

Declaration of Conformity



Manufacturer: Shenzhen Mindray Bio-Medical Electronics Co., Ltd.
Mindray Building, Keji 12th Road South, Hi-tech Industrial Park,
Nanshan, Shenzhen, 518057, P. R. China

Manufacturer SRN: CN-MF-000014156

EC-Representative Shanghai International Holding Corp. GmbH (Europe)
Eiffestraße 80 20537 Hamburg, Germany

Product: Endoscope Camera System

Model: UX5/UX5-TEC/UX5-NOR/UX5-SIM/UX3/UX3-TEC/
UX3-NOR/UX3-SIM/UX1/UX1-TEC/UX1-NOR/UX1-SIM/
UX4/UX410/UX420/UX430/UX450/UX460/UX470/UX5/
UX510/UX520/UX530/UX550/UX560/UX570/UX5-SIM/
UX5-NOR/UX5-TEC/UX7/UX7-TEC/UX7-NOR/ UX7-SIM

Basic UDI-DI: 69449040AB065100078D

Classification: I(According to Rule 13 of MDR Annex VIII)

Conformity Assessment Route: Article 52.7

EMDN code: Z120204

GMDN code: 35958

Supplementary information TV-300/TV-300T(220V)/TV-300T(110V)/TV-500/
TV-500T(220V)/TV-500T(110V)/TP-MFS Mobile Trolley;
CH5-SW100/CH5-SW110/CH5-SR100/CH5-SR110/
CH3-SW100/CH3-SW110 camera head

We declare that the above mentioned products meet the provisions of the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT. All supporting documentations are retained under the premises of the manufacturer. This declaration of conformity is issued under the sole responsibility of the manufacturer.

References to CS: /

Start of CE-Marking: 2023-6-29

I hereby am appointed as the authorized person to deal with all the registration and quality management affairs in my capacity as Deputy Director of Technical Regulation Department of Shenzhen Mindray Bio-Medical Electronics Co., Ltd, Effective immediately.

Place, Date of Issue: Shenzhen, 2024-12-31

Signature:

Name of Authorized Signatory:

Position Held in Company:

Mr. Wang Xinbing

Deputy Director, Technical Regulation

Applied Standards List

Product: Endoscope Camera System

Model:

UX5/UX5-TEC/UX5-NOR/UX5-SIM/UX3/UX3-TEC/
UX3-NOR/UX3-SIM/UX1/UX1-TEC/UX1-NOR/UX1-SIM/
UX4/UX410/UX420/UX430/UX450/UX460/UX470/UX5/
UX510/UX520/UX530/UX550/UX560/UX570/UX5-SIM/
UX5-NOR/UX5-TEC/UX7/UX7-TEC/UX7-NOR/ UX7-SIM

Standards Applied:

EN ISO 14971:2012/A11:2021	Medical devices – Application of risk management to medical devices
EN ISO 20417:2021	Medical devices - Information to be supplied by the manufacturer
EN ISO 15223-1:2021	Medical devices - Symbols to be used with information to be supplied by the manufacturer - Part 1: General requirements
EN 60601-1:2006/A2:2021	Medical electrical equipment -- Part 1: General requirements for basic safety and essential performance
EN 60601-1-2:2015/A1:2021	Medical electrical equipment -- Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests
EN 60601-1-6:2010/A2:2021	Medical electrical equipment - Part 1-6: General Requirements for basic safety and essential performance -Collateral standard: usability
EN 62304:2006/A1:2015	Medical device software - Software life-cycle processes
EN 62366-1:2015/A1:2020	Medical devices -- Application of usability engineering to medical devices
EN ISO 17664-1:2021	Processing of health care products - Information to be provided by the medical device manufacturer for the processing of medical devices - Part 1: Critical and semi-critical medical devices
EN 60601-2-18: 2015	Medical electrical equipment-Part 2-18:Particular requirements for the basic safety and essential performance of endoscopic equipment