

PD-730-165 EC Declaration of Conformity

The manufacturer IMAGE Information Systems Europe GmbH
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declares under its sole responsibility that the medical device stated as follows:

iQ-VIEW / PRO 3.1

is classified as **Class IIa** according to rules **10** and **16** of the Medical Device Directive 93/42/EEC, Annex IX.

The conformity assessment has been performed according to Annex II (4) of MDD 93/42/EEC based on the following elements:

- Conformity to the Essential Requirements according to Annex I of MDD 93/42/EEC
- Quality Management System for the products / product categories

Digital image processing systems

The license of certification is subject to surveillance by the Notified Body.

MEDCERT GmbH
Pilatuspool 2
20355 Hamburg, Germany
(Notified Body CE 0482)

Rostock, 2018-04-12


Dr. Arpad Bischof
Managing Director