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Operator's manual

lamp for operating theatre

TRIS-LED

Single Ceiling
Double Ceiling
(TRIS-LED+TRIS-LED)
Mobile
Wall



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Introduction

Dear user.

You are kindly invited to read this manual carefully before proceeding to use the Product in order to safeguard yourself and other people from any injuries.



This appliance is a Class 1 medical device pursuant to European Directives on medical devices (MDD) 93/42/EEC, Annex IX, and 2007/47/EC.

Conformity

The manufacturer declares that this product is in compliance with Annex I (essential requirements of Directive 93/42/EEC and certifies such conformity by affixing the CE mark.

The Product is classified in risk group 1 according to IEC 62471 standard (Photobiological Safety of Lamps).

Validity of manual

This operator's manual refers to the following Products:

- TRIS-LED single-ceiling version
- TRIS-LED double-ceiling version (TRIS-LED+TRIS-LED)
- TRIS-LED mobile version
- TRIS-LED wall version

Customer service

The customer service is at your disposal in case of Product details, information concerning its use, identification of spare parts being required and for any other queries you might have concerning the appliance, for ordering spares and for matters relating to assistance and warranty.

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Right to make changes Rimsa reserves the right to change, cancel or otherwise amend the data contained in this document at any time and for any reason without prior notice inasmuch as Rimsa is constantly seeking new solutions which lead to product evolution. Rimsa therefore reserves the right to make changes to the supplied Product in terms of shape, fittings, technology and performances.

Translations

With regard to translations into languages other than Italian, reference shall always be made to the Italian edition of this operator's manual.



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Manufacturer's declaration of conformity C€

The company:

RIMSA P. LONGONI S.r.I. Via Monterosa, 18/20/22 - 20831 SEREGNO (MB) - ITALY

Declares under its own responsibility that the Product (Medical lighting device for surgical and diagnosis use):

TRIS-LED

APPLICARE ETICHETTA

made by RIMSA P.LONGONI S.r.I., complies with Annex VII of Directive 93/42/EEC dated 14/05/1993, and subsequent amendments (including Directive 2007/47/EC dated 05/09/2007) and the following standards:

• IEC 60601-1 (Part 1: General requirements for basic safety and essential performance)

• IEC 60601-2-41 (Part 2: Particular requirements for basic safety and essential performance of surgical

luminaires and luminaires for diagnosis)

• IEC 60601-1-2 (Part 1: General requirements for basic safety and essential performance – Collateral

standard: Electromagnetic compatibility – Requierements and tests)

Classification with reference to article 9 and Annex IX of Directives 93/42/EEC and 2007/47/EC

DURATION: Short term (Par.1 "Definitions", art.1, sub-section 1.1, annex IX)

DESCRIPTION: Non-invasive medical device (Par.1 "Definitions", art.1, sub-section 1.2, annex IX)

Active medical device (Par.1 "Definitions", art.1, sub-section 1.4, annex IX)

My-P. LONGON [S.L.].

CLASS: I (Par.3 "Classification", art.1, sub-section 1.1 Rule 1, annex IX)

Technical reference file Code RIM-FT002.

 The conformity assessment is developed in reference to article 11 of the 93/42/EEC Directive and 2007/47/EC.

• The RIMSA Quality System conforms to the UNI EN ISO 9001 and UNI EN ISO 13485 regulations and is certified by the CSQ (certified CSQ n.9120.RMS1 and 9124.RMS2).

Name: Paolo Longoni Position: Managing Director



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1 Important information for the user

Product

The ME (Medical-Electrical) EQUIPMENT to which this manual refers is a LAMP for Operating Theatre or SYSTEM of LAMPS for Operating Theatre. For ease of description, this ME EQUIPMENT will be referred to in this manual with the name of "Product".

1.1 User qualification

Personnel

The Product and these operating instructions are intended for use by medical personnel and qualified technicians working in hospitals and medical surgeries who have acquired working skills by undergoing medical training and who are in possession of necessary authorisation where required.

Personal safety

Importance of personal safety. Before using the Product, read the safety

precautions in paragraphs 2.1, 3.4 and 3.5

Adjustments

The operations described in Chapter 6 "Adjustments" must be performed by a qualified technician of the Product operator in accordance with the safety rules and precautions indicated in this operator's manual.

Cleaning

Product cleaning can only be done by duly trained personnel.

Importance of manual

This manual is an integral part of the Product according to the provisions of the European Directives 93/42/EEC and 2007/47/EC.

Always keep this operator's manual close to the Product so as to be able to refer to it in case of doubts relating to lamp use, safety matters and other important information.

Never transfer the Product to another user or to other premises without its being accompanied by this operator's manual.

This manual must always accompany the Product.

These operating instructions must always be easily accessible to any Product user.

You are invited to carefully read this operator's manual before using the Product. This way, you can make best use of Product potential and protect yourself and others from any injuries.



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1.2 Precautions for safe appliance operation

Correct installation

This operator's manual is only valid after the correct installation of the Product, made in compliance with the valid installation instructions and with the correct startup by a professional installer. This operator's manual does not replace the obligation to instruct the user to carry out operations important for safety, operating, using and looking after the Product.

Safety provisions

The Product is made according to the current state of the art and its operation is safe, as long as it is used in compliance with all operating instructions and safety precautions.

Use of the Product can nevertheless be dangerous, especially if it is used by unqualified or inexpert persons or in an incorrect way, without abiding by the safety precautions contained in this operator's manual or in a way not in compliance with intended use.

To only be used in compliance with intended use

The Product is only designed to be used for the purposes indicated in this operator's manual. Any other use could cause mortal danger and/or hazards for the Product and the other material assets of the operator.

2 Precautions for the appliance operator

2.1 Technical safety specifications

Cleaning personnel

The Product cleaning and disinfecting operations described in Chapter 5 must only be performed by duly trained personnel.

Servicing personnel

The inspection and maintenance operations described in Chapter 6 must only be performed by professional technical personnel.

2.2 Personnel training obligation

Instructing users

Instruct personnel according to the operating instructions as regards controlling, cleaning and looking after the lamp.

The operator must provide such personnel with written instructions based on this manual.



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2.3 Warranty and liabilities

Rimsa disclaims all liability as regards unreliable Product operation in the following cases:

- assembly, changes and repairs not being made by a technician who has attended a training course on the Product organized by the manufacturer or by a professional technician,
- the Product not being used for the purposes for which it was intended, in compliance with the operating rules and instructions.

2.4 Structural changes or variations

Arbitrary changes

For safety reasons, no arbitrary structural changes or variations to the Product are acceptable. In case of changes or transformations of this kind, the manufacturer's Product warranty shall be invalidated. The manufacturer thus disclaims all liability for any damage or injuries caused by any arbitrary structural modifications or variations made or the use of non-original spare parts.

Only use original Rimsa spare parts

The use of parts not supplied by Rimsa or its dealers shall invalidate the warranty.

2.5 Disposal after use

Disposal at the end of life cycle

The used Product contains valuable materials which can be recycled.

Dispose of the used Product in an environment-friendly way and in compliance with applicable national directives on waste disposal.

3 Importance of personal safety

3.1 Intended use

Use in compliance with

standards The Product is made to light up the area occupied by the patient undergoing

surgery or observation and has been designed for use in operating theatres

or medical surgeries.

Field of work The Product correctly lights up the field of work from a distance of about 70

- 140 cm from the point of operation.



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Single lamp:

Definition

In compliance with the IEC60601-2-41 standard, a single lamp (TRIS-LED) is a secondary scialytic lamp for surgery and can only be used in operations where the interruption of lighting does not cause risks for the patient.

System of operating lamps:

Definition

In compliance with the IEC60601-2-41 standard, a system of lamps (TRIS-LED+TRIS-LED) made up of several lamp units can be used to locally light up the patient's body without any limitation. It is also suitable for continuous function.

It enables the surgeon to operate also in the most difficult conditions of visibility. It is intended to make treatment and diagnosis possible and to be used in operating theatres.

Undesired effects caused by superimposition of light fields If the light fields of several lamp units are superimposed, there will be an increase in heat in the patient area with consequent dehydration of tissues and, above all in the case of prolonged operation and reduced blood supply, considerable damage to tissues.

If reduced blood supply or the start of tissue dehydration occurs, reduce the light intensity.

3.2 Environmental conditions

- The Product is not suitable for use in explosion-risk areas.
- The Product is not suitable for use in the presence of inflammable mixtures of anesthetics with air, oxygen or NO₂ (laughing gas).
- During operation, the ambient temperature must be between 10°C and 40°C.
- Relative humidity must be between 30% and 75%.
- Atmospheric pressure must be between 700 and 1060hPa.

3.3 Use in combination with other medical products

- the Product can be equipped with appliances of other manufacturers.

 Refer to the operating instructions for such appliances.
- Only fit medical devices (e.g., LCD monitors) bearing the CE mark.



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3.4 Technical safety conditions

The safe use and proper operation of the Product is ensured if:

Safe fastening The lamp is safely fastened to the ceiling/wall from a static viewpoint and a

static stability test exists,

Wiring systems The wiring systems of the premises involved are in compliance with

applicable local regulations,

Authorised personnel Changes to the lamp or maintenance jobs are performed by personnel

trained by Rimsa or by a professional technician

Correct assembly

and start up

The Product has been installed following currently valid installation

instructions and has been started up by a professional installer,

Original spares With regard to assistance, repairs, structural changes and additional

accessories, only original Rimsa spare parts are used.

3.5 Other safety conditions (secondary effects)

Optical safety

- Do not direct the light source into the patient's and/or operator's eyes.
- Obligation to adequately protect the patient's eyes.

Failure to follow such precautions could cause glare and potential damage to

the retina.

Incorrect use

- Never place and/or hang anything on the Product.

Unless this precaution is taken, positioning will not be reliable and the

danger exists of such objects falling in the operating area.

- Never hang on the Product with the body weight of a person.

Unless this precaution is taken, mechanical parts of the Product could be

damaged.

Covering the heads

- Never cover the head of the Product during operation.

Failure to comply could prevent heat exchange with the environment and the

Product could overheat.

Knocks

- Avoid knocking the rocker arms and Product head.

A violent knock could damage the Product and pieces of paint could chip off

and fall onto the operating field in the patient area.



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3.6 Graphic symbols used in this manual

In these operating instructions and on the lamp itself, important indications are marked by means of symbols and notice words.

Notice words such as HAZARD, CAUTION or IMPORTANT indicate the classification of the risk of suffering injuries.

HAZARD indicates an immediately hazardous situation which could result in death or serious injuries.

CAUTION indicates a potentially hazardous situation that could result in death or serious injuries.

IMPORTANT indicates a potentially hazardous situation which could result in moderate or light injuries.

The following triangular symbol together with the explanation alongside indicates the type of hazard to be dealt with:



Electric shock, Mechanical hazard from sprung masses (quick break of a damped arm during installation)

3.7 Other graphic symbols used on the device

Below are the symbols to be found on the Product:



B-Type device. Indicates the level of protection against direct and indirect contact



Graphic symbol proving the EC marking of the product



Symbol indicating the manufacture date (month and year)



Fuses used by the device



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4 Lamp description and operation

4.1 Description of the Product

Version

The Product is available in different versions:

- mobile version
- wall version
- ceiling single version
- ceiling double version (scialytic lamp system)

See drawing 151

MOBILE version: wheel base (1), power supply unit (2), base cover (3), switching on base (4), vertical stem (5), oscillating arm (6), fork (7), lamp head (8), switching on cupola I/O (9), sterilizable handle (10), power supply plug (11).

See drawing 30

WALL version: wall plate (1), power supply unit (2), horizontal arm (3), oscillating arm (4), fork (5), lamp head (6), switching on cupola I/O (7), sterilizable handle (8), power supply plug (9).

See drawing 32

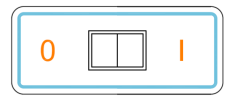
CEILING SINGLE version: ceiling cover (1), ceiling anchorage tube (2), horizontal arm (3), oscillating arm (4), fork (5), lamp head (6), switching on cupola I/O (7), sterilizable handle (8).

See drawing 79

CEILING DOUBLE version: ceiling cover (1), ceiling anchorage tube (2), horizontal arm (3), oscillating arm (4), fork (5), lamp head (6), switching on cupola I/O (7), sterilizable handle (8).

4.2 Description of the operation

The function control panel is applied to the cupola of the Product. Such panel allows to turn the lamp on and off by means of the switch I/O (1).



Light field

The mechanical regulation of the light field takes place through the rotation of the sterilizable handpiece. By rotating in a direction rather than the opposite one, the light field in the illuminated area (patient area) is enlarged or reduced.



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5 Cleaning and disinfection

5.1 Cleaning the Product



CAUTION – Electric shock hazard

Switch the Product off by means of the operating theatre main switch and make sure it cannot be switched back on.

Protect the Product against water spray and do not clean it/disinfect it with liquids.

Leave the lamp body to cool down. Only clean the lamp body when it is cold.

Clean with appropriate detergent with low alkaline content and chlorine free.

IMPORTANT Do not use abrasive products, petrol, paint thinners, alkaline detergents, acids, containing alcohol or aldehydes;

Dose the detergents so no liquids penetrate into the lamp bodies and into the support arm system.

Clean the Product with a damp but not wet cloth.

IMPORTANT

Non respecting the instructions of cleaning and disinfection could cause paint detachment with possible fall in the patient area, early deterioration of plastic parts and glass opacification.

5.2 Disinfecting



CAUTION – Electric shock hazard

Switch the Product off by means of the operating theatre main switch and make sure it cannot be switched back on.

Protect the Product against water spray and do not clean it/disinfect it with liquids.

Leave the lamp body to cool down. Only disinfect the lamp body when it is cold.

CAUTION

Disinfectants can contain substances which are harmful for the health: only use disinfectants in accordance with the rules on hygiene established by the hospital.



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The Product operator must comply with the rules established by the national commission for hygiene and disinfection.

IMPORTANT

To prevent damaging parts in stainless steel or aluminium, only use disinfectants which are chlorine and halogen free.

To prevent the plastic parts becoming fragile, use only disinfectants with low alcohol content.

Dose the disinfectants so no liquids penetrate inside the lamp bodies and into the support arm system.

Clean the Product with a damp but not wet cloth.

5.3 Sterilizing the handpieces



CAUTION – Hazard for the patient

Replace the handpieces as soon as these become cracked or deformed, as these could fall in the wound area.

The Product operator must comply with the rules of the national commission for hygiene and disinfection.

Handpiece fitting / removal:

- press the handpiece safety key and remove the handpiece.
- insert the handpiece up fast and turn it until it fastens on and rotation is blocked .

Cleaning, disinfecting and sterilizing the handpiece:

The handpieces are made of plastic material resistant to heat and knocks (PPSU).

They can be cleaned with a lightly-alkaline detergent free of active chlorine.

To disinfect the handpieces, we suggest using alcohol or aldehyde-based products. The disinfectants must be approved by the manufacturer for use on polyphenylsulfone (PPSU).

Before sterilizing, rinse the handpieces.

The handpieces can withstand about 300 steam sterilization cycles as follows:

steam sterilization at 121°C 1.3bar from 25 to 30 minutes,

or

- steam sterilization at 134°C 2.3 bar for 4 minutes.

Position the handpieces straight with open side downwards.

Do not exceed a sterilization temperature of 134°C.



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Avoid the handpieces coming into contact with other objects during the disinfection process.

Each Product, over time, is subject to a certain amount of wear. Product safety and operation must therefore be checked during inspection and maintenance intervals.

5.4 Yearly inspections by the keeper



ATTENTION

Keep to the yearly inspection schedules and inspect the product according to IEC 62353 standard.

5.5 Repairs



CAUTION – Unsuitable repairs

The Product must only be opened and repaired by a technician who has attended a course on the Product organised by the manufacturer or by a qualified technician in possession of the necessary technical skills.

6 Adjustments

6.1 Setting the rocker arm

See drawing 33

The Product is sold already balanced and does not require further adjustment. In the event of the swinging arm with spring balancing becoming stiff or loose over time, mechanical intervention is possible by regulating the compression of the internal spring.

Loosen the two stop dowels (1) which secure the cover (2) and move this forward. Fit a pin (3) with max diameter of 7 mm in the holes of the ring nut and turn in the direction indicated by the arrows to increase/decrease the load on the spring.

If the swing arm drops, this means the elastic force of the spring is insufficient:

turn the lever downwards and load the spring.



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If the swing arm continues to lift up, this means the elastic force of the spring is too high:

- turn the lever upwards and release the spring.

After making adjustments, return the covering to its original position.

6.2 Adjustment of the braking force

See drawing 33 The brakes are set during installation. As for all the mechanic parts, brakes

also are subject to wear and tear.

If the lamp body does not automatically keep the position in which it is put, it is necessary to adjust the braking force by acting on the screws of the

brakes.

Horizontal arm brakes Use a cut-suitable screwdriver to increase the braking force, rotating

clockwise the screws (4) and (5) of the horizontal arm.

Fork brakes To increase the head braking force, rotate clockwise the dowels (6 and 7) of

the brake with an Allen key.

6.3 Troubleshooting

No.	Problem	Solution	
1	The Product does not remain in position	Make sure the plate fitted on the wall (wall) is perfectly flat, that the stem is flat on the base (mobile) and that the tube secured to the ceiling (ceiling) is level. Further tighten the brakes on the joints so as to increase friction.	
2	The Product fails to work	Make sure fuses have been fitted inside the terminal board. Make sure the electrical connectors are fitted. Make sure there is power voltage in the lamp head (18/26VDC))	
3	The fuse continues to burn out	Check the specifications of the fitted fuses T1A (primary) and T6,3A (secondary) for 230Vac supply T2A (primary) and T6,3A (secondary) for 100Vac supply T2A (primary) and T10A (secondary) for battery 230Vac supply T4A (primary) and T10A (secondary) for battery 100Vac supply	
4	The light flickers and produces a stroboscopic effect	Contact the after sales service.	
5	The light beam on the operating field is not focalised (defective meeting of light fields)	Contact the after-sales service.	
6	The Product does not switch on	Check the supply power voltage and check the fuses.	



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6.4 Routine maintenance

no.	Period	Job	
1	Every 6 months	Inspect all the lamp joints and make sure they do not squeak. If they do, add white grease to the clutches involved. If the Product does not maintain the position, adjust the clutches. To determine which clutches to adjust, see point 6.2 .	
2	Once a year	Make sure the Tiges retention screws are tightened properly. Also check the 6 horizontal arm retention screws and the 3 swing arm screws. If these are not properly fastened, adequately tighten.	
3	Once a year	Check the integrity of the power cables. If they are not, proceed to correctly tighten. INTERRUPT THE POWER SUPPLY BEFORE PERFORMING THESE OPERATIONS.	
4	Once a year	Make sure the line voltage is correct. Make sure 24V are reaching the board.	
5	Once a year	Check the condition of the lamp paint. Make sure there are no paint pieces that could fall on the operating field.	

Description	Order code	
Sterilizable handle	Z200518	



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7 Technical data

Technical data on light	TRIS-LED TRIS-LED+TRIS-LED	
Illumination E _c at a distance of 1 m ± 10% [Lux] ± 10%	100.000 100.000+100.000	
Colour temperature [K] ± 5%	4.300	
Colour rendering index R _a [-]		96 / 96
R ₉		≥90
No. Leds	No.28	No.28 + No.28
Focus		Manual
Light field diameter adjustable from – to [cm]	26 - 38	26 - 38 (26 - 38)
Diameter of the light field d ₅₀ [mm]	120	120 + 120
Diameter of the light field d ₁₀ [mm]	240 240 + 240	
Illumination depth L1 + L2 at 60% [cm]	82	82+82
Maximum irradiation [W/m²]	280 280 + 280	
Irradiation / Illumination [mW/m²lx]	2,79 2,79 2,79	
Maximum irradiation in the UV [W/m²]	0,001 0,001 + 0,001	
Focusing by handle	Manual	
Data on electrical connection		
Primary alternating voltage [Volt ac]	100 ÷ 240	
Secondary continue voltage [Volt dc]	24 24	
Frequency [Hz]	50/60	
Power Absorbed [VA]	70 70 + 70	
Light source	n°28 LEDs n°28 + n°28 LEDs	
Led diode light source duration [h] (this datum can vary according to a power voltage higher than the specified one, voltage peaks and the frequency of use)	50.000	



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Light field adjustable from – to - [cm]		26-38	
General data			
Colour		RAL 9003	
Directive		2007/47/EC	
Standard		IEC 60601-2-41	
Electrical safety class		Class I	
Protection against direct and indirect contacts		B-type device	
Dimensions			
Lamp body diameter [cm]	40	40 + 40	
Light emission surface [cm²]	200	200 + 200	
Scialytic ceiling single, floor, wall, ceiling double TRIS- LED+TRIS-LED lamp weight [Kg]	35, 32, 27, 55	;	
Certificates			
CE	Complying wi	th directive 93/42/EEC and 2007/47/EC	

All lighting values are subject to a tolerance of \pm 6% due to manufacturing and metrological reasons.



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8 EMC compliance

The Product has been tested in accordance to EN60601-1-2 to ensure proper electromagnetic compatibility. Portable and mobile RF-communications equipment can affect the Product. Other products used in the vicinity of Product should also comply with this standard.

The Product is intended for use in the electromagnetic environment specified below. The customer or the user of the Product should assure that these are used in such an environment.

Emissions test	Compliance	Electromagnetic environment-guidance	
RF emissions CISPR 11	Group 1	The Product uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.	
RF emissions CISPR 11	Class A	The Product is suitable for use in all establishments other than domestic, and may be used in domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes, provided the following warning is heeded:	
Harmonic emissions IEC 61000-3-2	Class A		
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Compliance	Warning: This equipment/system is intended for use by healthcare professionals only. This equipment/system may cause radio interference or may disrupt the operation of nearby equipment. It may be necessary to take mitigation measures, such as re-orienting or relocating the Product or shielding the location	



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Immunity test	IEC 60601-1-2 test level	Compliance level	Electromagnetic environment-guidance
Electrostatic discharge (ESD) IEC 61000-4-2	+/- 6 kV contact +/- 8 kV air	+/- 6 kV contact +/- 8 kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%
Electrical fast transient / burst IEC 61000-4-4	+/- 2 kV for power supply unit +/- 1 kV for input/output lines	+/- 2 kV for power supply lines +/- 1 kV for input/output lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	+/- 1 kV differential mode +/- 2 kV common mode	+/- 1 kV differential mode +/- 2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	<5% U _T (>95% dip in U _T) For 0,5 cycle 40% of U _T (60% dip in U _T) For 5 cycles 70% of U _T (30% dip in U _T) For 25 cycles <5% U _T (>95% dip in U _T) For 5 sec	<5% U _T (>95% dip in U _T) For 0,5 cycle 40% of U _T (60% dip in U _T) For 5 cycles 70% of U _T (30% dip in U _T) For 25 cycles <5% U _T (>95% dip in U _T) For 5 sec	Mains power quality should be that of a typical commercial or hospital environment. If the user of the Product requires continued operation during power mains interruptions, it is recommended that the Product be powered from an uninterruptible power supply or battery.
Power frequency (50/60Hz) magnetic field IEC 61000-4-8	3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.



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Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment- guidance
Conducted RF IEC 61000-4-6 Radiated RF IEC 61000-4-3	3 Vrms 150 kHz to 80 MHz 3 V/m 80 MHz to 2,5GHz	3 Vrms 3 V/m	Portable and mobile RF communications equipment should be used no closer to any part of the Product, included cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter. Recommended separation distance $d = 1,2\sqrt{P}$ 150 KHz to 80 MHz $d = 1,2\sqrt{P}$ 80 MHz to 800 MHz $d = 2,3\sqrt{P}$ 80 MHz to 2,5 GHz where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacture and d is the recommended separation distance in meters (m). Field strengths from fixed transmitters, as determined by an electromagnetic site survey, should be less than the compliance leave in each frequency range. Interference may occur in the
			vicinity of equipment marked with the following symbol.



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Recommended separation distance between portable an mobile RF communications equipment and the Product

The Product is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the Product can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the Product as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter W	Separation distance according to frequency of transmitter m		
	150 kHz to 80 MHz d = 1,2P	80 MHz to 800 MHz d = 1,2P	800 MHz to 2.5 GHz $d = 2.3P$
0.01	0.12	0.12	0.24
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance *d* in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where *P* is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

Note 1: at 80 MHz and 800 MHz, the separation distance for the higher frequency range applies. **Note 2:** These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects an people.



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Notes

