CERTIFICATE OF CONFORMITY WITH EUROPEAN DIRECTIVE



Certificate No.: EU0904409 Order No.: 124470

We hereby certify that an examination has been carried out following the requirements of the national legislation "Regulation no. 1690 of 15thth December 2005 relating to medical devices pursuant to act no. 6 of 12th January 1995 relating to medical devices, transposing directive 93/42/EEC into Norwegian law to which the undersigned is subjected, confer EEA agreement, proposition no. 100 (1991-92) special appendix no. 2, volume 2A/3 A, goods, chapter XXX". We certify that the production quality system conforms to the relevant provisions of the Annex given below:

Name and address of the

manufacturer:

IBRAMED - Indústria Brasileira de Equipamentos Médicos Ltda

Rua Milão, 50 - Jardim Itália - CEP 13901-070

Amparo - SP

Brasil

Device category:

Ultrasound Therapy Equipment

GMDN code:

11248

Models:

See Appendix 1 to this certificate

Risk class as defined by the

manufacturer:

lla

Standards/provisions:

The audit of the quality system was based upon and assessed according

to the provisions in Annex II of the EC-Directive 93/42/EEC, with the

exemption of section 4.

Date of audit:

2009-04-28/29

Nemko EC notification No.:

0470

On this basis the manufacturer or the European authorised representative may draw up an EC / EEA Declaration of Conformity and affix the CE-marking as indicated below together with the Nemko EC notification number to each conforming product as long as the conformity audit and inspection procedure required by the EC directive will be fulfilled by the manufacturer and the factory. The product liability rests with the manufacturer or his representative in accordance with Council Directive 85/374/EEC.

Date of issue: 2009-07-28

April Stanjones

Date of verification: 2009-07-28

budgemall blad

signature: Frank Skarpsno Lead auditor/Principal Engineer signature: Arild R. Hansgård

Principal Engineer

C€ 0470

Nemko AS P.O. Box 73, Blindern N-0314 Oslo, Norway Office address Gaustadalléen 30 Oslo

Telephone +47 22 96 03 30

+47 22 960 550

Enterprise number: NO 9443522430

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Ultrasound Therapy Equipment

Appendix 1: Page 1 of 1.

The certificate referred to above includes the following devices/models:

Sonopulse 1 e 3MHz

Sonopulse Compact 1MHz

Sonopulse Compact 3MHz

Sonopulse III 1 e 3MHz

Date of issue: 2009-07-28

Date of verification: 2009-07-28

Arild Hanagard

Signature: Arild R. Hansgård
Principal Engineer

Signature: Frank Skarpsno Lead auditor /Principal Engineer