

Hunan Vathin Medical Instrument Co., Ltd. User's Manual

Name	User Manual of Digital Video Monitor	Page	0/30	No.	QP-DMR-DVM02-003
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# User's Manual Digital Video Monitor DVM-B1 DVM-B2



# **Rx only**

Caution: Federal law restricts this device to sale by or on the order of a physician.

- ◆ For use by trained clinicians/physicians only.
- ◆ For in-hospital use.
- ◆ For use with Vathin endoscope.
- ◆ Before use, thoroughly review this manual.
- ◆ Please keep all instruction manuals in a safe, accessible place.
- ◆ Have any questions or comments about this manual, contact with Vathin Medical.

# Hunan Vathin Medical Instrument Co., Ltd.

Version No.: A/0

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# 1. Important Information - Read Before Use

Please read these safety instructions carefully before using Digital Video Monitor. The instruction for use may be updated without further notice. Copies of the current version are available upon request. Please note that no clinical procedure is explained or discussed in this manual. The product can be operated only by a physician who has received clinical endoscopy training. Therefore, no clinical endoscopy procedure is explained or discussed in this manual only provides the basic operations and preventive measures for the Digital Video Monitor.

Before initial use of Digital Video Monitor, it is essential for operators to have received sufficient training in clinical endoscopic techniques and to be familiar with the intended use, warnings, cautions and contraindications mentioned in these instructions.

If, during the use of this device or as a result of its use, a serious incident has occurred, please report it to the manufacturer and to your national authority.

This Instruction for use only applies to the Vathin Digital Video Monitor. See relevant manuals for other products from Vathin Medical.

## 1.1. Intended Use

The equipment is specially designed to be used with medical endoscopes and other auxiliary equipment for the purposes of endoscopic diagnosis, treatment and video observation. Never use the product for any purposes other than those stated in this manual.

Digital video monitor is intended for patients requiring endoscopic diagnosis or treatment.



## 1.2. Contraindication

The product itself has no contraindications. If a doctor with appropriate qualifications believes that the use of this product will bring danger to the user, the product should not be used.

## 1.3. Repair and Refit

The product does not contain any user-repairable components. Never disassemble or refit or attempt to repair the product, which may cause injury to the patient or operator, damage to the product and/or failure to achieve the intended purpose. The product can be repaired only by the person authorized by Hunan Vathin Medical Instrument Co., Ltd.

## 1.4. Keywords

The following keywords are used throughout this manual:

# Warning

Indicates a potentially hazardous situation which, if not avoided, could result in death or serious injury.

# **A**Caution

Indicates a potentially hazardous situation which, if not avoided, could result in minor or moderate injury. It may also be used to warn of unsafe practices or potential damage to the product.

#### Note

Indicates other useful information.

## 1.5. Warnings and Cautions

Failure to observe these warnings or precautions may result in injury to the patient or damage to the product. Hunan Vathin Medical Instrument Co., Ltd. is not responsible for any system damage or patient injury caused by the improper use of the product.

# **A**Warning

As a BF-type application component, the endoscope connected to the product must not be directly
applied to the heart. The leakage current at the BF-type application component may be very dangerous, and
may cause ventricular fibrillation or severely affect the patient's heart function. Always observe the
following two points.

—Do not apply the endoscope connected with the product onto the heart or any area near the heart.

—Do not use endoscope treatment accessories or other endoscopes on or near the heart to make it contact the endoscope connected to the product.

- Do not install or use the product under the following conditions:
  - High oxygen concentration
  - Presence of oxidants (such as nitrous oxide) or flammable anesthetics in the atmosphere
- Do not use the product in an MRI environment.

• Do not use the product during the defibrillation process.

# **A**Warning

- To avoid the risk of electric shock, do not simultaneously touch the power socket, the docking connector or the bracket of the product when handling with the patients.
- To minimize the risk of contamination, always clean and disinfect the product as specified in Chapter 7 after each use.
- 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
- 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 the DVM-B1 & DVM-B2, including cables specified by the Vathin Medical. Otherwise, degradation of the performance of this equipment could result..
- Always observe the following precautions; otherwise, patients or medical personnel may be in danger.

—When using the product to examine patients, always prevent the metal parts of the endoscope or its accessories from touching any metal part of other components of the system.

-Keep all the electrical equipment away from any liquid. In case of any splash onto the product, stop it immediately and contact Vathin Medical.

- Wearing suitable protective filtering spectacles when the laser is used. Otherwise, eye damage may occur.
- When the lithium battery needs to be replaced, it needs to be returned to the manufacturer and replaced by professional personnel, otherwise it will cause certain risks

# **Caution**

- Always prepare an applicable and immediately available standby system to ensure that treatment procedure can be continued in the event of equipment failure.
- Always use the spare parts provided by Vathin Medical. Never refit any spare part.
- Always keep the product dry during preparation, usage and storage.
- The real-time imaging of the product may be affected by the use of high-frequency tools inside the connected endoscope. This phenomenon is not a malfunction. In such a case, just wait a few seconds to reset the image.
- The product is not intended to be repaired. In case that the product becomes defective, it should be discarded.

# 2. Symbols

Symbols	Expression	Symbols	Expression
CE	CE marking of conformity		Waste Bin symbol, indicating that waste mustbe collected according tolocal regulation and collection schemes fordisposal of electronic andelectrical waste (WEEE)
	General warning sign		Manufacturer
	Refer to instruction manual		Country of manufacture:CN(china) Production date, followed by YYYY-MM-DD.
₿ only	Indicates that the product can be used only with a prescription from a U.S. physician as required by the appropriate regulations of USA	LOT	Batch code
EC REP	Authorized representative in the European Community/European Union	Ť	Keep dry
<i>%</i>	Humidity limitation		Temperature limitation
( <b>*</b> )•( <b>*</b> )	Atmospheric pressure limitation	*	Keep away from sunlight
REF	Catalog number. Indicates the manufacturer's catalog number so that the medical device can be identifie	×	Type BF Applied Part
c <b>Al</b> us	UL Recognized Component Mark for Canada and the United States	15V <del></del> ⊖-€-⊕	Input voltage: 15V DC
$\bigcirc$	Application of endoscope Ports	HDMI	High definition multimedia interface Port
•	Universal serial bus Ports	品	wireless local area networks Port

	Only for indoor use	FC	Tested to comply with FCC Standards-medical Equipment
Ţ	Fragile, handle with care	#	Model number
MD	Medical device	UDI	Unique device identifier
	Protective earth (ground)		Direct current

# 3. System

The product is designed as the system controller of endoscopic image observation system to display, record and print endoscopic images. Certain functions described below can only be enabled when the necessary equipment is connected to the product. For information about the compatible endoscope and other equipment, please refer to their Instruction for use.

## 3.1. Digital Video Monitor

No.	Model	Specification of endoscope port	
1	DVM-B1	14PIN	
2	DVM-B2	16PIN	

## 3.2. Product Compatibility

The Digital Video Monitor has been designed to be used in conjunction with Vathin endoscope and medical monitor. The compatible equipments are listed below.

## 3.2.1. Compatible Endoscopes

No.	Model	Compatible endoscopes
1.	DVM-B1	Single-use Flexible Ureteroscope Single-use Flexible Cystoscope Single-use Flexible Video Rhinolaryngoscope
2.	DVM-B2	Single-use Flexible Video Rhinolaryngoscope

## 3.2.2. Compatible Monitor

Item	Recommended parameter
Screen brightness	≥ 300 cd/m²
Color gamut	BT.709 or SMPTE-C, 72% NTSC
Resolution	480/60 p,480/60 i, 576/50 p, 576/50 i, 720/60 p, 720/50 p, 1080/50 i, 1080/50 p, 1080/60 p, 1080/30 p, 1080/60 i
Monitoring technology	Bright LED backlight technology
Input signal type	DVI-D or HDMI Video bandwidth < 165 MHz TMDS: 600 mV for each differential line Input impedance: 50 ohm
Monitor size	8.5", 10", 15",19", 21",24", and 26"
Contrast	≥ 1000/1

# 4. Checking the Package Contents

## 4.1. Package Contents

Open the product packaging to ensure that all the components are provided. Check all the items in the package with the components shown below. Check each item for damage. Please contact Vathin Medical if any component is missing or damaged.

Items	Quantities	Pictures
DVM-B1 (or DVM-B2)Digital Video Monitor	1 pcs	
Monitor cable (HDMI-DVI)	1 pcs	
Power cable	1 pcs	
Power adapter (Type:HPU63A-106,manufactur er:Sinpro Electronics Co.,Ltd)	1 pcs	<i>.</i> 7

USB cable	1 pcs	
HDMI cable	1 pcs	
User's manual	1 pcs	C VERN We have and We have and We have and We have an and
Multilingual User's Manual (CD)	1 pcs	
Reset pin	1 pcs	

# 4.2. Optional Accessories

ltems	Quantities	Pictures
U Disk	1 pcs	
pylons	1pcs	



# 4.3. Structure and Function



No.	Part	Function
1.	Power button	Long press the power button to turn on the product, and press it to
		turn off the product.
2.	Power indicator	It is used to indicate the battery power status.

		The light is blue and keep on when the device is on. And it flashes
		when the device is being charged and when the battery is low.
3.	Display screen	It is used to display the image after the endoscope is connected,
		and it is a touch screen.
4.	LAN interface	It is used to connect to the local area network through the network
		cable.
5.	HDMI interface	It is used to connect an external high-definition monitor for display
		expansion.
6.	USB interface 3.0	It is used to connect external devices, such as U disk.
7.	USB interface 3.0	It is used to connect external devices, such as U disk.
8.	Power jack	It receives 15V DC input, and provides the power supply for the
		entire system.



No.	Part	Function
9.	USB 2.0	It can be connected to a PC for image display.
10.	Reset button	You can insert a pin into the hole and press the reset button in the
		hole to restore the factory settings.

		*It is recommended to use a $\phi$ 1 mm and 10 mm long pin.	
11.	Video endoscope	The video endoscope cable can be inserted to this socket to	
	cable connector	connect the video endoscope and the product.	
	socket		
12.	Ventilation holes	Cool hardware during use.	
13.	Bracket	It is used to support the product on a plane surface.	

# 5. Preparation and Operation

When using the product for the first time, please refer to this User's Manual and install the product according to the following steps.

# **A**Warning

- Do not get the power cable wet; otherwise, electric shock may be caused.
- Do not prepare, inspect or use the product with wet hands.
- Do not bend, pull or twist the power cable; Otherwise, electric shock, equipment damage or fire may be caused.
- The product can be operated only under the conditions that meets the requirements given in "Operating Environment", "Storage and Transportation Environment" of Chapter 9. Technical Specification; Otherwise, malfunction, impaired safety and/or equipment damage may be caused.
- Do not place any equipment on top of the product; otherwise, the product may be damaged.
- Do not install the product near any strong electromagnetic radiation source (e.g. microwave therapy equipment, shortwave therapy equipment, and MRI); Otherwise, the product may malfunction.
- Before each procedure, inspect the Digital Video Monitor as instructed below. Inspect other equipment to be used with this Digital Video Monitor as instructed in their respective instruction manuals. If any irregularity be observed, do not use the Digital Video Monitor.

# **A**Caution

- Do not use any sharp object to operate the screen of the product; otherwise, the screen may be damaged.
- Never apply excessive force to the connector; otherwise, the product may be damaged.

# 5.1. Preparation and Inspection

## <u> W</u>arning

 In order to avoid the risk of power supply, the power supply with medical certification specified in the manual should be selected. When the power supply needs to be connected, only the type and manufacturer as stated in the packing list is allowable. Otherwise, operator or patient injury may occur.

- To avoid the risk of electric shock, only the appropriately approved medical electrical equipment are allowed to be connected with the product.
- The product can only be connected to a medical display screen that has been qualified as per IEC 60601-1.
- 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.
- Do not use the product in case of any damage to it or any unacceptable item in the functional inspection. Otherwise, operator/patient injury may occur.

1. Prepare Digital Video Monitor, power cable, power adapter, cables and device used together. Carefully check the product and all the components for damage (and without wear). Never use the product in case of any damage.

2. Prepare and inspect the endoscope according to its instruction manual.

# 5.2. Installation of Digital Video Monitor

# 🚺 Warning

- To avoid risk of electric shock, this equipment must only be connected to a supply mains with protective earth. Electric shock and/or fire may be caused if the product is not properly grounded.
- All the system components shall be powered off before connecting them. Only appropriate cables can be used for the product;Otherwise, the product may be damaged or malfunction.

# **A**Caution

• Always place the power cable in the place where it is unlikely to be stepped on. Do not place any objects on the power cable.

1. Open the bracket (No. 12 in Section 4.3 Structure and Function) of the product and place it on a solid surface. Do not place the Digital Video Monitor in a place which is difficult to operate the disconnection device.

2. Connect the power cable to the power adapter, and connect the power adapter to the power jack (No.8 in Section 4.3 Structure and Function) of the Digital Video Monitor.

3. Connect the power cable to a supply mains with protective earth. The power supply should be stable. In case of severe fluctuations in the power supply, a power regulator or UPS power supply should be used for adjustment purpose.

#### 4. Connect the Digital Video Monitor with medical display or PC as needed.

**5.** Install the endoscope to the Digital Video Monitor by inserting the endoscope connector into the endoscope socket (No. 10 in Section 4.3 Structure and Function) properly as the arrow indicated.

# 5.3. Power-on and Setting

## 🔔 Warning

- When the loss of built-in power supply under normal working condition would result in unacceptable risks, the medical electrical equipment must be connected to an appropriate external power supply.
- If battery is used, check battery capacity before use. Otherwise patient injury may occur.
- 1. long press the power button to turn on the product. After the boot animation is over, the Basic

Function interface (Section 6.1 Basic Function Interface) will appear by default.

2. The real-time image will be displayed on the screen. Icons on the screen turn green and their function are available.

- 3. If battery will be used, check battery capacity of the equipment, and charge when the battery is low.
- 4. Set the parameters of Digital Video Monitor according to section 6.3 System Setting Interface.



#### Note

- The battery capacity icon in the lower left corner can display the remaining battery capacity, which changes depending on the remaining capacity and status of the battery.
- When the battery capacity is greater than 20%, this icon is green; when the battery capacity is less than 20%, it is orange; when the battery capacity is less than 10%, it is red, and a text prompt of insufficient

battery capacity appears on the screen.and a lightning symbol  $\overset{4}{\vee}$  appears in the icon when charging.

# 5.4. Inspection of the System

#### 🔔 Warning

• If any irregularity be observed during the inspection, do not use the Digital Video Monitor. Damage or irregularity may compromise patient or user safety and may result in more severe equipment damage.

1. Inspect photographing, recording function of the equipment. Please refer to Section 5.5.1 Photo-shooting and Section 5.5.2 Video Recording.

2. Inspect the endoscope function according to the instruction for use of the endoscope.

3. Before each use check to ensure the view observed through the display provides a live image and has the correct image orientation.

# 5.5. Operation of the Digital Video Monitor

#### 🔔 Warning

- When advancing or withdrawing a Vathin endoscope, always watch the real-time endoscopic image on the monitor.
- 1. Enter patient information: Enter the patient information according to Clause 6.1.1.
- 2. Adjust the white balance: Adjust the white balance according to Clause 6.1.2.
- 3. Perform examination: related function of the Digital Video Monitor please refer to Chapter 6 Function.

## 5.6. After Use

1. Disconnect the Vathin endoscope from the product. For information on how to dispose of the Vathin endoscope, refer to its user's manual.

2. Press the power button to turn off the product, and disconnect the power cable. To disconnect Digital Video Monitor from mains, remove the mains plug from the wall outlet.

3. Clean and disinfect the product according to Chapter 7 Cleaning and Disinfection.

## 6. Function

#### Note

- This product sets up user account management and authority management, and related functions are only open to users with authority. This product manual does not describe the authority of each user, and the authority of each user is subject to the actual authority provided by this product.
- Some key functions need to enter the user password when using, The initial user password of this product is 123456, you can reset the user password according to the instructions in the manual.
- The current interface can be exited by swiping from the right to the left of the screen.

# 6.1. Basic Function Interface



Description of basic function icons			
lcon	Name	Function	Reference section
	Basic function	Icon for the basic function interface	6.1
	File management	Icon for the file management interface.	6.2
8	System setting	Icon for the system setting interface	6.3
	Photo-shooting	Tap this button or the Vathin endoscope camera button to take the image and automatically save it in the patient file. The endoscope is connected. And function of the icons are available.	6.1.3
	Video recording	Tap this button to start recording video. Tap it again to stop recording and save the recorded video in the patient file.	6.1.4

	White balance	The white balance adjustment is required before surgery.	6.1.2
	Patient information	Tap it to enter the patient information mode.	6.1.1
	Patient information	Tap it to return to the basic function interface	/
	Photo preview	Photo preview zone, will be updated after the image being taken. Tap it to preview the latest images.	6.1.3
(((•	WiFi	Used to indicate the WiFi connection/disconnection	6.3.3
	Endoscope connection	Used to indicate the endoscope connection/disconnection upon on endoscope insertion status	/
	Memory	Used to indicate the available memory. A prompt will be given in case of insufficient memory.	/
	Memory	Storage space is not enough	/
	Battery capacity	Used to indicate the remaining battery capacity.	/
	Battery capacity	Low power capacity	/

## 6.1.1. Enter Patient Information

1. Tap to enter the Patient Information interface. Tap the input box on the screen to enter patient information through the keyboard that appears. Icon Can be used to enter the next input box.

Patient information		Patient information	
Patient ID: Please fill in the patient ID		Patient ID: Please fill in the patient ID	
Main Doctor: Please fill in the name of the doctor		Main Doctor: Please fill in the name of the doctor	
Name: Please fill in the patient name	·#·		*
Age: Please fill in the patient age	G	q w e r t y u i o j	p° 🖾
Gender: D Male D Female	<b>9</b>	as dfg hjkl	0
DETERMINE CANCEL CLEAR		\star z x c v b n m !	? *
		?123 ,	. ©

2. After the operation, tap clear to re-enter the patient information; tap "Determine" to confirm the operation, or tap "cancel" to cancel the operation and the Basic Function interface will appear automatically with the patient information displayed on the upper left corner.

3. Tap 😉 again to modify or clear the patient information being entered.

# **A**Caution

Once the patient information being entered, the photos and videos taken will be appended with the
patient information until the patient information is cleared.

#### Note

• The image showed in the image zone will be cleared after the patient ID is modified.

## 6.1.2. White Balance Adjustment

This adjustment procedure is used to display the correct image color on the monitor. Be sure to always adjust the white balance in the following cases:

- \* Before observation.
- \*After exchange of the endoscope
- \*Restart the Digital Video Monitor

\*When any abnormality can be seen on the color of the image even if white balance adjustment has been completed.

## <u> (</u>Warning

• When adjusting the white balance of the endoscope, use a white object such as a piece of gauze without bringing it in contact with the endoscope. Contact of the endoscope with a non-sterilized object may result in cross-contamination.

# **A**Caution

• When adjusting the white balance, make sure the light built in the distal end is on and take care not to expose the distal end of the endoscope to external light. Otherwise, it may cause an incorrect white balance adjustment.

1. The adjustment procedure is used to display the correct image color on the monitor. Be sure to always adjust the white balance before observation or when any abnormality can be seen on the color of the image even if white balance adjustment has been completed.

2. Hold the endoscope stable to avoid wash-out of the monitor image, monitoring a white object such as a piece of gauze in such a way that it does not contact the endoscope,, contact of the endoscope with a non-sterilized object may result in cross-contamination. Vathin Medical does not provide any accessories (e.g. white cap) for white

balance.

3. Maintaining the stable condition in step 1, tap the icon on the screen. Icon will appear in the upper right corner of the image. When the icon disappears, the white balance adjust is complete.

#### 6.1.3. Photo-shooting

1. There are two methods to take photos.

- \* Press the photographing button on the endoscope
- \* Tap Icon on the screen.
- 2. When taking photos, icon oppears in the upper right corner of the screen. The photo has been taken

when the icon disappears, and "Photo saved successfully" appears on the screen.

3. After taking a photo, the image area in the lower right corner will be updated accordingly. Click the updated image to preview the image.

#### Note

Keep the Vathin endoscope as still as possible to prevent obscure images during the photo-shooting.

## 6.1.4. Video Recording



The maximum allowable length of the video is 70 minutes (i.e. 4GB). Therefore, any video recording over 70 minutes will be automatically stored by splitting into two or more videos.

## 6.2. File Management Interface

#### 6.2.1. Enter the File Management Interface

Tap the File Management button <sup>1</sup> to enter the file management interface. This interface is used mainly for operations such as file viewing, folder editing & deleting, file moving, and medical record editing, as shown in

the figure below.



Description of file management-related icons			
lcon	Name	Function	Reference section
	File management	Icon for the file management interface.	6.2
	File editing	You can tap the icon or long press a folder/file to enter the selection mode, and then perform multiple selection, cutting, renaming, and deletion operation on the file or folder.	/
	USB storage device management	Used to indicate the connection of USB storage device. When connecting a storage device, you can choose to copy/cut files to a USB storage device. If no USB storage device is connected, the icon is gray, which means this function cannot be used.	1
	Medical record	Used to edit the medical records	6.2.3
	Save Medical record	Used to save pictures of medical records	6.2.3
	Print	Used to print the medical records, where Wifi connection is required	6.2.3
	Erase	Used to erase the current data	6.2.3

# 6.2.2. Photos/Videos Replay

In the File management interface, tap the icon 📁 to open the intended folder. Tap the corresponding image/video to replay.

#### Note

- The Photos/Videos taken are stored according to date and patient ID, and the file folder is named accordingly.For example, if the operation date is Sept. 17, 2020, patient ID is 12, the file path will be: >2020-09-17>12.
- If the patient ID is not set: A "pictures/medias" folder will be generated in the date folder after photo-shooting/video-recording, and the file path will be: >2020-09-17>picture/medias,
- The image are stored in PNG format, video are stored in MP4 format.

## 6.2.3. Medical Record Interface



# 6.3. System Setting Interface

Tap to enter the System Setting interface. This interface is used mainly for setting brightness, time, user permissions, network and language, and viewing the equipment information.



leen	Nama	Function	Reference
icon	Name	Function	section

	Screen adjustment	Used to adjust the brightness, sharpness, gamma, red saturation, green saturation, blue saturation	/
	Calendar	Used to set the date and time	/
	Equipment information	Used to view the equipment information	/
	User account	Used to set the user accounts	/
DELTE	Erase data	Used to erase internal files and user data	/
	Network setting	Used to set languages, Wifi and LAN	6.3.1

## 6.3.1. WiFi Setting

# **W**arning

- Please connect the Digital Video Monitor to a reliable network. An unreliable network may cause data loss or equipment failure.
- Related warning and information about the wifi please refer to Appendix 3. Information About Wifi.

On the area of wifi setting, turn on WiFi, and enter a valid password to connect to the network.

After the network is connected normally, the network connection symbol 🛜 appears in the status bar of the screen.

## 6.3.2. LAN setting

# **M**Warning

• When connecting to a network, make sure that the device is connected to a private local area network and that the private network is secure. Reliability of the network needs to be confirmed when network settings are updated. The unreliable network will cause a certain risk of data loss or function failure of the device.

Plug the end of the network cable into the Ethernet interface **b** of the device. After the network is connected normally, the network setting interface displays the network IP address.

#### Note:

- Digital Video Monitor uses WIFI direct connection to connect to mopria-supported WIFI printers for printing. The expected communication data involves medical and health data.
- Digital Video Monitor can be connected to the local Internet through the RJ45 interface to obtain the network synchronization time. The intended communication involves the interaction of network synchronized time data.

# 7. Cleaning and Disinfection

The product should be cleaned and disinfected before and after each use. It is recommended to clean and disinfect the product according to the following instructions before and after use. Vathin Medical has validated these instructions according to AAMI TIR 12 and 30. Any deviations from the instructions shall be assessed properly to determine its effectiveness and potential adverse consequences to ensure that it continues to achieve its intended purpose.

## <u> (</u>Warning

• Disconnect Digital Video Monitor from any mains power supply, remove any accessories and make sure the Digital Video Monitor is completely turned off before cleaning and disinfection.

You should take the following steps to clean and disinfect the product according to good medical practices:

# 7.1. Cleaning

- 1. Recommended detergent: enzymatic, mild pH of 7–9, and low foam (Enzol or equivalent).
- 2. Prepare the cleaning solution by using the standard enzymatic detergent prepared according to the manufacturer's recommendations.
- 3. Dip the sterile gauze into the enzyme solution and ensure that the gauze is moist without dripping.
- 4. Use the wet gauze to thoroughly clean the buttons and housing of the product. Well protect the product from being wet to avoid damage to its internal electronic components.
- 4. Use a sterile soft brush dipped in enzyme solution to clean the buttons until all dirt is removed.
- 5. Wait 10 minutes (or the duration recommended by the detergent manufacturer) for the enzyme to be activated.
- 6. Use the sterile gauze dipped in RO/DI water to clean the product to ensure that all traces of detergent are removed.

7. Repeat Steps 1 to 6 until the Digital Video Monitor is clean.

# 7.2. Disinfection

1. Disinfection Solution: isopropanol (alcohol) concentration 70-80%;

2. Preparation: add 80 cc of 95% isopropanol (alcohol) to 20 cc of purified water (PURW) (or use the EPA registered medical disinfectant wet wipes containing at least 70% isopropanol; all the appropriate safety precautions and manufacturer's instructions must be followed).

3. Use a piece of sterile gauze dipped in the alcohol mixture described above to wipe the surface of the product for about 15 minutes (approximately once every 2 minutes). The isopropanol should be handled according to appropriate safety procedures. The gauze should be moist without dripping because any liquid may impair the electronics inside the product. Always pay close attention to the buttons, housing, slots and gaps on the product. The sterile cotton swabs should be used for the disinfection in these areas.

## 8. Maintenance and Disposal

## 8.1. Storage

After being cleaned and disinfected, the product must be subjected to the pre-inspection procedures described in Chapter 5. The product must be stored during the period between two operations according to local guidelines and the storage environment requirements in this manual.

## 8.2. Battery Maintenance

In order to extend the service life of the battery, it is recommended to fully charge the product at least once every three months. The battery can last over 6 hours. The temperature range during charging should be controlled within 10–35 °C.

If the battery needs to be replaced, never refit any accessory of the product at will, and instead, contact your local dealer or Vathin Medical.

## 8.3. Return the Digital Video Monitor for repair

Vathin Medical shall have the right to request the technical department or equivalent at the customer's site to repair the product under the proper guidance of Vathin Medical. Any defective Vathin Digital Video Monitor must be disposed of by the person authorized by Vathin Medical. Circuit diagrams, component part lists, descriptions, calibration instructions, or other information are available on request by auhorized SERVICE PERSONNEL.

In order to prevent infection, it is strictly forbidden to transport contaminated medical equipment. The medical equipment must be disinfected on site before being shipped to Vathin medical. The cleaning and disinfection procedures described in Chapter 7 must be followed. Vathin Medical reserves the right to return contaminated medical equipment to the sender.

# 8.4. Disposal

After the product's service life expires, you should remove the monitor, and dispose of the battery and the monitor separately according to local regulations by using the disposal equipment that meets appropriate national and local regulations.

# 9. Technical Specification

## 9.1. Standard Applied

The Vathin<sup>®</sup> Digital Video Monitor meets the following standards:

- IEC 60601-1 Medical electrical equipment Part 1: General requirements for basic safety and essential performance
- IEC 60601-1-2 Medical Electrical Equipment Part 1–2 General requirements for basic safety and essential performance Collateral Standard: Electromagnetic disturbances Requirements and tests

# 9.2. Specification

Item	Specifications
Function	
White balance	Manual
Data record	Videos (MP4)/photos (png)
Memory	64G memory
Language setting	English/Français/日本語/中文简体/Deutsch/Español
Time setting	Manual/Internet-based self-calibration
Monitor	12.1"; 1280 × 800; touchscreen
Video output mode	HDMI
Screen brightness	Manual adjustment
Wifi	IEEE 802.11ac/a/b/g/n
LAN connection	RJ45,10/100/1000 Mbps
USB connection	A-type
Expected service life	5 years
Electrical power (Digital Video Monitor)	
Input Power	15V DC,4.2A max
Internal battery	11.1V DC 9000mAh, over 6 h endurance

Power adapter		
Power supply requirement	100–240 V AC; 47–63 Hz; 1.62–0.72 A	
Output Power	15V DC,4.2A max	
Protection against electric shock	class I	
Dimensions		
Length× width × thickness	310 mm × 231 mm × 39 mm	
Weight	2140g	
Storage and transportation		
Recommended storage temperature [°C, (°F)]	-10 °C to +40 °C (14–104 °F)	
Relative humidity [%]	10% to 80%	
Atmospheric pressure [kPa]	50 kPa to 106 kPa	
Operating environment		
Temperature [°C, (°F)]	+10 °C to +40 °C (50–104 °F)	
Relative humidity [%]	30% to 80%	
Atmospheric pressure [kPa]	86 kPa to 106 kPa	

# 10. Troubleshooting

If there is a problem with the product, please refer to this chapter to determine the cause and solve it. If the problem still exists after troubleshooting, please contact your local dealer.

Phenomenon	Possible cause	Recommended measures
The device cannot be turned on	The battery is low or the battery is protected	Connect the power adapter to charge the battery or wake up the protected battery, and then press the power button to start the machine
No image	The device cannot recognize the medical endoscope	Replacement of disposable medical endoscope, If there is still no image, please contact the agent for treatment
HDMI cannot output normally	The monitor is not compatible, the monitor is not set correctly	Follow the steps below: Reconnect the device to the monitor and set up the monitor correctly Replace the monitor of another model If not, please contact your local dealer

# **Appendix 1. Electromagnetic Compatibility**

#### **Essential Performance**

The product is designed to provide images for observation, excluding short-term automatically recoverable reduction caused by electromagnetic interference.

Cable	Max. cable length Shielded/unshielded		Qty.	Category
AC power cable	1.8 m	Unshielded	1 set	AC power supply
DC power cable	1.3 m	Unshielded	1 set	DC power supply
USB cable	2.8 m	Shielded	1 set	Signaling
HMDI-DVI cable	2.8 m	Shielded	1 set	Signaling
HDMI cable	2.8 m	Shielded	1 set	Signaling

The following cable information is given for EMC reference only.

#### Important information on electromagnetic compatibility (EMC)

The product shall be subjected to special precautions regarding EMC, and the EMC information provided in its user's manual shall apply when using it. The product complies with IEC 60601-1-2:2014 in terms of immunity and emission. However, the following special considerations shall be observed:

The equipment with above-mentioned ESSENTIAL PERFORMANCE is intended used in Professional healthcare facility environment, except for any places that is near the RF shielding room of the ME system for magnetic resonance imaging (MRI) with high electromagnetic interference intensity.

# 🔨 Warning

- It is required to avoid using the product next to or stacked with other equipment; Otherwise, improper
  operation may be caused. If you have to do so, the product and other equipment should be observed carefully
  to verify that they can operate properly.
- The use of accessories, transducers or cables other than those specified or provided by the manufacturer of the product may increase electromagnetic radiation or reduce electromagnetic interference, and cause improper operation.
- When using any portable radio frequency communication equipment (including antenna cables, external antennas and other peripheral equipment), such equipment should be placed over 30 cm (12 inches) away from any part of the product, including the cables specified by the manufacturer; Otherwise, the performance of such equipment may be impaired.

#### Statement

- The product is designed compatible with the high-frequency surgical equipment, i.e. it is allowed to keep working or standby near the high-frequency surgical equipment.
- In case of interrupted AC input voltage, the product will shut down, and once the power supply is restored, it can be manually restored by the operator. The degradation in such case is acceptable because it will neither cause unacceptable risks nor impair basic safety or essential performance.
- The degradation due to electrostatic discharge or electrical fast transient/burst is acceptable because it will neither cause unacceptable risks nor impair basic safety or essential performance.

The vertical-line flashing on the screen can automatically restore to the previous state.

## **EMI compliance**

Phenomenon	Compliance	Electromagnetic Environment
Radio frequency emission	CISPR 11 Group 1, Class A	Professional medical environment
Harmonic distortion	IEC 61000-3-2 Class A	Professional medical environment
Voltage fluctuation and Flicker	IEC 61000-3-3 Compliance	Professional medical environment

## Table 1. Emission

**Caution:** The emission characteristics of the product make it suitable for use in industrial areas and hospitals (CISPR 11 Class A). If being used in the residential environment (CISPR 11 Class B required usually), the product may not provide adequate protection for radio frequency communication services. The user may be required to take mitigation measures, for example, relocating or redirecting the product.

#### **EMS compliance**

## **Table 2. Enclosure Port**

Dhanamanan	Dhonomonon Basis EN/C standards	Immunity test level
Phenomenon	Dasic Elvic Standards	Professional medical environment
Electrostatic discharge	IEC 61000-4-2	±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air
Radiated, radio-frequency, electromagnetic field	IEC 61000-4-3	3 V/m 80 MHz–2.7 GHz 80% AM at 1 kHz
Fields near the radio-frequency wireless communication devices	IEC 61000-4-3	Refer to Table 3

Rated power frequency magnetic fieldIEC 61000-4-830 A 50 or	A/m or 60 Hz
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# Table 3. Fields near the radio-frequency wireless communication devices

Test frequency	Band	Band Immunity test level	
(MHz)	(MHz)	Professional medical environment	
385	380–390	Pulse modulation 18 Hz, 27 V/m	
450	430–470	FM, ±5 kHz deviation, 1 kHz sine, 28 V/m	
710			
745	704–787	Pulse modulation 217 Hz, 9 V/m	
780			
810			
870	800–960	Pulse modulation 18 Hz, 28 V/m	
930			
1720			
1845	1700–1990	Pulse modulation 217 Hz, 28 V/m	
1970			
2450	2400–2570	Pulse modulation 217 Hz, 28 V/m	
5240			
5500	5100–5800	Pulse modulation 217 Hz, 9 V/m	
5785			

# Table 4. Input AC power supply ports

Phenomenon	Desis FMC standards	Immunity test level
	Dasic Elvic Standards	Professional medical environment
Electrical fast transient/burst	IEC 61000-4-4	±2 kV 100kHz repetition frequency
Line-to-line surge	IEC 61000-4-5	±0.5 kV, ±1 kV
Line-to-ground surge	IEC 61000-4-5	±0.5 kV, ±1 kV, ±2 kV
Conducted disturbances induced by radio-frequency fields	IEC 61000-4-6	3 V, 0.15–80 MHz 6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz
		0% U <sub>T</sub> ; 0.5 cycle At 0º, 45º, 90º, 135º, 180º, 225º, 270º, and 315º
Voltage dips	IEC 61000-4-11	$0\% U_T$ ; 1 cycle and $70\% U_T$ ; 25/30 cycles Single phase: at $0^{\circ}$
Voltage interruptions	IEC 61000-4-11	0% U <sub>T;</sub> 250/300 cycles

Table 5.	Signal	input/	output	ports
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Dhanamanan	Basic EMC standards	Immunity test level
Phenomenon		Professional medical environment
Conducted disturbances induced by radio-frequency fields	IEC 61000-4-6	3 V, 0.15–80 MHz 6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz

# Appendix 2. Information About Wifi

# Warnings

Please take attention that changes or modification not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment.

This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions:

- (1) This device may not cause harmful interference, and
- (2) This device must accept any interference received, including interference that may cause undesired operation.

This equipment complies with FCC/IC RSS-102 radiation exposure limits set forth for an uncontrolled environment. This equipment should be installed and operated with minimum distance 20cm between the radiator & your body.

This device complies with Industry Canada licence-exempt RSS standard(s). Operation is subject to the following two conditions:

- (1) this device may not cause interference, and
- (2) this device must accept any interference, including interference that may cause undesired operation of the device.

Le présent appareil est conforme aux CNR d'Industrie Canada applicables aux appareils radioexempts de licence. L'exploitation est autorisée aux deux conditions suivantes :

- (1) l'appareil ne doit pas produire de brouillage, et
- (2) l'utilisateur de l'appareil doit accepter tout brouillage radioélectrique subi, même si le brouillage est susceptible d'en compromettre le fonctionnement.

(For licence-exempt equipment with detachable antennas, the user manual shall also contain the following notice in a conspicuous location)

Under Industry Canada regulations, this radio transmitter may only operate using an antenna of a type and maximum (or lesser) gain approved for the transmitter by Industry Canada. To reduce potential radio interference to other users, the antenna type and its gain should be so chosen that the equivalent isotropically radiated power (e.i.r.p.) is not more than that necessary for successful communication.

Conformément à la réglementation d'Industrie Canada, le présent émetteur radio peut

fonctionner avec une antenne d'un type et d'un gain maximal (ou inférieur) approuvé pour l'émetteur par Industrie Canada. Dans le but de réduire les risques de brouillage radioélectrique à l'intention des autres utilisateurs, il faut choisir le type d'antenne et son gain de sorte que la puissance isotrope rayonnée équivalente (p.i.r.e.) ne dépasse pas l'intensité nécessaire à l'établissement d'une communication satisfaisante.

If the distance from the product to the human body is greater than 20cm, the following warning is required (this requirement is not required for micro-power SRD devices

This equipment complies with FCC/IC RSS-102 radiation exposure limits set forth for an uncontrolled environment. This equipment should be installed and operated with minimum distance 20cm between the radiator & your body.

ce matériel est conforme aux limites de dose d'exposition aux rayonnements, FCC / CNR-102 énoncée dans un autre environnement.cette eqipment devrait être installé et exploité avec distance minimale de 20 entre le radiateur et votre corps.

The user manual for local area network devices shall contain instructions related to the restrictions mentioned in the above sections, namely that:

- (i) the device for operation in the band 5150-5250 MHz is only for indoor use to reduce the potential for harmful interference to co-channel mobile satellite systems;
- (ii) the maximum antenna gain permitted for devices in the bands 5250-5350 MHz and 5470-5725 MHz shall comply with the e.i.r.p. limit; and
- (iii) the maximum antenna gain permitted for devices in the band 5725-5825 MHz shall comply with the e.i.r.p. limits specified for point-to-point and non point-to-point operation as appropriate.
- (iv) the device does not be capable of transmitting in the band 5600-5650MHz.
- Les dispositifs fonctionnant dans la bande 5150-5250 MHz sont réservés uniquement pour une utilisation à l'intérieur afin de réduire les risques de brouillage préjudiciable aux systèmes de satellites mobiles utilisant les mêmes canaux.
- (ii) le gain d'antenne maximal autorisé pour les appareils dans les bandes 5250-5350 MHz et 5470-5725 MHz doivent respecter le pire limiter; et
- (iii) le gain d'antenne maximal autorisé pour les appareils dans la bande 5725-5825 MHz doivent respecter le pire limites spécifiées pour le point-à-point et l'exploitation non point à point, le cas échéant.
- (iv) le appareil n'est pas capable de transmettre dans la bande 5600-5650MHz.

Users should also be advised that high-power radars are allocated as primary users (i.e. priority users) of the bands 5250-5350 MHz and 5650-5850 MHz and that these radars could cause interference and/or damage to LE-LAN devices.

Les utilisateurs de radars de haute puissance sont désignés utilisateurs principaux (c.-à-d., qu'ils ont la priorité) pour les bandes 5250-5350 MHz et 5650-5850 MHz et que ces radars pourraient causer du brouillage et/ou des dommages aux dispositifs LAN-EL.

## frequency band

#### **FCC/IC Certification:**

1:2.412-2.462GHz 2:5.15-5.25GHz 3:5.25-5.35GHz

4:5.47-5.725GHz 5:5.725-5.875GHz

## CE Certification:

1:2.412-2.472GHz 2:5.15-5.25GHz 3:5.25-5.35GHz

4:5.47-5.725GHz 5:5.725-5.875GHz

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