



Operator's Manual RO MEDICAL-BASIC

Description: NRO-BAS-IFU-1001

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For the reverse osmosis type RO Medical-Basic, conformity according to EC directives is declared

Foreword

This Operator's Manual includes all information required for the installation and operation for the reverse osmosis model RO Medical-Basic.

Please keep this Operator's Manual readily available and near the unit.

This Operator's Manual applies for the units with the serial number:



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to Rm

1. General

1.1. Scope of supply

The scope of delivery includes the following parts:

1 reverse osmosis

1.2. Unit combinations

The unit model RO Medical-Basic may be combined with the following devices:

- Permeate tank
- City water tank

1.3. Notes for the Operator

The operator is responsible for:

- Competent and intended operation
- Compliance with work safety and accident prevention provisions
- Technical instruction of operating personnel

1.4. Laws and Standards

The following laws and standards are adhered to:

- □ Council Directive 93/42 EEC Medical Devices
- EN 60601
- DIN EN 1717 Protection of potable water against contamination

1.5. Symbols used in this Manual



Stands for a dangerous situation. Disregard can result in personal injury or material damage.



Stands for information and valuable tips.



1.6. Transport and Storage



Protect unit against frost and moisture



Protect against strong jolting and collisions.



Only move unit upright and with an appropriate lift.



The system may be stored for a maximum of 1 year.

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1.7. Model Plate



<u>!</u>	Attention, take note of accompanying documents	
C € ⁰²⁹⁷	CE mark with the number of the notified body. Here DQS	
IPX 4	Protection against the ingress of liquids. Here splash-water protection	
SN	Serial number	
	Manufacturer	
	Pay attention to manual	

1.8. Warning on the Unit.



Caution voltage. Turn mains switch off before opening housing. Fixed on control cabinet.

1.9. Shutdown

If a unit is shut down for more than 5 days, conservation will be necessary.



Please contact Nipro Pure Water before performing conservation.

1.10. Disposal

Regarding the WEEE guidelines of the European Union, the disposal of electronic devices and electronic sub-assemblies and parts into the general garbage is not lawful. These parts must be disposed environmentally appropriate:

If not appointed otherwise and no private disposal management is available, these devices or possibly other environmental hazardous items can be sent back.

The filters and membrane can be disposed via the general garbage.

1.11. Instruction / Further Documentation

The using personnel must be warned against the hazards during operation and must be warned against the hazards of misusing the product.

The personnel gets the instruction of operation and the specialties of usage. Instructed adult only are allowed to operate this device.

This instruction by the manufacturer or authorized personnel takes place during the commissioning of the device.

Further trainings are not necessary for this device.

For qualified personnel the following documents can be made available upon request.

Circuit diagrams Spare parts list

1.12. Duration of usage

The device is designed for a use of 10 years



2. Intended operation

The unit is designed for the treatment of potable water. The pure water (permeate) thus produced may be used for dialysis treatment.

Other applications are only possible after consulting the manufacturer and receiving their approval.



The unit can only be maintained by the manufacturer or technicians trained by the manufacturer.



Only original replacement parts may be used for maintenance and repairs.



Installation operations, modifications or reparations, are only allowed to be performed by persons authorised by the manufacturer and may only be done with original replacement parts. Improper performed reparations or modifications can lead to hazards to the user and/or may damage the device. The device may only be operated in perfect condition.



Before operating, check the following:

- Lose or defect parts
- Defect cables and/or isolations
- Serious soiling



The device may only be operated with the appropriate ring line.



The device does not produce water for injections.



The device has pressurized parts.



If the temperature sensor fails, the temperature in the permeate can increase. (Max 60° C)



The water treatment system RO Medical-Basic may only be used for permeate supply of dialysis devices, which have a temperature measurement (permeate temperature).



The device has no direct patient contact and no patient application part.

2.1 Contraindications / side effects

None

3. Safety

3.1 Risk Assessment

There will be no dangers associated with the reverse osmosis model RO Medical-Basic D if the operating instructions are followed.



The device can automatically start by way of an auto-start.

3.2 EMC

The device was developed and tested in accordance with current standards. Nevertheless, influence through electromagnetic fields cannot be completely excluded.

3.3 Emissions

The device does not produce dust or vibrations. The noise level is under 60 dB (A).



4. Technical Data

Permeate performance

Temperature	1 Membrane	2 Membranes	3 Membranes	4 Membranes
15° C	350 l/h	700 l/h	1050 l/h	1400 l/h

Inlet water

Quality	Potable Water
Hardness	< 1 °dH
Silicate	< 25 mg/l
Chlorine	< 0,1 ppm (mg/l)
Iron	< 0,1 ppm (mg/l)
Fouling Index (S.D.I)	< 3
Temperature	5-25°C

Connections

Water feed	1" internal thread
Pure water connection	Hose nozzle d20
Drain	HT 40

Electrical data

Supply voltage	220-230 V, 1 Phase, 50/ 60 Hz
Fuse	Automat 16 A-K, Fi Δl 30mA
Current consumption	9,8 A x
Degree of pollution	1

Ambient temperature

Storage / transport	3-40°C
Operation	10-35°C
Air pressure	795-1062 hPa

Display system

Conductivity	0-1000 µS/cm ±5%
pressure switch	0-10 bar ±5%
Flow (sight glass)	300-3000 l/h ±5% 100-1000 l/h

Size

RO Medical-Basic 1000x500x1640



5. Description of the device

5.1 Flow-Chart



- 1. Magnetic valve inlet
- 2. Pressure switch input
- 3. Disinfection point
- 4. Pump (pressure 10-13bar)
- 5. Manometer (pressure 10-13 bar)
- 6. Reverse osmosis membrane (1-4 pieces)
- 7. Permeate flow rate indicator

- 8. Temperature sensor
- 9. Conductivity sensor permeate flow
- 10. Permeate pressure switch
- 11. Connection flow hose nozzle d20
- 12. Connection back flow hose nozzle d20
- 13. Drain

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- 14. Shut-off valve permeate drain
- 15. Check valve
- 16. Flow indicator concentrate drain

5.2 Functional sequence

If the toggle switch is turned to the 'On' position, the magnetic valve (1) opens and water flows into the system. After a short delay, the pump (4) will start.

Now the water is pressed into the reverse osmosis membrane at a pressure of 10-15 bar. The flow is divided into a permeate and a concentrate part. The permeate content flows through the flow indicator (7), the temperature sensor (8), the conductivity sensor (9) and the pressure switch (10) into the ring line. The unused permeate flows back into the RO Medical-Basic via the connection (12).

To save water, the concentrate portion is divided up again, one portion is given into the drain via the flow indicator (16), the other is fed back in front of the pump.



5.3 Safety devices / Components



GIT+ TARM

6. Installation



The installation must be conducted by the manufacturer or by personnel trained and authorized by the manufacturer.

6.1 Environmental Condition

Conditions for the osmosis room:

- Relative air moisture < 90% at 20°C
- Room temperature between +10°C and +35°C (frost protected)
- Equipped with floor drain, water supply and electrical supply

6.2 Assembly

- Bring the device into the appropriate position
- Adjust machine feet until the device stands level and secure on the floor.



Do not store easily flammable materials in the vicinity of the device.



Do not store chemicals in the vicinity of the device.

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Only operate the device with the necessary water pre-treatment.



Room of osmosis may not be freely accessible. (Access for instructed personnel only)

6.3 Electrical installation



The installation may only be performed by a qualified electrician.



The device must be supplied by a permanent connection, connectors are not valid. The disconnection via the main switch at the control cabinet. The power cord must be provided with a strain relief.



Connection cable RO Medical-Basic



For protection against a re-start of the unit, the main switch can be locked with a padlock.

Safety class I



The device is equipped with a Protective earth terminal for prevention against high touch current For prevention of the hazard of an electric shock, this device may only be connected to a power supply with protective earth.



The power cord is fixed to the device and cannot be replaced.



6.4 Prefiltration (Example)

Install the necessary water pre-treatment equipment first! Only then connect the RO Medical-Basic and start up





Local water works regulations and DIN EN 1717 must be followed.



The water pre-treatment must be adapted to the local potable water quality.



6.5 Commissioning



Caution, device damage!

The device must be preconnected by a suitable pre-filter and a softening system as well as a pressure reducer.



1+1000

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Permeat back flow (3) Hose nozzle d20

6.6 Initial installation







- Connect and check the raw water connection (1), permeate flow (2), permeate back flow (3) and waste water (4)
- 2. Open the inlet valve manually. To do this, turn the white magnet coil 45° counter clockwise.

Now water is running into the device.

- 3. Carefully open the screw for the pump venting until a water outlet can be seen. Then close the screw again and reset the solenoid.
- 4. Make sure that the wall-mounted taps at the start and end of the ring are open.





- 5. Start the device using the toggle switch.
- 6. Check all connections for leaks.
- 7. Open the shut-off valve permeate to the drain.
- 8. Allow the device to discard permeate for at least 30 minutes.



7. Operation

7.1 Control Panel



This is used to switch the device on and off

- Fault light
 Lights up when there is a fault

 Diamony
- **3. Display** Display of conductivity and faults

th Key switch emergency operation The device can be switched to emergency operation here.

5th Main switch

With this the device can be switched off completely



7.2 System on / off without permeate tank



System on To start the device, turn the toggle switch to the right (on position). The green lamp lights up. The pump starts after a short delay



System off Reset toggle switch (position 0) The green lamp goes out. Pump stops.



If the device is switched off using the toggle switch, the rinse intervals are carried out as programmed. If no rinsing is to take place, the device must be switched off completely at the main switch. However, this is only recommended for decommissioning.



7.3 Device on / off with permeate tank



If the RO Medical-Basic is connected to a permeate tank, the toggle switch must be set to "Auto". Since the device switches off automatically when the tank is full, it is not necessary to reset it to position 0.



Caution, danger of overflow!

If the RO Medical-Basic is operated with a permeate tank, the system may only be started via "Auto". In the "On" position there is a risk of overflow.



Device on

To start the system, turn the knob switch to the right. If the permeate tank is empty, the system starts automatically. If the tank is already full, the osmosis waits to start until the level in the tank drops.



7.4 Emergency operation



Only use emergency operation if the automatic function fails. Have device repaired as soon as possible.

Attention!

There is no monitoring of the water inflow during the emergency operation. Therefore a continuous water inflow has to be guaranteed. Absent water causes the **destruction of the pump**.



1. Open the inlet valve manually. To do this, turn the white magnet coil 45° counter clockwise.

Now water runs into the device.



2. Set the key switch to position 1. Pump starts up



To turn off the device, turn switch to position 0.



In emergency operation all automatic functions are turned off. No cleaning cycle and no automatic start and/or stop will be conducted.



7.5 Operating displays



Device is off. Rinsing interval (target) is set to 90 minutes. 25:22 min have already passed. If the actual value reaches the set point, the system goes into rinsing.



Device is on. The current conductivities are displayed.



Device is being rinsed. The current conductivity is displayed.



8. Error messages / troubleshooting

8.1 Error messages

Display	Error Description	Troubleshooting
Fehler Schütz Pumpe ausgelöst 1Q5 prüfen	The motor protection switch of the pump has triggered. Check motor protection switch. If this alarm occurs frequently, the pump must be checked.	Turn the motor protection switch back to position 1.
Fehler Übertemperatur 16S13 prüfen	The temperature of the permeate has reached to 38 °C. The system switches off to protect the membranes.	The system must be cooled (see next page).
Fehler Überdruck 16S4 prüfen	The permeate pressure has exceeded 6 bar.	Check the start and end of the ring taps. Check setting of permeate pressure retention valve.
Fehler Wassermangel 16S5 prüfen	The pressure switch 1 has responded.	Check water inlet.
Fehler Leitfähigkeit	The conductivity has exceeded the limit of 100µS/cm.	There is probably a defect in the membranes. Call service.

8.2 Clear alarms

Reset the error message Follow the notes in Display.



Reset the alarm by pressing the F3 key



9. Maintenance and cleaning

9.1 External Cleaning

Stains and dust can be removed with a cloth and a commercially available cleaner.



Do not clean the device with solvents.



Stains from softening salts or disinfectants must be removed immediately.



9.2 Maintenance Intervals

Measure	period	Notes	user
Fill salt at softener	Daily		user
Chemical disinfection	lf needed		Manufacturer or persons authorized by manufacturer
Maintenance	Yearly		Manufacturer or persons authorized by manufacturer
Safety related check	Every 2 years		Manufacturer or persons authorized by manufacturer
Microbiological analysis	Every 3 months		User
Chemical analysis	Every 12 months		User



Not replacing the filter or replacing it too late can lead to damaging of the reverse osmosis.



9.3 Chemical Disinfection



A chemical disinfection should only be performed upon new installation or when high pathogen values are encountered



Disinfection may only be performed by **Nipro Pure Water** or by **instructed** persons.



Caution when handling disinfectants! Per acetic acids can cause damage to your health. Always **read** safety **guidelines** before handling.



Before the next dialysis each consumption point must be tested for disinfectant traces.



9.4 Microbiological Inspection

Necessary Values¹

- Pathogens < 100 CFU/ml no traces of Pseud. aeruginosa and E. coli
- Endotoxins < 0,25 EU/ml

Inspection Interval²

Inspection of permeate every 3-6 months.

Inspection method ²

Pathogen count determination:

Nutrient medium: TGEA (OXID Nr.CM 127), R2A Incubation temperature: 22°C ± 2°C

Endotoxins determination:

Method: GEL-Clot; Cromogen; Turbid metric



¹ According to the European Pharmacopoeia

² Recommendations according to the guideline for the practice of applied hygiene in treatment units for dialysis

10. Display / Parameter

10.1 Retrieval of the operating hours / conductivity records





10.2 Change the conductivity parameter



Press the F4 key and F2 key simultaneously.

The conductivity in the return is displayed.

Press the ESC key for 3 seconds, the first value is marked.

Press enter

Set the value with the arrow keys (up / down). Complete the entry with ENTER.

Press the ESC key.

Press the F4 and F2 keys simultaneously to return to the standard display



Both parameters (MAX1 and MAX2) must be set to the same value.



Standard value = 100 µS / cm



If the conductivity in the back flow reaches the value set here, the conductivity alarm is activated.



10.3 Change parameter temperature

SIEME	NS	LOG	D! TDE		
M	AX1 =	: 38 : 38			
-		-		L	
	SIEME M M	MAX1 = MAX2 =	MAX1 = 38 MAX2 = 38	SIEMENS LOGO! TDE MAX1 = 38 MAX2 = 38	SIEMENS LOGO! TDE MAX1 = 38 MAX2 = 38

Press the F4 key and F3 key simultaneously.

The temperature is displayed.

Press the ESC key for 3 seconds, the first value is marked.

Select corresponding value with the arrow keys.

Press enter

Set the value with the arrow keys (up / down). Complete the entry with ENTER.

Press the ESC key.

Press the F4 and F3 keys simultaneously to return to the standard display



Both parameters (MAX1 and MAX2) must be set to the same value.



Default value = 38°C



If the temperature reaches the value set here, the over temperature alarm is activated. CAUTION! Max. 40°C is allowed, higher temperatures damage the membranes.



Technical appendix



The settings and functions described below may only be carried out by technically trained personnel.



ATTENTION. DANGER TO PERSONS AND SYSTEM TECHNOLOGY! Incorrect settings can lead to hazards.



No service or maintenance work may be carried out during treatment.



11. Replacement of the reverse osmosis membrane



Caution pressure!

Membrane tubes are under pressure. Please open carefully.

The settings and functions described below may only be carried out by technically trained personnel.

1. Switch off the system at the main switch.



2. Open wing screw and take off the clip



3. Lift module cover by using a plier



4. Take off cover.



- 5. Pull out the membrane by using a plier
- Reinstall the new membrane in reverse order. Take care of the flow direction and position of the gasket



Rinse membrane! After the new membrane has been installed the mode "drain permeate" has to be started for 20 minutes.



12. Service Parameters



The settings and functions described below may only be carried out by technically trained personnel.



ATTENTION, SYSTEM DAMAGE! Incorrect settings can lead to damage.



Arrow key down and then press the ESC key. (Service mode)

Then Logo settings> ENTER> Program> ENTER> Set parameters> ENTER> select the appropriate parameter (B001-B040)

Parameter	Function	default value
T Pump	Pump start time delay	3sec
Time to Rinse	Rinse waiting time	60m
T Rinse	Rinse time	10m
T Conductivity	Conductivity alarm delay at start	2m
W Qty NOK	Dry running protection	5sec
T2 Conductivity	Conductivity alarm delay at operation	30sec
B025	Hours count	
T Temperatur	Temperature alarm delay	10sec
Conductivity	Conductivity adjustment (permeate)	
B040	Timer (auto start)	



12.1 Adjustment of the conductivity

- Select parameter Conductivity.
- "Ax" value = displays conductivity
- Select value "B" and confirm with Enter.
- Change the value with the arrow keys. (+00001 = current value +1)
- To change in minus, first enter the value and after the change the sign (-00001 = current value -1)
- press enter
- Press ESC several times (until time is displayed), then arrow key up.

12.2 Service point

The service point can be used for water withdrawal to check water temperature and conductivity.





Service point may only be opened, if operating pressure in the system is reached.

Service point must be closed prior to each system start. Service point may only be operated by **technically trained personnel**.



12.3 Change Temperature Parameter



Turn the locking ring to the unlocked position.

Turn both setting rings to the lowest setting.

This is important to ensure the setting accuracy.

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▶ Rotate SET ring to desired temperature.

ñ

Rotate RESET ring to desired temperature.

The RESET value must be lower than the SET value.

Minimum distance between SET and RESET = 3 K (= hysteresis).

Turn the locking ring to the locked position.

12.4 Setting date and time

- ▼ key arrow down
- Press ESC
- Select service mode (▼ key arrow down, ESC)
- LOGO Settings <ENTER> Setup <ENTER> Clock <ENTER> Set Clock <ENTER>
- Set the time and date with the arrow keys <ENTER
- To exit press ESC 5x and ▲ arrow key up

Start	Set Clock	Set Clock
Program +	S/W Time -	
Setup +-	Sync	F-1 #F-00
Network +		Fn. 15:30
Diagnostics +		2014-02-07
Card +		

12.5 Summer / winter time

- Select service mode (▼ arrow key down, ESC)
- LOGO Settings <ENTER> Setup <ENTER> Clock<ENTER>S/W Time Select the S/W Time

Start Program	BM AI NUM	Set Clock S/W Time	S/W Time
Setup >	Power-on Delay Msg Config	Sync	oEU
Diagnostics + Card +	Start Screen		

- "④": disables automatic S/W time conversion.

- "(5)" represents the start and end of European summertime.

- "6" represents the start and end of summertime in the United Kingdom.

- " $\overline{7}$ " represents the start and end of daylight-saving time (summertime) in the United States prior to 2007.

- "(8)" represents the start and end of daylight-saving time (summertime) in the United States in 2007 and later years.

- "(9)" represents the start and end of Australian summertime.

- "10" represents the start and end of Australian/ Tasmanian summertime.

- "(1)" represents the start and end of New Zealand summertime.

- "12)": Here you can enter any month, day and time zone difference.

To exit press ESC 5x and ▲ arrow key up

12.6 Auto Start (Timer)

Select service mode

Logo settings < ENTER > Program < ENTER > Set parameters < ENTER > Select parameter B040<ENTER>

To exit press ESC 5x and \blacktriangle arrow key up

B1 1/2	To set the on-/off-times:
D1 =MTWTFSS	-Move the cursor to one of the parameters of the timer.
On1 =06:30	-Press ENTER. The cursor is positioned on the day of the week.
Off1 =08:00	-Press up and down key to select one or several days of the week.
D2 =	-Press right arrow key to move the cursor to the first position of the
On2 =	on-time.
B1 2/2 Off2 =: D3 = Off3 =: Off3 =: Pulse =Off	-Set the on-time. Modify the value at the respective position, using the up and down keys and move to the cursor to the various positions, using the right and left arrow keys. At the first position, you can only select the value: (: means: No on-/off-times set). -Press right arrow key to move the cursor to the first position of the off-time. -Set the off-time -Confirm your entries with ENTER

The prefix "D=" (Day) has the following meaning: • M: Monday • T: Tuesday • W: Wednesday • T: Thursday • F: Friday • S: Saturday • S: Sunday

Uppercase letters indicate a specific day of the week. A "-" indicates no selection for the day of the week.

13. Disinfection



Disinfection may only be performed by **Nipro Pure Water** or by **instructed** persons.



Caution when handling disinfectants!

Per acetic acids can cause damage to your health. Always **read safety guidelines** before handling.



To be performed precisely!

Danger!

Ensure that no dialysis can be performed while disinfecting. Only approve thoroughly rinsed system for treatment use.



CAUTION!

While using chemicals. Wear safety gloves and safety goggles during the here described jobs.



ATTENTION!

Do not eat, drink or smoke during work.

Disinfectant: MINNCARE® Cold Sterilant (Artikelnr.:489)



Detection method: MINNCARE® Test Strips Residual (Artikelnr.:490) MINNCARE® Test Strips 1 Indication (Artikelnr.:491)











4. Start the device

5.



- Check for correct disinfectant concentration at the
 permeate outlet with test strips. Use Minncare Test Strips Indication 1% for this.
- 6. Let the disinfection pump run until sufficient disinfectant is detected.



7. Close the shut-off valve permeate to the drain.





- 8. Switch off the system
- 9. Restart the system after 20 minutes.
- 10. Open the shut-off valve permeate to the drain.





11. Let the system run until no more disinfectant can be detected at the permeate drain. Use Minncare residual test strips for this.



12. Close the shut-off valve permeate to the drain.

13. Check that all dialysis stations are free from disinfectants.



Switch the system off during disinfection with a conductivity alarm, clear the alarm and restart the system.



RISK OF DEATH !

Before the next dialysis, each sampling point must be tested again to ensure that it is free from disinfectants.



* **Free of disinfection means 0ppm-** no discoloration of the test strip. See the colour scale on the packaging of the test strip Residual.



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13.1 Disinfection Protocol

Desinfektionsprotokoll	/ Desinfection proto		WATER
Datum / date: Ersteller / creator:			
Kundendaten/ customer data:			
Zentrum / centre:			
Straße & Hsnr. / street & no.:			
PLZ & Stadt / zip code & city:		210	
Kunden-Nr/ client-no.:			
Anlagentyp / type of machine:	Phoenix One	Seriennummer / serial number:	
	Phoenix One+	Seriennummer / serial number:	
	Romedical	Seriennummer / serial number:	
	Desinfektion ohne Ringleitu	ng / desinfection without ring line	
	Desinfektion mit Ringleitung	/ desinfection with ring line	
Desinfektion wurde wie folgt durch	geführt/ desinfection carried o	ut as followed:	
Desinfektionsmittel / desinfectant:			
Einspülzeit / induction time:	min	Eingeimpfte Menge / quantity of odorant injected:	3
Ausspülzeit / rinsing time:	min	Wartezeit / waiting time:	min
Desinfektionsmittelfreiheit wurde an desinfectant traces was tested and	n allen Diatyseplätzen geprüft t confirmed at all consumption (ind wird hiermit bestätigt: oints:	
Patra (data		Interaction Constant New York Water	

Wurde eine Desinfektion durchgeführt, verpflichtet sich der Betreiber mit seiner Unterschrift zur nochmaligen Überprüfung aller Dialyseplätze auf Desinfektionsmittelfreihelt. Diese Überprüfung muss vor Beginn der ersten Dialyse durchgeführt werden.

If a desinfection was carried out, the operator commits to do a re-examination for desinfectant traces at each consumption points with his/her signature. This test has to be done before the beginning of the first dialysis.

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Datum / date

Name / Unterschrift Kunde name / signature customer

14. EMC manufacturer's declaration

Electromagnetic emissions and electromagnetic immunity

The RO Medical-Basic device is intended for use in electromagnetic environments as described below.

The customer or the operator of the RO Medical-Basic should ensure that the device is only used in such an environment.

This EMC manufacturer's declaration is based on the use of the power supply unit from Phoenix Contact.

The power supply is installed in the control cabinet.

The cable length between the power supply unit and the cable entry through the housing wall is 150 cm.

Warning

The use of other accessories, other power supply units and cables than specified can lead to increased emissions and/or reduced interference immunity of the RO Medical-Basic.

Requirements

During the interference immunity tests, the temperature accuracy and conductivity accuracy were checked.

Emission measurement	Compliance	Electromagnetic environment - Guidelines
RF emission in accordance with CISPR 11 / EN 5511	Group 1	The device only uses RF energy for its internal function. Its RF emissions are therefore very low and interference to nearby electronic devices is unlikely.
RF emission in accordance with CISPR 11 / EN 55011	Class B	The device is suitable for use at any location, including
Harmonics in accordance with IEC 61000-3-2	Class A	residential areas and facilities directly connected to the
Voltage fluctuations / flickers in accordance with IEC 61000-3-3	Fulfilled	public low-voltage grid for residential buildings.

Immunity test	Test level – IEC	Compliance	Electromagnetic
Discharge of static electricity (ESD) in accordance with EIC 61000-4-2	± 6 kV contact discharge ± 8 kV air discharge	± 6 kV contact discharge ± 8 kV air discharge	The floor should be made of wood, concrete, or of tiles. In case of synthetic flooring, relative air humidity should be at least 30%.
transient burst/immunity test in accordance with IEC 61000-4-4	± 2 kV for power cables ± 1 kV for input and output cables	± 2 kV for power cables ± 1 kV for input and output cables	should comply with that of a typical commercial or hospital environment.
Surge voltage in accordance with IEC 61000-4-5	± 1 kV outer conductor-outer conductor ± 2 kV outer conductor-ground	± 1 kV outer conductor-outer conductor ± 2 kV outer conductor-ground	The quality of supply voltage should comply with that of a typical commercial or hospital environment.
Voltage drops, short interruptions, and fluctuations in supply voltage in accordance with IEC 61000- 4-11	95% voltage drop for ½ period 60% voltage drop for 5 periods 30% voltage drop for 25 periods 95% voltage drop for 5 s	95% voltage drop for ½ period 60% voltage drop for 5 periods 30% voltage drop for 25 periods 95% voltage drop for 5 s	The quality of supply voltage should comply with that of a typical commercial or hospital environment. If the device is to continue functioning uninterruptedly in case of power interruptions, it is recommended that the device be operated via uninterrupted power supply or a battery.
Magnetic field at supply frequency (50/60 Hz) in accordance with IEC 61000- 4-8	3 A/m	3 A/m	In supply frequency, the magnetic fields should comply with the values characteristic of locations in a typical commercial or hospital environment.
Conducted RF disturbances in accordance with IEC 61000-4-6	3 V rms 150 kHz to 80 MHz	3 V rms 150 kHz to 80 MHz	When operating portable or mobile RF communication devices (transmitters), a safety distance should be observed to all parts of the device, including cables, calculated on the basis of one of the following equations depending on the transmission frequency. Recommended safety distance:
Radiated RF disturbances in accordance with IEC 61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m 80 MHz to 2.5 GHz	d = $1.2\sqrt{P}$ 150 kHz to 80 MHz d = $1.2\sqrt{P}$ 80 MHz to 800 MHz d = $2.3\sqrt{P}$ 800 MHz to 2.5 GHz

			Whereby P is the maximum nominal output of the respective transmitter specified by the manufacturer in Watts (W), and d is the recommended safety distance in meters (m). The field strength of stationary RF transmitters, which is definable via electromagnetic site survey a,
			should be below the
			compliance level of the
			individual frequency ranges b.
			Disturbances are possible near
			devices which bear the symbol
			below.
Note: These guide	values may not app	oly to all situations.	Spreading of electromagnetic
animals.			Juliulitys, iterits, persons, and
a. The field streng (mobile/cordless)	th of stationary tran and mobile land mo	ismitters (e.g. base bile radios, amateur	stations of mobile phones r radio stations, AM and FM
radio, and TV tran	smitters), cannot be	theoretically calcul	ated in advance. To identify
electromagnetic si	te survev should be	considered. If the f	ield strength identified at the
location at which t	he device is used ex	ceeds the RF comp	liance level specified above, the
device should be c	losely observed. It i alignment or transi	may be necessary to position of the devic	o take additional measures
(cigi chunging the	anginnent of transp		

b. Across the frequency range of 150 kHz to 80 MHz, the field strength should be less than 3 V/m.



Recommended minimum distances between portable and mobile RF communication devices and the RO

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

Max. output of the transmitter	Minimum distance in accordance with the frequency of the transmitter (m)				
(W)	150 kHz to 80 MHz 80 MHz to 800 MHz 800 MHz to 2.5 GHz				
	d=1.2 √P	d=1.2 √P	d=2.3 √P		
0.01	0.12	0.12	0.23		
0.1	0.38	0.38	0.73		
1	1.2	1.2	2.3		
10	3.8	3.8	7.3		
100	12	12	23		

For transmitters with a maximum output not specified above, the recommended distance d in meters (m) can be calculated in accordance with the equation appropriate for the frequency of the transmitter, whereby P is the maximum output of the transmitter in Watts (W) in accordance with the specifications of the manufacturer.

NOTE 1:

For 80 MHz and 800 MHz, the safety distance applies for the higher frequency range.

NOTE 2:

These guide values may not apply to all situations. Spreading of electromagnetic waves is also influenced by absorption and reflection via buildings, items, and persons.