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## **INSTALLATION, OPERATIONS & MAINTENANCE MANUAL UCI-OVAL**

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**NOTE:** MD use must be done only after having read all sections of the current operation instructions.

- Please follow all warnings in these instructions for use in the enclosed MD's

Device intended for use by lay users, the profane user: the patient, uses only the push-buttons to switch on/off the lighting and for the nurse call and uses the electrical sockets to connect mobile phone to recharge or connect a radio set, in case of different use they should consult a healthcare professional.

## 1) DESCRIPTION

Suspended unit designed to meet the requirements of different hospitalization, intermediate and critical areas

MODELS:

uci-oval: Suspended Bedhead unit equipped with electrical components, gas outlets +2 units lighting fittings



Representative image of model UCI-OVAL

## 2) INTENDED USE OF THE BED HEAD UNIT

Medical power supply units installed in intensive therapy rooms, operating theatres, destined to medical gas distribution.

These medical supply units are medical devices designed to provide close access of direct and indirect lighting, electrical sockets, data sockets, nurse call, diagnostic signals and alarms and, specifically for authorized medical personnel, medical gas outlets, necessary for the care and treatment of patients by authorized medical personnel.

### INTENDED USER PROFILE

The only form of direct contact between patient and device occurs during push the button on handle control panel for lights control and nurse call and/or switches and buttons for lighting control on device. the operator can be in contact to the aluminium structure, the polycarbonate covers and the push-button panels, to the switches, the electrical sockets, the buttons, the data sockets which are made of plastic material, then there are the gas sockets medical devices that are made of metal and equipotential sockets that can be metal or plastic and finally accessories made of plastic + metal.

**It is strongly recommended that visitors consult healthcare personnel before performing any operation, specially on hanging units for the operating room and intensive care (UCI-OVAL), because this medical device is intended for patients who can be under a critical health situation.**

Installation and maintenance must be carried out exclusively by personnel specialized authorized by the manufacturer.

The medical device is used by:

- authorized medical personnel, in detail:

- Turn on / off the direct and indirect lighting
- Use electrical outlets
- Nurse call
- Diagnostic signals and alarms
- Distribution of medical gases

- people visiting the intensive care rooms or patients who can be under a critical health situation, themselves “lay person” (not applicable for analysis laboratories): only non-medical functions, in detail:

- Turn on / off the direct and indirect lighting
- Use electrical outlets that are not currently being used by a medical device
- Nurse call

#### **Authorized medical personnel:**

The functions they perform are those of their position within the center hospitable. They are in charge of connecting the gas outlets present on the medical supply unit indicated for each patient and to connect the medical devices that will be powered by the medical supply unit.

To carry out these functions, the medical staff will have studies and basic knowledge in medicine and knowledge of electricity at the user level.

The medical personnel will have sufficient knowledge to read and understand the instructions for assembly that accompanies the headboard.

### **3) COMPONENTS**

The UCI-OVAL device can be delivered with a handset from which the user can control the lighting and send a warning signal to call for assistance. Minimal technical features:

- Casing in self-extinguishing polycarbonate, no propagation of toxic fumes. Designed for housing the electrical control units with recessed seats to prevent the units from being switched on accidentally and unwanted calls due to pressure on the pushbutton units caused by movements of patient’s body.
- Protection degree: IP43
- Safety plugs on fastening screws to prevent tampering by the patient.
- Silk-screened indications on control keys with symbols (designed for easy understanding (children, senior citizen, etc.)
- Flame-resistant white multi-wire cable

Components to be used with UCI-OVAL:

Internal Code	Manufacturer	Photo	Description and Technical data	Material	Manufacturer references	Class according MDR
J10TH2xx	ELLEDUE		Handle push button control to nurse call and light turn off/on	Self-extinguishing polycarbonate, no propagation of toxic fumes	255703 260101 260102 260203 265104 TH2401 TH2406 TH2507 TH2551 TH2557 TH2577 NX0884	I
J10THNX0884 J10THNX0884-SC J10THNX0883 J10THNX0883-SC J10THNX0881 J10THNX0881-SC	IBERNEX		Handle push button control to nurse call and light turn off/on	Self-extinguishing polycarbonate, no propagation of toxic fumes	NX0884/SC NX0883 NX0883/SC NX0881 NX0881/SC	I

**ACCOMPANYING COMPONENTS:**

Components to be used with UCI-OVAL:

Upon request, the unit can be supplied with carrousel with trays or drawers that move along their entire length by means of wheels with bearings, which run along a aluminium profile of the suspended unit. These carrousel suspended can turn at an angle of 300 degrees. They incorporate a handle that immobilizes them in their turning and translation movements.



HOV1101 Basic Carrousel  
 Can be moved longitudinally through the bedhead unit and can be twisted through 300°. It has an built-in blocking device for both movements  
 Maximum load. 100kg



H1106 tablet  
 Tablet with an usable surface of 640x380mm, fixed on the vertical bars of the carrousel and movable on vertical direction  
 Maximum load=60kg



**H1113 Half tablet**  
 Tablet with an usable surface of 380x320mm,  
 fixed on one of the vertical bars of the carousel  
 and movable on vertical direction  
 Maximum load=10kg



**H1114 Single drawer**  
 Dimensions: 630x380mm fixed under a tablet  
 (H1106)  
 Maximum load of the unit: 60kg  
 Maximum load of the drawer: 10kg



**H1115 Double drawer**  
 Unit formed by two drawers , each 630x380x80mm,  
 fixed under a 640x380mm plate  
 Maximum load of the unit: 60kg  
 Maximum load of the drawer: 10kg



**H1112 Straight rail**  
 30x10mm fixed to one of a vertical bar of the  
 carousel, movable on vertical direction and can be  
 twisted through 360°, provided with protection  
 caps on both ends.  
 Length of the rail:600mm  
 Maximum load: 5kg



**H1108 Compact support**

Fixed to one of a vertical bar of the carousel,  
movable on vertical direction.

There allows in this contour the fixing of clamps of  
30x10mm

Maximum load: 10kg



**H1107 Double rail**

Two paralell rails 30x10mm, fixed to the vertical  
bars of the carousel and movable on the vertical  
direction

Maximum load: 112kg



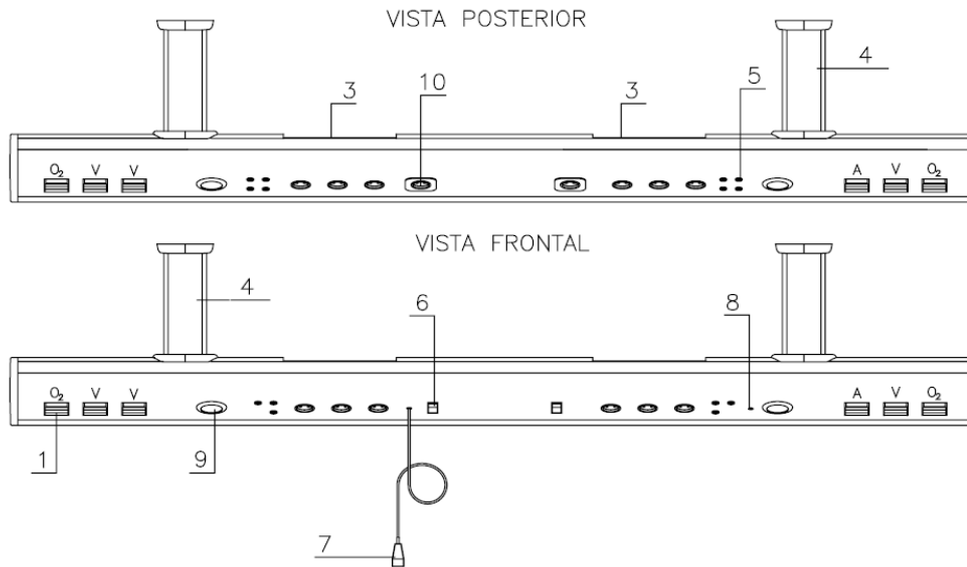
**H1023 Support for infusion bottles**

With 2 Maximum load: 3kg

or 4 hooks Maximum load: 6kg

#### 4) PARTS OF BEDHEAD UNIT

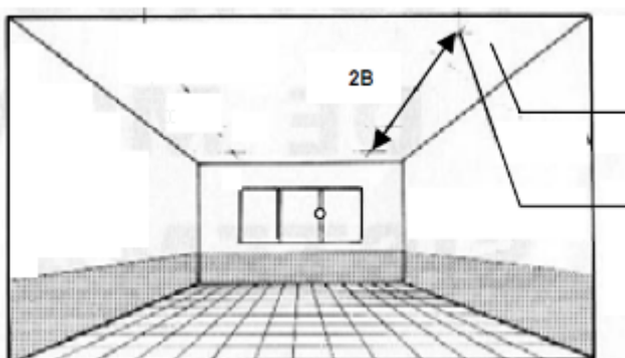
The UCI-OVAL unit has several separate compartments for electricity, for lighting and for medical gases, in accordance with international regulations and standards



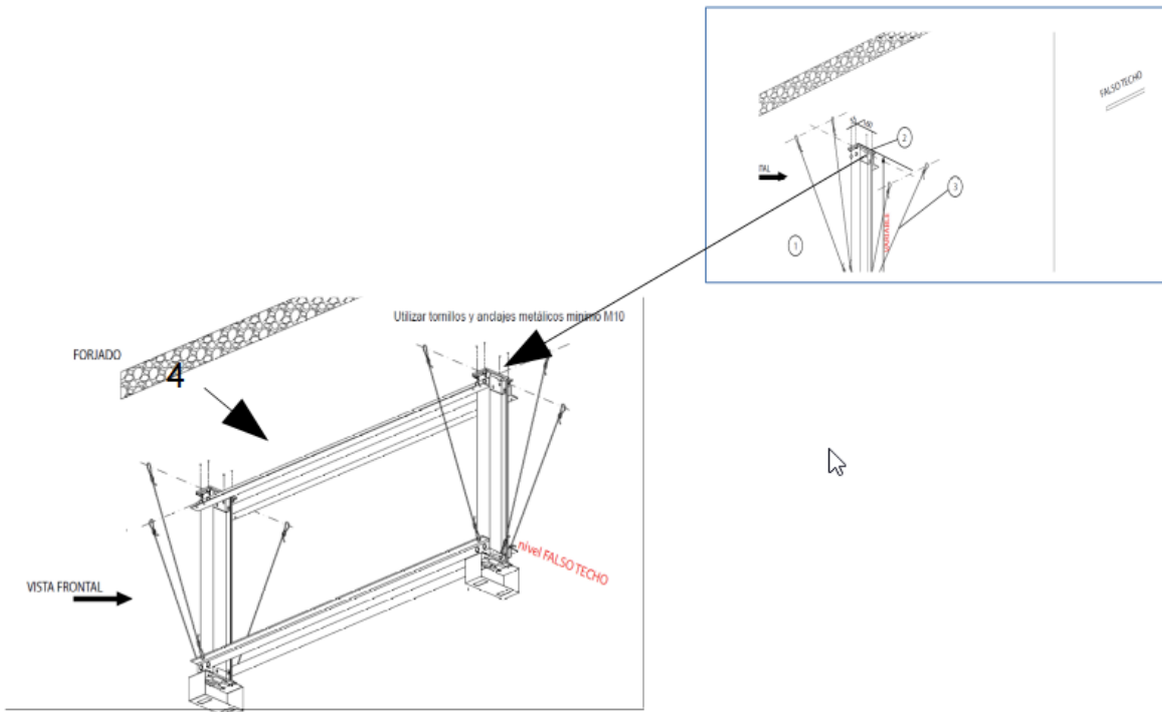
- 1- MEDICAL GAS OUTLETS
- 2- ELECTRICAL SOCKETS
- 3- INDIRECT LIGHTING
- 4- SUSPENSION
- 5- EQUIPOTENTIAL SOCKETS
- 6- SWITCHES
- 7- NURSE CALL
- 9- DIRECT LIGHTING

#### 5) INSTALLATION ON CEILING

A) First we will define the suspension points in the ceiling according to the distribution of units as indicated in the Project. The length between axes will be indicated in the drawings validated by the client. Make the holes in the ceiling considering the distances between bolts of the support bracket (55 and 160mm). Use M10 minimum screws and metal plugs

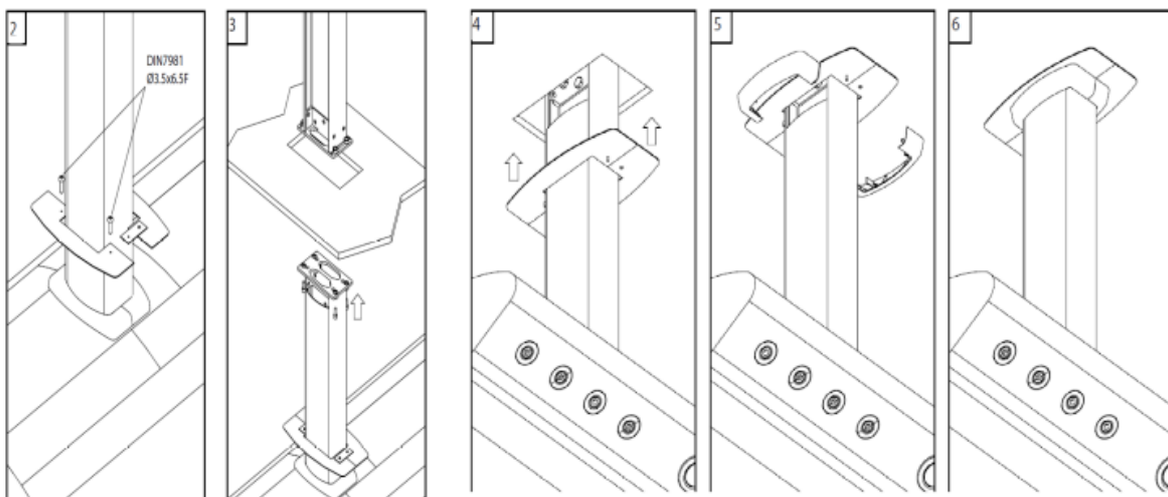


B) Assemble the vertical profiles by means of screws to the structural ceiling. Leave the screws untaught to be adjusted in order to avoid this alignment due to non-uniform ceiling.



C) Assemble the 4 profiles as an “L” and locate the 6 stabilizers in longitudinal and transversal directions respect to the unit. Once the stabilizers are located, check the horizontal alignment and correct the possible mistakes by adjusting the tensor wires and the joints between the suspensions and ceiling

D) Once the suspended ceiling is closed, fix the suspensions of the suspended beam to the ceiling supports (No. 3) by means of included screws (M10x25) and clip the cover pieces, if the cutouts has been made too large and the covers fails to cover it, they can place additional sheets (Nº2 and 4)



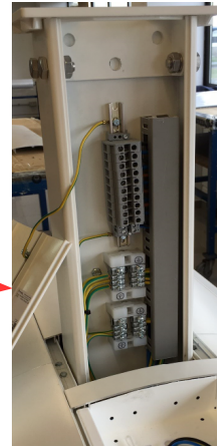
## 6) ELECTRICAL CONNECTIONS

Perform the electrical connection according to the wiring diagram placed inside the BHU



**ELECTRICAL CONNECTIONS ARE TO BE CARRIED OUT BY COMPETENT PERSONS ONLY**

Label indicating the wiring diagram are placed on the inner part of the plate who covers the electrical connections



**RISK OF INJURY DUE TO ELECTRICAL SHOCK!**

- Unit may contain electrical current.
- Please don't open system housing
- Before connecting to main power supply, pay attention to tension and frequency displayed on label
- Unit must be protected by circuit breaker differential switch
- To avoid electric shock danger, the unit must be wired with grounded connection
- Please don't exceed the electrical power medical unit as made for, which can be found on the labels applied on the device
- Please don't connect adapters between the device's electrical sockets and plugs

## 7) INSTALLATION OF MEDICAL GASES



**GAS CONNECTION AND TESTING TO BE CARRIED OUT BY CERTIFIED GAS INSTALLER**

The gas terminal units and pipeline system in the unit have been fully factory leak tested. After connection to the mains supply, the entire medical gas pipeline system is to be tested before being commissioned for use.

Always respect the regulations associated with the installation of medical gases, all this checks has to be documented, according to the instructions for testing and verification of distribution networks

- Analysis of Fluid
- Proof crossings
- Proof of obstructions



Do not use oil or grease on any of the gas terminal units or pipe work for any reason as this could lead to a fire or an explosion. Use only approved oxygen compatible lubricants.



**8) PROTECTION AGAINST FLAMMABLE MIXTURES**

Not protected - RISK OF INJURY DUE TO EXPLOSION – Not suitable for installation in places where can produce a flammable anesthetic mixture with air, oxygen or nitrous oxide



## 9) RISK OF INJURY DUE TO UNAUTHORIZED USE!

- UCI-OVAL Unit must not be tampered with or used differently from the intended use
- "ATTENTION": to avoid the risk of electrical shock, the bed head units must be connected to power with protective ground connection"
- do not connect the ground manifold of the bed head units to external equipment
- execute the fixing of equipment following the directions given on the layout and wiring diagram, using the appropriate anchors (brand Fischer, HILTI or similar)
- before connecting the device to main power have a particular attention to the plate reporting the voltage and frequency power
- don't exceed the electric power for which bed head units was built and labeled on the plate, don't connect adapters between the electrical sockets and plugs of device. the bed head units must be protected by appropriate differential switch.
- for the replacement of worn parts use components of same features
- when removing broken equipment, unpowered the equipment (using the switch on the general panel), and ask qualified personnel to replace the equipment
- the equipment must not be tampered or used for different purposes than providing issues
- the equipment unit must be installed and tested and used in accordance with the standards EN7396 by a qualified staff
- for the proper installation of gas outlets follow the instructions of manufacturer of gas outlet.
- equipment not sterilizable
- the user is not authorized to replace the components (such as transformers, relays, fuses, etc.), always use qualified personnel
- the bed head units requires special precautions regarding EMC (electromagnetic compatibility) and must be installed and put into service according to the information listed in the table 1 and 2, present in user manual, relative to the information on EMC (electromagnetic compatibility)
- the MD unit or MD system should not be used close to other equipment, if you were to use the MD system or the MD unit near or overlapping other equipment, it is necessary to keep checking the normal operation in the configuration in which it is used
- portable RF communications EQUIPMENT, including antennas, can effect medical electrical EQUIPMENT. The MD unit should be used no closer than 30cm (12 inches) to any part of portable RF communications EQUIPMENT, including antennas and cables specified by the manufacturer.
- other cables and accessories may negatively affect EMC performance.



## 10) RISK OF INJURY DUE TO SYSTEM PARTS FALLING DOWN

- Please don't overload medical supply unit's max load, listed on the related label.



## 11) RISK DUE TO UNQUALIFIED INSTALLATION!

- Medical supply unit installation must be done following indication listed on manufacturer installation manual.
- Electrical connection must be done following the manufacturer electrical scheme.  
Gas outlet distribution must be done by qualified personnel according to EN 7396-1 and EN 7396-2 regulations. Distribution medical gas system must be tested and prewired before delivery.



## 12) RISK DUE TO SPARE PARTS NOT AUTHORIZED BY THE MANUFACTURER!

- Please make sure to replace damaged parts with original parts only
- Please replace damaged screws with new ones with the same specification.

## 13) TECHNICAL SPECIFICATIONS

Regulation	The equipment should be considered class IIb
Classification according to the type of protection against electrical hazards	Class I device. The protection against electric shock is guaranteed by the metal parts of the ground protection
Classification by type of security against direct and indirect contact	The device is not equipped with applied parts
Classification according to the degree of protection against penetration of liquids and external agents	IP20
Classification according to the use and conditions	device for continuous operation
Noise level	Less than 35 dB
Testing in production for each single unit	For each unit the following tests are performed: -grounding impedance protection in accordance with 8.6 of standard CEI EN 60601-1 - standard measurement of leakage current and dielectric strength in accordance with 8.7 , 8.8, 16.6 of EN 60601-1 - Tests on medical gas and vacuum distribution in accordance with 12.3, 12.4, 12.5, 12.6 of EN ISO 7396-1 - Testing facilities on evacuation of anesthetic gases in accordance with 12.2, 12.3, 12.4 of EN ISO 7396-2 Tests carried out on end-of-line product form an integral part of this manual
Electromagnetic interference	The operation of other devices placed near the medical device (as portable equipment or furniture) can cause electromagnetic interference or other interference, always check with qualified personnel
Supply voltage	220-230 V 50/60z
Auxiliary voltage (usually)	12Vdc-12Vac, 24Vdc-24Vac
Power consumption - Lighting - Electrical socket	Max 150W Max 2000W for each socket
Replaceable components from technical support staff	All the electrical components present in the ""Parts list"
Protection provided on external power circuit	Provides adequate protection with circuit breakers or fuses dimensioned according to the power indicated on the label
Reference standards	ISO 11197 CEI EN 60601-1 EN ISO 14971 EN ISO 15223-1 EN 20417 EN ISO 7396-1 EN ISO 7396-2 EN ISO 5359
Documentation available on request by the user	The manufacturer is committed to providing the circuit diagrams, component parts lists, calibration instructions or other information that the technical assistance needs to repair parts of replaceable component
Registered office of the manufacturer and production site	Parque tecnológico de Asturias P10, 33428 Llanera-Asturias-España
Unit of Measures	[mm]

## 14) REPLACEMENT PARTS OF LIGHTING SYSTEM

The UCI-OVAL unit are equipped with high-efficiency Led modules with a useful life of 60,000 hours that guarantee operation without maintenance or replacements throughout the useful life of the medical device.

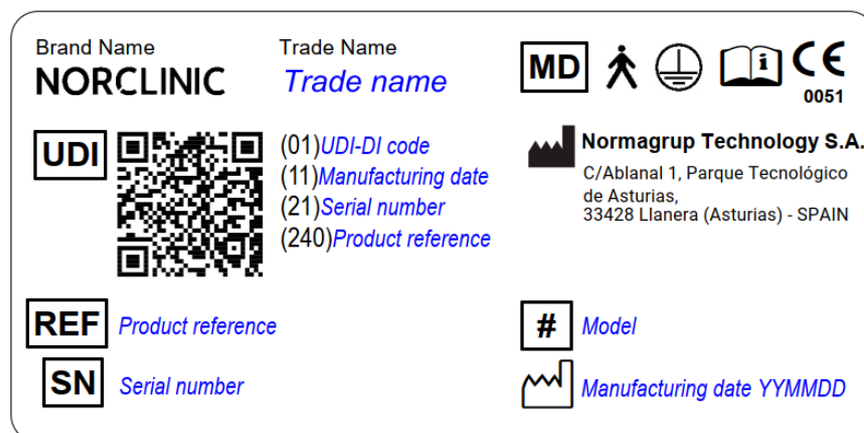
However if it is necessary to replace a module or power supply (ONLY BY PERSONNEL NORMAGRUP OR AUTHORIZED PERSONNEL BY NORMAGRUP) make sure that the external power supply is turned off, then remove the polycarbonate top. The reflector can be accessed as follows: the indirect and night light are found on top of this . By unplugging the connector, you can remove the lighting group in order to replace the Led modules or electrical components (driver).

## 15) LABELLING

A label with the following information is placed on all UCI-OVAL units at the top right of the profile



Below an example of label for model UCI-OVAL:



Indicate the serial number



Indicate the date of production



Indicate the reference of the product



Please read instructions before installation



CE mark, the product complies with regulation (EU) 2017/745. Notified body: IMQ S.p.A. (0051)



Medical Device



Protected earth connection



Electrical safety type B



Manufacturer

THE USER AND/OR PATIENT NEED TO REPORT ANY SERIOUS INCIDENT IN RELATION TO THE DEVICE TO THE MANUFACTURER AND THE COMPETENT AUTHORITY OF THE MEMBER STATE IN WHICH THE USER AND /OR PATIENT IS ESTABLISHED

## 16) AMBIENT CONDITIONS DURING TRANSPORT AND STORAGE

Ambient Temperature: -10 - +50°C  
 Relative humidity (non-condensing): 10-90%  
 Atmospheric pressure: 70-110kPa

Do not overlap more than 2 devices and do not place them vertically. In case of irreparable damages or end-of-life equipment, the device must be disposed following the existing rules. If the user does not follow the rules above, he assumes full responsibility for creating potential effects to the environment and to human health

## 17) AMBIENT CONDITIONS DURING OPERATION

Ambient Temperature: +10 - +40°C  
 Relative humidity (non-condensing): 10-90%  
 Atmospheric pressure: 70-110kPa

## 18) RECOMMENDED MAINTENANCE

Regular inspections must be carried out according to the following specifications

Functionality test and visual inspection of the whole supply unit	Every 6 months by specialized service personnel
Symbol and labels are complete and legible	
Inspection of all cables, electrical sockets, gas outlets, accessories..	

If inside the power unit a flexible prepping for medical, anaesthetic and VAC gas are installed, you can find inspection points on the units and the suspension structure, in order to guarantee a proper inspection. We recommend a regular inspection every 6 months. Flexible prepping replacement every 8 years is required. The new flexible prepping must be conform to standards laid down by EN ISO 5359.

After installation, tests must be perform in order to match standards laid down by EN ISO 7396-1 and EN ISO 7396-2.

## 19) CLEANING

The unit may be cleaned using a soft cloth with clean water mixed with mild non-abrasive detergent.  
 Do not use active solvents to clean the plastic parts.  
 Care should be taken not to expose the device excessively to fluids.

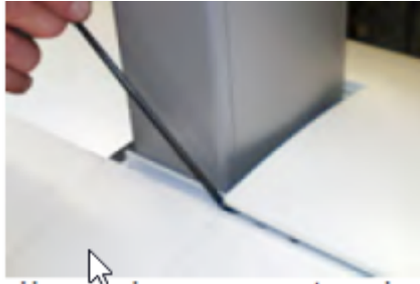
## 20) DISINFECTION OF APPLIED PART

The patient handset and cable may be cleaned and disinfected using a soft cloth with a surface disinfectant cleaner, we recommended for example Dismozon Plus manufacturer: Bode Chemie. with 1,6% Concentration. The application of this product showed good material compatibility and effectiveness.

## 21) INSTALLATION OF CARRIAGE



Separate the covers next to the suspension



Extract the upper covers on both sides of the suspension



Extract the end covers by removing the wingnuts



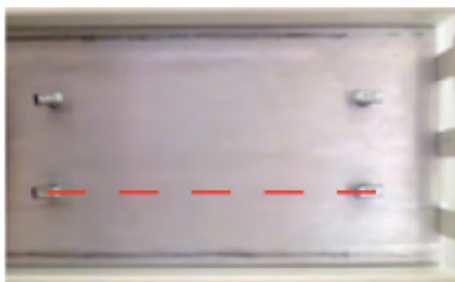
Remove the screws that fix the lower profile



Loosen the nuts on the inside face and leave the heads of the studs aligned to realise the profile



Introduce the carriage in to the profile



On the side where the profile is supported, check that the T-bolts are perfectly aligned



Fit the profile on the suspended unit and turn the T-bolts to retain it. Reassemble all the previous following the inverse process

## 22) GUIDANCE AND MANUFACTURER’S DECLARATION – ELECTROMAGNETIC EMISSIONS

Following Laboratory EMC Test (Emissions and immunity) are performed in accordance with this standards: EN 60601-1-2 (2015)+A1(2021), EN 55015 (2013) / A11 (2020) & EN 61547 (2009).

Table 1. Electromagnetic emissions

The model UCI-OVAL is intended for use in the electromagnetic environment especified below. The customer or the user of the UCI-OVAL model should assure that it is used in such a environment		
Emissions Test	Compliance	Electromagnetic environment – guidance
RF emissions CISPR11	Group 1	The device UCI-OVAL emit electromagnetic energy in order to perorm its intended function, for this its emissions are very low and are not likely to cause any interference in nearby electronic equipment
RF emissions CISPR11	Clase A	
Harmonic emissions IEC61000-3-2	Clase C	
Voltage fluctions/flicker emissions IEC61000-3-3	Comply	

NOTE: The EMISSIONS characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment

## 23) GUIDANCE AND MANUFACTURER'S DECLARATION – ELECTROMAGNETIC IMMUNITY

Following Laboratory EMC Test (Emissions and immunity) are performed in accordance with this standards: EN 60601-1-2 (2015)+A1(2021), EN 55015 (2013) / A11 (2020) & EN 61547 (2009).

Table 2. Electromagnetic immunity

The model UCI-OVAL is intended for use in the electromagnetic environment specified below. The customer or the user of the UCI-OVAL model should assure that it is used in such a environment		
IMMUNITY Test	test level and required electromagnetic environment	Electromagnetic environment – guidance
Electrostatic discharge (ESD) IEC61000-4-2	Contact discharge: $\pm 8\text{kV}$ Air discharge: $\pm 15\text{kV}$	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the humidity should be at least 30%
Fast transient electrical disturbances (bursts)(IEC 61000-4-4)	Power cable: $\pm 2\text{kV}$ Longer signal input lines/output lines: $\pm 1\text{kV}$	Mains power quality should be that of a typical commercial or hospital environment
Impulse voltages (surges) (IEC 61000-4-5)	Voltage, external conductor – external conductor: $\pm 1\text{ kV}$ Voltage, external conductor – protective groundconductor: $\pm 2\text{ kV}$	Mains power quality should be that of a typical commercial or hospital environment
Magnetic fields at mains frequency(IEC 61000-4-8)	50 Hz: 30 A/m	Power frequency magnetic fields should be at levels charateristic of a location in a typical commercial or hospital environment
Voltage dips and short interruptions in the supplyvoltage (IEC 61000-4-11)	Voltage dips of 100% and 30%, with 10ms, 20ms and 5s,different phase angles	Mains power quality should be that of a typical commercial or hospital environment
Radiated high-frequency disturbances(IEC 61000-4-3)	80 MHz to 2.7 GHz: 3 V/m	Mains power quality should be that of a typical commercial or hospital environment
Conducted high-frequency disturbances(IEC 61000-4-6)	150 kHz to 80 MHz: 3 V, ISM bands: 6 V	Mains power quality should be that of a typical commercial or hospital environment
Electromagnetic fields in the vicinity of wireless communication devices	Various frequencies from 385 MHz to 5785 MHz: $9\text{ V/m}$ to $28\text{ V/m}$	Mains power quality should be that of a typical commercial or hospital environment