



PAGE INTENTIONALLY LEFT BLANK

MANUAL PRESENTATION

Technical Name: Dental Lights
Commercial Name: Dental Light

Models:

Sirius G8 Sensor 5 LEDs Sirius G8 Sensor 3 LEDs Sirius G8 Sensor 5 (3X2) LEDs

Brand: Dabi Atlante

Basic UDI-DI: 78995813DentalLight013CT, 78995813DentalLight010CM, 78995813DentalLight014CV

Technical Responsible: Daniel R. de Camargo

CREA-SP: 5062199650

ANVISA Registration No.: 10069210069



Alliage S/A Industrias Médico Odontológica Rodovia Abrão Assed, Km 53 + 450m - CEP 14097-500 Ribeirão Preto - SP - Brazil Phone: +55 (16) 3512-1212



CINTERQUAL – Soluções de Comércio Internacional, Lda. Avenida Defensores de Chaves, Nº 4 Escritório Idea Spaces 1000-117 Lisboa, Portugal



77000000969 - Rev.: 06 - November/22

Document originally written in Portuguese.

TRADEMARKS

All terms mentioned in this manual that are known trademarks, registered trademarks or service marks have been appropriately labeled as such. Further products, services or terms that are mentioned in this manual may be registered marks, registered trademarks or service marks of their respective owners. Alliage S/A makes no claims regarding these trademarks. The use of a term in this manual should not be considered to influence the validity of any registered mark, registered trademark or service mark.

Copyright © 2019 Alliage S/A. All rights reserved.

The performance characteristics provided in this manual are for reference only and should not be considered as guaranteed specifications.

TABLE OF CONTENTS

1.	GENERAL INFORMATION	8
1.1.	DEAR COSTUMER	. 8
1.2.	INDICATONS FOR USE	. 8
1.3.	CONTRAINDICATION	. 8
1.4.	SYMBOLOGY	. 8
2.	WARNINGS, PRECAUTIONS AND RECOMMENDATIONS	11
3. SY	STEM GENERAL DESCRIPTION	16
3.1.	SYSTEM DESCRIPTION	16
3.2.	APPLICATION SPECIFICATION	16
3.2.1.	Operation Principles	16
3.2.2.	Significant physical characteristics	16
	User's profile	16
3.3.	MAIN PRODUCT COMPONENTS	18
3.3.1.	Light	
3.3.2	Arm	19
3.4.	APPLIED PARTS	20
3.5.	LABEL POSITIONING	20
3.6	SYSTEM REQUIREMENTS	20
3.6.1.	Place of Installation	20
4.	OPERATION	22
4.1	INITIAL PREPARATION	22
4.2.	ACTIVATING THE LIGHT	
4.3	LUMINOSITY SEQUENCE	23
4.4.	HEAD MOVEMENT	24
4.5.	HANDLE MOVEMENT	
5.	CLEANING AND DISINFECTION	26
6.	PROBLEMS DIAGNOSTIC	28
6.1.	PROBLEMS SOLUTION	
_	INSPECTION AND MAINTENANCE	
7. 7.1.		
7.1. 7.2.	PERIODIC INSPECTION	
7.2. 7.3.	CORRECTIVE MAINTENANCE	
7.3. 7.4.	ALLIAGE AUTHORIZED SERVICE NETWORK	
7.4.		
8.	WARRANTY	34
9.	RULES AND REGULATIONS	36
10.	TECHNICAL SPECIFICATIONS	38
	EQUIPMENT CLASSIFICATION	
	DEVICE INFORMATION	
	ENVIRONMENTAL CONDITIONS	
	LIGHT BEAM PROJECTION	
10.5	DENTAL LIGHT DIMENSIONS	

Owner's Manual

11.	ELECTROMAGNETIC COMPATIBILITY	48
11.1.	GUIDANCE AND DECLARATION FOR ELECTROMAGNETIC EMISSIONS	48
11.2.	GUIDELINES AND DECLARATION FOR ELECTROMAGNETIC IMMUNITY	49

1

GERENAL INFORMATION

1. GENERAL INFORMATION

1.1. DEAR COSTUMER

Congratulations on your excellent choice. When you buy ALLIAGE-quality equipment, you can be sure of purchasing products with technology compatible with the best in the world in their class. This manual offers you a general presentation of your equipment, describing important details that can guide you in its correct use, as well as in solving minor problems that may eventually occur. This manual must be read in full and kept for future reference.

1.2. INDICATONS FOR USE

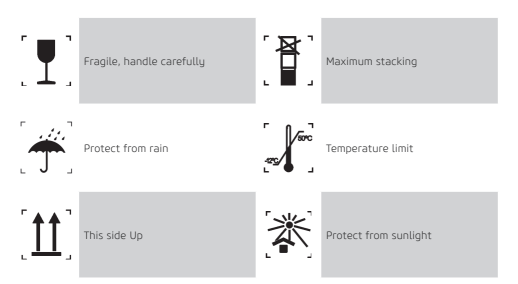
The Dental Light is intended to assist in the treatment of the patient and in the performance of dental practices, and is for dental use only.

1.3. CONTRAINDICATION

There are no known contraindications for this equipment.

1.4. SYMBOLOGY

The following symbols are used both throughout this manual and on the product. Make sure you fully understand each symbol and follow the instructions that come with it.





Recyclable



Indicates that the product must be taken to a special waste collection site at the end of its shelf-life. Applies to both device and accessories



Electrostatic sensitive (ESD) devices



Protection ground



Refer to the instructions manual



Mandatory action



Caution



Overall warning



Authorized representative in European Union



Indicates that the equipment complies with Directives 2011/65/EU and 2015/863/EU on the Restriction on the use of certain hazardous substances in electrical and electronic equipment.



Medical device



Catalog number



Model number



Serial number



Manufacturer



Manufacturing date

MODEL

Model

WARNINGS, PRECAUTIONS AND RECOMMENDATIONS

2. WARNINGS, PRECAUTIONS AND RECOMMENDATIONS

Overall warnings



Read and understand all instructions contained in these instructions for use before installing or operating this equipment.



Use the equipment only in perfect condition and protect yourself, patients and others against possible hazards.



This equipment must be installed and operated by personnel familiar with the necessary precautions.



The Dental Light has 3 different interactions with the user, namely:

- Identification label: Located on the side of the equipment;
- Safety symbols: Located in risk areas and on its identification tag;
- Sensor.

During transportation

The equipment must be transported and stored, observing the following:

- Handle carefully to avoid falls, excessive vibrations and impacts;
- The arrows on the packaging must be pointing upwards;
- To handle the package as a single unit, consider the center of gravity indicator;
- Do not stack above the quantity indicated on the packaging;
- Do not walk or stand above the packaging
- Protect against sunlight, humidity, water and dust;
- Observe the temperature, pressure and relative humidity limits.

During the installation of the equipment



Installation instructions can be found in the service manual, accessible only to authorized technicians.



The equipment must be installed only by the authorized technician. This technical procedure cannot be performed by the user.

Owner's Manual

- The equipment must be installed only by authorized technicians.
- The service manual's recommendations regarding the mandatory existence of protective grounding must be followed.
- Install the equipment in a location where it will not be in contact with humidity, water, plants and animals.
- Install the equipment in a location where it will not be damaged by pressure, temperature, humidity, direct sunlight, dust, salts or corrosive products.
- This equipment is not designed for use in the presence of steams from flammable anesthetic mixtures or nitrous oxide.
- Place any other external devices at least 1.5 meters away from the equipment, so that the patient cannot touch any other external devices while they are being attended to.
- The recommendations in this manual for EMC should be followed. Communication equipment and RF generating sources can affect the operation of the equipment.
- Equipment can cause radio interference or interrupt the operation of nearby equipment, making it necessary to take mitigating measures, such as reorientation, relocation of the equipment or shielding of the location.

Before using the equipment

In order to help ensure proper hygiene and protect against infectious diseases, before the first use, the equipment must be cleaned and disinfected according to the instructions contained in this manual.

While using the equipment

- Under no circumstances can the patient operate the equipment.
- The patient must not touch any parts other than those specific for their treatment.
- The equipment must be operated only by qualified health professionals.
- To operate the equipment, operating personnel must:
- Read and understand the user manual
- Be familiar with the basic structure and functions of this equipment.
- Be familiar with the emergency situation protocols for this equipment.
- Be able to recognize irregularities in the operation of the equipment and implement the appropriate measures, when necessary.
- This equipment was designed according to the electromagnetic compatibility standards, but in very extreme conditions, it can cause interference with other equipment. Do not use this equipment in conjunction with other devices that are very sensitive to interference or with devices that create high electromagnetic disturbances.
- If this product is exposed to water, humidity or foreign substances, turn it off immediately and contact an Alliage Authorized Service Center.
- In case of damage or defect, do not use the equipment and contact an Alliage Authorized Service Center.
- Do not use the equipment if any of its compartments or parts are damaged, loose or have been removed. Contact an Alliage Authorized Service Center and request repair or replacement of any damaged, loose or removed cabinets or parts of the equipment before using it again.
- Do not touch the equipment or use it if it is being repaired or if the equipment's cabinets have been removed.
- Do not open or remove any of the equipment's cabinets. No internal parts can be repaired by the user.
- In case of falling or impact of moving parts causing their breakage, be careful when handling

Owner's Manual

them, there may be sharp parts.

- This equipment does not produce physiological effects that are not obvious to the operator.
- The operator cannot come into contact with the patient when in contact with accessible connectors.
- The operator cannot use tools to open the equipment.

Cross-contamination prevention



Adequate cleaning and disinfection measures should be taken to avoid cross-contamination between patients, users and others.

• For each new patient, perform cleaning and disinfection procedures according to the instructions contained in this manual.

After using/operating the equipment

- Turn off the equipment if it is not in use for a long time
- All parts that have been in contact with the patient should be cleaned and disinfected for each new patient, in order to prevent the transmission of infectious agents that can cause serious illness.
- Perform cleaning and disinfection according to the instructions contained in this manual.
- Do not unplug the cable or other connections unnecessarily.
- Do not modify any part of the equipment.

Precautions in case of altered equipment operation

If the equipment shows any abnormality, check if the problem is included in any item listed in the "Problem diagnosis" topic of this user manual.

If it is not possible to solve the problem, turn off the equipment and contact an Alliage Authorized Service Center.



The manufacturer is NOT responsible for:

- The equipment being used for purposes other than those for which it was designed.
- Damage caused to the equipment, the operator and/or patient, as a result of incorrect installation and maintenance procedures that are not in accordance with the operating instructions accompanying the equipment.
- Inadequate equipment operation.

Precautions for reducing environmental impact

Alliage S/A aims to achieve an environmental policy to promote the supply of environmentally conscious medical and dental products that continually minimize environmental impact and are friendlier to the environment and human health. In order to maintain a minimal environmental impact, observe the recommendations below:

- After installation, send recyclable materials for recycling process.
- During the life cycle of the equipment, turn it off when it is not in use.
- To prevent environmental contamination, the disposal of waste and consumables must follow the usual procedure for biomedical waste.

Biomedical waste encompasses non-acute materials that may cause disease or suspicion of harboring pathogenic organisms that must be stored in a yellow bag properly labeled with a biohazard symbol, stored in a puncture-resistant, watertight container until collection and incineration.



The Equipment's packaging consists of wood, cardboard, plastic and expanded polyurethane (PU) which are 100% recyclable materials.

SIZES:

Main unit: $420 \times 420 \times 220$ mm/ MASS: Approximately: 2 Kg Arm: $910 \times 720 \times 140$ mm/ MASS: Approximately: 10 Kg

Precautions in case of unusable equipment

In order to avoid environmental contamination or improper use of the equipment, when it is unusable, it must be disposed of (in accordance with current legislation) in an appropriate place, as the materials inside can contaminate the environment.

For the European Economic Area (EEA), this product is subject to Directive 2012/19/EU, as well as the corresponding national laws. This directive requires that the product must be taken to a special waste collection site at the end of its shelf-life. Applies to both the device and accessories. Contact your salesperson if final product disposal is required.



This equipment must not be disposed of as household waste.

SYSTEM GENERAL DESCRIPTION

3. SYSTEM GENERAL DESCRIPTION

3.1. SYSTEM DESCRIPTION

Reflector for lighting for dental use with adjustable intensity.

Composed of a new light emission system, using LED technology. This is the acronym for Light Emitting Diode, a totally different way of emitting light, when compared to conventional halogen light devices. In addition to being infinitely more durable (over 50,000 hours) and with low energy consumption, LEDs have made the devices more compact, ergonomic and easy to install and transport.

With a rectangular light focus, it allows ample illumination of the operating field, minimizing the need for constant repositioning of the focus.

Enables the dental surgeon to choose the desired brightness according to the procedure used (brightness with white light or yellow light), which depends on the chosen version.

The use of white light is recommended for normal work procedures (illumination of the operating field).

The use of yellow light is recommended for work procedures with photo curable materials, as it does not interfere with their cure.

It has the following tupes of actuations:

- Through Optical Sensor and
- Control pedal for the dental chair.

Activation by optical sensor through hand approach, which provides practicality in the operation, and is a great ally in the control of cross infections.

Protector of the LEDs in resistant transparent material protecting them against aerosols.

Steel arm with vertical and horizontal movement with rounded corners. Steel structure with surface treatment through nanotechnology, smooth high-gloss, epoxy-based paint, polymerized in an oven at 250°C, resistant to corrosion and cleaning materials.

Head made of a resistant material, with 620° rotation, providing lightness, durability and ample mobility in all positions (conventional models). For models with "Orbital" arms, the head has full angular focus adjustment.

Removable and self-cleaning bilateral handles that allow isolation avoiding the risk of cross contamination.

3.2. APPLICATION SPECIFICATION

3.2.1. Operation Principles

The LED is a light emitting diode, that is, a semiconductor electronic component that has the property of transforming electrical energy into light.

This light emission is due to the fact that the electrons of the material's atoms are forced to change orbits. When an electron jumps from one orbit to another, it is forced to emit energy to reach the energy level of its new orbit and, under these conditions; the energy it discards appears in the form of light.

The light emitted by the LED is cold due to the absence of infrared in the light beam.

3.2.2. Significant physical characteristics

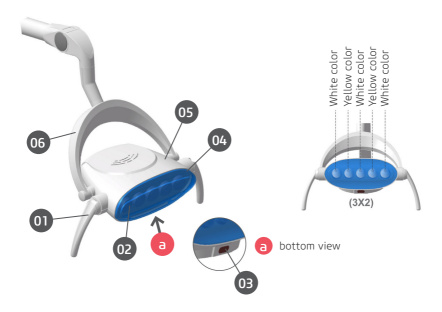
Composed of a new light emission system with adjustable intensity.

3.2.3. User's profile

The Dental Light can be used by both male and female, with the minimum level of literacy, and with the ability to read and understand images, symbols, icons, western characters (Arial font), and alphanumeric characters. People who use the Dental Light may not have a degree of visual imperfection for reading or seeing and an average degree of recent memory impairment, not being clearly able to perform the activities and functions of the product correctly in the profession. The user needs to be a healthcare professional that has been qualified and trained to perform the activities, functions frequently used in the application of the Dental Light and its primary operations functions.

3.3. MAIN PRODUCT COMPONENTS

3.3.1. Light



- 01 Handle
- 02 LED's
- 03 Optical Sensor
- 04 Protector
- 05 Reflective Body
- 06 Articulated arch

3.3.2 Arm



*Supports for coupling in the chair (allows the installation of the articulated arm)



*Support for floor/ ceiling coupling (allows the installation of the articulated arm)



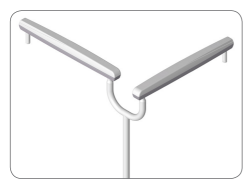
*Articulated arm



*Simple bench articulated arm



*Sleeve Kit - allows the fixation of the Light in offices with 38mm column



*double bench articulated arm



*Articulated arm with "orbital" movement

^{*} Optional items (Subject to commercial availability)

3.4. APPLIED PARTS

This equipment does not have applied parts

3.5. LABEL POSITIONING

The following figure illustrates the location of the labels on the equipment.



Illustrative picture





3.6 SYSTEM REQUIREMENTS

3.6.1. Place of Installation



In order to meet safety standards, do not operate non-medical equipment, such as personal computers, within the patient's area. Outside the patient's area, the presence of non-medical equipment is acceptable, provided that approved and certified computer equipment is used.

Computer equipment must be CE-approved and must comply with CE 60950-1: 2005 + AMD1: 2009 + AMD2: 2013 and low voltage 2014/35/EU and 2014/30/EU EMC guidelines.

4

OPERATION

4. OPERATION

4.1 INITIAL PREPARATION



The equipment must be cleaned and disinfected before use on a new patient, observing the instructions contained in this manual.

4.2. ACTIVATING THE LIGHT

Activating the light with sensor head on the "pedal"

To turn the light on or off, press one of the buttons as shown in the figures below:

- For the pedal with 11 keys, activate the keys (05) or (06), figure a.
- For the Chip Blower pedal, press the button (08), figure b.
- For the pedal with 07 keys, activate the key (07), figure c.
- For the Joystick pedal, press the key (03), figure d.
- For the pedal with 03 keys, activate the key (02), figure e.
- For the Joystick pedal with 07 keys, activate the key (02), figure f.



Activating the light through the "sensor" on the head

To turn the light on or off, run your hand over the sensor within a maximum distance of 10 cm. The pedal button must be turned on, as previously mentioned.

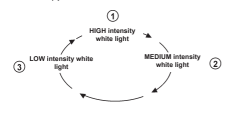
Programming the brightness of the sensor light

The sensor light, in addition to the on/off system, has the programming mode. With the light on, keep your hand near the sensor for 3 seconds. 1 Beep will be triggered and the light will enter the programming mode changing to the next intensity. Passing the sensor through the hand, the light will change the light intensity according to the light sequence below: High intensity, Medium intensity and low intensity. After choosing the desired intensity, wait for 5 seconds for it to be saved, a new Beep will be emitted.

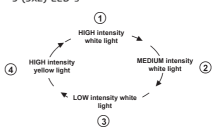


4.3 LUMINOSITY SEQUENCE

Sequence for light models: 5 LED's, / 3 LED's.



Sequence for light models: 5 (3x2) LED's





Yellow light is an exclusive function for the treatment of light-activated restorative materials. It should not be used for operational processes.

4.4. HEAD MOVEMENT

Head movement

The head movement is done through the handles (08), totally ergonomic, designed to provide absolute isolation.





Position the head 70 cm from the operating field.



When handling the equipment, be aware of the parts that may catch your fingers.

4.5. HANDLE MOVEMENT

The handle has a 360° movement in both clockwise and counterclockwise directions. By applying a light effort, remove the handle, without completely uncoupling it; perform the movement according to the desired need. Fit it back in.







Do not obstruct the air passages in the head.

CLEANING, DISINFECTION AND STERILIZATION

5. CLEANING AND DISINFECTION



Before starting the cleaning and disinfection procedure, turn off the main switch of the equipment to avoid permanent damage.



For your protection, during the equipment cleaning and disinfection process, use PPE such as disposable gloves and goggles.

The cleaning and disinfection process must be performed at each patient change.

When starting the process, check for visible dirt, such as blood or saliva.

For cleaning, use a clean, soft cloth moistened with mild soap and then dry with a clean, soft cloth or paper towel.

For the disinfection process of the equipment, use disinfectant detergent foam that has active components based on didecyldimethylammonium chloride.

Apply the disinfectant detergent foam on the surface or on a clean cloth and spread it on the surface to be treated. Respect the antimicrobial contact time specified by the manufacturer. After application, allow to dry. Do not rinse.



Do not spill disinfectant liquid on the equipment.



Do not use organic solvents, for example, thinner, to clean the equipment. In the event that the development solution is spilled on the panel, clean it immediately, as these solutions may compromise the equipment's paint.

PROBLEMS DIAGNOSTIC

6. PROBLEMS DIAGNOSTIC

6.1. PROBLEMS SOLUTION

In case any problem is found in the operation, follow the instructions below to check and fix the problem, and/or contact your representative.

Unforeseen events	Probable Cause	Solutions
- The light does not turn on;	- Lack of energy.- Fuse blown.- LED blown.	- Wait for power to be restored Turn off the chair power and request the presence of a Technician Request the presence of a technician.
- The sensor does not activate.	 Pedal off. Sensor blown. Hand positioning distance from the sensor greater than 10 cm. Lack of energy. Fuse blown. LED blown. 	 Switch on the pedal. Request the presence of a technician. Position your hand no more than 10 cm away from the sensor. Wait for power to be restored. Turn off the chair power and request the presence of a Technician. Request the presence of a technician.

If problems persist, contact the Alliage Service Department.

INSPECTION AND MAINTENANCE

7. INSPECTION AND MAINTENANCE



Maintenance or service procedures may only be carried out by a technical service authorized by the manufacturer.

All instructions for using the equipment as intended are provided in this user's guide. If a problem is detected and cannot be corrected with the instructions in the problem diagnostics section, contact the Alliage Service Department.

7.1. PERIODIC INSPECTION

This equipment must be regularly inspected to ensure operational safety and functional reliability. This inspection must be carried out by personnel familiar with the necessary precautions to avoid exposing the patient to risk.

Periodic inspection should be carried out at regular intervals (at least once a year) to ensure that the product is permanently safe and operational. All components subject to normal wear and tear should be checked and, if necessary, replaced.

The manufacturer and the assembler/installer are exempt from responsibility for the standard results not being compliant in cases where the user does not perform the maintenance recommended by the manufacturer.

Neither inspection nor service is part of the equipment's warrantu.

The maintenance performed must be documented and kept with the equipment.

Item	Inspection description	Recommended frequency
Security system	Collision, Warning Lights, and Interlock	Daily
Electric parts	Overheating/ Noise/ Burning smell	Monthly
Elevation	Operation/ Noise/ Vibration	Annually
Movement mechanism	Operation/ Noise/ Vibration	Annually
Pedal and controls	Operation/ Damage	Annually

If problems are detected during the inspection, contact the Alliage Service Department.

7.2. PREVENTIVE MAINTENANCE

In addition to the annual inspection, in order to ensure a long durability and the good operation of your equipment, it is important to carry out preventive maintenance for a maximum period of three (3) years.

 ${\it Please contact the Alliage Service Department about our periodic review and preventive maintenance program.}$

7.3. CORRECTIVE MAINTENANCE



To repair or replace any part, see instructions in the service manual.



Corrective maintenance cannot be performed by the user. Do not open the equipment or try to repair it yourself or with the help of someone without training or authorization. This can aggravate the problem or produce a failure that can compromise the safety of the equipment.



The power cables and electronic boards can be changed only by the authorized technician. See service manual for connection and anchoring information.



No maintenance or services can be provided to the equipment or any of its parts during use with a patient.



The equipment contains parts under high voltage. Risk of electric shock. Turn off the main switch before servicing.



Moving parts can cut or crush.



The service manual is only available for Authorized Technical Assistance.

Alliage declares that the provision of circuit diagrams, component lists or any other information that provides technical assistance on behalf of the user, can be requested as long as previously agreed between the user and Alliage. The warranty will be void if the original parts are removed/ replaced by unauthorized service technicians.

7.4. ALLIAGE AUTHORIZED SERVICE NETWORK

All services performed on the Alliage equipment must be performed by an Authorized Technical Assistant, as otherwise they will not be covered by the warranty. If you need to request electrical diagrams and or specification of components that is not stated in the user's manual, use the Alliage Customer Service to make the request.

Telephone: +55 (16) 3512-1212

Address: Rodovia Abrão Assed, Km 53 - Recreio Anhanguera - Ribeirão Preto-SP/ Brazil - Zip Code

14097-500.



8. WARRANTY

This equipment is covered by the warranty periods, terms and conditions contained in the Warranty Certificate that comes with the product.

9

RULES AND REGULATIONS

9. RULES AND REGULATIONS

This equipment has been designed and manufactured to meet the following standards:

	ABNT NBR IEC 60601-1:2010 Amendment 1:2016	Medical Electrical Equipment - Part 1: General requirements for basic safety and essential performance.
	ABNT NBR IEC 60601-1-2:2017	Medical Electrical Equipment, Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic interference - Requirements and tests.
	ABNT NBR IEC 80601-1-60:2015	General requirements for basic safety and essential performance of dental equipment
	ABNT NBR 60601-1-6:2011	Electro-medical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability;
	ABNT NBR IEC 62366:2016	Healthcare products - Application of usability engineering to healthcare products;
	IEC 60601-1-9:2014	Medical electrical equipment - Part 1-9: General requirements for basic safety and essential performance - Collateral standard: Requirements for environmentally conscious design
	IEC 62304:2006	Medical device software - Software lifecycle processes.
	ISO 9680:2014	Operating lights
	ISO 7494-1:2018	Dentistry – Dental units – Part 1: General requirements and test methods
	ISO 7494-2:2015	Dental units – Part 2: Air, water, suction and wastewater Systems
	ABNT NBR ISO 6875:2014	Dental patient chair
	ISO 9687:2015	Graphical symbols for dental equipment
	ISO 15223-1:2016	Graphic symbols for electrical equipment in medical practice
	EN 1041:2008+A1 2013	Information provided by the medical device manufacturer.
	ABNT NBR ISO 10993-1:2013	Biological assessment of medical devices - Part 1: Assessment and testing.
	ABNT NBR ISO 14971:2009	Medical devices - Application of risk management to medical devices.
	ABNT NBR ISO 13485:2016	Quality management systems - Requirements for regulatory purposes

10

TECHNICAL SPECIFICATIONS

10. TECHNICAL SPECIFICATIONS

10.1. EQUIPMENT CLASSIFICATION

Classification according to ANVISA

Class I

Classification according to CE/FDA

Class I

Equipment classification according to EN IEC 60601-1

Product classification for applied parts

N/A

Protection Against Electric Shock

Class I

Protection Against Harmful Water Penetration

IP00 - Product not protected against harmful penetration of water and particulate material

Degree of safety of application in the presence of a flammable anesthetic mixture with air, oxygen or nitrous oxide

Unsuitable equipment

Operation mode

Continuous operation

10.2. DEVICE INFORMATION

Power supply voltage (from the chair)

24 Vcc

Power consumption (from the chair)

Sirius G8 Sensor 5 LEDs: 60 VA Sirius G8 Sensor 3 LEDs: 40 VA

Sirius G8 Sensor 5 (3X2) LEDs: 40 VA

Net weight of light head

1 kg

Gross weight of light head

1,5 kg

INFORMAÇÕES ESPECIFICAS

Safety mode for compounds According to ISO 9680: 2014 item 5.2.10

Yellow Light

Illuminance 6500 lx

Treatment mode

White light

Correlated color temperature 4500K to 6400K

Power 5 LED's

60VA

Power 3 LED's

40VA

Brightness 5 LED's

High: 45.000 Lux Medium: 35.000 Lux Low: 25.000 Lux

Brightness 3 LED's

High: 30.000 Lux Medium: 20.000 Lux Low: 10.000 Lux

Brightness 3x2 LED's

High: 35.000 Lux Medium: 25.000 Lux Low: 15.000 Lux

Color reproduction index (Ra)

This light does not have this index because it is an LED operating light.

10.3.ENVIRONMENTAL CONDITIONS

Environmental conditions for transport and storage

Room temperature range for transport or storage

-12°C to +50°C

Relative humidity range for transport and storage

< 85% RH

Atmospheric pressure range

700 hPa to 1060 hPa (525 mmHg to 795 mmHg)

Environmental conditions for installation and operation

Room temperature range for operation

+10°C to +35°C

Operating relative humidity range (non-condensing)

< 75% RH

Atmospheric pressure range

700 hPa to 1060 hPa (525 mmHg to 795 mmHg)

Operating altitude

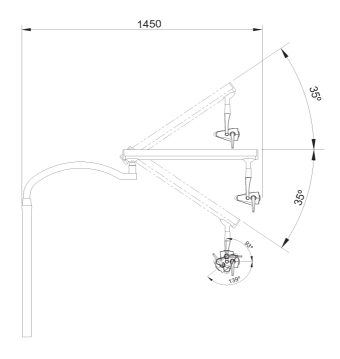
≤ 2000 m

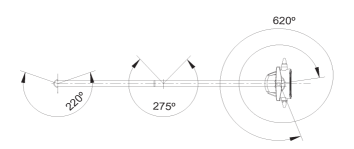
10.4. LIGHT BEAM PROJECTION

Projection	Perimeter (mm)	Area (mm²)
100%	503	15914
75%	475	14242
50%	459	12332
10%	457	8580

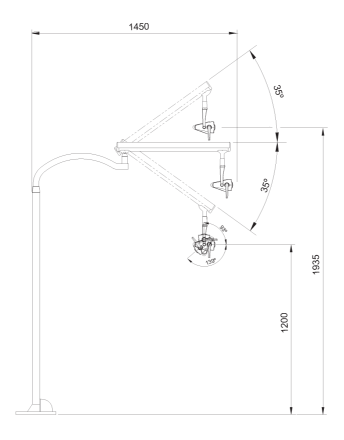
10.5 DENTAL LIGHT DIMENSIONS

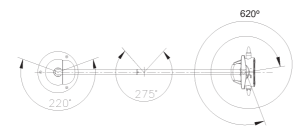
Conventional Arm (Coupled to the chair)



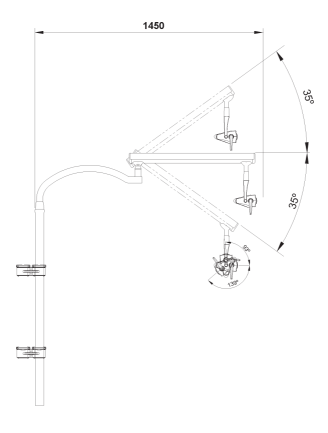


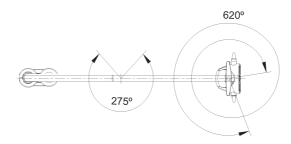
Floor



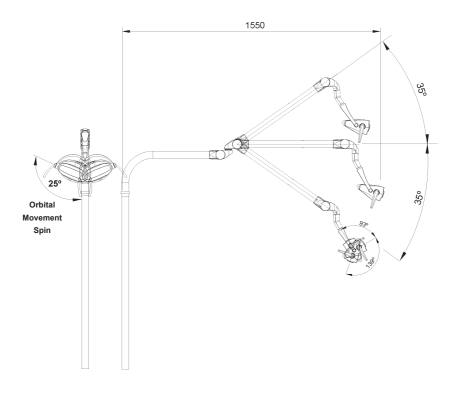


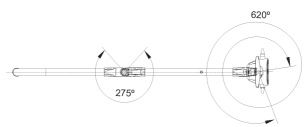
Coupled Arm



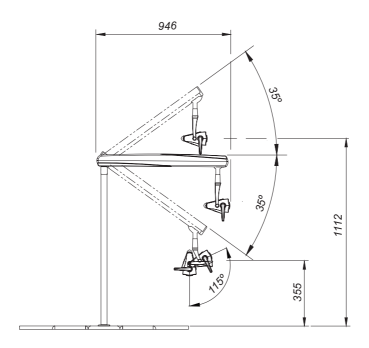


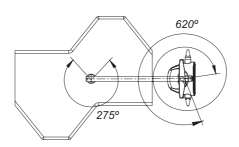
Orbital arm (coupled to the chair)



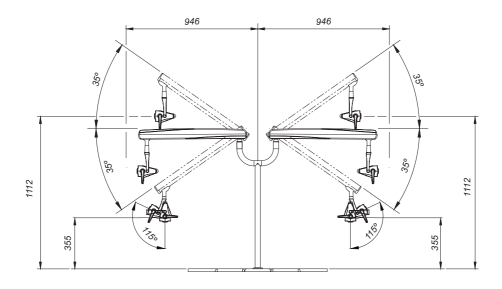


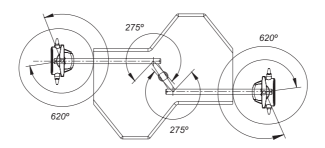
Simple bench





Double bench





11

ELECTROMAGNETIC COMPATIBILITY

11. ELECTROMAGNETIC COMPATIBILITY

The Dental Light is intended for use in the electromagnetic environment specified below. The buyer or user should ensure that it is used in such an environment.

The Dental Light is suitable for use in a professional healthcare environment, not including areas where there is sensitive equipment or sources of intense electromagnetic disturbances, such as the RF shielded room of a magnetic resonance imaging system, in operating rooms close to active AF surgical equipment, electrophysiology laboratories, armored rooms or areas where short wave therapy equipment is used.

The following tables provide information on the equipment's compliance with the ABNT NBR IEC 60601-1-2: 2017 standard.

11.1. GUIDANCE AND DECLARATION FOR ELECTROMAGNETIC EMISSIONS

Emission assay	Compliance	Electromagnetic Environments – guidelines
Emissions of RF CISPR 11	Group 1	The Dental Light uses RF energy only for its internal functions. Therefore, its RF emissions are very low and are unlikely to cause any interference in nearby electronic equipment.
Emissions of RF CISPR 11	Class A	The Dental Light is suitable for use in
Harmonic emissions IEC 61000-3-2	Class A	all facilities, except domestic ones and those directly connected to the public
Voltage fluctuation/ Scintillation emissions IEC 61000-3-3	Complies	low voltage power supply network that supplies buildings used for domestic purposes.

Note: The emission characteristics of this equipment make it suitable for use in industrial areas and hospitals (IEC/CISPR 11, Class A). If used in a residential environment (for which IEC/CISPR 11, Class B is normally required), this equipment may not provide adequate protection for radio frequency communication services. The user may need to take mitigation measures, such as relocating or redirecting equipment.

11.2. GUIDELINES AND DECLARATION FOR ELECTROMAGNETIC IMMUNITY

Phenomenon	Basic EMC standard or test method	Immunity assay level	Compliance level	
Electrostatic discharge	IEC 61000-4-2	±8 KV contact ± 2 KV, ± 4 KV, ± 8 KV, ± 15 KV air	±8 KV contact ± 2 KV, ± 4 KV, ± 8 KV, ± 15 KV air	
Radiated RF EM fields	IEC 61000-4-3	3 V/m 80 MHz – 2,7 GHz 80 % AM to 1 kHz	3 V/m 80 MHz – 2,7 GHz 80 % AM to 1 kHz	
Fields close to RF wireless communication equipment	IEC 61000-4-3	Refer to table	Refer to table	
Fast/ saved electrical	IEC 61000-4-4 power supply a.a.	± 2 kV 100 kHz repetition frequency	± 2 kV 100 kHz repetition frequency	
transients	IEC 61000-4-4 signal input/output	± 1 kV 100 kHz repetition frequency	± 1 kV 100 kHz repetition frequency	
Outbreak line-line	IEC 61000-4-5	± 0,5 kV, ± 1 kV	± 0,5 kV, ± 1 kV	
Outbreak line-earth	IEC 61000-4-5	± 0,5 kV, ± 1 kV, ± 2 kV	± 0,5 kV, ± 1 kV, ± 2 kV	
Conducted disturbances induced by RF fields	IEC 61000-4-6	3 V 0.15 MHz – 80 MHz 6V in ISM bands between 0.15 MHz and 80 MHz 80 % AM to 1 kHz	3 V 0.15 MHz - 80 MHz 6V in ISM bands between 0.15 MHz and 80 MHz 80 % AM to 1 kHz	
Magnetic fields at the stated supply frequency	IEC 61000-4-8	30 A/m 50 Hz or 60 Hz	30 A/m 50 Hz or 60 Hz	
Voltage drops	IEC 61000-4-11	0 % UT; 0.5 cycle A 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycles Monophase: at 0°	0 % UT; 0.5 cycle A 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycles Monophase: at 0°	
Voltage interruptions	IEC 61000-4-11	0 % UT; 250/300 cycles	The device will shut down or restart if the power is interrupted for five seconds.	

NOTE 1 At 80 MHz and 800MHz, the higher frequency range is applicable.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

NOTE 3 UT is the mains voltage c.a. before applying the test level.

Fields close to RF wireless communication equipment

Testing fre- quency (MHz)	Band (MHz)	Service	Modulation	Maximum power(W)	Distance (m)	Immunity assay le- vel (V/m)
385	380-390	TETRA 400	18Hz pulse modulation	1.8	0.3	27
450	430-470	GMRS 460, FRS 460	FM deviation of ± 5 kHz Sinusoidal 1kHz	2	0.3	28
710	704-787	LTE Band	217 Hz pulse	0.2	0.3	9
745		13, 17	modulation			
7480						
810	800-960	GSM 800 /		2	0.3	28
870		900 , TETRA modulation 800 , IDEN 820 , CDMA 850 , LTE 5 Band				
930						
1720	1700 -1990			2	0.3	28
1845			modulation			
1970		DECT; LTE Band 1, 3, 4, 25; UMTS				
2450	2400-2570	Bluetooth, W L A N 802.11	217 Hz pulse modulation	2	0.3	28
5240	5100 - 5800			0.2	0.3	9
5500		2450, LTE 7 Band	7 modulation			
5785		555				

List of cables used

Cables	Description	Length
Supply	Tripolar Power Cable Gauge 3x2.50 mm², 250V AC, Male Plug 20A NBR 14136 2P+T, no female plug, Inmetro. (From the Chair).	3 m



The Dental Light is intended to assist the healthcare professional, and it is for dental use only. In case of EMC disturbances, the operator may experience loss of communication between the equipment and controls.



Compliance with EMC and EMI standards cannot be guaranteed by using cables that have been altered or that do not comply with the same standards for which the equipment has been validated.



Use of this equipment adjacent to other equipment should be avoided, as it may result in improper operation. If this use is necessary, it is advisable that this and the other equipment be observed to verify that they are operating normally.



Do not use accessories, transducers, internal parts of components and cables other than those previously specified by the manufacturer. This can result in increased emission or decreased electromagnetic immunity and result in improper operation.



Portable RF communication equipment (including peripheral items, such as antenna cables and external antennas) should not be used within 30 cm away from any part of the equipment, including cables specified by the manufacturer. Otherwise, performance degradation of this equipment may occur.



In order to maintain basic safety from electromagnetic disturbances during the expected shelf-life, always use the equipment in the specified electromagnetic environment and follow the maintenance recommendation described in this manual.



The pins, connector sockets or elements that carry the ESD warning symbol must not be touched or connected without ESD protection measures.



NUM. REG. ANVISA: 10069210069



