



# CERTIFICATE OF EUROPEAN UNION AUTHORISED REPRESENTATIVE



*This is to certify that Trans Lite LLC.*

has duly registered the following relevant product types with the UK Competent Authority through its Appointed Representative in accordance with *Article 14* of the Council Directive 93/42/EEC as revised by Council Directive 20/47/EC concerning medical devices (The “Medical Devices Directive”) (UK Medical Devices Regulation 1994: Regulation 14).

\*\*\*\*\* Applicable ANNEX \*\*\*\*\*

Annex VII

\*\*\*\*\* Scope of Supply \*\*\*\*\*

Product Class I non-sterile. Product Family: Veinlite

Competent Authority Registration Reference CA 008352

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In accordance by self-declaration with *Article 11* and *Annex VII* for Class I devices may apply the CE Mark

\*\*\*\*\* Appointment \*\*\*\*\*

We certify that M. Devices Group was appointed as the Authorised Representative on the 5<sup>th</sup> August 2004

Signature  
Authorised Representative

Date: 31 August 2016



Certificate No. MDG—AR-40 Valid to 4-August-2017

Health & Education Centre, The Church  
Portland Street, Southport PR8 1HU  
United Kingdom