



# EU Declaration of Conformity

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|---|---|
| <b>Manufacturer</b>                     | <b>Etac Immedia A/S</b><br>Parallevej 3<br>DK-8751 Gedved<br>Denmark<br>www.etac.com  |
| <b>SRN</b>                              | DK-MF-000019241   |
| <b>Statement</b>                        | This EU declaration of conformity is issued under the sole responsibility of the manufacturer. The device(s) covered by present declaration is/are in conformity with EU Regulation 2017/745 on medical devices.  |
| <b>Basic UDI-DI</b>                     | 57080121013LR   |
| <b>Device description</b>               | Positioning Horizontal  |
| <b>EMDN</b>                             | Y122799   |
| <b>Intended purpose</b>                 | The assistive device is intended for alleviation of, or compensation for, a functional impairment due to an injury or disability. The device is designed for an individual requiring reduction of friction at pressure points due to reduced mobility or physical strength. |
| <b>Device name(s)</b>                   | MultiGlide Single Patient Use   |
| <b>Risk class of the device</b>         | Class I, rule I   |
| <b>Harmonized/Established Standards</b> | Separate list available upon request  |
| <b>Place</b>                            | Gedved, Denmark   |
| <b>Date of issue</b>                    | 15. January 2024  |
| <b>Name and function</b>                | Michael Bruun, Senior Vice President  |

Signature, on behalf of Etac Immedia A/S