EU Declaration of Conformity

EndyMed Medical Systems Ltd
12 Leshem Street,
Caesarea, 3088900,
Israel

We, EndyMed, manufacturers of RF Skin Treatment Systems, as detailed hereunder, that are placed in the European market, declares that our products conforms and meets the essential requirements set out in Annex I of the Low Voltage Directive (LVD 2014/35/EU).

The EU Directives covered by this Declaration

2014/35/EU Low Voltage Equipment Directive

The Products Covered by this Declaration

- Newa™
  Model: Collagen Visage

The manufacturer hereby declares under his sole responsibility that the products identified above comply with the principal elements of the safety objectives of the Low Voltage Equipment directive according to the following standard

EN 60601-1 Electrical Safety Testing of Medical Equipment
EN 60601-1-2 Electromagnetic Compatibility
EN 60601-1-11 Electrical Safety for Medical Equipment and Medical Electrical Systems Used In Home Health Care Environment
EN 60335-2-23 Household and similar electrical appliances - Safety - Part 2-23: Particular requirements for appliances for skin or hair care

Within these requirements we prepared the required technical documentation to demonstrate that the product meets the requirements of the Low Voltage Directive has been complied by the signatory below and is available for inspection by the relevant enforcement authorities. We have appointed QsiteEU,
Gerrit van der Veenstraat 84HS,
1077 EL Amsterdam,
The Netherlands
to act as our Authorized Representative in the European Community.
This CE mark was first applied in: 2014.
The products described above comply with the essential requirements of the directives specified.

**Name:** Rebecca Feldman  
**Date:** 01/01/2017– 31/12/2018  

**Location:** Caesarea, Israel  
**Signature:** …………………