

[Issue 1]

## Declaration of Conformity EU

<b>Manufacturer Details:</b>	Silvalea Limited, Units 1-4 Silverhills Buildings, Silverhills Road, Decoy Industrial Estate, Newton Abbot, Devon, UK, TQ12 5LZ
<b>Single Registration Number:</b>	GB-MF-000003832
<b>Authorised Representative:</b>	Advena Limited, Tower Business Centre, 2nd Floor, Tower Street, Swatar, BKR 4013. Malta
<b>Authorised Representative SRN:</b>	MT-AR-000000234
<b>Basic UDI-DI:</b>	506040783
<b>Name of the Device:</b>	Disposable Patient Lifting Slings
<b>Intended Purpose:</b>	Disposable Slings are intended to be used to transfer patients between bed, chair and/or wheelchair and provide head and torso support to patients.
<b>Device variants:</b>	No variants
<b>Classification:</b>	Class 1
<b>GMDN code:</b>	37408

Silvalea Limited are governed and conducts its manufacturing and design towards the guidelines set out in the International Standard ISO EN 10535 'Hoists for the transfer of disabled persons – Requirements and test methods,' EN ISO 3758:2012, LOLER:1998.

This standard sets out the requirements for testing both the materials and the overall design of a sling to ensure conformity of the standards exacting requirements. As a UK manufacturer our guidelines are set out under the BS EN ISO 10535:2006 testing requirements.

The other harmonised standards used to comply with the Medical Device Regulation (EU) 2017/745 are ISO 13485:2016, ISO 14971:2019 and EN 15223-1:2016.

Silvalea are pleased to advise that this international standard also covers the Australian Standard AS/NZS ISO 10535:2011. Silvalea are registered with the FDA, registration number: 3008720499. The conformity assessment route followed by Silvalea involved drawing up the Technical Documentation according to Annex II and Annex III of the (EU) 2017/745, issuing the Declaration of Conformity according to Annex IV and finally assigning the CE marking.

### Declaration:

This declaration of conformity is issued under the sole responsibility of Silvalea Limited. We hereby declare that the medical device specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices. This declaration is supported by the Quality System approval to ISO:9001. All supporting documentation is retained at the premises of the manufacturer.

Approved by:



**Gary Bevan**  
Director, UK. On Behalf of Silvalea Limited

**Date: 24.05.2021**