

EC-Declaration of Conformity

according to Directive 93/42/EEC, Annex II excluding section (4)

MiE medical imaging electronics GmbH

represented by the presidents Thomas Kühl and Günther W. Kühl

declares in own responsibility that following described system fulfills the requirements of Annex I of the Directive 93/42/EEC.

Manufacturer:

MiE medical imaging electronics GmbH

Hauptstr. 112,

D-23845 Seth, Germany

Tel.: +49 4194 9977-0 Fax: +49 4194 9977-55 mie@miegermany.de www.mie-scintron.com

Product Name:

System PICOLA - SCINTRON

Product Description:

Anger type gamma-camera with integrated energy- and linearity

correction and Nuclear Medicine Workstation

Classification:

Class IIa

(according to Rule 10 of Annex IX to Council Directive 93/42/EEC)

Validity:

This Declaration of Conformity is valid until 26.05.2024

Name and Address of Notified

Body:

DEKRA Certification GmbH

Handwerkstraße 15

D-70565 Stuttgart, Germany

Notified Body number:

0124

Quality Management System:

EN ISO 13485:2016/AC:2016

Medical Devices - Quality Management Systems -

Requirements for regulatory purposes

Seth, 20.05.2021

Date

Signature President