

DECLARATION OF CONFORMITY  
TO COUNCIL DIRECTIVE 93/42/EEC of 14 JUNE 1993  
CONCERNING MEDICAL DEVICES

**Manufacturer:** Candela Corporation  
530 Boston Post Road  
Wayland, MA 01778-1886

**European Representative:** Scanlan Group B.V.  
Aalsmeerderweg 610  
1437 EJ Schiphol-Rozenburg  
The Netherlands

**Medical Device:** GentleLASE Pro (Model No. 9914-00-9015)  
GentleLase Pro LE (Model No. 9914-00-9040)

**Type of Equipment:** Dermatology Laser System

**Classification – ANNEX IX:** CLASS IIb, RULE 9

**Conformity Assessment Route:** ANNEX II, CLAUSE 3

We, the Manufacturer, herewith declare that the stated medical devices meet the transposition into national law, the provisions of Council Directive 93/42/EEC of 14 June 1993 concerning medical devices – as amended by Directive 2007/47/EC. All supporting documentation is retained at the premises of the manufacturer.

**Standards Applied:** SEE ATTACHED LIST OF (HARMONIZED – EN)  
STANDARDS FOR WHICH DOCUMENTED EVIDENCE FOR  
COMPLIANCE CAN BE PROVIDED

RoHS compliance is ensured under the sole responsibility of the manufacturer.

**Notified Body:** TÜV SÜD Product Service GmbH  
Ridlerstrasse 65  
D-80339 München  
Germany

**Identification Number:** 0123

**(EC) Certificate(s):** G1 14 01 34736 021

**Start of CE-Marking:** April 2011 (GentleLASE Pro)  
August 2012 (GentleLase Pro LE)

**Place and Date of Declaration:** Candela Corporation, Wayland, MA, USA  
September 30, 2014

**Signature:** 

**Sam Wade**  
Vice President, Regulatory Affairs/Quality Assurance

DECLARATION OF CONFORMITY  
TO COUNCIL DIRECTIVE 93/42/EEC of 14 JUNE 1993  
CONCERNING MEDICAL DEVICES

EN 60601-1:2006/A1:2013  
EN 60601-1-2:2007  
EN 60601-1-6:2010  
IEC 60601-2-22:2007  
EN 60825-1:2007  
EN 62304:2006/AC:2008  
EN 62366:2008  
EN ISO 13485:2012  
EN ISO 14971:2012  
EN ISO 10993-1:2009  
EN 1041:2008  
MEDDEV 2.7.1