

Document No-LaboAmerica/MDR/24-04-1.0

**EU DECLARATION OF CONFORMITY****Labo America**

Declare under sole responsibility that

Our medical devices: Surgical Microscopes Dentistry

Models: Prima Lite/ Prima Pro/Magna/Prima DNT/ Prima Mu DNT

Model Nos.: 6181501,6181502,6181503/6183601,6183602,6183603/
6129100,6129200,6129300/6138000,6137101,6137201,6137500/6211000

Basic UDI-DI 81017855DNTSURGICALCW

Are AC powered devices intended for use in dentistry, general & plastic surgery
to provide a magnified view of the region of interestHave been subjected to conformity assessment procedure complying with Article 52 (7) and technical
documentation accordance to Annexure II and III of council regulation (EU) 2017/745.These Devices are Class 1 (without a measuring function) per Rule 13 annex VIII
of the regulation MDR 2017/745Labo America designs and manufactures all products under Quality Systems registered to meet
the requirements of ISO 9001, ISO 13485, 21CFR 820 and MDR 2017/745. In case of
alteration of the device, not agreed upon by us, this declaration will lose its validity.

Applied harmonized standards, national standards and other normative documents:

EN/ISO: 15223-1 EN
EN /IEC 60601-1-1
EN /IEC 60601-1-2
EN/ISO: 14971
EN/ISO: 10936
EN 62366EN 61010-1
EN/IEC 61326-1
EN/IEC 61000-3-3
EN/IEC 61000-3-2
CISPR11/EN 5501**AUTHORIZED SIGNATORY**Gautam Aggarwal
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