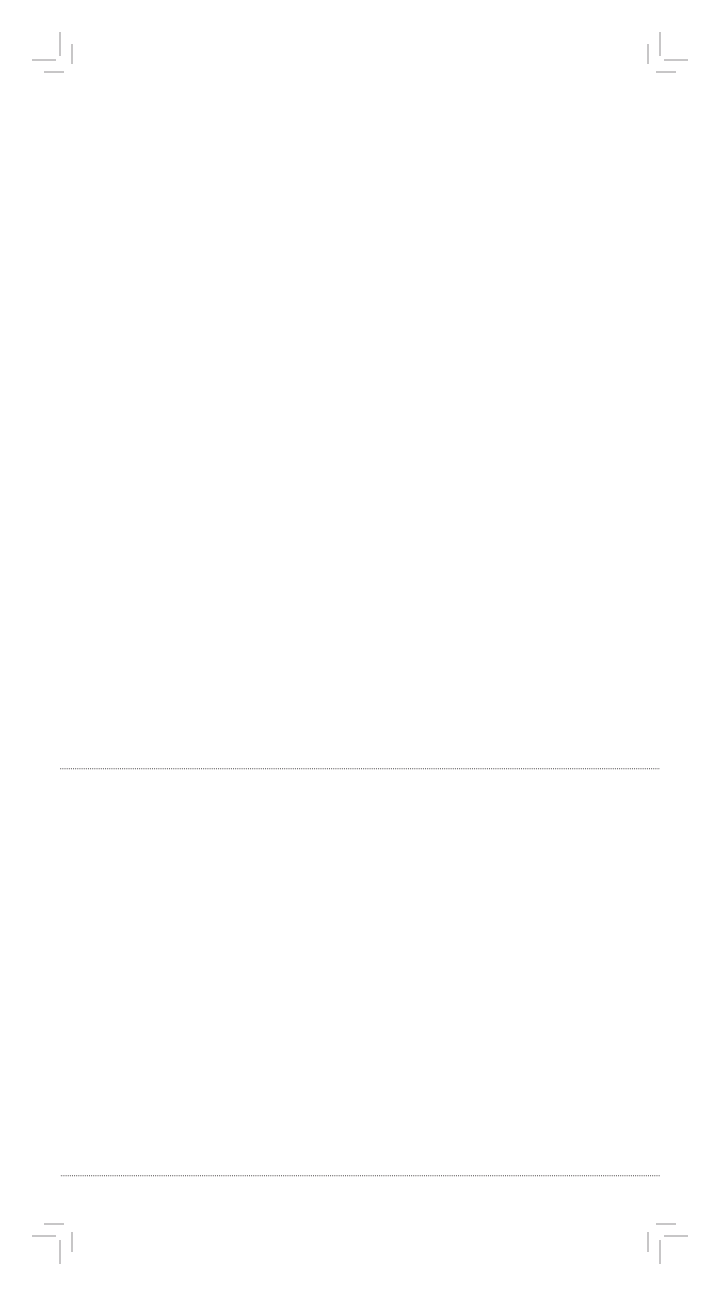
Back Cover Page Placement (See Seperate File for Artwork)

Front Cover Page Placement (See Seperate File for Artwork)



INSTRUCTIONS FOR USE

MODE D'EMPLOI

BEDIENUNGSANLEITUNG

ISTRUZIONI PER L'USO

INSTRUCCIONES DE USO

GEBRUIKSAANWIJZING

INSTRUÇÕES DE UTILIZAÇÃO

ΟΔΗΓΙΕΣ ΧΡΗΣΗΣ

BRUGSVEJLEDNING

BRUKSANVISNING

КÄҮТТÖОНЈЕ

MATTIOONS

BRUKSANVISNING

HASZNÁLATI UTASÍTÁS

NÁVOD K POUŽITÍ

INSTRUKCJA UŻYTKOWANIA

O. M.O. MILE

KULLANIM TALİMATLARI



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1 Overview

1.1 Indications For Use

The Site~Rite* 5 Ultrasound System with associated probe and accessories provide ultrasound guidance for placement of needles and catheters in vascular structures. Ultrasound guidance may occur intraoperatively or percutaneously. Ultrasound imaging of vascular structures, various organs and structures of the body may also be performed.

1.2 Site~Rite* 5 Ultrasound System and Components

The Site~Rite* 5 Ultrasound System is an easy to use and portable ultrasound scanner.

Site~Rite* 5 Ultrasound System and authorized accessories include:

- Site~Rite* 5 Ultrasound System scanner
- Site~Rite* 5 Ultrasound System vascular access probe
- Site~Rite* 5 Ultrasound System standard roll stand
- Site~Rite* needle guides
- Site~Rite* 5 Ultrasound System disposable probe covers
- Combination Site~Rite* 5 Ultrasound System A/C adapter and battery charger
- Site~Rite* 5 Ultrasound System roll stand mounted battery
- · Ultrasound gel
- · Ultrasound vessel phantom
- USB storage device with no external power connection (e.g. USB flash drive)

Contact your Bard Access Systems' Sales Representative or Customer Service at (800) 545-0890 to order.

1.3 Warnings, Precautions and Notes

Warnings

Warning:

Warning:	Do not rem	ove outer	protective	covers	from the	Site~Rite*	5 Ultrasound System scanner.
	1.1	To the		1 .	4.1.4	4.1	

Hazardous voltages exist at several points within the system.

Warning: Do not operate the Site~Rite* 5 Ultrasound System or the Site~Rite* 5 Ultrasound System A/C

adapter and battery charger in the presence of flammable anesthetics or gases. Explosion may

resul

Warning: Do not use for ophthalmic indications. Ophthalmic use may cause patient injury.

Warning: Misuse of the Site~Rite* 5 Ultrasound System may result in damage to the equipment or personal injury.

injury.

Warning: Use only the combination Site~Rite* 5 Ultrasound System A/C adapter and battery charger to charge Site~Rite* 5 Ultrasound System. Use of any other device to charge Site~Rite* 5 Ultrasound System battery packs may damage the battery packs and will void your warranty.

Only connect a Site~Rite* 5 Ultrasound System combination A/C adapter and battery charger

unpredictable operation, may damage the system and will void your warranty.

Warning: If a probe is damaged in any way, discontinue use immediately. Damage to the scanner may occur.

to the Site~Rite* 5 Ultrasound System. Use of any other A/C adapter may cause intermittent or

Warning: Avoid subjecting the probe to excessive mechanical shock. Damage to the probe may occur.Warning: Use only Bard Access Systems probes with this system. Use of unapproved probes may result in

patient injury or equipment damage.

Warning: When using Site~Rite* Needle Guides on the Site~Rite* 5 Ultrasound System probe, use only sterile plastic probe covers that are 1 mil (0.001 inch or 0.0254 mm) thick.

Warning: Do not allow liquid to enter the scanner, combination A/C adapter and battery charger, probe

connector or probe port. Damage to equipment may occur.

Warning: Do not attempt to sterilize the Site~Rite* 5 Ultrasound System scanner or probes with ethylene

oxide or heat sterilization methods. Damage to the equipment may occur.

Warning: Always properly dispose of dead battery packs in accordance with local regulations. Improper

disposal may present an environmental hazard.

Irning: Only qualified personnel should attempt to service this equipment. The Site~Rite* 5 Ultrasound

System contains static sensitive components and circuits. Failure to observe proper static control procedures may result in damage to the system.

Warning: The following actions void the warranty of the Site~Rite* 5 Ultrasound System and/or may result in patient injury or equipment damage:

- Opening or servicing the scanner or the probe housing by anyone other than Bard Access Systems authorized service personnel.
- Removal of system labels by anyone other than by Bard Access Systems authorized service personnel.



- Opening or servicing the battery pack or the combination A/C adapter and battery charger by anyone other than Bard Access Systems authorized service personnel.
- Connecting the Site~Rite* 5 Ultrasound System scanner to any power source other than the Site~Rite* 5 Ultrasound System combination A/C adapter and battery pack.
- Connecting the Site~Rite* 5 Ultrasound System scanner to any A/C adapter other than the one
 provided with the scanner.
- Connecting the Site~Rite* 5 Ultrasound System to any unauthorized accessory.
 Refer to Section 1.2.

Warning: Inspect A/C adapter and battery cord for damage. If any of the prongs are damaged, use battery power until replacement cord is obtained.

Warning: Verify that all accessories attached to the system comply to 60601 safety standards. Non-compliance may result in increased patient risk.

Warning: Use only IEC or ISO approved safety devices outside the patient environment. Failure to do so may damage the equipment.

Warning: Equipment that relies on basic insulation only should not be used with this system. Failure to comply could result in increased patient risk.

Warning: Do not overtighten screws when attaching to the VESA roll stand mount. Doing so may damage the scanner.

Warning: Use only screws provided in packaging. Ensure the unit is secure against the VESA roll stand mount. Failure to do so may cause the scanner to disconnect from the VESA roll stand mount.

Warning: Do not use the probe with high frequency surgical equipment. Doing so may damage the equipment.

Warning: Do not pull on probe cable. Doing so may cause the system to tip.

Precautions

Caution: The adverse biological effects of ultrasound on tissue appear to be threshold effects. When tissue

is repeatedly exposed to ultrasound, with intervals in between, there will likely be no cumulative biological effect. If, however, a certain threshold has been passed biological effects may occur. While the Site~Rite* 5 Ultrasound System acoustic output parameters fall well below all FDA thresholds for adverse biological effects, any given Ultrasound Procedure should be performed using the principle of ALARA (As Low As Reasonably Achievable). The licensed medical practitioner should limit the time of patient exposure to ultrasonic radiation using the principle of ALARA.

Caution: Federal (U.S.A.) law restricts this device to sale by or on the order of a physician.

Caution: Do not pull the cable to disconnect the probe connector from the scanner. Pulling the cable may damage the cable, cable connection or scanner.

Caution: Do not twist or bend the probe cable in excess of that required during normal use of the probe. Excessive twisting or bending of the cable may cause failure, intermittent or unpredictable operation.

Caution: When disinfecting the probe with a liquid disinfectant, do not soak the probe cable, cable bend relief, probe connector or probe buttons. Doing so may damage the probe.

Caution: Only apply commercially available ultrasonic couplant, which has been specifically formulated for use in medical applications, to the acoustic window (or face) of the probe.

Caution: Use water or rubbing alcohol and a soft cloth to remove couplant from the acoustic window (or face) of the probe. Failure to do so may scratch the acoustic window.

Caution: Do not to allow ultrasonic couplant to dry on the acoustic window (or face) of the probe. If the couplant should dry, use water or rubbing alcohol and a soft cloth to remove it. Never use a tool of any kind to remove dry couplant from the acoustic window (or face) of the probe.

Caution: Some commercially available probe covers contain latex. Natural rubber latex may cause allergic reactions. Refer to the US FDA alert titled: "Medical Alert: Allergic Reactions to Latex-Containing Medical Devices", issued March 29, 1991. Bard Access Systems distributes sterile probe covers and needle quide kits that do not contain latex.

Caution: Do not force the probe connector. Damage to the connector and system could result.

Caution: Always snap the needle guides on to the probe hook. Do not slide the needle guide on to the needle guide hook, as the sterile sheath may tear.

Caution: Do not subject the probe to excessive vibration. Vibration may dislodge sensitive components and cause intermittent or unpredictable operation.

Caution: Prior to each use please inspect the integrity of all power cords and connectors as well as the integrity of the unit itself. If any problems are found please discontinue use immediately and contact an authorized service representative. Use of a damaged power cord could damage the machine.

Caution: Unapproved extension cords should not be used with this system. Doing so may increase patient risk and/or damage the system.

Caution: During use, the AC connector needs to be easily accessible. In case of emergency remove the power cord as soon as possible.

Caution: To avoid unnecessary strain on the user, use the device in a comfortable manner.

Caution: Attach the power source in such a way as to prevent damage. Improper installation may damage power cords.

Caution: Inspect the probe prior to each use. If damage to the cable or transducer face is noted, do not use the probe. Damage to the system may occur.



inglish]

Caution: Hot water (in excess of 113° F or 45° C) may damage the probe.

Caution: Use only Bard Access Systems cleaning and disinfection procedures. Failure to do so may damage

the device

Notes

Note: When cleaning the system and components, it is important to remove all particles or other matter

from all surfaces and crevices.

Note: For 240 V applications use only center tapped 240 VAC single phase power.

2 Assembling the Site~Rite* 5 Ultrasound System

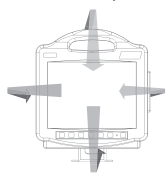
Unpack the Site~Rite* 5 Ultrasound System and verify the contents against the packing slip. Inspect all parts for damage.

2.1 Assembling the Roll Stand

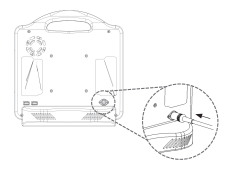
Refer to roll stand assembly instructions.

2.2 Adjusting the Scanner

Tilt and Swivel Adjustment



2.3 Attaching the Power Source and Charging the Battery



Warning: Use only the combination Site-Rite* 5 Ultrasound System A/C adapter and battery charger to charge Site-Rite* 5 Ultrasound System. Use of any other device to charge Site-Rite* 5 Ultrasound System battery packs may damage the battery packs and will void your warranty.

Warning: Only connect a Site-Rite* 5 Ultrasound System combination A/C adapter and battery charger to the Site-Rite* 5 Ultrasound System. Use of any other A/C adapter may cause intermittent or unpredictable operation, may damage the system and will void your warranty.

Warning: Always properly dispose of dead battery packs in accordance with local regulations. Improper disposal may present an environmental hazard.

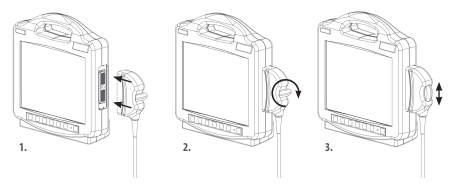
Warning: Inspect A/C adapter and battery cord for damage. If any of the prongs are damaged, use battery power until replacement cord is obtained.

When the Site~Rite* 5 Ultrasound System is attached to AC power, the battery charges automatically.

Caution: Attach power source in such a way as to prevent damage. Improper installation may damage power cords.



2.4 Connecting and Disconnecting Site~Rite* 5 Ultrasound System Probes



Warning: Use only Bard Access Systems probes with this system. Use of unapproved probes may result in patient injury or equipment damage.

Caution: Do not pull the cable to disconnect the probe connector from the scanner. Pulling the cable may damage the cable, cable connection or scanner.

Caution: Do not twist or bend the probe cable in excess of that required during normal use of the probe. Excessive twisting or bending of the cable may cause failure, intermittent or unpredictable operation.

2.5 Powering on the Site~Rite* 5 Ultrasound System

To power on the Site~Rite* 5 Ultrasound System

- 1. Verify that the probe is connected to the Site~Rite* 5 Ultrasound System scanner.
- 2. Depress and release the power button located on the front of the Site~Rite* 5 Ultrasound System scanner.
- 3. Wait approximately 20 seconds for the display screen to illuminate.
- 4. To power off the Site~Rite* 5 Ultrasound System, depress and release the power button again. The system is powered off when the display screen darkens.

2.6 Resetting the Site~Rite* 5 Ultrasound System

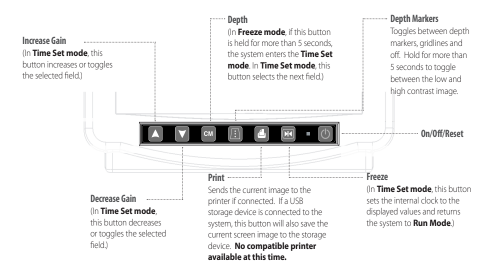
Should the scanner display ever appear blank or show an exclamation mark (!), the system may need to be reset.

To reset the system:

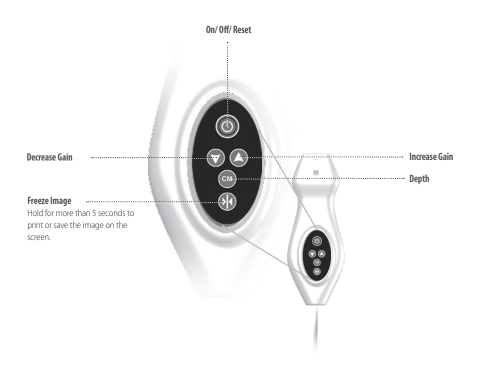
- 1. Depress and release the power button located on the front of the Site~Rite* 5 Ultrasound System scanner.
- 2. Wait approximately 60 seconds for the unit to reset itself.
- 3. Power the unit back on per instructions in section 2.5.

3 Site~Rite* 5 Ultrasound System Information

3.1 Front Panel Controls



3.2 Probe Controls





3.3 Display Screen Information

Probe Orientation Marker -----

Time and Date

The time is shown in 24 hour "00:00" format and the date is displayed in YYYY-MM-DD format.

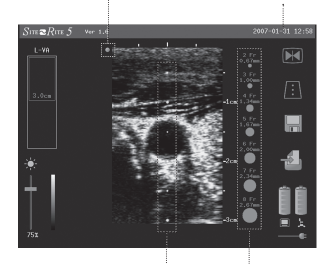
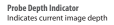


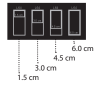
Image Depth Markers

When enabled, the depth markers are shown as green dots superimposed on the image at 1.0 cm pitch with smaller dash marks at 0.5 cm.

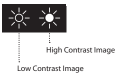
Catheter Size Icons

Displayed in proportion to the vessel image at a selected depth. Icons assist clinicians in determining the appropriate catheter size for the vessel being imaged.

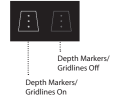




Contrast Indicator



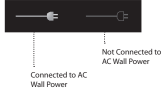
Depth Marker Indicator

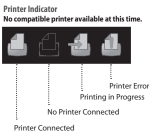


Run/Freeze Indicator

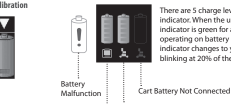


AC Power Indicator









There are 5 charge levels on each battery indicator. When the unit is recharging, the indicator is green for all 5 levels. When the unit is operating on battery power, the console battery indicator changes to yellow at 40% and red/blinking at 20% of the remaining battery life.

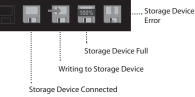


No Storage Device Con

nected



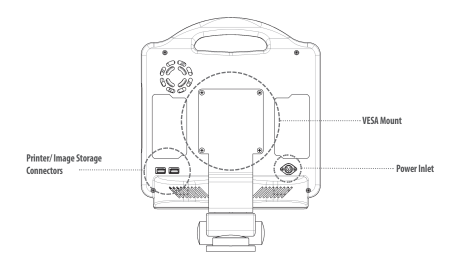
Storage Device Indicator



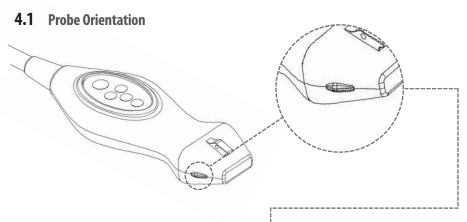


English]

3.4 Back Panel



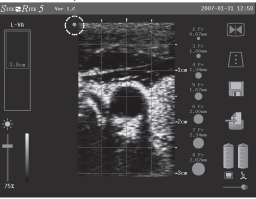
4 Using the Site~Rite* 5 Ultrasound System Probe



The Site~Rite* 5 Ultrasound System vascular access probe includes an orientation bump on the left side of the probe. This bump corresponds to the probe orientation mark located on the Site~Rite* 5 Ultrasound System display screen.

When using the Site~Rite* 5 Ultrasound System probe; hold the probe so that the side with the needle guide hook points away from the heart.

Warning: If the probe is damaged in any way, discontinue use immediately. Damage to the scanner may occur.



4.2 Draping the Probe for Sterile Use

When using the Site~Rite* 5 Ultrasound System probe in a sterile environment, the probe and part of the probe cable must be covered with a sterile, acoustically transparent plastic probe cover.

Warning: Use only sterile, legally marketed plastic probe covers that are 1 mil (0.001 inch or 0.0254 mm) thick.

Caution: Some commercially available probe covers contain latex. Natural rubber latex may cause allergic reactions. Refer to the US FDA alert titled: "Medical Alert: Allergic Reactions to Latex-Containing Medical Devices", issued March 29, 1991. Bard Access Systems distributes sterile probe covers and needle guide kits that do not contain latex.

To purchase sterile plastic probe covers, contact Bard Access Systems' Customer Service at:

Customer Service: (800) 545-0890

www.bardaccess.com



To drape the probe for sterile use:

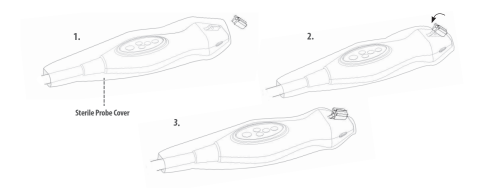
- 1. Place the probe in the side arm probe holder on the roll stand.
- 2. Apply a layer of non-sterile ultrasonic coupling gel on the acoustic window of the probe head.
- 3. Make sure that the probe cover is fully rolled up.
- 4. Place the probe cover over the probe head, being careful not to wipe off the coupling gel.
- 5. Cover the probe and cable with the probe cover.
- 6. Smooth the probe cover over the acoustic window of the probe head to remove any air bubbles or folds in the sheath.
- 7. Use the latex free poly-bands to hold the probe cover in place.
- 8. Apply a layer of sterile coupling gel to the covered acoustic window.

4.3 Ultrasound Guidance for Vascular Access

To use the Site~Rite* 5 Ultrasound System for vascular access:

- 1. Drape the probe for sterile use. Refer to instructions in Section 4.2.
- 2. Attach a needle guide to the probe.

Caution: Always snap the needle guides on to the probe hook. Do not slide the needle guide on to the needle guide hook, as the sterile sheath may tear.



- 3. Slide the appropriately sized needle, beveled edge facing the probe, into the channel on the guide.
- 4. Place the probe against the skin, perpendicular to the target vessel.
- 5. Hold the probe so that the side with the needle guide hook points away from the heart.
- 6. Center the dot markers on the target vessel.
- 7. While keeping the dot markers centered on the target vessel, slowly advance the needle while looking at the screen of the Site~Rite* 5 Ultrasound System scanner. When the needle approaches the target vessel, you should see the anterior wall indenting. Once puncture occurs, the vessel returns to normal shape.
- 8. Hold the needle, then gently rock the probe away from the needle for a smooth separation. The needle guide channel opens, and the needle smoothly disengages from the guide.

To purchase Needle Guides and sterile plastic Probe Covers, contact Bard Access Systems' Customer Service at:

Customer Service: (800) 545-0890 www.bardaccess.com

For instructions on the proper use of the Site~Rite* Needle Guides, refer to the Site~Rite* Needle Guide Kits & Ultrasound Probe Cover Kits Instructions for Use.

Warning: When using Site~Rite* Needle Guides on the Site~Rite* Ultrasound System probe, use only sterile, legally marketed plastic probe covers that are 1 mil (0.001 inch or 0.0254 mm) thick.

5 Settings

5.1 Date, Time and Image Size

To modify date, time and image size settings:

- Press the Freeze button (►I◄) to pause the system.
- 2. Press and hold the Depth button (**cm**) to enter the time/image size mode. The year field will then be highlighted. Release the Depth button.
- 3. Press the Depth button (cm) to select the desired date or time field to be modified.
- 4. Press the Gain buttons (∇ , \triangle) to change the selected date or time values.
- 5. After setting the date and time, press the Depth button (**cm**) to enter the image size fields. The current image size will be highlighted.
- 6. Press the Gain buttons (∇ , \triangle) to select the desired image size.
- 7. Press the Freeze button (> 1 d) to store the updated settings and resume normal operation.

5.2 Image Parameters

The Site~Rite* 5 Ultrasound System image may be changed from the factory default settings to a higher contrast image setting.

To change contrast settings, press and hold the Depth Marker Indicator (/:1) for five seconds. The image will switch between a low contrast and high contrast image setting.

5.3 Image Gridlines

Gridlines can be viewed over the ultrasound image to provide an effective tool to estimate vessel diameter. To toggle the image gridlines on and off, press the Depth Marker button (/:\).

5.4 Image Depth

The Site~Rite* 5 Ultrasound System image depth may be changed to image structures at different depths. Adjusting the depth also adjusts the focus of the ultrasound. Adjusting the depth to place the structure of interest at the appropriate focus will improve the ultrasound image.

Image Depth Setting (cm)	Focal Depth (cm)
1.5 cm	0.6 cm
3.0 cm	1.5 cm
4.5 cm	3.0 cm
6.0 cm	5.0 cm

- 1. To change the depth setting, press the depth button (cm) on the front panel control or probe.
- 2. Select the image depth that has the focal depth closest to that of the target structure.

5.5 Image Gain

The image gain can be adjusted to amplify the signal returned to the ultrasound machine.

Adjusting the gain effects the entire image. Increasing gain will amplify the signal from the target structure along with non-targeted structures.

- 1. To change the gain, press the gain up or down buttons ($\blacktriangle \blacktriangledown$) on the front panel or probe.
- 2. Select the gain that provides the best ultrasound image for the targeted structure.

6 Cleaning and Disinfection

6.1 Cleaning Procedures

To clean the scanner, probe and combination A/C adapter and battery charger:

- 1. Turn off the system.
- 2. Dampen a nonabrasive cloth with either warm water or rubbing alcohol.
- 3. Gently wipe the dampened cloth over exterior surfaces requiring cleaning.

6.2 Disinfection Procedures

The Site~Rite* 5 Ultrasound System probe may be liquid disinfected by soaking it in Cidex* plus 28 day solution. Follow the solution manufacturer's recommendations for soak time necessary to achieve the desired germicide level of activity.

Warning: Do not allow liquid to enter the scanner, combination A/C adapter and battery charger, probe connector or probe port. Damage to equipment may occur.

Warning: Do not attempt to sterilize the Site~Rite* 5 Ultrasound System scanner or probes with ethylene oxide or heat sterilization methods. Damage to the equipment may occur.

Caution: When disinfecting the probe with a liquid disinfectant, do not soak the probe cable, cable bend relief, probe connector or probe buttons. Doing so may damage the

Caution: Hot water (in excess of 113°F or 45°C) may damage the probe.

Caution: Use only Bard Access Systems cleaning and disinfection procedures. Failure to do so may damage the device.

Note: When cleaning the system and components, it is important to remove all particles or other matter from all surfaces and crevices.

For a list of disinfectants recommended for use on the Site~Rite* 5 Ultrasound System and probe, contact Bard Access Systems for the "Site~Rite* Ultrasound System Compatible Disinfectants" document.



7 Troubleshooting & Error Screens

[English]



Cause: Scanner does not recognize or identify a probe or probe not attached. **Solution:** Ensure that a Site-Rite 5* Ultrasound System probe is properly connected to the system.

stem alfunction

Cause: Scanner is not operating within normal parameters.

Solution: Discontinue use immediately. Return to authorized repair facility.

Display White Screen

Cause: Display malfunction.

Solution: Most display malfunctions can easily be corrected by resetting the system. To do so, power off the device, wait 60 seconds, then power the system back on. If the display malfunction is not resolved by resetting the system, discontinue use and return to authorized repair facility.

Battery Empty

Cause: Battery empty.

Solution: Connect system to AC outlet for operation and battery recharge.

Battery Malfunction

Cause: Battery malfunction.

Storage Device Indicator

Solution: Send system to authorized repair facility for battery replacement.

Cause: Storage device error.

Solution: Replace storage device.

Printer Indicator

Poor Image Quality

Solution: Check paper or service printer.
No compatible printer available at this time.

Cause: Incorrect Settings.

Solution: Refer to Section 5.

Cause: Scanner is not operating within normal parameters. Solution: Return to authorized repair facility.

8 Upgrading Software

The Site~Rite* 5 Ultrasound System allows the software to be upgraded through the USB connectors located on the rear of the scanner.

To install software:

- 1. Press the Freeze button (▶I◄) to pause the system.
- 2. Insert the USB drive containing the software upgrade into one of the USB connectors located on the rear of the Site~Rite* 5 Ultrasound System scanner.
- 3. Wait until the storage device icon is illuminated before proceeding.
- 4. Simultaneously press and hold the Gain buttons ($\blacktriangle \Psi$) until the configuration screen appears.
- 5. When the configuration screen appears, simultaneously press and hold the Depth button (cm) and Depth Marker button (/A) until the software installation process begins.

 $Note: The \ Site \sim Rite * 5 \ Ultrasound \ System \ will \ automatically \ upload \ the \ software \ from \ the \ attached \ USB \ drive.$

Note: The screen may appear blank and/or inactive during part of the software installation process.

- 6. When the system displays a message that the software update is successful, power off the Site~Rite* 5 Ultrasound System scanner.
- 7. Disconnect the USB drive.
- 8. Power on the Site~Rite* 5 Ultrasound System scanner.
- 9. Verify that the correct software version appears on the upper left hand corner of the screen.
- 10. Resume normal operation.

9 Calibrating the Rechargeable Battery

The Site~Rite* 5 Ultrasound System batteries may occasionally require calibration to ensure the battery power meter is accurate. The following icon indicates that the Site~Rite* 5 Ultrasound System rechargeable batteries require calibration.

To calibrate the Site~Rite* 5 Ultrasound System Rechargeable Batteries:

- 1. Disconnect the Site~Rite* 5 Ultrasound System from A/C power.
- Power on the Site~Rite* 5 Ultrasound System scanner and operate on battery power until the system powers off.
- 3. Connect the Site~Rite* 5 Ultrasound System to A/C power to recharge the batteries.

Note: At least five hours of charge time is recommended to fully charge the Site~Rite* 5 Ultrasound System batteries.

- 4. Disconnect the Site~Rite* 5 Ultrasound System from A/C power.
- 5. Power on the Site~Rite* 5 Ultrasound System and operate on battery power until the system powers off.

Note: The batteries are now calibrated.

6. Connect the Site~Rite* 5 Ultrasound System scanner to A/C power to recharge the batteries and continue normal use.







10 Warranty

The manufacturer, Bard Access Systems, warrants this product against defects in material and workmanship for a period of one year from the date of original purchase, and during such period agrees to repair, or at Bard Access Systems' discretion, replace any defective unit free of charge. The warranty on the repaired or replaced unit continues from the purchase date of the original unit. This warranty does not cover damages resulting from misuse, abuse, modification, or alteration of the Site~Rite* 5 Ultrasound System.

The following actions void the warranty of the Site~Rite* 5 Ultrasound System:

- Opening or servicing the scanner or the probe housing by anyone other than Bard Access Systems authorized service personnel.
- Removal of system labels by anyone other than by Bard Access Systems authorized service personnel.
- Opening or servicing the battery pack or the combination A/C adapter and battery charger by anyone other than Bard Access Systems authorized service personnel.
- Connecting the Site~Rite* 5 Ultrasound System scanner to any power source other than the Site~Rite* 5 Ultrasound System combination A/C adapter and battery pack.
- Connecting the Site~Rite* 5 Ultrasound System scanner to any A/C adapter other than the one provided with the scanner.
- Connecting the Site~Rite* 5 Ultrasound System to any unauthorized accessory.

TO THE EXTENT ALLOWABLE BY APPLICABLE LAW, THIS LIMITED PRODUCT WARRANTY IS IN LIEU OF ALL OTHER WARRANTIES, WHETHER EXPRESS OR IMPLIED, INCLUDING, BUT NOT LIMITED TO, ANY IMPLIED WARRANTY OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE.

IN NO EVENT WILL Bard Access Systems BE LIABLE TO YOU FOR ANY INDIRECT, INCIDENTAL OR CONSEQUENTIAL DAMAGES RESULTING FROM YOUR HANDLING OR USE OF THIS PRODUCT.

Some states/countries do not allow an exclusion of implied warranties, incidental or consequential damages. You may be entitled to additional remedies under the laws of your state/country.

11 Service and Repair

Warning:

Only qualified personnel should attempt to service this equipment. The Site~Rite* 5 Ultrasound System contains static sensitive components and circuits. Failure to observe proper static control procedures may result in damage to the system.

For servicing information or to return your Site~Rite* 5 Ultrasound System for repair, please contact Bard Access Systems Technical / Clinical Support at (800) 443-3385.

12 Technical Specifications

12.1 Operating and Storage Conditions

Operating Temperature: 59°F to 100°F (15°C to 38°C) **Storage Temperature:** 50°F to 100°F (10°C to 38°C)

 Operating Humidity:
 5% to 85% Relative Humidity (non-condensing)

 Storage Humidity (packaged):
 5% to 95% Relative Humidity (non-condensing)

 Storage Humidity (unpackaged):
 5% to 85% Relative Humidity (non-condensing)

12.2 Scanner Specifications

Dimensions: 12" W x 13" H x 5" D

Weight: 10 lbs.

Power Sources: AC adapter, Internal and External DC Battery Pack

Power Consumption:84 Watts MaximumMonitor Size:12.1" diagonal

IEC 60601-1: Class II, Type BF Applied Part, Continuous Operation, Internally Powered

Equipment, Not Category AP or APG Equipment, Not protected against

ingress of water.

12.3 Probe Acoustic Output Specifications

Description of Probe	Operating Mode	I _{spta.} X (X denotes statistically determined maximum)	FDA l-spta-3 Published Values	MI X (X denotes statisti- cally determined maximum)	FDA MI Published Values
			Peripheral Vessel < 720 m W/cm ²		Peripheral Vessel < 1.9
L-VA	В	49.003mW/cm ²	Cardiac < 430 mW/cm ²	.885	Cardiac < 1.9
			Fetal Imaging & Other** < 94 mW/cm ²		Fetal Imaging & Other** <



**Abdominal, Intraoperative, Pediatric, Small Organ (breast, thyroid, testes, etc.), Neonatal Cephalic, Adult Cephalic.
All measurements were conducted in accordance with the measurement procedures of the NEMA Standard Publications UD-2 and UD-3, and following the calibration procedures given in Appendices B, C, D and E of the 1985 FDA 510(k) Guide, and Part A, Sections Ill-IV, and Appendices A, B, C and D of the 1989 FDA 510(k) Guide, and the Track 1 and Track 3 reporting requirements of the September 30, 1997 Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers

[English]

Caution: The adverse biological effects of ultrasound on tissue appear to be threshold effects. When tissue is repeatedly exposed to ultrasound, with intervals in between, there will likely be no cumulative biological effect. If however a certain threshold has been passed biological effects may occur. While the Site~Rite* 5 Ultrasound System acoustic output parameters fall well below all FDA thresholds for adverse biological effects, any given Ultrasound Procedure should be performed using the principle of ALARA (As Low As Reasonably Achievable). The licensed medical practitioner should limit the time of patient exposure to ultrasonic radiation using the principle of ALARA.

12.4 Probe Specifications

L-VA: Linear Vascular Access Probe

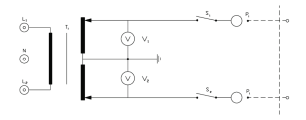
Frequency:5 - 10 MHzElevation Focus:1.8 cmMaximum Scan Depth:6.0 cmScan Width:1.9 cm

Lateral Foci:

Image Depth	Focal Depth
1.5 cm	0.6 cm
3.0 cm	1.5 cm
4.5 cm	3.0 cm
6.0 cm	5.0 cm

12.5 Power Supply Specifications

Note: For 240 V applications use only center tapped 240 VAC single phase power as shown below.



A/C Adapter Specifications

Input Voltage: 100-240 VAC, 50/60 Hz.
Input Current (Max): 2 Amps

Output Voltage: 15 VDC
Output Current (Max): 6 Amps

Internal Battery Pack Specifications

Battery Chemistry: Lithium Ion
Nominal Output Voltage: 10.8 VDC
Output Current (Max): 6 Amps
Output Current (Max): 6 Amps
Output Current (Max): 1.0 Hours
Charge Time (Full): 1.75 Hours

Site~Rite* 5 Ultrasound System Roll Stand Mounted Battery

Nominal Battery Output Voltage: 10.8 VDC Battery Output Current (Max): 6 Amps Battery Chemistry: Lithium Ion Output Power (Full Charge): 95 Wh System Run Time on Full Charge: 2.5 Hours Battery Charge Time (Full): 3 Hours





12.6 Standards Information

The Site~Rite* 5 Ultrasound System is designed to comply with applicable sections of the following International

- UL 60601-1: 2003, Medical Electrical Equipment, Part 1: General Requirements for Safety
 CAN/CSA C22.2 No. 601.1-M90, Medical Electrical Equipment Part 1: General Requirements for Safety
 IEC 60601-1: 1988, Medical Electrical Equipment Part 1: General Requirements for Safety
 EN 60601-1: 1990, Includes Amendments A1:1993, A11:1993, A12:1993, A2:1995 and A13:1996, Medical Electrical Equipment Part 1:
 General Requirements for Safety
 IEC 60601-1-1: 2000, Medical Electrical Equipment Part 1-1: General Requirements for Safety Collateral Standard: Safety Requirements for Medical Electrical Equipment Part 1-2: General Requirements for Safety Collateral Standard: Electromagnetic Compatibility Requirements and Texts
- Compatibility Requirements and Tests
 IEC 60601-1-2: 2001, Medical Electrical Equipment Part 1-2: General Requirements for the Safety of Ultrasonic Medical Diagnostic and Monitoring Equipment
 IEC 60601-1-4: 2000, Medical Electrical Equipment Part 1-4: General Requirements for the Safety of Ultrasonic Medical Diagnostic and Monitoring Equipment
 IEC 60601-1-4: 2000, Medical Electrical Equipment Part 1-4: General Requirements for Safety Collateral Standard: Programmable

- Electrical Medical Systems

 NEMA UD-2: 2004, Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment

 NEMA UD-3: 2004, Standard for Real-Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment
- EN 55011: 1998, Group 1, Class A Industrial, Scientific, and Medical (ISM) Radio-Frequency Equipment Radio Disturbance Characteristics-Limits and Methods of Measurement



13 Disposal Information

To return the Site~Rite* 5 Ultrasound System for end of life recycling, please contact your nearest Bard sales or distributor office in the country of purchase.

Warning: Always properly dispose of dead battery packs in accordance with local regulations. Improper disposal may present an enviromental hazard.

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Bard Limited

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An issued or revision date for these instructions is included for the users information. In the event two years have elapsed between this date and product use, the user should contact Bard Access Systems, Inc. to see if additional product information is available.

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