

Transvaginal Electronic Convex Probe UST-9124 Instruction Manual MN1-1162 Rev.21

Notes for operators and responsible maintenance personnel

★ Please read through this Instruction Manual carefully prior to use.

★ *Keep this Instruction Manual together with the ultrasound diagnostic instrument for any future reference.*

CE₀₁₂₃

Hitachi, Ltd.

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Introduction

This is an instruction for model UST-9124, an ultrasound probe.

Read the manual carefully before using the instrument. Take special note of the items in section 1, "Safety Precautions."

Keep this manual securely for future reference.

The CE mark on the probe indicates that this probe is valid when it is connected to equipment bearing the CE mark that is specified as available in section 2 of this document. Therefore, if a probe bearing the CE mark is connected to equipment that is specified as available but does not have a CE mark, part of this instruction manual may not apply.

Symbols used in this document

The terms below are used in the safety information provided to prevent hazards and injuries to the operator or patients. The severity of the hazard and injury that can occur when failing to observe the displayed safety information are indicated in four levels: "Danger," "Warning," "Caution," and "Note."

⚠ Danger

Indicates an imminently hazardous situation which, if not avoided, will result in death or serious injury to the operator or patient.

⚠ Warning

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

\triangle Caution

Indicates a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the operator or patient, or property damage only.

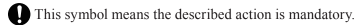
[▲] Note

Indicates a strong request concerning an item that must be observed in order to prevent damage or deterioration of the equipment and also to ensure that it is used efficiently.

The type of safety information is indicated by the symbols below.

This symbol means attention is required.

This symbol means that the described action is prohibited.



CONTENTS

1. Safety Precautions

1-1. Intended use	1
1-2. Usage precautions	1
1-2-1. Warnings and safety information	
1-2-2. Option usage precautions	
1-2-3. Cleaning, disinfection and sterilization precautions	
1-2-4. Labels	
2. Specifications and Parts name	
2-1. Principles of operation	
2-2. Specifications	
2-3. Performance	
2-4. Names of each parts	
2-5. Environmental conditions	
2-5-1. Operating environmental conditions	
2-5-2. Storage environmental conditions	
2-6. Classification of ME equipment	
3. Preparations for Use	
3-1. Startup check	
3-1-1. Visual check	
3-1-2. Verification of cleaning, disinfection and sterilization	
3-1-3. Verification of operation	
4. Usage	
4-1. Operation	
4-2. Mounting of rubber boot	
4-3. Removal of rubber boot	
4-4. Connecting to ultrasound diagnostic instrument	
4-5. Removing from the ultrasound diagnostic instrument	
4-6. Precautions when performing puncture operations	
4-7. Actions to be taken when an abnormal state is detected	
4-7-1. Ensuring safety of patients	
4-7-2. Handling the instrument	
5. Cleaning, disinfection and sterilization	
5-1. Precautions for cleaning, disinfection and sterilization	
5-2. Reprocessing instruction according to ISO 17664	

5-3. Point of use (Pre-cleaning)	29
5-4. Containment and transportation	29
5-5. Manual cleaning and disinfection	
5-5-1. Manual cleaning	31
5-5-2. Manual disinfection	32
5-5-3. Cable and connector	
5-6. Automated cleaning and disinfecting	
5-7. Applicable cleaners and disinfectants / Suppliers List	34
5-8. Drying	
5-9. Maintenance, inspection and testing	
5-10. Packaging	
5-11. Sterilization	37
5-11-1. Ethylene oxide (EtO) gas sterilization	37
5-11-2. STERRAD [®] sterilization	
5-11-3. Liquid sterilization (USA only)	
5-11-4. Autoclave sterilization	
5-12. Storage	
6. Storage	
6-1. Actions before storing the probe	41
6-2. Environmental conditions for storage	41
7. Moving and Transporting	
7-1. Moving and transporting	43
7-2. Preparing the probe and accessories for moving	43
7-3. Packing for transportation	43
7-4. Environmental conditions during transportation	43
8. Periodic Inspection	
8-1. Safety tests	45
8-2. Testing of measurement tolerances	46
8-2-1. Conducting tests	46
8-2-2. Result judgement	46
9. Configuration	
9-1. Standard configuration	47
9-2. Options	47
10. Disposal of the Device	49
This Instruction Manual contains the main body of 50pages and 5pages until the CONT	TENTS.

1. Safety Precautions

1-1. Intended use

This probe is intended for use by a doctor or other qualified operator for inserting into a human vagina and making ultrasonic observations of the uterus and surrounding organs.

\triangle Caution

Do not use this equipment for other than its intended purpose. Use for other purposes can cause burns or other injuries to the patient or operator.

1-2. Usage precautions

The terms below are used in the safety information provided to prevent hazards and injuries to the operator or patients. The severity of the hazard and injury that can occur when failing to observe the displayed safety information are indicated in four levels: "Danger," "Warning," "Caution," and "Note."

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⚠ Note

Indicates a strong request concerning an item that must be observed in order to prevent damage or deterioration of the equipment and also to ensure that it is used efficiently.

The type of safety information is indicated by the symbols below.



A

This symbol means attention is required.

This symbol means that the described action is prohibited.



This symbol means the described action is mandatory.

1-2-1. Warnings and safety information

⚠ Warning		
0	Follow the information in this manual and the documentation supplied with any equipment used together with this probe. Use that is not in accordance with the supplied documentation can result in a serious or moderate injury, equipment breakdown, or physical damage that impairs operation.	
0	Be sure to preparations for use. Use of the probe while failing to notice an abnormal condition can result in injury to the operator or patient. If any abnormalities are noted on the probe in the startup inspection, immediately stop using it and contact one of our offices and/or distributor's offices listed on the back cover. See section 3-1 "Startup inspection" for the startup inspection content and procedure.	
0	Do not use on the eyes. This probe is not intended for use on the eyes. The acoustic output can have an adverse effect on the eyes.	
\bigcirc	Do not attempt to disassemble, modify, or repair the probe. Electric shock or other unforeseen accidents could result. Contact one of our offices and/or distribu- tor's offices listed on the back cover to request repair.	
0	Clean, disinfect and sterilize before using the probe. Perform proper cleaning, disinfection and sterilization after use. Otherwise, there is a risk of infection. Note that the probe is not sterilized at the factory. Before using the probe first, be sure to clean, disinfect and sterilize it.	
0	Wear medical gloves during examination. Conducting examinations with the bare hands can expose the operator to a risk of infection.	
0	Dispose the probe used for patients with Creutzfeldt-Jakob disease. Otherwise, there is a risk of infection to the operator or patient. Our ultrasound probe is not compatible with any disinfection/sterilization method for Creutzfeldt-Jakob disease.	
0	When using ultrasound contrast agent, follow the supplied documentation. Unexpected accidents could result. Check the state of the patient and take appropriate precautions to avoid side effects.	
0	Do not use the probe fallen on to floor. Ohterwise, there is a risk of infection. Stop the operation, and perform the procedure in section 8-1 "Safety tests," section 5 "Cleaning, disinfection and sterilization" and section 3-1 "Startup inspection."	

▲ Caution		
0	Constantly check for anything abnormal about the patient's condition and probe. Continued use without noticing that an abnormal condition has occurred can result in an electric shock and injury to the operator or patient. If an abnormal condition occurs, immediately move the probe away from the patient and stop use of the probe.	
0	The probe is vulnerable to damage by impact. Therefore, handle it with care. There is a risk of damage to the probe when the probe is fallen or hit somewhere.	
\otimes	Do not use this probe with other equipment except for those specifically approved in the manual. Use with unapproved equipment can result in an electric shock, burn, or other injury to the patient or operator and damage to the probe and the other equipment.	

0	Scan for the minimum length of time necessary for the diagnosis and at the lowest suitable output. Overuse can adversely affect the internal tissues of the patient. For details about the acoustic output, please refer to the documentation supplied with the ultrasound diagnostic instrument.
0	Regularly perform maintenance inspection and safety tests of the ultrasound diagnostic instrument and probe. If you use equipment for a long period of time, it can reduce the performance, or cause smoke or fire. If anything unusual occurs, immediately stop using it and contact one of our offices and/or distributor's offices listed on the back cover.

Use, move, and transport the probe under the environmental conditions specified in this manual. Otherwise, it may be damaged.

0

See section 2-5 "Environmental conditions" and section 7-4 "Environmental conditions during transportation."

1-2-2. Option usage precautions

⚠ Warning		
•	Use by covering the rubber boot over the insertion portion. If the rubber boot is not used, residual pathogens on the probe could infect the patient. Also, the puncture guide tube could become loose during puncturing, resulting in puncturing of an unintended body part.	
0	Use Hitachi-approved rubber boots only. Use of an item lacking biocompatibility can cause an adverse reaction of the human body.	
0	Check that the rubber boot is sterilized. Use of an infective item could spread infection to the patient.	
\bigcirc	Do not reuse the rubber boot. Use of an infective item could spread infection to the patient.	
\bigcirc	Do not apply unsterilized acoustic medium to the outer surface of the rubber boot. Use of an acoustic medium that is contaminated by a pathogen can cause an infection on the patient.	
\bigcirc	Do not use on patients who may have an allergic reaction to latex products. Use of the rubber boot for these types of patients could result in anaphylactic shock. Ask the patient about allergy history beforehand.	

≜ Caution

Ω

Check the rubber boot for abnormalities before use.

Store the rubber boots in a cool, dry location not exposed to direct sunlight, and do not use rubber boots that have exceeded their expiration date (for items where the expiration date is not displayed, 2 years from the displayed sterilization date). Also do not use the rubber boot when there are severe discolorations, cracks or other visible effects.

Check that the acoustic medium has no air bubbles inside the rubber boot that is covering the probe. Air bubbles inside the rubber boot can result in misdiagnosis caused by overlooking or misinterpreting lesions due to poor image quality or improper rendering.

1-2-3. Cleaning, disinfection and sterilization precautions

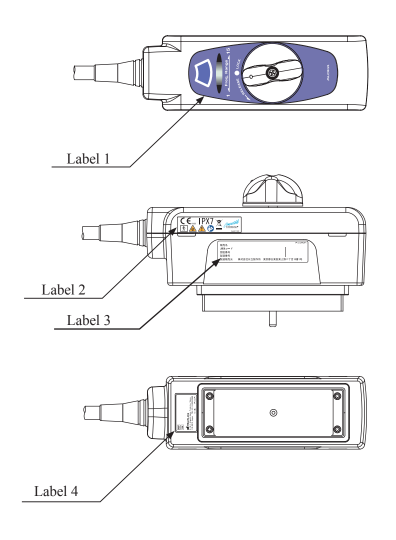
⚠ Warning		
0	Wear protective gloves and other protective gear during cleaning, disinfection and sterilization. Handling of the probe with your bare hands before disinfection or sterilization can result in an infection.	
0	After soaking in cleaning agents, thoroughly wash the probe with running water. Residual cleaning agents can cause an adverse reaction on the bodies of the operator or patient.	
0	After soaking in a disinfectant, throughly wash the equipment with deionized water. Leavings of the disinfectant can cause an adverse reaction on the bodies of the operator or patient.	
0	Perform aeration completely after gas disinfection and sterilization. Residual gas can cause an adverse reaction on the bodies of the operator or patient.	
\oslash	Do not clean, disinfect or sterilize using procedures other than those specified in this manual. Infection could result due to incomplete cleaning, disinfection and sterilization. It can also result in damage to the probe or reduced performance. The probe cannot withstand autoclave sterilization or boiling and other types of sterilization at temperatures exceeding 60°C (140°F).	
0	For details on the usage conditions of chemicals and sterilization procedures, refer to the documentation supplied with the respective chemical or sterilization equipment. Infection could result due to incomplete disinfection or sterilization. This could also cause deterioration of the probe.	

\triangle Caution

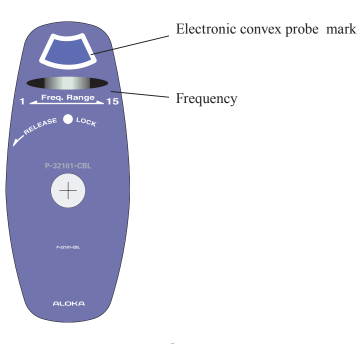
O Do not immerse the probe in any liquids beyond the range of IPX7 shown in section 2-2 "Specifications". Use when liquid has gotten inside the connector can result in a risk of electric shock to the operator or patient. If liquid gets inside the connector, immediately stop use and contact one of our offices and/ or distributor's offices listed on the back cover.

1-2-4. Labels

(1) Probe unit



Label 1



Label 2





This instrument complies with Directive 93/42/EEC relating to Medical Device and Directive 2011/65/EU relating to RoHS.

IPX7



Type BF applied part

See section 2-2, "Specifications".

IPX7 mark



Do not waste the instrument as general waste. Comply with a local regulation. See section 10.



STERRAD sterilization compatibility mark See section 5.



Safety warning sign



Biohazard See section 5.



Follow the instruction manual to operate this instrument. If not avoided, may result in injury, property damage, or the equipment trouble.

Label 3(Examples)

		P-12102-M3-1
販売名		
JMDNコード		
認証番号		
製造番号		
製造販売元	株式会社日立製作所	東京都台東区東上野二丁目16番1号

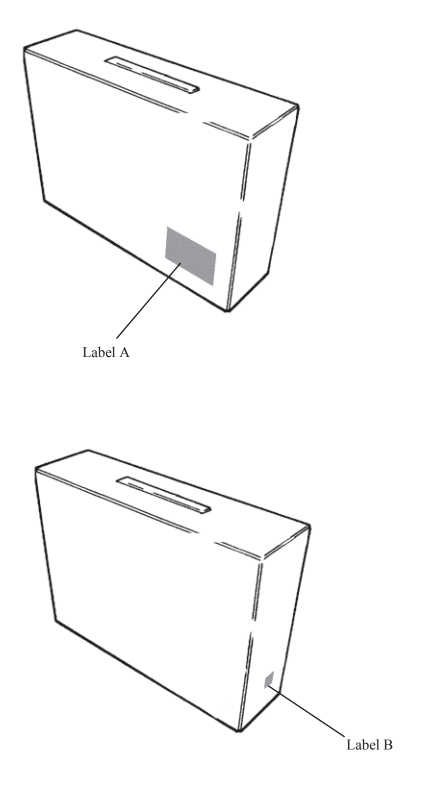
Label 4(Examples)

REF		
Hitachi, Lt 2–16–1, Higashi-U	eno, Taito-	
110-0015, Japan		3-6284-3668 P-1212V-1

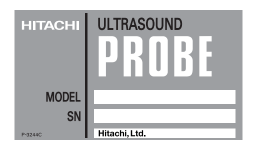
Manufacturer Model, Serial No.

This label may not be affixed depending on the destination.

Country of manufacture: JAPAN, CHINA or INDIA Model Serial No. Manufacturer Address Rx Only: By prescription only. U.S. Federal Law restricts this device to sale on order of a physician only. (2) Storage case



Label A(Examples)



Model Serial No.

Label B(Examples)





This instrument complies with Directive 93/42/EEC relating to Medical Device and Directive 2011/65/EU relating to RoHS.



DATE OF MANUFACTURE (in case of 2016-09)



MANUFACTURER



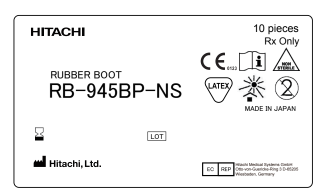
AUTHORISED REPRESENTATIVE IN EUROPEAN COMMUNITY

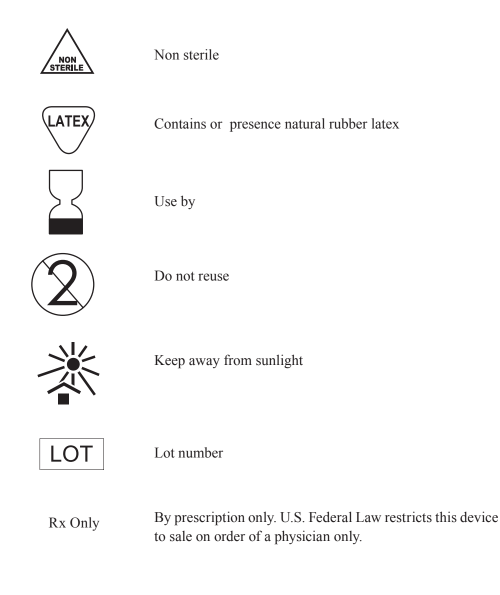
(3) Rubber boot

Label for 1 piece



Label for 10 pieces





2. Specifications and Parts name

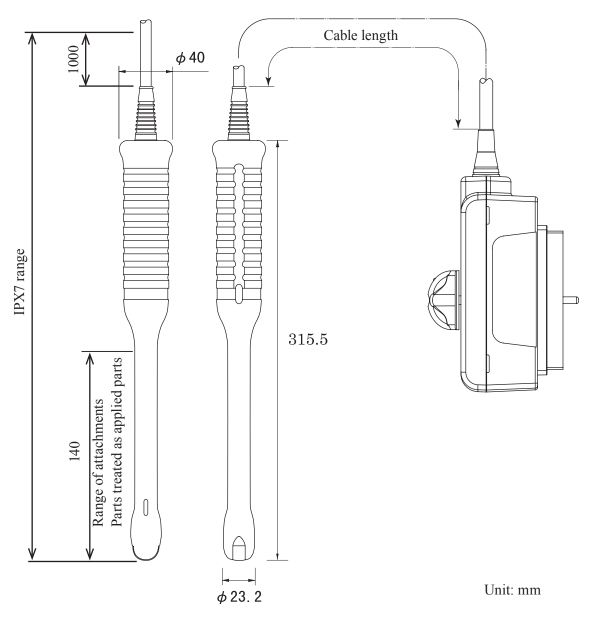
2-1. Principles of operation

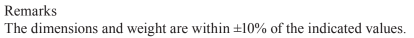
This probe and the ultrasound diagnostic instrument enable image diagnosis using ultrasonic waves. These instruments operate under the principles described below.

- (1) When an electric pulse signal is applied from the transmitter to the transducer of the probe, the transducer operates by converting electrical vibrations to mechanical vibration energy for emitting pulse-shaped ultrasonic waves into the body part contacting the transducer or into liquid or other medium.
- (2) The emitted ultrasonic waves are reflected by boundaries with different acoustic characteristics (acoustic impedance) within the body.
- (3) The transducer is also used to receive reflected ultrasonic waves. The transducer vibrates mechanically due to the received ultrasonic vibrations and uses an electro-mechanical conversion operation to convert the received mechanical vibrations to electric energy. The received echo is also converted to electric signals, and a brightness modulation operation is used to convert the electric pulses to shades of brightness for forming an image.

2-2. Specifications

Application regions:	Obstetric and gynecological areas
Form of application to patient:	Transvaginal
Connectable instruments:	SSD-4000, SSD-3500, SSD-1000, ProSound 6, ProSound α 6, F37, F31
Field of view:	180 °
Frequency:	3.0 to 8.5 MHz
Cable length:	2.5 m
Weight:	1,095 g
Service life:	Three years
Range of applied part	As shown in the figure below.
Parts treated as applied parts	As shown in the figure below
IPX7 range	As shown in the figure below.
External dimensions:	As shown in the figure below.

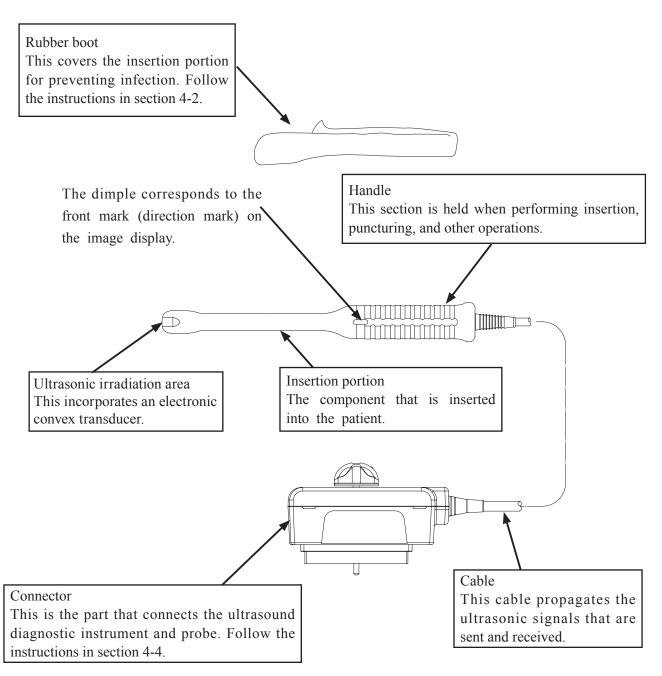


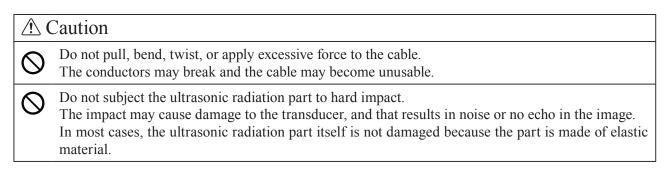


2-3. Performance

For measurement tolerances, operating tolerances, and other data, refer to the instruction manual for the ultrasound diagnostic instrument.

2-4. Name of each parts





2-5. Environmental conditions

Use and store the probe under the following conditions.

2-5-1. Operating environmental conditions

Ambient temperature:	10°C to 40°C
	50°F to 104°F
Relative humidity:	30% to 75%
Atmospheric pressure:	700 hPa to 1060 hPa
Altitude:	3,000 m or less

2-5-2. Storage environmental co	onditions
Ambient temperature:	−10°C to 50°C
	14°F to 122°F
Relative humidity:	10% to 90%
Atmospheric pressure:	700 hPa to 1060 hPa

	Avoid operating or storing the probe in the following locations.
	• Locations exposed to water or other liquids
	 Locations subject to adverse conditions such as air pressure, temperature, humidity, ventilation, direct sunlight, dust, or air containing salt, sulfur, or other corrosive substances Locations where chemical substances are stored or where gases are generated Storage in these locations can result in a breakdown or reduced performance.
)	Avoid rapid temperature change which may cause condensation. Avoid using in locations where condensation or water droplets can form. Condensation can occur when moving the probe from a cool location to a warm one. Use when condensation has occurred can result in a breakdown or reduced performance.

2-6. Classification of ME equipment

- Classification based on degree of protection against electric shock . Type BF applied Part
- Classification for protection against ingress of liquids IPX7 (Watertight equipment)

For the range of applied parts, parts treated as applied parts and the range of IPX7, see section 2-2.

3. Preparations for Use

3-1. Startup check

3-1-1. Visual check

Visually check the insertion portion, handle, cable, and connector. If any holes, indentations, abrasion, cracks, deformation, looseness, discoloration, or other abnormalities are found, do not use the equipment.

- 3-1-2. Verification of cleaning, disinfection and sterilization Verify that cleaning, disinfection and sterilization are conducted according to the intended use.
- 3-1-3. Verification of operation

Connect to the ultrasound diagnostic instrument by following the instructions in section 4-2, "Connecting to the ultrasound diagnostic instrument," and check that the selected probe match the covex display and the displayed frequency, and check the image for errors.

Remarks

For details on the displayed screens, see the documentation supplied with the ultrasound diagnostic instrument.

If the probe is operated in still air, brightness on the top of the image may be non uniform, but this does not affect the performance of the probe.

⚠ Warning

Be sure to preparations for use.

Using the probe without noticing an abnormal condition can result in injury to the operator or patient. If an inspection finds an abnormal condition in the probe, immediately stop use, and contact one of our offices and/or distributor's offices listed on the back cover.

A Caution

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Do not use the probe if the selected probe and image do not match the frequency. An incorrect acoustic output can result in burns or other injuries to the patient. Contact one of our offices and/or distributor's offices listed on the back cover.

4. Usage

4-1. Operation

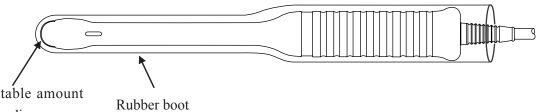
Check that the rubber boot is mounted, and insert the probe into the vaginal cavity. An image of the region of interest is displayed on the monitor of the ultrasound diagnostic instrument. For details on displaying and adjusting the screens, see the documentation supplied with the ultrasound diagnostic instrument.

	▲ Caution						
\bigcirc	Do not move the probe with excessive force. Pressing down with more force than necessary can cause injury to the patient.						
0	Scan for the minimum length of time necessary for the diagnosis and at the lowest suitable output. There is the possibility that the patient's internal tissues could be affected. For details about the acoustic output, please refer to the documentation supplied with the ultrasound diagnostic instrument.						
\bigcirc	Do not touch the connector terminal pin of the probe. The probe may deteriorate or be damaged due to electrostatic discharge.						
\bigcirc	Do not touch the electronic probe connecting socket of the diagnostic instrument and the patient at the same time. It can cause electric shock to the patient.						

4-2. Mounting of rubber boot

Apply a suitable amount of acoustic medium to the ultrasonic irradiation area of the probe, and then cover it with the rubber boot.

Remove the air bubbles or wrinkles formed on the ultrasonic irradiation area of the probe.



Apply	a suita	ble amou	11
of acou	istic me	dium.	

	Warning
0	Use by covering the rubber boot over the insertion portion. If the rubber boot is not used, residual pathogens on the probe could infect the patient. Also, the puncture guide tube could become loose during puncturing, resulting in puncturing of an unintended body part.
0	Use Hitachi-approved rubber boots only. Use of an item lacking biocompatibility can cause an adverse reaction of the human body.
0	Check that the rubber boot is sterilized. Use of an infective item could spread infection to the patient.
\bigcirc	Do not reuse the rubber boot. Use of an infective item could spread infection to the patient.
\bigcirc	Do not apply unsterilized acoustic medium to the outer surface of the rubber boot. Use of an acoustic medium that is contaminated by a pathogen can cause an infection on the patient.
\oslash	Do not use on patients who may have an allergic reaction to latex products. Use of the rubber boot for these types of patients could result in anaphylactic shock. Ask the patient about allergy history beforehand.

	Caution
0	Check the rubber boot for abnormalities before use. Store the rubber boots in a cool, dry location not exposed to direct sunlight, and do not use rubber boots that have exceeded their expiration date (for items where the expiration date is not displayed, 2 years from the displayed sterilization date). Also do not use the rubber boot when there are severe discolorations, cracks or other visible effects.
0	Check that the acoustic medium has no air bubbles inside the rubber boot that is covering the probe. Air bubbles inside the rubber boot can result in misdiagnosis caused by overlooking or

4-3. Removal of rubber boot

- 1. Wrap the rubber boots in tissue paper and remove it from the probe.
- 2. Dispose used tissue paper and rubber boots using infection prevention procedures based on the rules of your facility.

\triangle Caution

0

Before disposing the equipment, disinfect or take other infection-prevention measures. Disposal of the equipment without taking the proper preventative measures can lead to infection.

4-4. Connecting to ultrasound diagnostic instrument

The lock lever of the connector moves over the range shown in the figure at right.

Align the \circ mark with the LOCK or RELEASE position, and lock or release the probe connector.

Connect the probe to the electronic probe connecting socket of the diagnostic instrument (probe connector) by following the procedure below.

release gnostic o mark Lock lever

• Connection procedure

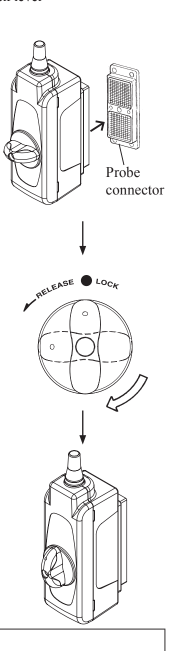
The probe is connected when in one of the following states.

- The power switch is set to OFF.
- The image displayed on the ultrasound diagnostic instrument is frozen.

Before inserting the probe into the probe connector, check that the connector pins are not bent.

- 1. Turn the connector lock lever to align the \circ mark on the lever with the RELEASE position.
- 2. Firmly insert the connector into the probe connector on the ultrasound diagnostic instrument.
- 3. Turn the lock lever clockwise by 1/4 turn until the \circ mark is aligned with the LOCK position.
- 4. Check that the connector is firmly inserted into the probe connector on the instrument.

This completes connection of the probe.



\triangle Caution

If there is resistance when trying to turn the lock lever when connecting the connector, do not forcibly try to connect it. Instead, correctly perform the steps for connecting the connector and firmly insert it into the probe connector.

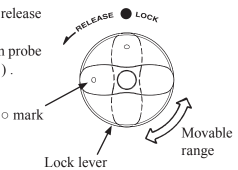
Forcibly turning the lever may damage the connector and the probe connector on the instrument.

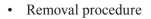
4-5. Removing from the ultrasound diagnostic instrument

The lock lever of the connector moves over the range shown in the figure at right.

Align the \circ mark with the LOCK or RELEASE position, and lock or release the probe connector.

Use the procedure below to remove the probe from the electronic scan probe connector of the ultrasound diagnostic instrument (probe connector).



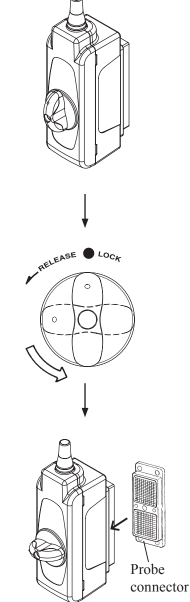


The probe is removed when in one of the following states.

- The power switch is set to OFF.
- The image displayed on the ultrasound diagnostic instrument is frozen.
- 1. Turn the connector lock lever to align the \circ mark on the lever with the RELEASE position.
- 2. Firmly grasp the connector unit, and pull it out from the probe connector of the ultrasound diagnostic instrument.

This completes the removal of the probe.

After use, perform cleaning, disinfection, and sterilization of the probe by following the procedure in section 5 "Cleaning, disinfection and sterilization." If the probe will not be used for an extended period of time, store it by following the instructions in section 6 "Storage."



4-6. Precautions when performing puncture operations

A Warning
Carefully read the usage precautions in the documentation supplied with the puncture guide tube. Be sure that the preparations for use are completed before using.
Puncturing must be performed by a skilled doctor. Improper puncturing can injure the patient. Puncturing operations must be performed by a doctor who fully understands the characteristics of ultrasound diagnostics and who is skilled and has a thorough knowledge of puncture operations under an ultrasound guide.
The puncture guide tube must be properly mounted on the probe during use. Puncturing with the puncture guide tube improperly mounted or the puncture guide tube uninstalled can result in the puncture guide tube coming off during puncturing or puncturing of an unintended body part, causing injury to the patient. For details about the puncture guide tube mounting procedure, see the documentation supplied with the puncture guide tube.
Be sure that the puncture guide tube and the needle are sterilized before use. Use of unsterilized items can cause an infection. For details concerning the puncture guide tube sterilization procedure, see the documentation supplied with the puncture guide tube.
For the acoustic medium, use sterilized physiological saline. Using an unsterilized ultrasound medium can cause an infection on the patient.
Use a compatible puncture needle size. Use of a puncture needle that is not a compatible size can result in the puncture guide tube coming off during puncturing or puncturing of an unintended body part, causing injury to the patient. For the compatible puncture needle sizes, see the documentation supplied with the puncture guide tube.
Always use a straight needle. Puncturing of an unintended body part can cause injury to the patient.
 During the puncture operation, display a suitable puncture guideline on the screen of the ultrasound diagnostic instrument. Puncturing of an unintended body part can cause injury to the patient. Display the puncture guideline on the screen referring to the documentation supplied with the ultrasound diagnostic instrument, to use it as an aid in determining the puncturing direction.
 Constantly check the safety in the needle insertion direction using the needle echo rendered by the ultrasonic wave. A bent puncturing needle can result in puncturing of an unintended body part and cause injury to the patient.
Check that no other organs lie in the puncture path. If another organ lies in the puncture path, an unintended body part can be punctured and cause injury to the patient. Before puncturing, carefully check the body parts, and constantly confirm the needle echo during the operation.
Do not try to forcibly perform operations. If excessive force is applied in a direction other than the insertion direction of the puncturing needle, the puncturing needle can come off the guideline, resulting in puncturing of an unintended body part, causing injury to the patient.

\triangle Caution

0

Handle the needle carefully to ensure that the probe or puncture guide tube is not damaged. Using a probe or puncture guide tube that has been damaged by a needle can result in an injury to the operator or patient.

4-7. Actions to be taken when an abnormal state is detected

4-7-1. Ensuring safety of patients

Immediately move the probe away from the patient and quit operation. Keep the patient in safe condition, and administer the required medical treatment.

4-7-2. Handling the instrument

Turn off the ultrasound diagnostic instrument, remove its plug from the AC socket, and sterilize if it is contaminated. For details, refer to the instruction manual for the ultrasound diagnostic instrument.

\triangle Caution

 \bigcirc

Do not use a probe where a problem has been found.

Using a probe in an abnormal state can cause injury to the patient. Contact one of our offices and/or distributor's offices listed on the back cover.

5. Cleaning, disinfection and sterilization

Applicable cleaning, disinfection and sterilization methods for each product are listed in the Table 1. The detail of each method is described in Chapter 5-2.

in II		· · · · · ·		Disinfection		Sterilization					
Model	Refer the corresponded items Chapter 5-3, 5-5, 5-6, 5-8 and 5	Manual	Automated *1	Manual	Automated *1	EtO	STERRAD [®]	Liquid *2	Autoclave	STERIS®	Waterproof cover (MP-2790)
UST-9124	A	Х		Х		Х	Х	Х			

Table 1 Applicable	cleaning.	disinfection	and s	terilization	methods
	····				

Accessories Washing brush Size(M)L-Ki-266, Size(L)L-Ki-265	В	x	x	x	x	x		x				
---	---	---	---	---	---	---	--	---	--	--	--	--

Note: X means "Applicable"

*1: Automated Need waterproof cover

*2: Liquid sterilization USA only

5-1. Precautions for cleaning, disinfection and sterilization

The following warnings and cautions must be observed when cleaning, disinfecting and sterilizing the probe and accessories.

<u>^</u>	Warning
0	Wear protective gloves and other protective gear during cleaning, disinfection and sterilization. Handling of the probe with your bare hands before sterilization can result in an infection.
0	After finishing soaking the probe in cleaning agents, thoroughly wash it with running water. Residual cleaning agents can cause an adverse reaction to the operator or the patient.
0	After chemical sterilization, thoroughly wash the probe with sterile water. Residual chemicals can cause an adverse reaction to the operator or patient. (USA only)
0	After disinfecting the probe, throughly wash the probe with deionized water. Leavings of the disinfectant can cause an adverse reaction on the bodies of the operator or patient. (EU only)
0	Perform full aeration after gas sterilization. Residual gas can cause an adverse reaction to the operator or patient.
\oslash	Do not clean or sterilize using procedures other than those specified in this manual. Failure to clean and sterilize the equipment can result in an infection. It can also result in damage to the probe or reduced performance. The probe is not compatible with autoclave sterilization or boiling and other types of sterilization at temperatures exceeding 60°C [140°F].
0	For details on the usage conditions of chemicals and sterilization procedures, refer to the documentation supplied with the respective chemical or sterilization equipment. Infection can be resulted due to incomplete sterilization. Wrong sterilization procedure could cause deterioration of the probe.

	Caution
\oslash	Do not immerse the probe into any liquid beyond the range of IPX7. The range is indicated in the section 2-2 "specification". If any liquid enters the connector, immediately stop using the probe and contact one of our offices and/or distributor's offices listed on the back cover. Liquid in the connector could cause electric shock to the operator or patient.
\bigcirc	Do not wipe the ultrasonic radiation part with alcohol. Alcohol could damage the part.
\bigcirc	Do not use organic solvent such as thinner for cleaning to prevent the probe from damage.
\bigcirc	Do not use hard or sharp objects to remove residue on the probe. Such objects may damage the probe.

Additional information:

The Instructions provided above have been validated by the medical device manufacturer as being CAPABLE of preparing a medical device for re-use. It remains the responsibility of the processor to ensure that the processing as actually performed using equipment, material and personnel in the processing facility achieve the desired result. This requires validation and routine monitoring of the process. Likewise any deviation by the processor from the instructions provided should be properly evaluated for effectiveness and potential adverse consequences.

5-2. Reprocessing instruction according to ISO 17664

Take care about clean circumstances before using the probe on the next patients. If processors reprocess this equipment, refer to these instructions.

WARNINGS	 The probe is delivered unsterile. Prior to the first use, reprocess the probe. Do not exceed 60 °C [140 °F]. Probe connector has no water resistance. 	
Limitations on reprocessing	The probe is not completely submergible (Do not immerse the probe into any liquid beyond the range of IPX7. The range is indicated in the section 2-2 "specification".) Parts which are not submergible can only be disinfected by wipe disinfection.	
Transportation before using	* I Sterile Subdiv Department (USSD) to operating room. Be careful that no dama	

The level of processing required depends on the type of equipment and its use.

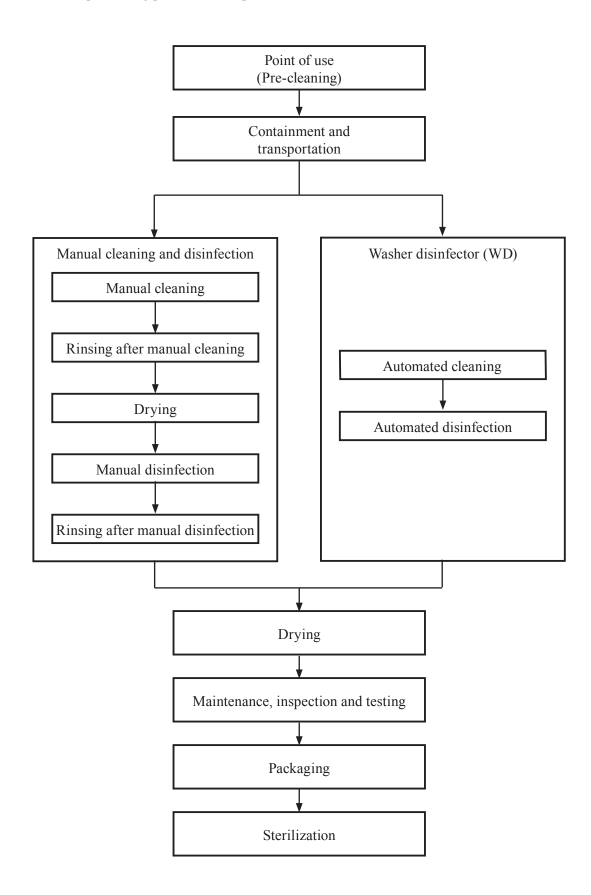
The CDC (Centers for Disease Control and Prevention) in the USA and the RKI (Robert Koch Institute) in Germany classify medical devices according to their use. For each classification, they specify the level of disinfection/sterilization processing that is required before use. Table 3 summarizes this information.

Table 2	
---------	--

Classification	Definition	Processing
Noncritical	Application part only contacts intact and uninjured skin	Cleaning ↓ Disinfection [in the USA, low-level disinfection]
Semicritical	Application part contacts mucosa (intracavitary application)	Cleaning ↓ Disinfection (Disinfectant with bactericidal, fungicidal and virucidal effect) [in the USA, high-level disinfection or sterilization]
Critical	Application part contacts intracorporeal tissue directly (intraoperative application)	Cleaning ↓ Disinfection ↓ Sterilization *1

*^{1.} When sterilization is not possible, the FDA in the USA recognize that disinfection (in the USA, highlevel disinfection) and the use of a sterile gel and sterile transducer cover, as described in the instructions provided with the transducer cover, is an accepted method of infection control for probe.

Flowchart of reprocessing process of this probe and accessories is as follows:



NOTE: Only the accessories are compatible with automated reprocessing according to the flowchart above.

5-3. Point of use (Pre-cleaning)

In the operating room after use of the probe

A). Probe

- 1) Remove any accessories from the probe like puncture guide tube and rubber boot.
- 2) Flush patient's blood or fluid by tap water directly after use until the surface looks visually clean.
- 3) Wipe the whole surface of the probe by gauze pad and remove superficial visible impurities until the surface looks visually clean.
- B). Accessories
 - 1) Clean the accessory of all patient's blood or fluid with running tap water until the surface of the accessory looks visually clean.
 - 2) Wipe the whole surface of the accessory by gauze pad.

5-4. Containment and transportation

Putting the contaminated equipment into exclusive shock and damage proof container for transportation is recommended. It is recommended that instruments are reprocessed as soon as possible and not later than 4 hours after usage.

\triangle Caution

Do not immerse the probe in any liquids beyond the range of IPX7 shown in section 2-2 "Specifications". Use when liquid has gotten inside the connector can result in a risk of electric shock to the operator or patient. If liquid gets inside the connector, immediately stop use and contact one of our offices and/ or distributor's offices listed on the back cover.

5-5. Manual cleaning and disinfection

Prepare following items before manual cleaning and disinfection.

A). Probe

- 1) Detergent: ENZOL[®]/Cidezyme[®] (Johnson & Johnson, #2258) or another cleaning agent with approved material compatibility for this medical device.
- 2) Disinfectant: Cidex[®] OPA (Johnson & Johnson, # 20391) or another disinfectant with approved material compatibility for this medical device.
- 3) 2 tanks, 1 for cleaning and 1 for disinfection optional: 1 additional tank for rinsing with deionized/ tap water. (sufficient size for immersion of the submergible part of the probe at full length)
- 4) Soft, fluff free cloth or single use towel
- 5) Personal protective equipment (gloves, water repellent protective skirt, face protection mask or protective glasses see also instructions of the manufacturer for the detergent and the disinfectant)
- B). Accessories
 - 1) Detergent: ENZOL[®]/Cidezyme[®] (Johnson & Johnson, #2258) or another cleaning agent with approved material compatibility for this medical device.
 - 2) Disinfectant: Cidex[®] OPA (Johnson & Johnson, # 20391) or another disinfectant with approved material compatibility for this medical device.
 - 3) 2 tanks, 1 for cleaning and 1 for disinfection optional: 1 additional tank for rinsing with deionized/ tap water. (sufficient size for immersion of the accessory at full length)
 - 4) Soft, fluff free cloth or single use towel
 - 5) Personal protective equipment (gloves, water repellent protective skirt, face protection mask or protective glasses, see also instructions of the manufacturer for the detergent and the disinfectant)

5-5-1. Manual cleaning

A). Probe

- The temperature of the detergent solution should be between 15-30 °C [59-86 °F], concentration is 1.6%. Please note the minimum contact time of the detergent in the manufacturer's instruction. If a differing detergent is used, please also consider the approved material compatibility for this probe.
- 2) Immerge the submergible part of the probe (see figure) without connector into the detergent.
- 3) Wipe the submergible part of the probe under the surface of the detergent solution with a single-use, fluff free soft cloth to remove all visible soil. Be sure that all grooves of the probe are implemented during the cleaning process. If necessary use an appropriate cleaning brush for this purpose.
- 4) Wipe the non-submergible parts of the probe with a soft cloth dipped with a detergent.
- 5) Rinse the submergible part of the probe with running tap water for 1 minute.
- 6) Alternatively to step 5 suspend the submergible part of the probe in a tray filled with deionized water/tap water for 5 min.
- 7) Visually check the outer surface of the probe for cleanness. If necessary, use magnifying glass for visually check. If there is still soil visible, repeat all above steps.

B). Accessories

- The temperature of the detergent solution should be between 15-30 °C[59-86°F], concentration is 1.6%. Please note the minimum contact time of the detergent in the manufacturer's instruction. If a differing detergent is used, please also consider the approved material compatibility for this medical device.
- 2) Immerge the accessory into the detergent.
- 3) Wipe the accessory under the surface of the detergent solution with a single-use, fluff free soft cloth to remove all visible soil. Be sure that all grooves of the accessory are implemented during the cleaning process.
- 4) Rinse the accessory with running tap water for 1 minute.

(Alternatively, immerse the accessory in a tray filled with deionized water/tap water for 5 min.)

5) Visually check the outer surface of the accessory for cleanness. If necessary, use magnifying glass for visually check. If there is still soil visible, repeat all above steps.

5-5-2. Manual disinfection

- A). Probe
 - Before immersing the equipment, it is recommended to test the concentration of disinfectant solution before each usage. The solution Cidex[®] OPA is ready for use and does not need to be diluted. Test strips to verify that the appropriate concentration of Cidex[®] OPA is correct are available by manufacturer. Test strips will indicate a concentration above the Minimum Effective Concentration (MEC). Temperature of disinfectant solution should be minimum 20 °C[68 °F]. The minimum contact time is 5 minutes. If a differing disinfectant is used follow the manufacturer's instructions. Please also consider the material compatibility for the medical device.
 - 2) Wipe the non-submergible parts of the probe with a soft and fluff free cloth with disinfectant.
 - 3) Immerge the submergible part of the probe (see figure) into the disinfectant. Set a clock to insure the recommended contact time is observed.
 - 4) Rinse the submergible part of the probe with running deionized water for 1 minute.
 - 5) Alternatively to step 4 suspend the submergible part of the probe in a tray filled with deionized water for 5 min.
 - 6) Visually check the outer surface of the probe for that there are no leavings of the disinfectant. If necessary, repeat the rinsing.

\triangle Caution

Do not wipe the ultrasonic radiation part with alcohol. Alcohol could damage the part.

- B). Accessories
 - Before immersing the equipment, it is recommended to test the concentration of disinfectant solution before each usage. The solution Cidex[®] OPA is ready for use and does not need to be diluted. Test strips to verify that the appropriate concentration of Cidex[®] OPA is correct are available by manufacturer. Test strips will indicate a concentration above the Minimum Effective Concentration (MEC). Temperature of disinfectant solution should be minimum 20 °C[68 °F]. The minimum contact time is 5 minutes. If a differing disinfectant is used follow the manufacturer's instructions. Please also consider the material compatibility for the medical device.
 - 2) Immerge the accessory into the disinfectant. Set a clock to insure the recommended contact time is observed.
 - 3) Rinse the accessory with running deionized water for 1 minute.
 - 4) Alternatively to step 4 suspend the accessory in a tray filled with deionized water for 5 min.
 - 5) Visually check the outer surface of the accessory for that there are no leavings of the disinfectant. If necessary, repeat the rinsing.

A Warning

After finishing soaking the probe in the cleaning agent or disinfectant, thoroughly rinse it with running water (after cleaning) and deionized water (after disinfection). Residual agent can cause an adverse reaction to the operator or patient.

5-5-3. Cable and connector

Wipe the cable in 20 cm intervals with gauze dipped in ethyl alcohol or water, and dry it after wiping. Clean the connector with gauze dipped in ethyl alcohol, and dry it after cleaning.

Clean the other parts of the probe which must not be soaked in liquid in the same manner as the connector.

[▲] Note

If the entire length of the cable is wiped at once, a part of the cable may be wrinkled. If this occurs, pull the wrinkled part in the opposite direction to smooth it.

5-6. Automated cleaning and disinfecting

A). Probe

🗥 Warning

The probe cannot withstand automated cleaning and disinfecting.

B). Accessories

The following items must be provided prior to automated cleaning and disinfection:

a) Washer disinfector: according to ISO 15883 with chemo-thermal program (temperature: max 60 °C[140 °F])

- b) Detergent: Korsolex Endo-Cleaner (Bode Chemie; # 972 020)
- c) Disinfectant: Korsolex Endo-Disinfectant (Bode Chemie; # 972 030)
- 1) The parameters of the cleaning and disinfection of the device are as follows:

Program step	Water (401)	Dosage (ml/l)	Temp. (°C)/(°F)	time (min)
Pre-Rinse	Cold water			5
Cleaning	Deionized water	5 (0.5%)	50/122	10
Rinse	Deionized water			1
Disinfection	Deionized water	10 (1%)	55/131	5
Rinse	Deionized water			1
Rinse	Deionized water		55/131	1
Drying			55/131	15

2) After closing the door, start the chemo-thermal program.

3) Open the door after the end of the program.

4) Check whether the accessory is dry. If not, proceed as described under drying.

5-7. Applicable cleaners and disinfectants / Suppliers List

The applicable chemical solutions are listed below.

General name	Trade name	Manufacturer
Enzyme cleaning agent	ENZOL [®] /Cidezyme [®] Practical liquid 0.8V/V%	ADVANCED STERILIZATION PRODUCTS [®] A Johnson & Johnson company Division of Ethicon, Inc.
Alkylpolyalkylenglykolether	Korsolex [®] Endo-Cleaner	BODE Chemie GmbH

General name	Trade name	Manufacturer	
Glutaral	CIDEX [®] Solution 2.4%	ADVANCED STERILIZATION PRODUCTS [®]	
Ortho-phthalaldehyde	CIDEX [®] OPA Solution 0.55%	A Johnson & Johnson company Division of Ethicon, Inc.	
Glutaral	Cidex plus [®]		
Glutaral	STERIHYDE [®] * Practical liquid 2W/V%	Maruishi Pharmaceutical Co., Ltd.	
Benzethonium chloride	Hyamine [®] * Practical liquid 0.1W/V%	DAIICHI SANKYO Co., Ltd.	
Didecyl dimethylammonium chloride	Cleanisept [®] Wipes * Solution 7.5%	Dr. Schumacher GmbH	
Dimethyl-dioctyl- ammonium-chloride	Gigasept [®] AF forte * Solution 2.0%	Schülke & Mayr	
Glutaral	Korsolex extra *	BODE Chemie GmbH	
Glutaral	Korsolex Endo- Disinfectant	BODE Chemie GmbH	

Note: * indicates that the marked disinfectant is not applicable in Canada.

General name	Trade name	Manufacturer	
Hydrogen peroxide	PERASAFE ^{TM*} Practical liquid 1.62W/V%	ANTEC INTERNATIONAL	
Peracetic acid	Acecide® * Solution 6%	Saraya Co., Ltd.	
Glutaraldehyde	WAVICIDE [®] -01 * Solution 2.65%	Medical Chemical Corporation	
Glutaraldehyde	STERANIOS * Solution 2.0%	Laboratoires ANIOS	
Glutaral	Cidex plus® Solution 3.4%	ADVANCED STERILIZATION PRODUCTS ⁴ A Johnson & Johnson company Division of Ethicon, Inc.	

High-level disinfection

Note: * indicates that the marked disinfectant is not applicable in Canada.

⚠ Warning

0

After disinfection, thoroughly rinse the probe with deionized water. Residual disinfectant can cause an adverse reaction to the operator or patient.

5-8. Drying

A). Probe

- 1) Wipe the probe with single use, fluff free wipe or towel for removing moisture on the surface of the equipment.
- 2) If using drying heater for medical equipment, the temperature limit is a maximum of 60 °C [140 °F]. Dry until no visible moisture is left.
- 3) If using natural drying, temperature range should be between 15-30°C[59-86°F] for a minimum time of 4 hours.

B). Accessories

- 1) Wipe the accessory with a single-use, fluff-free wipe or towel to remove moisture from the surface of the accessory.
- 2) Dry the accessory naturally in an ambient temperature between 15-30°C[59-86°F] for a minimum of 4 hours. Alternatively the accessory can be dried using a drying heater at a temperature of less than 60°C[140°F]

5-9. Maintenance, inspection and testing

Confirm following items

- 1) the function of mechanical moving parts
- 2) the image performance when the probe is connected to the scanner
- 3) there are no abnormal exterior damages such as cracks on the surface of the equipment
- 4) Safety tests (See section 8-1)

5-10. Packaging

Store the disinfected probe in a dustproof environment until next application. Before sterilization it is necessary to pack all parts in a pouch suitable for sterilization, or in a tray with wrap according to ISO 11607-1 and ISO 11607-2 "Packaging for terminally sterilized devices" and ISO/TS 16775 "Packaging for terminally sterilized medical devices - Guidance on the application of ISO 11607-1 and ISO 11607-2" or the local hospital procedure. Follow the pouch manufacturer's specifications or the local regulations for how to pack and seal the pouches. Check the sealing seam after heat sealing for any defects. In case of processing mistakes or defects the package has to be opened again and the device has to be packed and sealed again.

5-11. Sterilization

See "Table 1. Applicable cleaning, disinfection and sterilization methods" for available sterilization methods Follow the instructions of the sterilizer manufacturer regarding usage, temperature and sterilization-time etc. Handling and maximum input to chamber of sterilizer should be according to operation manual of the sterilizer.

5-11-1. Ethylene oxide (EtO) gas sterilization

Sterile conditions of applicable sterilization methods are as follows.

Regarding the operation of the sterilizer, refer to the documentation supplied with the sterilizer.

Perform sterilization in the following conditions:		
Gas Type: 10% EO/ 90% HCFC		
Temperature:	50 - 60°C	
	122 - 140°F	
Exposure Time:	More than 120 minutes	
Pressurization:	162 - 200kPa	
Depressurization:	13 - 8kPa	
Relative humidity:	40 - 90%	
Aeration is minimum	12 hours	

RB-945BP-NS

Perform sterilization in the following conditions:		
Gas Type:	30% EO/ 70% CO2	
Temperature:	45 - 55°C	
	113 - 131°F	
Exposure Time:	More than 240 minutes	
Pressurization:	89 - 108kPa	
Depressurization:	13 - 8kPa	
Relative humidity:	30 - 60%	
Aeration is minimum	12 hours	

⚠ Warning

Perform full aeration after gas sterilization. Residual gas can cause an adverse reaction to the operator or patient.

5-11-2. STERRAD[®] sterilization

Sterile conditions of applicable sterilization methods are as follows. The applicable gas is listed below.

General name	Trade name	Manufacturer
Hydrogen peroxide (58% density)	STERRAD [®] Sterilization system (STERRAD [®] 50, 100S, 200, NX or 100NX)	ADVANCED STERILIZATION PRODUCTS [®] A Johnson & Johnson company Division of Ethicon, Inc.

Regarding the operation of the sterilizer, refer to the documentation supplied with the sterilizer.

Remark:

Some discoloration of the probe may occur, but this does not affect performance or safety.

	Caution
\otimes	Do not sterilize the probe using the STERRAD system if the probe is not compatible with the STERRAD system. STERRAD compatibility is shown by the STERRAD label on the connector. Perform STERRAD sterilization only for STERRAD compatible probes, otherwise it can cause damage or deterioration to the probe.
0	Do not put the probe directly into the sterilization pouch*. Otherwise the pouch sticks to the cable and results in damage to the cable. Completely wrap the entire probe (including the probe tip, cable and connector) with sterilization wraps* before putting it into the sterilization pouch*. *: A Johnson & Johnson company Division of Ethicon, Inc. product

5-11-3. Liquid sterilization (USA only)

• Applicable chemical solution for sterilization The applicable sterilants are listed below.

General name	Trade name	Manufacturer	
Hydrogen peroxide	PERASAFE ^{®*} Practical liquid 1.62W/V%	ANTEC INTERNATIONAL	
Peracetic acid	Acecide [®] * Solution 6%	Saraya Co., Ltd.	
Glutaraldehyde	WAVICIDE®-01 * Solution 2.65%	Medical Chemical Corporation	
Glutaraldehyde	STERANIOS * Solution 2.0%	Laboratoires ANIOS	
Glutaral	Cidex plus® Solution 3.4%	ADVANCED STERILIZATION PRODUCTS [®] A Johnson & Johnson company Division of Ethicon, Inc.	

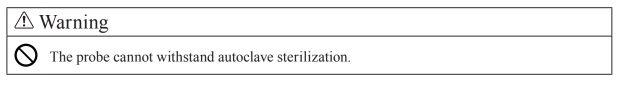
Note: * indicates that the marked sterilant is not applicable in Canada.

⚠ Warning

After chemical sterilization, thoroughly rinse the probe with sterile water. Residual sterilant can cause an adverse reaction to the operator or patient.

5-11-4. Autoclave sterilization

A). Probe



B). Accessories

A Warning	
The washing brush cannot withstand	autoclave sterilization.

5-12. Storage

For details about the storage environmental conditions, see section 2-5-2 "Storage environmental conditions".

6. Storage

6-1. Actions before storing the probe

When the probe will not be used for an extended period of time, perform the procedures described in section 5 "Cleaning, disinfection and sterilization," and then store it in its storage case.

6-2. Environmental conditions for storage

For details about the storage environmental conditions, see section 2-5-2 "Storage environmental conditions."

7. Moving and Transporting

7-1. Moving and transporting

In this section, *moving* refers to "carrying of the probe within a facility," and *transporting* refers to "transferring using a vehicle or sending the probe for repairs."

7-2. Preparing the probe and accessories for moving

Store in the storage case after performing the procedure in section 5 "Cleaning, disinfection and sterilization."

7-3. Packing for transportation

Store in the storage case after performing the procedure in section 5 "Cleaning, disinfection and sterilization," and then put the storage case in a cardboard box for additional protection.

7-4. Environmental conditions during transportation

Ambient temperature:	-10°C to 50°C	
	14°F to 122°F	
Relative humidity:	10% to 90%	
Atmospheric pressure:	700 hPa to 1060 hPa	

⚠ Note

The probe is a precision equipment and is vulnerable to physical impact. Protect it by packing it properly for transportation.

Contact one of our offices and/or distributor's offices listed on the back cover when transporting the probe.

8. Periodic Inspection

8-1. Safety tests

The safety tests should be conducted at least once a year by a qualified technician. The test record should be stored for future reference.

Remarks 1

Qualified technician: personnel for conducting safety tests of medical electrical equipment. If the user requires an appropriate qualified technician, service personnel trained by us can conduct a test at the user's expense. Contact one of our office written on the back cover.

Remarks 2

Make a copy of the Safety Inspection Data Sheet provided in the instruction manual of the ultrasound diagnostic instrument. Use the sheet as a test record.

Procedure for periodic safety tests and judgment

(1) Test of patient leakage current from the patient connection to earth

Using the measuring instruments which usable to the requirement of IEC 60601-1 :2005, conduct the test as shown in Fig. 15 of IEC 60601-1 :2005.

Soak the insertion portion and handle in saline solution and measure the leakage current between the applied part and earth.

Do not soak probes in saline solution beyond the "IPX7 range" provided in section 2-2.

(2) Test of patient leakage current caused by an external voltage on the patient connection of an F-type applied part.

Using the measuring instruments which usable to the requirement of IEC 60601-1 :2005, conduct the test as shown in Fig. 16 of IEC 60601-1 :2005.

Soak the insertion portion and handle in saline solution and measure the leakage current between the applied part and earth.

Do not soak probes in saline solution beyond the "IPX7 range" provided in section 2-2.

Item	Normal condition	Single fault condition
(1) Patient leakage current from the patient con- nection to earth		
DC	10 µA or less	50 µA or less
AC	100 μ A or less	500 µA or less
(2) Patient leakage current caused by an external voltage on the patient connection of an F-type applied part		5000 µA or less

Table. Standard Values for Periodic Safety tests (Extract from IEC 60601-1 :2005)

⚠ Warning

Ω

Perform a safety tests at least once a year, and keep a record of the inspection results. Failure to notice an abnormal condition while using the probe can result in injury to the operator or patient. If an inspection finds an abnormal condition in the probe, immediately stop use, and contact one of our offices and/or distributor's offices listed on the back cover.

8-2. Testing of measurement tolerances

Perform the measurements specified below using an ultrasonic phantom* at least once per year. The test record should be stored for future reference.

- Sensitivity
- Resolution

Remarks

Make a copy of the Measurement accuracy inspection data sheet provided in the instruction manual for the ultrasound diagnostic instrument. Use the sheet as a test record.

* The ultrasonic phantom is made of a substance which is similar to human tissue in terms of its response to ultrasonic waves.

Regions with different textures and targets spaced at preset intervals are embedded in the phantom. Some phantoms contain a mechanism for Doppler measurement. The phantom is used to check the performance of the probe and ultrasonic diagnostic instrument, as well as to adjust the image settings.

8-2-1. Conducting tests

Some types of ultrasonic phantoms have targets with narrow gaps between them for confirming the resolution.

This enables you to check the level of detail that images can be viewed on the display. For phantoms with no targets, the resolution determines the fineness of the displayed textures. The sensitivity can be determined by examining the luminance of ultrasonic images. Other factors that affect the resolution include the type of connected probe, gain, focus, and recording instrument. The specific testing conditions must be recorded in detail to enable proper comparison at the next inspection.

8-2-2. Result judgement

Compare the currently-obtained value with the value recorded at the last test. If there is a significant difference between the two values, the current value is considered to be abnormal.

It is important to note that the resolution varies depending on the type of ultrasonic phantom, and phantoms generally deteriorate over time.

A Caution

Do not use a probe or ultrasound diagnostic instrument where a problem has been found. This can result in an incorrect diagnosis. Contact one of our offices and/or distributor's offices listed on the back cover.

9. Configuration

9-1. Standard configuration

Probe	UST-9124	1 set
Storage case	CB-UST1-P1 or CB-UST5-P1	1 set
Instruction manual	MN1-1162	1 copy

9-2. Options

• Puncture guide tube

The following puncture guide tubes are provided for various puncture needle sizes.

MP-2748-G14	(for 14G needles)
MP-2748-G16	(for 16G needles)
MP-2748-G17	(for 17G needles)
MP-2748-G18	(for 18G needles)
MP-2748-G19	(for 19G needles)
MP-2748-G22	(for 22G needles)

- Washing brush (L) L-Ki-265 (for MP-2748-G14) Washing brush (M) L-Ki-266 (for MP-2748-G16 to G19)
- Puncture guide tube storage tray MP-2718

• Puncture guide tube set MP-2748-SET	
Puncture guide tube	MP-2748-G18
Washing brush (M)	L-Ki-266
Puncture guide tube storage tray	MP-2718

• Rubber boot RB-945BP-NS (unsterilized)

10. Disposal of the Device

Recycle or dispose this equipment properly in compliance with the Waste Management and Public Cleansing Law.



V

Before disposing the equipment, disinfect or take other infection-prevention measures. Disposal of the equipment without taking the proper preventative measures can lead to infection.

Waste Electrical and Electronic Equipment (WEEE) Directive

The illustration on the right is required by the EU WEEE Directive to appear on all electrical and electronic equipment.

For proper disposal of this product in an EU nation, contact an EU office or agency and observe appropriate local and national regulations and laws.



Manufacturer

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Contact

+81-3-6284-3668

http://www.hitachi.com/businesses/healthcare/index.html

Overseas Offices:

EC REP

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