EU Declaration of Conformity

acc. to Regulation (EU) 2017/745, Annex IV

We, the manufacturer

IMAGE Information Systems Europe GmbH

SRN: DE-MF-000004979

Lange Str. 16 18055 Rostock, Germany Tel.: +49 381 496 58 20

www.image-systems.biz | info@image-systems.biz

declare, under our sole responsibility, the conformity of the medical device specified below with all applicable requirements of the Medical Device Regulation (MDR) 2017/745.

Device/trade name(s)	iQ-SYSTEM PACS		
Device version	v1		
Basic UDI-DI	++B403IQSYSTEMPACS1YG		
Intended purpose	A picture archiving and communication system (PACS) for the management, retention, routing, analysis and review of medical images stored in proper DICOM format.		
Risk class	IIb		
Applicable classification rules	11-2		
References to applied CS to which conformity is declared	N/A		
Associated device components	(Trade) Name(s)	Version	UDI-DI
	iQ-WEB	7.2.9	+B403IQWEB72906
	iQ-4VIEW	2.4	+B403IQ4VIEW240-
	iQ-VIEW	4.0	+B403IQVIEW400U
	iQ-ROUTER	5.0	+B403IQROUTER5001

The medical device stated above fulfills the General Safety and Performance Requirements according to Annex I of MDR 2017/745.

Conformity has been established by means of the conformity assessment procedures according to Annex IX of MDR 2017/745 based on the following elements:

- Assessment of the Quality Management System
- Assessment of the Technical Documentation

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

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

The EU Quality Management System Certificate as issued by the Notified Body is valid until 2028-02-14.

This declaration is valid with the date of the signature.

Rostock, 2024-07-11

DocuSigned by:

288892E72746439.. Dr. Arpad Bischof

PRRC | Managing Director