation; MDR (EU) 2017/745. We explicitly</p> | | |



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designate MDSS GmbH to act as our sole Authorized Representative in the European Union for the above indicated products.

For United Kingdom:

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 medical devices referenced in this declaration comply with the UK Medical Devices Regulation 2002 (SI 618) as subsequently amended by the EU Exit Regulations of 2019 (SI 791) and 2020 (SI 1478). We explicitly designate Emergo Consulting (UK) Limited to act as our sole Responsible Person in the UK for the above indicated products.

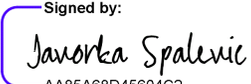
For Australia (Non-sterile products):

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.

For Switzerland:

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 medical devices referenced in this declaration comply with the applicable parts of the MedDO. These are class I medical devices that declare conformity to Regulation (EU) 2017/745 (MDR). We explicitly designate MDSS CH GmbH to act as our sole Authorized Representative in Switzerland for the above indicated products.

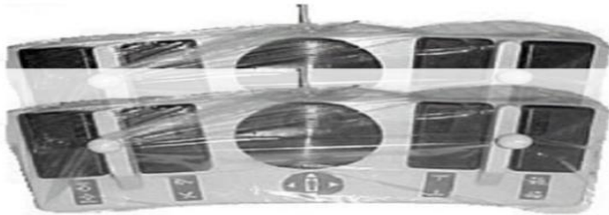
Signed on behalf of TIDI Products LLC, in Neenah, WI. 54956

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| Name of TIDI Representative; Javorka Spalevic | Title, Function | Date |
| Approval:  <small>Signed by: AA85A68D45604C2...</small> | Approval: Regulatory Affairs Compliance Manager | January 28, 2026 |

| Product Name, Model Number, GMDN, and UDI-DI to which this declaration applies | | | |
|--|--|-------|----------------|
| Model Number (REF) | Product Name | GMDN | UDI-DI |
| C000-0492 (22918) | Accessory Cover | 12535 | 00618125155699 |
| CFI-956 N/S | X-Ray Cassette Cover | 12535 | 00618125102495 |
| CFI-959 N/S | X-Ray Cassette Cover | 12535 | 00618125140848 |
| 5406 | Footswitch Cover | 12535 | 00618125140046 |
| 5421 | Wide Footswitch Cover | 12535 | 00618125139880 |
| 00-900410-02-OEC, E9100AD (20764) | Gray Footswitch Cover All OEC® System | 12535 | 00618125152193 |
| 00-900941-01-OEC, E9100BC (20783) | OEC UroView® 2600, Disposable Wide Footswitch Cover | 12535 | 00618125112371 |

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| Product Name, Model Number, GMDN, and UDI-DI to which this declaration applies | | | |
|--|--|-------|----------------|
| Model Number (REF) | Product Name | GMDN | UDI-DI |
| 00-900940-01-OEC, E9100AZ (20780) | All OEC UroView®/Urofast Systems Disposable Collection Bag 5 gallon/19 liter | 61677 | 00618125184019 |
| 00-902099-01, E9100BJ (21185) | OEC UroView® 2800, Disposable Wide Footswitch Cover | 12535 | 00618125152230 |
| 5300 N/S | Uro-Catcher Bag | 61677 | 00618125141111 |
| 5313 N/S | Uro-Catcher Drain Bag | 61677 | 00618125141128 |
| 5442 N/S | Urology Drain Bag | 61677 | 00618125140985 |
| 5477 | 6" Extender Hose with Adapter | 61677 | 00618125142989 |
| 5010 | Contain Drain Urology Drain Collection Bag | 61677 | 00618125139989 |
| C000-0612A (25340) | Urology Drain Collection Unit | 61677 | 00618125155705 |
| C000-1111 (32586) | Urology Drain Bag | 61677 | 00618125155712 |
| 3075-8000-00000-000 (32065) | G2 Drain Bag | 61677 | 00618125167975 |
| 31780 N/S (BF459) | Urology Drain Bag | 61677 | 00618125192274 |

| Product Name | Photo (if appropriate) |
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| Wide Footswitch Cover (5421) |  |
| All OEC UroView®/Urofast Systems Disposable Collection Bag 5 gallon/19 liter (20780) |  |

| Glossary of Global Medical Device Nomenclature (GMDN) Terms | |
|---|--|
| GMDN | Term |
| 12535 | Medical equipment/instrument drape, single-use |
| 61677 | Urological Fluid Funnel |