

## **EC Declaration of Conformity**

This declaration of conformity is issued under the sole responsibility of ENRAF-NONIUS B.V., Vareseweg 127, 3047 AT Rotterdam, The Netherlands

Quality Management System according to EN ISO 13485:2016 and Annex II (excl. section 4) of European Medical Devices Directive 93/42/EEC.

Registration No. HD 1274285-1

Notified body: TÜV Rheinland LGA Products GmbH (0197), Tillystraße 2, 90431 Nürnberg, Germany

Device name:

Physisonic ES-7

Description:

**Ultrasound Therapy Equipment** 

Article number(s):

1631931 / 1631932 / 1631933

1631961 / 1631962 / 1631963 / 1631964

Classification:

Ila (according to rule 9, in subclause 3.1 of Annex IX of MDD

93/42/EEC)

Record of conformity:

026-400-273-45 ER

Device marking:

CE0197

We hereby declare that the above-mentioned device complies with the European Medical Devices Directive 93/42/EEC

This declaration of conformity is valid in combination with the test certificate of the device. Any modifications to the product not authorized by Enraf-Nonius will invalidate this declaration.

Rotterdam, 10 March 2023

Signature:

The Doodkorte
Commercial Director