



CERTIFICATE OF EUROPEAN UNION AUTHORISED REPRESENTATIVE



This is to certify that 3 Gen 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 (revised by 2007/47/EC) concerning medical devices

(The “Medical Devices Directive”) (UK Medical Devices Regulation 1994: Regulation 14).

***** Applicable ANNEX *****

Annex VII

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

DERMLITE Class I non-sterile

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 15th Nov 2003

Signature
Authorised Representative

Date 14Nov2012



Certificate No. MDG-1034-AR Valid to 14 Nov 2015

Marlborough House, Riding Street,
Southport, PR8 1EW, England.