

3Gen Inc

31521 Rancho Viejo Rd, Suite #104
San Juan Capistrano, CA 92675, USA

Declaration of Conformity

Annex of the Medical Device Directive under which Declaration is made: **VII**

Date of Issuance: April 16, 2014 (Supersedes Declaration dated: March 18, 2014)

Manufacturer:
3Gen, Inc.
31521 Rancho Viejo Rd., #104,
San Juan Capistrano, CA 92675

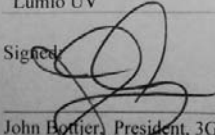
EC Representative:
M. Devices Group
Marlborough House, Riding Street,
Southport, PR8 1EW, UK
Tel: +44 1704 544 944

This Declaration is applicable to all products listed and manufactured after the Date of Issuance of this Declaration of Conformity.

We hereby declare under our sole responsibility, that the following products, comply fully with the requirements of the Medical Device Directive (93/42/EEC, amended by directives 98/79/EC, 2000/70/EC and 2007/47/EC).

Model Number	Device Class
DermLite DL100	I
DermLite Platinum	I
DermLite Pro DP-R	I
DermLite Foto	I
DermLite II Pro (DL2Pro)	I
DermLite II Multispectral (DL2MS)	I
SkinLite ALT100-0033	I
DermLite II Pro HR	I
DermLite II Fluid	I
DermLite II Hybrid	I
Lumio	I
SkinLite II	I
Carbon	I
Alumina	I
DermLite 3	I
DermLite I	I
Lumio S	I
DLCam	I
DL3N	I
Foto II Pro (DLF2-PRO)	I
Lumio UV	I

Signed


John Botjer, President, 3Gen Inc.
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