

EU Declaration of Conformity (DoC)

NOTICE: Sections bracketed with three plus signs (+++) may not be changed or removed without approval from a Quality Director or designee within the Entity and/or function (do not delete the text in this header).

Straps	Document Number: NPD39197; Version: 3.0																										
Manufacturer Name and Address: Liko AB and Nedre vagen 100, 975 92 Lulea, Sweden, +46 (0)920 474700																											
Manufacturer Single Registration Number (SRN): SE-MF-000001404																											
Authorised Representative Name and Address: Not Applicable, Registered place of business is within European union																											
Authorised Representative Single Registration Number (SRN): Not Applicable																											
+++ We as Manufacturer declare, under our sole responsibility, that the product(s) listed below conform to the applicable provisions of the Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on Medical Devices, and the following Directive(s), Regulation(s) and Common Specification(s). +++																											
Other relevant Directives, Regulations and Union Legislations that the device is in conformity with: Not Applicable																											
Common Specifications Applied: Not Applicable																											
Product/Trade Name and Product Code or REF. number: MultiStraps and FixStraps																											
<table border="1"><thead><tr><th>Reference Number</th><th>Description</th><th>UDI</th></tr></thead><tbody><tr><td>1. 3696002</td><td>1. FIXSTRAP 260 CM</td><td rowspan="11">0887761GMN000038UD</td></tr><tr><td>2. 3696002-10</td><td>2. FIXSTRAP 260 CM</td></tr><tr><td>3. 3695115</td><td>3. MULTISTRAP STANDARD</td></tr><tr><td>4. 3695116</td><td>4. MULTISTRAP WIDE</td></tr><tr><td>5. 3695117</td><td>5. MULTISTRAP ULTRA</td></tr><tr><td>6. 3695915</td><td>6. SOLO MULTISTRAP STANDARD</td></tr><tr><td>7. 3695915-2</td><td>7. SOLO MULTISTRAP STANDARD</td></tr><tr><td>8. 3695916</td><td>8. SOLO MULTISTRAP WIDE</td></tr><tr><td>9. 3695916-2</td><td>9. SOLO MULTISTRAP WIDE</td></tr><tr><td>10.3696001</td><td>10.FIXSTRAP 135 CM</td></tr><tr><td>11.3696001-10</td><td>11.FIXSTRAP 135 CM</td></tr></tbody></table>	Reference Number	Description	UDI	1. 3696002	1. FIXSTRAP 260 CM	0887761GMN000038UD	2. 3696002-10	2. FIXSTRAP 260 CM	3. 3695115	3. MULTISTRAP STANDARD	4. 3695116	4. MULTISTRAP WIDE	5. 3695117	5. MULTISTRAP ULTRA	6. 3695915	6. SOLO MULTISTRAP STANDARD	7. 3695915-2	7. SOLO MULTISTRAP STANDARD	8. 3695916	8. SOLO MULTISTRAP WIDE	9. 3695916-2	9. SOLO MULTISTRAP WIDE	10.3696001	10.FIXSTRAP 135 CM	11.3696001-10	11.FIXSTRAP 135 CM	
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Intended Purpose/Use: The MultiStrap are used for lifting and holding parts of the patient's body to facilitate examination, care and treatment. The MultiStrap can also be used for turning a patient in bed and supporting the patient in a lateral position.																											

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Solo MultiStrap are intended for individual use. The solo MultiStrap are used for lifting and holding parts of the patient's body to facilitate examination, care and treatment. The solo MultiStrap can also be used for turning a patient in bed and supporting the patient in a lateral position.

It is intended for use in following environments: Health care, Intensive care, Emergency ward, Rehabilitation, Habilitation and Home healthcare environment.

Device Risk Class: Class I

MDR EU Certificate(s) No.: Not Applicable

Conformity Assessment Description/Annexes: Annex II and III

Notified Body Name and Address: Not Applicable as it is class I Product

Notified Body Identification Number: Not Applicable as it is class I Product

+++ This Declaration is made on the following basis:

- For devices with a MDR EU Certificate issued by a Notified Body:
 - The validity of this document shall not start earlier than the validity date of the corresponding MDR EU Certificate.
 - The DoC declares conformity to all product lots released within the validity period/dates of the corresponding MDR EU Certificate.
- For Class I devices (*that are non-sterile, have no measurement function or are not reusable surgical instruments*) the DoC declares conformity to the product lots released after the date of signature.
- Compliance to standards and regulations as defined in the Technical Documentation and General Safety and Performance Requirements (GSPR).
- Additional information may be attached/appended to this template, such as common specifications, compliance to other union regulations/registrations, product code list or any other supporting information. +++

Authorised Signatory:	
Name and Title:	Sofie Nybom
Function:	QMR
Place of Issue:	Luleå, Sweden
Date of Issue:	27-NOV-2023
Signature:	