

Manufacturer

UAB "Medinstrus" SRN LT-MF-000022328 Krucių g. 9, Krucių k., Mažeikių r.

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EU DECLARATION OF CONFORMITY FOR MEDICAL DEVICES

acc. to Article 19 of Regulation (EU) 2017/745 on Medical Devices

On our sole responsibility, we hereby declare that the product(s)

Equipment type: Chair

Model: Patient chair

Type: ChairS ,ChairSR, ChairM, ChairG, ChairE, ChairI, ChairD, ChairDR

(Basic UDI 477905507CHAIRF9)

comply with the relevant provisions of the following Regulation(s) and Directive(s),

harmonised standards:

DIRECTIVE 2014/30/EU OF THE EUROPEAN PARLIAMENT

AND OF THE COUNCIL of 26 February 2014 on the harmonisation of the laws of the Member States relating to

electromagnetic compatibility (recast)

Directive 2011/65/EU of the European Parliament and of the

Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (recast)

Text with EEA relevance

Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April

2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No

178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives

90/385/EEC and 93/42/EEC (Text with EEA relevance.)

Medical electrical equipment – Part 1: General requirements for basic safety and

essential performance EN 60601-1:2006 (including IEC 60601-1:2005/A1:2012; IEC

 $60601-1:2005/A2:2020; \ EN\ 60601-1:2006/AC:2010; \ EN\ 60601-1:2006/A12:2014; \ EN\ 60601-1:2006/A12$

60601-1:2006/A1:2013/AC:2014;

ISO 14971:2019 Medical devices — Application of risk management to medical devices ISO

14971:2019

Classification (acc. to Annex VIII of MDR): According to Rule 13, Class I.

We herewith declare that the above mentioned products,

- Do not contain a CMR (substances which are carcinogenic, mutagenic or toxic to reproduction ('CMR') and/or endocrine-disrupting substances,
- Do not contain a phthalates,

2014/30/EU

2011/65/EU

2017/745 EU

EN 60601-1:2006

- Do not contain materials of biological origin of human or animal origin,
- Do not contain nanomaterial.

The declaration of conformity is prepared based on the requirements of Regulation (EU) 2017/745 of the European Parliament and of the Council on medical devices.

Date of issue 2023-11-27 Place Mažeikiai, Lithuania



Signed on behalf of UAB Medinstrus CEO Vitalis Balčytis