DECLARATION OF CONFORMITY

EC COUNCIL DIRECTIVE 93/42/EEC FOR MEDICAL DEVICES

Manufacturer:  LM-Instruments Oy
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Herewith we declare that all the dental hand instruments we are producing as well as LM-Cello, LM-EndoMax, LM-ErgoHold3, LM-Multiseptor, LM-Servo, LM-ServoMax, extraction instruments, orthodontic appliances and all accessories for above-mentioned products as well as LM-ProPower air polishing powders, LM-Sodium B, LM-Calcium C and LM-Glycine Neutral, comply with the relevant regulations of the guideline 93/42/EEC for medical devices as they have been modified with the directive 2007/47/EC. The products are classified in: Medical Devices, Class I.

Date, place: 21.10.2015, Parainen

[Signature]

Kari Lehtonen
Development Manager
LM-Instruments Oy