

SGS

Certificate GB17/873380

The management system of

Outside In (Cambridge) Limited also trading as Lumie

3 The Links, Trafalgar Way, Bar Hill, Cambridge, CB23 8UD, UK

has been assessed and certified as meeting the requirements of

ISO 13485:2003 EN ISO 13485:2012

For the following activities

**Manufacture and service of light therapy devices for the treatment
of acne vulgaris and SAD (seasonal affective disorder),
and the symptoms of SAD.**

This certificate is valid from 17 October 2017 until 31 March 2019
and remains valid subject to satisfactory surveillance audits.
Re certification audit due before 31 March 2019
Issue 1. Certified since 17 October 2017

Authorised by



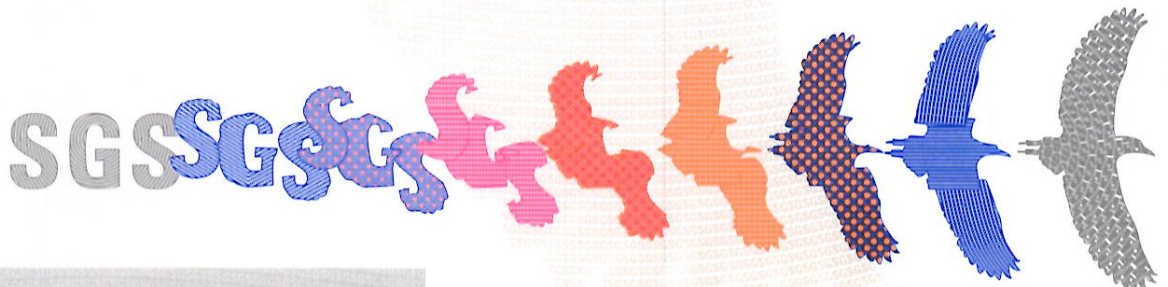
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SGS 13485 2003 0117

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The management system of

Outside In (Cambridge) Limited also trading as Lumie

3 The Links, Trafalgar Way, Bar Hill, Cambridge, CB23 8UD, UK

has been assessed and certified as meeting the requirements of

Directive 93/42/EEC on medical devices, Annex V

For the following products

The scope of registration appears on page 2 of this certificate

This certificate is valid from 17 October 2017 until 17 October 2022
and remains valid subject to satisfactory surveillance audits.
Re certification audit due before 13 October 2020
Issue 9. Certified since 29 November 1999

Certification is based on reports numbered GB/PC 228934

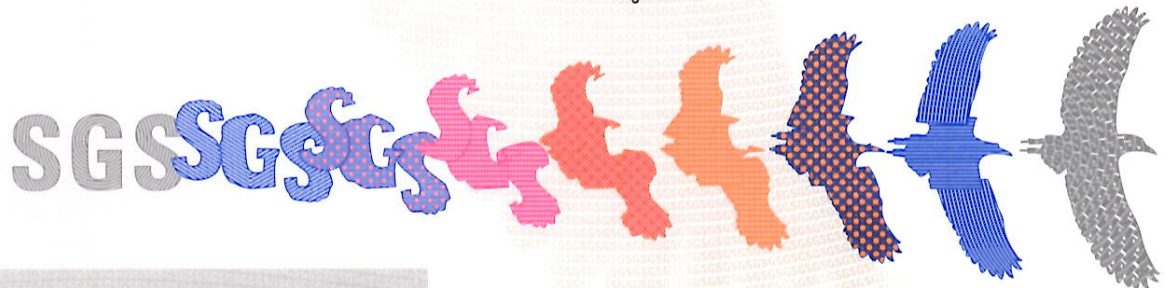
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Outside In (Cambridge) Limited also trading as Lumie

Directive 93/42/EEC on medical devices, Annex V

Issue 9

Detailed scope

Light therapy devices for the treatment of Seasonal affective disorder(SAD):

- Bright Spark BS2
- LUMIE Arabica LBA
- Lumie Desklamp LDL2
 - LUMIE Zest
 - LUMIE Brazil
- Light therapy device for the treatment of Acne Vulgaris
 - LUMIE CLEAR.

Where the above scope includes class IIb or class III medical device(s), a valid EC Type Examination Certificate according to Annex III is a mandatory requirement for each device in addition to this certificate to place that device on the market.