EC CERTIFICATE

Number: 2079177CE01

Full Quality Assurance System

Directive 93/42/EEC on Medical devices, Annex II excluding (4)

(Devices in Class IIa, IIb or III)

Manufacturer:

Philips Medical Systems Nederland B.V. Veenpluis 4-6 5684 PC Best The Netherlands

For the product category(ies)

Diagnostic X-Ray systems and associated software and hardware for the area of interventional and surgical procedures

Note that the address is also known as: "Veenpluis 6", which refers to the same physical location.

DEKRA grants the right to use the EC Notified Body Identification Number illustrated below to accompany the CE Marking of Conformity on the products concerned conforming to the required Technical Documentation and meeting the provisions of the EC-Directive which apply to them:

0344

Documents, that form the basis of this certificate:

Certification Notice 2079177CN, initially dated 25 January 2005 Addendum, initially dated 1 February 2005

DEKRA hereby declares that the above mentioned manufacturer fulfils the relevant provisions of 'Besluit Medische Hulpmiddelen', the Dutch transposition of the Council Directive 93/42/EEC of June 14, 1993 concerning Medical devices, including all subsequent amendments. The manufacturer has implemented a quality assurance system for design, manufacture and final inspection for the above mentioned product category in accordance to the provisions of Annex II of Council Directive 93/42/EEC of June 14, 1993 and is subject to periodical surveillance. For placing on the market of Class III devices an additional EC design examination certificate according to Annex II (4) is mandatory. The necessary information related to the quality management system of the manufacturer, including facilities and the reference to the relevant documentation, of the products concerned and the assessments performed, are stated in the Certification Notice which forms an integrative part of this certificate.

This certificate is valid until: 1 February 2023 Issued for the first time: 1 February 2005 Revised: 7 May 2020 Reissued: 7 March 2018

DEKRA Certification B.V.

B.T.M. Holtus Managing Director J.A. van Vugt Certification Manager

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ADDENDUM

Belonging to certificate: 2079177CE01

CE MARKING OF CONFORMITY MEDICAL DEVICES

Diagnostic X-Ray systems and associated software and hardware for the area of interventional and surgical procedures

Issued to:

Philips Medical Systems Nederland B.V.

Veenpluis 4-6 5684 PC Best The Netherlands

This certificate covers the following product(s):

This certificate covers the activities of the Business Unit Image Guided Therapy - Systems of the certification holder.

Allura Xper Series (class IIb):

Allura Xper FD10

Allura Xper FD20

Allura Xper FD10/10

Allura Xper FD20/10

Allura Xper FD20/15

Allura Xper FD20/20

Allura Xper OR Table Series (class Ilb):

Allura Xper FD10 OR Table

Allura Xper FD20 OR Table

Allura Xper FD10/10 OR Table

Allura Xper FD20/10 OR Table

Allura Xper FD20/15 OR Table

Allura Xper FD20/20 OR Table

BV Pulsera (class IIb)

BV Endura (class IIb)

Veradius (class IIb)

Veradius Unity (class IIb)

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CE MARKING OF CONFORMITY **MEDICAL DEVICES**

Diagnostic X-Ray systems and associated software and hardware for the area of interventional and surgical procedures

Issued to:

Philips Medical Systems Nederland B.V.

Veenpluis 4-6 5684 PC Best The Netherlands

Zenition 50 Zenition 70

Interventional tools (class IIa): Interventional Workspot 3D Roadmap

MR-CT Roadmap **XperCT Dual**

XperGuide

Stentboost

2D Perfusion

EchoNavigator

EmboGuide

HeartNavigator

EP navigator

VesselNavigator

AneurysmFlow

2D Quantitative Analysis

StentBoost Live

Dynamic Coronary Roadmap

3Ď-RA

Surgical Navigation

ViewForum

SmartPerfusion

DEKRA Certification B.V.

B.T.M. Holtus **Managing Director**

J.A. van Vugt Certification Manager

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ADDENDUM

Belonging to certificate: 2079177CE01

CE MARKING OF CONFORMITY MEDICAL DEVICES

Diagnostic X-Ray systems and associated software and hardware for the area of interventional and surgical procedures

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Veenpluis 4-6 5684 PC Best The Netherlands

Azurion Series (Class IIb):

Azurion 7 M12

Azurion 7 M20

Azurion 3 M12

Azurion 3 M15 Azurion 7 B12

Azurion 7 B20

UNIQ Series (class IIb):

UNIQ / UNIQ Clarity FD10

UNIQ / UNIQ Clarity FD20

UNIQ / UNIQ Clarity FD10/10

UNIQ / UNIQ Clarity FD20/10

UNIQ / UNIQ Clarity FD20/15

UNIQ / UNIQ Clarity FD20/20

All certified devices are manufactured in the facility:

Philips Medical Systems Nederland B.V., Veenpluis 4-6, 5684 PC Best, The Netherlands

DEKRA Certification B.V.

B.T.M. Holtus Managing Director

J.A. van Vugt Certification Manager

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The BV Endura, BV Pulsera, Veradius Unity and Zenition 50 and Zenition 70 devices are also manufactured in the subcontractor facility:

Philips India Limited., Plot no. B-79, MIDC, Phase-II, Chakan, Valuka-khed, Village – Savardari, District: Pune, Maharashtra 410 501, India

The following Class IIb products, which are no longer in production, are still subject to refurbishment:

- Integris Allura 9C (last produced: 2007)
- Integris Allura 9F (last produced: 2007)
- Integris Allura 9 Biplane (last produced: 2005)
- Integris Allura 12 & 15 Monoplane (last produced: 2008)
- Integris Allura 12 & 15 Biplane (last produced: 2006)
- OmniDiagnost Eleva (last produced: 2010)
- UroDiagnost Eleva (last produced: 2007)
- Allura CV20 (last produced: 2013)
- MultiDiagnost Eleva (class IIb) (last produced: 2015)
- BV Libra (class IIb) (last produced: 2015)
- Allura Centron (class IIb) (last produced: 2018)

Initial date: 1 February 2005 Revision date: 7 May 2020

DEKRA Certification B.V.

B.T.M. Holtus Managing Director J.A. van Vugt Certification Manager

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