

KOSMOS on iOS

User Guide



Table of Contents

CHAPTER 1	Getting Started 1
	What's new in this release? 1
	Packaged contents 1
	Intended users 1
	Intended use/indications for use 2
	Clinical applications and modes of operation for Kosmos on iOS 2
	<i>Clinical applications</i> 2
	<i>User guide</i> 3
	<i>Symbols in this user guide</i> 4
	<i>User guide conventions</i> 4
	Contraindications 4
	General warnings and cautions 5
	EchoNous customer support 6
CHAPTER 2	Kosmos Overview 7
	What is Kosmos? 7
	Kosmos clinical applications 8
	Training 9
	Kosmos classifications 9
	Patient environment 9
CHAPTER 3	Using Kosmos 10
	System overview 10
	<i>Device requirements</i> 10
	Kosmos hardware 11
	<i>Kosmos Torso-One</i> 12
	<i>Kosmos Lexsa</i> 12
	<i>Kosmos Link</i> 12
	Getting started 13
	<i>Downloading the EchoNous Kosmos ultrasound app</i> 13
	<i>Connecting Kosmos probes</i> 13
	Kosmos Link for iOS 13
	<i>Setting up Kosmos Link</i> 14
	<i>How to remove tablet from Kosmos Link</i> 14
	<i>Charging Kosmos Link</i> 14
	General interaction 15
	<i>Home screen: Kosmos Torso-One</i> 15
	<i>Home screen: Kosmos Lexsa</i> 15
	<i>Learn</i> 16
	Settings 16
	<i>Imaging preferences</i> 16

P008465-002 Rev C

March 2025

*Apple licenses the "iOS" trademark from Cisco

© 2015 to 2025 EchoNous, Inc., or its affiliates. All rights reserved.

	About	17
	DICOM	17
	Managing MWL	20
	USB export	21
	Report settings	21
	Wireless Networking Functions	21
	Connection specifications	21
CHAPTER 4	Performing an Exam	22
	Overview	22
	Primary exam workflows	22
	Exam workflows	23
	Standard workflow	23
	Quick workflow	24
	AI-assisted EF workflow	25
	Kosmos Bladder AI Workflow	26
	Managing exams	26
	Starting an exam	26
	Searching for an exam	27
	Deleting exams	27
	Acquiring images and clips	27
	Completing exams	28
	Managing patient data	28
	Adding a new patient	28
	Accessing patient information using MWL	28
	Searching for a patient	28
	Changing to another patient	29
	Editing a patient record	29
	Merging two patient records	29
	Deleting patient records	30
	Organ presets	30
	Imaging modes and features	30
	2D/B-mode	31
	M-mode	31
	Color Doppler	32
	Color Power Doppler	33
	Pulsed-Wave Doppler	34
	Tissue Doppler Imaging	35
	Continuous-Wave Doppler	36
	Auto Preset	37
	Auto Doppler	38
	Image mode controls	38
	Using the Kosmos AI-assisted EF workflow	39
	Calculating EF with the AI-assisted EF workflow	40
	Reviewing/adjusting the ED/ES frames and LV contours	41
	Recommendations for acquiring optimal A4C and A2C clips for accurate EF calculations	42
	Error conditions and system notifications for Kosmos AI-assisted EF workflow	43
	Kosmos cardiac measurements	44

Kosmos AI FAST	46
Using Kosmos AI for FAST Exam	46
Kosmos Bladder AI	47
Accessing the Bladder Preset	48
Pre Void Volume	48
Post Void Volume	52
Kosmos vascular calculations	56

CHAPTER 5	Reviewing an Exam	57
	Starting an exam review	57
	Annotating images and clips	57
	Navigating to the Edit Image screen	57
	Annotation tools	58
	Measuring with the caliper tool	58
	Deleting annotations	59
	Managing images and clips	59
	Filtering images and clips	59
	Selecting images and clips	59
	Trimming and saving images and clips	60
	Deleting images and clips	60
	Reviewing and editing a report	60
	Opening a report	60
	Editing a report	60
	Exporting images and clips to a USB drive	61
	Completing an exam review	62
	Archiving an exam to a PACS server	63
	Deleting an exam	63

CHAPTER 6	Kosmos Probes	65
	Kosmos probe sheaths	65
	Ultrasound transmission gels	65
	Kosmos probe storage	66
	Daily storage	66
	Storage for transport	66
	Transducer element check	66

CHAPTER 7	Kosmos Maintenance	67
	Cleaning and disinfecting	67
	General cautions	67
	Tablet	67
	Kosmos Link	68
	Kosmos probes	69
	Guidelines for AR (automated reproprocessors)	73
	Recycling and disposal	73
	Troubleshooting	74
	Preventive inspection, maintenance, and calibration	74

CHAPTER 8	Safety	75
	Electrical safety	75

	References	75
	Labeling symbols	76
	Contact information	80
	Biological safety	83
	ALARA education program	83
	Kosmos Torso-One acoustic output tables	86
	Kosmos Lexsa maximum acoustic output summary	93
	Measurement accuracy	99
	Control effects	101
	Related references	101
	Transducer surface temperature rise	101
	Ergonomics	102
	Basic safety	102
	Electromagnetic compatibility	103
	Electromagnetic emissions	104
	Electromagnetic immunity	105
	Separation distances	107
	Standards	107
	HIPAA	107
	DICOM	107
CHAPTER 9	Specifications	108
	System specifications	108
	Environmental operating and storage conditions for Kosmos probes, Kosmos Link and compatible tablets	108
	<i>Kosmos probes and tablets: operating, charging, transport, and storage condition ranges</i>	108
	<i>Kosmos Link: operating, charging, transport, and storage condition ranges</i>	109
	Mode of operation	109
	Kosmos Link electrical specifications	109
	Output	109
	Internal batteries	109
	Power supply	109
CHAPTER 10	IT Network	110
	Wireless networking	110
	Functions	110
	Security	110
	Network for connecting the device	110
	IT network failure recovery measures	111
	Glossary	113
APPENDIX A	Auto EF Clinical Performance and Non-Clinical Testing	117
	Auto EF Clinical Performance Testing	117
	Study Design	117
	Results	117
	Software Verification and Validation Testing	118
	Algorithm Testing	118

APPENDIX B	AI FAST Clinical Performance and Non-Clinical Testing	119
	AI FAST Clinical Performance Testing	119
	Study Design	119
	Results	119
	Software Verification and Validation Testing	120
	Algorithm Testing	120
APPENDIX C	Kosmos Bladder AI Clinical Performance and Non-Clinical Testing	121
	Kosmos Bladder AI Clinical Performance Testing	121
	Study Design	121
	Results	121
	Software Verification and Validation Testing	122
	Algorithm Testing	122

Getting Started

What's new in this release?

New features and changes for v4.0 software for Kosmos® iOS include:

- Kosmos Bladder AI Workflow

	For electronic versions of the user guides, please visit the EchoNous website at echonous.com/product/resources .
	Not all features are available in all markets. Please check with your local representative for availability in your region.

Packaged contents

For Kosmos on iOS users, the Kosmos box contains the following items:

- Kosmos Torso-One and/or Kosmos Lexsa
- Kosmos probe connector guard (optional accessory) with installation instructions
- Kosmos Platform Quick Start Guide
- Kosmos Welcome Letter
- Chemical Compatibility
- USB flash drive containing:
 - Kosmos on iOS User Guide
 - Kosmos AI Station 2 User Guide

Intended users

Kosmos is intended to be used by qualified and trained healthcare professionals that are legally authorized by law in the country, state, or other local municipality in which they practice to use the device. The list of the potential users includes but is not limited to (based on title/geographical location): Medical specialists, primary care physicians, point-of-care (POC) users, sonographers, medical healthcare technicians, nurses, nurse practitioners, physician assistants, and medical students.

Users may or may not be working under supervision or authority of a physician.

Intended use/indications for use

	To help ensure the diagnostic quality of the images obtained, all patient images must be obtained by qualified and trained healthcare professionals.
---	--

Kosmos is intended to be used by qualified and trained healthcare professionals in the clinical assessment of the cardiac and pulmonary systems and the abdomen by acquiring, processing, displaying, measuring, and storing ultrasound images.

Kosmos is intended to be used in clinical care and medical education settings on adult and pediatric patient populations.

Kosmos includes the AI-assisted automated ejection fraction software, known as Auto EF, which is used to process previously acquired transthoracic cardiac ultrasound images, to store images, and to manipulate and make measurements on images using the Kosmos. Auto EF provides automated estimation of left ventricular ejection fraction. This measurement can be used to assist the clinician in a cardiac evaluation. Auto EF is indicated for use on adult patients only in healthcare facilities.

Kosmos includes the Auto Anatomical Structure Labeling and View Identification, also referred to as AI FAST software, which is intended for use only by qualified and trained medical professionals for automatic real-time detection and labeling of anatomical structures during image acquisition during cardiac, thoracic/lung, or abdominal ultrasound imaging. This feature is only indicated for use on adult patients in healthcare facilities.

Kosmos includes the Bladder Biplane Caliper Volume software, also referred to as Kosmos Bladder AI, which is intended for use only by qualified and trained medical professionals to obtain ultrasound imaging of the bladder that is used to automatically determine bladder volume.

The device is non-invasive, reusable, and intended to be used on one patient at a time.

With respect to its ultrasound imaging capabilities, Kosmos is a general purpose diagnostic ultrasound system used in the following clinical applications and modes of operation:

Clinical applications and modes of operation for Kosmos on iOS

Clinical applications

- **Torso-One:** Cardiac, Thoracic/Lung, Abdominal, and Bladder

- **Lexsa:** Lung, Vascular/Peripheral Vascular, Musculoskeletal, Nerve and Image Guidance for Needle/Catheter Placement (includes needle/catheter placement, fluid drainage, and nerve block)
- **Modes of operation:** B-mode, M-mode, Color Doppler, Color Power Doppler, combined modes of B+M and B+CD, PW Doppler, CW Doppler, TDI, Harmonic Imaging, and Kosmos Bladder AI

TABLE 1-1. Modes of operation and purchasable features for Kosmos on iOS

Mode	Torso-One iOS	Lexsa iOS	Purchasable Features
B-mode	x	x	
M-mode	x	x	
B + CD (Color Doppler)	x	x	
Harmonic Imaging	x		
AI-assisted EF Workflow	x		x
PW Doppler	x	x	x
TDI	x		x
CW Doppler	x		x
AI FAST	x		x
Color Power Doppler		x	
Auto Preset	x		x
Auto Doppler	x		x
Kosmos Bladder AI	x		x

User guide

This user guide is intended to assist you with the safe and effective operation of Kosmos. Before attempting to operate Kosmos, read this user guide and strictly observe all the included warnings and cautions. Also, pay special attention to the information in the chapter titled **Safety**.

Not all software versions include all the features described in this guide. Reference the software version on your device.

This user guide and any digital media (and the information they contain) is the proprietary and confidential information of EchoNous and may not be reproduced, copied in whole or in part, adapted, modified, disclosed to others, or disseminated without the prior written permission of the EchoNous legal department. This document or digital media is intended to be used by customers and is licensed to them as part of their EchoNous purchase. Use of this document or digital media by unauthorized persons is strictly prohibited. This user guide is also available through the EchoNous website, or a paper copy may be supplied on request.

Symbols in this user guide

	Warning	A warning describes precautions to prevent injury or loss of life.
	Caution	A caution describes precautions to prevent damage to the device.
	Note	A note provides supplemental information.

User guide conventions

The following style conventions are used in this guide:

- Numbered and lettered steps must be performed in a specific order.
- Bulleted items are lists in no specific order.
- Kosmos touch screen icons and buttons are indicated in bold, such as **SCAN**.
- The word:
 - **Tap** refers to touching the screen quickly with your finger
 - **Double tap** refers to touching the screen two times in quick succession with your finger
 - **Drag** refers to touching the screen with your finger and then moving your finger across the screen
 - **Swipe** refers to moving your finger across the screen quickly
 - **Pinch** refers to moving two fingers in a pinch motion or pinch release motion across the screen
 - **Check** refers to tapping a check box to enable the associated function
 - **Clear** refers to tapping a check box to disable the associated function
 - **Select** refers to tapping a menu item from a menu list
- Links to other sections within the guide appear bold and colored, such as the cross reference, see **“Imaging modes and features” on page 30**.

Contraindications

Kosmos is designed for transcutaneous scanning and trans thoracic echocardiography only.

Kosmos is not intended for ophthalmic use or any use causing the acoustic beam to pass through the eye.

	Show care when scanning near a wound to avoid damaging or further injuring the affected area.
	Federal (United States) law restricts this device to sale by or on the order of a physician.

General warnings and cautions

	System users are responsible for image quality and diagnosis
	Kosmos is not MRI compatible and should not be used in an MRI suite.
	Kosmos is not for use in oxygen-rich environments.
	To avoid the risk of electrical shock, do not allow any part of Kosmos (except for Kosmos probe lens) to touch the patient.
	To avoid the risk of electrical shock or injury, do not open the tablet or Kosmos probe enclosures for any reason. All internal adjustments and replacements (such as the battery) need to be made by a qualified Kosmos technician.
	To avoid the risk of electrical shock and fire hazard, inspect the power supply, AC power cords, cables, and plugs on a regular basis to ensure that they are not damaged.
	The Kosmos system is not defibrillation proof. To prevent injury to the operator/bystander, Kosmos probes must be removed from patient contact before the application of a high-voltage defibrillation pulse.
	Before using Kosmos for needle guidance procedures, you must have training in the applicable interventional procedures in addition to training in the use of ultrasound imaging for needle guidance. Well known limitations of ultrasound physics may lead to an inability to visualize the needle or differentiate the needle from acoustic artifacts. Serious injury or complications may result from attempting an interventional procedure without proper training.
	As a precaution, be careful when scanning near a wound or over a dressing.
	Do not use Kosmos for intracavity imaging.
	Kosmos uses Bluetooth wireless communication technology.
	Keep power cords away from trafficked areas.
	No modifications to this equipment shall be made without written consent of manufacturer, EchoNouS, Inc.
	Do not charge tablet while scanning a patient unless it is connected to the Kosmos Link with the Globtek P005974 power supply.
	Do not connect any unauthorized equipment while using the Kosmos system.
	Only use tablets that have been approved compatible by EchoNouS.
	Certain tablets require the Kosmos Link to operate Kosmos. Please check with your EchoNouS representative or visit the EchoNouS website for more information.

EchoNouS customer support

Contact customer support:

Phone: 844-854-0800

Fax: 425-242-5553

Email: info@echonous.com

Web: www.echonous.com

Resources: echonous.com/product/resources

- End of section -

Kosmos Overview

What is Kosmos?

Kosmos consists of Kosmos Torso-One or Kosmos Lexsa connected by cable to a compatible tablet which runs the EchoNous Kosmos Ultrasound App. When the display is connected to a Kosmos probe, the combination is configured as a medical electrical system. The current list of compatible tablets can be found on the EchoNous website at echonous.com/product/device-compatibility.

The following probes are available for the Kosmos System:

- Kosmos Torso-One:
 - A phased array ultrasound-only probe with a smaller, more streamlined form factor to help fit in between intercostal spaces.
 - Provides portable ultrasound imaging and supports non-invasive Cardiac, Thoracic/Lung, Abdominal imaging.
- Kosmos Lexsa:
 - A linear array ultrasound probe.
 - Provides portable ultrasound imaging and supports non-invasive Lung, Vascular/Peripheral Vascular, Musculoskeletal and interventional guidance (includes needle/catheter placement, fluid drainage, and nerve block).

Kosmos uses pulse-echo ultrasound to generate real-time ultrasound images. This process involves transmitting high-frequency acoustic pulses into the body from the probe and detecting the returned signals and processing the return echoes through analog and digital processing to form real-time images of anatomy (B-mode and M-mode) and blood flow (Color Doppler). Reference **TABLE 4-3, "Modes of operation and features for Kosmos on iOS," on page 31** for more information about which modes are applicable for each Kosmos Probe.

Kosmos Link can be used as an optional accessory to provide extended scanning time for all imaging modes when used with compatible iOS tablets. The Link also provides for connecting multiple probes, which the user can select on the screen of the tablet. Please visit the EchoNous website for more information.

Kosmos provides optional wireless connectivity, allowing remote storage.

Kosmos also includes the AI-assisted EF Workflow, AI FAST, and Kosmos Bladder AI tools.

Kosmos Overview

Kosmos uses ultrasound imaging to permit a clinical assessment of the key cardiac structures, including the heart chambers, heart valves, and major heart vessels for adult and pediatric patients. As part of this clinical assessment, Kosmos permits visualization of blood flow using color Doppler technology.

The Kosmos AI-assisted EF workflow can help guide you through the calculation of left ventricular (LV) ejection fraction (EF). Kosmos uses a guided workflow to record the necessary clips. The recorded clips are then used by AI to provide an initial calculation of the EF and stroke volume (SV) with results that you can review and adjust if you need to.

More specifically, Kosmos AI provides an initial calculation of the EF, which is based on identifying the end-diastolic (ED) and end-systolic (ES) frames, along with the corresponding LV contours. Those ED/ES frames and LV contours can then be adjusted (as necessary) or accepted as is.

While reviewing these frames, you can adjust them based on your analysis, while Kosmos (using your adjustments) computes the EF and stroke volume (SV).

Kosmos AI FAST can help guide you through a FAST exam by identifying views and labeling key anatomical structures in real-time.

Kosmos Bladder AI can help determine the bladder volume by placing calipers on images acquired during a biplane bladder exam.

	SV is calculated as ED LV volume minus ES LV volume.
	Features vary by software version. For more information on available features for your device, please contact your EchoNous representative.

Kosmos clinical applications

Kosmos is for non-invasive imaging of the human body and is intended for the following applications by probe:

Torso-One:

- Cardiac
- Thoracic/Lung
- Abdominal
- Bladder

Lexsa:

- Lung
- Vascular/Peripheral Vascular
- MSK
- Nerve

Training

Kosmos is intended to be used by clinicians with appropriate professional qualifications and clinical training.

All users should read the generic ALARA education program supplied with Kosmos (see *ISBN 1-932962-30-1, Medical Ultrasound Safety* on the USB flash drive or the Health Canada *Guidelines for the Safe Use of Diagnostic Ultrasound* available on the Health Canada website). This program outlines the guiding principle for diagnostic ultrasound, where the qualified user keeps ultrasound exposure to "as low as reasonably achievable" while performing a diagnostic examination.

In addition to the above, users intending to use the ultrasound imaging function must have appropriate training in ultrasound. Appropriate information on training may be obtained by contacting EchoNous or your local professional body.

Kosmos classifications

- Kosmos Torso-One and Kosmos Lexsa are Type BF Applied Parts. The Applied Parts include:
 - The lens (front surface) of the Kosmos probe
- Kosmos Torso-One and Kosmos Lexsa are IPx7
- Kosmos Link with an approved power supply and an approved tablet classifies as a medical electrical system.
- Kosmos Link is IP32 rated

Patient environment

Kosmos is intended to be used in a medical facility. The Link and tablet may be charged in the patient environment using the Globtek P005974 power supply.

 Do not charge tablet while scanning a patient unless it is connected to the Kosmos Link with the GlobTek P005974 power supply.

- End of section -

Using Kosmos

System overview

Use this section to acquaint yourself with the ultrasound system and components.

Device requirements

For a list of devices that EchoNous has tested and determined to be compatible with the Kosmos app, visit the Kosmos website at echonous.com/product/device-compatibility.

The EchoNous Kosmos Ultrasound App can only be downloaded and installed on the supported tablets listed on the EchoNous website. The key requirements met by the supported tablets are listed below:

iOS:

- Minimum of 50 MB of storage space (plus more for patient data storage)
- Color display, minimum 203mm (8 in)
- Touch interface
- Internally mounted speakers
- IEC 60950-1-compliant or IEC 62386-1 compliant
- Only one USB port
- Date/time configuration
- Full compliance with USB On-The-Go standard
- 2560 x 1600 resolution (minimum)
- iOS 15 later operating system
- Wireless or cellular networking capability
- Audio capability
- Front- and rear- facing cameras

Please review all safety considerations in the chapter titled **Safety**. The tablet must have the corresponding ratings to be used within specified environmental conditions.

Kosmos hardware

 Contact EchoNous or your local representative for a list of accessories available from or recommended by EchoNous.

The figures in the following sections point out key features on Kosmos Torso-One, Kosmos Lexsa and the Link.

- **"Kosmos Torso-One" on page 12**
- **"Kosmos Lexsa" on page 12**
- **"Kosmos Link" on page 12**

Kosmos Torso-One



Kosmos Lexsa



Kosmos Link



Charge with Globtek P005974 power supply.

Getting started

Downloading the EchoNouS Kosmos ultrasound app

Getting Started with Kosmos on iOS

1. Connect iOS tablet to Wi-Fi.
2. If applicable, delete the previously installed version of the Kosmos App from the tablet.

	Ensure you have archived data before deleting the previously installed version of the Kosmos App from the tablet.
---	---

3. Download the EchoNouS **Kosmos Ultrasound App** from the Apple App Store.
4. Open the Kosmos App. From the **Home** screen, tap **Enable drivers**. You will be directed to the tablet's settings. Toggle each driver to the on position.

Connecting Kosmos probes

	Before each use, inspect Kosmos Torso-One and/or Kosmos Lexsa for damage, such as cracks, splitting, or sharp edges. If damage is evident, discontinue using the Kosmos probe(s) and contact your EchoNouS representative.
	Only use devices and accessories recommended by EchoNouS.

To connect Kosmos Torso-One or Kosmos Lexsa to approved iOS tablets:

1. Plug the Kosmos probe cable into the USB-C port on the side of the tablet.
 - To register your transducer and licensed features for the first time, the probe must be connected to the device and your device must be connected to the internet. This step may take a few minutes.
2. When ready to start scanning, tap the preset of your choice to begin.

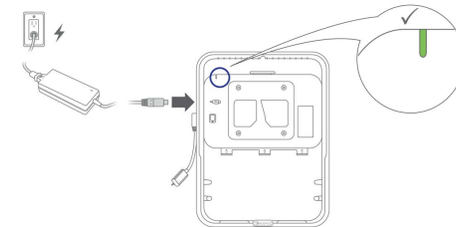
Kosmos Link for iOS

Kosmos Link is a power source that allows the use of all features on approved iOS tablets and provides extended scanning time with Kosmos probes. Please visit echonous.com/product/device-compatibility for an updated list of compatible tablets.

Setting up Kosmos Link

	The Link is intended to be used only with compatible iOS tablets. Please contact your EchoNouS representative for additional details.
	Ensure Link is placed such that the probe connection port, charging port, and wall outlet are accessible.
	For more detailed Link instructions, please refer to the Kosmos Link Quick Guide (P008154).
	Ensure Kosmos Link is securely attached to the tablet prior to use.
	Ensure Link is securely mounted on the stand or safely placed on a tabletop with the kickstand fully extended prior to use.

1. Charge Kosmos Link prior to use, until the LED shows green
2. To install the tablet onto the Link, bring the tablet / bracket assembly to the front face of the Link.
3. Slide the tablet downward, ensuring that it is moving along the rubber seal on the front of the Link. The orange slider button (under the rubber cap) will move over, then snap back to its original position. This indicates that the tablet is secure on the Link.
4. Connect the Link USB-C cable to the tablet USB-C port.



How to remove tablet from Kosmos Link

- * To remove the tablet, pull the orange slider button and then move the tablet upward until it is free of the Link.

Charging Kosmos Link

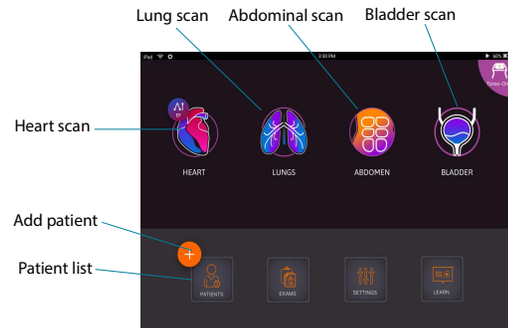
1. Probes may stay connected during charging.

- Connect the charger to the Kosmos Link. Once connected, the LED on the Link will indicate the general battery power level. White is low, blue is mid-range, and green is full.

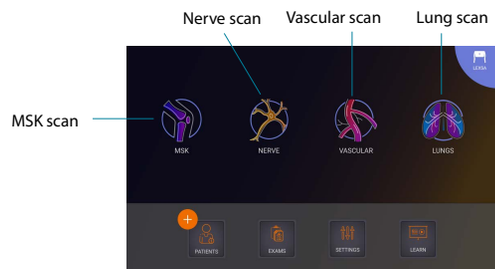
Battery Status	Battery Level		
	0%-20%	20%-80%	80%-100%
Not charging	Solid White	Solid Blue	Solid Green
Charging	Flashing White	Flashing Blue	Flashing Green

General interaction

Home screen: Kosmos Torso-One



Home screen: Kosmos Lexsa



Learn

To access the how-to-videos available on YouTube, ensure your device is connected to Wi-Fi and tap **Learn**.

Settings

Once you have configured your system settings, they remain as you set them whenever you log back on to the Kosmos App.

Imaging preferences

The **Imaging Preferences** screen is where you can customize the information displayed on the **Imaging** screen.

To set the imaging preferences:

- From the **Home** screen, tap **Settings**.
- Tap **Imaging Preferences**.
- To have certain information display in the top bar of the Imaging screen, tap one of the following options under **Customize information**:
 - Name of facility** Displays the name of your organization in the top bar of the imaging screen.
 - Patient name** Displays the patient name in the top bar of the imaging screen.
 - Patient ID** Displays the patient ID in the top bar of the imaging screen.
- To set how long the clips record, select a time from the **Clip Duration** area.
- To configure the way Kosmos records clips, select **Prospective** or **Retrospective** under the **Record Clip**:
 - Prospective** Captures frames after you tap the **Record Clip** icon. Kosmos captures frames for the selected **Clip Duration**.
 - Retrospective** Captures frames from the cine buffer when you tap the **Record Clip** icon. Kosmos captures cine buffer frames for the selected **Clip Duration**.

	Once a selection is made, a corresponding p or r will be present on the video button while live scanning.
	During an exam, if you tap the Record Clip icon again, you can finish the recording earlier than the clip duration defined here.

- To adjust the horizontal screen split between **M-mode** and **B-mode**, select from the following options under **M-Mode layout**:
 - 1:2** Tap this option to adjust the screen split so the M-mode area is twice as big as B-mode.

- **1:1** Tap this option to adjust the screen split so that the M-mode and B-mode areas are equal.
7. From the **Thermal index display** area, select from the following:
 - **TIS** Thermal index for soft tissue.
 - **TIB** Thermal index with bone near the focus.
 8. Select the **cardiac imaging orientation** preset:
 - Select **Left** or **Right** orientation.
 9. To enable **Auto Functionality** features, tap the toggle to switch to the on position.
 - **Auto Doppler** When scanning in cardiac PW and TDI modes, use **Auto Doppler** for AI-assisted auto placement of PW and TDI sample gates.
 - **Auto Preset** When scanning in Heart, Lung and Abdomen presets, the AI-assisted **Auto Preset** feature will recognize anatomy and automatically transition to the appropriate preset.
 10. For **PW** and **CW** modes, select from the following:
 - **Synchronized** focal point/gate and color box.
 - **Decoupled** focal point/gate and color box.

About

The **About** section is where you will find essential information about your device, such as the Kosmos software version, model number, device registration status, and licensed features. You will also be able to access transducer information, perform a transducer element check, and find the contact information for support.

1. From the Kosmos app **Home** screen, go to **Settings** --> **About**.
2. If you have not registered Kosmos, tap **Register**. This will connect your Kosmos device to the EchoNous cloud. Make sure your device is connected to the internet.
3. To run the transducer element check, tap **Check**.

DICOM

Manage your modality worklist (MWL) and PACS archive from the DICOM section.

- New systems do not come with any configured profiles.
- You cannot have two PACS profiles active at the same time; when you add a new profile, the current one is deactivated.

Adding a profile

To add a PACS profile:

1. From the **Home** screen, tap **Settings**.

2. Tap **DICOM** --> **PACS archive**.
3. Tap **Add Profile**.

If you are adding a new PACS-SCP profile and already have an existing one, the system deactivates the existing profile. However, all the jobs in the existing queue and any scheduled archives must first be completed.

4. Type the following information in the **DICOM connection** area:
 - **Station AE title** Kosmos' Application Entity title
 - **Server AE title** Archive server's Application Entity title
 - **Server IP address** Archive server's unique identifier
 - **Server port number** Archive server's port number
5. To make sure the connection is working on an active profile, tap one of the following:
 - **PING** to test the network connection between Kosmos and the PACS archive.
 - **Verify** to check the availability of the active PACS archive.

The results are displayed on-screen.
6. In the **Profile nickname** box, type a unique name to display in the PACS profile list.
7. In the **Archival options** area, you have three options:
 - **Prompt options every time** Switched on by default; each time you tap the **Archive** button from the Exam review screen, a pop-up menu with different options displays. If you turn the switch off, Kosmos does not display the pop-up menu.
 - **Attach report** Switched off by default. If you turn it on, Kosmos attaches a report to the archive.
 - **Attach DICOM SR report** Switched off by default. When selected, Kosmos will attach the DICOM SR report to the archive.
8. In the **Auto archive** area, select from the following options:
 - **On/Off** The auto archive is switched off by default. This means that all the controls (except the on/off switch) are disabled and cannot be edited. If you turn the switch on, all the controls are enabled and can be edited.
 - **Archival frequency**
 - **Completion of exam** The archival time selector is disabled.
 - **Daily** Only the time section of the archival time selector is enabled.
 - **Weekly** The complete archival time selector is enabled
 - **Archival time** Select a daily time and day to archive exams. In the **Retry interval (in seconds)** area, select **60**, **300**, or **600**.

If you turn on auto archive, make sure the Kosmos App is always running in the background. Closing the Kosmos App will pause the archives. Go to Job Queue to resume or retry if job(s) are not successfully archived.

- In the **Maximum retries** area, select **1,2**, or **3**.
- To have the system automatically retry failed jobs, keep the switch set to **On**; otherwise, slide it to **Off**.

Deactivating a profile

- To activate or deactivate a profile, in the **PACS archive** list, tap the switch to toggle between **Active** and **Inactive**.

TLS SETTING FOR DICOM:

- On the **active profile** page, tap **Settings**.
- Tap **DICOM** --> Scroll down to **TLS Encryption** section and turn on **TLS Encryption**.
- Select **SCU Security**. The options are **Anonymous** or **Authenticated**.
- Next set the SCP Certificate for the profile. Select the option **Select TLS Certificate** or **Select TLS Certificate from Device**.
- Clicking the **Select TLS Certificate** option initiates the selection of a new certificate. This option displays the file explorer for the user to select the certificate provided by the administrator.
- Clicking the **Select TLS Certificate from Device** option displays the list of certificates already set up in the application.

Deleting a profile

To delete a PACS profile:

Deleting a PACS profile also deletes all configurations of the profile. There must be an active PACS profile before you can archive any exams.

- From the Home screen, tap **Settings**.
- Tap **DICOM** --> **PACS archive**.
- From the list of profiles, tap to slide the arrow to the left of the profile you would like to delete.
- Tap the **Delete**  icon.

Managing MWL

New systems do not come with any configured profiles.
You cannot have two MWL profiles active at the same time; when you add a new profile, the current one is deactivated.

Adding a profile

To add a MWL profile:

- From the **Home** screen, tap **Settings**.

- Tap **DICOM** --> **MWL**.
- Tap **Add Profile**.

If you are adding a new MWL profile and already have an existing one, the system deactivates the existing profile.

- Type the following information in the **DICOM connection** area:
 - Station AE title** Kosmos' Application Entity title.
 - Server AE title** Archive server's Application Entity title.
 - Server IP address** Archive server's unique identifier.
 - Server port number** Archive server's port number.
- To make sure the connection is working on an active profile, tap one of the following:
 - PING** to test the network connection between Kosmos and the MWL server.
 - Verify** to check the availability of the active MWL server.
 - The results are displayed on-screen.
- In the **Profile nickname** box, type a unique name to display in the MWL profile list.

Deactivating a profile

- To activate or deactivate a profile, in the **MWL** list, tap the switch to toggle between **Active** and **Inactive**.

Deleting a profile

To delete a MWL profile:

Deleting a MWL profile also deletes all configurations of the profile.

- From the **Home** screen, tap **Settings**.
- Tap **DICOM** --> **MWL**.
- From the list of profiles, tap to slide the arrow to the left of the profile you would like to delete.
- Tap the **Delete**  icon.

USB export

To configure USB export preferences:

- From the Kosmos app **Home** screen, go to **Settings** --> **USB export**.
- Check the box to enable exporting exams to USB drive.
- Select the file type.

Report settings

To customize the measurements and metrics of the report settings:

1. From the Kosmos app **Home** screen, go to **Settings** --> **Report Settings**.
2. For each cardiac measurement, select from the following:
 - **Last** measurement taken
 - **Average (Avg)** measurement
 - **Maximum (Max)** measurement
3. Select the metrics for distance and velocity.

Wireless Networking Functions

You can connect Kosmos to an IT network to perform the following:

- Storing exam data (static images and clips) acquired by Kosmos in Picture Archiving and Communication System (PACS) by DICOM communication.
- Setting Kosmos time correctly by inquiring the network time service.

Connection specifications

Hardware specification

802.11 a/b/g/n/ac, Bluetooth 4.0 or later.

Software specification

Kosmos is connected to PACS by the DICOM standard. For details, refer to the DICOM Conformance Statement that is available on the EchoNous website.

Use restriction

This device is restricted to indoor use when operating in the 5150 to 5350 MHz frequency range. This restriction applies in: AT, BE, BG, CH, CY, CZ, DE, DK, EE, EL, ES, FI, FR, HR, HU, IE, IS, IT, LI, LT, LU, LV, MT, NL, NO, PL, PT, RO, SE, SI, SK, TR, UK.

-- End of section --

Performing an Exam

Overview

	Before using Kosmos for a critical procedure, such as needle guidance, make sure it is fully charged. You do not want the procedure interrupted by a drained battery, which may cause harm to the patient.
	The maximum temperature of a Kosmos probe scan head may be greater than (41°C) but is less than (43°C) when in contact with the patient for normal use. Special precautions should be considered when using the transducer on children or on other patients who are sensitive to higher temperatures.
	To reduce the risk of infection, use sterile sheaths when conducting needle procedures.
	To avoid a mix-up of patient data, complete the exam before examining a new patient.
	Not all features are available in every market and vary by regionally released software versions. For more information on available features for your device, please contact your EchoNous representative.

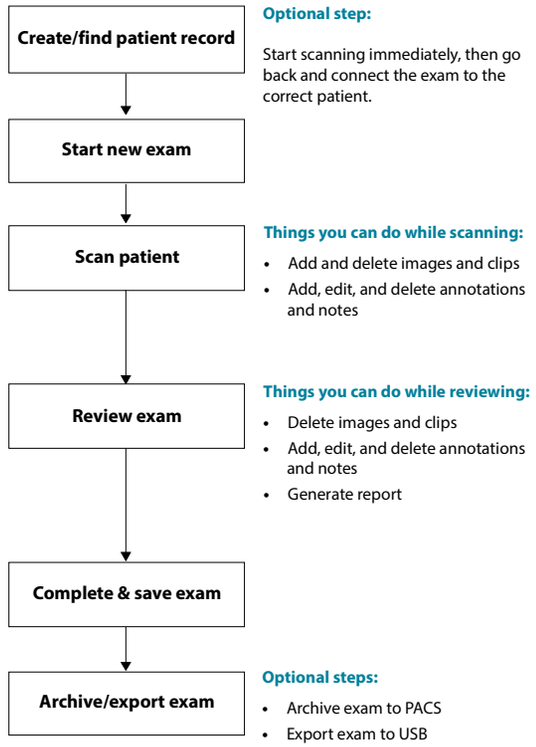
Primary exam workflows

With Kosmos, there are three primary exam workflows; click one of the links to go to that workflow:

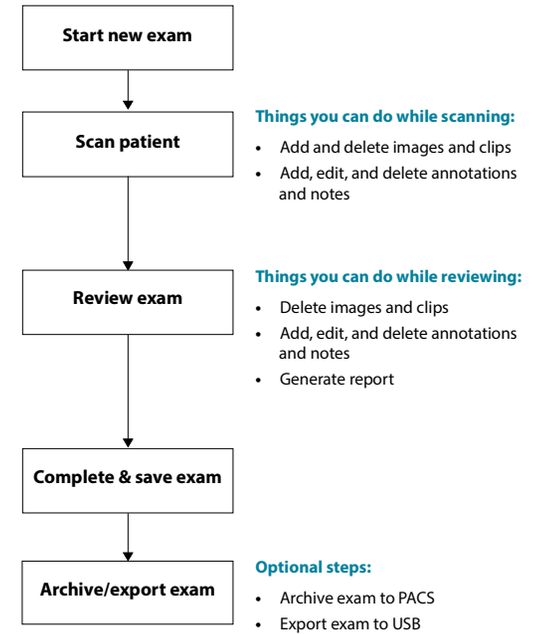
- **“Standard workflow”** starts with either creating a patient or searching for an existing patient.
- **“Quick workflow”** starts with scanning a patient.
- **“AI-assisted EF workflow”** uses AI to perform initial EF calculations.
- **“Kosmos Bladder AI Workflow”** uses AI to place calipers to measure bladder volume.

Exam workflows

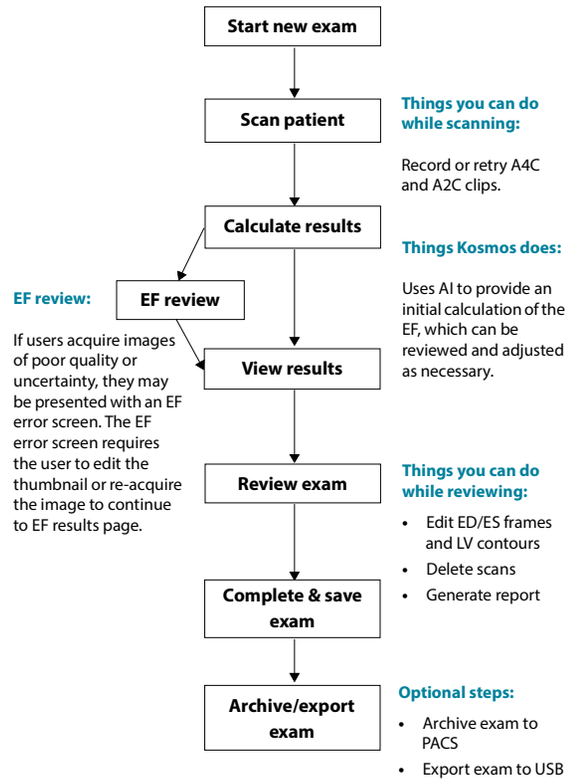
Standard workflow



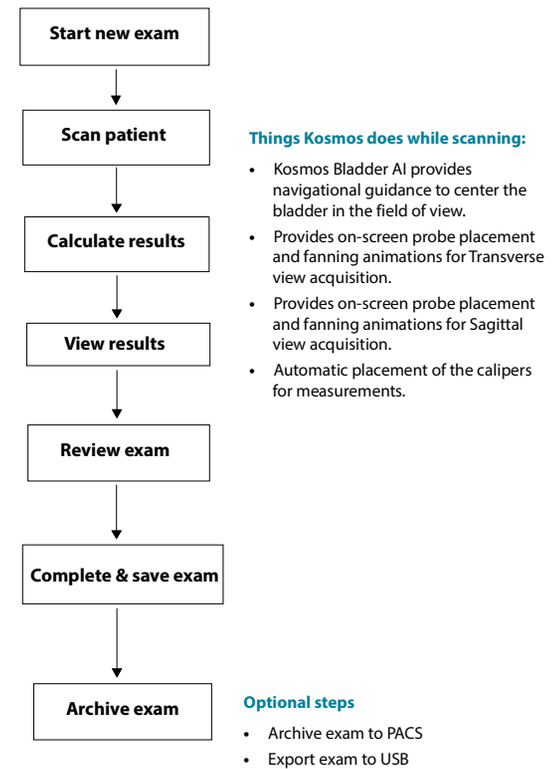
Quick workflow



AI-assisted EF workflow



Kosmos Bladder AI Workflow



Managing exams

Starting an exam

There are several ways you can start an exam:

- To start scanning immediately, from the **Home** screen, tap a preset and begin scanning.

- When you save the exam, Kosmos automatically generates a temporary ID and saves the images/clips to the temporary ID.
- From the **Home** screen --> **PATIENTS** --> **NEW PATIENT** --> **SCAN**.
 - Use the **Add**  icon as a shortcut to add a new patient.
- For existing patients, from the **Home** screen --> **PATIENTS** --> Select a patient from patient list --> **SCAN**.
- From the **Home** screen --> **EXAMS** --> **NEW PATIENT** or look up an existing patient--> **SCAN**.

Searching for an exam

To search for an exam:

1. From the **Exam** screen, tap the **Search**  icon.
2. Type the search criteria, such as date, patient name, DOB, or MRN.
3. From the list of search results, tap the exam you want to view. Each exam listed shows the number of scans taken, as shown below.



Deleting exams

To delete one or more exams:

1. From the list of exams, tap one or more circles to the left of the exam. The circle turns into a check mark, showing it is selected.
2. Tap the **Trash**  icon.
3. At the prompt, tap **OK**.

To delete all the empty exams (those without images/clips):

1. From the list of exams, tap the **More options**  icon.
2. Tap **Delete all empty exams**.
3. At the prompt, tap **OK**.

Acquiring images and clips

To acquire an image:

- * From the **Imaging** screen, tap the **Save image**  icon.

To acquire a clip:

- * From the **Imaging** screen, tap the **Save clip**  icon.

Completing exams

To avoid mixing up images and clips saved from multiple patients, make sure to complete an exam.

To complete an exam:

1. From the **Imaging** screen, tap the **Exam review**  icon.
2. Tap **Complete**.
3. At the prompt, tap **OK**.

If you do not tap **Complete** from the **Exam Review** screen, Kosmos automatically completes the exam:

- When you start a new exam
- When you archive the in-progress exam
- When the app is closed

Managing patient data

Adding a new patient

To add a new patient from the **Home** screen:

1. From the **Home** screen, tap the **Add**  icon on the **Patients** button.
2. Enter the patient information.
3. Optionally, you can enter exam information.
4. Tap **Scan** when you are done.

Accessing patient information using MWL

If you are connected to a healthcare information system and MWL is set up on your Kosmos, you can access patient information.

1. From the **Home** screen, tap the **Patients** button.
2. Tap the **MWL** button. Tap the  icon to see the entire list.
3. Tap the  icon to search for a specific patient.
4. Tap **Scan** to start scanning.

Searching for a patient

To search for a patient:

1. From the **Home** screen, tap **Patients**.
2. Tap the **Search**  icon.

3. Type the search criteria for the patient you are looking for, such as name, date of birth, or medical record number.
4. Select the patient from the search result list, and tap **Done**.

Changing to another patient

To change to or add a new patient when you have already started an exam:

1. From the **New Exam** screen, tap **Change**.
2. Do one of the following:
 - To change to another patient, tap **Add New**, and complete the patient form.
 - To look for an existing patients, tap **Search History**, use the search tool to find the patient, and tap the patient name from the list.

Editing a patient record

To edit a patient record:

1. From the **Home** screen, tap **Patients**.
2. From the Patient list, double-tap the patient record you want to edit.
3. Enter the patient information, and tap **Save** when you are done.

Merging two patient records

If you have saved multiple patients with the same name, and they are actually the same patient, you can merge all the exams of that patient into one patient record so it is easier to keep track of that patient.



In order to merge two patients, make sure the following fields are complete:

- **First name**
- **Last name**
- **DOB**
- **Gender**

To merge two patient records:

1. From the **Home** screen, tap **Patients**.
2. Tap to select one of the patients.
3. From the **Patient review** screen, tap the **More options**  icon.
4. Tap **Merge to patient**.
5. From the list, tap the other patient you want to merge.
6. Tap **Next**.

7. Tap the fields to keep for the patient.
8. Tap **Merge**, then tap **OK**.

Deleting patient records

To delete all patient records without exams:

1. From the **Home** screen, tap **Patients**.
2. Tap the **More options**  icon.
3. Tap **Delete all patients without exams**.

To delete selected patient records:

1. From the **Home** screen, tap **Patients**.
2. Tap one or more patient names from patient list.
3. Tap the **Trash**  icon.

Organ presets

TABLE 4-2 provides an overview of the organ presets that are available for each Kosmos probe.

TABLE 4-2. Organ presets by Kosmos probe

Organ	Torso-One	Lexsa
Heart	x	
Lung	x	x
Abdomen	x	
Bladder	x	
Vascular		x
Nerve		x
MSK		x

Imaging modes and features

For an overview of the applicable imaging modes for each Kosmos probe, reference **TABLE 4-3**.

TABLE 4-3. Modes of operation and features for Kosmos on iOS

Mode	Torso-One iOS	Lexsa iOS
B-mode	x	x
M-mode	x	x
B + CD (Color Doppler)	x	x
Harmonic Imaging	x	
AI-assisted EF Workflow	x	
PW Doppler	x	x
TDI	x	
CW Doppler	x	
AI FAST	x	
Kosmos Bladder AI	x	
Color Power Doppler		x
Cardiac Calculations	x	
Vascular Calculations		x
Auto Preset	x	
Auto Doppler (for Cardiac preset in PW and TDI modes)	x	

2D/B-mode

2D/B-mode is the system's default imaging mode. The system displays echoes in two dimensions by assigning a brightness level based on the echo signal amplitude.

The 2D/B-mode controls are hidden when in Doppler modes. You can switch between 2D/B-mode and Doppler mode controls.

- ★ To view the 2D/B-mode controls, tap **2D**.

M-mode

M-mode is also known as Motion Mode. It provides a trace of the image displayed over time. A single beam of ultrasound is transmitted, and reflected signals are displayed as dots of varying intensities, which create lines across the screen.

When M-mode is turned on, the screen splits to show B-mode as well as M-mode. You can adjust depth, and gain (similar to B-mode) along with M-mode specific controls like M-line and sweep speed.

While scanning with Lexsa probe, M-mode is only available in the Lung preset.

- ★ To start M-mode, tap the **M-mode**  icon.

M-Line

- ★ To move the M-Line, use your finger to change to M-mode, tap and drag the M-Line to the location you want.

Sweep speed

You can change the sweep speed to isolate individual motions.

- ★ To change the M-mode sweep speed, tap **Speed** and select: **25, 50, 75** or **100** mm/sec.

Color Doppler

Color Doppler is used to visualize the presence, velocity, and direction of blood flow in a wide range of flow states.

When using Kosmos, you can turn Color Doppler on and off without it interfering with the system's color acquisition.

- ★ To turn Color Doppler on and off, tap the **Color**  icon.

Color box

You can move and resize the color box during imaging. The maximum axial and lateral size of the box may be limited depending on the organ, depth, or other settings.

- To move the color box, select the side of the color box and drag it to another position.
- To resize the color box, select one of the corners to adjust the size.

Scale

Scale changes the pulse repetition frequency that defines the velocity scale with the range shown at the top and bottom of the color map.

- ★ To change the scale, tap on the **Scale**.

Sensitivity

Three sensitivity range selections are available to optimize for low, medium, and high range.

- ★ To change the sensitivity, tap **Sensitivity**, and select an option.

Wall filter

The wall filter is set on the highest filter which blocks low frequency noise.

- ★ To change the wall filter, tap **wall filter**, and select the appropriate option.

Steer

Steer changes the steering angle of the color ROI. There are 5 angles to choose from.

- ★ To select desired angle, tap **Steer**.

Steer is only available in Lexsa Color Doppler mode.

Artery

Artery enables Artery/Vein selection. Artery should be selected for arterial flow and Vein should be selected for venous flow.

- ★ For Artery/Vein selection, tap **Artery**.

Artery is only available in Lexsa Color Doppler mode.

Color Map

To change the heart color map:

1. Tap the **More options**  icon next to the color map on the right side of the screen.
2. Select the color map you like.
3. To invert the color map, select the check box, and tap **OK** to save the changes.

Color Power Doppler

Color Power Doppler (CPD) is used to measure the amplitude of blood flow. CPD is more sensitive to lower blood velocities and smaller vessels.

- ★ To turn Color Power Doppler on and off, tap the **CPD** icon .

Color Power Doppler is available in Vascular, Nerve and MSK presets while scanning with Kosmos Lexsa.

Pulsed-Wave Doppler

Pulsed-Wave Doppler (PW) mode uses short bursts of ultrasound with a process called range gating to facilitate signal analysis from a small area at a specified depth from the transducer.

PW mode is available in the abdomen and heart preset while scanning with Kosmos Torso-One.
PW mode is available in the Vascular, Nerve and MSK presets while scanning with Kosmos Lexsa.

- ★ To start PW Doppler, tap the **PW mode** icon.

Duplex screen

- ★ Tap the **Update** button for the **duplex** screen. The frozen B-mode image will be displayed on the top with the live Doppler trace on the bottom.

Focal point and Doppler line

- ★ Adjust the focal point and the Doppler line by moving the dotted circle. In the abdomen preset, tap the focal point to see view and set the angle adjust line. If color mode is on, moving the circle will also move the color box. The circle and the color box can be decoupled by going to **Settings** --> **Imaging Preferences**.

Baseline

- ★ Tap and move the baseline up and down in the Doppler trace.

Live display

- ★ Tap the **Live display** to toggle between PW live and B live modes. In the B live mode, Doppler trace is frozen.

Wall filter

Wall filter helps to filter out echoes from low frequency signals.

- ★ Tap icon to select the strength of the filter: **Low, Medium, High**.

Invert

- * To invert the Doppler spectrum, tap on the **Invert** button.

Scale

Scale changes the velocity scale.

- * To change the scale, tap **Scale**.

Doppler Gain

Gain controls the brightness/strength of the Doppler spectrum.

- * To adjust Doppler gain, tap **Gain**.

Audio gain

Audio Gain controls strength of the audio volume.

- * To adjust Audio gain, tap **Audio gain**.

Sweep speed

Four sweep speed selections are available.

- * To change the sweep speed, tap **Speed** and select: **25, 50, 75** or **100** mm/sec.

Tissue Doppler Imaging

Tissue Doppler Imaging (TDI) mode uses Doppler to measure the velocity of myocardial motion throughout the cardiac cycle.

- * To start TDI mode, tap on the **TDI mode** icon. TDI is available in **B-mode** and **Color (B+C) mode** screens.

<p>TDI mode is only available in abdomen and heart presets while scanning with Kosmos Torso-One.</p>
--

Continuous-Wave Doppler

Continuous-Wave Doppler (CW) mode uses continuous transmission and reception of ultrasound waves to measure blood velocities.

<p>When CW is used for a prolonged period, auto-freeze goes into effect to manage probe temperature. A 60-second timer appears every time before the auto-freeze.</p>
<p>CW mode is only available in the abdomen preset and heart preset while scanning with Kosmos Torso-One.</p>

- * To start CW Doppler, tap the **CW mode** icon.

Duplex screen

- * Tap the **Update** button for the **duplex** screen. The frozen B-mode image will be displayed on the top with the live Doppler trace on the bottom.

Focal point and Doppler line

- * Adjust the focal point and the Doppler line by moving the dotted circle. In the abdomen preset, you can tap the focal point to see and set the angle adjust line. If Color mode is on, moving the circle will also move the color box. The circle and the color box can be decoupled by going to **Settings** --> **Imaging preferences**.

Baseline

- * Tap and move the baseline up and down in the Doppler trace.

Live display

- * Tap the **Live display** to toggle between CW live and B live modes. In the B live mode, Doppler trace is frozen.

Wall filter

Wall filter helps to filter out echoes from low frequency signals.

- * Tap icon to select the strength of the filter: **Low, Medium, High**.

Invert

- * To invert the Doppler spectrum, tap on the **Invert** button.

Scale

Scale changes the velocity scale.

- * To change the scale, tap **Scale**.

Doppler Gain

Gain controls the brightness/strength of the Doppler spectrum.

- * To adjust Doppler gain, tap **Gain**.

Audio Gain

Audio gain controls the strength of the audio volume.

- * To adjust Audio gain, tap **Audio gain**.

Sweep speed

Four sweep speed selections are available.

- * To change the sweep speed, tap **Speed** and select: **25, 50, 75** or **100** mm/sec.

Save clips and images

- * Tap **Freeze** to review or directly save images and clips. Audio will also be saved in clips.

Auto Preset

When scanning in a selected preset, the Auto Preset feature will recognize anatomy and automatically transition to the appropriate preset. This feature is only available for Torso-One.

- * To enable Auto Preset, go to **Settings --> Imaging Preferences** and use toggle to enable feature.
 - Users are provided 3 seconds to reject the transition from the selected preset to the auto adjusted preset.

 If the user rejects the transition to the auto adjusted preset, Auto Preset will be disabled for the rest of the exam. Users have the ability to turn Auto Preset back on by selecting the Preset drop down menu.

- Please reference **TABLE 4-4, "Auto Preset scenarios," on page 38** for a list of Auto Preset scenarios.

TABLE 4-4. Auto Preset scenarios

User Selected Preset	Anatomy Scanned	Kosmos Auto Adjusted Preset
Abdomen	Lung	Lung
Abdomen	PLAX, PSAX, (AV, MV, PM, Apex), A4C, A2C, A3C, A5C, SSN, RVOT, RVIT	Heart
Lung	RUQ, LUQ, SUP, Abdominal Aorta (sagittal view), Aortic Sweep	Abdomen
Lung	PLAX, PSAX, (AV, MV, PM, Apex), A4C, A2C, A3C, A5C, SSN, RVOT, RVIT, IVC, Subcostal 4C	Heart
Heart	RUQ, LUQ, SUP, Abdominal Aorta (sagittal view), Aortic Sweep	Abdomen
Heart	Lung	Lung

Auto Doppler

The Auto Doppler feature will automatically place the Doppler gate in selected views. This feature is only available in PW and TDI modes for Torso-One in the Cardiac preset.

- * To enable Auto Doppler, go to **Settings --> Imaging Preferences** and use toggle to enable feature.
 - Users will still have the option to place the gate manually when Auto Doppler feature is enabled.
 - Please reference **TABLE 4-5** for a list of Auto Doppler gate placements.

TABLE 4-5. Auto Doppler gate placement by mode

Mode	Gate Placement	View
PW	Mitral Valve	A4C
PW	LV Outflow Tract	A5C
PW	Tricuspid Valve	A4C
PW	Pulmonary Valve	RVOT, PSAX-AV
TDI	MV Septal Annulus	A4C
TDI	MV Lateral Annulus	A4C
TDI	TV Lateral Annulus	A4C

Image mode controls

Flipping an Image

You can only flip an image from right to left when you are scanning the heart.

- ★ To flip the image, double-tap the orientation marker.

Adjusting Depth and Gain

To adjust depth:

- ★ To increase or decrease the displayed depth, tap **Depth**, and move the Depth wheel up and down.

To adjust gain:

- ★ To adjust gain in Color Doppler mode and B-mode, tap **Gain**, and move the slider up and down.

To adjust near and far gain:

- ★ Tap **TGC**, and move the sliders left and right. Notice the gain values automatically update as you adjust the sliders.

Zooming In and Out

- While scanning, use two fingers to pinch and expand the image area.
- To return to the default image size, tap the magnifying glass.
- Notice that the zoom factor is shown near the magnifying glass as well as the orange color of the depth scale along the side image area.
- You can freeze while zoomed (and can zoom out and zoom in while frozen).

Freezing an image

- ★ To freeze an image, tap the **Freeze**  icon. The annotation tools automatically display on the left side of the screen (see [“Annotating images and clips” on page 57](#) for more information).

Using the Kosmos AI-assisted EF workflow

The AI-assisted EF workflow guides you through the steps of data acquisition followed by an AI-based initial EF calculation which is based on the American Society of Echocardiography (ASE)-recommended modified Simpson’s method of disks (Lang 2005, 2015). The initial LV contours are produced with AI algorithms that have been trained on expert-annotated LV contours (Ronneberger 2015). You can then review the initial AI results (which include the ED/ES frames along with the corresponding LV contours), and adjust them, as necessary.

Calculating EF with the AI-assisted EF workflow

To calculate EF:

1. From the **Home** screen, tap the **Heart AI** icon.

Tap to start the AI-assisted EF workflow

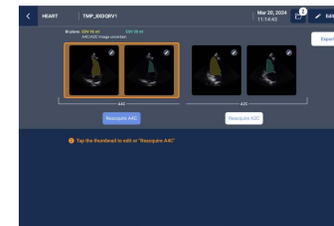


	When you tap the Heart AI icon, Kosmos creates a new exam that includes this EF scan.
	Do not rely upon EF calculation as the sole diagnostic criteria. Whenever possible, use EF calculation in conjunction with other clinical information.

2. After you have a good A4C view of the patient, tap **A4C** to acquire a clip.
3. If you are not satisfied with the recorded clip, tap **Try again** to acquire a new clip, or tap **Accept** to proceed (after four seconds, Kosmos automatically accepts the clip).
4. Tap **Skip** to see the A4C results, or continue with the A2C acquisition.

	We recommend that you take both A4C and A2C clips for more accurate calculations.
---	---

5. After you acquire images, the algorithm will assess the quality and uncertainty of the clip and users may be presented with the **EF error** screen. To proceed to your results, the **EF error** screen requires that you edit the thumbnail or re-acquire the image.



- After you have a good A2C view of the patient, tap **A2C** to acquire a clip.
- If you are not satisfied with the recorded clip, tap **Try again** to acquire a new clip, or tap **Accept** to see the A4C/A2C (biplane) results (after four seconds, Kosmos automatically accepts the clip).

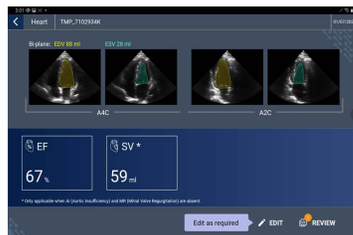
Note that when the A4C and A2C clips are recorded and accepted, the system selects the ED and ES frames, draws the corresponding LV contours, and calculates the biplane EF using the modified Simpson's method of disks (20 disks are used in the calculation).

Reviewing/adjusting the ED/ES frames and LV contours

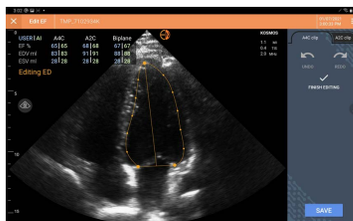
When reviewing the initial AI calculations for ED/ES frames and LV contours, you can adjust just the frames, the LV contours, or both before saving the results. If you do not make any changes, the AI calculations become the final result.

To adjust the ED/ES frames:

- From the **Results** screen, tap **Edit** or one of the thumbnail images. You can also tap **Review** to review previously acquired scans.



- Depending on which clip you'd like to edit, tap the **A4C clip** or **A2C clip** tab.
- To set a different ED or ES frame, move the orange Seek button to the desired location, and tap **SET ED** or **SET ES**.



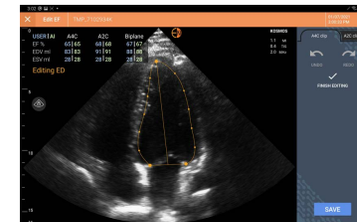
- To return to the original AI calculations, tap the **More options** icon and then **Reset**.

- If desired, make changes to the other clip (A4C or A2C), and tap **Save**.

To adjust the LV contours:

	If you are wearing gloves while editing the LV contours, make sure they are snug against your fingertips/nails.
	Having gel on your fingers may hinder using the touchscreen effectively. Make sure to wipe the touchscreen regularly.

- From the **Results** screen, tap one of the four images to go to that image. If you don't specify which image you want, Kosmos defaults to the A4C frame.
- Depending on which clip you'd like to adjust, tap the **A4C clip** or **A2C clip** tab.
- Tap the **A4C clip** or **A2C clip** tab to select an ED or ES frame.
- Tap the LV contour. The LV contour becomes adjustable, and the color changes to orange.



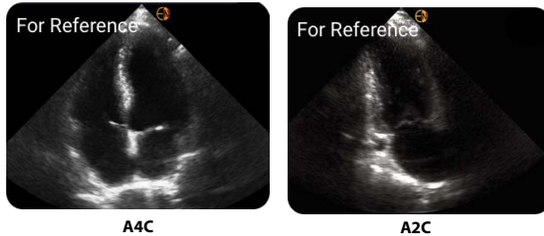
- Select one or more control points and move them.
Notice the calculations are updated as you change the contour.
- After you are done editing, tap **Finish editing**.
- If desired, make more changes.
- Tap **Save**.

Recommendations for acquiring optimal A4C and A2C clips for accurate EF calculations

EchoNous recommends the following:

- The patient should be lying on their side in the left lateral position (the left side of the patient is touching the scanning table).

Shown below are examples of clinically acceptable A4C and A2C reference images on the top left of the **Imaging** screen:



- For an A4C clip, ensure all four cardiac chambers (left ventricle, left atrium, right ventricle, and right atrium) are captured in the ultrasound image (see the A4C reference image above).
- For an A2C clip, ensure both left ventricle and left atrium are captured in the ultrasound image (see the A2C reference image above). Ensure the endocardial border of the LV is clearly visible with the best possible contrast. Use the **Gain** settings to achieve a clear definition of the LV endocardial border.
- Adjust the depth so that the atria are near the bottom of the ultrasound image yet still visible (see the A4C and A2C reference images above).
- Avoid truncating the LV.
- Avoid foreshortening the LV.
- For an A4C clip, ensure the intraventricular septal wall (the wall between the left and right ventricles) is vertical (see the A4C reference image above).
- For an A4C clip, ensure that the orange marker on Kosmos Torso-One is pointed towards the scanning table to avoid acquiring a mirrored view.
- Once you have obtained a proper A4C view, rotate the probe 90 degrees counterclockwise to find the A2C view.
- Ask the patient to hold their breath while recording the clip.
- Make sure to review the results for correctness of ED/ES frames and LV contours and, using the Kosmos editing tool, adjust as needed.

Error conditions and system notifications for Kosmos AI-assisted EF workflow

- If the resulting EF scan (initial and/or with edits) is out of the 0%-100% range, you will not be able to save the EF result in the report or export/archive the scan.

You will first need to edit the ED/ES frames and corresponding LV contours to produce a valid EF. Then you will be able to save the results and export/archive the scan.

- Kosmos will prompt you to edit the results or scan again if any of the following conditions are met:
 - ESV > 400 ml
 - EDV > 500 ml
 - Difference between A4C and A2C EF is more than 30%

Kosmos cardiac measurements

⚠ Do not rely upon Kosmos cardiac measurements as the sole diagnostic criteria. Whenever possible, use Kosmos Cardiac measurements in conjunction with other clinical information.

Kosmos Cardiac Calculations package provides the tools to assess cardiac structure and function. Kosmos cardiac measurements are available in B-mode, Doppler and M-mode.

While in Exam Review, cardiac calculations and annotation tools can be used to perform cardiac measurements.

To access the Cardiac Calculation tools:

- * From the **Exam Review** screen, tap **Calc**.

To access the Annotation tools:

- * From the **Exam Review** screen, tap **Annotate**.

For a list of measurements, reference **TABLE 4-6, "Cardiac measurements by mode," on page 46**.

While reviewing the Doppler cine, you can:

1. Perform Doppler measurements
 - VTI: When you tap on **VTI**, you will have the option to select **Auto** or **Manual** VTI trace.
 - If you select **Auto**, tap the signal that you want to trace and the device will trace the signal automatically.
 - If you select **Manual**, you will be prompted to manually trace the signal with your finger.
 - Edit the VTI trace by moving the control points.
 - Choose a different peak by double tapping it.

📖 Please note that auto trace is not available for Mitral Valve VTI in PW and CW tracing. Auto tracing is only available in Annotations or for LVOT VTI (PW) and AV VTI (CW).

- PHT and Delta Velocity: Move to two end points of the calipers to the appropriate location on the Doppler spectrum.
- Velocity and PG: Move the cursor to the desired location.

- You can perform three PHT, three Velocity, and three VTI measurements per image/clip.
 - Only three frames in 2D cine loops can be placed.
 - Only three VTI measurements at a time.

 You will receive a notification that the measurement is full in the report if you try to place a 4th measurement. You can delete a measurement in the report to make room for a new measurement.

- Add Annotations:
 - Text
 - Marker
- Move the baseline
- Invert the Doppler Spectrum
- View measurements by tapping the **Report** icon 
 - When viewing the report, the last measurement taken is the default measurement. However, by clicking Last, the device will calculate the average value or provide the maximum value of each measurement.

TABLE 4-6. Cardiac measurements by mode

2D Measurements	
PLAX	RVIDd, IVSd, LVIDd, LVPWd, LVIDS, LA diam, LVOTd
Right Heart	RV basal, RV mid, RV length
Mitral Valve	MV Annulus diameter
Aortic Valve	Annulus, Sinus, ST junction, Ascending AO, Vena Contracta, LVOT diameter
IVC	IVC min, IVC max, RAP
Doppler Measurements	
PW	Right Heart: PV AcT (Acceleration Time) Mitral Valve: MV VTI (PW), E wave Velocity, Deceleration Time, A wave Velocity Aorta: LVOT VTI (PW) Diastology: E wave Velocity (PW), A wave Velocity, Deceleration Time (PW) Aortic Valve: LVOT VTI (PW)
CW	Right Heart: TR (CW), PAEDP (CW), PR (CW) Mitral Valve: MV VTI (CW), Pressure Half Time (CW) Aortic Valve: AV VTI (CW), Peak AV Velocity, Pressure Half Time (CW) Diastology: TR (CW)
TDI	Right Heart: TV annulus s' Mitral Valve: e' -point (m/s), a' -point (m/s) Diastology: e' -point (m/s), a' -point (m/s)
M-mode Measurements	
M-mode	EPSS, TAPSE, MAPSE, IVC min, IVC max, HR, RAP
PLAX-M-Mode	RVIDd, IVS, LVIDd, LVPW, LVIDS, AO dist, LA dist

Kosmos AI FAST

 Do not rely solely on the AI FAST tool for diagnostic purposes. Kosmos AI FAST assists users by providing a quick orientation to the anatomy of the abdomen. Users should exercise their judgment to ensure annotations are correct.

Using Kosmos AI for FAST Exam

Kosmos AI FAST provides automated anatomical labeling and view identification for the FAST exam in real time. The labels that appear while scanning are only there while you are scanning; after you save the image or clip, the labels will no longer be there.

Reference **TABLE 4-7** for a list of anatomical structures in each FAST Exam imaging views.

TABLE 4-7. Anatomical structures for FAST Exam

FAST View	Anatomical Structures
RUQ	Liver, right kidney, diaphragm, gallbladder, IVC Potential fluid space: hepatorenal space, pleural
LUQ	Spleen, left kidney, diaphragm Potential fluid space: splenorenal space, pleural space
SUB	Heart, diaphragm, liver Potential fluid space: pericardium
A5	Liver, transverse aorta, transverse IVC
IVC	Liver, sagittal IVC
Aorta	Liver, sagittal aorta
A4C	Heart
A2C	Potential fluid space: pericardium
PLAX	
PSAX	Heart
SUB2	Liver, heart, IVC, aorta Potential fluid space: pericardium

To enable Kosmos AI FAST:

- * In Abdominal preset, tap **AI**.

When scanning with Torso-One probe, Kosmos AI FAST feature is only available in the Abdomen preset.

Kosmos Bladder AI

- Kosmos Bladder feature assists users by automatically estimating bladder volume following Transverse and Sagittal view acquisition. Kosmos Bladder provides instructions to identify the bladder, assists users with orientation and real-time probe animations, and automatically calculates estimated bladder volume.

⚠ Do not rely solely on Kosmos Bladder AI for diagnostic purposes.

- Using the Bladder Preset, the depth can be adjusted by tapping the **Body Type** icon located in the left hand side of the screen. The gain can also be adjusted by tapping the **Gain** icon which is next to the **Body Type** icon.

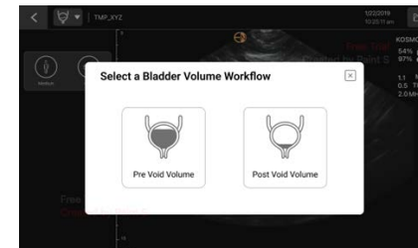
- The system walks through acquiring a Transverse view, first, then a Sagittal view.
- On screen directions are located at the bottom of the screen, below the live ultrasound image.
- When the system is ready to move to the next step, the reference videos will automatically play then minimize to the upper left hand corner of **Imaging** screen.

Accessing the Bladder Preset

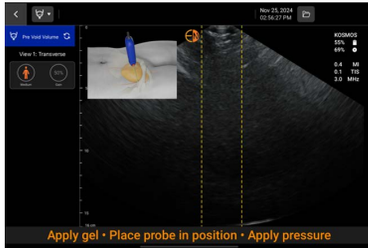
1. From the **Home** Screen, tap the **Bladder icon**  to begin the Bladder Volume Workflow.
2. When prompted, select the appropriate workflow from the dialog box by tapping on **Pre Void Volume** or **Post Void Volume**. For Pre Void Volume, follow the next set of instructions. Otherwise, skip to the **Post Void Volume** section for instructions

Pre Void Volume

1. When prompted, tap **Pre Void Volume**.

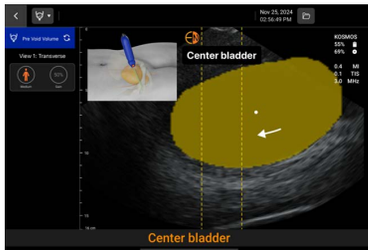


- Following the on-screen instructions, apply ultrasound gel, place the probe into position, and apply pressure.



The reference video in the upper left corner of the screen demonstrates proper placing of the probe in relation to the patient's body position.

- Once the system detects a bladder, follow the on-screen instructions and center the bladder. This can be done by moving the probe so the white dot lies in between the vertical lines.



- If the system cannot detect a bladder, it will prompt you to reposition the probe and try again. To keep scanning, tap **Keep Scanning**.
- If the bladder cannot be detected, using the Pre Void Volume workflow, tap **Pre Void Volume** in the upper left hand side of the screen to select **Post Void Volume**. Follow the instructions located in Post Void Volume.

- Once the bladder is centered, the color will change from yellow to green.

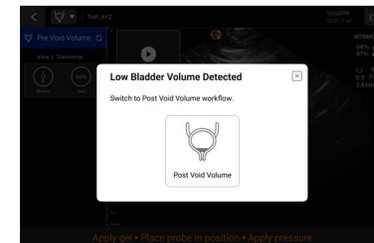
- Follow the on screen instructions to fan the probe until the green ring completely surrounds the center white ring.



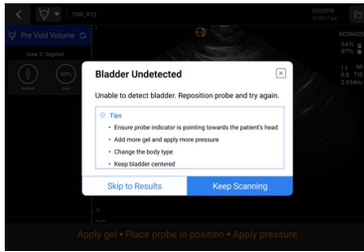
- Once the Transverse view has been acquired, follow the video show on on-screen prompt and rotate the probe by 90 degrees.



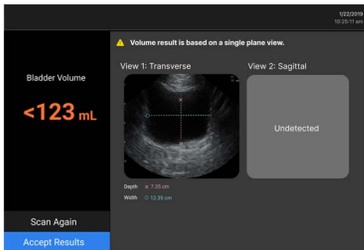
- Repeat steps 1-6 to acquire the sagittal view.
 - If the system detects a low Bladder volume, it will prompt you to use the Post Void Volume workflow. To change the workflow, tap the **Post Void Volume** icon shown in the prompt. To continue with Pre Void Volume, tap the "X" located in the upper right hand corner of the dialog box.



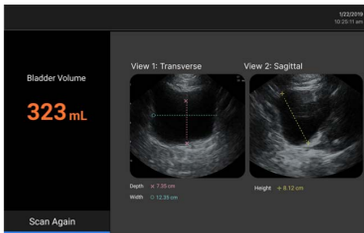
- b. If the Bladder cannot be detected, the system will prompt you to skip or keep scanning.
To skip, tap **Skip to Results**. Otherwise, tap **Keep Scanning**.



- c. By skipping to the **Results** screen, the system will show the approximate volume with a caution statement indicating the result only used a single view.
To save the results, tap **Accept Results** then **Save**.
To repeat scanning, tap **Scan Again**.

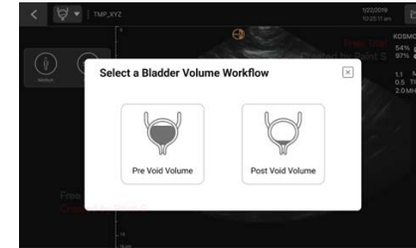


7. After acquiring the Sagittal view, the system will automatically show the **Results** screen. To save the results, tap **Accept Results** then **Save**. To repeat scanning, tap **Scan Again**.

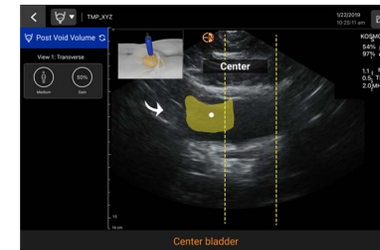


Post Void Volume

1. When prompted, tap **Post Void Volume**.

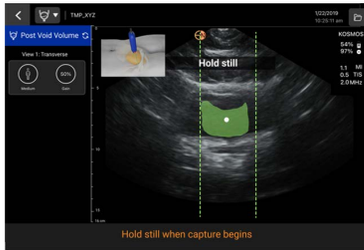


2. Following the on-screen instructions, apply ultrasound gel, place the probe into position, and apply pressure.



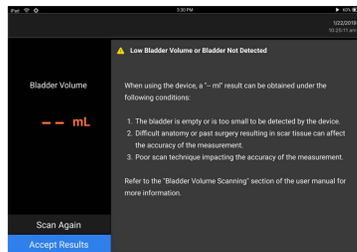
The reference video in the upper left corner of the screen demonstrates proper placing of the probe in relation to the patient's body position.

- Once the system detects a bladder, follow the on-screen instructions and center the bladder. This can be done by moving the probe so the white dot lies in between the vertical lines.



If the system cannot detect a bladder, it will prompt you to reposition the probe and try again. To skip, tap **Skip to Results**. Otherwise, tap **Keep Scanning**.

- If **Skip to Results** is tapped, the system will show the volume as "-- mL" and a caution statement indicating a low bladder volume or bladder not detected. To save the results, tap **Accept Results** then **Save**. To repeat scanning, tap **Scan Again**.



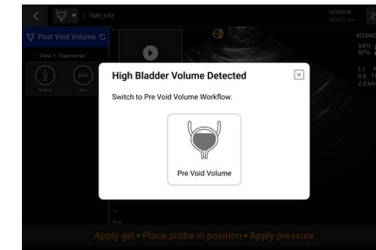
- Once the bladder is centered, the color will change from yellow to green.
- Follow the on screen instructions to hold the probe until the green ring completely surrounds the center white ring.

- Once the Transverse view has been acquired, follow the video show on-screen prompt and rotate the probe by 90 degrees.

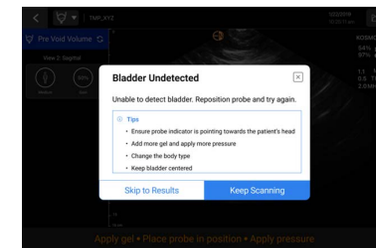


- Repeat Steps 1-6 to acquire the Sagittal view.

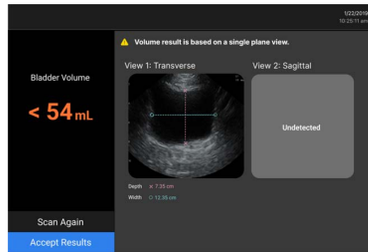
- If the system detects a high Bladder volume, it will prompt you to use the Pre Void Volume workflow. To change the workflow, tap the **Pre Void Volume** icon shown in the prompt. To continue with Post Void Volume, tap the "X" located in the upper right hand corner of the dialog box.



- If the Bladder cannot be detected, the system will prompt you to skip or keep scanning. To skip, tap **Skip to Results**. Otherwise, tap **Keep Scanning**.



- c. By skipping to the **Results** screen, the system will show the approximate volume with a caution statement indicating the result only used a single view. To save the results, tap **Accept Results** then **Save**. To repeat scanning, tap **Scan Again**.



- 8. After acquiring the Sagittal view, the system will automatically show the **Results** screen. To save the results, tap **Accept Results** then **Save**. To repeat scanning, tap **Scan Again**.

Auto Bladder Imaging Controls

Auto Bladder is only available in 2D/B-mode.

Body Type

Body Type controls the image depth.

- * To adjust the body type, tap on the icon.

Gain

Gain controls the brightness of the image.

- * To adjust the gain, tap on the icon.

Kosmos vascular calculations

Do not rely upon Kosmos vascular measurements as the sole diagnostic criteria. Whenever possible, use Kosmos Vascular measurements in conjunction with other clinical information.

Kosmos Vascular Calculations package provides the tools to assess vascular structure and function. Kosmos vascular measurements are only available in 2D mode and PW Doppler mode while scanning with Kosmos Lexsa.

Reference **TABLE 4-8, "Vascular measurements and calculations by mode,"** on page 56 for a list of vascular measurements.

Please note that DICOM SR is not available for the Vascular Calculations Report.

TABLE 4-8. Vascular measurements and calculations by mode

Measurements & Calculations in 2D and PW Doppler Modes	
Venous	Peak Systolic, End Diastolic, Reflux Time, Vessel Diameter, Temporal Average Max, Temporal Average Mean, VTI (grafts)
Arterial	Peak Systolic, End Diastolic, VTI, Vessel Diameter, Temporal Average Max, Temporal Average Mean
Calculations	S/D Ratio, Pulsatility Index, Resistance Index, Flow Volumes

-- End of section --

Reviewing an Exam

Once you have completed an exam, you cannot add any images to it; however, before archiving the exam, you can add, edit, and delete any annotations you have saved.

Once the archive process begins, you will not be able to make edits to the exam.

Starting an exam review

- To start a review during an exam, tap the **Exam review**  icon.
- To start a review for a completed exam, do one of the following:
 - From the **Home** screen, tap **Exams**, then tap the exam you would like to review.
 - From the list of patients, find the patient, then tap the exam you would like to review.

Annotating images and clips

You can add annotations during the exam when the image is frozen or after you have completed the exam. All annotations are saved as overlays on the image or clip.

 Once you have archived an image or clip, you cannot annotate it.

Navigating to the Edit Image screen

While scanning a patient:

1. Tap the **Freeze**  icon.
2. Add your annotations.
3. Tap the **Save image**  or **Save clip**  icon.

After scanning a patient:

1. Tap the **Exam review**  icon.
2. Tap the image/clip you want to annotate.
3. Tap the **Edit**  icon.

Reviewing an Exam

From the **Home** screen:

1. Tap **Exam**.
2. Tap the exam row that you want to edit.
3. Tap the clip you want to annotate.
4. Tap the **Edit**  icon.

From the **Patient** screen:

1. Tap a patient from the list.
2. Tap the exam.
3. Tap the image/clip you want to annotate.
4. Tap the **Edit**  icon.

Annotation tools

Annotations can be added to individual images and clips.

When you add an annotation (text, measurements, arrow, area) to a clip or a cine, they persist through all frames.

You can also hide the overlay of the annotations you make by tapping the **Hide overlay**  icon on saved images and clips.

Measuring with the caliper tool

You can add up to two calipers per image/clip.

When a caliper is not selected and you start dragging one of the two end points of the caliper, the caliper will become selected and will resize based on where you are dragging it.

To place a measurement:

1. From the **Edit image** or **Edit clip** screen, tap **Distance**, and a caliper appears in the center of the image or clip.
2. Tap to select the **caliper**.

 Notice that the distance of the caliper displays in the legend on the upper left side of the screen. If you have multiple calipers, they display in different colors.

3. To resize the caliper, tap and drag one of its end points.
4. To move the caliper, tap anywhere on the caliper except the two end points.
5. To clear the caliper, tap an empty area outside it.

Zooming in and out

Use two fingers to pinch and expand the image area. To return to “normal” tap the **magnifying glass**. Also, zoom factor is shown near magnifying glass as well as orange color of depth scale along the side. You are able to freeze the image while zoomed (and can zoom out and zoom in frozen state).

Deleting annotations

- * To delete one annotation, tap the annotation to select it, then tap **Delete**.
- * To delete all the annotations you have made, tap **Clear ALL**.

Managing images and clips

Filtering images and clips

When you review an exam, all the images and clips, regardless of the scan type (lung, heart, abdomen) are visible in the thumbnail list.

You can filter images and clips in the following ways:

- Drag and pull the thumbnail list down to reveal the filter options.
- Tap the **Filter** icon on top of the thumbnail list to reveal the filter options.
- Tap the **More options**  icon in the title bar, and tap **Filter images and clips**. When the filter options are visible, a **blue check** icon will be shown next to **Filter images and clips**.

When you select a filter, only the tagged images/clips are visible in the thumbnail list. You can tag images/clips by tapping the **star** icon under each image/clip in the thumbnail list so the star turns yellow.

To dismiss the filters you have selected, tap the **More options**  icon, then tap the **Filter images and clips** again to remove the filters.

Selecting images and clips

To select images and clips:

1. Tap the **More options**  icon, and tap **Select images and clips**.
2. Select the images and clips you want. A gray check will appear in the top right corner of the thumbnail.
3. Optionally, tap the check on the thumbnail; it turns red, and a numbered circle displays to indicate how many images and clips you have selected. To clear the red check, tap it again.
4. To clear the selections, tap the **More options**  icon, and tap **Select images/clips**.

Trimming and saving images and clips

To trim and save a clip:

1. Tap the **Freeze**  icon.
2. Move the right and left end points of the cine clip.
3. Tap the **Clip**  icon.

To trim and save an image:

1. From the Exam Review screen, find the saved clip.
2. Tap **Edit**.
3. Move the right and left end points of the image.
4. Tap **Save**.

Deleting images and clips

To delete selected images and clips:

1. Tap the **More options**  icon, and tap **Select images/clips**.
2. Select the images and clips you want to delete.
3. Tap **Delete** and, when prompted, tap **OK**.

Reviewing and editing a report

 Reports are not yet encapsulated in the DICOM file; you can only see images and clips at this review step.

The exam report lets you review patient and exam information, text notes, audio notes, pictures that were taken, images, and clips in the exam report.

Opening a report

- * To open a report, tap **Report**.

Editing a report

Once you've opened the report, each section is expanded for your review. You can collapse each section by tapping the arrow button. Just tap the arrow button to expand the section again.

You can edit each section of the report with the exception of the patient information. This is read-only and cannot be changed.

Editing exam information

The exam information section displays the exam related information that was entered before the scan.

To edit the exam information:

1. Tap the **Edit**  icon.
2. Make any necessary updates to the section.

Adding a text note

You can add text notes that will display under each scan.

To add a text note:

1. Tap the **Add text note** icon. A text box, date and time label appear under the last text note.
2. Using the keyboard, type the note.
3. Tap **Done**.

Editing a text note

To edit a text note:

1. Tap an existing text note. A text box containing the existing note and the keyboard displays.
2. Using the keyboard, edit the text note.
3. Tap **Done**.

Deleting a text note

To delete a text note:

1. Long press an existing text note. A delete button displays.
2. Tap **Delete** and, when prompted, tap **OK**.

Exporting images and clips to a USB drive

When exporting an images and clips, use a micro USB or adapter.

You can export images and clips from one exam or multiple exams.

 To protect patient data, take appropriate precautions when exporting patient data to a USB drive.

To export images and clips from one exam to a USB drive:

1. From the **Home** screen, tap **Exams**.
2. Tap a row to select an exam.
3. Tap the **bookmark** icon under each of the thumbnails you would like to export. (This is an optional step and only useful if you would like to export some but not all images and clips.)
4. Connect the USB drive using the USB-c adapter.
5. Tap **Export**. A dialog box appears.
6. Select the file type and whether you want all images and clips exported or only the tagged images and clips.
7. Tap **OK** to start exporting to USB drive.

To export images and clips from multiple exams to a USB drive:

1. From the Home screen, tap **Exams**.
2. Tap the circles next to each exam you would like to export.
3. Connect the USB drive using the USB-c adapter.
4. Tap the **Export**  icon on the top of the screen. A dialog box appears.
5. Select the file type and whether you want all images and clips exported or only the tagged images and clips.
6. Tap **OK** to start exporting to USB drive.

The following table is a legend for the exporting icons.

	Exam is waiting to be exported.
	Export is in progress
	Export is complete.
	Export failed.

Completing an exam review

To complete an exam:

1. Tap **Complete**.
2. When prompted, click **OK**.

Archiving an exam to a PACS server

After completing an exam, you can archive it to a PACS server. Once an exam is archived, you cannot edit it.

For more information about setting up a PACS server, see ["DICOM" on page 17](#).

For each EF scan, multiple images/clips are archived and exported.

The following table is a legend for the archiving icons.

	Exam is waiting to be archived.
	Archive is in progress.
	Archive is complete.
	Archive failed.

You can archive an exam either from the **Exam list** or the **Exam review** screens.

To archive an exam from the **Exam list** screen:

1. From the **Exam List** screen, tap to select the completed exam(s) you want to archive.
2. Tap the **Archive**  icon. The complete exam is archived according to the default archive options. For more information, see ["DICOM" on page 17](#).

To archive an exam from the **Exam review** screen:

1. From the **Exam review** screen, tap **Archive**.
2. From the Archive exam to PACS server screen, select which images and clips you want to archive and if you would like to include a report.
3. Click **OK** and, when prompted, click **OK** again.

Deleting an exam

To delete an exam from the **Exam list**:

1. Tap the **left** icon next to the exam you would like to delete. The icon turns into a **check mark** .
2. Tap the **Trash**  icon.
3. When prompted, tap **OK**.

To delete an exam while reviewing it:

1. Tap the **More options**  icon.
2. Tap **Delete the exam**.
3. When prompted, click **OK**.

-- End of section --

Kosmos probe sheaths

Where fluid contamination is possible, cover the probe being used (Kosmos Torso-One or Kosmos Lexsa) with an appropriate sterile sheath from CIVCO, which will promote asepsis and minimize cleaning.

▲	Be aware that some patients have a latex allergy. Some commercially available Kosmos probe covers contain latex.
▲	To prevent cross-contamination, use sterile transducer sheaths and sterile coupling gel for clinical applications contacting compromised skin.
▲	Some sheaths contain natural rubber latex and talc, which can cause allergic reactions in some individuals.
▲	Use market-cleared sheaths for clinical applications when a Kosmos probe is likely to be splashed or splattered with blood or other bodily fluids.
▲	Use market-cleared, sterile sheaths and sterile coupling gel to prevent cross-contamination. Do not apply the sheath and coupling gel until you are ready to perform the procedure. After use, remove and discard the single-use sheath, and clean and disinfect the Kosmos probe using an EchoNous-recommended high-level disinfectant.
▲	After inserting the Kosmos probe into the sheath, inspect the sheath for holes and tears.

Ultrasound transmission gels

▲	Some ultrasound gels may cause an allergic reaction in some individuals.
▲	To prevent cross-contamination, use single-use gel packs.

EchoNous recommends the use of:

- Aquasonic 100 Ultrasound Gel, Parker
- Aquasonic Clear Ultrasound Gel, Parker
- SCAN Ultrasound Gel, Parker

Kosmos probe storage

▲	To prevent cross-contamination or unprotected exposure of personnel to biological material, containers used to transport contaminated Kosmos probes should carry an ISO biohazard label.
---	--

Daily storage

Kosmos is intended to be used and stored in normal ambient conditions inside a medical facility. In addition, the packaging provided with the device may be used for long-term storage.

Storage for transport

Kosmos is intended to be hand held for easy transport. Users may use the packaging supplied with the device for transport. Consult your EchoNous sales representative for information on approved bags and other accessories.

Transducer element check

Every time a Kosmos probe is connected, a test is run automatically to check for the integrity of the transducer elements. The test reports to the user whether all transducer elements are functioning properly (successful test), or whether failures were detected.

The same test runs automatically when the Kosmos App boots up with Kosmos probe connected.

-- End of section --

Cleaning and disinfecting

General cautions

	The provided cleaning instructions are based on requirements mandated by the U.S. Food and Drug Administration. Failure to follow these instructions may result in cross contamination and patient infection.
	Cleaning and disinfection instructions must be followed when using a transducer cover or sheath.
	Some reprocessing chemicals may cause an allergic reaction in some individuals.
	Ensure that cleaning and disinfecting solutions and wipes are not expired.
	Do not allow cleaning solution or disinfectant into the tablet or Kosmos probe connectors.
	Wear the appropriate personal protective equipment (PPE) recommended by the chemical manufacturer, such as protective eye wear and gloves.
	Do not skip any steps or abbreviate the cleaning and disinfecting process in any way.
	Do not spray cleaners or disinfectants directly on tablet surfaces or on the tablet and Kosmos probe connectors. Doing so may cause solution to leak into Kosmos, damaging it and voiding the warranty.
	Do not attempt to clean or disinfect the tablet, Kosmos probes, or Kosmos probe cable using a method that is not included here or chemical not listed in this guide. Doing so can damage Kosmos and void the warranty.
	Do not pull the cable of the Kosmos probe while holding or disinfecting the device. Pulling on the cable may cause damage to the probe.

Tablet

	The tablet is not sterile when shipped; do not attempt to sterilize it.
	To avoid electrical shock, before cleaning, turn off the tablet and disconnect it from the power supply.

Cleaning

Avoid spraying the cleaning and disinfection solutions directly onto the tablet. Instead spray onto a non-abrasive cloth and then gently wipe. Ensure that all excess solution is wiped off and not left on the surface after cleaning. The following cleaning and disinfection method must be followed for the tablet.

1. Disconnect the Kosmos probe from the tablet.
2. Remove any accessories, such as Kosmos Link or power supply.
3. Using a wipe carefully clean the screen and all other areas of the tablet. Choose an EchoNous-approved wipe from the list in **TABLE 7-1, "Presaturated wipes," on page 69.**
4. If necessary, clean the tablet with additional wipes to remove all visible contaminants.

Kosmos Link

	The Link is not sterile when shipped; do not attempt to sterilize it.
	To avoid electrical shock, before cleaning, disconnect the Link and disconnect it from the power supply.

Avoid spraying the cleaning and disinfection solutions directly onto the Link. Instead spray into a non-abrasive cloth and gently wipe. Ensure that all excess solution is wiped off and not left on the surface after cleaning. The following cleaning and disinfection method must be followed for the Link.

1. After each use, disconnect the USB cable from the tablet.
2. Disconnect the probes from the underside of the Link.
3. Using a wipe from an approved presaturated disinfectant wipe, carefully wipe all the areas of the Link. Choose an EchoNous approved wipe from the list provided in **TABLE 7-1, "Presaturated wipes," on page 69.**
4. If necessary, clean the Link with additional wipes to remove all visible contaminants.

	After disinfection, examine the Link for cracks, and if damage exists, discontinue use of the Link and contact EchoNous Customer Support.
---	---

TABLE 7-1. Presaturated wipes

Product	Company	Active Ingredients	Contact Condition
Duo ULT	Tristel	Chlorine Dioxide 100% (Proprietary formulation)	30 seconds wet contact time for disinfection
Sani-Cloth Super	PDI Inc.	Iso Propyl Alcohol 55.5% Quaternary Ammonium compounds, C12-18-alkyl[(ethoxyphenyl) methyl] dimethyl, chlorides 0.25% n-alkyl dimethyl benzyl ammonium chloride 0.25%	5 minutes wet contact time for disinfection

▲	Do not use a chlorine dioxide-based agent, such as Tristel Duo ULT, on Kosmos Bridge or Kosmos Link because it may corrode the aluminum housing.
■	A complete guide to compatible cleaning and disinfection agents can be found online at www.echonous.com/resources/mediatype-chemical-compatibility-guides/

Kosmos probes

Cleaning

The following cleaning instructions must be followed for Kosmos Torso-One and Kosmos Lexsa. Kosmos probes must be cleaned after each use. Cleaning Kosmos probes is an essential step before effective disinfection.

Before cleaning Kosmos Torso-One and Kosmos Lexsa, read the following warnings and cautions.

▲	Always disconnect the probe from the tablet or Link before cleaning and disinfecting.
▲	After cleaning, you must disinfect Kosmos probes by following the appropriate instructions.
▲	Always wear protective eye wear and gloves when cleaning and disinfecting any equipment.
▲	Use only EchoNous-recommended wipes. Using a non-recommended wipe can damage the Kosmos probe and void the warranty.
▲	When cleaning and disinfecting Kosmos probes, do not allow any fluid to enter electrical connections or metal portions of the USB connector.
▲	The use of a cover or sheath does not preclude proper cleaning and disinfecting of a Kosmos probe. When choosing a cleaning and disinfecting method, treat Kosmos probes as if a cover was not used in the procedure.

To clean probes:

1. Disconnect the Kosmos probe from the tablet.
2. Remove any accessories attached to, or covering the Kosmos probe, such as a sheath.
3. At point of use, wipe Kosmos probe with an approved presaturated wipe.
4. Prior to disinfecting the Kosmos probe, remove all ultrasound gel from the Kosmos probe face by using an approved presaturated disinfectant wipe. Choose an EchoNous-approved wipe from the list in **TABLE 7-1**.
5. Using a new wipe, remove any particulate matter, gel, or fluids that remain on the Kosmos probe using a new presaturated wipe from **TABLE 7-1**, "Presaturated wipes," on page 69.
6. If necessary, clean the Kosmos probe with additional wipes to remove all visible contaminants.
7. Before continuing with disinfection, ensure the Kosmos probe is visibly dry.

Disinfecting (intermediate-level)

Use the following steps to disinfect a Kosmos probe whenever it has not come into contact with non-intact skin, or intact mucous membranes (non-critical use). Before performing the following steps, read the following warnings and cautions.

▲	For low-level and intermediate-level disinfection, EchoNous validated its disinfection with intermediate-level disinfection.
▲	Always disconnect the Kosmos probes before cleaning and disinfecting.
▲	Always use protective eye wear and gloves when disinfecting any equipment.
▲	Before disinfecting, clean Kosmos probes by following the appropriate instructions to remove all gels, fluids, and particulates that may interfere with the disinfection process.
▲	Use only EchoNous-recommended disinfectants. Using a non-recommended disinfecting wipe can damage the Kosmos probe and void the warranty.

To disinfect Kosmos Probes (intermediate-level):

1. After cleaning, choose an intermediate-level disinfectant from the list in **TABLE 7-1**, "Presaturated wipes," on page 69, and observe the recommended minimum wet contact time.
2. With a new wipe, clean the cable and the Kosmos probe, starting from the exposed cable, wiping toward the Kosmos probe head.
3. Observe the required wet contact time. Monitor the Kosmos probe for wet appearance. Use at least three wipes to ensure effective disinfection.

- Before reusing the Kosmos probe, ensure the Kosmos probe is visibly dry.

▲	Check the Kosmos probe for damage, such as cracks, splitting, or sharp edges. If damage is evident, discontinue using the Kosmos probe, and contact your EchoNous representative.
---	---

Disinfecting (high-level)

Use the following steps to high-level disinfect the Kosmos probes whenever it has come into contact with intact mucous membranes, or non-intact skin (semi-critical use). High-level disinfection of Kosmos probes typically uses an immersion method with high-level disinfectants or chemical sterilant.

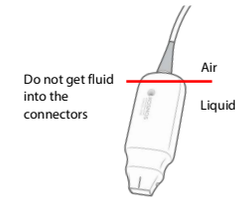
Before performing the following steps, read the following warnings and cautions.

▲	Always disconnect Kosmos probes from tablet during cleaning and disinfection.
▲	Before disinfection, clean the Kosmos probe by following the appropriate cleaning instructions in Cleaning to remove all gels, fluids, and particulates that may interfere with the disinfection process.
▲	Always use protective eye wear and gloves when disinfecting any equipment.
▲	When disinfecting Kosmos probes, do not allow any fluid to enter electrical connections or metal portions of the USB.
▲	Do not attempt to disinfect Kosmos probes using a method that is not included in these instructions. This can damage the Kosmos probe and void the warranty.
▲	Use only EchoNous-recommended disinfectants. Using a non-recommended disinfecting solution or incorrect solution strength can damage the Kosmos probe and void the warranty.
▲	If the Kosmos probe has come into contact with intact mucous membranes or non-intact skin (semi-critical use), use the high-level cleaning and disinfection procedure.

To disinfect Kosmos probes (high level):

- After cleaning, choose a high-level disinfectant that is compatible with Kosmos probes. For a list of compatible disinfectants, see **TABLE 7-1, "Presaturated wipes," on page 69.**
- Test the solution strength by using a Cidex OPA test strip. Ensure that the solution is not older than 14 days (in an open container) or 75 days (from a just opened storage container).
- If a pre-mixed solution is used, be sure to observe the solution expiration date.

- Immerse Kosmos probe into the disinfectant as shown below. Kosmos probes may be immersed only up to the immersion point shown. No other part of the Kosmos probe, such as cable, strain relief, or connectors should be soaked or immersed in fluids.



- Refer to **TABLE 7-1, "Presaturated wipes," on page 69** for duration of immersion and contact temperature.
- Do not immerse Kosmos probe longer than the minimum time needed for semi-critical level of disinfection.
- Rinse Kosmos probe for at least one minute in clean water up to the point of immersion to remove chemical residue. Do not soak or immerse any other part of the Kosmos probe, such as the cable, strain relief, or connector.
- Repeat, rinsing three times to ensure proper rinsing.
- Air dry or use a soft sterile cloth to dry the Kosmos probe until visibly dry.
- Wipe the strain relief and first 18 inches (45 cm) of the Kosmos probe cable with an approved wipe from the list in **TABLE 7-1, "Presaturated wipes," on page 69.**
- Examine the Kosmos probe for damage such as cracks, splitting, or sharp edges. If damage is evident, discontinue using the Kosmos probe, and contact your EchoNous representative.

TABLE 7-2. Disinfectant solutions for Kosmos probe immersion

Product	Company	Active Ingredients	Contact Condition
Cidex OPA Solution	Advanced Sterilization Product	Products 0.55% ortho phthaldehyde	12 minutes at 20°C

- Check the expiration date on the bottle to ensure the disinfectant has not expired. Mix or check that the disinfection chemicals have the concentration recommended by the manufacturer (for example, a chemical strip test).
- Check that the temperature of the disinfectant is within the manufacturer's recommended limits.

Guidelines for AR (automated reprocessors)

▲	Always disconnect the Kosmos probe before cleaning and disinfecting.
▲	Ensure cable insulation is intact before and after cleaning.
■	The EMC suppressor on probes should be inside the trophon2 chamber below the cable clamp during disinfection.

All Kosmos probes are compatible with Nanosonic™ Trophon2 System. Refer to the Trophon®2 user guide for detailed instructions related to disinfection of ultrasound probes.

For questions related to compatibility with other AR systems, contact EchoNous support.

Recycling and disposal

▲	Do not incinerate or discard Kosmos Link in general waste at end of life. The lithium battery is a potential environmental and fire safety hazard.
▲	The lithium ion battery inside Kosmos Link may explode if exposed to very high temperatures. Do not destroy this unit by incinerating or burning. Return the unit to EchoNous or your local representative for disposal.

The system should be disposed of in an environmentally responsible manner in compliance with federal and local regulations. EchoNous recommends taking Kosmos probes and Kosmos Link to a recycling center which specializes in the recycling and disposal of electronic equipment.

In cases where a Kosmos probe or Kosmos Link have been exposed to biologically hazardous material, EchoNous recommends using biohazard containers and in compliance with federal and local regulations. Kosmos probes and Kosmos Link should be taken to a waste center which specializes in the disposal of biohazard waste.

Troubleshooting

Preventive inspection, maintenance, and calibration

- Kosmos does not require any preventative maintenance or calibration.
- Kosmos does not contain any serviceable parts.

■	If Kosmos is not functioning as designed and intended, contact EchoNous customer support.
▲	Do not open Kosmos Link enclosure.

--End of section --

Safety

Electrical safety

References

IEC 60601-2-37: 2015 Medical electrical equipment – Part 2-37: *Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment*

ANSI AAMI ES 60601-1: 2012 Medical electrical equipment. Part 1: *General requirements for basic safety and essential performance* – IEC 60601-1:2012, Edition 3.1

IEC 60601-1-2:2021: AMD1:2020 Medical electrical equipment – Parts 1-2: *General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests*

IEC 62304:2015 Medical device software - *Software life-cycle processes*

ISO 14971:2021 Medical devices - *Application of risk management to medical devices*

10993-1:2020 Biological evaluation of medical devices - *Part 1: Evaluation and testing within a risk management process*

Lang, Roberto M., et al. *Recommendations for chamber quantification: a report from the American Society of Echocardiography's Guidelines and Standards Committee and the Chamber Quantification Writing Group, developed in conjunction with the European Association of Echocardiography, a branch of the European Society of Cardiology*. Journal of the American Society of Echocardiography 18.12 (2005): 1440-1463.

Lang, Roberto M., et al. *Recommendations for cardiac chamber quantification by echocardiography in adults: an update from the American Society of Echocardiography and the European Association of Cardiovascular Imaging*. European Heart Journal-Cardiovascular Imaging 16.3 (2015): 233-271.

Ronneberger, Olaf, Philipp Fischer, and Thomas Brox. *U-net: Convolutional networks for biomedical image segmentation*. International Conference on Medical image computing and computer-assisted intervention. Springer, Cham, 2015.

Labeling symbols

Symbol	EchoNous Description	SDO Title Reference Number Standard
	Indicates device manufacturer. Includes name and address of the manufacturer	Manufacturer Ref. No. 5.1.1 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	Tested to comply with FCC standards	None
	Probes are tested to Type BF protection	TYPE BF APPLIED PART Refer to D1.20 IEC 60601-1 Medical Electrical Equipment - Part 1: General requirement for basic safety and essential performance
	Class II equipment	Class II equipment Ref. No. D.1-9 IEC 60601-1 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
	Safety cautions are identified with this mark on the device.	Caution Ref. No D1.10 IEC 60601-1 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
	Consult instructions for use	Operating instructions Ref. No. D.1-11 IEC 60601-1 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance

Symbol	EchoNous Description	SDO Title Reference Number Standard
	Do not dispose of this product in normal trash or landfill; refer to local regulations for disposal	Separate collection Annex IX Waste Electrical and Electronic Equipment (WEEE) Directive 2012/19/EU of the European Parliament
IPX7	Kosmos Torso-One and Kosmos Lexsa are protected against temporary immersion in water.	IP Code for degree of protection IEC 60529 Degrees of protection provided by enclosures (IP Code)
IP32	The Kosmos Link is protected against ingress of a solid foreign object greater than or equal to 2.5mm in diameter and protected against access to hazardous parts with a finger and protected against direct sprays of water up to 15 degrees from vertical.	IP Code for degree of protection IEC 60529 Degrees of protection provided by enclosures (IP Code)
REF	Part or model number	Catalog number Ref. No. 5.1.6 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
SN	Serial number	Serial number Ref. No. 5.1.7 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements

Symbol	EchoNous Description	SDO Title Reference Number Standard
	Date of manufacture	Date of manufacture Ref. No. 5.1.3 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	Acceptable temperature range XX is generic placeholder for specified temperatures	Temperature limit Ref. No. 5.3.7 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	Acceptable humidity range XX is generic placeholder for specified percentages	Humidity limitation Ref. No. 5.3.8 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	Acceptable atmospheric pressure range XX is generic placeholder for specified kPa	Atmospheric pressure limitation Ref. No. 5.3.9 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	Stack box this way up	This way up Ref. No. 13 ISO 780 Packaging - Distribution packaging - Graphical symbols for handling and storage of packages

Symbol	EchoNouS Description	SDO Title Reference Number Standard
	Indicates direct current	Direct current Ref. No. D.1-4 IEC 60601-1 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
	Indicates alternating current	Alternating current Ref. No. D.1-1 IEC 60601-1 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
LOT	Batch code	Batch code Ref. No. 5.1.5 ISO 15223-1 Medical devices - Symbols to be used with medical device labels, labeling and information to be supplied - Part 1: General requirements
	UL Classified. Medical - General medical equipment as to electrical shock, fire and mechanical hazards only in accordance with ANSI/AAMI ES 60601-1 (2005) + AMD (2012) / CAN/ CSA-C22.2 No. 6060-1 (2008) + (2014). E509516	None
Rx Only	Caution: Federal law restricts this device to sale by or on the order of a physician.	Reference: USA FDA 21 CFR 801.109
	A manufacturer's indication that a device is in conformity with the applicable requirements set out in EU MDR 2017/745 for CE marking, and the Notified Body reference number.	CE marking of conformity Article 20, Annex V EU MDR 2017/745

Symbol	EchoNouS Description	SDO Title Reference Number Standard
	Medical device	Symbol for Medical Device in compliance with EU MDR directive
	UK Conformity Assessed	Symbol for UK Conformity Assessed. MHRA Department of Business, Energy & Industrial Strategy December 31, 2020
	Switzerland Representative	Symbol for Switzerland Representative MU600_00_016e_MB

Contact information

United States



EchoNouS Inc.
8310 154th Avenue NE
Building B, Suite 200
Redmond, WA 98052

Technical Support (toll free): 844-854-0800

Sales (toll free): 844-854-0800

Email (support): support@EchoNouS.com

Web: www.EchoNouS.com

Phone: 844-854-0800

Fax: 425-242-5553

Email (corporate): info@echonouS.com

European Economic Area

Authorized Representative
Advena Ltd
Tower Business Centre
2nd Flr, Tower Street
Swatar, BKR 4013

Malta**Switzerland Authorized Representative**

QUNIQUE GmbH
Bahnhofweg 17
5610 Wohlen
Switzerland

UK Responsible Person

Qserve Group UK, Ltd
49 Greek St, London W1D 4EG,
United Kingdom

Australia Sponsor

LC & Partners Pty Ltd
Level 32, 101 Miller Street
North Sydney, NSW, 2060
Australia
Tel: +61 2 9959 2400

Brazil Authorized Representative

Detentor da Notificação:
VR Medical Importadora e Distribuidora de Produtos Médicos Ltda
Rua Batataes no 391, conjuntos 11, 12 e 13 - Jardim Paulista
São Paulo - SP - 01423-010

CNPJ: 04.718.143/0001-94

SAC: 0800-7703661

Farm. Resp: Cristiane Ap. de Oliveira Aguirre – CRF/SP: 21.079

Notificação ANVISA no: 80102519147

Suporte ao cliente da EchoNous
Entre em contato com o suporte ao cliente:

Telefone: 844-854-0800

Fax: 425-242-5553

E-mail: info@echonous.com

Web: www.echonous.com

Fabricante:

EchoNous, Inc.
8310 154th Ave NE, Edifício B, Suíte 200
Redmond, WA 98052
Estados Unidos da América

Pais de Origem: Estados Unidos da América

ANATEL: 00430-22-14521

Designated Marketing Authorization Holder:

販売名：超音波画像診断装置 KOSMOS Series Plus

管理医療機器

特定保守管理医療機器

一般の名称：汎用超音波画像診断装置（JMDNコード：40761000）

認証番号：306AIBZI00001000

外国指定高度管理医療機器製造等事業者：ECHONOUS, INC. (米国)

選任製造販売業者：有限会社ユーマンネットワーク

2-7-4 Aomi, Koto-ku, the SOHO

Tokyo, 135-0064 Japan

TEL: 03 (5579) 6773

Biological safety

ALARA education program

The guiding principle for the use of diagnostic ultrasound is defined by the “as low as reasonably achievable” (ALARA) principle. The decision as to what is reasonable has been left to the judgment and insight of qualified personnel (users). No set of rules can be formulated that would be sufficiently complete to dictate the correct response to every circumstance. By keeping ultrasound exposure as low as possible, while obtaining diagnostic images, users can minimize ultrasonic bioeffects.

Since the threshold for diagnostic ultrasound bioeffects is undetermined, users are responsible for controlling total energy transmitted into the patient. Reconcile exposure time with diagnostic image quality. To ensure diagnostic image quality and limit exposure time, Kosmos provides controls that can be manipulated during the exam to optimize the results of the exam.

The ability of the user to abide by the ALARA principle is important. Advances in diagnostic ultrasound, not only in the technology but in the applications of that technology, have resulted in the need for more and better information to guide users. The output display tables are designed to provide that important information.

There are a number of variables which affect the way in which the output display tables can be used to implement the ALARA principle. These variables include index values, body size, location of the bone relative to the focal point, attenuation in the body, and ultrasound exposure time. Exposure time is an especially useful variable, because it is controlled by the user. The ability to limit the index values over time supports the ALARA principle.

A generic ALARA education program is supplied with Kosmos (see enclosed ISBN 1-932962-30-1, Medical Ultrasound Safety).

Applying ALARA

The Kosmos imaging mode used depends upon the information needed. B-mode imaging provides anatomical information, while Color-mode imaging provides information about blood flow.

Understanding the nature of the imaging mode being used allows users to apply the ALARA principle with informed judgment. Additionally, the Kosmos probe frequency, setup values, scanning techniques, and experience allow users to meet the definition of the ALARA principle.

The decision as to the amount of acoustic output is, in the final analysis, up to the user. This decision must be based on the following factors: type of patient, type of exam, patient history, ease or difficulty of obtaining diagnostically useful information, and the potential localized heating of the patient due to transducer surface temperatures. Prudent use of Kosmos occurs when patient exposure is limited to the lowest index reading for the shortest amount of time necessary to achieve acceptable diagnostic results.

Although a high index reading does not mean that a bioeffect is actually occurring, a high index reading should be taken seriously. Every effort should be made to reduce the possible effects of a high index reading. Limiting exposure time is an effective way to accomplish this goal.

There are several system controls that the operator can use to adjust the image quality and limit the acoustic intensity. These controls are related to the techniques that a user might use to implement ALARA.

Output display and display accuracy

OUTPUT DISPLAY

Kosmos displays the two bioeffect indices prescribed by IEC 60601-2-37. Medical electrical equipment. Part 2-37: Particular requirements for the safety of ultrasonic medical diagnostic and monitoring equipment.

The thermal index (TI), provides a measure of the expected temperature increase.

THERMAL INDEX

TI is an estimate of the temperature increase of soft tissue or bone. There are three TI categories: TIS, TIB, and TIC. However, since Kosmos is not intended for transcranial applications, the TI for cranial bone at the surface (TIC) is not available for display on the system. The following TI categories are available for display:

- TIS: Soft tissue thermal index. The main TI category. Used for applications that do not image bone.
- TIB: Bone thermal index (bone located in a focal region).

MECHANICAL INDEX

MI is the estimated likelihood of tissue damage due to cavitation. The absolute maximum limits of the MI is 1.9 as set by the *Guidance for Industry and FDA Staff - Marketing Clearance of Diagnostic Ultrasound Systems and Transducers (2019)*.

ISPTA

The Ispta is the Spatial Peak Temporal Average Intensity. The absolute maximum limit of Ispta is 720 mW/cm² as set by the *Guidance for Industry and FDA Staff - Marketing Clearance of Diagnostic Ultrasound Systems and Transducers (2019)*.

OUTPUT DISPLAY ACCURACY

Output display accuracy of the bioeffect indices, MI and TI, is dependent on the uncertainty and precision of the measurement system, engineering assumptions within the acoustic model used to calculate the parameters, and variability in the acoustic output of the systems. EchoNous also compares both internal and third-party acoustic measurements and confirms that both measurements are within recommended display quantization of 0.2 as outlined by the standards.

All MI and TI values displayed on Kosmos will not exceed the maximum global values (listed in the Track 3 acoustic output tables) by more than 0.2.

The accuracy of the MI and TI indices are as follows:

- MI: accurate to within ±25% or +0.2, whichever value is larger.
- TI: accurate to within ±30% or +0.2, whichever value is larger.

See Kosmos Torso-One and Kosmos Lexsa acoustic output tables, **TABLE 8-1** through **TABLE 8-14**.

Kosmos Torso-One acoustic output tables

TABLE 8-1. Transducer: Kosmos Torso-One, operating mode: B-mode, combined acoustic output table: reportable mode 1 (B-mode) cardiac, body type 2, 16 cm

Index label	MI	TIS		TIB	
		At surface	Below surface	At surface	Below surface
Maximum index value	1.11	0.56		0.56	
Index component value		1: 0.30 2: 0.26	1: 0.30 2: 0.26	1: 0.30 2: 0.26	1: 0.30 2: 0.26
Acoustic parameters	$p_{r,\alpha}$ at z_{pII} (MPa)	1: 1.58			
	P (mW)		1: 41.03 2: 37.03		1: 41.03 2: 37.03
	P_{Tx1} (mW)		1: 30.42 2: 27.46		1: 30.42 2: 27.46
	z_s (cm)		1: 4.27 2: 4.23		
	z_b (cm)				1: 3.93 2: 3.87
	z_{MI} (cm)	1: 4.20			
	$z_{pII,\alpha}$ (cm)	1: 4.20			
	f_{DWF} (MHz)	1: 2.03	1: 2.03 2: 2.03		1: 2.03 2: 2.03
	p_{rr} (Hz)	1: 1589.5			
	s_{rr} (Hz)	1: 28.4			
Other information	n_{pps}	1: 1			
	$p_{a,\alpha}$ at $z_{pII,\alpha}$ (W/cm ²)	1: 91.28			
	$I_{spIa,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm ²)	25.13			
	I_{spIa} at z_{pII} or z_{sII} (mW/cm ²)	42.50			
	p_r at z_{pII} (MPa)	1: 2.13			
Operating Control Conditions	Exam	Cardiac			
	BMI Setting	2			
	Depth	16 cm			

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 Information need not be provided regarding TIC for an TRANSDUCER ASSEMBLY not intended for transcranial or neonatal cephalic uses.
 NOTE 4 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB or TIC.
 NOTE 5 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 6 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 7 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-2. Transducer: Kosmos Torso-One, operating mode: M-mode, acoustic output reporting table: reportable mode 3 M-mode (cardiac, body type: medium, 12 cm depth)

Index label	MI	TIS		TIB	
		At surface	Below surface	At surface	Below surface
Maximum index value	0.43	5.32E-02		0.11	
Index component value		5.32E-02	2.15E-02	5.32E-02	0.11
Acoustic parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	0.70			
	P (mW)		4.55		4.55
	P_{TxT} (mW)		4.11		4.11
	z_s (cm)		5.37		
	z_b (cm)				4.80
	z_{MI} (cm)	5.37			
	$z_{pII,\alpha}$ (cm)	5.37			
	f_{doff} (MHz)	2.72	2.72		2.68
	p_{rr} (Hz)	800			
	s_{rr} (Hz)	N/A			
Other information	n_{pps}	1			
	$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm^2)	52.08			
	$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm^2)	16.71			
	I_{spta} at z_{pII} or z_{sII} (mW/cm^2)	31.29			
	p_r at z_{pII} (MPa)	45.72			
Operating Control Conditions					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-3. Transducer: Kosmos Torso-One, operating mode: M-mode, acoustic output reporting table: reportable mode 4 M-mode (cardiac, body type: medium, 14 cm depth)

Index label	MI	TIS		TIB	
		At surface	Below surface	At surface	Below surface
Maximum index value	0.39	5.33E-02		9.70E-02	
Index component value		5.33E-02	2.12E-02	5.33E-02	9.70E-02
Acoustic parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	0.63			
	P (mW)		4.60		4.60
	P_{TxT} (mW)		4.14		4.14
	z_s (cm)		5.50		
	z_b (cm)				4.97
	z_{MI} (cm)	5.50			
	$z_{pII,\alpha}$ (cm)	5.50			
	f_{doff} (MHz)	2.70	2.70		2.67
	p_{rr} (Hz)	800			
	s_{rr} (Hz)	N/A			
Other information	n_{pps}	1			
	$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm^2)	41.86			
	$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm^2)	13.64			
	I_{spta} at z_{pII} or z_{sII} (mW/cm^2)	38.22			
	p_r at z_{pII} (MPa)	1.06			
Operating Control Conditions					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-4. Transducer: Kosmos Torso-One, operating mode: BC-Mode (max MI, 12cm depth, small ROI, image top)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	1.56	0.37		0.37		0.64
Index component value		1: 6.47E-02 2: 0.30	1: 6.47E-02 2: 0.30	1: 6.47E-02 2: 0.30	1: 6.47E-02 2: 0.30	
Acoustic parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	2: 2.50				
	P (mW)		1: 5.89 2: 27.52	1: 5.89 2: 27.52	1: 5.89 2: 27.52	1: 5.89 2: 27.52
	P_{1x1} (mW)		1: 5.02 2: 24.07	1: 5.02 2: 24.07		
	z_s (cm)		1: N/A 2: N/A			
	z_b (cm)				1: N/A 2: N/A	
	z_{MI} (cm)	2: 1.91				
	$z_{p_{ii},\alpha}$ (cm)	2: 2.00				
	f_{awf} (MHz)	2: 2.65	1: 2.71 2: 2.65	1: 2.71 2: 2.65		
	p_{rr} (Hz)	2: 1248.9				
	s_{rr} (Hz)	2: 31.2				
Other information	n_{pps}	2: 10				
	$I_{pa,\alpha}$ at $z_{p_{ii},\alpha}$ (W/cm ²)	2: 282				
	$I_{spta,\alpha}$ at $z_{p_{ii},\alpha}$ or $z_{s_{ii},\alpha}$ (mW/cm ²)	160.04				
	I_{spta} at $z_{p_{ij}}$ or $z_{s_{ij}}$ (mW/cm ²)	233.06				
	p_r at $z_{p_{ij}}$ (MPa)	2: 2.85				
Operating Control Conditions	Component 1: UTP 4					
	Component 2: UTP 275					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths $z_{p_{ii}}$ and $z_{p_{ii},\alpha}$ apply to NON-SCANNING MODES, while the depths $z_{s_{ii}}$ and $z_{s_{ii},\alpha}$ apply to SCANNING MODES.

TABLE 8-5. Transducer: Kosmos Torso-One, operating mode: BC-Mode (max TIS/TIB, ISPTA, 12cm depth, large ROI, image top)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.98	0.96		0.96		1.74
Index component value		1: 5.66E-02 2: 0.90	1: 5.66E-02 2: 0.90	1: 5.66E-02 2: 0.90	1: 5.66E-02 2: 0.90	
Acoustic parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	2: 1.58				
	P (mW)		1: 5.15 2: 86.25	1: 5.15 2: 86.25	1: 5.15 2: 86.25	1: 5.15 2: 86.25
	P_{1x1} (mW)		1: 4.39 2: 72.84	1: 4.39 2: 72.84		
	z_s (cm)		1: N/A 2: N/A			
	z_b (cm)				1: N/A 2: N/A	
	z_{MI} (cm)	2: 4.24				
	$z_{p_{ii},\alpha}$ (cm)	2: 4.24				
	f_{awf} (MHz)	2: 2.59	1: 2.71 2: 2.59	1: 2.71 2: 2.59	1: 2.71 2: 2.59	1: 2.71 2: 2.59
	p_{rr} (Hz)	2: 3824.6				
	s_{rr} (Hz)	2: 25.5				
Other information	n_{pps}	2: 10				
	$I_{pa,\alpha}$ at $z_{p_{ii},\alpha}$ (W/cm ²)	2: 153				
	$I_{spta,\alpha}$ at $z_{p_{ii},\alpha}$ or $z_{s_{ii},\alpha}$ (mW/cm ²)	69.29				
	I_{spta} at $z_{p_{ij}}$ or $z_{s_{ij}}$ (mW/cm ²)	151.32				
	p_r at $z_{p_{ij}}$ (MPa)	2: 2.23				
Operating Control Conditions	Component 1: UTP 4					
	Component 2: UTP 277					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths $z_{p_{ii}}$ and $z_{p_{ii},\alpha}$ apply to NON-SCANNING MODES, while the depths $z_{s_{ii}}$ and $z_{s_{ii},\alpha}$ apply to SCANNING MODES.

TABLE 8-6. Transducer: Kosmos Torso-One, acoustic output reporting table, operating mode: PW Doppler (max MI, TIS, TIB)

Index Label	MI	TIS		TIB	
		At surface	Below surface	At surface	Below surface
Maximum index value	0.42	3.04		3.04	
Index component value		0.49	3.04	3.04	3.04
Acoustic Parameters					
$P_{r,\alpha}$ at z_{pII} (MPa)	0.59				
P (mW)		50.93		50.93	
P_{TxI} (mW)		37.76		37.76	
z_s (cm)		1.93			
z_b (cm)					1.87
z_M (cm)	1.93				
$z_{pII,\alpha}$ (cm)	1.93				
f_{Dop} (MHz)	2.03	2.03		2.03	
Other Information					
prf (Hz)	14468				
srr (Hz)	N/A				
n_{pps}	1				
$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm ²)	12.14				
$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm ²)	429.69				
I_{spta} at z_{pII} or z_{sII} (mW/cm ²)	553.54				
p_r at z_{pII} (MPa)	0.68				
Operating Control Conditions					
PRF	14468 Hz				
Gate Size	4mm				
Focal Depth	20mm				

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-7. Transducer: Kosmos Torso-One, acoustic output reporting table, operating mode: CW Doppler (max MI, TIS, TIB)

Index Label	MI	TIS		TIB	
		At surface	Below surface	At surface	Below surface
Maximum index value	0.07	0.49		0.49	
Index component value		0.47	0.49	0.47	2.43
Acoustic Parameters					
$P_{r,\alpha}$ at z_{pII} (MPa)	0.0976				
P (mW)		62.48		62.48	
P_{TxI} (mW)		50.17		50.17	
z_s (cm)		1.27			
z_b (cm)					1.27
z_M (cm)	0.9				
$z_{pII,\alpha}$ (cm)	1.27				
f_{Dop} (MHz)	1.95	1.95		1.95	
Other Information					
prf (Hz)	N/A				
srr (Hz)	N/A				
n_{pps}	1				
$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm ²)	N/A				
$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm ²)	279.77				
I_{spta} at z_{pII} or z_{sII} (mW/cm ²)	331.51				
p_r at z_{pII} (MPa)	0.10				
Operating Control Conditions					
Focal Depth	4cm				
CW Mode					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

Kosmos Lexsa maximum acoustic output summary

TABLE 8-8. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: B-mode (max MI, ISPTA, MSK, 3cm depth)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.77	5.39E-03		5.39E-03		1.25E-02
Index component value		5.39E-03	5.39E-03	5.39E-03	5.39E-03	
Acoustic Parameters						
$p_{r,\alpha}$ at Z_{Ml} (MPa)	2.01					
P (mW)		0.52		0.52		0.52
P_{Tx1} (mW)		0.15		0.15		
Z_s (cm)		1.57				
Z_b (cm)				1.57		
Z_{Ml} (cm)	1.43					
$Z_{pil,\alpha}$ (cm)	1.57					
f_{awf} (MHz)	6.77	7.44		7.44		7.44
p_{rr} (Hz)	1820.0					
s_{rr} (Hz)	28.0					
n_{pps}	1					
$I_{pa,\alpha}$ at $Z_{pil,\alpha}$ (W/cm^2)	1.7E+02					
$I_{spta,\alpha}$ at $Z_{pil,\alpha}$ or $Z_{sil,\alpha}$ (mW/cm^2)	1.62					
I_{spta} at Z_{pil} or Z_{sil} (mW/cm^2)	3.58					
p_r at Z_{pil} (MPa)	2.24					
Other Information						
Operating Control Conditions						
UTP 71						

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths Z_{pil} and $Z_{pil,\alpha}$ apply to NON-SCANNING MODES, while the depths Z_{sil} and $Z_{sil,\alpha}$ apply to SCANNING MODES.

TABLE 8-9. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: B-mode (max TIS, TIB, MSK, 10cm depth)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.19	9.16E-03		9.16E-03		2.05E-02
Index component value		9.16E-03	9.16E-03	9.16E-03	9.16E-03	
Acoustic Parameters						
$p_{r,\alpha}$ at Z_{Ml} (MPa)	0.53					
P (mW)		0.85		0.85		0.85
P_{Tx1} (mW)		0.25		0.25		
Z_s (cm)		1.63				
Z_b (cm)				1.63		
Z_{Ml} (cm)	1.63					
$Z_{pil,\alpha}$ (cm)	1.63					
f_{awf} (MHz)	7.69	7.69		7.69		7.69
p_{rr} (Hz)	1300.0					
s_{rr} (Hz)	20.0					
n_{pps}	1					
$I_{pa,\alpha}$ at $Z_{pil,\alpha}$ (W/cm^2)	17.0					
$I_{spta,\alpha}$ at $Z_{pil,\alpha}$ or $Z_{sil,\alpha}$ (mW/cm^2)	1.36					
I_{spta} at Z_{pil} or Z_{sil} (mW/cm^2)	3.23					
p_r at Z_{pil} (MPa)	0.82					
Other Information						
Operating Control Conditions						
UTP 87						

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths Z_{pil} and $Z_{pil,\alpha}$ apply to NON-SCANNING MODES, while the depths Z_{sil} and $Z_{sil,\alpha}$ apply to SCANNING MODES.

TABLE 8-10. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: BC, CPD-Mode (max MI, vascular, 4cm depth, large ROI)

Index label	MI	TIS		TIS		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	1.37	7.72E-02		7.72E-02		0.29
Index component value		1: 2.35E-03 2: 7.48E-02	1: 2.35E-03 2: 7.48E-02	1: 2.35E-03 2: 7.48E-02	1: 2.35E-03 2: 7.48E-02	
Acoustic Parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	2: 2.88				
	P (mW)		1: 0.26 2: 11.93	1: 0.26 2: 11.93	1: 0.26 2: 11.93	1: 0.26 2: 11.93
	P_{Tx1} (mW)		1: 6.90E-02 2: 3.56	1: 6.90E-02 2: 3.56	1: 6.90E-02 2: 3.56	1: 6.90E-02 2: 3.56
	z_s (cm)			1: N/A 2: N/A		
	z_b (cm)				1: N/A 2: N/A	
	z_{MI} (cm)	2: 0.96				
	$z_{pII,\alpha}$ (cm)	2: 1.57				
	f_{awf} (MHz)	2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42
	p_{rr} (Hz)	2: 8236.4				
	s_{rr} (Hz)	2: 21.4				
Other Information	n_{pps}	2: 12				
	$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm^2)	2: 23.3				
	$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm^2)	29.58				
	I_{spta} at z_{pII} or z_{sII} (mW/cm^2)	48.42				
	p_f at z_{pII} (MPa)	2: 0.95				
Operating Control Conditions	Component 1: UTP 225					
	Component 2: UTP 339 (16V)					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-11. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: BC, CPD-Mode (max ISPTA, vascular, 4cm depth, small ROI, image top)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	1.37	6.50E-02		6.50E-02		7.98E-02
Index component value		1: 3.23E-03 2: 6.18E-02	1: 3.23E-03 2: 6.18E-02	1: 3.23E-03 2: 6.18E-02	1: 3.23E-03 2: 6.18E-02	
Acoustic Parameters	$p_{r,\alpha}$ at z_{MI} (MPa)	2: 2.88				
	P (mW)		1: 0.36 2: 2.94	1: 0.36 2: 2.94	1: 0.36 2: 2.94	1: 0.36 2: 2.94
	P_{Tx1} (mW)		1: 9.49E-02 2: 2.94	1: 9.49E-02 2: 2.94	1: 9.49E-02 2: 2.94	1: 9.49E-02 2: 2.94
	z_s (cm)			1: N/A 2: N/A		
	z_b (cm)				1: N/A 2: N/A	
	z_{MI} (cm)	2: 0.96				
	$z_{pII,\alpha}$ (cm)	2: 1.57				
	f_{awf} (MHz)	2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42	1: 7.15 2: 4.42
	p_{rr} (Hz)	2: 2026.6				
	s_{rr} (Hz)	2: 28.1				
Other Information	n_{pps}	2: 12				
	$I_{pa,\alpha}$ at $z_{pII,\alpha}$ (W/cm^2)	2: 23.3				
	$I_{spta,\alpha}$ at $z_{pII,\alpha}$ or $z_{sII,\alpha}$ (mW/cm^2)	48.65				
	I_{spta} at z_{pII} or z_{sII} (mW/cm^2)	79.44				
	p_f at z_{pII} (MPa)	2: 0.95				
Operating Control Conditions	Component 1: UTP 225					
	Component 2: UTP 339 (16V)					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pII} and $z_{pII,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sII} and $z_{sII,\alpha}$ apply to SCANNING MODES.

TABLE 8-12. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: BC, CPD-Mode (max TIS, TIB)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.94	0.10		0.10		0.29
Index component value		1: 1.91E-03 2: 0.10	1: 1.91E-03 2: 0.10	1: 1.91E-03 2: 0.10	1: 1.91E-03 2: 0.10	
Acoustic Parameters						
$p_{r,\alpha}$ at z_{MI} (MPa)	2: 2.34					
P (mW)		1: 0.22 2: 11.60		1: 0.22 2: 11.60		1: 0.22 2: 11.60
$P_{I,xj}$ (mW)		1: 5.62E-02 2: 3.46		1: 5.62E-02 2: 3.46		
z_s (cm)			1: N/A 2: NA			
z_b (cm)					1: N/A 2: NA	
z_{MI} (cm)	2: 0.93					
$z_{pili,\alpha}$ (cm)	2: 1.40					
f_{awf} (MHz)	2: 6.22	1: 7.15 2: 6.22		1: 7.15 2: 6.22		1: 7.15 2: 6.22
Other Information						
pr_r (Hz)	2: 8830.3					
srr (Hz)	2: 17.8					
n_{pps}	2: 16					
$I_{pa,\alpha}$ at $z_{pili,\alpha}$ (W/cm^2)	2: 73.7					
$I_{spta,\alpha}$ at $z_{pili,\alpha}$ or $z_{sili,\alpha}$ (mW/cm^2)	29.56					
I_{spta} at z_{pji} or z_{sji} (mW/cm^2)	54.39					
p_r at z_{pji} (MPa)	2: 1.51					
Operating Control Conditions						
Component 1: UTP 225						
Component 2: UTP 161						

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pji} and $z_{pili,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sji} and $z_{sili,\alpha}$ apply to SCANNING MODES.

TABLE 8-13. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: PW Doppler (max MI)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.35	0.19		0.47		0.26
Index component value		0.19	0.06	0.19	0.47	
Acoustic Parameters						
$p_{r,\alpha}$ at z_{MI} (MPa)	0.88					
P (mW)		6.45		6.45		6.45
$P_{I,xj}$ (mW)		6.45		6.45		
z_s (cm)		2.6				
z_b (cm)				2.6		
z_{MI} (cm)	1.22					
$z_{pili,\alpha}$ (cm)	1.24					
f_{awf} (MHz)	6.26	6.26	6.26	6.26	6.26	6.26
pr_r (Hz)	15625					
srr (Hz)	N/A					
n_{pps}	1					
$I_{pa,\alpha}$ at $z_{pili,\alpha}$ (W/cm^2)	23.9					
$I_{spta,\alpha}$ at $z_{pili,\alpha}$ or $z_{sili,\alpha}$ (mW/cm^2)	338.3					
Other Information						
I_{spta} at z_{pji} or z_{sji} (mW/cm^2)	575.2					
p_r at z_{pji} (MPa)	1.14					
Operating Control Conditions						
PRF	15625					
Gate Size	5mm					
Gate Focal Depth	10mm					

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{pji} and $z_{pili,\alpha}$ apply to NON-SCANNING MODES, while the depths z_{sji} and $z_{sili,\alpha}$ apply to SCANNING MODES.

TABLE 8-14. Transducer: Kosmos Lexsa acoustic output reporting table, operating mode: PW Doppler (max TIS, TIB, TIC)

Index label	MI	TIS		TIB		TIC
		At surface	Below surface	At surface	Below surface	
Maximum index value	0.15	0.66	0.66	1.64	1.64	0.64
Index component value		0.66	0.26	0.66	1.64	
$p_{r,\alpha}$ at z_{MI} (MPa)	0.38					
P (mW)			22.23		22.23	22.23
P_{1x1} (mW)			22.23		22.23	
z_s (cm)			2.6			
z_b (cm)					2.6	
z_{MI} (cm)	2.58					
$z_{p1i,\alpha}$ (cm)	2.58					
f_{DWT} (MHz)	6.25	6.25	6.25	6.25	6.25	6.25
p_{rr} (Hz)	7621					
s_{rr} (Hz)	N/A					
n_{pps}	1					
$I_{pa,\alpha}$ at $z_{p1i,\alpha}$ (W/cm^2)	5.42					
$I_{spt,\alpha}$ at $z_{p1i,\alpha}$ or $z_{s1i,\alpha}$ (mW/cm^2)	127.8					
$I_{spt,\alpha}$ at z_{p1i} or z_{s1i} (mW/cm^2)	539.19					
p_r at z_{p1i} (MPa)	0.73					
Operating Control Condition	PRF	7621				
	Gate Size	5mm				
	Gate Focal Depth	50mm				

NOTE 1 Only one operating condition per index.
 NOTE 2 Data should be entered for "at surface" and "below surface" both in the columns related to TIS or TIB.
 NOTE 3 If the requirements of 201.12.4.2a) are met, it is not required to enter any data in the columns related to TIS or TIB.
 NOTE 4 If the requirements of 201.12.4.2b) are met, it is not required to enter any data in the column related to MI.
 NOTE 5 Unshaded cells should have a numerical value. The equipment setting related to the index has to be entered in the operating control section.
 NOTE 6 The depths z_{p1i} and z_{s1i} apply to NON-SCANNING MODES, while the depths z_{p1i} and z_{s1i} apply to SCANNING MODES.

Measurement accuracy

Measurement accuracy for distance and area in B-mode images are as follows:

- Axial measurement accuracy: Axial distance measurements in 2D imaging modes shall be accurate to within +/- 2% of the displayed value (or 1 mm, whichever is larger).
- Lateral distance measurement accuracy: Lateral distance measurements in 2D imaging modes shall be accurate to within +/- 2% of the displayed value (or 1 mm, whichever is larger).
- Diagonal measurement accuracy: Diagonal distance measurements in 2D imaging modes shall be accurate to within +/- 2% of the displayed value (or 1 mm, whichever is larger).

- Area measurement accuracy: Area measurement accuracy in 2D imaging modes shall be +/-4% of the nominal value.

Measurement accuracy for distance and time in M-mode images are as follows:

- M-mode distance measurement: M-mode distance measurements shall be accurate to within +/- 3% of the displayed value.
- M-mode time measurement accuracy: M-mode time measurements shall be accurate to within +/- 2% of the displayed value.

Kosmos AI-assisted EF Workflow measurements accuracy:

- The accuracy of the Kosmos EF calculations depends on the correct selection of ED/ES frames and accurate tracing of the LV endocardial border. It is important to review the initial ED/ES frames and LV contours provided by the Kosmos AI algorithms, confirm their accuracy, and edit them, as required.
 - Ensure that the selected ED/ES frames accurately represent the corresponding end-diastolic and end-systolic cardiac phases in the A4C and A2C clips. Use the editing tool to select a more appropriate frame, as required.
 - Ensure that the LV contours accurately follows the LV endocardium. Use the editing tool to properly trace and adjust the LV contours.
- When possible, acquire both A4C and A2C clips to obtain a biplane A4C/A2C EF, which is more accurate than the single plane A4C EF.
- The following table shows the results of comparing Kosmos EF calculations, without any user adjustments, to the average of manual expert measurements performed by two independent Echo Core Labs on the same A4C/A2C clips. Subjects across a wide variety of age, gender orientation, race, body habitus, and health were scanned with Kosmos AI-assisted EF workflow in a clinical point-of-care ultrasound setting. The EFs of the subjects scanned ranged from 20% to 80%. The results below include both A4C/A2C biplane and A4C single-plane acquisitions, with the majority being biplane (A4C single-plane acquisition was sufficient when an adequate A2C view could not be obtained within a reasonable amount of time).

TABLE 8-15. EF comparison metrics

EF Metrics	EF Percentage Units (iOS)
RMSD ¹	6.70 (p-value<0.0001)
Bias	-3.41
95% limits of agreement ²	-14.67 / 7.91

¹ Root-mean-square deviation (RMSD) is a metric of the deviation between Kosmos EF calculations (without any user adjustments), and the average manual expert measurements.

² 95% limits of agreement are expected to include approximately 95% of the differences between Kosmos EF calculations (without any user adjustments) and the average manual expert measurements.

Kosmos Bladder AI Workflow Volume Estimation Accuracy:

- $\pm 3\text{mL}$ for volumes under 100mL and $\pm 3\%$ for volumes between 100mL and 600mL. The accuracy specification assumes the system is being used per the instructions in the Kosmos User Guide while scanning a tissue-equivalent phantom.

Control effects

Kosmos does not provide the user with direct control of acoustic output power. Kosmos has been designed to automatically adjust the output to ensure that acoustic limits are not exceeded in any imaging mode. Since there is no direct user control for output, the user should rely on controlling exposure time and scanning technique to implement the ALARA principle.

Related references

- U.S. Dept. of Health and Human Services, Food and Drug Administration, Guidance for Industry and FDA Staff - Marketing Clearance of Diagnostic Ultrasound Systems and Transducers (2023).
- IEC 60601-2-37:2015 Medical electrical equipment - Part 2-37: Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment
- IEC 62359:2017 Ultrasonics - Field characterization - Test methods for the determination of thermal and mechanical indices related to medical diagnostic ultrasonic fields
- NEMA UD 2-2004 (R2009) Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment Revision 3

Transducer surface temperature rise

TABLE 8-16 summarizes the expected maximum temperature rise for Kosmos. The values are based on a statistical sample test of production-equivalent systems and were measured in accordance with IEC 60601-2-37. The values listed in the table are determined with 90% confidence, that 90% of the systems will result in a temperature rise less than or equal to that stated in the table.

TABLE 8-16. Surface temperature rise

Test	Temperature rise (°C)
Still air	16.02
Simulated use	9.85

Ergonomics

	<p>Repetitive ultrasound scanning may cause occasional discomfort in your thumbs, fingers, hands, arms, shoulders, eyes, neck, back, or other parts of your body. However, if you experience symptoms such as constant or recurring discomfort, soreness, pain, throbbing, aching, tingling, numbness, stiffness, burning sensation, muscle fatigue/weakness, or limited range of motion, do not ignore these warning signs. Promptly see a qualified health professional. Symptoms such as these can be linked with Work Related Musculoskeletal Disorders (WRMSDs). WRMSDs can be painful and may result in potentially disabling injuries to the nerves, muscles, tendons, or other parts of the body. Examples of WRMSDs include bursitis, tendonitis, tenosynovitis, carpal tunnel syndrome, and De Quervain syndrome.</p> <p>While researchers are not able to definitively answer many questions about WRMSDs, there is a general agreement that certain factors are associated with their occurrence, including preexisting medical and physical conditions, overall health, equipment, and body position while performing work, frequency of work, and duration of work.</p>
--	---

Kosmos is intended for quick-look applications by qualified health professionals. It is not intended for continual use in radiology or other departments. If you need to use the device for a continual period, take the following precautions:

- Position yourself comfortably, either with a chair with appropriate lower-back support or by sitting or standing upright.
- Minimize twisting, relax your shoulders, and support your arm with a cushion.
- Hold Kosmos Torso-One or Kosmos Lexsa lightly, keep your wrist straight, and minimize the pressure applied to the patient.
- Take regular breaks.

Basic safety

The transducer and software, along with the Apple iPad Pro 12.9" (A2436), have been verified as compliant with IEC 60601-1. Refer to EchoNouS Tablet compatibility list available on the EchoNouS website at echonous.com/ **product/device-compatibility** for all supported configurations. For maximum safety, observe these warnings and cautions:

	Devices that are compliant with IEC 60950-1 and 62368-1 have not been evaluated for compliance with IEC 60601-1 temperature limits for patient contact.
	Do not operate this system in the presence of flammable gases or anesthetics. Explosion can result. The system is <i>not</i> compliant in AP/ APG environments as defined by IEC 60601-1.
	Do not bring the tablet into contact with the patient. Contact of the tablet with the patient could result in electric shock and risk of burn.

▲	Only charge the tablet and Link with the GlobTek P005974 power supply.
▲	Only use devices and accessories recommended by EchoNous.

It is up to the responsible organization to check leakage current of the tablet used with EchoNous probes in the patient environments to ensure it meets 60601-1 requirements.

Electromagnetic compatibility

▲	The System complies with the Electromagnetic Compatibility requirements of AS/NZ CISPR 11:2015 and EN IEC 60601-1-2:2014: AMD1:2020. However, electronic and mobile communications equipment may transmit electromagnetic energy through air and there is no guarantee that interference will not occur in a particular installation or environment. Interference may result in artifacts, distortion, or degradation of the ultrasound image. If the System is found to cause or respond to interference, try re-orienting the System or the affected device, or increasing the separation distance between the devices. Contact EchoNous customer support or your EchoNous distributor for further information.
▲	EchoNous does not recommend the use of high-frequency electromedical devices in proximity to its systems. EchoNous equipment has not been validated for use with high-frequency electrosurgical devices or procedures. Use of high-frequency electrosurgical devices in proximity to its systems may lead to abnormal system behavior or shutdown of the system. To avoid the risk of a burn hazard, do not use Kosmos probes with high-frequency surgical equipment. Such a hazard may occur in the event of a defect in the high-frequency surgical neutral electrode connection.
▲	The System contains sensitive components and circuits. Failure to observe proper static control procedures may result in damage to the System. Any faults should be reported to EchoNous customer support or your EchoNous distributor for repair.

The **System** is intended for use in the electromagnetic environment specified below. The user of the **System** should assure that it is used in such an environment.

Electromagnetic emissions

TABLE 8-17. Guidance and manufacturer's declaration: electromagnetic emissions

Emissions test	Compliance	Electromagnetic environment: guidance
RF emissions CISPR 11	Group 1	The System uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class A	
Harmonic emissions IEC 61000-3-2	Class A	The System is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	

The **System** has Class A compliance in meaning it is suitable for use in all establishments other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes. If the **System** is found to cause or respond to interference follow the guidelines in the warning section above.

Electromagnetic immunity

TABLE 8-18. Guidance and manufacturer's declaration: electromagnetic immunity

Immunity test	Compliance level	Electromagnetic environment: guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±8kV contact ±2kV, ±4kV, ±8kV, ±15kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2 kV at 100 kHz repetition frequency on Power Supply Lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±0.5 kV, ±1 kV Line to Line ±0.5 kV, ±1 kV, ±2 kV Line to ground	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	0% U_T ; 0.5 Cycle at 0 deg, 45 deg, 90 deg, 135 deg, 180 deg, 225 deg, 270 deg and 315 deg. 0% U_T ; 1 cycle and 70% U_T 25/30 cycles single phase at 0 deg	Mains power quality should be that of a typical commercial or hospital environment.
Power frequency (50/60 Hz) magnetic field IEC 61000-4-8	8 A/m at 30 kHz in CW modulation 65 A/m at 134.2 kHz in 2.1 kHz pulse modulation 75 A/m at 13.56 MHz in 50 kHz Pulse modulation	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
^{1,2} Conducted RF IEC 61000-4-6	3 Vrms ⁵ 0.15 MHz - 80 MHz 6Vrms in ISM and Amateur radio bands between 0.15 MHz -80 MHz 80% AM at 1 kHz	Portable and mobile RF communications equipment should be used no closer to any part of the system , including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter Recommended separation distance $d = 1.2 \sqrt{P}$

TABLE 8-18. Guidance and manufacturer's declaration: electromagnetic immunity

Radiated RF IEC 61000-4-3	3 V/m 80 MHz- 2.7 GHz 80% AM at 1 kHz	$d=1.2 \sqrt{P}$ 80MHz to 800MHz $d=2.3 \sqrt{P}$ 800MHz to 2.5GHz Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separations distance in meters (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey ³ , should be less than the compliance level in each frequency range ⁴ . Interference may occur in the vicinity of equipment marked with the following symbol. 
------------------------------	---	--

- 1 UT is the AC mains voltage prior to application of the test level
- 2 At 80MHz and 800 MHz, the higher frequency range applies
- 3 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
- 4 Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the system is used exceeds the applicable RF compliance level above, the system should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orientating or relocating the system.
- 5 Over the frequency range 150kHz to 80MHz, field strengths should be less than 3V/m.

	When using the optional mobile stand, the System can be susceptible to ESD and may require manual intervention. If ESD results in a System error, unplug the probe and plug back in to restore operation.
	Using cables, or accessories other than those specified for use with the system may result in increased emissions or decreased immunity of the system.

Separation distances

TABLE 8-19. Separation distances

Recommended separation distances between portable and mobile RF communications equipment and the EchoNous System			
Rated maximum output power of transmitter W	Separation distance according to frequency of transmitter		
	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2,5 GHz
	$d=1.2 \cdot \sqrt{P}$	$d=1.2 \cdot \sqrt{P}$	$d=2.3 \cdot \sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.
 NOTE 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.
 NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Standards

HIPAA

Kosmos includes security settings that help you to meet the applicable security requirements listed in the HIPAA standard. Users are ultimately responsible for ensuring the security and protection of all electronic protected health information collected, stored, reviewed, and transmitted on the system.

The Health Insurance Portability and Accountability Act, Pub.L. No. 104-191 (1996). 45 CFR 160, General Administrative Requirements.

45 CFR 164, Security and Privacy

DICOM

Kosmos conforms to the DICOM standard as specified in the Kosmos DICOM Conformance Statement, available at echonous.com. This statement provides information about the purpose, characteristics, configuration, and specifications of the network connections supported by the system.

--End of section--

Specifications

System specifications

Feature	Height (mm)	Width (mm)	Depth (mm)	Weight (grams)	Cable (m)	Operating Frequency (MHz)	Scanning Depth (cm)
Kosmos Torso-One	150 *	56	35	267 (with ferrite equipped cable)	1.5	1.5 - 4.5	4 - 30
Kosmos Lexsa	155	56	35	280 (with cable)	1.5	3 - 10.5	1 - 10
Kosmos Link	295	225	31	800	0.1	-----	-----

*Excluding cable (the hard plastic housing length)

Environmental operating and storage conditions for Kosmos probes, Kosmos Link and compatible tablets

Kosmos probes and Kosmos Link are intended to be used and stored in normal ambient conditions inside a medical facility.

Kosmos probes and tablets: operating, charging, transport, and storage condition ranges

	Operating	Transport/Storage
Temperature (°C)	0°C to +40°C	-20°C to +60°C
Relative humidity (non-condensing)	15% to 95%	15% to 95%
Pressure	62 kPa to 106 kPa	62 kPa to 106 kPa

Kosmos Link: operating, charging, transport, and storage condition ranges

	Operating	Transport/Storage
Temperature (°C)	0°C to +40°C	-20°C to +60°C
Relative humidity (non-condensing)	15% to 95%	15% to 95%
Pressure	70 kPa to 106 kPa	70 kPa to 106 kPa

Mode of operation

▲	After storage at extreme temperatures, check the Kosmos probe surface temperature before applying to a patient. A cold or hot surface may burn the patient.
▲	Only operate, charge, and store Kosmos within the approved environmental parameters.
▲	When used in high ambient temperatures (such as 40°C), the Kosmos safety feature may disable scanning to maintain safe touch temperature.

Kosmos enforces scanning limits to maintain safe user contact temperatures.

Kosmos Link electrical specifications

Output

- Tablet: USB PD 5-12Vdc @ 0-3A
- Kosmos Probes: 5 Vdc ±5%, Max 2.5 A

Internal batteries

- Li-ion battery: 7.2V, 4.04Ah
- Battery charging time: The time to charge the battery from 0% to 90% is ~2 hours.
- Battery life: A fully charged Link will provide 3-8 hours of uninterrupted scanning (performance may vary based on scanning modes used).

Power supply

- Globtek P005974
- Input: 100-240 V~, 50-60 Hz, 1.5 A
- Output: 5-11.9 Vdc, 0.4A, 47.6 W

--End of section --

IT Network

Wireless networking

Functions

Connection to the IT network is required for the following functionality.

- Storing exam data (static images and clips) acquired by Kosmos in Picture Archiving and Communication System (PACS) by DICOM communication. For details, refer to the DICOM Conformance Statement that is on the EchoNous website.
- Setting Kosmos time correctly by inquiring the network time service.

Security

Patient data protection

It is your responsibility to configure your iOS device to comply with your local security policies and regulatory requirements. EchoNous recommends that you protect patient data by encrypting your device and setting a passcode for device access. The Kosmos app does encrypt the patient database as an added level of security.

Wireless networking

Refer to the documentation that accompanies the EchoNous approved tablet for information regarding configuring your device for wireless networking. Consult your IT security department to ensure that your device is configured in a manner that complies with all applicable security requirements.

Network for connecting the device

To ensure safety, use an IT network that is isolated from the external environment by a firewall.

IT network failure recovery measures

Connection to an IT network may become, at times, unreliable, and this may lead to failure to perform the functions described in **"Functions"**. As a result, the following hazardous situations may occur:

Network failure	Impact on equipment	Hazard	Countermeasures
IT network becomes unstable	Unable to transmit exam data to PACS	Delay of diagnosis	Kosmos has internal memory, and exam data is stored in it. After the IT network has returned to stable, the user can re-initiate the transfer of data.
	Delay of transmission to a PACS		
	Incorrect data transmitted to a PACS	Misdiagnosis	Integrity of the data is ensured by the TCP/IP and DICOM protocols used by Kosmos.
	Unable to get the time from a time server	Incorrect exam data	Kosmos has the capability of entering data and time manually.
	Incorrect time data		Kosmos always indicates the date and the time on the main screen.
Firewall has broken down	Attack via network	Manipulation of exam data	Kosmos closes unnecessary network ports.
	Infection by computer virus	Leak of exam data	Kosmos prevents a user from loading software and executing it.

- Connection of equipment to an IT network that includes other systems could result in previously unidentified risks to patients, operators, or third parties. Before connecting the equipment to an uncontrolled IT Network, make sure that all potential risks resulting from such connections were identified and evaluated, and suitable countermeasures were put in place. IEC 80001-1:2010 provides guidance for addressing these risks.

- When a setting of the IT network to which Kosmos is connected has been changed, check that the change does not affect it, and take measures, if necessary. Changes to the IT network include:
 - Changing the network configuration (IP address, router, and so on)
 - Connecting additional items
 - Disconnecting items
 - Updating equipment
 - Upgrading equipment

- Any changes to the IT network could introduce new risks requiring additional evaluation to be performed.

-- End of section --

Glossary

Term	Description
A2C	Apical 2 chamber.
A4C	Apical 4 chamber.
ACEP	American College of Emergency Physicians
Annotation	Annotations are text notes, arrows, and/or measurements that a clinician may add to an image or clip. An annotation appears as an overlay on the image/clip.
Archive	After a report is generated, the patient information is updated in the hospital's EMR/PACS system. The device needs to have a secure connection for data transfer. Once an exam is archived, it cannot be edited. At this point, it is safe to purge the exam from KOSMOS to create more room for new studies.
Arrow	An arrow is an arrow icon that a clinician may put on a certain location of an image/clip to highlight something. This displays as an overlay on the image/clip.
BMI	Body mass index.
B-mode	Kosmos Torso-One array scans a plane through the body and produces a 2D image on the screen. This is also called B-mode imaging.
Calculation	Calculations are estimations made from specific sets of measurements.
Caliper	You perform most measurements by using calipers that you drag into position. The active caliper has a round highlighted handle.
Cine	A cine is a period of images, stored digitally as a sequence of individual frames. It is recorded at high frame rates and may contain more frames than were displayed during the examination.
Clip	A clip is a short sequences of multiple frames like a movie.

Term	Description
Completed exam	Once an exam is completed, you won't be able to add images to the exam. You can add/edit/delete any annotations that have been saved as overlays on images/clips until the exam is archived. Once archived, you cannot edit anything. If the clinician does not complete an exam, KOSMOS will automatically complete the exam when KOSMOS is turned off.
DICOM	Digital Imaging and Communications in Medicine. DICOM is the most universal and fundamental standard in digital medical imaging. It's an all-encompassing data transfer, storage, and display protocol built and designed to cover all functional aspects of contemporary medicine. PACS functionality is DICOM driven.
ED	End-diastolic.
EDV	End-diastolic volume.
EF	Ejection fraction, calculated as (a percentage): $EF = (EDV - ESV) / EDV * 100$
ES	End-systolic.
ESV	End-systolic volume.
Exam	An exam contains all the objects, images, clips, and reports that are saved during a clinical examination of a patient with KOSMOS, which usually maps to a patient's visit.
FOV	Field of view is the two-dimension space of B-mode image acquisition.
Frozen state	The state KOSMOS gets into when you tap the Freeze button in live imaging. During the frozen state, you can add annotations to one frame of the cine and save the still image. The measurements only stay on one frame of the cine, but the annotations will persist in the whole cine. When you save a clip from the cine, annotations are saved as overlays on the clip, but the measurement won't be saved in the clip. That is because usually measurements are relevant to only one frame of a cine instead of the whole series of frames.
HR	Heart rate.
Image	An image is a single frame of an ultrasound view captured by KOSMOS.
LV	Left ventricle.
M-line	A line that appears in B-mode for which M-mode provides the trace.

Term	Description
Measurement	A measurement is a distance or area measurement on images with no inference to underlying anatomy. A measurement overlay shows the tool (such as a caliper or ellipse) and the measured values.
MWL	Modality Worklist
PACS	Picture Archiving and Communication Systems. PACS refer to medical systems (hardware and software) built to run digital medical imaging. The main components of PACS include digital image acquisition devices, digital image archives, and workstations. The PACS settings in this document refer to the settings of connecting to digital image archives.
Physical coordinates	The position in the field of view expressed in terms of physical dimensions either in millimeters or radians with respect to a designated point of reference.
Picture	You can use the KOSMOS camera to take pictures of a wound or injury as part of the exam.
PIMS	Patient Information Management Systems.
Ping test	A ping test is used to test a TCP/IP connection. If the test is successful, the connection between the KOSMOS and PACS archive is working.
Report	A report consists of details of an exam, along with the notes entered by the clinician.
Review	This is the state of KOSMOS where you can review and edit patient data if it has not been archived.
ROI	Region of Interest. The ROI refers to the bounded region in the field of view where color flow information is depicted.
Scan	A scan is a system preset where system parameters are optimized for scanning a certain organ, such as heart or lungs. Scans can include multiple images, clips, and reports that you can save. The scan preset drives calculations, measurements, and reports.
Snackbar	The snackbar is a brief message that displays on the bottom of many KOSMOS screens. You don't have to act on the messages, and they automatically go away after a short period of time.

Term	Description
Study	A study is a collection of one or more series of medical images and presentation states that are logically related for diagnosing a patient. Each study is associated with one patient. A study may include composite instances that are created by a single modality, multiple modalities, or by multiple devices of the same modality. In KOSMOS, the term "exam" means "study" in the DICOM world. An exam contains all the objects, images, clips, and reports that are saved during a clinical examination of a patient with KOSMOS, which usually maps to a patient's visit.
SV	Stroke volume, calculated as: SV=EDV-ESV
TLS	Transport Layer Security
Verify	This is used to conduct a DICOM C-Echo, which sends a signal to the PACS archive using a DICOM protocol to confirm that the PACS archive is working and available on the network.

-- End of section --

Auto EF Clinical Performance and Non-Clinical Testing

Auto EF Clinical Performance Testing

A prospective study was conducted to evaluate the difference between the left ventricle ejection fraction (LVEF; % EF) automatically generated by Kosmos' Auto EF algorithm and manually calculated by cardiologists from clips acquired by cardiac sonographers.

Study Design

A total of 153 participants were scanned by the cardiac sonographers and the two views were obtained using a Kosmos ultrasound system with Auto EF software. The A2C and A4C views were used to calculate the % EF, using Simpson's biplane method, at point-of-care. A range of body-mass index (BMI) was included in the dataset, with 22.8% of the patients overweight ($25 \leq \text{BMI} < 30 \text{ kg/m}^2$) and 31.6% obese ($\text{BMI} > 30 \text{ kg/m}^2$). In addition, 19% of patients had reduced EF ($30\% \leq \text{EF} < 53\%$), and 22% of the patients had severely reduced EF ($\text{EF} < 30\%$). All studies were traced by three (3) sonographers using the biplane Simpson's method to establish the reference standard.

The study continued enrollment until four sonographers had completed scans of 17 patients each. Enrolled patients were evenly stratified into four groups based on BMI to ensure a sufficient distribution of patients by sex and BMI. Following the patient exams, three (3) independent readers manually calculated % EF from scans containing both A4C and A2C views ($N = 141$) which acted as the ground truth.

The performance goal is the Root Mean Square Deviation (RMSD) $< 10\%$ EF between LVEF manually measured by experts and Kosmos' Auto EF on biplane scans (A4C and A2C) acquired by cardiac sonographers.

Results

Results of the hypothesis testing of Kosmos Auto EF software evaluating the performance of the algorithm against its objective performance goal of $< 10\%$ RMSD is shown below. The performance goal was met.

Endpoint	RMSD (95% CI)	p-value	Pearson Correlation	Bias (95% CI)
LVEF calculations	4.57% EF (5.129, 5.153)	< 0.0001	0.96	1.54% EF (6.90, -9.98)

Software Verification and Validation Testing

Software documentation generated as part of EchoNous' design process include:

1. Software/Firmware Description
2. Device Hazard Analysis
3. Software Requirement Specifications
4. Architecture Design Chart
5. Software Design Specifications
6. Traceability
7. Software Development Environment Description
8. Verification and Validation Documentation
9. Revision Level History
10. Unresolved Anomalies
11. Cybersecurity

A comprehensive risk analysis was generated for the software with detailed description of the hazards, their causes and severity, as well as acceptable methods for control of the identified hazards. EchoNous developed a description, with test protocols including pass/fail criteria and report of results, of acceptable verification and validation activities at the unit, integration, and system level.

Algorithm Testing

Comprehensive non-clinical performance testing of the deep-learning algorithms used in the device was performed to support their clinical performance. Specifically, the performance testing evaluated the performance of the following software functionality:

- EF Workflow

Overall, the non-clinical performance testing results provide evidence in support of the functionality of Auto EF's algorithm.

- End of Section

AI FAST Clinical Performance and Non-Clinical Testing

AI FAST Clinical Performance Testing

A prospective study was conducted to evaluate the sensitivity and precision of AI FAST's view identification and object labeling algorithms.

Study Design

A total of 32 subjects were recruited across two sites (16 subjects per site). Each subject was scanned by a cardiac sonographer and a sonographer with abdominal experience using the following views: SUP (suprapubic), RUQ (right upper quadrant), LUQ (left upper quadrant), AS ([transverse] aortic sweep), IVC ([longitudinal] inferior vena cava), Aorta, SUB (subcostal 4-chamber), SUB2 (subcostal 2-chamber), A4C (apical 4-chamber), A2C (apical 2-chamber), PLAX (parasternal long axis), PSAX (parasternal short axis), and Lung.

The participants in this study comprised of a well-rounded representation of demographic factors such as age, gender, BMI, ethnicity, and race. Enrolled patients were evenly stratified into four groups based on BMI to ensure a sufficient distribution of patients by sex and BMI.

Five (5) radiologists, from a pool of nine (9), independently and collectively reviewed the algorithm's predictions on anatomical structure labeling and view identification using extracted frames. These radiologists were used to establish ground truth to compare the algorithm's performance and were blinded to assessments from others on the panel. The results from the expert-panel reads were used for the statistical analysis.

Two (2) prospectively defined primary endpoints were evaluated to demonstrate the efficacy of the anatomical object labeling and view identification of AI FAST's algorithms in clinical settings.

Results

Results of the hypothesis testing of Kosmos AI FAST software evaluating the performance of the algorithm against its objective performance goal of < 20% false detection rate (FDR) is shown below. The performance goal was met.

Endpoint	FDR (95% CI)	Recall	Precision
Object Labeling	0.9% (0.0 – 1.5%)	93.8%	99.6%
View Identification	3.4% (0.0 – 4.3%)	89.9%	96.6%

Note, recall is defined as how well the algorithm can identify positive instances, also known as sensitivity. Precision is how well the algorithm can correctly predict positive instance.

Software Verification and Validation Testing

Software documentation generated as part of EchoNous' design process include:

1. Software/Firmware Description
2. Device Hazard Analysis
3. Software Requirement Specifications
4. Architecture Design Chart
5. Software Design Specifications
6. Traceability
7. Software Development Environment Description
8. Verification and Validation Documentation
9. Revision Level History
10. Unresolved Anomalies
11. Cybersecurity

A comprehensive risk analysis was generated for the software with detailed description of the hazards, their causes and severity, as well as acceptable methods for control of the identified hazards. EchoNous developed a description, with test protocols including pass/fail criteria and report of results, of acceptable verification and validation activities at the unit, integration, and system level.

Algorithm Testing

Comprehensive non-clinical performance testing of the deep-learning algorithms used in the device was performed to support their clinical performance. Specifically, the performance testing evaluated the performance of the following software functionality:

- Abdominal Object Detection and View Identification

Overall, the non-clinical performance testing results provide evidence in support of the functionality of AI FAST's algorithms.

- End of Section -

Kosmos Bladder AI Clinical Performance and Non-Clinical Testing

Kosmos Bladder AI Clinical Performance Testing

A prospective study was conducted to evaluate the correlation between manual bladder volume determination and Kosmos Bladder Biplane Caliper Volume AI, also referred to as Kosmos Bladder AI, algorithm in a clinical setting.

Study Design

A total of 146 participants, 2 abdominal sonographers, and 4 nurses were enrolled in the study. Each participant underwent scanning by an assigned pair of healthcare professionals, comprised of a sonographer and a nurse. Three (3) independent sonographers were recruited to perform manual labeling of the bladder calipers on acquired videos that were used for evaluating the performance of Kosmos Bladder AI workflow. The participants in this study comprised of a well-rounded representation of demographic factors such as age, gender, BMI, ethnicity, and race. This diversity enriched the dataset and provided comprehensive insights.

Each acquired bladder video (transverse and sagittal) was labeled by three (3) sonographers. The average of these 3 measurements was considered the ground truth for assessment.

- Primary Endpoint: correlation coefficient ≥ 0.90

Results

Results of the hypothesis testing of Kosmos Bladder AI software evaluating the performance of the algorithm against its objective performance goal of a correlation coefficient ≥ 0.90 . The performance goal was met.

Endpoint	Result (95% CI)	p-value
Correlation Coefficient	0.988 (0.986 - 0.99)	< 0.0001

Software Verification and Validation Testing

Software documentation generated as part of EchoNous' design process include:

1. Software/Firmware Description
2. Device Hazard Analysis
3. Software Requirement Specifications
4. Architecture Design Chart
5. Software Design Specifications
6. Traceability
7. Software Development Environment Description
8. Verification and Validation Documentation
9. Revision Level History
10. Unresolved Anomalies
11. Cybersecurity

A comprehensive risk analysis was generated for the software with detailed description of the hazards, their causes and severity, as well as acceptable methods for control of the identified hazards. EchoNous developed a description, with test protocols including pass/fail criteria and report of results, of acceptable verification and validation activities at the unit, integration, and system level.

Algorithm Testing

Comprehensive non-clinical performance testing of the deep-learning algorithms used in the device was performed to support their clinical performance. Specifically, the performance testing evaluated the performance of the following software functionality:

- Kosmos Bladder Biplane Caliper Volume Algorithm

Overall, the non-clinical performance testing results provide evidence in support of the functionality of Kosmos Bladder AI's algorithms.

- End of Section -