Manufacturer Disclosure Statement for Medical Device Security -- MDS2 FUJIFILM SonoSite, Inc. Sonosite PX D24031-02 Au

August, 2023

QUESTION		NOTES
Manufacturer Name	FUJIFILM SonoSite, Inc.	
Device Description	Ultrasound	
Device Model	Sonosite PX	_
Document ID	D24031-02	
Manufacturer Contact Information	FUJIFILM SonoSite Technical	
	Support	
	Phone: 877-657-8118	
	Email: ffss-service@fujifilm.com	
Intended use of device in network-connected	DICOM based communications	_
environment:	including but not limited to:	
	Ultrasound Image Storage, Modality	
	Worklist, Print, Storage	
	Commitment,	
	Modality Performed Procedure Step	
Document Release Date	August, 2023	
Coordinated Vulnerability Disclosure: Does the	Yes	https://www.sonosite.com/support/security
manufacturer have a vulnerability disclosure		
program for this device?		
ISAO: Is the manufacturer part of an Information	Yes	
Sharing and Analysis Organization?		
Diagram: Is a network or data flow diagram available	Yes	
that indicates connections to other system		
components or expected external resources?		
SaMD: Is the device Software as a Medical Device	No	_
(i.e. software-only, no hardware)?		
	N/A	
Does the SaMD rely on an owner/operator provided	N/A	
operating system?		
Is the SaMD hosted by the manufacturer?	N/A	
·		
Is the SaMD hosted by the customer?	N/A	
	Device Description Device Model Document ID Manufacturer Contact Information Intended use of device in network-connected environment: Document Release Date Coordinated Vulnerability Disclosure: Does the manufacturer have a vulnerability disclosure program for this device? ISAO: Is the manufacturer part of an Information Sharing and Analysis Organization? Diagram: Is a network or data flow diagram available that indicates connections to other system components or expected external resources? SaMD: Is the device Software as a Medical Device (i.e. software-only, no hardware)? Does the SaMD contain an operating system? Does the SaMD rely on an owner/operator provided operating system?	Device Description Device Model Document ID Manufacturer Contact Information Manufacturer Contact Information Intended use of device in network-connected environment: Intended use of device in network-connected environment: Dicom based communications including but not limited to: Ultrasound Image Storage, Modality Worklist, Print, Storage Commitment, Modality Performed Procedure Step Document Release Date August, 2023 Coordinated Vulnerability Disclosure: Does the manufacturer have a vulnerability disclosure program for this device? ISAO: Is the manufacturer part of an Information Sharing and Analysis Organization? Diagram: Is a network or data flow diagram available that indicates connections to other system components or expected external resources? SaMD: Is the device Software as a Medical Device (i.e. software-only, no hardware)? Does the SaMD contain an operating system? Does the SaMD rely on an owner/operator provided operating system? Is the SaMD hosted by the manufacturer? N/A

	MANAGEMENT OF PERSONALLY IDENTIFIABLE		NOTES
	INFORMATION		
	Can this device display, transmit, store, or modify	Yes	Along with ultrasound images and clips, the device
	personally identifiable information (e.g. electronic		has the ability to store and transmit the following
	Protected Health Information (ePHI))?		ePHI items: Full Patient Name, DOB, Gender,
			Patient ID, Accession Number and Indications.
MPII-1			
	Does the device maintain personally identifiable	Yes	
MPII-2	information?		
	Does the device maintain personally identifiable	Yes	_
	information temporarily in volatile memory (i.e.,		
MPII-2.1	until cleared by power-off or reset)?		
	Does the device store personally identifiable	Yes	
MPII-2.2	information persistently on internal media?		
	Is personally identifiable information preserved in	Yes	
	the device's non-volatile memory until explicitly		
MPII-2.3	erased?		
	Does the device store personally identifiable	Yes	_
MPII-2.4	information in a database?		
	Does the device allow configuration to automatically	Yes	
	delete local personally identifiable information after		
	it is stored to a long term solution?		
MPII-2.5			
	Does the device import/export personally	Yes	_
	identifiable information with other systems (e.g., a		
	wearable monitoring device might export personally		
MPII-2.6	identifiable information to a server)?		
	Does the device maintain personally identifiable	Yes	
	information when powered off, or during power		
MPII-2.7	service interruptions?		

	Does the device allow the internal media to be	Yes	
	removed by a service technician (e.g., for separate		
MPII-2.8	destruction or customer retention)?		
	Does the device allow personally identifiable	Yes	
	information records be stored in a separate location		
	from the device's operating system (i.e. secondary		
	internal drive, alternate drive partition, or remote		
MPII-2.9	storage location)?		
	Does the device have mechanisms used for the	Yes	
	transmitting, importing/exporting of personally		
MPII-3	identifiable information?		
	Does the device display personally identifiable	Yes	
MPII-3.1	information (e.g., video display, etc.)?		
	Does the device generate hardcopy reports or	Yes	_
	images containing personally identifiable		
MPII-3.2	information?		
	Does the device retrieve personally identifiable	Yes	
	information from or record personally identifiable		
	information to removable media (e.g., removable-		
	HDD, USB memory, DVD-R/RW,CD-R/RW, tape,		
MPII-3.3	CF/SD card, memory stick, etc.)?		
	Does the device transmit/receive or import/export	No	
	personally identifiable information via dedicated		
	cable connection (e.g., RS-232, RS-423, USB,		
MPII-3.4	FireWire, etc.)?		
	Does the device transmit/receive personally	Yes	
	identifiable information via a wired network		
MPII-3.5	connection (e.g., RJ45, fiber optic, etc.)?		
	Does the device transmit/receive personally	Yes	
	identifiable information via a wireless network		
	connection (e.g., WiFi, Bluetooth, NFC, infrared,		
MPII-3.6	cellular, etc.)?		
	Does the device transmit/receive personally	No	
	identifiable information over an external network		
MPII-3.7	(e.g., Internet)?		
	Does the device import personally identifiable	Yes	PII can be imported via a barcode scanner.
MPII-3.8	information via scanning a document?		
	Does the device transmit/receive personally	No	
MPII-3.9	identifiable information via a proprietary protocol?		
	Does the device use any other mechanism to	No	
	transmit, import or export personally identifiable		
MPII-3.10	information?		
Management of Privat	te Data notes:		

	AUTOMATIC LOGOFF (ALOF)		NOTES
	The device's ability to prevent access and misuse by		
	unauthorized users if device is left idle for a period of		
	time.		
	Can the device be configured to force	Yes	Inactivity timer to enter sleep mode configurable to
	reauthorization of logged-in user(s) after a		off, 5 minutes or 10 minutes.
	predetermined length of inactivity (e.g., auto-logoff,		Inactivity timer to power down configurable to off,
	session lock, password protected screen saver)?		15 minutes or 30 minutes.
ALOF-1			
	Is the length of inactivity time before auto-	Yes	Inactivity timer to enter sleep mode configurable to
	logoff/screen lock user or administrator		off, 5 minutes or 10 minutes.
	configurable?		Inactivity timer to power down configurable to off,
ALOF-2			15 minutes or 30 minutes.

	AUDIT CONTROLS (AUDT)		NOTES
	The ability to reliably audit activity on the device.		
	Can the medical device create additional audit logs	Yes	
AUDT-1	or reports beyond standard operating system logs?		
AUDT-1.1	Does the audit log record a USER ID?	Yes	
	Does other personally identifiable information exist	No	
AUDT-1.2	in the audit trail?		
	Are events recorded in an audit log? If yes, indicate	Yes	
	which of the following events are recorded in the		
AUDT-2	audit log:		

	Ta	I	
AUDT-2.1	Successful login/logout attempts?	Yes	
AUDT-2.2	Unsuccessful login/logout attempts?	Yes	
AUDT-2.3	Modification of user privileges?	Yes	
AUDT-2.4	Creation/modification/deletion of users?	Yes	
	Presentation of clinical or PII data (e.g. display,	Yes	
AUDT-2.5	print)?		
AUDT-2.6	Creation/modification/deletion of data?	Yes	
	Import/export of data from removable media (e.g.	Yes	
AUDT-2.7	USB drive, external hard drive, DVD)?		
	Receipt/transmission of data or commands over a	Yes	_
AUDT-2.8	network or point-to-point connection?		
AUDT-2.8.1	Remote or on-site support?	N/A	
	Application Programming Interface (API) and similar	N/A	
AUDT-2.8.2	activity?		
AUDT-2.9	Emergency access?	Yes	
AUDT-2.10	Other events (e.g., software updates)?	Yes	
AUDT-2.11	Is the audit capability documented in more detail?	Yes	
	Can the owner/operator define or select which	No	
AUDT-3	events are recorded in the audit log?		
	Is a list of data attributes that are captured in the	Yes	
AUDT-4	audit log for an event available?		_
AUDT-4.1	Does the audit log record date/time?	Yes	
	Can date and time be synchronized by Network Time	Yes	
AUDT-4.1.1	Protocol (NTP) or equivalent time source?		
AUDT-5	Can audit log content be exported?	Yes	
AUDT-5.1	Via physical media?	Yes	
	Via IHE Audit Trail and Node Authentication (ATNA)	No	
AUDT-5.2	profile to SIEM?		_
	Via Other communications (e.g., external service	No	
AUDT-5.3	device, mobile applications)?		_
	Are audit logs encrypted in transit or on storage	Yes	All logs are encrypted locally on the device.
AUDT-5.4	media?		This logo are error ypical rodaily on the devicer
	Can audit logs be monitored/reviewed by	Yes	
AUDT-6	owner/operator?		
AUDT-7	Are audit logs protected from modification?	Yes	
AUDT-7.1	Are audit logs protected from access?	Yes	
AUDT-8	Can audit logs be analyzed by the device?	No	
7,001.0	can addit logs be analyzed by the device:	110	

	AUTHORIZATION (AUTH)		NOTES
	The ability of the device to determine the authorization of users.		
	Does the device prevent access to unauthorized users through user login requirements or other	Yes	
AUTH-1	mechanism?		
AUTH-1.1	Can the device be configured to use federated credentials management of users for authorization (e.g., LDAP, OAuth)?	Yes	_
AUTH-1.2	Can the customer push group policies to the device (e.g., Active Directory)?	No	_
AUTH-1.3	Are any special groups, organizational units, or group policies required?	No	_
AUTH-2	Can users be assigned different privilege levels based on 'role' (e.g., user, administrator, and/or service, etc.)?	Yes	Individual user accounts are required when the device is configured for Secure mode. Accounts can be created for Clinical users, Administrators and Guest users.
AUTH-3	Can the device owner/operator grant themselves unrestricted administrative privileges (e.g., access operating system or application via local root or administrator account)?	No	_
AUTH-4	Does the device authorize or control all API access requests?	N/A	_
AUTH-5	Does the device run in a restricted access mode, or 'kiosk mode', by default?	Yes	_

CYBER SECURITY PRODUCT UPGRADES (CSUP)	INOTES

	I=1 1111 6 11 1 15	T	T
	The ability of on-site service staff, remote service		
	staff, or authorized customer staff to install/upgrade		
	device's security patches.		
	Does the device contain any software or firmware	Yes	FUJIFILM SonoSite will provide system updates to
	which may require security updates during its		deploy any applicable security patches. FUJIFILM
	operational life, either from the device manufacturer		SonoSite performs
	or from a third-party manufacturer of the		regular security scans on their ultrasound systems.
	software/firmware? If no, answer "N/A" to		,
CSUP-1	questions in this section.		
C301 1	Does the device contain an Operating System? If yes,	Voc	FILIEUM ConoCito systems run on a closed
		res	FUJIFILM SonoSite systems run on a closed
00.15.0	complete 2.1-2.4.		proprietary operating system which includes
CSUP-2			components from Windows 10 LTSC.
	Does the device documentation provide instructions	Yes	_
	for owner/operator installation of patches or		
CSUP-2.1	software updates?		
	Does the device require vendor or vendor-	No	
	authorized service to install patches or software		_
CSUP-2.2	updates?		
C301 Z.Z	Does the device have the capability to receive	No	There is no remote assess to the device
		NO	There is no remote access to the device
	remote installation of patches or software updates?		
CSUP-2.3			
	Does the medical device manufacturer allow security	No	_
	updates from any third-party manufacturers (e.g.,		
	Microsoft) to be installed without approval from the		
CSUP-2.4	manufacturer?		
000. 2	Does the device contain Drivers and Firmware? If	Yes	
CSUP-3		163	—
C3UP-3	yes, complete 3.1-3.4.	, , , , , , , , , , , , , , , , , , ,	
	Does the device documentation provide instructions	Yes	
	for owner/operator installation of patches or		
CSUP-3.1	software updates?		
	Does the device require vendor or vendor-	No	
	authorized service to install patches or software		
CSUP-3.2	updates?		
	Does the device have the capability to receive	No	
		140	—
00.15.0.0	remote installation of patches or software updates?		
CSUP-3.3			
	Does the medical device manufacturer allow security	No	
	updates from any third-party manufacturers (e.g.,		
	Microsoft) to be installed without approval from the		
CSUP-3.4	manufacturer?		
	Does the device contain Anti-Malware Software? If	No	
CSUP-4	yes, complete 4.1-4.4.		
	Does the device documentation provide instructions	N/A	
	•	IN/A	
CCLID 4.4	for owner/operator installation of patches or		
CSUP-4.1	software updates?		
	Does the device require vendor or vendor-	N/A	_
	authorized service to install patches or software		
CSUP-4.2	updates?		
	Does the device have the capability to receive	N/A	
1	remote installation of patches or software updates?		
CSUP-4.3	and the second of the second o		
	Does the medical device manufacturer allow security	N/A	
		IV/A	_
	updates from any third-party manufacturers (e.g.,		
	Microsoft) to be installed without approval from the		
CSUP-4.4	manufacturer?		
	Does the device contain Non-Operating System	Yes	
	commercial off-the-shelf components? If yes,		
CSUP-5	complete 5.1-5.4.		
		Yes	
	for owner/operator installation of patches or		
CSLID-5 1			
CSUP-5.1	software updates?	N	
	Does the device require vendor or vendor-	No	-
	authorized service to install patches or software		
CSUP-5.2	updates?		
	Does the device have the capability to receive	No	
	remote installation of patches or software updates?		
CSUP-5.3	and the second s		
	1		

Does the medical device manufacturer allow security	No	
updates from any third-party manufacturers (e.g.,		
Microsoft) to be installed without approval from the		
manufacturer?		
Does the device contain other software components	No	_
(e.g., asset management software, license		
management)? If yes, please provide details or		
reference in notes and complete 6.1-6.4.		
Does the device documentation provide instructions	N/A	
for owner/operator installation of patches or		
software updates?		
Does the device require vendor or vendor-	N/A	
authorized service to install patches or software		
updates?		
Does the device have the capability to receive	N/A	
remote installation of patches or software updates?		
Does the medical device manufacturer allow security	N/A	
updates from any third-party manufacturers (e.g.,		
Microsoft) to be installed without approval from the		
manufacturer?		
Does the manufacturer notify the customer when	Yes	
updates are approved for installation?		
Does the device perform automatic installation of	No	
software updates?		
Does the manufacturer have an approved list of third-	N/A	
party software that can be installed on the device?		
Can the owner/operator install manufacturer-	No	
approved third-party software on the device		
themselves?		
Does the system have mechanism in place to prevent	Yes	
installation of unapproved software?		
Does the manufacturer have a process in place to	Yes	
assess device vulnerabilities and updates?		
Does the manufacturer provide customers with	Yes	
review and approval status of updates?		
Is there an update review cycle for the device?	Yes	
	updates from any third-party manufacturers (e.g., Microsoft) to be installed without approval from the manufacturer? Does the device contain other software components (e.g., asset management software, license management)? If yes, please provide details or reference in notes and complete 6.1-6.4. Does the device documentation provide instructions for owner/operator installation of patches or software updates? Does the device require vendor or vendorauthorized service to install patches or software updates? Does the device have the capability to receive remote installation of patches or software updates? Does the medical device manufacturer allow security updates from any third-party manufacturers (e.g., Microsoft) to be installed without approval from the manufacturer? Does the manufacturer notify the customer when updates are approved for installation? Does the device perform automatic installation of software updates? Does the manufacturer have an approved list of third-party software that can be installed on the device? Can the owner/operator install manufacturer-approved third-party software on the device themselves? Does the system have mechanism in place to prevent installation of unapproved software? Does the manufacturer have a process in place to assess device vulnerabilities and updates? Does the manufacturer provide customers with review and approval status of updates?	Microsoft) to be installed without approval from the manufacturer? Does the device contain other software components (e.g., asset management software, license management)? If yes, please provide details or reference in notes and complete 6.1-6.4. Does the device documentation provide instructions for owner/operator installation of patches or software updates? Does the device require vendor or vendor-authorized service to install patches or software updates? Does the device have the capability to receive remote installation of patches or software updates? Does the medical device manufacturer allow security updates from any third-party manufacturers (e.g., Microsoft) to be installed without approval from the manufacturer? Does the manufacturer notify the customer when updates are approved for installation? Does the device perform automatic installation of software updates? Does the manufacturer have an approved list of third-party software that can be installed on the device? Can the owner/operator install manufacturer-approved third-party software on the device themselves? Does the system have mechanism in place to prevent installation of unapproved software? Does the manufacturer have a process in place to assess device vulnerabilities and updates? Does the manufacturer have and updates? Does the manufacturer provide customers with review and approval status of updates?

	HEALTH DATA DE-IDENTIFICATION (DIDT)		NOTES
	The ability of the device to directly remove		
	information that allows identification of a person.		
	Does the device provide an integral capability to de-	Yes	The device can be configured to obfuscate PHI on
	identify personally identifiable information?		the display screen and has the ability to de-identify
DIDT-1			patient data prior to USB export.
	Does the device support de-identification profiles	Yes	
	that comply with the DICOM standard for de-		
DIDT-1.1	identification?		

	DATA BACKUP AND DISASTER RECOVERY (DTBK)		NOTES
	The ability to recover after damage or destruction of		
	, , , , , , , , , , , , , , , , , , , ,		
	device data, hardware, software, or site		
	configuration information.		
	Does the device maintain long term primary storage	No	_
	of personally identifiable information / patient		
DTBK-1	information (e.g. PACS)?		
	Does the device have a "factory reset" function to	Yes	
	restore the original device settings as provided by		
DTBK-2	the manufacturer?		
	Does the device have an integral data backup	Yes	
DTBK-3	capability to removable media?		
	Does the device have an integral data backup	Yes	
DTBK-4	capability to remote storage?		
	Does the device have a backup capability for system	Yes	
	configuration information, patch restoration, and		
DTBK-5	software restoration?		
	Does the device provide the capability to check the	No	
DTBK-6	integrity and authenticity of a backup?		

	EMERGENCY ACCESS (EMRG)		NOTES
	The ability of the device user to access personally		
	identifiable information in case of a medical		
	emergency situation that requires immediate access		
	to stored personally identifiable information.		
	Does the device incorporate an emergency access	Yes	
EMRG-1	(i.e. "break-glass") feature?		

	HEALTH DATA INTEGRITY AND AUTHENTICITY (IGAU)		NOTES
	How the device ensures that the stored data on the		
	device has not been altered or destroyed in a non-		
	authorized manner and is from the originator.		
	Does the device provide data integrity checking	Yes	
	mechanisms of stored health data (e.g., hash or		
IGAU-1	digital signature)?		
	Does the device provide error/failure protection and	No	_
	recovery mechanisms for stored health data (e.g.,		
IGAU-2	RAID-5)?		

	MALWARE DETECTION/PROTECTION (MLDP)		NOTES
	The ability of the device to effectively prevent, detect		
	and remove malicious software (malware).		
	Is the device capable of hosting executable software?	No	_
MLDP-1			
	Does the device support the use of anti-malware	No	FUJIFILM SonoSite ultrasound systems feature
	software (or other anti-malware mechanism)?		whitelist software, which prevents third-party
	Provide details or reference in notes.		software from being installed and/or executed on
			the product.
			No third party software can be installed and/or
			executed on FUJIFILM SonoSite ultrasound systems.
MLDP-2			·
	Does the device include anti-malware software by	No	
MLDP-2.1	default?		
	Does the device have anti-malware software	No	_
MLDP-2.2	available as an option?		
	Does the device documentation allow the	No	
	owner/operator to install or update anti-malware		
MLDP-2.3	software?		
	Can the device owner/operator independently (re-	N/A	
MLDP-2.4)configure anti-malware settings?		
	Does notification of malware detection occur in the	N/A	_
MLDP-2.5	device user interface?		
	Can only manufacturer-authorized persons repair	No	
MLDP-2.6	systems when malware has been detected?		
MLDP-2.7	Are malware notifications written to a log?	No	
	Are there any restrictions on anti-malware (e.g.,	N/A	
MLDP-2.8	purchase, installation, configuration, scheduling)?		
	If the answer to MLDP-2 is NO, and anti-malware	Yes	FUJIFILM SonoSite ultrasound systems feature
	cannot be installed on the device, are other		whitelist software, which prevents third-party
	compensating controls in place or available?		software from being installed and/or executed on
MLDP-3			the product.
	Does the device employ application whitelisting that	Yes	
	restricts the software and services that are permitted		
MLDP-4	to be run on the device?		
	Does the device employ a host-based intrusion	No	
MLDP-5	detection/prevention system?		
	Can the host-based intrusion detection/prevention	No	
MLDP-5.1	system be configured by the customer?		
	Can a host-based intrusion detection/prevention	No	
MLDP-5.2	system be installed by the customer?		

NODE AUTHENTICATION (NAUT)	NOTES
The ability of the device to authenticate	
communication partners/nodes.	

	Does the device provide/support any means of node	Yes	When optionally configured for DICOM based
	authentication that assures both the sender and the		communications, the modality (sender) and the
	recipient of data are known to each other and are		recipient must be identified
	authorized to receive transferred information (e.g.		
NAUT-1	Web APIs, SMTP, SNMP)?		
	Are network access control mechanisms supported	Yes	Connections limited to pre defined DICOM server.
	(E.g., does the device have an internal firewall, or use		
NAUT-2	a network connection white list)?		
	Is the firewall ruleset documented and available for	No	
NAUT-2.1	review?		
	Does the device use certificate-based network	Yes	Wireless network communication, along with
	connection authentication?		remote worksheets and Secure LDAP can be
			configured to use certificate based network
NAUT-3			authentication.

	CONNECTIVITY CAPABILITIES (CONN)		NOTES
	All network and removable media connections must		
	be considered in determining appropriate security		
	controls. This section lists connectivity capabilities		
	that may be present on the device.		
	Does the device have hardware connectivity	Yes	
CONN-1	capabilities?		
CONN-1.1	Does the device support wireless connections?	Yes	
CONN-1.1.1	Does the device support Wi-Fi?	Yes	
CONN-1.1.2	Does the device support Bluetooth?	No	
	Does the device support other wireless network	No	
CONN-1.1.3	connectivity (e.g. LTE, Zigbee, proprietary)?		
	Does the device support other wireless connections	No	
	(e.g., custom RF controls, wireless detectors)?		
CONN-1.1.4			
CONN-1.2	Does the device support physical connections?	Yes	
	Does the device have available RJ45 Ethernet ports?	Yes	
CONN-1.2.1			
CONN-1.2.2	Does the device have available USB ports?	Yes	
	Does the device require, use, or support removable	Yes	
CONN-1.2.3	memory devices?		
	Does the device support other physical connectivity?	No	
CONN-1.2.4			
	Does the manufacturer provide a list of network	Yes	
	ports and protocols that are used or may be used on		
CONN-2	the device?		
	Can the device communicate with other systems	Yes	_
CONN-3	within the customer environment?		
	Can the device communicate with other systems	No	
	external to the customer environment (e.g., a service		
CONN-4	host)?		
CONN-5	Does the device make or receive API calls?	No	
	Does the device require an internet connection for	No	_
CONN-6	its intended use?		
	Does the device support Transport Layer Security	Yes	_
CONN-7	(TLS)?		
CONN-7.1	Is TLS configurable?	Yes	
	Does the device provide operator control	No	
	functionality from a separate device (e.g.,		
CONN-8	telemedicine)?		

	PERSON AUTHENTICATION (PAUT)		NOTES
	The ability to configure the device to authenticate		
	users.		
	Does the device support and enforce unique IDs and	Yes	There are no default service accounts on the
	passwords for all users and roles (including service		device.
PAUT-1	accounts)?		
	Does the device enforce authentication of unique IDs	Yes	
	and passwords for all users and roles (including		
PAUT-1.1	service accounts)?		
	Is the device configurable to authenticate users	Yes	MS Active Directory and LDAP.
	through an external authentication service (e.g., MS		
PAUT-2	Active Directory, NDS, LDAP, OAuth, etc.)?		

	<u>, </u>		
	Is the device configurable to lock out a user after a	Yes	
PAUT-3	certain number of unsuccessful logon attempts?		
	Are all default accounts (e.g., technician service	Yes	
	accounts, administrator accounts) listed in the		
PAUT-4	documentation?		
PAUT-5	Can all passwords be changed?	Yes	
	Is the device configurable to enforce creation of user	Yes	
	account passwords that meet established		
PAUT-6	(organization specific) complexity rules?		
	Does the device support account passwords that	Yes	
PAUT-7	expire periodically?		
	Does the device support multi-factor authentication?	No	
PAUT-8			
PAUT-9	Does the device support single sign-on (SSO)?	No	
	Can user accounts be disabled/locked on the device?	Yes	
PAUT-10			
PAUT-11	Does the device support biometric controls?	No	
	Does the device support physical tokens (e.g. badge	No	
PAUT-12	access)?		
	Does the device support group authentication (e.g.	Yes	
PAUT-13	hospital teams)?		
	Does the application or device store or manage	Yes	_
PAUT-14	authentication credentials?		
PAUT-14.1	Are credentials stored using a secure method?	Yes	_

	PHYSICAL LOCKS (PLOK)		NOTES
	Physical locks can prevent unauthorized users with		
	physical access to the device from compromising the		
	integrity and confidentiality of personally identifiable		
	information stored on the device or on removable		
	media		
	Is the device software only? If yes, answer "N/A" to	No	
PLOK-1	remaining questions in this section.		
	Are all device components maintaining personally	Yes	
	identifiable information (other than removable		
	media) physically secure (i.e., cannot remove		
PLOK-2	without tools)?		
	Are all device components maintaining personally	N/A	
	identifiable information (other than removable		
	media) physically secured behind an individually		
PLOK-3	keyed locking device?		
	Does the device have an option for the customer to	N/A	Media is none removable.
	attach a physical lock to restrict access to removable		
PLOK-4	media?		

	ROADMAP FOR THIRD PARTY COMPONENTS IN		NOTES
	DEVICE LIFE CYCLE (RDMP)		
	Manufacturer's plans for security support of third-		
	party components within the device's life cycle.		
	Was a secure software development process, such as	Yes	
	ISO/IEC 27034 or IEC 62304, followed during product		
RDMP-1	development?		
	Does the manufacturer evaluate third-party	Yes	
	applications and software components included in		
RDMP-2	the device for secure development practices?		
	Does the manufacturer maintain a web page or	Yes	https://www.sonosite.com/support/sonosite-
	other source of information on software support		product-retirement-schedule
RDMP-3	dates and updates?		
	Does the manufacturer have a plan for managing	Yes	
RDMP-4	third-party component end-of-life?		

	SOFTWARE BILL OF MATERIALS (SBoM)		NOTES
	A Software Bill of Material (SBoM) lists all the		
	software components that are incorporated into the		
	device being described for the purpose of		
	operational security planning by the healthcare		
	delivery organization. This section supports controls		
	in the RDMP section.		
SBOM-1	Is the SBoM for this product available?	Yes	

	Does the SBoM follow a standard or common	Yes	
SBOM-2	method in describing software components?		
SBOM-2.1	Are the software components identified?	Yes	
	Are the developers/manufacturers of the software	Yes	
SBOM-2.2	components identified?		
	Are the major version numbers of the software	Yes	
SBOM-2.3	components identified?		
SBOM-2.4	Are any additional descriptive elements identified?	Yes	
	Does the device include a command or process	No	_
	method available to generate a list of software		
SBOM-3	components installed on the device?		
SBOM-4	Is there an update process for the SBoM?	Yes	

	SYSTEM AND APPLICATION HARDENING (SAHD)		NOTES
	The device's inherent resistance to cyber attacks and		
	malware.		
	Is the device hardened in accordance with any	Yes	All ports and services not needed for the device to
	industry standards?		operate as intended have been disabled or
SAHD-1			removed
	Has the device received any cybersecurity	No	_
SAHD-2	certifications?		
	Does the device employ any mechanisms for	Yes	System and Integrity checking is performed during
SAHD-3	software integrity checking		boot up
	Does the device employ any mechanism (e.g., release	Yes	System and Integrity checking is performed during
	specific hash key, checksums, digital signature, etc.)		boot up
	to ensure the installed software is manufacturer-		
SAHD-3.1	authorized?		
	Does the device employ any mechanism (e.g., release	Yes	System and Integrity checking is performed during
	specific hash key, checksums, digital signature, etc.)		boot up
	to ensure the software updates are the		
SAHD-3.2	manufacturer-authorized updates?		
	Can the owner/operator perform software integrity	No	_
	checks (i.e., verify that the system has not been		
SAHD-4	modified or tampered with)?		
	Is the system configurable to allow the	Yes	_
	implementation of file-level, patient level, or other		
SAHD-5	types of access controls?		
	Does the device provide role-based access controls?	Yes	
SAHD-5.1			
	Are any system or user accounts restricted or	No	_
SAHD-6	disabled by the manufacturer at system delivery?		
	Are any system or user accounts configurable by the	Yes	_
SAHD-6.1	end user after initial configuration?		
	Does this include restricting certain system or user	Yes	Individual user accounts are required when the
	accounts, such as service technicians, to least		device is configured for Secure mode. Accounts can
	privileged access?		be created for Clinical users, Administrators and
SAHD-6.2			Guest users.
	Are all shared resources (e.g., file shares) which are	Yes	_
	not required for the intended use of the device		
SAHD-7	disabled?		
	Are all communication ports and protocols that are	Yes	_
	not required for the intended use of the device		
SAHD-8	disabled?		
	Are all services (e.g., telnet, file transfer protocol	Yes	_
	[FTP], internet information server [IIS], etc.), which		
	are not required for the intended use of the device		
SAHD-9	deleted/disabled?		
	Are all applications (COTS applications as well as OS-	Yes	_
	included applications, e.g., MS Internet Explorer,		
	etc.) which are not required for the intended use of		
SAHD-10	the device deleted/disabled?		
	Can the device prohibit boot from uncontrolled or	Yes	_
	removable media (i.e., a source other than an		
SAHD-11	internal drive or memory component)?		
	Can unauthorized software or hardware be installed	No	_
	on the device without the use of physical tools?		
SAHD-12			

	Does the product documentation include	No	
	information on operational network security		
SAHD-13	scanning by users?		
	Can the device be hardened beyond the default	Yes	Administrator can disable the following:
	provided state?		Ethernet, WiFi and USB devices, export to USB
			devices.
SAHD-14			Admin. can configure password complexity rules.
	Are instructions available from vendor for increased	No	
SAHD-14.1	hardening?		
	Can the system prevent access to BIOS or other	Yes	
SHAD-15	bootloaders during boot?		
	Have additional hardening methods not included in	Yes	
SAHD-16	2.3.19 been used to harden the device?		

	SECURITY GUIDANCE (SGUD)		NOTES
	Availability of security guidance for operator and		
	administrator of the device and manufacturer sales		
	and service.		
	Does the device include security documentation for	Yes	
SGUD-1	the owner/operator?		
	Does the device have the capability, and provide	Yes	
	instructions, for the permanent deletion of data		
SGUD-2	from the device or media?		
	Are all access accounts documented?	Yes	
SGUD-3			
	Can the owner/operator manage password control	Yes	
SGUD-3.1	for all accounts?		
	Does the product include documentation on	Yes	
	recommended compensating controls for the		
SGUD-4	device?		

	HEALTH DATA STORAGE CONFIDENTIALITY		NOTES
	(STCF)		
	The ability of the device to ensure unauthorized		
	access does not compromise the integrity and		
	confidentiality of personally identifiable information		
	stored on the device or removable media.		
	Can the device encrypt data at rest?	Yes	The device uses AES-256 bit encryption to protect
STCF-1			data at rest.
STCF-1.1	Is all data encrypted or otherwise protected?	Yes	
	Is the data encryption capability configured by	Yes	
STCF-1.2	default?		
	Are instructions available to the customer to	Yes	Factory reset or secure erase patient drive will
STCF-1.3	configure encryption?		generate a new encryption key.
	Can the encryption keys be changed or configured?	Yes	Factory reset or secure erase patient drive will
STCF-2			generate a new encryption key.
	Is the data stored in a database located on the	Yes	
STCF-3	device?		
	Is the data stored in a database external to the	No	
STCF-4	device?		

	TRANSMISSION CONFIDENTIALITY (TXCF)		NOTES
	The ability of the device to ensure the confidentiality		
	of transmitted personally identifiable information.		
	Can personally identifiable information be	No	_
	transmitted only via a point-to-point dedicated		
TXCF-1	cable?		
	Is personally identifiable information encrypted prior	Yes	
	to transmission via a network or removable media?		
TXCF-2			
	If data is not encrypted by default, can the customer	Yes	_
TXCF-2.1	configure encryption options?		
	Is personally identifiable information transmission	Yes	DICOM fixed list
TXCF-3	restricted to a fixed list of network destinations?		
TXCF-4	Are connections limited to authenticated systems?	Yes	

TXCF-5	Are secure transmission methods supported/implemented (DICOM, HL7, IEEE 11073)?	Yes	
	TRANSMISSION INTEGRITY (TXIG)		NOTES
	The ability of the device to ensure the integrity of		
	transmitted data.		
	Does the device support any mechanism (e.g., digital	Yes	The device provides the option to encrypt data with
	signatures) intended to ensure data is not modified		a FIPS 140-2 validated encryption algorithm prior to
TXIG-1	during transmission?		wireless transmission.
	Does the device include multiple sub-components	No	_
TXIG-2	connected by external cables?		

	REMOTE SERVICE (RMOT)		NOTES
	Remote service refers to all kinds of device		
	maintenance activities performed by a service		
	person via network or other remote connection.		
	Does the device permit remote service connections	No	_
RMOT-1	for device analysis or repair?		
	Does the device allow the owner/operator to	N/A	_
	initiative remote service sessions for device analysis		
RMOT-1.1	or repair?		
	Is there an indicator for an enabled and active	N/A	_
RMOT-1.2	remote session?		
	Can patient data be accessed or viewed from the	N/A	_
RMOT-1.3	device during the remote session?		
	Does the device permit or use remote service	No	_
RMOT-2	connections for predictive maintenance data?		
	Does the device have any other remotely accessible	No	
	functionality (e.g. software updates, remote		
RMOT-3	training)?		

OTHER SECURITY CONSIDERATIONS (OTHR)		
NONE		
Notes:		