Алматы (7273)495-231 Ангарск (3955)60-70-56 Архангельск (8182)63-90-72 Астрахань (8512)99-46-04 Барнаул (3852)73-04-60 Белгород (4722)40-23-64 Благовещенск (4162)22-76-07 Брянск (4832)59-03-52 Владивосток (423)249-28-31 Владикавказ (8672)28-90-48 Владимир (4922) 49-43-18 Владимир (4922) 49-43-16 Волгоград (844)278-03-48 Вологда (8172)26-41-59 Воронеж (473)204-51-73 Екатеринбург (343)384-55-89 Ижевск (3412)26-03-58 Иваново (4932)77-34-06 Иркутск (395)279-98-46 азань (843)206-01-48 Калининград (4012)72-03-81 Калуга (4842)92-23-67 Кемерово (3842)65-04-62 Киров (8332)68-02-04 Коломна (4966)23-41-49 Котрома (4942)77-07-48 Кострома (4942)77-07-48 Краснодар (861)203-40-90 Красноярск (391)204-63-61 Курск (4712)77-13-04 Курск (4712)50-90-47 Липецк (4742)52-20-81

Магнитогорск (3519)55-03-13 Москва (495)268-04-70 Мурманск (8152)50-54-93 Набережные Челны (8552)20-53-41 Нижний Новгород (831)429-08-12 Новокузнецк (3843)20-46-81 Ноябрьск (3496)41-32-12 Новосибирск (383)227-86-73 Ноябрьск (3496)41-32-12 Омск (3812)21-46-40 Орел (4862)44-53-42 Оренбург (3532)37-68-04 Пенза (8412)22-31-16 Петрозаводск (8142)55-98-37 Псков (8112)59-10-37

Пермь (342)205-81-47 Ростов-на-Дону (863)308-18-15 Рязань (4912)46-61-64 Самара (846)206-03-16 Саранск (8342)22-96-24 Саратов (845)249-38-78 Саратов (845)249-38-78 Севастополь (8692)22-31-93 Симферополь (3652)67-13-56 Смоленск (4812)29-41-54 Сочи (862)225-72-31 Сочи (852)225-72-31 Ставрополь (8652)20-65-13 Сыктывкар (8212)25-95-17 Сургут (3462)77-98-35 Тамбов (4752)50-40-97

Казахстан (772)734-952-31

Тверь (4822)63-31-35 Тольяти (8482)63-91-07 Томск (3822)98-41-53 Тула (4872)33-79-87 Тюмень (3452)66-21-18 Улан-Удэ (3012)59-97-51 Ульяновск (8422)24-23-59 Уфа (347)229-48-12 Хабаровск (4212)92-98-04 Чебоксары (8352)28-53-07 Челябинск (351)202-03-61 Череповец (8202)49-02-64 Чита (3022)38-34-83 Якутск (4112)23-90-97 Ярославль (4852)69-52-93

Киргизия (996)312-96-26-47

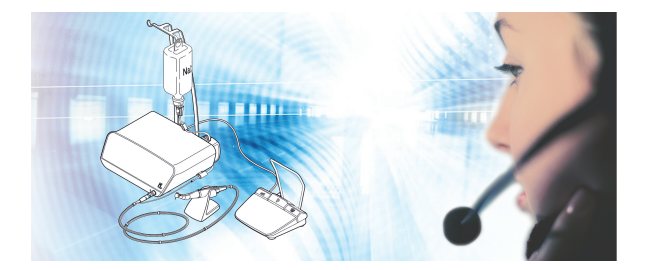
https://kavo.nt-rt.ru || koc@nt-rt.ru

Россия (495)268-04-70

# Физиодиспенсер EXPERTsurg LUX

# Instructions for use

# **EXPERTsurg LUX** REF 1.008.3500



### 1.4 Terms and conditions of warranty

KaVo provides the final customer with a warranty that the product specified in the delivery note functions properly and is free of defects in the material or workmanship for a period of 12 months from the date of purchase, subject to the following conditions: Upon justified complaints of flaws or a short delivery, KaVo will make good its warranty by replacing the product free of cost or repairing it according to the customer's wishes. Other claims of any nature whatsoever, in particular with respect to compensation, are excluded. In the event of default and gross negligence or intent, this shall only apply in the absence of mandatory legal regulations to the contrary.

KaVo shall not be held liable for defects and their consequences due to natural wear, improper cleaning or servicing, non-compliance with operating, servicing or installation instructions, calcification or corrosion, contaminated air or water supplies or chemical or electrical factors deemed abnormal or impermissible in accordance with factory specifications.

The warranty does not usually cover bulbs, glassware, rubber parts and the colourfastness of plastics.

Defects or their consequences that can be attributed to interventions on or changes made to the product by the customer or a third party are excluded from the warranty. Claims from this warranty can only be asserted when the delivery note of the product has been sent to KaVo, and the original can be presented by the operator or user.

# 1.5 Transportation and storage

### 1.5.1 Currently valid packaging regulations



### Note

Only valid for the Federal Republic of Germany.

Dispose of and recycle the sales packaging appropriately in accordance with current packaging regulations, employing waste management or recycling companies. Comply with the comprehensive return system. KaVo has had its sales packaging licensed for this purpose. Please comply with the regional public waste-disposal system.

### 1.5.2 Damage in transit

### In Germany

If the packaging is visibly damaged on delivery, please proceed as follows:

- 1. The recipient of the package must record the loss or damage on the delivery receipt. The recipient and the representative of the shipping company must sign this delivery receipt.
- 2. Leave the product and packaging in the condition in which you received it.
- 3. Do not use the product.
- 4. Report the damage to the shipping company.
- 5. Report the damage to KaVo.
- 6. Consult with KaVo first, before returning a damaged product.
- 7. Send the signed delivery receipt to KaVo.

If the product is damaged but there was no discernable damage to the packaging upon delivery, proceed as follows:

1 User instructions | 1.5 Transportation and storage

- 1. Report the damage to the shipping company immediately and no later than 7 days after delivery.
- 2. Report the damage to KaVo.
- 3. Leave the product and packaging in the condition in which you received it.
- 4. Do not use a damaged product.



If the recipient fails to comply with any of the above-mentioned obligations, the damage will be considered to have arisen after delivery (in accordance with the General German Freight Forwarders' Terms and Conditions, Art. 28)

# **Outside Germany**



### Note

Note

KaVo shall not be held liable for damage arising from transportation. The shipment must be checked on arrival.

If the packaging is visibly damaged on delivery, please proceed as follows:

 The recipient of the package must record the loss or damage on the delivery receipt. The recipient and the representative of the shipping company must sign this delivery receipt.
 Without this ovidence, the recipient will not be able to assort a claim for damages.

Without this evidence, the recipient will not be able to assert a claim for damages against the shipping company.

- 2. Leave the product and packaging in the condition in which you received it.
- 3. Do not use the product.

If the product is damaged but there was no discernable damage to the packaging upon delivery, proceed as follows:

- 1. Report any damage to the shipping company either immediately or no later than 7 days after delivery.
- 2. Leave the product and packaging in the condition in which you received it.
- 3. Do not use a damaged product.



### Note

If the recipient fails to comply with any of the above-mentioned obligations, the damage will be considered to have arisen after delivery (in accordance with CMR law, Chapter 5, Art. 30).

### 1.5.3 Information on the packaging: Storage and transportation



### Note

Please keep the packaging in case you need to return the product for servicing or repair.

The symbols printed on the outside are for transportation and storage, and have the following meaning:

<u><b>†</b></u>	Transport upright with the arrows pointing upwards!
Y	Fragile - protect against impact!
Ť	Protect from moisture!

1 User instructions | 1.5 Transportation and storage

kg max	Permissible stacking load
Ĵ, Ĵ	Temperature range
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Humidity
hPa hPa	Air pressure

2 Safety | 2.1 Description of safety instructions

# 2 Safety

# 2.1 Description of safety instructions

# 2.1.1 Warning symbol



# 2.1.2 Structure



The introduction describes the type and source of the hazard.

- This section describes potential consequences of non-compliance.
- The optional step includes necessary measures for hazard prevention.

# 2.1.3 Description of hazard levels

Safety instructions distinguishing between three hazard levels are used in this document to prevent personal and property damage.



### CAUTION

# 

indicates a hazardous situation that can cause damage to property or mild to moderate injuries.



# WARNING

indicates a hazardous situation that can lead to serious or fatal injury.



### DANGER

\Lambda DANGER

indicates a maximal hazard due to a situation that can directly cause death or fatal injury.

# 2.2 Information about electromagnetic compatibility

### Note

Based on IEC 60601-1-2 (DIN EN 60601-1-2) concerning the electromagnetic compatibility of electrical medical devices, we must draw your attention to the following points:

• Medical electrical devices are subject to special precautions concerning the electromagnetic compatibility and must be installed and operated in accordance with the KaVo assembly instructions.

• High-frequency communications devices may interfere with electrical medical devices.

### See also:

12 Information about electromagnetic compatibility, Page 64



2 Safety | 2.3 Disposal of electronic and electrical devices



### Note

KaVo cannot guarantee the compliance of accessories, cables, and other components not supplied by KaVo with the EMC requirements of IEC 60601-1-2 (DIN EN 60601-1-2).

2.3 Disposal of electronic and electrical devices



For final disposal, please contact:

### In Germany

To return an electrical device, you need to proceed as follows:

- 1. On the homepage www.enretec.de of enretec GmbH, you can download a form for a disposal order under the menu item eom. Download the disposal order or complete it as an online order.
- Enter the corresponding information to complete the order, and submit it as an online order or by fax +49 (0) 3304 3919-590 to enretec GmbH. The following contact options are also available for questions and for initiating a disposal order:

Phone: +49 (0) 3304 3919-500 Email: eom@enretec.de and Postal address: enretec GmbH, Geschäftsbereich eomRECYCLING® Kanalstraße 17 D-16727 Velten

A unit that is not permanently installed will be picked up at the office.
 A permanently installed unit will be picked up at the curb at your address on the agreed date.

The owner or user of the device will have to bear the cost of disassembly, transportation and packaging.

### International

For country-specific information on disposal, contact your dental supplier.

### 2.4 Safety instructions

### 

Use of un-authorised accessories or un-authorised modifications of the product. Accessories that have not been approved and/or inadmissible modifications of the product could lead to hazards and/or personal injury or material damage.



- Only use accessories that have been approved for the combination with the product by the manufacturer or are equipped with standardised interfaces (e. g. MUL-TIflex couplings, INTRAmatic).
- Do not make any modifications to the device unless these have been approved by the manufacturer of the product.

2 Safety | 2.4 Safety instructions



### Electrical sparks in the product.

Explosion and/or fire.

- Do not use product in areas subject to an explosion hazard.
- Do not operate the product in an oxygen-enriched atmosphere.



# 

# Damaged mains cable / missing protective conductor.

Electrical shock.

 Check the mains cable before use. The socket outlet must have a protective contact and meet the respective national guidelines.

### Damage by liquids.

Faults on electrical components.

Inadvertent penetration of liquids.

Protect openings of the product from any ingress of liquids.

# 



### Electrical shock.

- Do not place the product in a tub-like container.
- Check the coolant containers and lines for absence of leakage. If any liquid is detected on the device, do not touch the device and disconnect the device from the mains supply without delay. Make sure that the surface of the device is completely dry before plugging the main plug back in the socket.



### 

#### Rotating parts while the pump is operating Injuries

• Do not stick anything in the pump. Turn off the device when the pump is open.

### Risks from electromagnetic fields.

Electromagnetic fields might interfere with the functions of implanted systems (such as pacemakers).

Ask patients if they have a cardiac pacemaker or other system implanted before you start the treatment!



### Impact of power failure.

Failure of the voltage supply or other errors can cause the surgical motor to come to a standstill.

Make sure that the power supply is working.

# 

#### 3 Product description | 3.1 Purpose - Intended use

### **3 Product description**

### 3.1 Purpose – Intended use



#### Note

The EXPERTsurg LUX is approved for use in surgical theatres.

This KaVo product is intended only for use in the field of dentistry, for surgery to expose and dissect oral tissue structures or endodontic treatments (e.g. periodontal gap, gingiva, bone, jaw, extractions, implantations) and must be used by expert medical staff only. Any other type of use is not permitted.

"Proper use" includes compliance with all information in the Instructions for Use and ensuring that all inspections and service tasks are performed.

The overarching guidelines and/or national laws, national regulations and the rules of technology applicable to medical devices for start-up and use of the KaVo product for the intended purpose must be applied and followed.

The user must ensure that the unit works properly and is in satisfactory condition before each use.

The applicable national legal regulations must be observed during the use of the device, in particular the following:

- Applicable regulations governing the connection and start-up of medical devices.
- Current occupational safety regulations.
- Current accident prevention regulations.

Users have a duty to:

- Only use equipment that is operating correctly
- to protect himself, the patient and third parties from danger.
- to avoid contamination from the product

To guarantee the consistent readiness for use and to preserve the value of the KaVo product, the recommended maintenance services must be carried out in 2 year intervals.

The following persons are authorised to repair and service the KaVo product:

- Technicians of KaVo branch offices after appropriate product training.
- Specifically KaVo-trained technicians of KaVo franchised dealers.



### Note

The permitted work is described in the Technician's Instructions available to the trained service staff.

In Germany, operators, equipment managers and users are obliged to operate their equipment in accordance with the MPG regulations.

The services encompass all the test tasks required in accordance with § 6 of the medical devices operator ordinance (Medizinprodukte-Betreiberverordnung, MPBetreibV). After servicing, interventions, and repairs of the device, the device must be tested according to IEC 62353 (according to the state of the art) before re-use.



#### Note

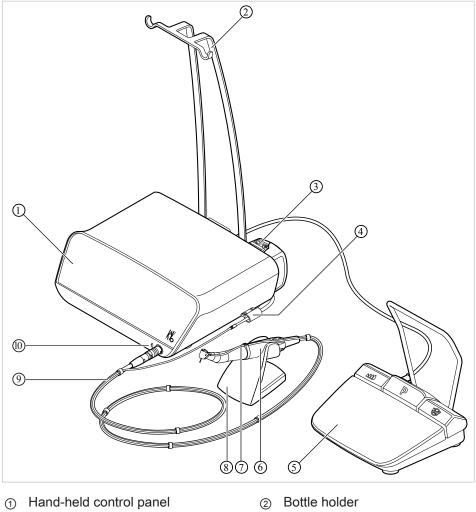
The product must be cleaned and serviced according to instructions if it is not to be used for an extended period of time.



Any waste which is generated must be recycled or disposed of in strict compliance with all applicable national regulations in a manner which is safe both for people and the environment.

If you have any questions regarding proper disposal of the KaVo product, please contact the KaVo branch.

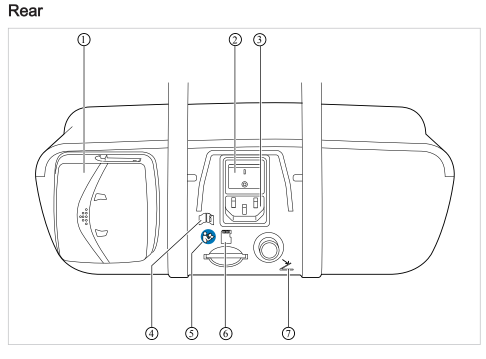
# 3.2 EXPERTsurg LUX



- ③ Hose pump
- ⑤ Foot control
- Coolant hose 7
- Motor cable 9

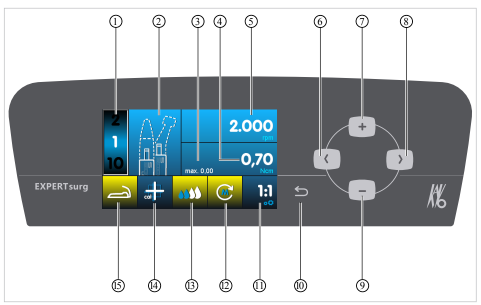
- ④ Hose fixation
- Surgical motor 6
- Handpiece tray 8
- Symbol of type B application part 10

### 3 Product description | 3.2 EXPERTsurg LUX



- Hose pump locking mechanism
- ③ Mains plug
- 5 Follow the instructions for use
- ⑦ Foot control electrical outlet
- On-button
- ④ Please note the instructions for use
- 6 SD card slot

# 3.3 Controls



- ① Program step
- ③ Maximal torque reached
- ⑤ Speed
- ⑦ Plus key, increase value
- Minus key, decrease value
- ① Transmission ratio
- (3) Coolant pump settings
- (5) Foot control status indicator / Service check request

- Display of the activity
- ④ Torque limit
- 6 Left arrow key
- ⑧ Right arrow key
- Back key
- ② Direction of motor rotation
- Activation of one-touch calibration



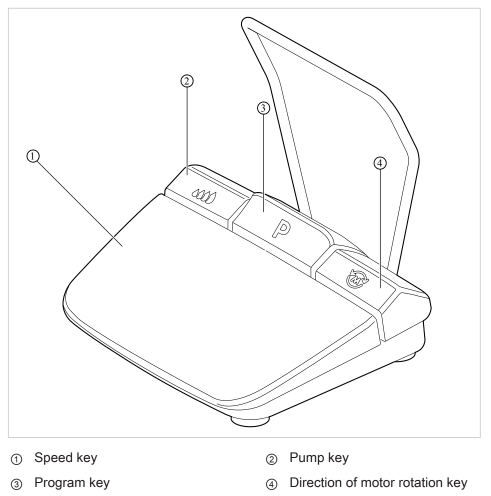
The back key has two functions. Pressing the back key briefly opens the selection of program steps. Pressing the back key long opens the device settings.

Parameters can be selected using the arrow keys.

Use the plus key to increase the value.

Use the minus key to decrease the value.

# 3.4 Foot control



# 3.5 Rating plates of EXPERTsurg LUX and foot control

The rating plates of EXPERTsurg and foot control are affixed on the underside of the housing and include the following symbols:

CE	CE mark
(DE)	VDE mark
	CSA mark
*	Classification, type B
$\land$	Please note the instructions for use
	Please note the electronic instructions for use
	Follow the instructions for use

пг	Operating mode: continuous operation with intermittent load
ЦЦ	
$\sim$	Alternating current (AC)
V	Supply voltage
	Protection class II
	Manufacturer
SN	YYYY = Year manufactured XXXXXXX = Serial number
REF	Material number
Type:	Device type
X	For disposal information, see Intended use
	GOST R certification
	HIBC Code

# 3.6 Technical Specifications of the EXPERTsurg LUX

Width	265 mm
Depth	255 mm
Height	100 mm
Weight	approx. 1.9 kg
Weight of foot control	approx. 1.1 kg
Weight of motor	approx. 125 g
Input voltage	100 - 240 V ~
Input frequency	50/60 Hz
Speed	300 – 40,000 rpm
Max. torque on the motor	5.5 Ncm
Pump delivery rate	30 - 110 ml/min
Foot control: Class of protection	IPX8
Foot control: cable length	2.5 m
Length of motor cable	6.5 ft (2 m)
Operating mode Continuous operation with intermittent load	30 sec. of operation/ 9 min. pause



The maximal motor load is 30 seconds operating time / 9 minutes pause (full load at maximal speed).

### Transportation and storage conditions

Ambient temperature	-20 °C - +50 °C
Relative humidity	5% - 95%
Air pressure	700 hPa - 1,060 hPa

### **Operating environment**



# MARNING Inappropriate operating conditions.

Impairment of the electrical safety of the device.

It is essential to comply with the operating conditions specified in the "Technical Specifications" chapter.

Ambient temperature	+10 °C - +35 °C
Relative humidity	15% - 80%
Air pressure	700 hPa - 1,060 hPa
Max. elevation for operation	up to 3,000 m

# 3.7 Scope of delivery

The scope of delivery of the EXPERTsurg LUX includes the following:

- EXPERTsurg LUX unit
- Foot control
- Surgical motor INTRA LUX S600 LED
- Motor cable S600
- Handpiece tray
- Hose set sterile S600 (5 units)

# 4.1 Unpacking



### Note

You need to keep the cardboard box and all packaging materials to be able to safely ship the unit in the future.

- Open the cardboard box.
- Remove the hose boxes.
- Take out the foot control and additional equipment.
- ► To take out the unit, pull it vertically upward and place it on a level surface.

# 4.2 Installing the bottle holder

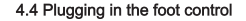


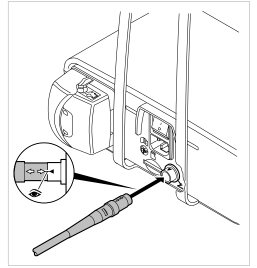
- ► Slide the bottle holder ① in the guide on the underside of the unit.
- $\Rightarrow$  The bottle holder ① snaps into place audibly and is then affixed.

# 4.3 Dismantling the bottle holder

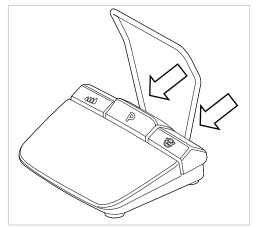


Press down the click-stop knob on the bottom side of the unit and pull off the bottle holder ① towards the back.





Insert the plug of the foot control in the electrical outlet for the foot control. Make sure that the marker arrows on the plug and the socket are aligned towards each other.



• Slide the bracket into the designated recesses.

# 4.5 Connecting the surgical motor

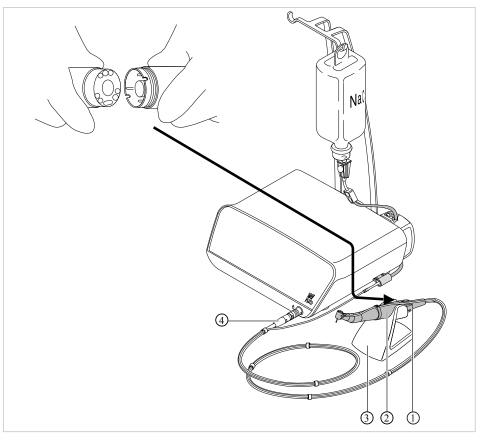


### Note

The delivered parts are not sterile (except for the coolant hose). Before the first treatment of a patient, the surgical motor, motor cable, and the handpiece tray need to be reprocessed.

#### See also:





- ① Motor coupling ② Surgical motor
- Instrument trayPlug of motor cable
- Plug the surgical motor ② into the motor coupling ① and secure it with a union nut. Please note the separate instructions for use of the motor.
- ▶ Place the surgical motor on the handpiece tray ③.
- Insert the plug of the motor cable ④ into the connector on the device, align the marker points, and insert the plug until it snaps into place.

### 4.6 Connecting the coolant container and hose set



- Running, open hose pump. Risk of injury.
- Turn off the device before opening the hose pump.

### 



Danger of tipping due to the coolant containers being too heavy. Malfunctions.

- Use coolant containers with a maximal volume of 1 litre only.
- Check the stability.



### Note

The coolant must be selected to suit the planned application.

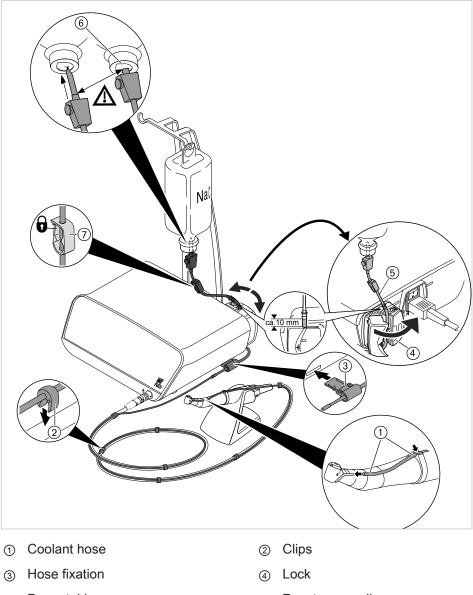


The hose set sterile S 600 (10 pcs.) (Mat. no. 1.009.8757) must be changed after each application.



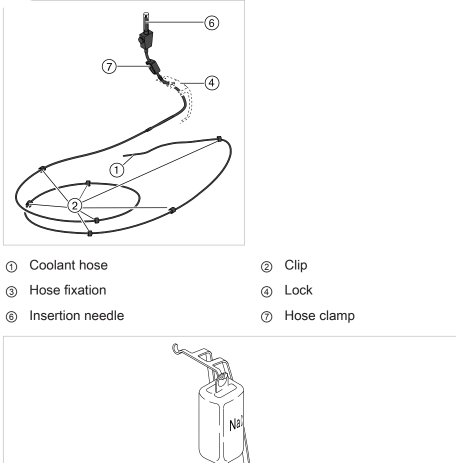
### Note

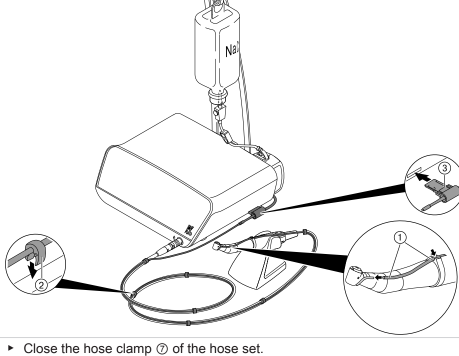
Check the integrity of the hose set before use. If product or packaging are damaged, the product needs to be discarded.



- ⑤ Pump tubing
- ⑦ Hose clamp

6 Puncture needle

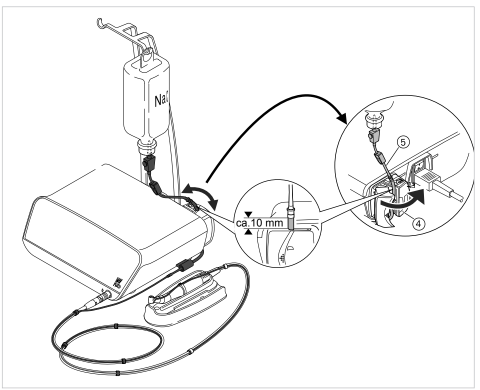




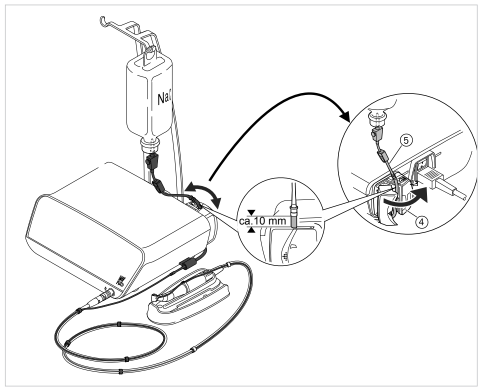
- Attach the coolant hose ① to the straight or contra-angle handpiece.
- Place the coolant hose ① tightly, without loops or kinks, against the outside of the motor cable and attach it in regular intervals using the enclosed clips ②.
- ▶ Plug the hose fixation ③ into the device until the hose fixation ③ snaps in.



The unit recognises the hose adaptor. If the unit fails to recognise the hose adaptor or if it is not plugged in, the coolant supply symbol is shown with a yellow back-ground.



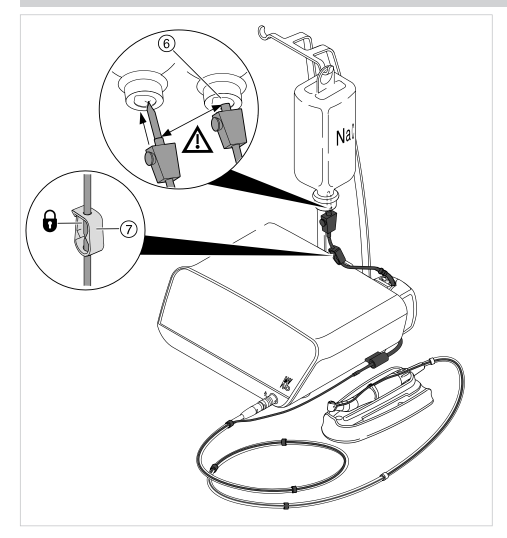
• Open the lock ④ and insert the pump hose ⑤.

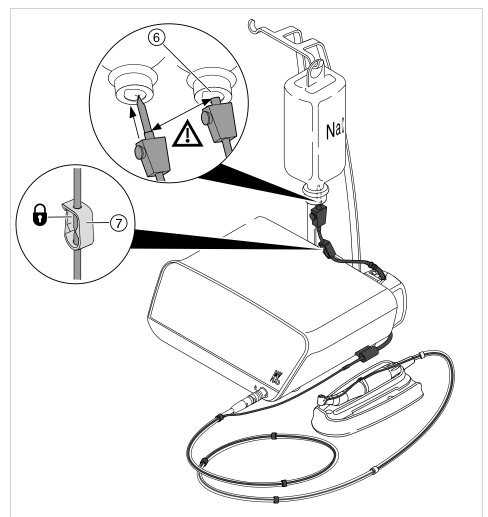


► Close the lock ④.



Make sure to place the pump hose in the pump appropriately such that the pump hose does not get clamped or pinched by the lock. Route all hoses relaxed and without tension.





 Stick the puncture needle into the coolant container and hook-in the coolant container on the bottle holder.

- Check the sealing and firm seating of the puncture needle 6. Prevent fluid from leaking above the device.
- ► If you use a glass bottle, open the ventilation on the puncture needle .
- ► If you use a bag, keep the ventilation on the puncture needle ⑥ closed.
- Open the hose clamp ⑦ before start-up.



### Note

Using a new hose, it may take up to approx. 10 seconds for the coolant to exit on the handpiece, depending on the feed rate.

# 4.7 Electrical connection



# ▲ CAUTION Damaged mains cable / missing protective conductor.

Electrical shock.

Check the mains cable before use. The socket outlet must have a protective contact and meet the respective national guidelines.

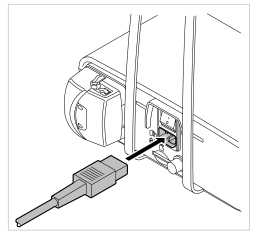


The unit must be set-up appropriately such that the mains plug and the electrical outlet are easily accessible.



### Note

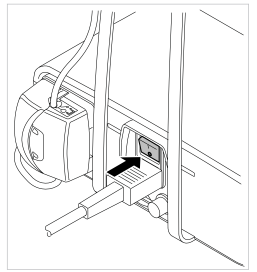
The protective earth conductor is used as functional earthing (FE) rather than as protective earthing (PE).



 Plug the mains cable first into the mains plug on the device and then the other end of the mains cable into the electrical outlet of the supply mains.

# 5 Operation

# 5.1 Switching the device on



- Turn the device on.
- $\Rightarrow$  The device runs a self test.

### Note

Unless the unit is monitored, KaVo recommends turning it off for safety and energysaving reasons.

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### Note

### Auto-Off function

After 10 minutes of inactivity, the light on the handpiece, the pump and the motor on the unit are turned off.

### 5.2 Device settings

The following device settings can be made or displayed:

- Setting the language
- Setting the LUX brightness
- Setting the LUX afterglow time
- Setting the operating mode of the foot pedal
- Adjusting the time of day
- · Setting the date
- Setting the LCD brightness
- Setting the volume
- Setting the key sound volume
- Exporting settings
- Importing settings
- Factory settings
- Version



- Press the back key for a long time, until the device settings are shown.
- Press the plus or minus keys to select the desired setting.
- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change selected and activated settings.
- Press the left arrow key to return to the selection of device settings.
- $\Rightarrow$  Changed values are stored automatically.
- Press the back key for a long time in order to exit from the device settings.

# 5.2.1 Setting the language



- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

### 5.2.2 Setting the LUX brightness

The LUX brightness determines the brightness of the LEDs on the handpiece. The brightness can be set in 4 steps ranging from off to maximal brightness.



	>/
+	-
<	5

- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

### 5.2.3 Setting the LUX afterglow time

The LUX afterglow time determines how long the handpiece LEDs keeps glowing after each motor stop. The afterglow time can be set from 0 to 10 seconds.

+	+
LUX delay	<b>&gt;</b> - <mark>, - , - , - , - , 3</mark> ₅
_	_

- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

### 5.2.4 Setting the operating mode of the foot control

The motor can be triggered in either of two operating modes:

+	+
Foot pedal 🖌	
_	_



The motor starts at full speed when the foot pedal is actuated.



The motor speed can be adjusted continuously up to the maximal level using the foot pedal.

- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

# 5.2.5 Setting the clock time

The time can be set in hours, minutes and seconds:

+	+	+	
Time	11	15	21
	I	-	

- Press the left arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

# 5.2.6 Setting the date

The date can be set in terms of day, month and year:

+	+	+	+
Date 🕻	27	03	2014
_	-	-	1

<	
+	-
<	5

- Press the left arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

# 5.2.7 Setting the LCD brightness

The LCD brightness determines the brightness of the display. The brightness can be set in 3 steps ranging from dark to maximal brightness.

+	+
LCD brightness	> -☆-
_	_

- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

### 5.2.8 Setting the volume

The volume level determines the volume of signal sounds. The volume can be set in 3 steps ranging from quiet to maximal volume. The volume cannot be turned off for safe-ty reasons.



Press the right arrow key to activate the selected setting.



>

- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.
- ⇒ Changed values are stored automatically.

# 5.2.9 Setting the key sound volume

The key sound determines the volume of the sound indicating that a key is pressed. The volume can be set in three steps or switched off.



- Press the right arrow key to activate the selected setting.
- Press the plus and minus keys concurrently to change the selected setting.
- Press the left arrow key or back key to return to the selection of device settings.

⇒ Changed values are stored automatically.

### 5.2.10 Exporting settings

The export of device settings allows the user to export all device settings to the SD card in a <SURG\_EX.SET> file. KaVo recommends exporting the settings as soon as the individual basic settings have been made.

+	
Export setup	Write data
_	

# 5.2.11 Importing settings

The import of device settings allows the user to import all device settings from the SD card in a <SURG\_EX.SET> file. This allows a defined status to be restored, e.g. if settings were changed or deleted inadvertently.

5 Operation | 5.2 Device settings



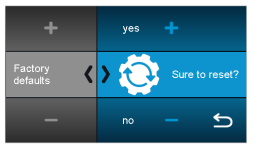
# 5.2.12 Factory settings

Factory settings can be used to re-set the unit to its condition at the time of delivery. All program steps and device settings are re-set to their default values.



Press the right arrow key to activate the selected setting.

In the window, you will be asked whether or not you wish to carry out the action.



- Press the plus key to carry out the action.
- Press the minus key or back key to discontinue the action.

# 5.2.13 Version

+	Software Version: 01.10
Version	

Display of software version

• Press the back key for a long time in order to exit from the device settings.

+

# 5.3 Surgical Motor INTRA LUX S600 LED

### Note

Following instructions for use, service instructions and installation instructions in the motor, handpiece and contra-angle handpiece packaging.

#### See also:

Instructions for use INTRA LUX S600 LED

### 5.3.1 Attaching the straight or contra-angle handpiece



Damage from changing the straight and contra-angle handpieces during operation. Wear to the catch on the straight and contra-angle handpiece and motor.

Unbalanced motor axis.

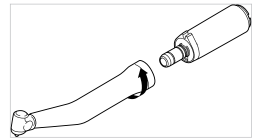
 Change the straight and contra-angle handpieces only when the motor is not running.



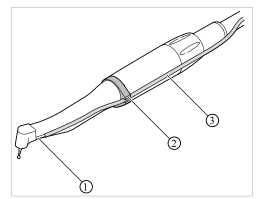
### Note

Following instructions for use, service instructions and installation instructions in the motor, handpiece and contra-angle handpiece packaging.

All straight and contra-angle handpieces with INTRAmatic connection ISO 3964 can be attached.



- Place the KaVo handpiece on the motor, lightly press it against the motor while turning it slightly in the direction of the arrow until the guide stud can be heard to lock into place.
- ▶ Pull on the KaVo handpiece to make sure that it is securely attached to the motor.



Route the coolant hose ③ from the unit along the motor cable (clips) and connect it to the straight or contra-angle handpiece ①. Place the coolant hose ③ into the holding ring ② for this purpose.



# 5.3.2 Removing the straight or contra-angle handpiece

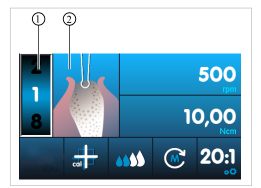


**Damage from changing the straight and contra-angle handpieces during operation.** Wear to the catch on the straight and contra-angle handpiece and motor. Unbalanced motor axis.

- Change the straight and contra-angle handpieces only when the motor is not running.
- ▶ Pull the coolant hose off the straight or contra-angle handpiece.
- Twist the straight or contra-angle handpiece slightly to pull it off.

### 5.4 Setting and executing program steps

The EXPERTsurg LUX is based on program steps and associated activities and can be operated intuitively using the graphical user interface.



### Program step 1: Marking

The current program step is shown on the display as number ① and the corresponding activity is shown as symbol ②. Each program step can be assigned to any activity by selecting the corresponding symbol.

Visualising the activity is an easy means for checking if the activity set on the device is the same as the current treatment step. Maloperation can thus be largely prevented. Default values have been set at the factory for the parameters, speeds, torques, transmission ratios and coolant flow rate for every activity according to application. The parameters can be changed only within a reasonable range for the specific activity. In the activity, "Free use", all available values can be set. The table below lists the ranges of values and factory settings.

A treatment sequence can consist of 4 to 10 program steps and/or activities. The treatment sequence can be designed individually through any arrangement of the activities. Navigate with the foot control during the sequence such that the device does not have to be touched again during the intervention.

Changed values are saved automatically and are then available for the next use.

### 5.4.1 Factory settings

The following program steps are pre-set at the factory:

Program step	Symbol	Activity	Speed [rpm]	Torque [Ncm]	Transmission ratio	Coolant flow
1		Marking	200 – 2,000 500 (D)	5 – 20 10 (D)	16:1 27:1 20:1 (D)	0 – 4 2 (D)
2		Pilot drilling	200 – 2,000 500 (D)	5 – 20 10 (D)	16:1 27:1 20:1 (D)	0 – 4 2 (D)
3		Template drill- ing	200 – 2,000 500 (D)	5 – 20 10 (D)	16:1 27:1 20:1 (D)	0 – 4 2 (D)
4		Thread cutting	15 – 50 20 (D)	5 – 80 25 (D)	16:1 27:1 20:1 (D)	0 – 4 2 (D)
5		Placing im- plant	15 – 50 20 (D)	5 – 80 25 (D)	16:1 27:1 20:1 (D)	0 – 4 0 (D)
6		Setting closure cap	15 – 50 20 (D)	5 – 15 8 (D)	16:1 27:1 20:1 (D)	0 – 4 0 (D)
7	A/7	Free use	300 - 40,000 40,000 (D) 20 - 2,500 15 - 2,000	0.15 - 5.5 3 (D) 5 - 80 5 - 80	1:1 16:1 20:1	0 – 4 2 (D)
			15 – 1,200	5 – 80	27:1	-
8		Rinsing func- tion	-	-	-	-

Program step	Symbol	Activity	Speed [rpm]	Torque [Ncm]	Transmission ratio	Coolant flow
9		Treatment completed (can be set from program step 4)	-	-	-	-

(D) = factory setting (default setup)

### Note

The listed indications are only examples. In order to prevent risks, it is essential to comply with the manufacturer recommendations concerning implants, handpieces, and tools.

# 5.4.2 Examples of program set sequences

Step	1	2	3	4	5	6	7	8
Activity	Marking	Pilot drilling		Thread cut- ting	-	Setting clo- sure cap		Treatment completed (can be set from pro- gram step 4)
Symbol							<u>∧</u> /7 ⊢1	

### Example 1: Factory setting

Example 2: Program steps that do not include the activity, "Cut thread", and include the "Rinsing function" activity

Step	1	2	3	5	4	5	7
Activity	Marking	Pilot drilling		Rinsing func- tion	Placing im- plant		Treatment completed (can be set from pro- gram step 4)
Symbol							

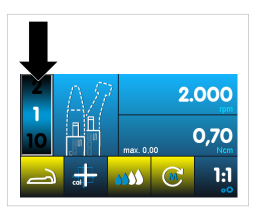
Step	1	2	3	4	5
Activity	Free use	Marking	Pilot drilling		Treatment com- pleted (can be set from program step 4)
Symbol					

# Example 3: Activity "Free use" as step 1, screw implant in manually

# 5.4.3 Selecting the program steps

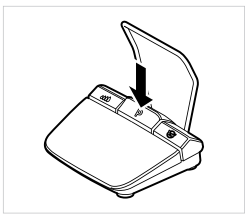


Press the arrow keys until the program step display is highlighted.



+

- Press the plus and minus keys to select the desired program step.
- $\Rightarrow$  The program step is saved automatically.



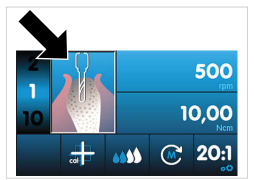
The program steps can be selected during the treatment using the program key of the foot control. After the last program step follows the first step again. Press the program key long to select the previous program step.

### See also:

5.8 Foot control , Page 48

### 5.4.4 Selecting activities

• Press the arrow keys until the activities display is highlighted.



Press the plus or minus keys to select the desired activity.

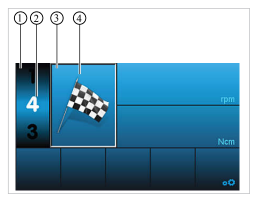


 $\Rightarrow$  The activity is saved automatically.



### 5.4.5 Limiting the program steps

The number of program steps can be limited. Program steps that are not needed are deleted from the display.





- ► Press the arrow keys until the program step display ① is highlighted.
- Press the plus and minus keys to select the program step ② that terminates the treatment (can be set from program step 4).
- ▶ Press the arrow keys until the activities display ③ is shown.



- ▶ Press the plus and minus keys until the flag symbol ④ is displayed.
- ⇒ The settings are saved automatically and the treatment is terminated with the selected program step.
- ⇒ Navigating through the program steps with the wireless foot control, the step with the flag symbol is skipped.

#### Undo the limitation of program steps

• Select the program step with the flag symbol and assign different activity.

## 5.5 Changing default values

The default values set at the factory can be changed within a given range. If the selected activity is Free use, the values can be set freely.

The following values can be changed:

- Maximal speed
- Torque limit
- Coolant flow
- Direction of motor rotation
- Transmission ratio
- Press the arrow keys to select the desired value.
- ⇒ The selected value is shown highlighted.
- Press the plus and minus keys to set the desired value. The step width depends on the transmission ratio and the range of values.
- $\Rightarrow$  The value is saved automatically.

#### 5.5.1 Setting the maximum speed

Press the arrow keys until the speed display is highlighted.



- Press the plus and minus keys to set the maximal speed. The step width depends on the corresponding transmission ratio setting of the surgical motor.
- $\Rightarrow$  The value is saved automatically.

## 5.5.2 Setting the torque limit



#### Note

The EXPERTsurg LUX reduces the power to prevent the maximal torque setting from being exceeded. This may lead to the motor coming to a standstill if the rotating handpiece is blocked.

▶ Press the arrow keys until the torque display is highlighted.





#### Note

The torque values can deviate by max.  $\pm$  10 % with the KaVo contra-angle handpieces SURGmatic S201. Larger deviations are possible with other contra-angle handpieces.

- Press the plus and minus keys to set the desired value. The step width depends on the transmission ratio and the range of values.
- $\Rightarrow$  The value is saved automatically.

The maximal torque value reached is displayed during the treatment in the activities "Cut thread", "Place implant" and "Set closure cap". The value is re-set as soon as the motor starts again.



## 5.5.3 Setting the coolant flow



Coolant dosed incorrectly.

Tissue damage.

- Please note the instructions for use of the attachment tool.
- Set the coolant flow sufficiently high.

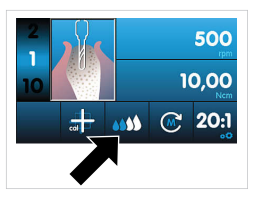
The coolant flow rate can be set to 4 levels or switched off:

- Off
- Level 1 = 32 ml/min
- Level 2 = 50 ml/min

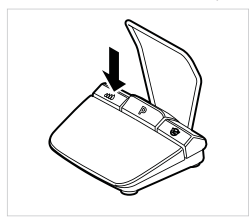
- Level 3 = 76 ml/min
- Level 4 = 110 ml/min



Press the arrow keys until the coolant display is highlighted.



- Press the plus and minus keys to set the desired value. The step width depends on the transmission ratio and the range of values.
- $\Rightarrow$  The value is saved automatically.



The coolant flow can be set during the treatment using the pump key of the foot control. The changed value is shown on the display and is then available for the next use.

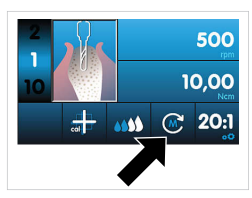
#### See also:

5.8 Foot control , Page 48

#### 5.5.4 Changing the direction of motor rotation

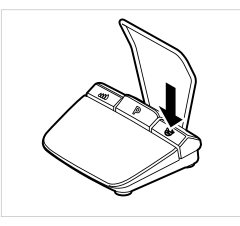


 Press the arrow keys until the display of the direction of motor rotation is highlighted.



+

 Press the plus and minus keys concurrently to change the direction of motor rotation.



The direction of motor rotation can be changed during the treatment using the direction of motor rotation key of the foot control. The changed direction of motor rotation is shown on the display. For safety reasons, running in counterclockwise direction is not saved.

#### See also:

5.8 Foot control , Page 48



Three audible signals indicate counterclockwise rotation. When the motor starts up, 3 more audible signals are issued.

#### 5.5.5 Setting the transmission ratio



Press the arrow keys until the transmission ratio display is highlighted.



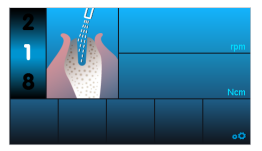
- +
- Press the plus and minus keys to set the desired value. The step width depends on the transmission ratio and the range of values.
- $\Rightarrow$  The value is saved automatically.

#### See also:

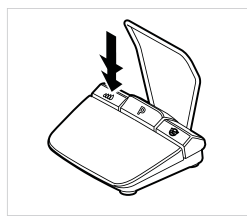
5.4.1 Factory settings, Page 37

## 5.6 Rinsing function

## 5.6.1 Manual rinsing function

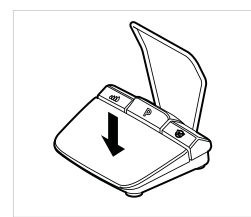


The rinsing function serves to feed liquid and to start-up the illumination on the handpiece. The motor is not activated during this process. The rinsing function can be called up manually at any time.

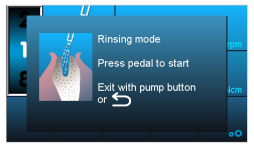


Press the pump key on the foot control for an extended time.

5 Operation | 5.7 Activating the one-touch calibration



Press the speed key on the foot control to start the rinsing function and control the coolant flow.



► To terminate the rinsing function, press the back key or the pump key.

## 5.6.2 Program step Rinsing function

The rinsing function serves to feed liquid and to start-up the illumination on the handpiece. The motor is not activated during this process. The rinsing function can be defined as an activity in the course of the program.

Defining and executing program steps.

#### See also:

Examples of program step sequences (example 2)

#### 5.7 Activating the one-touch calibration

The one-touch calibration automatically compensates for torque deviations of the motor that may be caused, e.g., by aging processes. When the handpiece is attached, the unit detects if the handpiece runs sluggish or is defective. The one-touch calibration thus provides for a more accurate torque on the contra-angle handpiece.

#### Note

The handpiece must be attached for calibration.

One-touch calibration should be carried out only with KaVo surgical handpieces with a transmission ratio of 16:1, 20:1 or 27:1.

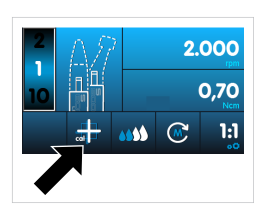
The one-touch calibration cannot be carried out with third-party handpieces or handpieces with different transmission ratios.

The calibration must be repeated whenever the handpiece is changed.



Press the arrow keys until the one-touch calibration is highlighted.

5 Operation | 5.8 Foot control



- Press the plus or minus keys to start the one-touch calibration.
- ⇒ The display shows "Press foot control".



#### The motor starts at full speed. Risk of injury.

- Hold the motor firmly or put it in a safe holder during the calibration.
- Press the foot control and hold it down until the display shows that the calibration has been successful by displaying the message, "Measurement done".

- If you release the foot control before the display shows that the calibration was successful, press the foot control again until the display shows that the calibration was successful.
- Press the back key to terminate the calibration and to return to the selection of device settings.

If an unsuitable or defective handpiece was used in the calibration, the calibration is discontinued and the error message, "Measurement failed - Non-permissible current", is shown.

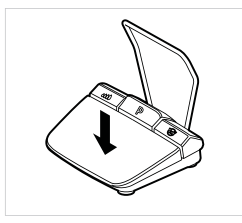
Press the back key to terminate the failed calibration.

#### See also:

8 Troubleshooting, Page 58

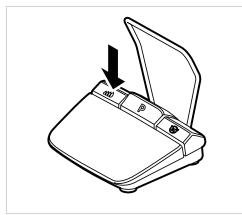
#### 5.8 Foot control

#### 5.8.1 Changing the speed, coolant flow, and direction of motor rotation

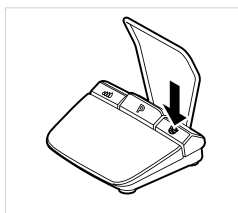


5 Operation | 5.8 Foot control

• Press the speed key with your foot to start the motor and increase the speed.



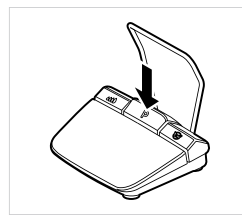
- Press the pump key of the foot control to set the coolant flow.
- Press the pump key of the foot control for an extended time to activate the rinsing function.



Press the direction of motor rotation key of the foot control to set the direction of motor rotation.

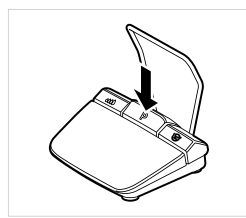
## 5.8.2 Selecting the program steps

Program steps can be selected during the treatment using the program key of the foot control.



• Press the program key of the foot control **briefly** to select the next program step.

5 Operation | 5.9 Changing the coolant container

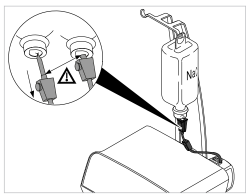


Press the program key of the foot control long to select the previous program step.

## 5.9 Changing the coolant container

The coolant container can be changed as follows:

Close the hose clamp.



- ► Pull the hose and puncture needle out of the empty coolant container.
- Replace the empty coolant container by a full coolant container.

#### See also:

Connecting the coolant container and hose set

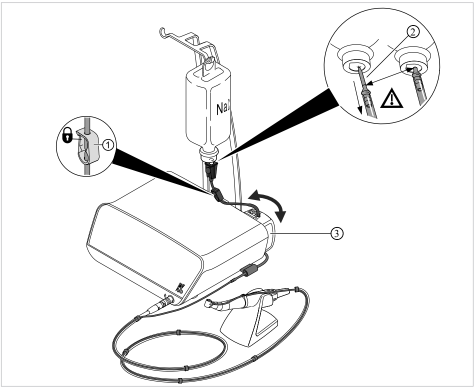
6 Disposal | 5.9 Changing the coolant container

## 6 Disposal

Note



The coolant hose with accessories needs to be exchanged and discarded after each treatment.



- Close the hose clamp ①.
- ▶ Pull the puncture needle ② out of the coolant container.
- Open the lock ③ and remove the hose.
- Remove the hose set from the unit and discard it.

#### 7 Reprocessing steps in accordance with DIN EN ISO 17664



#### Note

The reprocessing steps for surgical motors with motor cable and for the straight and contra-angle handpieces are described in the corresponding Instructions for use.

# 1

#### Note

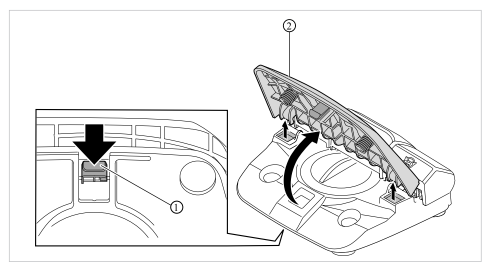
The instructions for cleaning and sterilisation have been validated by the manufacturer. Any departure from the instructions provided must be checked by the user for efficacy and possible detrimental consequences.

## 7.1 Cleaning

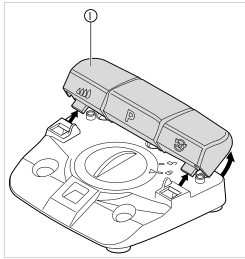
#### 7.1.1 Manual cleaning

Use a moist disposable cloth to wipe down all visible surfaces of the unit, bottle holder, foot control surfaces, and connecting cables.

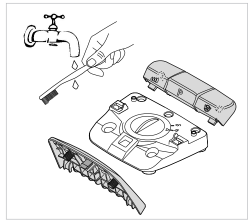
## Cleaning the foot control



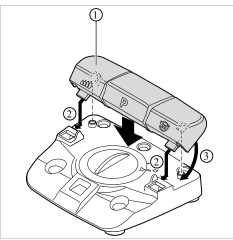
On the underside of the foot control, press down the snap-in lug ① of the speed button ② and take the speed button ③ off the foot control.



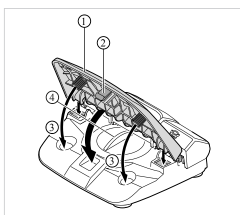
 Pull the button bar ① including the pump button, program button, and motor direction button slightly upwards and take it off the foot control. 7 Reprocessing steps in accordance with DIN EN ISO 17664 | 7.1 Cleaning



 Clean the individual parts of the foot control under running water and then dry them.



Plug the button bar ① onto the foot control ② and press it on lightly until the button bar ① snaps into place ③.



Plug the speed button ① onto the foot control and press it on lightly until the snap-in lug ② snaps in. Make sure that the pedal springs are situated in the recesses of the housing ③.



## Note

The lid needs to stay closed while the foot control is being cleaned.

## 7.1.2 Machine cleaning



#### Damage by liquids.

Faults on electrical components.

Do not machine-clean the device.

The following parts of the unit are released for machine-based cleaning:

- Handpiece tray
- Motor and motor cable



#### Note

Please comply with the corresponding Instructions for use when you re-process the motor and motor cable.

The re-processing steps for the handpiece tray are as follows:

KaVo recommends thermodisinfectors in accordance with EN ISO 15883-1, which are operated with alkaline cleaning agents with a pH value of max. 10 (e.g. Miele G 7781/G 7881 – Validation was carried out with Programme "VARIO-TD", cleaning agent "neodisher® mediclean", neutralisation agent "neodisher® Z" and rinsing agent "neodisher® mielclear" and only applies to the material compatibility with KaVo products).

 For program settings as well as cleansers and disinfectants to be used, please refer to the Instructions for Use of the thermodisinfector (complying with max. pH value of 10).

## 7.2 Disinfection



#### Note

After each treatment of a patient, the surfaces near the patient that may have been contaminated by contact or aerosol need to be disinfected. All disinfection measures need to be carried out by wipe disinfection.



#### Note

Please comply with the corresponding Instructions for use during the re-processing of the motor and motor cable.

#### 7.2.1 Manual disinfection

- Use a soft disposable cloth and an approved disinfectant for disinfection by wiping down all visible surfaces of the unit, bottle holder, foot control surfaces, and connecting cables. Make sure that all surfaces are wetted.
- Let the disinfectant act for the prescribed time.
- Dry the surfaces.

Permissible disinfectants (application range in accordance with the available manufacturer's instructions and national guidelines. Please note material safety data sheets.) KaVo recommends the following products based on the compatibility of the materials. The microbiological efficacy must be ensured by the disinfectant manufacturer.

- FD 322 Dürr
- Microcide AF Liquid (Schülke & Mayr)

7 Reprocessing steps in accordance with DIN EN ISO 17664 | 7.2 Disinfection

CaviCide made by Metrex

## 7.2.2 Automated disinfection

The following parts of the unit are released for machine-based disinfection:

- Handpiece tray
- Motor and motor cable



#### Note

Please comply with the corresponding Instructions for use when you re-process the motor and motor cable.

The re-processing steps for the handpiece tray are as follows:



KaVo recommends thermodisinfectors in accordance with EN ISO 15883-1, which are operated with alkaline cleaning agents with a pH value of max. 10 (e.g. Miele G 7781/G 7881 – Validation was carried out with Programme "VARIO-TD", cleaning agent "neodisher® mediclean", neutralisation agent "neodisher® Z" and rinsing agent "neodisher® mielclear" and only applies to the material compatibility with KaVo products).

 For program settings as well as cleansers and disinfectants to be used, please refer to the Instructions for Use of the thermodisinfector (complying with max. pH value of 10).

## 7.2.3 Drying



#### Note

Coolant hose with accessories is intended for single use only and is not to be disinfected and sterilised. No drying required.

Allow all disinfected and sterilised parts to dry fully exposed to room air before using them again.

## Automatic Drying

The drying procedure is usually a part of the cleaning program of the thermodisinfector.

► Follow the instructions for use of the thermodisinfector.

## 7.2.4 Service, inspection and testing after preparation

#### Note

It is essential to comply with the hygiene requirements (sterility) during the test after reprocessing. If sites of fracture and clear changes of the surface are visible, the parts need to be checked by the Service.

Check for cleanliness, intactness, servicing, and repair as described in the following:

- Check the adjustable functions of the unit and the motor function.
- Check the hose pump for sufficient coolant flow rate.
- Check the control commands on the foot control.

## 7.3 Packaging



#### Note

The quality and use of the sterilisation packaging must comply with applicable standards and be suitable for the sterilisation procedure!

## i

#### Note

If potentially infectious liquids and particles can contact the products, it is recommended to cover and protect these areas with sterile disposable products.

Seal the handpiece tray and motor cable in a sterilisation pouch.

## 7.4 Sterilisation

Sterilisation by moist heat in accordance with ISO 17665-1 in a steam steriliser (autoclave)



Damage to device due to imprope	r sterilisation.

Damage to the sterile device.

No hot air sterilisation, no chemical cold sterilisation, do not sterilise with ethylene oxide!



## 

Product damage Contact corrosion

 Remove the sterilised item from the autoclave immediately after sterilising and drying.

## Note

Treating patients who are possibly afflicted by an acute, critical infectious disease, be sure to comply with the hygienic measures cited in applicable publications and reports. If possible, use suitable disposable devices to avoid the transmission of critical pathogens. This concerns the protection of the user, patients and all participants in the surgery.

All dental and medical materials considered to be contaminated must be suitably processed and sufficiently identified after cleaning and sterilisation before returning them.



#### Note

The user is responsible for observing the regulations and conditions for sterility. The coolant container needs to be disposed and the hoses need to be changed after each patient.



KaVo medical devices released for sterilisation are temperature-resistant up to 138  $^{\circ}$ C (280.4  $^{\circ}$ F).

The following parts are released for sterilisation:

- Motor cable
- Instrument tray
- Autoclave with 3-fold fractionated pre-vacuum:
  - at least 3 minutes at 134 °C -1 °C / +4 °C (273 °F -1.6 °F / +7.4 °F)

7 Reprocessing steps in accordance with DIN EN ISO 17664 | 7.4 Sterilisation

- Drying time: 20 min.
- Autoclave using the gravitation method
  - at least 10 minutes at 134 °C -1 °C / +4 °C (273 °F -1.6 °F / +7.4 °F)
- Drying time: 30 min.



#### Note

Allow the sterilised items to cool to room temperature before using them again.

## 7.4.1 Storage

Observe all necessary measures for hygiene when storing sterile goods. Store protected from dust and in a dry place, release with identification on the packaging. Evaluate the duration of storage.

## 8 Troubleshooting



#### Note

If malfunctions cannot be located or eliminated using this troubleshooting guide, a technician trained by KaVo must be commissioned to eliminate the problem.



## Note

The permitted work is described in the Technician's Instructions available to the trained service staff.

In case of malfunction, the display names the malfunction directly or shows an error number.

Malfunction	Cause	Remedy
Non-functional device.	The unit is switched off.	<ul> <li>Switch-on the mains switch on the rear of the unit.</li> </ul>
	Neither end of the power cable is plugged in.	<ul> <li>Plug in the power input cable.</li> </ul>
	Unknown.	<ul> <li>Turn the unit off and on.</li> </ul>
	Blown fuse.	<ul> <li>Contact customer serv- ice.</li> </ul>
No coolant in the hand- piece.	No coolant flow pre-selec- ted.	<ul> <li>Pre-select coolant flow.</li> </ul>
	Pump is off.	5.5.3 Set the coolant flow, Page 43
	Hose fixation is not plug- ged in.	<ul> <li>Plug in the hose fixation.</li> <li>See also:         <ul> <li>4.6 Connect the coolant container and hose set, Page 22</li> </ul> </li> </ul>
	Bottle is empty.	<ul> <li>Connect new bottle.</li> </ul>
	Hose clamp is closed.	<ul> <li>Open the hose clamp.</li> </ul>
	Pump locking mechanism is not closed.	<ul> <li>Check and close, if nee- ded, the closing mecha- nism.</li> </ul>
	Hose is kinked.	<ul> <li>Check hose and remove the kink, if any.</li> </ul>
	Glass bottle containing the coolant is not ventilated.	<ul> <li>Open the cap on the ventilation valve of the puncture needle.</li> </ul>
Insufficient coolant flow in the instrument.	Spray nozzles are crusty or soiled.	<ul> <li>Clean the spray nozzles with the nozzle needle or re-process the part.</li> </ul>
		See also: <ul> <li>SURGmatic Instructions for use</li> </ul>
	Glass bottle containing the coolant is not ventilated.	<ul> <li>Open the cap on the ventilation valve of the puncture needle.</li> </ul>

8 Troubleshooting | 7.4 Sterilisation

Malfunction	Cause	Remedy	
The motor makes a grind- ing noise or does not run smoothly.	The motor is not correctly plugged on or screwed on.	<ul> <li>Firmly insert the motor hose into the housing.</li> <li>Firmly screw on the mo- tor hose to the motor.</li> <li>Check if all the connec- tions and couplings are firmly seated.</li> </ul>	
No light on the straight or contra-angle handpiece	The light is not turned on.	<ul> <li>Turn on the light.</li> <li>See also:         <ul> <li>5.2.2 Setting the LUX brightness, Page 30</li> </ul> </li> </ul>	
	The straight and contra-an- gle handpiece is improper- ly attached.	<ul> <li>Attach the straight and contra-angle handpiece until the catch audibly locks.</li> </ul>	
	Defective LED.	<ul> <li>Replace the LED.</li> <li>See also:         <ul> <li>Instructions for use INTRA LUX S600 LED</li> </ul> </li> </ul>	
	Not a suitable straight and contra-angle handpiece.	<ul> <li>Use a suitable light, straight and contra-angle handpiece.</li> </ul>	

## Error message from software

Malfunction	Cause	Remedy
Motor symbol has a yellow background.	No motor attached.	<ul> <li>Connect the motor.</li> </ul>
Pump symbol has a yellow background.	Hose fixation is not plug- ged in.	<ul> <li>Plug in the coolant hose.</li> </ul>
Warning E3: Release the foot pedal	Foot control was activated while the unit started up.	<ul> <li>Release the foot control.</li> </ul>
Warning E4: Data initialisation	Setting data newly initial- ised.	<ul> <li>Confirm message and check and correct the program settings if nee- ded.</li> <li>If the problem persists, notify service engineer</li> </ul>
Message E5: Internal communication er- ror	Internal system error.	<ul> <li>Turn the unit off and on. If the problem persists, notify service engineer.</li> </ul>
Message E9: Setting the date and time	Time was not reset after re-start.	<ul> <li>Setting the date and time.</li> <li>If the problem persists, have a service engineer replace the battery of the real-time clock.</li> </ul>

#### 8 Troubleshooting | 7.4 Sterilisation

Malfunction	Cause	Remedy
Foot control symbol has a yellow background	Foot control malfunction.	<ul> <li>Check if the foot control plug is plugged in.</li> </ul>
		<ul> <li>Check if the foot control is installed correctly.</li> </ul>
		<ul> <li>If the problem persists, notify service engineer.</li> </ul>
One-touch calibration failed.	Handpiece is too sluggish.	<ul> <li>Perform a run in the ab- sence of the handpiece.</li> </ul>
		If no error message is displayed, the handpiece runs too sluggish.
		<ul> <li>Use a different hand- piece.</li> </ul>
	If the run in the absence of	<ul> <li>Use a different motor;</li> </ul>
	the handpiece again pro-	possibly return the motor
	duces an error message,	for repair.
	the motor torque is too low.	
Hardware error	Internal system error.	<ul> <li>Turn the unit off and on. If the problem persists, notify service engineer.</li> </ul>
Service symbol is green	Service is due soon.	<ul> <li>Arrange a precautionary appointment at a KaVo subsidiary or with a Ka- Vo authorised dealer.</li> </ul>
Service symbol is yellow	Service period is expired	<ul> <li>Arrange an appointment at a KaVo subsidiary or with a KaVo authorised dealer.</li> </ul>
Service symbol is red	Service overdue: > 4 months	<ul> <li>Arrange an appointment immediately at a KaVo subsidiary or with a Ka- Vo authorised dealer.</li> </ul>
Warning E36 and E37:	Overtemperature on stator.	► Let the motor rest. If the
Motor overload	Over-current > nominal current.	problem persists, notify service engineer.
Error message "SD card defective" during software update	SD card is formatted incor- rectly or SD card is defec- tive.	<ul> <li>Format SD card in FAT16 or FAT32 format or use new CD card. Then repeat the updating process.</li> </ul>

9 Run a software update | 7.4 Sterilisation

#### 9 Run a software update

Please proceed as follows to update the software:

- Download the current firmware file from www.kavo.de/produkte.
- ► Copy the firmware file to an SD card (storage capacity 1 32 GB, FAT format).
- Turn the device off.
- Insert the SD card with the new firmware file into the unit. Make sure that only a single firmware file with the .bin file extension (the downloaded current file) is stored on the SD card.
- ► Turn the device on.
- $\Rightarrow$  The update process starts automatically.



## Note

The unit must not be turned off during the update process.

After the update process, the unit starts using the updated software.



#### Malfunctions of the unit.

The software version displayed on the start screen or in the version display must coincide with the software version that was downloaded. If these do not coincide or if there is any other error, please contact Customer Service.

#### See also:

5.2.13 Version, Page 35



#### Note

All program and device settings remain unchanged.

## 10 Safety checks ("STK")

The EXPERTsurg LUX must be subjected to a service check including safety check ("STK") every 2 years. The safety check may only be done by a professional trained by KaVo or in a shop trained by KaVo. Perform the safety check ("STK") as described in the KaVo technician's instructions.

The urgency of the service check is indicated on the display by a symbol in "traffic light colours".

Symbol	Description	
<u></u>	Service check is soon due.	
	•	Arrange a precautionary appointment at a Ka- Vo subsidiary or with a KaVo-authorised deal-
green		er.
	Service check is due.	
	•	Arrange an appointment at a KaVo subsidiary or with a KaVo-authorised dealer.
yellow		
6	Service check is over-due.	
	•	Arrange an appointment immediately at a Ka- Vo subsidiary or with a KaVo-authorised deal-
red		er.

## 11 Accessories

The following accessories are approved for the EXPERTsurg LUX:

- Hose set sterile S600 (10 pcs.) (Mat. no. 1.009.8757)
- Handpiece tray (Mat. no. 1.009.3411)
- Motor INTRA LUX S600 LED (Mat. no. 1.008.8000)
- Motor cable S600 (Mat. no. 1.009.1700)
- KaVo SURGmatic straight and contra-angle handpieces

## 12 Details on electromagnetic compatibility

## 12.1 Guidelines and manufacturer's declaration - electromagnetic transmission

The EXPERTsurg LUX is designed for operation in an environment like the one described below. The EXPERTsurg LUX customer or user must ensure that the unit is used in an environment matching the description.

Measurements of emitted interfer- ence	Conformance	Electromagnetic environment - Guidelines
HF emissions according to CISPR 11	Group 1	The EXPERTsurg LUX uses HF en- ergy for its internal functions exclu- sively. Therefore, the HF emission of the device is very low and inter- ference with adjacent electronic de- vices is unlikely.
HF emissions according to CISPR 11	Class B	The EXPERTsurg LUX is designed for use in all facilities including resi- dential ones, and facilities that are directly connected to a public power supply that also supplies residential buildings.
Emission of harmonics according to IEC 61000-3-2	Class A	The EXPERTsurg LUX is designed for use in all facilities including resi- dential ones, and facilities that are directly connected to a public power supply that also supplies residential buildings.
Emission of voltage fluctuations/ flicker according to IEC 61000-3-3	complies	The EXPERTsurg LUX is designed for use in all facilities including resi- dential ones, and facilities that are directly connected to a public power supply that also supplies residential buildings.



## Note

The device or system may not be used or stacked directly next to other devices. If it has to be used close to or stacked next to other devices, the device or system must be monitored to ensure that it is used properly in the existing arrangement.



#### Note

The immunity test levels required in IEC 60601-1-2 (DIN EN 60601-1-2) are met.

## 12.2 Guidelines and manufacturer's declaration - electromagnetic resistance to jamming

The EXPERTsurg LUX is designed for operation in an environment like the one described below. The EXPERTsurg LUX customer or user must ensure that the unit is used in an environment matching the description. 12 Details on electromagnetic compatibility | 12.3 Guidelines and manufacturer's declaration - electromagnetic resistance to jamming

Interference immunity tests	IEC 60601 test levels	Compliance level	Electromagnetic environ- ment - Guidelines
Electrostatic discharge (ESD) according to IEC 61000-4-2	± 6 kV contact discharge ± 8 kV atmospheric dis- charge	± 6 kV contact discharge ± 8 kV atmospheric dis- charge	Floors should be made of wood or concrete or be fit- ted with ceramic tiles. If the floor is fitted with synthetic material, the relative hu- midity must be at least 30 %.
Fast transient electrical in- terference / bursts accord- ing to IEC 61000-4-4	± 2 kV for power lines	± 2 kV for power lines	The quality of the supply voltage should correspond to that of a typical business or hospital environment.
Surges according to IEC 61000-4-5	± 1 kV push-pull voltage (symmetrical) ± 2 kV common mode volt- age (unsymmetrical)	± 1 kV push-pull voltage (symmetrical) ± 2 kV common mode volt- age (unsymmetrical)	The quality of the supply voltage should correspond to that of a typical business or hospital environment.
Voltage interruptions, short-term interruptions, and fluctuations of the sup- ply voltage according to IEC 61000-4-11	< 5 % $U_T$ for ½ period (> 95 % interruption) 40 % $U_T$ for 5 periods (60 % interruption) 70 % $U_T$ for 25 periods (30 % interruption) < 5 % $U_T$ for 5 s (> 95 % interruption)	< 5 % $U_T$ for ½ period (> 95 % interruption) 40 % $U_T$ for 5 periods (60 % interruption) 70 % $U_T$ for 25 periods (30 % interruption) < 5 % $U_T$ for 5 s (> 95 % interruption)	The quality of the supply voltage should correspond to that of a typical business or hospital environment. If the user of the EXPERT- surg LUX needs uninter- rupted function of the unit even when the power sup- ply is interrupted, it is rec- ommended to supply the EXPERTsurg LUX from an uninterruptible power sys- tem or a battery.
Magnetic field at a supply frequency (50/60 Hz) ac- cording to IEC 61000-4-8	3 A/m	3 A/m	Magnetic fields at the mains frequency should correspond to typical val- ues in a business and hospital environment.

Note: U  $_{\scriptscriptstyle T}$  is the alternating mains voltage before the application of the test level.

## 12.3 Guidelines and manufacturer's declaration - electromagnetic resistance to jamming

The EXPERTsurg LUX is designed for operation in an environment like the one described below. The EXPERTsurg LUX customer or user should ensure that the unit is used in an environment matching the description. 12 Details on electromagnetic compatibility | 12.4 Recommended safe distance between portable and mobile HF telecommunications equipment and the EXPERTsurg LUX

Interference immunity tests	IEC 60601 test levels	Compliance level	Electromagnetic environment - Guidelines
Wire-based HF interfer- ence according to IEC 61000-4-6 Wireless HF interfer- ence according to IEC 61000-4-3	3 V <sub>eff</sub> 150 kHz to 80 MHz 3 V/m 80 MHz to 2.5 GHz	3 V <sub>eff</sub> 3 V/m	Portable and mobile radio devices should not be used closer to the EXPERTsurg LUX, including the wires, than the recommended safe distance calculated using the equa- tion for the transmission frequency. Recommended safe distance: $d = [3.5/3]\sqrt{P} = 1.17\sqrt{P}$ $d = [3.5/3]\sqrt{P} = 1.17\sqrt{P}$ for 80 MHz to 800 MHz $d = [7.0/3]\sqrt{P} = 2.33\sqrt{P}$ for 800 MHz to 2.5 GHz where P is the maximal nominal power of the transmitter in watts (W) as specified by the transmitter manufacturer and d is the recom- mended safe clearance in metres (m). The field strength of stationary wire- less radio transmitters as measured locally <sup>a</sup> should be lower than the conformance level at all frequen- cies. <sup>b</sup> Interference is possible in the vicini- ty of devices that bear the following symbol. <sup>((w))</sup>

Note 1: At 80 MHz and 800 MHz, the higher frequency range applies.

Note 2: These guidelines may not be applicable in every case. The propagation of electromagnetic waves is subject to absorption and reflection by buildings, objects, and people.

<sup>a</sup>The field strength of stationary transmitters such as base stations of mobile telephones and land mobile radio devices, amateur radio stations, AM and FM radio and television broadcasting stations cannot be determined based on theoretical considerations. A site study should be considered to determine the electromagnetic environment in terms of stationary transmitters. If the field strength measured at the site, at which the EXPERTsurg LUX is used, exceeds the compliance levels shown above, the EX-PERTsurg LUX should be monitored to demonstrate proper function. Should unusual performance features be observed, additional measures may be required, such as, e.g., a different alignment or different location for the EXPERTsurg LUX.

 $^{\rm b}$  In the frequency range of 150 kHz to 80 MHz, the field strength should be less than  $3V_{\rm eff}$  V/m.

## 12.4 Recommended safe distance between portable and mobile HF telecommunications equipment and the EXPERTsurg LUX

The EXPERTsurg LUX is designed for operation in an electromagnetic environment like the one described below. The customer or user of the EXPERTsurg LUX can help prevent electromagnetic interference by keeping the minimum safe distance between portable and mobile HF telecommunication devices (transmitters) and the EXPERT-surg LUX – depending on the output cable of the communication device - as given below.

12 Details on electromagnetic compatibility | 12.4 Recommended safe distance between portable and mobile HF telecommunications equipment and the EXPERTsurg LUX

Rated power of the trans- mitter in W			800 MHz to 2.5 GHz d=2.33 $\sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.37	0.37	0.74
1	1.17	1.17	2.33
10	3.70	3.70	7.37
100	11.70	11.70	23.30

For transmitters whose maximum rated power is not included in the above table, the recommended safe distance d in metres (m) can be calculated using the equation for the respective column, where P is the maximum rated power of the transmitter in Watts (W) as specified by the manufacturer.

Note 2: These guidelines may not be applicable in every case. The spread of electromagnetic waves is absorbed and reflected by buildings, objects and people. Comment 1: To calculate the recommended safe distance from transmitters with a frequency range of 80 MHz to 2.5 GHz, an additional factor of 10/3 was used to reduce the probability that a mobile/portable communication unit that is inadvertently brought into the patient area would cause malfunction.

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