# MEDCAPTAIN

### NAVI-30/NAVI-60

Vein Illuminator

### **Operation Manual**

Before using the NAVI-30/NAVI-60 vein illuminator, please read this Manual carefully and follow the safety precautions and operating instructions contained herein.

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MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.

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MEDCAPTAIN is responsible for the safety, reliability and performance of this product, only if:

- The product is used in accordance with this Manual.
- All installations, replacements, tests, modifications and repairs are conducted by technicians authorized to do so by MEDCAPTAIN.
- All replacement components and accessories are provided by MEDCAPTAIN.
- All maintenance service records are kept.

#### Illustrations

• All the illustrations provided in this operation manual are for your reference only. The settings or data on the illustrations may differ from the actual settings or data of the product.

### **Version Information**

#### V1.0

- Version of operation manual: First release
- Software release version: V1
- Issued on: August 2018

#### V1.1

- Updated some parameters of the built-in battery.
- Issued on: April 2019

### **After-Sales Service**

Thanks for purchasing our vein illuminator.

- MEDCAPTAIN provides limited warranty for the product. That is, we provide free after-sales services for the product within the warranty period. The specific warranty period is stipulated on the sales contract. For details, please contact your local distributor. However, a product damage or fault is not covered by the warranty if it is caused by:
  - Operator error;
  - Improper use;
  - Out-of-range grid voltage;
  - Force majeure such as natural disasters;
  - Replacement with or use of any component, accessory or consumable other than authorized by MEDCAPTAIN; or
  - Other damages/faults not caused by the product.
- After the warranty period expires, MEDCAPTAIN shall continue to provide paid maintenance service within the service life of the product.
- Feel free to contact us or your local distributor if you have any problem in using the product.
- Contact our after-sales service department:

#### After-sales service provider: MEDCAPTAIN MEDICAL TECHNOLOGY CO., LTD.

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### Contents

1	Overview	1
2	Safety	3
3	Product Specifications	8
4	Product Description	10
5	Installation Description	15
6	Operating Instructions	18
7	Common Faults	24
8	Cleaning and Disinfection	
9	Maintenance	30
A	ppendix A	35

#### **1** Overview

#### 1.1 Intended Use

The NAVI-30/NAVI-60 vein illuminator is used to observe and locate the superficial veins, and to assist medical personnel in venipuncture.

### 1.2 Contraindications

Direct eye exposure to the light emitted by this product is forbidden.

#### **1.3 Product Features**

As a portable professional medical device for vein imaging, the NAVI-30/NAVI-60 vein illuminator detects veins beneath skin by using infrared light and then indicates the vein positions on the skin surface above the veins by using light during operation. By observing the vascular system projected on the skin surface, qualified medical personnel can find a vein of appropriate size and position for venipuncture.

- Veins beneath skin are detected by using the near-infrared light and then projected on the body surface by using the high-definition vein illuminator, implementing real-time and accurate display of vein positions.
- This device can be placed at any place above the detection area to project the vein positions accurately. It does not need to be fastened at a specific position above the detection area for accurate projection.
- Easy to operate. Users can press to power on/off the device.
- Lightweight structural design, easy to take along.

### Overview

- Handheld device for medical personnel. Ergonomic design, comfortable to hold it.
- Five projection colors are available for selection. The corresponding inverse color mode of each color can also be selected.
- The projection brightness can be adjusted to adapt to different operation environments.
- The size of the projection window can be adjusted to adapt to different patients and different detection positions.
- This device can be used to detect and display the depth of the vein beneath skin.
- The NAVI-60 vein illuminator is equipped with a display screen to display various operation information like the remaining battery capacity, mode, color, and projection size.
- A lithium battery is built in to allow long-time portable use.
- This device can be used while it is connected to an external power supply for battery charging.
- This device can be used together with a support or trolley.

#### 1.4 Model Differences

Product Model	Difference	
NAVI-30	Without a display screen	
NAVI-60	With a display screen	

#### 2.1 Warnings and Cautions

In this Manual, the precautions are classified by importance into warnings and cautions as defined below:

## **WARNING:**

The precautions related to safety and effectiveness. Failure to follow them may cause personal injuries.

# **A**CAUTION:

The precautions related to guidance and suggestions. Failure to follow them may affect the normal use of the product.

Please read all warnings and cautions contained herein carefully.



- The vein illuminator must be operated by professional medical personnel or under the guidance of clinicians. In addition, the operator must receive a training related to the use of this device.
- High-frequency surgical equipment, mobile phones, wireless devices, and defibrillators may cause interference on the vein illuminator. Therefore, keep the vein illuminator away from these devices when using the vein illuminator.
- To avoid the risk of electric shock, connect the vein illuminator to only the power supply system with protective earth. In case that the power supply system does not have protective earth, disconnect the power cable of the vein illuminator from the power supply system and use the built-in battery to power the vein illuminator.

- The vein illuminator does not have a patient connection circuit. Prevent the patient from touching the vein illuminator.
- This vein illuminator can only identify veins, and it cannot effectively evaluate arteries.
- Do not use this product as diagnostic equipment or for any treatment purposes.
- This product is a class I continuous-operation device without waterproof protection design. Do not splash or drip any liquid onto or into the device, and do not immerse any part of the main unit in liquid. If any liquid splashes into the product during charging or operation of the product, immediately power off the product and stop using it.
- Only use the dedicated accessories provided by MEDCAPTAIN. Before use, check the power cable, power adapter, and other accessories. In case of any damage, stop using the product and contact the after-sales service department of MEDCAPTAIN.
- This product has a removable battery inside. Operate, store, and transport this product in strict accordance with the operation manual. Do not use the product in flammable and explosive environment.
- The lithium battery provided by MEDCAPTAIN is the original battery. If this product will not be used for a long time, fully charge it for storage. The service life or shelf life of the battery is one year.
- Do not discard the retired vein illuminator at will. The product contains a battery and therefore it must be disposed of according to the Technical Policy for Discarded Batteries Pollution Prevention in a professional manner.

- Do not disassemble or try to repair the vein illuminator. Otherwise, serious hazards may be incurred. The manufacturer and distributor shall not be responsible for any vein illuminator that has been disassembled, modified or used for any purpose other than its intended purpose.
- Do not apply this product on the skin with scar, tattoo, skin disease, or lots of hairs because they may interfere with the imaging of the product.
- This product must only be used for assisting medical personnel in vein locating or for training and teaching medical personnel how to use it. This product is not the sole method of vein locating and cannot completely substitute other vein locating methods that are based on reliable medical judgments or sight and touch judgments.
- If the vein illuminator falls to the ground or it is affected by an external force, stop using vein illuminator even if it appears normal. Contact your local distributor and have an inspection performed to judge whether the vein illuminator is operating properly.
- Do not service or maintain the vein illuminator or its accessory when it is being used on a patient.
- Do not try to upgrade the software of the vein illuminator. To upgrade the software, please contact your local distributor for help. The software upgrade must be executed by trained technicians. Otherwise, an error of the vein illuminator may occur. After software upgrade, the vein illuminator must be validated by trained technicians before use.

# **A**CAUTION:

- Do not touch the display screen by using sharp objects. Otherwise, the display screen may get damaged.
- Ensure that the vein illuminator is placed beyond the reach of the patient and other unauthorized persons.
- Ensure that the battery is always installed in the vein illuminator during use.
- If the vein illuminator fails to act as specified herein for unknown reason, power it off and report the conditions when the fault occurs to your local distributor or the after-sales service department of MEDCAPTAIN.
- Do not disassemble or reconstruct the vein illuminator without permission.
- This product requires maintenance by authorized personnel. The authorized personnel can ask for such materials as the service manual and list of spare parts from the manufacturer.
- The power adapter is a part of the vein illuminator.

Symbol	Description	
$\triangle$	CAUTION	
•••	Manufacturer	
$\langle$	Alternating current	
*	Refer to instruction manual/booklet	
Â	General warning sign	

#### 2.2 Symbol Description

$\sim$	Date of manufacture	
	Direct current	
EC REP	Authorized representative in the European Community	
SN	Serial number	
C€	CE Mark: conforms to essential requirements of the Medical Device Directive 93/42/EEC	
X	DISPOSAL: Do not dispose of this product as unsorted municipal waste. Separate collection of such waste for special treatment is necessary.	
	Fragile, handle with care	
×	Keep away from sunlight	
-310-	Temperature limit	
×®	Stacking limit by number	
<u><u>†</u>†</u>	This way up	
Ť	Keep dry	
6UTkP2	Atmospheric pressure limitation	
	Humidity limitation	

### **3** Product Specifications

Name	Vein illuminator	
Model	NAVI-30/NAVI-60	
Dimensions	224 (W) x 68 (H) x 64 (D) mm	
Weight	About 0.5kg	
	Input: 100-240Vac 50/60Hz 1.5A Max	
Power Adapter	Output: 12V === 3.5A	
	Model of the power adapter:LXCP52-012	
	Built-in lithium battery: 7.3V, 2750mAh	
	Battery model: 18650-2S1P	
	Continuous operation duration of the lithium	
	battery: not shorter than 2.5 hours	
Battery	Time required for fully charging an exhausted	
Duttery	lithium battery: not longer than 4 hours (the	
	device is powered off during the charge)	
	Charge mode of the lithium battery: The	
	battery can be charged using a power adapter	
	when AC input is available.	
Display Mode	Projection mode	
Light Source	No	
Туре		
Infrared		
Wavelength	850nm dual light sources	
Optimal Focus	210 20	
Position	210mm±30mm	

### **Product Specifications**

Depth of Field of Imaging	h of Field of >30mm		
Infrared Radiation Energy	$\leq 0.6 \text{mW/m}^2$		
Operating Conditions	Temperature: 5 °C~40 °C Humidity: 20%~90% RH, non-condensing Pressure altitude: 70~106.0kPa		
Storage and Shipping Conditions	Temperature: -20 ℃~+55 ℃ Humidity: 10%~95% RH, non-condensing Pressure altitude: 61.7~107.4kPa		
Service Life	5 years		
Classification Date of	<ol> <li>Class I/Internally powered equipment;</li> <li>IPX0;</li> <li>Not sterilized;</li> <li>Not category AP/APG equipment;</li> <li>Mode of operation: continuous</li> </ol> See the product label.		
Manufacture Main Safety Standards	IEC60601-1:2012MedicalElectricalEquipment, Part 1: General Requirements for basic safety and essential performanceElectricalIEC60601-1-2:2014MedicalElectricalEquipment - Part1-2: General requirements for basic safety and essentialessentialfor basic safety and basic safety and essentialstandard:Electromagnetic compatibility-Requirements and testsstandard:		

### 4.1 Structural Composition

This product mainly consists of the infrared light source, image sensor (CCD), image processing chip, projection module, internal optical path module, power supply, and support. It does not come into contact with the patient during the use.

### 4.2 Operating Principles

Compared with the superficial skin, the hemoglobin in veins has a stronger absorption of near-infrared light. The vein illuminator is designed based on this principle. During use of the vein illuminator, the reflected infrared light is perceived by the CCD. After a series of digital image processing by the image processing chip, the vein illuminator shows the outline image of the veins and projects the image on the skin surface to reveal the vein distribution. Qualified medical personnel can observe and locate the vein for venipuncture or blood drawing according to the vein distribution image projected on the skin surface.

### 4.3 Main Unit

### 4.3.1 Front View



1 – Display screen	2 - Running indicator
3 – Battery indicator	4 - Power indicator
5 - Brightness button	6 – Size button
7 – Mode button	8 – Power button

- Display screen (only NAVI-60 has a display screen): Displays various operation information like the remaining battery capacity, mode, color, and projection size.
- Running indicator: This indicator is steady green when the device is in power-on state and extinguished when the device is in power-off state.
- Battery indicator:

- When the device is connected to an external power supply and the battery is in charge state, the battery indicator is steady green. After the battery is fully charged, the battery indicator is extinguished.
- When the battery is powering the device, the battery indicator is slowly blinking green. In case of low battery, the battery indicator is quickly blinking green. When the battery is exhausted, the battery indicator is extinguished.
- Power indicator: The power indicator is steady blue when the device is connected to an external power supply. When the device is in power-on state and not connected to any external power supply, the power indicator is slowly blinking blue.
- Power button: A user can press the power button to power on the device. When the device is in running state, a user can press the power button to power off the device. If the device is powered by the battery, it will be automatically powered off if no operation is performed within 4min.
- Brightness button: A user can press this button to adjust the projection brightness. Four brightness levels are available for selection.
- Size button: A user can press this button to switch the size of the projection image.
- Mode button: A user can press this button to switch between the basic mode, color mode, and depth mode. In addition, a user can press and hold this button and then release to switch the current display color.
- Basic mode: This mode is suitable for the users with different visual feelings.

- Green light mode, red light mode, blue light mode, and light purple mode: The noise is weakened to accurately show the vein distribution.
- Depth mode: The vein depth can be identified to assist medical personnel in inserting a needle.



#### 4.3.2 Rear View

1–Projection window: Used to receive infrared light and project an image.

2–Infrared lamp: Used to emit infrared light.

3-Battery compartment: rear cover of battery.

### 4.3.3 Bottom View



• Power interface: Used to connect to the power adapter.

#### 4.4 Accessories

No.	Accessory Name	
1	AC power cable	
2	Power adapter	
3	Accuracy test card	



 All accessories of this device must be provided by the manufacturer. Otherwise, this device may be damaged, an electric shock may be caused, or the device may fail to reach specifications asserted in the operation manual.

#### **5** Installation Description

#### 5.1 Environment Requirements

To ensure normal operation of the vein illuminator, please ensure that the installation environment meets the following requirements:

- If a support is used, the installation workbench must be smooth and steady.
- No power supply interference exists.
- No corrosive or flammable gas should be present.
- No flammable and explosive materials should be present.

#### 5.2 Open Package Inspection

Before opening the package, please inspect the packaging box carefully. In case of any damage, please contact your local distributor or the after-sales service department of the manufacturer immediately.

- 1. Take the vein illuminator and accompanied accessories out of the packaging box.
- 2. Check whether the accessories in the packaging box are consistent with those on the packing list, and check whether there is any mechanical damage on the device or its accessories. In case of any doubts, please contact the local distributor or the after-sales service department of the manufacturer immediately.

#### 5.3 Connecting the Power Supply

Place the vein illuminator in an environment meeting the requirements stipulated in section 5.1, and connect the device to an external power supply.

 Use the AC power cable and power adapter provided by the manufacturer.

### **Installation Description**

- 2. Connect the AC power cable and power adapter.
- 3. Connect one end to the power interface at the bottom of the vein illuminator.
- 4. Connect the other end to a matching AC power socket.





- Do not touch the power plug with a wet hand. If any liquid or liquid residue exists on or around the power plug or power socket, remove this liquid or liquid residue before plugging in the device. Otherwise, an accident may occur.
- Use the power cable provided by the manufacturer to ensure that the device is properly grounded. If the device is not properly grounded, the safety performance cannot be guaranteed and an electric shock may occur.

# **A**CAUTION:

• The plug of the AC power cable must be firmly and fully inserted into the power socket.

### **Installation Description**

• Do not install the vein illuminator at a place where the power plug is difficult to be disconnected from the power socket.

#### 5.4 Checking the Device Accuracy

Since the vein illuminator is a precise device, before powering on it for the first time or after transporting it, the operator must check the accuracy of the device using an accuracy test card.

Place the accuracy test card in the projection area of the device, and check the positional deviation of the projection image and test card image at the optimal focus position (the projection hole is 210±30mm away from the test card). If the deviation is smaller than 1mm, the device can be used properly. If the deviation is larger than 1mm, stop using the device and contact the local distributor or the after-sales service department of MEDCAPTAIN immediately.



#### 6.1 Powering On the Vein Illuminator

After installing the vein illuminator, power on the vein illuminator according to the following steps:

- 1. Press the power button to power on the device. The running indicator is illuminated, and the device starts to project 6s later.
- 2. Check the accuracy of the device according to section 5.4. After that, the device can be used to observe the superficial veins. During actual use, the user can adjust the projection distance until the English letters at both sides of the projection image achieve the optimal effect (or become the clearest).

### 6.2 Setting the Mode

The vein illuminator is suitable for the users with different visual feelings. A user can press the mode button to set the mode. Seven modes are available for selection: basic mode, green light mode, red light mode, blue light mode, light purple mode, depth mode, and inverse color mode. In any mode except depth mode, a user can press and hold the mode button and then release to switch to the inverse color mode of the current mode. In inverse color mode, the vein color and skin color in the projection image are inversed.

In depth mode, align the long side of the red cross in the middle of the projection with the vein to be detected. Then, you can observe that the green indicator is illuminated to indicate the vein depth information.

Mode	Image	Remarks
Basic mode		The image is monochrome.
Green light mode	RTHA	The image is green.
Red light mode		The image is red.

Blue light mode	THAT	The image is blue.
Light purple mode		The image is light purple.
Depth mode	Excitation Ventility	1 bar of green light: The vein depth is 0~2mm (shallow).
	H	2 bars of green light: The vein depth is 2~4mm (relatively shallow).



### 6.3 Adjusting the Brightness

Four levels of projection brightness are available for selection to match different use environments. A user can press the brightness button to adjust the brightness. See the following figures.

Level 1 brightness (darkest)	Level 2 brightness	
KTHA	KTYKA	
Level 3 brightness	Level 4 brightness (brightest)	
KTA	KTYA	

### 6.4 Adjusting the Size

Three different sizes of projection image are designed to make the device be suitable for different groups of people, especially for children and infants. A user can press the size button to switch the size. See the following figures.

Minimum Size	Middle Size	Normal Size
		KIN
360*192 pixel	360*288 pixel	720*576 pixel

### 6.5 Powering Off the Vein Illuminator

After use, press the power button to power off the device.

#### 7 Common Faults

• The device cannot be powered on after being connected to the power adapter.

#### **Possible cause:**

- a. The device is not properly connected to the external power supply or the external power supply has no voltage.
- b. The power board in the device is damaged because an incorrect power adapter is used.

#### Check method:

- a. Check whether the device is properly connected to the power adapter by checking whether the power indicator is steady on.
- b. Check that the device can be powered on when it is powered by battery and cannot be powered on when it is connected to the power adapter.
- The projection image is blurring or obvious spots can be found in the image.

Possible cause: The lens is dirty.

**Solution**: Clean the lens according to the method described in section 8.2.

• The device cannot be powered on when battery is used to supply power.

**Possible cause**: The battery is exhausted, or the battery is damaged because it was stored in a humid environment.

### **Common Faults**

**Solution**: Connect the **device** to the power adapter for charging the battery (the battery indicator is extinguished when the battery is fully charged), or directly use the external power supply to power the device. If the device still cannot be powered on, it may be abnormal. In this case, contact the after-sales service department of MEDCAPTAIN.

• The continuous operation duration of the battery is insufficient.

**Possible cause**: The battery is not fully charged, or the continuous operation duration of the battery decreases due to natural deterioration.

**Check method**: Connect the device to the power adapter. If the battery indicator is blinking, the battery is not fully charged. If the battery indicator is not illuminated, the battery is already fully charged. The decrease in the continuous operation duration of the battery is caused by increase in number of charge/discharge times, which is natural and inevitable.

**Solution**: If the continuous operation duration of the battery is too short, the battery needs to be replaced. Please consult the after-sales service department of MEDCAPTAIN about battery replacement.

• The device does not respond or breaks down occasionally. Possible cause: The device has been used for a long time and the internal temperature is too high.

**Solution**: Power off the device for a moment to cool down the device.

### 8 Cleaning and Disinfection

It is highly recommended that the materials and methods listed in this chapter be used for cleaning and disinfection of the device. If other materials or methods are used, the device may be damaged or its service life may be shortened.



- In case of any doubts about the use of the detergent or disinfectant, please consult the local distributor.
- Please dispose of the wastes generated after the cleaning and disinfection according to the relevant regulations of the local hospital.

### 8.1 Preparations

- 1. Before cleaning and disinfection, power off the device and disconnect the power cable from the device.
- 2. Wear a pair of rubber gloves and a gauze mask to prevent contaminants from splashing onto your skin during the cleaning and disinfection.
- 3. You are not allowed to disassemble this device for cleaning and disinfection. To disassemble this device for further cleaning and disinfection, please contact the local distributor.
- Prepare several pieces of soft medical gauze, a detergent container, and a disinfectant container.

### 8.2 Cleaning WARNING:

- Do not immerse the device in the detergent solution.
- Prevent the solution from seeping into the device.
- Do not use halogenated solvent, petroleum-based solvent, glass detergent, acetone, or other irritant detergents.
- Only manual cleaning is allowed to be adopted for this device.
   Do not adopt the automatic cleaning mode for this device.

Cleaning procedure:

- Completely immerse a piece of soft medical gauze in neutral or slightly alkaline detergent solution, wring out the gauze, and then use the gauze to wipe the device surface.
- 2. Wipe all the surfaces of the device in sequence until all the contaminants are removed from the device surface.
- 3. Drip several drops of absolute ethyl alcohol on a piece of lens paper and use this lens paper to gently wipe and clean the lens surface along the same direction.
- 4. Ensure that all the edges and corners of the device are completely cleaned.
- After the cleaning, use a piece of dry medical gauze to remove the residual detergent solution.

The following table lists the detergents recommended for the device.

Detergent Name	<b>Cleaning Method</b>
Clean water	Wipe
Soapy water (pH value: 7.0~10.5)	Wipe

Table 8-1	Recommended	detergents
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<sup>27/44</sup> 

### **Cleaning and Disinfection**

#### 8.3 Disinfection

### **M**WARNING:

- Do not immerse the device in the disinfectant solution.
- Prevent the solution from seeping into the device.
- Use the disinfectant according to its operation manual.
- Do not autoclave the device.
- Only manual disinfection is allowed to be adopted for this device. Do not adopt the automatic disinfection mode for this device.

Disinfection procedure:

- 1. Before the disinfection, clean the device according to the method provided in section 8.2.
- Completely immerse a piece of soft medical gauze in the intermediate-efficiency or high-efficiency disinfectant solution, wring out the gauze, and then use the gauze to wipe the device surface.
- Wipe all the surfaces of the device in sequence. For the contact time of the disinfectant, see the operation manual of the disinfectant.
- 4. Drip several drops of disinfectant solution on a piece of lens paper and use this lens paper to gently wipe and clean the lens surface along the same direction.
- Ensure that all the edges and corners of the device are completely disinfected.
- 6. After the disinfection, immerse another piece of soft medical gauze in clean water, wring out the gauze, and then use the gauze to wipe the device surface for removing the residual disinfectant solution.

### **Cleaning and Disinfection**

The following table lists the disinfectants recommended for the device and the required contact time for the disinfection.

|--|

Disinfectant Solution	Contact Time	Disinfection Method
Name		
75% alcohol	3min	Wipe
70% isopropanol	3min	Wipe
3% hydrogen peroxide	30min	Wipe

### 8.4 Air Drying and Transportation WARNING:

- Do not dry the device by using a drying machine or similar products.
- Connect the device to the power supply again after the device is completely dry.
- After cleaning and disinfection, place the device in a shady, cool, and ventilated environment for air drying.
- 2. If you are not going to use the device soon after air drying, place the device in its original package for storage and transportation.

### 9 Maintenance

#### 9.1 Regular Maintenance

To ensure safe use and lengthen the service life of the vein illuminator, please conduct regular maintenance and check. Table 9-1 lists the maintenance plan.

Table 9-1 Maintenance plan

Maintenance Item Frequency		Maintenance Method
Appearance check	Before each use	See section 9.1.1.
Power adapter and	Before each use	See section 9.1.2.
power cable check		
Accuracy check	Every two years	See section 9.1.3.
Electrical safety test	Every two years	See section 9.1.4.

### 9.1.1 Appearance Check

- Appearance check: Check that no crack or damage exists.
- Button operation: Check that the buttons can be smoothly pressed and function properly.
- Check that all the sealing parts and the installation of the vein illuminator are normal and no crack exists on any materials.

### 9.1.2 Power Adapter and Power Cable Check

- Check the appearance of the power adapter and power cable. If a surface damage or poor contact between plug and socket is found, contact the distributor for replacement in time.
- If the AC/DC power indicator is not illuminated after the vein illuminator is connected to an AC/DC power supply or the vein illuminator cannot be started, contact the distributor for maintenance in time.

### 9.1.3 Accuracy Check

Check the device accuracy according to section 5.4. If the accuracy exceeds the reference range, contact the local distributor or the after-sales service department of MEDCAPTAIN for calibration of the device accuracy.

### 9.1.4 Electrical Safety Test

To ensure safety, please conduct a dielectric strength test, leakage current test, and ground impedance test according to the method stipulated in IEC60601-1.

### 9.2 Battery Maintenance

### 9.2.1 Battery Overview

The NAVI-30/NAVI-60 vein illuminator is equipped with a built-in battery to ensure normal operation of the vein illuminator in case of an external power failure. The battery starts to be charged when the vein illuminator is connected to an external power supply. In case of a sudden power failure, the system automatically switches to the battery supply mode without interrupting the operation of the vein illuminator. If the external power supply recovers from the failure before the built-in battery supply mode to external power supply mode to ensure uninterrupted operation of the vein illuminator.



• If you have any doubts about the integrity or wire of the protective earth, unplug the device for the battery to power the device.

#### 9.2.2 Using the Battery

• Before using the device for the first time or using the device after the device is not used for a long time

Before using the vein illuminator for the first time, charge the built-in battery. Power off the vein illuminator and connect it to an external power supply for at least 10 hours until the battery is fully charged. After that, you are allowed to use the vein illuminator.



Battery optimization

- 1. Power off the vein illuminator.
- 2. Connect the vein illuminator to an AC power supply to charge the battery for over 10 hours uninterruptedly.
- 3. Disconnect the vein illuminator from the AC power supply for the battery to power the vein illuminator until the battery is exhausted.
- 4. Connect the vein illuminator to the AC power supply again to charge the battery for over 10 hours uninterruptedly.



- Before using the built-in battery, check the battery to ensure that sufficient power is available. Recharge the battery if required.
- Improper use of the battery may shorten the service life of the battery.
- Frequent use of the battery will shorten the service life and continuous operation duration of the battery.
- If this product will not be used for a long time, fully charge it for storage.

#### 9.2.3 **Replacing the Battery**

1. Remove the rubber plugs from the retaining screws on the battery compartment.

### Maintenance

- 2. Remove the retaining screws, and take down the battery compartment.
- 3. Take out the old battery, and install a new battery in the battery compartment.
- 4. Insert the battery plug into the socket in the direction shown in the following figure. Hold the socket when inserting or removing the battery. Do not hold the battery and pull the socket.



# **A**CAUTION:

- The service life or shelf life of the battery is one year. It is recommended that the battery be replaced in time.
- Aging may shorten the continuous operation duration of the battery. Please check and replace the battery at regular intervals.
- Replace the battery when the vein illuminator is not in use.
- Remove the battery if the vein illuminator is not likely to be used for some time.
- Battery replacement must be finished by trained technicians. Otherwise, a danger may be incurred.

### 9.3 Storage

- Avoid water spill.
- Do not store the vein illuminator in a hot and humid place.
- Store the vein illuminator far away from excessive vibration, dust, and corrosive gas.
- Store the vein illuminator out of direct sunlight and ultraviolet ray to avoid color fading.

#### 9.4 Transportation

The vein illuminator is allowed to be transported using a common vehicle, but it must be protected from the drastic impact, vibration, and rain and snow splash during the transportation. In addition, the vein illuminator must be transported in accordance with the requirements specified in the order contract.

### 9.5 Environmental Protection and Recycling

Contact your local distributor to recycle the retired vein illuminator, or otherwise dispose of it and its battery in accordance with the local laws and regulations.

### Appendix A Electromagnetic Compatibility

#### (EMC)



- The NAVI-30/NAVI-60 vein illuminator complies with EMC standard IEC 60601-1-2:2014.
- Users must install and use the NAVI-30/NAVI-60 vein illuminator based on the EMC information provided in the accompanying document.
- Portable and mobile RF (Radio-Frequency) communication devices may affect the performance of the NAVI-30/NAVI-60 vein illuminator. Avoid strong electromagnetic interference during the use, for example, stay away from mobile phone and microwave oven.
- For the declaration of emissions CLASS and group and Immunity level, please see the Appendix.
- The NAVI-30/NAVI-60 vein illuminator is suitable for Professional healthcare facilities environment, e.g. hospitals except for near active HF surgical equipment and the RF shielded room of an ME SYSTEM for magnetic resonance imaging, where the intensity of electromagnetic disturbances is high. Due to conducted interference and radiated interference, it may be difficult to ensure electromagnetic compatibility in other environments.
- To assure that the NAVI-30/NAVI-60 vein illuminator remains safe with regard to electromagnetic disturbances throughout the expected service life:
  - Conduct periodic maintenance based on the recommended maintenance/service interval and method provided in the operation manual.

- After each maintenance, ensure that the internal structure, shielding system, and grounding system of the device remain complete and effective.
- The Emissions characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.

### **WARNING**:

• Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.

Only the following cables provided by the manufacturer allowed to be used to meet the electromagnetic emission and anti-interference requirements.

No.	Cable Name	Length	Shielded
1	AC power cable	2.5	No
2	Power adapter	1.6	No

- Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation.
- Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no

closer than 30 cm (12 inches) to any part of theNAVI-30/NAVI-60 vein illuminator, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.

Guidance and manufacturer's declaration – electromagnetic emissions The NAVI-30/NAVI-60 vein illuminator is intended for use in the electromagnetic environment specified below.

The customer or the user of the NAVI-30/NAVI-60 vein illuminator should assure that it is used in such an environment.

Emission test	Compliance	Electromagnetic environment –	
Emission test	Compliance	guidance	
RF emission	Group 1	The NAVI-30/NAVI-60 vein	
CISPR 11		illuminator uses RF energy only for	
		its internal function. Therefore, its	
		RF emissions are very low and are	
		not likely to cause any interference	
		in nearby electronic equipment.	
Radio-frequency	Class A	The NAVI-30/NAVI-60 vein	
emission		illuminator is suitable for use in all	
CISPR 11		establishments other than domestic	
Harmonic	Class A	and those directly connected to the	
emission		public low-voltage power supply	
IEC61000-3-2		network that supplies buildings	
Voltage	Complies	used for domestic purposes.	
fluctuation and			
flashing			
IEC 61000-3-3			

Guidance and manufacturer's declaration - electromagnetic immunity

The NAVI-30/NAVI-60 vein illuminator is intended for use in the electromagnetic environment specified below. The customer or the user of the NAVI-30/NAVI-60 vein illuminator should assure that it is used in such an environment.

IMMUNI	IEC60601 test	Compliance	Electromagnetic
TY test	level	level	environment
			-guidance
Electrostat	±8 kV contact	±8 kV contact	Floors should be
ic	±2 kV,±4 kV,±8	±2 kV,±4	wood, concrete or
discharge	kV,	kV,±8 kV,	ceramic tile. If floors
(ESD)	±15 kV air	±15 kV air	are covered with
IEC			synthetic material,
61000-4-2			the relative humidity
			should be at least
			30%.
Electrical	±2 kV 100KHz	±2 kV	Mains power quality
fast	AC power cable	100KHz	should be that of a
transient	±2 kV 100KHz	AC power	typical commercial or
(EFT)	DC power	cable	hospital environment.
IEC61000	cable(>3m)		
-4-4	±1 kV 100KHz		
	SIP/SOP		
	cable(>3m)		
Surge	±0.5 kV, ±1 kV	±0.5 kV, ±1	
IEC	Line-to-line	kV	
61000-4-5	±0.5 kV, ±1	Line-to-line	
	kV,±2 kV	±0.5 kV, ±1	

	Line-to-ground	kV,±2 kV	
	AC power cable	Line-to-groun	
	DC power	d	
	cable(>3m)	AC power	
	±2 kV	cable	
	Line-to-ground		
	SIP/SOP		
	outdoor cable		
The	0% 0.5 cycle	0% 0.5 cycle	Mains power quality
voltage	At 0 °, 45 °, 90 °,	At 0 °, 45 °, 90	should be that of a
dips, and	135 °, 180 °,	°,135°,180°,	typical commercial or
interruptio	225 °, 270 °and	225 °,270 °	hospital environment.
ns	315°;	and 315 °;	If the user of the
IEC		0% 1 cycle	NAVI-30/NAVI-60
61000-4-1	0% 1 cycle	And	vein illuminator
1	And	70% 25/30	requires continued
	70% 25/30	cycles	operation during
	cycles	Single phase:	power mains
	Single phase: at	at 0°	interruptions, it is
	0 °	0% 300 cycle	recommended that
	0% 300 cycle		the
			NAVI-30/NAVI-60
			vein illuminator be
			powered from an
			uninterruptible power
			supply or a battery.
Power	30 A/m 50 Hz or	30 A/m 50 Hz	Power frequency
frequency	60 Hz	or 60 Hz	magnetic fields

magnetic		should be at levels
fields		characteristic of a
(50/60Hz)		typical location in a
(PFMF)		typical commercial or
IEC		hospital environment
61000-4-8		

NOTE: UT is the a.c. mains voltage prior to application of the test level.

Guidance and manufacturer's declaration – electromagnetic immunity The NAVI-30/NAVI-60 vein illuminator is intended for use in the electromagnetic environment specified below. The customer or the user of theNAVI-30/NAVI-60 vein illuminator should assure that it is used in such an environment.

Immuni	IEC 60601	Compliance	Electromagnetic
ty Test	test level	level	environment –guidance
Conduc	3 Vrms	3 Vrms	Portable and mobile RF
ted RF	150 kHz to	150 kHz to	communications equipment
IEC610	80MHz;	80MHz;	should be used no closer to
00-4-6		6 Vrms in	any part of the
		ISM bands	NAVI-30/NAVI-60 vein
	6 Vrms in	Between	illuminator, including
	ISM bands <sup>a</sup>	0.15MHz and	cables, than the
	Between	80 MHz;	recommended separation
	0.15MHz	80% AM at 1	distance calculated from the
	and 80	kHz	equation applicable to the
	MHz;		frequency of the
	80% AM at		transmitter.

Radiate3 V/m3 V/mdistanced RF80 MHz -80 MHz - 2.7 $d = 1.2\sqrt{P}$ IEC6102.7 GHz;GHz; $d = 1.2\sqrt{P}$		1 kHz		Recommended separation
00-4-380% AM at 1 kHz80% AM at 1 kHz $d = 1.2\sqrt{P}$ 1 kHzkHz80M~800MHz $d = 2.3\sqrt{P}$ 27V/m:380-3927V/m:380-39 390MHz;80M~800MHz $d = 2.3\sqrt{P}$ 390MHz;0MHz;800M~2.7GHz28V/m:430-28V/m:430-47where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m) <sup>b</sup> .28V/m:170028V/m:2400-2 28V/m:2400Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey <sup>c</sup> , a should be less than the compliance level in each frequency ranged. Interference may occur in the vicinity of equipment marked with the following symbol:	Radiate d RF IEC610 00-4-3	3 V/m 80 MHz – 2.7 GHz; 80% AM at 1 kHz 27V/m:380- 390MHz; 28V/m:430- 470MHz; 9V/m:704-7 87MHz; 28V/m:800- 960MHz; 28V/m:1700 -1990MHz; 28V/m:2400 -2570MHz; 9V/m:5100- 5800MHz;	3 V/m 80 MHz – 2.7 GHz; 80% AM at 1 kHz 27V/m:380-39 0MHz; 28V/m:430-47 0MHz; 9V/m:704-787 MHz; 28V/m:800-96 0MHz; 28V/m:1700-1 990MHz; 28V/m:2400-2 570MHz; 9V/m:5100-58 00MHz;	distance $d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P}$ $d = 1.2\sqrt{P}$ 80M~800MHz $d = 2.3\sqrt{P}$ 800M~2.7GHz where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m) <sup>b</sup> . Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey <sup>c</sup> , a should be less than the compliance level in each frequency range <sup>d</sup> . Interference may occur in the vicinity of equipment marked with the following symbol: ((()))

NOTE 1: At 80 MHz and 800 MHz, the higher frequency range applies. NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

<sup>a</sup> The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6,765 MHz to 6,795 MHz; 13,553 MHz to 13,567 MHz; 26,957 MHz to 27,283 MHz; and 40,66 MHz to 40,70 MHz.

<sup>b</sup> The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2,7 GHz are intended decrease the likelihood mobile/portable to that communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance for transmitters in these frequency ranges.

<sup>c</sup> Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which theNAVI-30/NAVI-60 vein illuminator is used exceeds the applicable RF compliance level above, theNAVI-30/NAVI-60 vein illuminator should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating theNAVI-30/NAVI-60 vein illuminator.

 $^{\rm d}$  Over the frequency range 150 kHz to 80 MHz, field strengths should be less than [3] V/m

Recommended separation distances between portable and mobile RF communications equipment and the NAVI-30/NAVI-60 vein illuminator The NAVI-30/NAVI-60 vein illuminator is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the NAVI-30/NAVI-60 vein illuminator can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the NAVI-30/NAVI-60 vein illuminator as recommended below, according to the maximum output power of the communications equipment.

Rated	Separation distance according to frequency of transmitter m			
maximum output power of transmitte r W	$150 \text{ kHz to } 80$ MHz outside ISM bands $d = 1.2\sqrt{P}$	150 kHz to 80 MHz in ISM bands $d = 1.2\sqrt{P}$	$80M \sim 800MH$ $d = 1.2\sqrt{P}$	$800M \sim 2.7G$ Hz $d = 2.3\sqrt{P}$
0.01	0.12	0.12	0.12	0.23
0.1	0.38	0.38	0.38	0.73
1	1.2	1.2	1.2	2.3
10	3.8	3.8	3.8	7.3
100	12	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance din meters (m) can be determined using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE 2: The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6,765 MHz to 6,795 MHz; 13,553 MHz to 13,567 MHz; 26,957 MHz to 27,283 MHz; and 40,66 MHz to 40,70 MHz. NOTE 3: An additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2,7 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 4: These guidelines may not apply in all situations.

Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

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