





# starlight s+

**INSTALLATION MANUAL** 

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### ΕN

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#### 1 INTRODUCTION

Carefully read this manual before proceeding with installation operations, use, maintenance or other intervention operations on the device. Always keep this manual at hand.

Important: To avoid damages to persons or things, read with particular attention all the paragraphs "Safety Requirements" present in the manual.

According to the degree of severity the safety requirements are classified with the following indications:

**MARNING:** Always referred to damage to persons

**CAUTION:** Referring to possible damages to things

The aim of this manual is to make the operators aware of the safety regulations, installation procedures, instructions for proper use and maintenance of the device and its accessories.

Use of this manual for aims other than those strictly linked to the installation, use and maintenance of the device is prohibited.

The information and illustrations in this manual were updated on the edition date shown on the last page.

MECTRON is engaged in continuously updating its products with possible changes to the components of the device.

In case you encounter discrepancies between the descriptions found in this manual and the device in your possession you can:

- check for any available updates in the section MANUALS of MECTRON website<sup>1</sup>;
- ask clarifications to Your Dealer:
- contact MECTRON After Sales Service.

#### 1.1 Intended Use

Polymerisation of photo-hardening dental materials with a photoinitiator that can be activated in the wavelength band comprised between 440 and 480 nm.

Although most composite materials are activated within this wavelength range, in case of uncertainty consult the specifications of the composite material or contact the manufacturer. This device may be used only in a dentist's surgery or out-patient's department where there are no inflammable gases (anaesthetic mixtures, oxygen, etc.).

## 1.2 Description of the Device

starlight s+ is an device for polymerising photo-hardening composites.

The light source used is a very high-efficiency monochromatic LED with a dominant wavelength between 440 nm and 465 nm.

Unlike traditional halogen lamps, therefore, all the light being emitted by starlight s+ is used to activate the camphorquinone photoinitiator.

### 1.2.1 Patient Group Directions

This medical device is designed to be used with the following patient population:

- · Children;
- Adolescents:
- Adults:
- · Elderly.

This medical device can be used on any patient (if applicable) of any age, weight, height, gender and nationality.

#### 1.2.2 Patient Selection Criteria

The use of the device is not recommended in the following cases:

- Patients with active implantable medical devices (for example: pacemakers, hearing aids and/or other electromagnetic prostheses) without the prior authorization of their doctor;
- Patients with a history of light stimulation, for example in photoexposure dermatitis and/ or porphyrias, etc. or who are being treated with photosensitizing drugs. In all cases of possible risk, consult a specialist doctor;
- Patients whose medical history shows pathologies of the retina must first consult the ophthalmologist to receive authorization for treatment with the Mectron curing light.

MARNING: Adopt strict safety measures for patients who have undergone cataract surgery and are therefore particularly sensitive to light (for example, safety glasses that filter out blue light).

All models of curing lights are intended for professional use only. Therefore, the user is the only person able to decide if and how to treat their patients.

MARNING: Contraindications. In all cases of potential risk, a specialist doctor must be consulted.

#### 1.2.3 Indications for Use

The use of the device is indicated for all the intended patients (see *Chapter 1.2.1* on page 2) for whom a polymerization treatment of light-curing dental materials is prescribed, by the treating physician, within the intended use of the device (see *Chapter 1.1* on page 2).

#### 1.2.4 Users

The device must be used only by specialised and properly trained personnel, such as the dentist and/or assistant, adults of any weight, age, height, gender and nationality, able-bodied. No specific training activities are required for the use of the device.

#### 1.3 Disclaimer

The manufacturer MECTRON disclaims all responsibility, express or implied, and cannot be held responsible for direct or indirect personal injury and/or property damage, occurring as a result of incorrect procedures linked to the use of the device and its accessories.

The manufacturer MECTRON cannot be held responsible, expressly or by implication of any type of injury to persons and/or damage to things, carried out by the user of the product and its accessories and happened by way of example and not of limitation, in the following cases:

- Misuse or use during procedures other than those specified in the destination of use of the product;
- The environmental conditions for preservation and storage of the device are not complying with the requirements indicated in Chapter 8 on page 15;
- The device is not used in accordance with all the instructions and requirements described in this manual;
- The electrical system of the places where the device is used do not comply with the laws in force and the related regulations;
- Assembly operations, extensions, re-adjustments, upgrades and repairs of the device are carried out by personnel not authorized by MECTRON;
- Misuse, abuse, abnormal use, negligent use, intentional misconduct or use exceeding the limits of the device indicated and allowed and/or normal wear or deterioration, illtreatment and/or incorrect interventions:
- · Any attempt to tamper with or modification of the device under every circumstance;
- Breach of the requirements and the information contained in Chapter 5 on page 11 of the Use and Maintenance manual;
- Unauthorized repairs in accordance with the indications contained in Chapter 9 on page 24 of this manual.

## 1.4 Safety Requirements

**CAUTION:** No alterations to this device are permitted.

**CAUTION:** The electrical system of the premises in which the device is installed and used must comply with the rules in force and the relevant requirements of electrical safety.

#### **MARNING:** Qualified and specialised personnel.

The device must be used exclusively by specialized personnel with proper medical culture; no training activities are foreseen for the use of the device. The use of the device does not cause side effects if it is used correctly.

#### ⚠ WARNING: Intended use.

Use the device solely for the purpose for which it is intended (see *Chapter 1.1 on page 2*). Failure to comply with this requirement could lead to serious harm to the patient and/or to the operator and/or damage to/failure of the device.

#### **⚠ WARNING: Contraindications.**

Do not use the device on patients with Pacemakers or other implantable electronic devices. This regulation also applies to the operator.

#### ⚠ WARNING: Point the beam of light directly at the material to be polymerised.

Do not point the beam of light on the gums or other soft tissues (if necessary these parts should be suitably shielded). The effect of the light should be limited to that part of the oral cavity to be clinically treated.

## riangle WARNING: Never point the beam of light on the eyes.

The effect of the light should be limited to that part of the oral cavity to be clinically treated.

#### WARNING: Contraindications.

Do not use this device for patients who have a case history of positive reaction to stimulation by light, e.g. urticaria solaris and/or porphyria, etc. or who are receiving treatment with photosensitising drugs. In all cases of possible risk consult a specialised physician.

#### **MARNING: Contraindications.**

Adopt strict safety measures for patients who have undergone cataract surgery and who are therefore particularly sensitive to light (e.g. protective goggles able to filter out blue light).

#### ♠ WARNING: Contraindications.

Patients who have a case history of diseases of the retina should consult their optician beforehand and be specifically authorised to receive treatment with the starlight s+.

 $\triangle$  CAUTION: Photobiological safety of the curing lights and lamp systems IEC 62471.

According to the Standard IEC 62471, the device results in risk class 2 (moderate risk) concerning a retinal risk from blue light or thermal retinal risk.

The following CAUTION indications are applied to the device package.

#### Risk Group 2



**CAUTION.** Possibly hazardous optical radiation emitted from this product. Do not stare at operating lamp. May be harmful to the eye

Product tested against IEC62471

MARNING: Cleaning, disinfection and sterilisation of new or repaired products. Before treatment, all new or repaired products should be cleaned and disinfected and, if suitable for this treatment, autoclave sterilised following the instructions provided in Chapter 5 on page 11 of the Use and Maintenance manual strictly.

#### MARNING: Infection control.

In order to ensure maximum safety for both the patient and the operator, clean, disinfect and sterilise the optical fibre and the optical protection before each treatment. Follow the instructions provided in *Chapter 5 on page 11* of the Use and Maintenance manual.

MARNING: Use only original Mectron accessories and spare parts.

### ⚠ WARNING: Checking the condition of the device before treatment.

Before each treatment always check that the device is in proper working order and that the accessories are efficient. Do not carry out the treatment if any problems are encountered in operating the device. If the problems concern the device contact an authorised technical service centre.

### **⚠ WARNING: Risk of explosions.**

The device cannot operate in environments where there are saturated atmospheres of flammable gases (anaesthetic mixtures, oxygen, etc.).

⚠ **CAUTION:** In case the final user, operating in their own medical room or surgery, in order to comply with mandatory requirements, must periodically inspect the device present in the surgery, the test procedures to apply to medical electrical equipment and medical electrical systems for the safety assessment must be carried out following the standard EN 62353 'Medical electrical equipment - Periodic inspections and tests to be carried out after repair of medical electrical equipment'. The interval for periodic checks, in the intended operating conditions and described in this "Use and Maintenance" manual, is one year or 2000 hours of use, depending on which of these two conditions occurs first.

MARNING: If an adverse event and/or serious incident attributable to the device occurs during correct and intended use, it is recommended to report it to the Competent Authority and to the manufacturer indicated on the product label.

## 2 IDENTIFICATION DATA

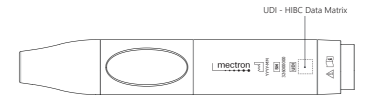
A correct description of the model and of the serial number of the device will allow the After Sales Service to provide fast and effective answers.

Always provide this information every time that you contact MECTRON After Sales Service.

## 2.1 Identification Data of the Handpiece

On the handpiece are laser marked the traceability data, UDI code included.

**NOTE:** The complete list of symbols and their description are shown in *Chapter 6 on page 12.* 





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## 3 DELIVERY

The packaging of the device cannot undergo strong impacts as contains electronic components, therefore the transport and the storage must be carried out with particular care.

All the material shipped by MECTRON is controlled at the time of dispatch.

The device is shipped appropriately protected and packaged.

Upon receipt of the device, check for any possible damage caused during transport and in case any damage and/or defects is found, complain to the transporter.

Keep the packaging in case there is a necessity to send any item to a MECTRON Authorized Service Centre and to store the device during long periods of inactivity.

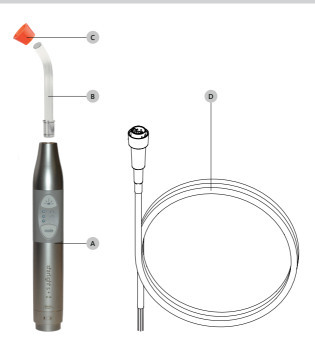
## 3.1 List of Components

Refer to following Figure:

- A. 1 starlight s+ handpiece;
- B. 1 Optical fibre;
- C. 1 Optical protection;
- D. Cord and connector.

These components can also be ordered separately.

**NOTE:** The components included in the package may vary in occasion of promotional campaigns.



## EN

## 4 INSTALLATION

## 4.1 Safety Requirements in the Installation Phase

**WARNING:** The electrical system of the premises in which the device is installed and used must comply with the rules in force and the relevant requirements of electrical safety.

⚠ **WARNING: Risk of explosion.** The device cannot operate in environments where there are saturated atmospheres of flammable gases (aesthetic mixtures, oxygen, etc.).

 $\triangle$  **WARNING:** Install the device in a safe place protected from impact or accidental water or liquid spray.

**MARNING:** Do not install the device above or near sources of heat. Arrange during installation for a suitable circulation of air around the device.

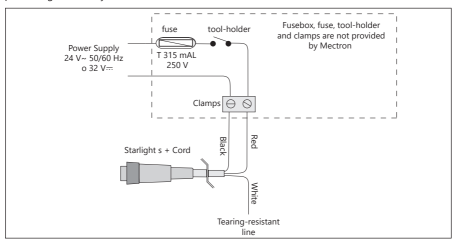
**CAUTION:** Do not expose the device to direct sunlight or sources of UV light.

**MARNING:** The device electricity supply voltage must comply with the plate characteristics.

**CAUTION:** If long supply connections are necessary, use appropriate section wires of at least 0.5 mm<sup>2</sup>.

#### 4.2 Device Connection

The device shall be installed on a dental chair. We recommend to apply the wiring scheme as shown in the following figure, that is featured with 24  $V_{\sim}$  or 32  $V_{\sim}$  supplied by a power supplier with double insulation compliant to IEC/EN 60601-1, controlled by the switch on the quiver and protecting the line by a fuse.



**NOTE:** The device is not equipped with a disconnection device from the power network according to clause 8.11.1 of IEC/EN 60601-1, as it is a component that will be installed on a dental chair in its final configuration. Such a device must be designed for the final installation and the manufacturer of the dental chair shall take care of assuring the compliance to this requirement.

## 4.3 Descriptions of Commands and Signalling

For the description of commands and signalling, refer to following figure and tables.

Ref.	Name	Description	
А	Start button	Starts or stops a polymerisation cycle.	
В	<b>Mode</b> button	Allows selecting the exposure type. By repeatedly pressing the button, in the order are selected: FAST, SLOW RISE, SOFT	000
С	Green Mode LED	Indicates the selected exposure. From left to right: FAST, SLOW RISE, SOFT	



Selected function via Mode button	Signalling	Exposure Type	Exposure Time
FAST Polymerisation	Left LED ON	Maximum power emitted.	10 sec. 20 sec.
SLOW RISE Polymerisation	Middle LED ON	Increased emission during first two seconds. Maximum emission power	10 sec. 20 sec.
SOFT Polymerisation	Right LED ON	Emission 70% of maximum power for whole cycle	10 sec. 20 sec.

## 5 DISPOSAL PROCEDURES AND PRECAUTIONS

- · This device must be disposed of and treated as waste requiring separate collection;
- At the end of the life-cycle of this device, the purchaser is entitled to return the device
  to the dealer supplying new device. Instructions for disposal are available from Mectron
  S.p.A.;
- Failure to comply with the foregoing points may entail punishment in accordance with Directive about waste of electrical and electronic equipment WEEE.

#### **⚠ WARNING: Hospital waste.**

Treat the following items as hospital waste:

- · Optical fibre, when worn or broken;
- Optical protection, when worn or broken.

## 6 SYMBOLS

Symbol	Description	Symbol	Description
C€	Class I device (torque wrench) compliant with Regulation (EU) 2017/745.	Clectrical Safety ES 60601-1	Nemko Mark UL - CSA standards compliance
MD	Medical Device	<u> </u>	Caution!
[]i	Consult instructions for use or consult electronic instruction for use		Manufacturer
	Date of manufacture	SN	Serial Number
LOT	Batch Number	REF	Catalogue number
NON	Non-sterile	135°C	Sterilizable up to a max. temperature of 135°C
$\sim$	Alternating current	===	Direct current
	The appliance and its accessories must not be disposed of or treated as municipal solid waste	1/2	Starts or stops a polymerisation cycle
*	Type B applied part		Temperature limit
<u></u>	Humidity limitation	<b>\$•</b> \$	Atmospheric pressure limitation
<u> </u>	Generic warning signal <sup>a)</sup>	IP 44	Protection degree provided by the casing (IEC/EN 60529)
UDI	Unique Device Identifier	#	Model number
HIBC	Health Industry Bar Code	1	Fragile
<b>*</b>	Keep dry		

a) The symbol is represented by a yellow warning triangle and a black graphical symbol.

## 7 TROUBLESHOOTING

## 7.1 Quick Troubleshooting Guide

If the device appears not to be working correctly, read the instructions again and then check the following Table.

Problem	Possible Cause	Solution
An acoustic signal (3 beeps) is produced during the exposure cycle and at the end of the cycle starlight s+ will not allow any further treatment to be carried out. The selected function LED flashes.	The thermal protection has been activated.	It will be possible to use the equipment only after it has cooled down. Wait approximately 10 minutes before re-using the device.
When the 'start' button is pressed, an acoustic warning sounds (4 beeps) and the device does not emit light.	The temperature control circuit reports a failure.	Contact a Mectron service centre.
When the 'start' button is pressed, the mode selection LED flashes. The device does not emit light, or the intensity of the light emitted is unsuitable.	The control circuit reports an LED failure.	Contact a Mectron service centre.
The polymerisation is insufficient.	The terminal surface of the tip of the optical fibre is soiled.	See Chapter 5 on page 11 of the Use and Maintenance manual.

## 7.2 Diagnostic System and Possible Solutions

The device is provided with a diagnostic circuit that allows identifying the main operating issues.

Signalling Type	Description	Solution
4 Веер	The temperature control circuit is failed. The device does not emit any light.	Contact a Mectron service centre.
3 Веер	The control circuit indicates that the maximum LED temperature has been reached. The device does not emit any light.	Place the device back in its housing and wait about 10 minutes before using it again.
Mode LED flashing	The control circuit reports an LED failure. The device does not emit any light, or the intensity of the emitted light is very low.	Contact a Mectron service centre.

## 7.3 How to Prepare the Device and the Accessories Before Sending them to Repair

In the event that there is a need to send the device, the optical fiber and accessories to an authorized Mectron service center, we invite our kind customers to respect the good code of conduct reported in the following:

- Clean the device, optic fibre and accessories according to the instructions reported in the Chapter 5 on page 11 of the Use and Maintenance manual;
- 2. Sterilse the sterilisable parts according to the instructions reported in the Chapter 5 on page 11 of the Use and Maintenance manual:
  - · Optic fibre;
  - Optical protection.
- 3. Leave the sterilised parts in their sterilization pouches that certify the sterilization completion;
- 4. If the device is still under warranty, attach a copy of the purchase document;
- 5. Whenever possible, send all the parts in their original packaging or, alternatively, in proper packages, to avoid any damage during the transportation.

The above requirements (points 1 and 2) are in compliance with the current requirements concerning the protection of health and safety in the workplace, as per Italian legislative decrees 81/08 and subsequent amendments.

In the event that the customer does not comply with the requirements (points 1 and 2), Mectron reserves the right to charge him/her for cleaning and sterilization costs or to reject the items for repair received in unsuitable conditions.

## 8 TECHNICAL SPECIFICATIONS

Device compliant with Regulation (EU) 2017/745	Class I
Classification under the IEC/EN 60601-1	The class definition is delegated to the manufacturer of the dental chair incorporating the device.  Applied part type B (Optical Fibre) IP 44 (Device)
Essential performances	According to IEC 80601-2-60 Standard, the device does not provide essential performances.
Handpiece for intermittent operation	60" ON 60" OFF - Max 3 consecutive cycles
Voltage	Power supply compliant to IEC/EN 60601-1 with double insulation. 24 V $\sim$ 50/60 Hz or 32 V $\stackrel{\text{\tiny FFF}}{=}$
Power Consumption	11 VA
Light source	High power LED with optic. Dominant wavelength: 440 - 465 nm LED in Class 2 (IEC 62471) retinal risk from blue light or thermal retinal risk.
Optical fibre	Diameter 8 mm. Composition: Drawn coherent fibres surfused with transparent quartz. Autoclave sterilisable (max. temp. 135 °C for 20 minutes - max. 500 cycles).
Exposures	FAST: Exposure time 10/20-seconds Acoustic signal at the beginning and at the end of an exposition cycle; in case of 20 seconds there will be also a signal after 10 seconds of exposition.  SLOW RISE: Exposure time 10/20-seconds Gradual increase during the first 2 seconds. Acoustic signal at the beginning and at the end of an exposition cycle; in case of 20 seconds there will be also a signal after 10 seconds of exposition.  SOFT: Exposure time 10/20-seconds Light emission at 70% of maximum power. Acoustic signal at the beginning and at the end of an exposition cycle; in case of 20 seconds there will be also a signal after 10 seconds of exposition. The cycles can be stopped or repeated at any time.
Operating conditions	from 10 °C to 35 °C Relative Humidity from 30% to 75% Air pressure P: 800 hPa/1060 hPa

Transport and storage conditions	from -10 °C to 60 °C Relative Humidity from 10% to 90% Air pressure P: 500 hPa/1060 hPa		
Altitude	less than or equal to 2000 meters		
Weights and dimensions	Handpiece: Weight 102 g L 141 mm Max. Ø 23 mm		

## 8.1 Electromagnetic Compatibility IEC/EN 60601-1-2

#### **⚠ WARNING: Contraindications. Interference with other equipment**

Even if compliant with standard IEC/EN 60601-1-2, starlight s+ may interfere with other devices in the vicinity. starlight s+ must not be used near or stacked with other devices. However, if this is necessary, the correct operation of the device in that configuration, and of all equipment, must be checked and monitored before starting the intervention.

**WARNING:** Portable and mobile radio communication equipment may influence the correct operation of the device.

#### MARNING: Contraindications. Interference from other equipment

An electrosurgical scalpel or other electrosurgical units positioned near the starlight s+ device may interfere with the correct operation of the device itself.

MARNING: The device requires particular EMC precautions and must be installed and put into service according to the EMC information provided in this chapter.

**WARNING:** The use of cables and accessories not supplied by MECTRON may adversely affect the EMC performances.

## 8.2 Guide and Manufacturer's Declaration - Electromagnetic Emissions

starlight s+ is designed to operate in the electromagnetic environment specified below. The purchaser or user of starlight s+ should ensure that it is used in such an environment.

<b>Emissions Test</b>	Compliance	Electromagnetic Environment Guidance
RF Emissions CISPR 11	Group 1	starlight s+ uses RF energy only for its internal operation. Therefore, its RF emissions are very low and probably do not cause any interference with nearby electronic devices.
RF Emissions CISPR 11	Class B	starlight s+ is suitable for use in all buildin-
Harmonic emissions IEC 61000-3-2	Class A	gs, including domestic buildings, and those directly connected to the public low-voltage
Emissions of fluctuations voltage/flicker IEC 61000-3-3	Compliant	power supply network that supplies buildings used for domestic purposes.

## 8.3 Accessible Parts of the Casing

starlight s+ is designed to operate in the electromagnetic environment specified below. The purchaser or user of starlight s+ should ensure that it is used in such an environment.

Phenomenon	Essential EMC standard or test method	Immunity test values	Electromagnetic Environment Guidance
Electrostatic discharge (ESD)	IEC 61000-4-2	±8 kV on contact ±2 kV, ±4 kV, ±8 kV, ±15 kV in air	The floor must be made of wood, concrete or ceramic tiles. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Radiated RF EM fields <sup>a)</sup>	IEC 61000-4-3	3 V/m <sup>f)</sup> 80 MHz - 2,7 GHz <sup>b)</sup> 80% AM a 1 kHz <sup>c)</sup>	Portable and mobile RF communication devices should not be used near
Proximity fields from RF wireless communications equipment	IEC 61000-4-3	See Chapter 8.5 on page 22	any part of the product, including cables, except when they respect the recommended and calculated distances from the equation applicable at the frequency of the transmitter.
RATED power frequency magnetic fields <sup>d)</sup>	IEC 61000-4-8	30 A/m 50 Hz or 60 Hz	Power supply frequency magnetic fields should have levels characteristic of a typical location in a commercial or hospital environment.
Proximity magnetic fields	IEC 61000-4-39	See Chapter 8.6 on page 23	Portable and mobile RF communication devices shall be used with a separation distance of at least 0,15 m from the field sources.

- a) The interface between the PATIENT physiological signal simulation, if used, and the device shall be located within 0,1 m of the vertical plane of the uniform field area in one orientation of the device.
- b) The device that intentionally receives RF electromagnetic energy for the purpose of its operation shall be tested at the frequency of reception. Testing may be performed at other modulation frequencies identified by the RISK MANAGEMENT PROCESS. This test assesses the BASIC SAFETY and ESSENTIAL PERFORMANCE of an
- intentional receiver when an ambient signal is in the passband. It is understood that the receiver might not achieve normal reception during the test.
- Testing may be performed at other modulation frequencies identified by the RISK MANAGEMENT PROCESS
- Applies only to devices with magnetically sensitive components or circuitry.
- e) Void
- f) Before modulation is applied.

## 8.4 Guide and the Manufacturer's Declaration - Electromagnetic Immunity

## 8.4.1 Power Connection A.C. Input

starlight s+ is designed to operate in the electromagnetic environment specified below. The purchaser or user of starlight s+ should ensure that it is used in such an environment.

Phenomenon	Essential EMC standard or test method	Immunity test values	Electromagnetic Environment Guidance
Electrical fast transients / bursts <sup>I) (o)</sup>	IEC 61000-4-4	±2 kV on contact 100 KHz repetition frequency	The quality of the network voltage should be that of a typical commercial or hospital environment.
Surges <sup>b) j) o)</sup> Line-to-line	IEC 61000-4-5	± 0.5 kV, ± 1 kV	The quality of the network voltage should be that of a typical commercial or hospital environment.
Surges b) j) k) o) Line-to- ground	IEC 61000-4-5	± 0.5 kV, ± 1 kV, ± 2kV	The quality of the network voltage should be that of a typical commercial or hospital environment.
Conducted disturbances induced by RF fields () () ()	IEC 61000-4-6	3 V <sup>m)</sup> 0.15 MHz - 80 MHz 6 V <sup>m)</sup> in the ISM bands between 0.15 MHz and 80 MHz <sup>n)</sup> 80 % AM at 1 KHz <sup>e)</sup>	Portable and mobile RF communication devices should not be used near any part of the product, including cables, except when they respect the recommended and calculated distances from the equation applicable at the frequency of the transmitter.
Voltage dips	IEC 61000-4-11	0% UT; 0,5 cycle <sup>9</sup> At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°	The quality of the network voltage should be that of a
9 P) 9		0 % UT; 1 cycle e 70 % UT; 25/30 cycle <sup>h)</sup> Single phase: at 0°	typical commercial or hospital environment.
Voltage interruptions	IEC 61000-4-11	0% UT; 250/300 cycle <sup>h)</sup>	The quality of the network voltage should be that of a typical commercial or hospital environment.

ΕN

- a) Void.
- b) All device cables are attached during the test.
- c) Calibration for current injection clamps shall be performed in a 150  $\Omega$  system.
- d) If the frequency stepping skips over an ISM or amateur band, as applicable, an additional test frequency shall be used in the ISM or amateur radio band. This applies to each ISM and amateur radio band within the specified frequency range.
- Testing may be performed at other modulation frequencies identified by the RISK MANAGEMENT PROCESS.
- f) ME EQUIPMENT and ME SYSTEMS with a d.c. power input intended for use with a.c.-to-d.c. converters shall be tested using a converter that meets the specifications of the MANUFACTURER of the ME EQUIPMENT or ME SYSTEMS. The IMMUNITY TEST LEVELS are applied to the a.c. power input of the converter.
- g) Applicable only to the device connected to singlephase a.c. mains.
- E.g. 10/12 means 10 periods at 50 Hz or 12 periods at 60 Hz.
- i) ME EQUIPMENT and ME SYSTEMS with RATED input current greater than 16 A / phase shall be interrupted once for 250/300 cycles at any angle and at all phases at the same time (if applicable). The device with battery backup shall resume line power operation after the test. For ME EQUIPMENT and ME SYSTEMS with RATED input current not exceeding 16 A, all phases shall be interrupted simultaneously.

- ME EQUIPMENT and ME SYSTEMS that does not have a surge protection device in the primary power circuit may be tested only at ± 2 kV line(s) to earth and ± 1 kV line(s) to line(s).
- k) Not applicable to CLASS II device.
- l) Direct coupling shall be used.
- m) R.M.S., before modulation is applied.
- n) The ISM (industrial, scientific and medical) bands between 0.15 MHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz. The amateur radio bands between 0.15 MHz and 80 MHz are 1.8 MHz to 2.0 MHz, 3.5 MHz to 4.0 MHz, 5.3 MHz to 5.4 MHz, 7 MHz to 7.3 MHz, 10.1 MHz to 10.15 MHz, 14 MHz to 14.2 MHz, 18.07 MHz to 18.10 MHz, 21.0 MHz to 21.4 MHz, 24.89 MHz to 24.99 MHz, 28.0 MHz to 29.7 MHz and 50.0 MHz to 54.0 MHz.
- Applicable to ME EQUIPMENT and ME SYSTEMS with RATED input current less than or equal to 16 A / phase and ME EQUIPMENT and ME SYSTEMS with RATED input current greater than 16 A / phase.
- Applicable to ME EQUIPMENT and ME SYSTEMS with RATED input current less than or equal to 16 A / phase.
- q) At some phase angles, applying this test to ME EQUIPMENT with transformer mains power input might cause an overcurrent protection device to open. This can occur due to magnetic flux saturation of the transformer core after the voltage dip. If this occurs, the ME EQUIPMENT shall provide BASIC SAFETY during and after the test.
- r) For ME EQUIPMENT and ME SYSTEMS that have multiple voltage settings or auto ranging voltage capability, the test shall be performed at the power input voltage specified in Table 1 - "Power input voltages and frequencies during the tests" of the IEC 60601-1-2:2014/AMD1:2020.

#### 8.4.2 Points of Contact with the Patient

starlight s+ is designed to operate in the electromagnetic environment specified below. The purchaser or user of starlight s+ should ensure that it is used in such an environment.

Phenomenon	Essential EMC standard or test method	Immunity test values	Electromagnetic Environment Guidance
Electrostatic discharges (ESD) c)	IEC 61000-4-2	±8 kV on contact ±2 kV, ±4 kV, ±8 kV, ±15 kV in air	The floor must be made of wood, concrete or ceramic tiles. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Conducted disturbances induced by RF fields <sup>a)</sup>	IEC 61000-4-6	3 V <sup>b)</sup> 0.15 MHz - 80 MHz 6 V <sup>b)</sup> in ISM bands between 0.15 MHz and 80 MHz 80 % AM at 1 KHz	Portable and mobile RF communication devices should not be used near any part of the product, including cables, except when they respect the recommended distances, calculated from the equation applicable at the frequency of the transmitter.

- a) The following apply:
  - All PATIENT-COUPLED cables shall be tested, either individually or bundled
  - PATIENT-COUPLED cables shall be tested using a current clamp unless a current clamp is not suitable.
     In cases where a current clamp is not suitable, an EM clamp shall be used.
  - No intentional decoupling device shall be used between the injection point and the PATIENT COUPLING POINT in any case.
  - Testing may be performed at other modulation frequencies identified by the RISK MANAGEMENT PROCESS.
  - Tubes that are intentionally filled with conductive liquids and intended to be connected to a PATIENT shall be considered to be PATIENT-COUPLED cables.
  - If the frequency stepping skips over an ISM or amateur radio band, as applicable, an additional test

- frequency shall be used in the ISM or amateur radio band. This applies to each ISM and amateur radio band within the specified frequency range.
- The ISM (industrial, scientific and medical) bands between 0.15 MHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz. The amateur radio bands between 0.15 MHz and 80 MHz are 1.8 MHz to 2.0 MHz, 3.5 MHz to 4.0 MHz, 5.3 MHz to 5.4 MHz, 7 MHz to 7.3 MHz, 10.1 MHz to 10.15 MHz, 14 MHz to 14.2 MHz, 18.07 MHz to 18.17 MHz, 21.0 MHz to 22.0 MHz, 24.89 MHz to 24.99 MHz, 28.0 MHz to 29.7 MHz and 50.0 MHz to 54.0 MHz.
- b) R.M.S., before modulation is applied.
- c) Discharges shall be applied with no connection to an artificial hand and no connection to PATIENT simulation. PATIENT simulation may be connected after the test as needed in order to verify BASIC SAFETY and ESSENTIAL PERFORMANCE.

#### 8.4.3 Parts Accessible to the Input / Output Signals

starlight s+ is designed to operate in the electromagnetic environment specified below. The purchaser or user of starlight s+ should ensure that it is used in such an environment.

Phenomenon	Essential EMC standard or test method	Immunity test values	Electromagnetic Environment Guidance
Electrostatic discharges (ESD) <sup>e)</sup>	IEC 61000-4-2	±8 kV on contact ±2 kV, ±4 kV, ±8 kV, ±15 kV in air	The floor must be made of wood, concrete or ceramic tiles. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transients / bursts <sup>b) f)</sup>	IEC 61000-4-4	±1 kV on contact 100 KHz repetition frequency	The quality of the network voltage should be that of a typical commercial or hospital environment.
Surges Line- to-ground <sup>a)</sup>	IEC 61000-4-5	± 2kV	The quality of the network voltage should be that of a typical commercial or hospital environment.
Conducted disturbances induced by RF fields d) g) j) k)	IEC 61000-4-6	3 V <sup>h)</sup> 0.15 MHz - 80 MHz 6 V <sup>h)</sup> in the ISM bands between 0.15 MHz and 80 MHz <sup>i)</sup> 80% AM a 1 KHz <sup>c)</sup>	Portable and mobile RF communication devices should not be used near any part of the product, including cables, except when they respect the recommended distances, calculated from the equation applicable at the frequency of the transmitter.

- This test applies only to output lines intended to connect directly to outdoor cables.
- b) SIP/SOPS whose maximum cable length is less than 3 m in length are excluded.
- Testing may be performed at other modulation frequencies identified by the RISK MANAGEMENT PROCESS.
- d) Calibration for current injection clamps shall be performed in a 150  $\Omega$  system.
- e) Connectors shall be tested per 8.3.2 and Table 4 of IEC 61000-4-2:2008. For insulated connector shells, perform air discharge testing to the connector shell and the pins using the rounded tip finger of the ESD generator, with the exception that the only connector pins that are tested are those that can be contacted or touched, under conditions of INTENDED USE, by the standard test finger shown in Figure 6 of the general standard, applied in a bent or straight position.
- f) Capacitive coupling shall be used.
- g) If the frequency stepping skips over an ISM or amateur radio band, as applicable, an additional test frequency shall be used in the ISM or amateur radio

band. This applies to each ISM and amateur radio band within the specified frequency range.

- h) R.M.S., before modulation is applied.
- i) The ISM (industrial, scientific and medical) bands between 0.15 MHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz. The amateur radio bands between 0.15 MHz and 80 MHz are 1.8 MHz to 2.0 MHz, 3.5 MHz to 4.0 MHz, 5.3 MHz to 5.4 MHz, 7 MHz to 7.3 MHz, 10.1 MHz to 10.15 MHz, 14 MHz to 14.2 MHz, 18.07 MHz to 18.17 MHz, 21.0 MHz to 21.4 MHz, 24.89 MHz to 24.99 MHz, 28.0 MHz to 29.7 MHz and 50.0 MHz to 54.0 MHz.
- j) See IEC 61000-4-6:2013, Annex B, for modified start frequency versus cable length and equipment size.
- k) SIP/SOPS whose maximum cable length is less than 1 m are excluded.

# 8.5 Specifications of the tests for the Immunity of the Accessible Parts of the Casing to the Wireless RF Communications Device

starlight s+ is designed to operate in an electromagnetic environment in which radiated RF disturbances are under control. The purchaser or operator of starlight s+ can help prevent electromagnetic interferences by guaranteeing a minimum distance between the mobile and portable RF communication devices (transmitters) and starlight s+, as recommended below, in relation to the maximum output power of the radio communication devices.

Test Freq. (MHz)	Band (MHz)	Service	Modulation	Max power (W)	Distance (m)	Immunity test level (V/m)
385	380 - 390	TETRA 400	Pulse modulation <sup>b)</sup> 18 Hz	1.8	0.3	27
450	430 - 470	GMRS 460 FRS 460	FM <sup>c)</sup> ± 5 kHz deviation 1 kHz sine	2	0.3	28
710			Pulse			
745	704 - 787	LTE band 13, 17	modulation <sup>b)</sup>	0.2	0.3	9
780		.,	217 Hz			
810		GSM 800/900				
870	800 - 960	TETRA 800 iDEN 820	Pulse modulation <sup>b)</sup>	2	0.3	28
930	000 300	CDMA 850 Band LTE 5	18 Hz		0.5	20
1720		GSM 1800				
1845		CDMA 1900 GSM 1900	Pulse			
1970	1700 - 1990	DECT LTE Band 1, 3, 4, 25 UMTS	modulation b) 217 Hz	2	0.3	28
2450	2400 - 2570	Bluetooth WLAN 802.11 b/g/n RFID 2450 Band LTE 7	Pulse modulation <sup>b)</sup> 217 Hz	2	0.3	28
5240			Pulse			
5500	5100 - 5800	WLAN 802.11 a/n	modulation <sup>b)</sup>	0.2	0.3	9
5785		302 4,71	217 Hz			

For some services, only uplink frequencies are included.

The carrier shall be modulated using a 50 % duty cycle square wave signal.

c) As an alternative to FM modulation, the carrier may be pulse modulated using a 50 % duty cycle square wave signal at 18 Hz. While it does not represent actual modulation, it would be worst case.

**NOTE**: If necessary to achieve the IMMUNITY TEST LEVEL, the distance between the transmitting antenna and the device may be reduced to 1 m. The 1 m test distance is permitted by IEC 61000-4-3.

⚠ WARNING: Portable RF communication equipment (including peripheral devices such as antenna cables and external antennas) must not be used closer than 30 cm to any part of the starlight s+ device, including the cables specified by the manufacturer. Otherwise, there may be a performance degradation of these devices.

## 8.6 Immunity to Proximity Magnetic Fields in the Frequency Range 9 kHz to 13,56 MHz

The following table reports the test specifications for ENCLOSURE PORT IMMUNITY to proximity magnetic fields in the frequency range 9 kHz to 13,56 MHz.

Test Frequency	Modulation	Immunity test level (A/m)
30 kHz <sup>a)</sup>	CW	8
134,2 kHz	Pulse modulation b) 2,1 kHz	65 <sup>c)</sup>
13,56 MHz	Pulse modulation <sup>b)</sup> 50 kHz	7,5 <sup>c)</sup>

This test is applicable only to devices intended for use in the HOME HEALTHCARE ENVIRONMENT.

- The carrier shall be modulated using a 50% duty cycle square wave signal.
- c) r.m.s., before modulation is applied.

### 9 WARRANTY

Before being marketed, all MECTRON devices, are subjected to a thorough final check that verifies the full functionality.

MECTRON warrants its products, purchased new from a MECTRON dealer or importer, against defects in material and workmanship for a period of 2 (TWO) YEARS from the date of purchase. During the period of validity of the warranty, MECTRON undertakes to repair (or, at his free choice, replace) free of charge those parts of products that in their opinion prove being defective. Complete replacement of Mectron products is excluded.

MECTRON disclaims any responsibility for direct or indirect damage to people or things, in the following cases:

- The device is not used according to the intended use for which it is provided;
- The device is not used in accordance with all the instructions and requirements described in this manual;
- The electrical system of the places where the device is used do not comply with the laws in force and the related regulations;
- Assembly operations, extensions, re-adjustments, modifications, replacements and repairs
  are carried out by personnel not authorized by Mectron or in breach of what is provided
  in this manual also in regard to the origin of the authorised material;
- The environmental conditions for preservation and storage of the device are not complying with the requirements indicated in Chapter 8 on page 15;
- The installation or the transport of the device is not performed as specified in this manual
  or in other documentation provided by MECTRON, or in any case available on the website
  of the latter:
- The device or the component thereof is purchased from a subject not authorised by MECTRON:
- The device, including its subcomponents, parts or assemblies, are altered or modified with respect to what is provided in this manual.
- Accident, misuse, abuse, abnormal use, negligent use, misconduct or intentional use
  exceeding the limits recommended and allowed by the device or in the case of normal
  wear or deterioration of the same.
- If the defect or non-conformity are not promptly and readily communicated in writing to MECTRON as specified in this manual.
- If the damage, costs or expenses are caused by events of force majeure.
- The connection of the device has been carried out at a voltage different from the one envisaged, including WARNING lights, knobs and all the accessories.

In any case, MECTRON case will not recognize indemnity or compensation for loss of use, inconvenience, loss of profits, loss of business, business opportunities lost, damage to reputation, and any incidental or consequential damages arising out of or relating to the device.

The expected service life of the device is 5 years, minimum.

The service life/duration does not define a limit of use; the service life of the device defines the period of time, subsequent to installation and/or commissioning, during which the original performances or, in any case, adequate for the intended use are guaranteed, without any degradation occurring such as to compromise its functionality and reliability.

The service life is a minimum qualitative target of the design, therefore, it is not excluded that single parts or components guarantee performances and reliability higher than those declared by the manufacturer.

The service life is intended in compliance with the maintenance plans provided for in this manual, it does not include components normally subjected to "wear" and it is independent of the warranty period: the service life period does not establish any implicit or explicit extension of the warranty period.

#### CAUTION

The warranty starts from the date of purchase of the device, which evidence is given by the delivery note/purchase invoice issued by the Dealer / Importer or, in case of device with activation code, from the date of activation of the same.

In order to avail of the warranty service, the customer must return, at its own expense, the device to be repaired to the MECTRON Dealer / Importer from which they purchased the product. The device must be returned together with the original packaging, accompanied by all the accessories and by a form containing:

- · The data of the owner and telephone number;
- The data of the Dealer / Importer;
- Photocopy of the delivery note/purchase invoice of the device by the owner where are reported the date, the name of the device and the serial number;
- · Description of the failure.

The transport and the damage caused by transport are not covered by the warranty.

ΕN

	ill: mectron@mectron.com +39 0185 351374						
MECTRON S.p.A. Via Loreto 15/A 16042 Carasco (Ge), Italy							
Í	C /	our address / Ihre Adresse / Votre adresse / Вашият адрес / Vaše adresa Din adresse / Η διεύθυνσή σας / Su dirección / Teie aadress / Vaša adresav Cín Vostro indirizzo / Jūsų adresas / Jūsu adrese / Uw adres / Państwa adres / Seundereço / Adresa dumneavoastră / Din adress					
EN	Please send me, free of charged below):	ge, a copy of the Instructions for Use of the following product (please complete					
DE	Bitte senden Sie mir eine kost	enfreie Gebrauchsanweisung des folgenden Produktes zu (bitte unten ausfüllen)					
FR	Veuillez me faire parvenir gr dessous) :	atuitement une notice d'utilisation pour le produit suivant (veuillez remplir ci					
BG	Моля, изпратете ми безпла долу):	тно ръководство за употреба за следния продукт на (моля, попълнете по					
CS	Zašlete mi prosím zdarma ná	vod k použití následujícího výrobku (vyplňte laskavě dole):					
DA	Send mig venligst en gratis brugsvejledning til efterfølgende produkt (udfyld nedenfor):						
EL	Παρακαλώ να μου στείλετε (συμπληρώστε κάτω):	ε δωρεάν οδηγίες χρήσης και συναρμολόγησης του ακόλουθου προϊόντο					
ES	Rogamos nos envíen gratuita producto (por favor, rellenar	amente una copia impresa del manual de instrucciones para uso del siguiento abajo):					
ET	Palun saatke mulle tasuta kas	sutusjuhend järgmise toote kohta (palun täitke altpoolt):					
HR	Molim, pošaljite mi besplatne	e upute za uporabu sljedećeg proizvoda (ispuniti u nastavku):					
HU	Kérem, küldjenek ingyenes h	asználati utasítást a következő termékről (kérjük, töltse ki):					
IT	Vogliate inviarmi gratuitamer	nte le istruzioni per l'uso del seguente prodotto (compilare la parte sottostante)					
LT	Atsiųskite man nemokamą ši	o gaminio naudojimo instrukciją (užpildykite apačioje):					
LV	Lūdzu atsūtīt man produkta k	pezmaksas lietošanas instrukciju (aizpildīt zemāk):					
NL	Stuur mij a.u.b. een gratis gel	oruikshandleiding van het volgende product (a.u.b. hieronder invullen):					
PL	Proszę o przysłanie mi bezpła	atnej instrukcji obsługi następującego produktu (proszę uzupełnić na dole):					
PT	Enviem-me gratuitamente u baixo):	m exemplar das Instruções de utilização do seguinte produto (preencher en					
RO	Vă rog să îmi trimiteți un exe să completați datele de mai j	emplar gratuit din instrucțiunile de utilizare pentru următorul produs (vă rugăn os):					
SV	Skicka en kostnadsfri bruksar	nvisning för följande produkt (fyll i nedan):					
		ione del prodotto / Important information for product ordering: escription (e.g. combi touch)					
REF	(e.g. 05120065)						
SN	(e.g. 423001234)						







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