

EC Declaration of Conformity

We, **Kaz Europe Sàrl**, Place Chauderon 18, 1003 Lausanne, Switzerland declare under our sole responsibility that the product to which this declaration relates is in conformity with the following standard(s) or other normative document(s) and proves the conformity of the designated product with the provisions of the below Directive(s):

- **MDD - Council Directive 93/42/EEC of 14 June 1993 concerning medical devices**
- **RoHS - 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**

Name

Braun ThermoScan 7 /ThermoScan 5 / Thermoscan 6 /
Thermoscan 7+:IRT6520, IRT6030, IRT6020, IRT6515 and
IRT6525 series

Type or model

IRT6520	IRT6020NOEE
IRT6520MNLA	IRT6020MNLA
IRT6520WE	IRT6030
IRT6520BWE	IRT6515NOEE
IRT6520NOEE	IRT6515MNLA
IRT6520LA	
IRT6525	
IRT6525NOEE	
IRT6525MNLA	
IRT6525WE	
IRT6525KO	
IRT6525AP	
IRT6525AU	

IRT6520WEGP (IRT6520WE + a Toy thermometer)
IRT6520NOEEGP (IRT6520EEE + a Toy thermometer)
IRT6520MNLAP (IRT6520MNLA + a Toy thermometer)

Note: IRT6520WEGP, IRT6520NOEEGP, IRT6520MNLAP are sold with a Toy thermometer into the packaging. The toy is covered by his own EC Declaration of Conformity under the Toy Safety Directive with Lechner Ges.m.b.H, Österreich as manufacturer.

Standards Applied:

Standard Reference	Edition	Title
EN ISO 13485	2016	Medical devices — Quality management systems — Requirements for regulatory purposes
EN 60601-1	2006 +A1:2013	Medical electrical equipment - Part 1: General requirements for safety and essential performance.
EN 60601-1-2	2015	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/A1:2013	Medical electrical equipment — Part 1-6: General requirements for basic safety and essential performance — Collateral Standard: Usability.
EN 60601-1-11	2015	Medical electrical equipment – Part 1-11: General requirements for basic safety and essential performance - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment
EN ISO 14971	2019	Medical devices — Application of risk management to medical devices.
EN 62304	2006 A1:2008	Medical device software - Software life-cycle processes

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EN ISO 10993-1	2009/AC:2010	Biological evaluation of medical devices - Part 1: Evaluation and testing.
EN ISO 10993-5	2009	Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity
ISO 10993-10	2010	Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization
EN 62366-1	2015	Medical devices - Application of usability engineering to medical devices.
EN ISO 80601-2-56	2017	Medical electrical equipment - Part 2-56: Particular requirements for basic safety and essential performance of clinical
EN 12470-5	2000+A1:2009	Clinical thermometers - Part 2: Phase change type (dot matrix) thermometers
EN 1041	2008/A1:2013	Information supplied by the manufacturer with medical devices
EN ISO 15223-1	2016	Symbols to be used with medical device labels, labeling, and information to be supplied - Part 1 - General requirements
ASTM E1965-98	2016	Standard Specification for Infrared Thermometers for Intermittent Determination of Patient Temperature
EN ISO 14155	2011/AC:2011	Clinical investigation of medical devices for human subjects - Good clinical practice

The Technical Documentation is the responsibility of: Kaz Europe Sàrl, Place Chauderon 18, CH-1003 Lausanne, Switzerland

Additional information:

For Medical Device Directive 93/42/EC	
Regulatory class (MDD, Annex IX):	class IIa (Annex IX rule 10)
Conformity assessment procedure:	Annex V
GMDN	17887
UMDNS	17-887
Notified Body	DQS Medizinprodukte GmbH August Schanz Str. 21 D-60433 Frankfurt, Germany Registration number: 0297
EC Certificate	381008 MR5
EN ISO 13485 Certificate	381008 MP2016

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This declaration of conformity is valid until May 26, 2024.

Michael Burke

Lausanne December 8, 2021

General Manager EMEA

Legally binding signature

Place

Date

Company Stamp:



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