STORZ-ENDOSKOPE

en Instructions for use IMAGE1 S





10-2022

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1 General information

1.1 Read the instructions for use

If the instructions for use are not followed, patients, users, and third parties may be injured or the product may be damaged.

- ▶ Read the instructions for use carefully and follow all the safety notes and warnings.
- Read the reprocessing instructions carefully and follow all the safety notes and warnings. The reprocessing instructions can be downloaded from www.karlstorz.com/ifu by entering the item number.
- ▶ Keep the instructions for use and reprocessing instructions in a safe place.

1.2 Read the instructions for use of compatible products

If the instructions for use of compatible products are not followed, patients, users, and third parties may be injured or the product may be damaged.

- Read the instructions for use of the compatible products carefully and follow all the safety notes and warnings.
- Read the reprocessing instructions of the compatible products carefully and follow all the safety notes and warnings.

1.3 Scope

These instructions for use apply to the following products up to and including software version 4.4:

Product name	Item number
IMAGE1 S CONNECT	TC200
IMAGE1 S CONNECT II	TC201
IMAGE1 S H3-LINK	TC300
IMAGE1 S X-LINK	TC301
IMAGE1 S D-LINK	TC302
IMAGE1 S 4U-LINK	TC304
IMAGE1 S D3-LINK extension cable	TC015

The products listed here may not yet be available in all countries due to differences in approval requirements.

1.4 General signs and symbols

The signs and symbols used in this document have the following meaning:

Practical tip

 (\mathbf{i}) This sign refers to useful and important information.

Actions to be performed

Action to be carried out by several steps:

- \checkmark $\,$ Prerequisite that must be met before carrying out an action.
- 1. Step 1
 - ⇒ Interim result of an action

General information



- 2. Step 2
- ⇒ Result of a completed action

Actions in safety notes or in the case of a single step:

▶ Step 1

Lists

- 1. Numbered list
- Unnumbered list, 1st level
 - Unnumbered list, 2nd level

1.5 Description of warning messages

To prevent any injury to persons or damage to property, the warnings and safety notes in the instructions for use must be observed. The warnings use the following levels of danger:

WARNING

Designates a possible imminent risk. If this is not avoided, it could lead to death or serious injuries.

CAUTION

Designates a possible imminent risk. If this is not avoided, it could lead to minor injuries.

NOTICE

Designates a possibly harmful situation. If this is not avoided, the products could be damaged.



2 Normal use

2.1 Intended use

IMAGE1 S

The IMAGE1 S is a camera control unit for use with camera heads or videoscopes and is used for visualizing and documenting endoscopic and microscopic procedures. The IMAGE1 S has no direct contact with the human body.

Extension cable

The extension cable is used for extending the camera head cable for connection to the IMAGE1 S camera control unit. The extension cable has no direct contact with the human body.

2.2 Indications

The camera control units do not come into direct contact with the patient; instead, they are used in conjunction with the corresponding accessories (videoendoscopes, camera, light source, and monitor) and the applied parts for the purpose of visualizing and documenting endoscopic and exoscopic procedures. Use is not restricted to certain medical disciplines.

2.3 Contraindications

No contraindications relating directly to the medical device are currently known. The responsible physician must decide whether the foreseen application is admissible based on the general condition of the patient.

2.4 Target user populations

The medical device may only be used by doctors and medical assistants with a relevant specialist qualification.

2.5 Patient population

There are no restrictions in terms of patient groups for this product.



3 Safety and warning

A WARNING

Danger due to non-observance of warnings and safety notes

This chapter contains warnings and safety notes structured according to hazards and risks.

- 1. Carefully read and observe all warnings and safety notes.
- 2. Follow the instructions.

3.1 Serious incidents

A 'serious incident' includes incidents which, directly or indirectly, had, could have had or could have any of the following consequences:

- Death of a patient, user, or another person
- Temporary or permanent serious deterioration in the medical condition of a patient, user, or another person
- A serious threat to public health
- ► The manufacturer and appropriate authority must be notified of all serious incidents.

3.2 Observing ambient condition

If the device is operated in an environment which is not suitable, patients, users and third parties may be injured.

► Always store and operate the product in the prescribed ambient conditions.

3.3 Combination with other components

The use of unauthorized devices and components or unauthorized changes to the product can result in injuries.

Additional devices connected to electrical medical equipment must comply with the relevant IEC or ISO standards. Furthermore, all configurations must comply with the requirements for medical electrical systems.

- Only combine the product with devices and components that are approved for combined use by the manufacturer.
- Only use devices and components that have standardized interfaces and do not breach the intended use of the product.
- ► Do not modify this equipment without authorization of the manufacturer.
- ► Do not use multiple socket outlets with this device.
- ► If the use of a multiple socket outlet cannot be avoided: Only use a multiple socket outlet with an isolation transformer in accordance with IEC 60601-1.

3.4 Damage due to ingress of liquid in electrical components

In the case of electrical products, individual components or the product itself may be live. Liquid ingress into an electrical product may result in a short circuit or an unintentional transfer of current. The product is damaged as a result and patients, users and third parties may be injured.

- ► Do not store liquids near the product or on the product.
- ► If liquid has entered the product, pull out the plug and allow the product to dry completely.



3.5 Dangers from electrical current

An improper power supply may cause an electric shock and injure patients, users, or third parties. All electrical installations in the operating room in which the product is connected and used must meet all applicable IEC standards.

- > Do not touch the output jacks of the product and the patient at the same time during use
- Do not open the product.
- ► Use either the power cord supplied by KARL STORZ or a power cord which has the same properties and which bears a national mark of conformity.
- ▶ Make sure that the plug is completely clean and dry.
- ▶ Do not use more than one extension cable.
- ▶ The product's ground line should be installed by a qualified electrician.
- ► Ensure potential equalization according to the applicable national rules and regulations.
- ► To ensure reliable protective earth grounding, connect the product to a properly installed socket that is approved for use in the operation room. Routinely inspect the electrical plug and cord and do not use if the inspection reveals damage.
- ► To avoid the risk of electric shock, this product must only be connected to a supply mains with protective earth.

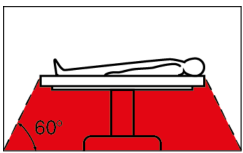
If several products supplied with energy are used simultaneously, the patient leakage currents accumulate. These leakage currents can exceed the limit values and injure patients.

▶ The patient applied parts of the simultaneously used products must be type BF or type CF.

3.6 Risk of explosion and fire

The product can generate sparks, which cause combustible or flammable gases and liquids to ignite or explode. This may cause injuries to patients, users, and third parties.

▶ When using explosive narcotic gases: Operate the product outside of the hazard zone.



- ▶ Do not use the product in the presence of flammable anesthetics.
- ► The product must not be operated in oxygenated environments.
- Only connect or disconnect the power plug to or from the power supply outside explosive atmospheres.

3.7 Electromagnetic interference

Medical electrical products are subject to special precautions regarding electromagnetic compatibility and must be installed and commissioned according to the tables on electromagnetic compatibility. If other products (e.g. for MRT, CT, diathermy, electrocautery or RFID) emit electromagnetic radiation, the function of the product may be impaired. High-frequency communication equipment can affect electrical medical products and impair their performance.

 Observe the information on electromagnetic compatibility, see chapter *Electromagnetic* compatibility [p. 60].



The discharge of a defibrillator can cause image loss.

▶ If the image has not recovered after 4 seconds, switch the camera control unit off and on again.

3.8 Product not clean

The product is not clean when delivered. The use of products that have not been cleaned poses a risk of infection for patients, users, and third parties.

 Reprocess the product in line with the reprocessing instructions before initial use and every subsequent use.

3.9 Failure of products

The product may fail during use.

- ▶ Perform an equipment test before use, see chapter *Preparation* [p. 25].
- ► If the image fails during the procedure, remove the camera from the endoscope and continue the procedure optically.
- ► If the surgery cannot be continued optically, determination of how to further the procedure is at the physician's discretion based on the surgical circumstances.
- ► Have a replacement system ready for each application.



4.1 Description of operation

The product offers the following functions:

- IMAGE1 S Visualization modes to support the user with contrast enhancement and/or consistent brightness perception.
- Light Source Control for automatic adjustment of the light source.
- Integrated image and video capture for simple documentation.

4.2 Product overview

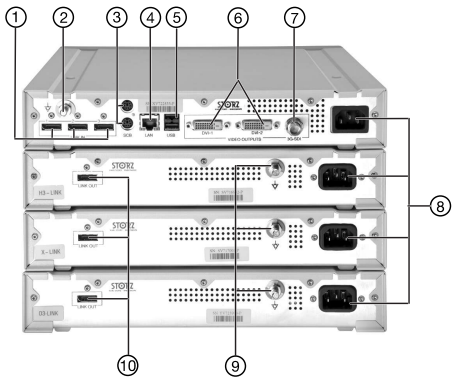


IMAGE1 S CONNECT, front

- 1 Power switch
- 2 White balance
- 3 USB port
- 4 H3-LINK camera connection
- 5 X-LINK camera connection

- 6 D3-LINK camera connection
- 7 IMAGE1 S D3-LINK (TC302)
- 8 IMAGE1 S X-LINK (TC301)
- 9 IMAGE1 S H3-LINK (TC 300)
- 10 IMAGE1 S CONNECT (TC200)





Rear side of IMAGE1 S CONNECT

- 1 LINK input ports
- 2 Potential equalization connector
- 3 SCB input/output sockets (KARL STORZ Communication Bus)
- 4 LAN interface, KS HIVE interface
- 5 USB interface

- 6 HD DVI-D video output sockets
- 7 3G-SDI video output
- 8 Mains sockets
- 9 Potential equalization connector
- 10 LINK output



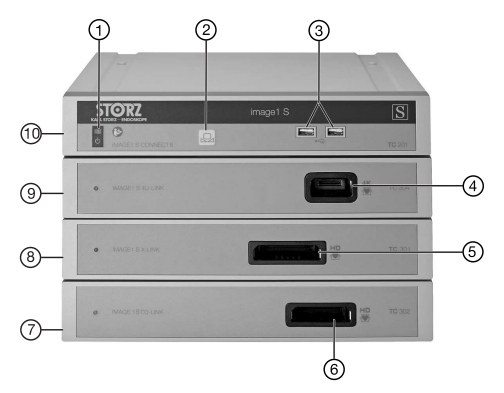
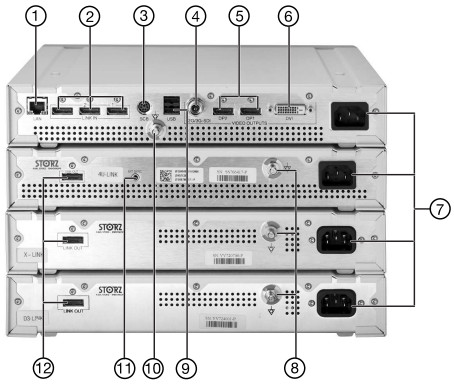


IMAGE1 S CONNECT II, front

- 1 Power switch
- 2 White balance
- 3 USB port
- 4 4U-LINK camera connection
- 5 X-LINK camera connection

- 6 D3-LINK camera connection
- 7 IMAGE1 S D-LINK (TC302)
- 8 IMAGE1 S X-LINK (TC301)
- 9 IMAGE1 S 4U-LINK (TC304)
- 10 IMAGE1 S CONNECT II (TC201)





Rear side of IMAGE1 S CONNECT II

- 1 LAN interface, KS HIVE interface
- 2 LINK input ports
- 3 SCB input/output sockets (KARL STORZ Communication Bus)
- 4 12G/3G SDI video output
- 5 DisplayPort video output sockets
- 6 HD DVI-D video output socket

- 7 Mains sockets
- 8 Potential equalization connector
- 9 USB interface
- 10 Potential equalization connector
- 11 EXT SYNC connection
- 12 LINK output

4.3 Technical data

IMAGE1 S CONNECT

Description	Value
HD video outputs	2x DVI-D
	1x 3G-SDI
Signal output format	1,920 x 1,080p, 50/60 Hz
LINK input ports	3x
KS HIVE interface	RJ45
USB interface	4x USB
Image format	JPEG
Video format	MPEG 4
SCB interface	2x 6 pin Mini DIN



IMAGE1 S CONNECT II

Description	Value
4K and HD video outputs	2x DP
	1x 12G/3G SDI
	1x DVI-D
Signal output format	3,840 x 2,160p, 50/60 Hz for DP and 12G-SDI
	1,920 x 1,080p, 50/60 Hz for 3G-SDI and DVI-D
LINK input ports	3x
KS HIVE interface	RJ45
USB interface	4x USB
Image format	JPEG
Video format	MPEG 4
SCB interface	1x 6 pin Mini DIN
Power supply (AC) and frequency require- ments	Value
IMAGE1 S CONNECT	100–120 V, 50/60 Hz
	200–240 V, 50/60 Hz
IMAGE1 S CONNECT II	100–240 V, 50/60 Hz
IMAGE1 S H3-LINK	100–240 V, 50/60 Hz
IMAGE1 S X-LINK	100–240 V, 50/60 Hz
IMAGE1 S D3-LINK	100–240 V, 50/60 Hz
IMAGE1 S 4U-LINK	100–240 V, 50/60 Hz
Dimensions	Value (W x H x D)
IMAGE1 S CONNECT	320 x 54 x 305 mm
IMAGE1 S CONNECT II	320 x 54 x 305 mm
IMAGE1 S H3-LINK	320 x 54 x 305 mm
IMAGE1 S X-LINK	320 x 54 x 305 mm
IMAGE1 S D3-LINK	320 x 54 x 305 mm
IMAGE1 S 4U-LINK	320 x 54 x 305 mm
Weight	Value
IMAGE1 S CONNECT	2.1 kg
IMAGE1 S CONNECT II	3.1 kg
IMAGE1 S H3-LINK	1.86 kg
IMAGE1 S X-LINK	1.86 kg
IMAGE1 S D3-LINK	1.86 kg
IMAGE1 S 4U-LINK	1.86 kg



Power input	Value
IMAGE1 S CONNECT	62 VA
IMAGE1 S CONNECT II	100–135 VA
IMAGE1 S H3-LINK	80 VA
IMAGE1 S X-LINK	80 VA
IMAGE1 S D3-LINK	80 VA
IMAGE1 S 4U-LINK	86 VA

4.4 Possible combinations

It is recommended that the suitability of the products for the intended procedure be checked prior to use. Please note that the products listed here may not yet be available in all countries due to differences in approval requirements.

The IMAGE1 S camera control unit (CCU) consists of the IMAGE1 S CONNECT (TC200) or IMAGE1 S CONNECT II (TC201) and 1–3 additional LINK modules. The various modules serve as interfaces to camera heads and videoendoscopes.

The following LINK modules are available:

- IMAGE1 S H3-LINK (TC300)
- IMAGE1 S X-LINK (TC301)
- IMAGE1 S D3-LINK module (TC302)
- IMAGE1 S 4U-LINK module (TC304)
- (i) Only compatible products may be combined with the CONNECT modules and LINK modules, see chapter *Accessories* [p. 52].

The following products are required for navigation and input:

- Keyboard
- Mouse

The following products are available as options for navigation:

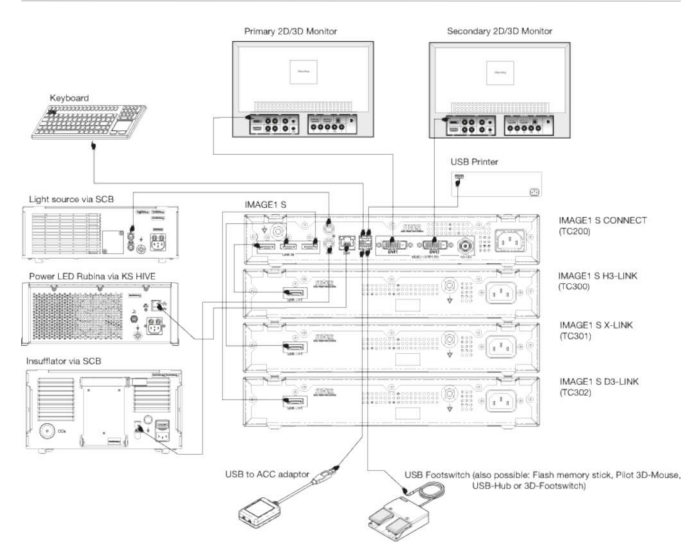
- Compatible camera heads and videoendoscopes
- IMAGE1 PILOT (TC014)
- Microscope footswitch (TC019)

Do not connect network devices (Ethernet, Bluetooth, or WiFi dongles) to the USB port of the IMAGE1 S. The connection may impair operation of the IMAGE1 S.

4.4.1 Installation overview of IMAGE1 S CONNECT

(i) If multiple KS HIVE-compatible peripheral devices are used, the OR1 HIVE router is required.

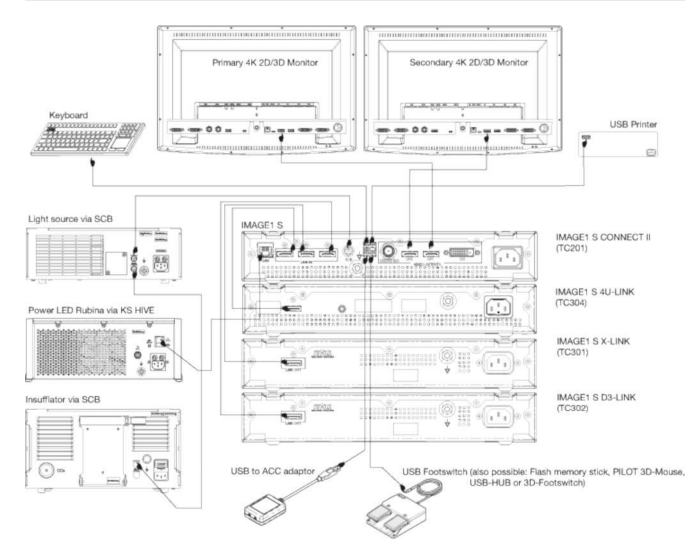




4.4.2 Installation overview of IMAGE1 S CONNECT II

- (i) To use OPAL1 fluorescence imaging, the IMAGE1 S 4U-LINK (TC304) and Power LED Rubina (TL400) must also be connected via a sync cable when using the TIPCAM1 Rubina.
- (i) If multiple KS HIVE-compatible peripheral devices are used, the OR1 HIVE router is required.





4.5 Symbols employed

4.5.1 Symbols on the packaging

Symbol	Meaning
	Manufacturer
	Date of manufacture
MD	Medical device
REF	Article no.
SN	Serial number



Symbol	Meaning
QTY	Number of products in the product packaging
UDI	Unique Device Identifier
ī	Consult the printed or electronic instructions for use
Ţ	Fragile, handle with care
Ť	Keep dry
X	Temperature limit
<u>%</u>	Humidity limit
MR	Not MR safe
Rx only	Federal (USA) law restricts this device to sale by or on the order of a physician.
CE	CE marking With this marking, the manufacturer declares the conformity of the product with the applicable EU directives. A code number after the CE mark indicates the responsible notified body.
	The EU directives relevant to the product can be found in the EU Declaration of Conformity, which can be requested from KARL STORZ.
<u>11</u>	This side up



4.5.2 Symbols on the product

Symbol	Meaning
\$	Follow the instructions for use. The color may differ on the product. The symbol is black/white on the packag- ing label.
\bigcirc	ON/OFF (standby)
∙<⁺	USB
	White balance
	Applied part type CF defibrillation protection
\forall	Potential equalization connector
0	Prevention of pollution by electronic devices
	Separate collection of electrical and electronic devices. Do not dispose of in household refuse.
ĺ	Consult the printed or electronic instructions for use

4.5.3 Symbols on the user interface

Symbol	Meaning
H3-LINK	H3-LINK The button appears once the H3-LINK module is correctly connected.
	X-LINK The button appears once the X-LINK module is correctly connected.
D3-LINK	D3-LINK The button appears once the D3-LINK module is correctly connected.
4U-LINK	4U-LINK The button appears once the 4U-LINK module is correctly connected.



Symbol	Meaning
	Insufflator
	Printer
Se	Light source
	Memory/Eject
0	Patient
00	Presets
×	Setup menu
?	Info
	Brightness
	Print
	Video
• O •	Image



Symbol	Meaning
R	Image flip
S	Enhance
= <u>*</u>	Parameter regions
Î a	Patient data privacy
	Dashboard
	Live menu
%	Image
í	Software license
X	Exit
	Frequency
	Display
	Contact information





Symbol	Meaning
Date + Time	Date and time
	Camera buttons
	White balance
OPAL1	OPAL1
5	Network
Ö	IMAGE1 PILOT
	Setup
	Update
<i></i>	Remote Service

4.6 Ambient conditions

Storage and transport cond	litions
Temperature	-10°C +60°C (14°F 140°F)
Relative humidity (non-condensing)	20 – 95%
Air pressure	500 – 1,060 hPa



Operating conditions	
Temperature	0°C 40°C (32°F 104°F)
Relative humidity (non-condensing)	20 – 85%
Air pressure	700 – 1,060 hPa
Max. operating altitude	3,000 m



5 Preparation

5.1 Unpacking the product

- 1. Carefully remove the product and accessories from the packaging.
- 2. Check the delivery for missing items and possible damage.
- 3. In the case of damage, hidden defects, and short deliveries, document their nature and extent and contact the manufacturer or supplier immediately.
- 4. Keep packaging for further transport.

5.2 Inspecting the product

- 1. Check the product and the camera head or videoendoscope for external damage.
- 2. Ensure that the camera head cable or videoendoscope cable is not damaged or kinked.
- 3. Point the camera head or the videoendoscope toward an object to check the display quality on the monitor.
- 4. When using 3D videoendoscopes, check to make sure the monitor's display is correct.
- 5. Be sure to use the correct polarized eyewear for the monitor.

5.3 Setting up the product

Blocked air inlets and air outlets! Risk of fire!

If the air inlets and outlets are blocked, the product may overheat. This can cause the product to fail and start a fire. Users, patients, and third parties may be injured.

- ► Remove any blockages in the air inlets and outlets when setting up the product.
- 1. Set the product down on a horizontal, flat surface or in a video cart. Make sure that the power cord can be unplugged at any time.
- 2. Ensure adequate air circulation.
- 3. For best thermal performance, position the product at the top of the device stack.

5.4 Connecting the product

1. Connect the potential equalization cable.







- 2. Connect the potential equalization line to the connector of the treatment room.
- 3. Connect the numbered inputs in ascending order.
- 4. Use the LINK cable to connect the LINK module to the CONNECT module.



- 5. Pull back the protection device of the SCB connector and insert the connector into one of the SCB sockets.
- 6. Connect the other end of the cable to a KARL STORZ SCB control device (KARL STORZ Communication Bus).





7. Insert the OR1 patch cable into the KS HIVE/LAN interface.



- Connect the other end of the cable to a KS HIVE-compatible product, e.g., Power LED Rubina (TL400), Power LED Saphira (TL410), ARTIP CRUISE (UR500B, UR500C), AIDA (WD300, WD350), OR1 HIVE router (WO10240, WO10307).
- 9. Connect the IMAGE1 S 4U-LINK (TC304) and Power LED Rubina (TL400) to a sync cable if the products are being used with the TIPCAM1 Rubina.
- 10. Connect the monitor. For the IMAGE1 S CONNECT (TC200), connect the primary work screen to DVI 1.



- 11. Ensure that the monitor supports a resolution of 1,920 x 1,080 pixels.
- 12. If the D3-LINK module (TC302) is used in conjunction with a 3D videoendoscope, connect a compatible 3D monitor.
- 13. For the IMAGE1 S CONNECT II (TC201) connect the primary work screen to DP 1.
- 14. Ensure that the monitor supports a resolution of 3,840 x 2,160 pixels.



15. Connect the power cord. Push the power plug fully into the power socket.



16. Push the power plugs fully into the power sockets of the LINK modules.



- 17. Connect the power plugs to the power sockets.
- (\mathbf{i}) Use the network device only in a protected network with firewall and router.

5.5 Putting the product into operation

Hot surfaces! Risk of burns!

The surfaces can get hot due to locked ventilation fans or blocked air intakes. There is a possible risk of a burn if touched.

▶ If the surfaces get hot, switch off the product.

HF electrodes and laser probes! Risk of injury!

If HF electrodes and laser probes are activated in the instrument channel, the patient may be seriously injured.

 Only apply power to the HF electrodes and laser probe once the active parts have left the instrument channel and are in the user's field of vision, and the site of the operation is clearly recognizable.



Image and function malfunctions!

Incorrectly connected camera connections can cause image interference and malfunctioning.

- Ensure that the camera connector is always attached properly and securely before starting the procedure.
- 1. Turn the product on using the power switch.

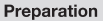


- \Rightarrow The connected LINK modules start automatically.
- 2. Format the USB storage device as FAT 32.
- 3. Connect the USB storage device to the IMAGE1 S CONNECT to store still images or videos. Do not connect more than 1 USB storage device.



- 4. Check the values set for the primary source and the secondary source.
- 5. Check the layout.
- 6. Insert the camera head connector into the socket of the relevant LINK module.







5.6 Performing the white balance

- (1) The white balance must be performed again after replacing the endoscope.
- (1) Failure to perform proper white balance of system may result in color distortion.
- 1. Ensure that the imaging instrument is connected and a live image appears on the monitor.
- 2. Connect the light source to the imaging instrument via the light cable.
- 3. Point the tip of the imaging instrument at a white and clean surface, e.g., white gauze.
- 4. Make sure that the light source emits enough light to sufficiently illuminate the surface.
 ⇒ The monitor displays a completely white and well illuminated live image.
- 5. Press the White Balance button on the IMAGE 1 CONNECT or the button in the Live menu.
 ⇒ The white balance is performed.
- 6. Check the monitor to see whether the white balance was successful.
- 7. Repeat the white balance if required.



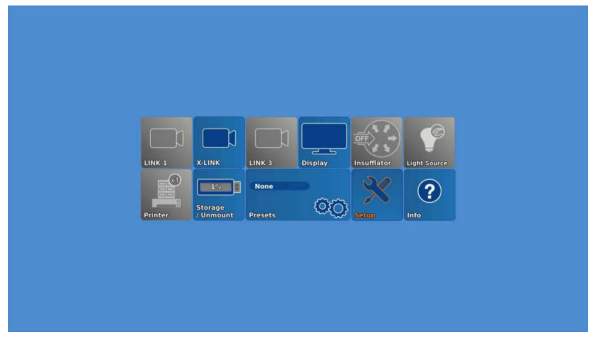
6 Application

6.1 Dashboard

The dashboard displays the LINK modules, peripheral devices, and recommended accessories connected to the LINK module. Unconnected peripheral devices, recommended accessories, and the LINK modules are shown in gray.

If more than one LINK module is used, the IMAGE1 S with the Auto Source Detection function automatically recognizes the module to which a camera head is connected. Provided that the secondary source is configured with *None* or S-Technologies. If the secondary source is configured with a LINK module, the IMAGE1 S behaves according to the configuration.

Depending on the configuration, individual icons may be deactivated or unavailable. The configuration can be changed in the Setup menu, see chapter *Setup menu* [p. 37].



The dashboard is shown when no camera head or videoendoscope is connected.

- 1. Tap the **Display** button.
- 2. Set the primary source, secondary source, PiP layout, and grid.
- 3. Tap the **Insufflator** button.
- 4. In the **Insufflator** menu, adjust the settings for gas, flow, and pressure.
- 5. Tap the Light Source button.
- 6. Turn the light source on or off and set the light intensity and the mode.
 - ⇒ If two or more light sources, camera heads, videoendoscopes are connected at the same time, no light source settings can be made. Exception: The light source tile remains active if one OPAL1-compatible and one non OPAL1-compatible camera head are used.
- 7. Tap the **Printer** button.
- 8. The number of images per page and the number of copies can be set in the **Printer** menu.
- 9. On the **Memory/Eject** button, read off the amount of used storage space on the connected USB storage device.



10. Tap the **Memory/Eject** button to eject the USB storage device. Data may be lost if the USB storage device is not ejected.

 \Rightarrow The USB storage device can be removed.

- 11. Tap the **Presets** button.
- 12. If presets have been defined in the Setup menu, select a preset.
- 13. Tap the **Setup** button to open the Setup menu. Alternatively, you can press the SHIFT+ENTER keys.
- 14. Tap the **Info** button to see contact information.
- 15. Tap the **Patient** button.
- 16. Create, select, modify, or delete a patient information record. Up to 50 patient information records can be created.

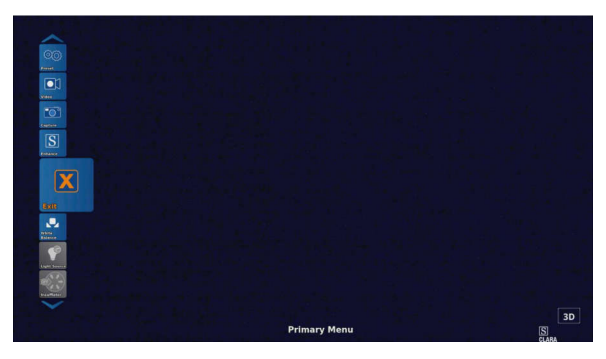
6.2 Live menu

(1) Using an electrosurgical unit in ICG color light mode can cause image distortion.

The Live menu allows you to choose between different camera functions and change the settings of connected devices. Unconnected devices are shown in gray.

The Live menu can be opened using the following keys/buttons:

- The Enter key on the keyboard.
- The menu button on the camera head or videoendoscope.
- The left side button on the IMAGE1 PILOT.
- Be sure not to touch the 3D wheel or the buttons while the IMAGE1 PILOT is connecting to the IMAGE1 CONNECT.



- 1. Tap the Brightness button to set the brightness of the LINK module.
- 2. Tap the **Print** button to immediately print all images in the print queue.
- \Rightarrow Printouts with patient data remain in the queue until sufficient ink and paper are available.
- 3. Switch the IMAGE1 S off and back on again to clear the queue.



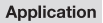
- 4. On the **Memory/Eject** button, read off the amount of used storage space on the connected USB storage device.
- 5. Tap the **Memory/Eject** button to eject the USB storage device. Data may be lost if the USB storage device is not ejected.
 - \Rightarrow The USB storage device can be removed.
- 6. Tap the **Presets** button.
- 7. To select a preset, define presets in the Setup menu.
- 8. When using presets in conjunction with 3D settings, switch manually between 2D and 3D if necessary.
- 9. Tap the Video button to start or stop a video recording.
- 10. Tap the Image button to capture a still.
- 11. Connect a USB storage device to save the images and videos.
- 12. Connect a printer to print the images.
- 13. Connect AIDA to display and save the images and videos.
- 14. Tap the OPAL1 button to set the fluorescence imaging, see chapter OPAL1 [p. 35].
- 15. Tap the **Enhance** button to set the S-Technologies, see chapter *Enhance (S-Technologies)* [p. 35].
- 16. Tap the White Balance button to perform the white balance.
- 17. Tap the Light Source button.
- 18. Turn the light source on or off and set the light intensity and the mode.
- 19. If the light sources are connected via KS HIVE, activate KS HIVE device integration, see chapter *Network* [p. 41].
 - ⇒ If two or more light sources, camera heads, or videoendoscopes are connected at the same time, no light source settings can be made.
 - ⇒ The light source tile remains active if one OPAL1-compatible and one non-OPAL1-compatible camera head are used.
- 20. Tap the **Insufflator** button.
- 21. In the Insufflator menu, adjust the settings for Gas, Flow, and Pressure.
- 22. Tap the Display button, see chapter Display (Live menu) [p. 36].



23. Tap the **Zoom** button to enlarge or reduce the display.

\sim			
X			
Fait			
White Balance			
Y			
Light-Source			
AT .			
insuffator/			
Display			
	1		
	Zoom		
	Adaptive Zoom	Enabled	
	Adaptive 200m	Enabled	
Zoom	Exit		
R			
Image Filip			
Head Button			
			Delener
			Primary Menu

- 24. Select **Adaptive Zoom** in the *Zoom* menu. The menu is available if the following products are connected:
- IMAGE1 S CONNECT II (TC201) and a 4U-LINK in combination with 4U camera heads (TH120 or TH121)
- IMAGE1 S CONNECT (TC200) or IMAGE1 S CONNECT II (TC201) and a D3-LINK in combination with a 4 mm TIPCAM1 S 3D
- IMAGE1 S CONNECT (TC200) or IMAGE1 S CONNECT II (TC201) and a X-LINK in combination with a CMOS-based X-Line videoendoscopes
- IMAGE1 S CONNECT (TC200) or IMAGE1 S CONNECT II (TC201) and an X-LINK in combination with HX camera heads (TH110, TH111, TH112, TH113)
- 25. Select Enabled.
 - ⇒ The system detects the size of the endoscopic image. With small telescopes, the endoscopic image is zoomed to vertically fill the display. If the endoscopic image crosses the vertical boundaries, the display zooms to full screen.
 - ⇒ In the case of camera heads, the adaptive zoom function is used by means of the white balance function.
- 26. Tap the Image Flip button to flip the image vertically or horizontally or rotate it through 180°.
- 27. Tap the **Camera Keys** button to assign functions to the camera head or videoendoscope buttons.
- 28. Tap the IMAGE1 PILOT button to assign functions to the IMAGE1 PILOT buttons.
- 29. Tap the **Exit** button to exit the Live menu. Alternatively, press the ESC key or right side button on the IMAGE1 PILOT.





6.2.1 Enhance (S-Technologies)

- ► Tap the **Enhance** (S-Technologies) button.
 - \Rightarrow The following settings are possible:
- CLARA: A specially optimized algorithm reduces disruptive shadows and ensures more even brightness perception of the endoscopic image.
- CHROMA: Improves the perception of fine structures via contrast enhancements, particularly in the red area. The original colors are retained; the color intensity is changed.
- SPECTRA A, SPECTRA B: Based on color inversions, a false color image with enhanced contrast is generated.
- Fiber filters: The digital fiberscope filter can be set to two levels: Filter A: anti-moiré, Filter B: anti-grid. Interfering artifacts are reduced, such as the moiré effect or the display of the fiberscope grid. Filter A is suitable if the honeycombed grid structures are small. Filter B is suitable for larger honeycomb structures; for example, when an optical zoom is used.

6.2.2 OPAL1

- \checkmark The light source and camera head are connected to the FI support.
- ► Tap the OPAL1 button to define the settings for the fluorescence imaging (FI).
 - \Rightarrow The following settings are possible:
- FI switch: It is possible to switch between the FI application and white light in the menu, via the camera head buttons, or via a footswitch.
- Overlay color: The NIR/ICG signal can be displayed in blue or green.
- FI intensity (only available for PDD): Adjustable in 3 levels. When the FI intensity is increased, the exposure time is extended and the live image is delayed.

Fl Mode	Overlay 🔽
FI Switch	White Light
Overlay Color	Blue 🔽
Exit	

FI mode	Description
Overlay	The overlay image combines the white light image and the detected NIR signal. The NIR signal is displayed in either blue or green together with the white light signal.
Intensity Map	Intensity Map combines the white light image and the detected NIR signal. The NIR information is displayed in a color scale to differentiate between strong and weak NIR signals. The yellowish display represents stronger signals, the blue display weaker signals. The color scaling shown in the image depends on the position of the optics relative to the tissue. The display cannot be used to quantify the NIR signal.
Monochromatic	The monochromatic visualization displays the NIR signal in grayscale. No white light information is displayed.
Overlay Opacity (for TH121)	Set the overlay opacity of the NIR signal. The white light information remains unchanged.



6.2.3 Display (Live menu)

1. Tap the **Display** button in the Live menu.

X				
Willie .				
Constanti	Display Mode	20 20 30		
-	Reset Orientation	ОК		
Concentration of Concen	Adjust Horizon			
	Reset Horizon to 0			
Regimy	Primary Source	4U-LINK 🔽		
	Secondary Source (PiP)	None 🔽		
Marrel (PTP Layout			
R	Grid	off 🔽		
	Griti Visibility	100% 💟		
	Exit			

- 2. Select one of the 4 display modes for the TIPCAM1 Rubina in **Display Mode**:
- 2D: 2D image without horizon control
- 2D Autorotation: 2D image with digital horizon control
- 3D: 3D image without horizon control
- 3D/2D Autoswitch: Automatic switch between 3D image and upright 2D image, depending on the rotation angle of the TIPCAM1 Rubina. The angle for the image switch can be adjusted.
- 3. If the position of the horizon indicator is incorrect, tap the **Reset Orientation** button to restore the correct position.
- 4. In Adjust Horizon, tap the Plus or Minus button to rotate the digital image by the set angle.
- 5. Tap the **OK** button in **Reset Horizon to 0** to reset the horizon.
- 6. Open the Primary Source drop-down menu and select a LINK module.
 - ⇒ If the secondary source is configured with None or S-Technologies, the IMAGE1 S automatically recognizes the module to which a camera head is connected, even if the LINK used is not the primary source.
- 7. Open the **Secondary Source (PiP)** drop-down menu and select another LINK module or an S-Technology.
- 8. In PIP Layout, select a layout to adapt the picture-in-picture (PiP) function display.
 - ⇒ If S-Technology is selected and a visualization mode (CLARA or CHROMA) is selected, the standard white light image and the applicable visualization mode are displayed next to one another as PiP.
 - ⇒ If a D3-LINK or 4U-LINK is selected as the primary source and 3D equipment is connected, the secondary source is deactivated.
- 9. Activate the **Grid** option to fade a grid into the camera image.
- 10. Set the grid transparency in Grid Visibility.



6.3 Setup menu

In the Setup menu, the general settings and the LINK-specific settings can be changed and saved as a preset. If the 2D image is displayed, no menu structure appears in the image.

- 1. Open the Setup menu using one of the following options:
- 2. Press the Setup button on the dashboard.
- 3. Press and hold the center camera head button in the Live menu for 7 seconds.
- 4. Press the SHIFT+ENTER shortcut on the keyboard.
- 5. Press and hold the left side button on the IMAGE1 PILOT in the Live menu for 7 seconds.
- 6. Tap the **Presets** button to perform the following actions:
- Store the current settings as presets
- Load stored presets
- Delete stored presets
- Apply default settings
- 7. To amend the LINK-specific settings, tap a displayed LINK module.
- 8. Tap the **Camera Keys** button to assign functions to the camera head or videoendoscope buttons.
- 9. Tap the Setup button and set which visualization modes are to appear in the S-menu.



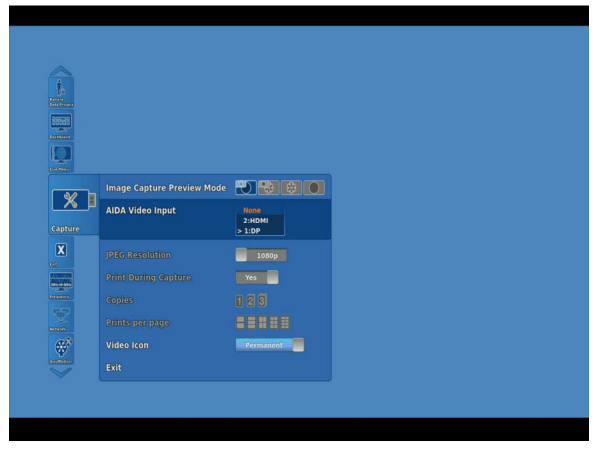
10. Tap the **Module Information** button to call up information about the CONNECT module, the connected LINK modules, and the camera heads or videoendoscopes.



- 11. Tap the General button to apply the general settings.
- 12. Tap the **Parameter Regions** button to adjust the display of additional information from connected SCB devices and KS HIVE devices.
- 13. Tap the **Patient Data Privacy** button to configure the settings for the **Patient Data Privacy**, see chapter *Patient data privacy* [p. 43].
- 14. Tap the **Dashboard** button and select the icons to be displayed on the dashboard here.
- 15. Tap the Live Menu button and select the icons to be displayed in the Live menu.



16. Tap the Capture button to apply settings for capturing the live image.



- 17. Use Image Capture Preview Mode to set how the frozen image or live image is to be displayed.
- 18. If AIDA is connected, select full screen.
- 19. If AIDA is connected via KS HIVE, select the video input being used in **AIDA Video Input** (available with AIDA software version 1.7 and higher). To do this, turn on KS HIVE device integration, see chapter *Network* [p. 41].
- 20. Set the resolution of the captured image in JPEG Resolution.
- 21. In Freeze/Capture Preview Time, set whether and how long the image is to remain frozen.
- 22. In Print During Capture, set whether to create a printout during capture.
- 23. In Copies, set the number of printouts.
- 24. In Images per page, set the number of images to be printed per page.
- 25. In Video Icon, set a permanent or temporary display of a video icon during video recording.
- 26. Tap the Frequency button to select the image refresh rate between 50 and 60 Hz.
- 27. Tap the **Network** button to call up the network parameters and configure the network settings, see chapter *Network* [p. 41].
- 28. Tap the **Remote Service** button to adjust software update settings and to perform software updates.
- 29. Tap the **Insufflation** button and, in the dashboard, select the insufflator functions that can be set in the Live menu.
- 30. Tap the **Display** button to configure the display settings, see chapter *Display (Setup menu)* [p. 40].
- 31. Tap the **Date + Time** button to set the date, time, and display format.



- 32. Tap the **Contact Information** button to store the institution name and contact information. The contact information can be accessed via the Help function on the dashboard.
- 33. Tap the **Software Licenses** button to view a list of the open source software included in the product, and the open source software license terms.

6.3.1 Display (Setup menu)

- 1. Tap the **Display** button in the Setup menu.
 - ⇒ The following menu appears:

X	Primary Source	4U-LINK	
	Autoswitch Angle Interface	One line on approach	
Roman y	Autoswitch Angle	- 45 +	
÷1	Hysteresis Angle		
tarate	Horizon Indicator	Enabled	
*	Adjust Horizon	Enabled	
and	DP-1 Output Source	30	
	DP-2/DVI Output Source		
Deploy	SDI Output Source	30	
Đ	SDI Output Resolution	1080p 🔽	
La dien	Secondary Source (PiP)	None 🔽	
	User Interface Aspect Ratio	Widescreen	
innian General Land	Display Language	English 🔽	
1 AND	Keyboard Language	English 🔽	
	Exit		

- 2. In **PIP Layout**, select a layout to adapt the picture-in-picture (PiP) function display.
- 3. Open the Primary Source drop-down menu and select a LINK module.
 - ⇒ If the secondary source is configured with None or S-Technologies, the IMAGE1 S automatically recognizes the module to which a camera head is connected, even if the LINK module used is not the primary source.
- 4. If the IMAGE1 S CONNECT (TC200) is being used, select one of the following modes for emitting the 3D signal:
- DVI 1
- DVI 1 / DVI 2
- DVI 1 / 3G-SDI
- DVI 1, DVI 2 / 3G-SDI

All outputs that are not selected transmit a 2D image. The 3D signal is always present at the DVI 1 (see label on the IMAGE1 S CONNECT (TC200)).

- 5. If the IMAGE1 S CONNECT II (TC201) is used with connected D3-LINK, 4U-LINK, and 3D equipment, select outputs DP-1, DP-2/DVI, and SDI as 2D or 3D mode.
- 6. Open the **SDI Output Resolution** drop-down menu and select the resolution of the signal outputs (available only with IMAGE1 S CONNECT II (TC201)).



- 7. Open the **Secondary Source (PiP)** drop-down menu and select another LINK module or an S-Technology.
 - ➡ If S-Technology is selected and a visualization mode (CLARA, CHROMA, or SPECTRA) is selected, the standard white light image and the applicable visualization mode are are displayed next to one another as PiP.
 - If a D3-LINK or 4U-LINK is selected as the primary source and 3D equipment is connected, the secondary source is deactivated.
- 8. In User Interface Aspect Ratio, select Widescreen (16:9) or 4:3.
- 9. Select the Display Language and the Keyboard Language.
- 10. Open the **Autoswitch Angle Interface** drop-down menu and select one of the following settings for the autoswitch limits display behavior when a TIPCAM1 Rubina is connected:
- One line on approach
- Four lines on approach
- Four lines permanently
- 11. In **Autoswitch Angle**, tap the **Plus** or **Minus** button to set the angle between +/- 30° and +/- 60° in 5° increments. The autoswitch angle specifies the TIPCAM1 Rubina rotation angle at which the image changes from 3D to the upright 2D mode.
- 12. In **Hysteresis Angle**, tap the **Plus** or **Minus** button to set the angle between +/- 0° and +/- 15° in 3° increments.
 - ⇒ The autoswitch limit of the active area moves by the specified number of degrees. Performing work at the autoswitch limit prevents frequent switching.
- 13. Activate or deactivate the horizon indicator.
- 14. In Adjust Horizon, select whether the adjustment option for the Live menu is activated or deactivated.

6.3.2 Network

The following network parameters can be called up:

- Internet Protocol Version 4
- Network address
- Netmask
- MAC address
- DNS server



1. Tap the Network button.

⇒ The following information appears:

IPv4	Auto IP 🔽
Network Address	169.254.91.45
Subnet Mask	255.255.0.0
MAC Address	F4:5E:AB:EF:D1:1B
Gateway	0.0.0.0
DNS Server 1	0.0.0.0
DNS Server 2	0.0.0.0
KS HIVE Device Integration	Yes
KS HIVE Device Info	
Apply and Exit	
Exit	
	Network Address Subnet Mask MAC Address Gateway DNS Server 1 DNS Server 2 KS HIVE Device Integration KS HIVE Device Info

- 2. In **Patient Data Privacy**, enter the password to access the **Network** menu, see chapter *Patient data privacy* [p. 43].
- 3. Alternatively, enter the password when opening the **Network** menu for the first time.
- 4. In **KS HIVE Device Integration**, switch the KS HIVE communication interface on or off as required. KS HIVE device integration enables bidirectional communication between the IMAGE1 S and KS HIVE-compatible peripheral devices.
- 5. To change the KS HIVE settings, switch the CONNECT module off and back on again.



Menu			
x p			
ture	IPv4	Auto IP 🔽	
X	Network Address	KS HIVE Status Information	8
€ + + 100Hz	Subnet Mask	KS HIVE Certificate Derived from Developer Roote KS HIVE Peer Connected LED_light_source-EESIMULATED Yes	CA
juency	MAC Address		
	Gateway		
etwork	DNS Server 1	Cancel	
**	DNS Server 2	0.0.0	
fflation	KS HIVE Device Integra	ation Yes	
	KS HIVE Device Info		3
L)	Apply and Exit		
e + Time	Exit		-

6. In **KS HIVE Device Info**, call up information about KS HIVE with the connected devices, their availability and their status.

6.4 Patient data privacy

- (i) Do not record sensitive patient information including patient's face, name, social security number, etc.
- (1) KARL STORZ assumes no legal liability or responsibility for the loss, accuracy, damage, or disclosure of patient data that has been managed using IMAGE1 S.
- (i) The patient data administration functions (e.g. entry, access, display, storage and transfer) must be activated manually. Upon activating the patient data administration functions, the user assumes responsibility for all patient-related data which are managed using IMAGE1 S as per the applicable national data protection laws.

In the **Patient Data Privacy** settings, you can edit all settings related to patient information. Patient information is available only if the corresponding settings have been made. The system does not contain patient data as standard.

- 1. Tap the **Patient Data Privacy** button.
- 2. The **Patient Data Privacy** menu is password-protected as standard. Enter a personal password that fulfils the following criteria:
- At least 8 characters
- At least 1 uppercase letter
- At least 1 lowercase letter



- At least 1 numeral
- At least 1 special character
- 3. Repeat the password in the second line and confirm it.

	o Modify Settings	Below Activate			
reased Path					
aplay Durati		Set Password			
ow Patlent 1		And a state of the	_		
able the priv	New Password	Password accepted.			
d Patient Na	Verify	*******			
tient Data S		Password accepted.			
ve logged ev	Cancel		Confirm		
turn For Servic	e / Reset Passwor	d Reset		J	
it					
	aplay Duratik ow Patient T able the prin d Patient Na tient Data S ve logged ev turn For Servic	aplay Duratin ow Patient 1 able the prin d Patient Na Verify tient Data S ve logged en Cancel turn For Service / Reset Passwor	apiay Duration ow Patient T able the print d Patient Ns Verify Password accepted. d Patient Ns Verify Password accepted. tent Data S ve logged et turn For Service / Keset Password Accepted.	apiay Durativ ow Patient 1 able the prive d Patient Na Verify •••••••• ve logged er Cancel Confirm turn For Service / Keset Password accepted.	apiay Duratii ow Patient T able the prin d Patient Na Verify ve logged er turn For Service / Keset Password Reset Reset

- \Rightarrow A prompt appears.
- $\Rightarrow\,$ The menu can be used once the password has been entered again.
- 4. Activate the Increased Patient Data Privacy setting to apply the cybersecurity presets.
 - ⇒ If this setting is activated, all sub-settings for Patient Data Privacy will be deactivated and the default settings restored.



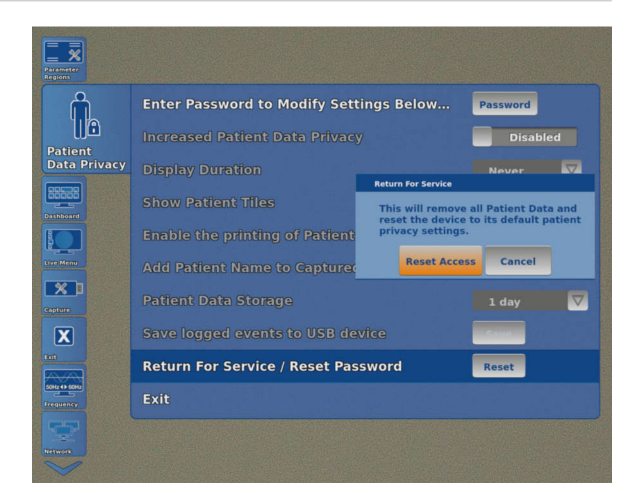
Application

Parameter Regions		
Â	Enter Password to Modify Settings Below	Password
UA Patient	Increased Patient Data Privacy	Disabled
Data Privacy	Display Duration	Never 🔽
Dashboard	Show Patient Tiles	Yes
	Enable the printing of Patient Data on Prints	Yes
Live Menu	Add Patient Name to Captured File Name	Yes
Capture	Patient Data Storage	1 day 🔽
X	Save logged events to USB device	Same
Exit	Return For Service / Reset Password	Reset
50Hz () 60Hz Frequency	Exit	
2		
Network		

Setting	Description	Default setting
Display Duration	Specify the display duration for the pa- tient data.	Never
Show Patient Tiles	Specify whether the Patient tile is to appear on the dashboard and in the Live menu.	No
Enable the printing of Patient Data on Prints	Specify whether prints will be labeled with patient information or not.	No
Add patient name to the name of the recorded file	Specify whether media files stored will be labeled with the patient information.	No
Patient Data Storage	Set the data storage period on the de- vice.	1 day
Save Logged Events to USB storage device	The audit log can be viewed via Open Log . Audit logs can be exported via Save As .	-

The **Return for Service / Reset Password** option is not affected by the **Increased Patient Data Privacy** setting. The option is available at any time and provides access to the settings of the **Patient Data Privacy** menu when the password is reset. All patient data is cleared and the default settings are restored. This function is recommended before returning the device for service.





6.5 Remote Service

- 1. Make sure that the product is registered for Remote Service. If necessary, contact the local contact person.
- 2. Make sure that software version 4.4 or higher is installed.
- 3. Make sure that IPv4 is set to DHCP in the network settings, see chapter Network [p. 41].
- 4. Check whether the date and time are set correctly, Setup menu.
- 5. Using an OR1 patch cable, connect the product (TC200, TC201) to a network connection to connect to the Internet. If necessary, use an OR1 HIVE router.



6.5.1 Setup

- 1. Tap the **Remote Service** button in the **Setup** menu.
 - \Rightarrow The following window appears:

~				
SSEE				
XI				
		Enter Password		
	Current Password	*******		
Tronsoner Retwork	Cancel		Confirm	
Ramota Starvica				

- 2. Enter the password to access the **Patient Data Privacy** and **Network** menu, see chapter *Patient data privacy* [p. 43].
- 3. Tap the **Setup** button.

Cipitale X			
	Remote Update Disabled		
البيني الم	Email Notification Disabled		
Satup	Exit		

- 4. If necessary, activate the **Remote Update** option to enable online software updates, see chapter *Performing an update* [p. 48].
- 5. If necessary, activate the **E-mail Notification** option to receive notifications about available software updates via e-mail.





6.5.2 Performing an update

- (i) The installation process may not be interrupted.
- (1) The product cannot be used during the update.
- 1. Touch the **Update** button.
 - \Rightarrow Both the installed and currently available software are displayed.
- 2. Tap the **Description** button for information on the software.
- 3. When updating the CONNECT modules and LINK modules, make sure that no camera heads or videoendoscopes are connected.
- 4. Tap the **Update** button to install the software which is currently available. Make sure that the update is performed first for the CONNECT module (TC200, TC201).
- 5. Follow the instructions on the screen.

Extract 0% Link 1 H3-LINK (TC300) Installed 4.3.dev Available 4.3.1 Link 2 X-LINK (TC301) Installed 4.3.dev Available 4.3.1 Link 3 4U-LINK (TC304) Installed 4.3.dev Available 4.3.1	CONNECT IMAGE1 S II (TC20	1) Installed 4.3.dev	Available 4.3.1
California (1998) Link 2 X-LINK (TC301) Installed 4.3.dev Available 4.3.1 California (1998)		Extract 0%	<u></u>]
	Link 1 H3-LINK (TC300)	Installed 4.3.dev	Available 4.3.1
Link 3 4U-LINK (TC304) Installed 4.3.dev Available 4.3.1	Link 2 X-LINK (TC301)	installed 4.3.dev	Available 4.3.1
	Link 3 4U-LINK (TC304)	Installed 4.3,dev	Available 4.3.1

- 6. Update the LINK module software.
- 7. If the update fails, restart the system and repeat the update.



6.6 Switching off the product

1. Switch the product off with the power switch.





7 Maintenance, servicing, repairs, and disposal

7.1 Maintenance

The following maintenance intervals are recommended:

Interval	Activity	To be performed by
annually	Safety test	KARL STORZ service techni-
		cians

7.2 Safety inspection in accordance with IEC 62353

Risk of injury due to product degradation!

Patients, users and third parties may be injured as a result of product and accessory degradation.

- Shut down the product.
- ► Have the deficiencies repaired by persons authorized by KARL STORZ.

Regardless of the national accident prevention regulations and testing intervals for medical devices, for this device safety checks must be performed as repeat inspections according to IEC 62353 and recorded by a qualified electrician at least once a year. Detailed specifications regarding the scope and execution of the safety inspection can be found in the service manual.

7.2.1 Visual inspection

- 1. Check the product and accessories for any mechanical damage.
- 2. Check labels for readability.

7.2.2 Electric measurements

- (i) Limit values for electrical measurements can be found in the current IEC 62353.
- 1. Measure the protective ground resistance.
- 2. Measure the earth leakage current.
- 3. Measure the touch current.
- 4. Measure the patient leakage current.

7.3 Repairing the product

Repair work may only be performed by KARL STORZ or by a company authorized by KARL STORZ. The maintenance steps described in this instruction manual are exempt from this rule.

 Please contact your local [Product/Brand variable] subsidiary or authorized dealer (see the list of subsidiaries).

Contaminated devices may not be shipped. To prevent contact infections and airborne infections, products must first be decontaminated. KARL STORZ reserves the right to send back contaminated products.

7.4 Disposing of the product

The product meets the requirements of the Directive on Waste Electrical and Electronic Equipment (WEEE).



Within the scope of application of this directive, KARL STORZ SE & Co. KG is responsible for the proper disposal of this product.

- 1. The product must be disposed of in accordance with the applicable national laws and regulations at a suitable collection point for the reprocessing of electrical and electronic equipment.
- 2. Contact KARL STORZ SE & Co. KG, a KARL STORZ branch or an authorized dealer to find out the address of the collection point in your area.



8 Accessories and spare parts

8.1 Accessories

Basic equipment

Article	Order no.
Videoendoscope adaptor 0°	TC001
Videoendoscope adaptor 90°	TC013
IMAGE 1 FULL HD extension cable, length 520 cm, for use with all IMAGE 1 HD camera heads – except H3-M	22220072
USB to ACC adaptor	TC009
ACC connecting cable, to control peripheral devices, length 180 cm	20221070
Power cord, length 300 cm	400A
Power cord, US version, 200 cm	400B
OR1 patch cable CAT6a 2.0 m UL	WO10275
Two-pedal footswitch	TC010
USB silicone keyboard with touchpad, CH	20040240CH
USB silicone keyboard with touchpad, DE	20040240DE
USB silicone keyboard with touchpad, ES	20040240ES
USB silicone keyboard with touchpad, FR	20040240FR
USB silicone keyboard with touchpad, IT	20040240IT
USB silicone keyboard with touchpad, PT	20040240PT
USB silicone keyboard with touchpad, RU	20040240RU
USB silicone keyboard with touchpad, SE	20040240SE
USB silicone keyboard with touchpad, US	20040240US
USB flash drive, 32 GB	20040282
DVI connecting cable, length 200 cm	20040086
DVI connecting cable, length 300 cm	20040089
DisplayPort cable, length 3 m	TC003
3G-SDI cable, length 10 m	TC005
Synch connecting cable, 3.5mm, 100 cm	TL006
12G-/3G-SDI cable, length 3 m	TC007
12G-/3G-SDI cable, length 10 m	TC008
LINK cable, length 30 cm	TC011
IMAGE1 PILOT	TC014
Extension cable IMAGE1 S D3-LINK	TC015
OR1 HIVE router	WO10240
HIVE router US	WO10307



Accessories and spare parts

Article	Order no.
USB silicone keyboard with touchpad, DE	20040241DE
USB silicone keyboard with touchpad, CH	20040241CH
USB silicone keyboard with touchpad, ES	20040241ES
USB silicone keyboard with touchpad, FR	20040241FR
USB silicone keyboard with touchpad, IT	20040241IT
USB silicone keyboard with touchpad, PT	20040241PT
USB silicone keyboard with touchpad, RU	20040241RU
USB silicone keyboard with touchpad, SE	20040241SE
USB silicone keyboard with touchpad, US	20040241US
Microscope footswitch	TC019
E-Box for X-LINK	TC028

Compatible camera heads for IMAGE1 S H3-LINK with S-Technologies

Item	Order no.
IMAGE1 S H3-Z three-chip FULL HD camera head	TH100
IMAGE1 S H3-Z three-chip FULL HD camera head	TH101
IMAGE1 S H3-Z FI three-chip FULL HD camera head	TH102
IMAGE1 S H3-P three-chip FULL HD pendulum camera head	TH103
IMAGE1 S H3-ZA three-chip FULL HD camera head	TH104
IMAGE1 S H3-M COVIEW three-chip FULL HD camera head	TH106

Compatible camera heads for IMAGE1 S H3-LINK without S-Technologies

Item	Order no.
IMAGE 1 H3-Z three-chip FULL HD camera head	22220055-3
IMAGE1 H3-ZI three-chip FULL HD inline camera head	22220053-3
IMAGE1 H3-Z FI three-chip ICG FULL HD camera head	22220085-3
IMAGE1 H3-P three-chip FULL HD pendulum camera head	22220056-3
IMAGE1 H3-ZA three-chip FULL HD camera head	22220061-3
IMAGE1 H3-FA three-chip FULL HD camera head	22220060-3

Compatible camera heads for IMAGE1 S X-LINK with S-Technologies

Item	Order no.
IMAGE1 S D1, 50 Hz	TH115
IMAGE1 S D1, 60 Hz	TH116
IMAGE1 S HX-FI	TH112
IMAGE1 S HX-P-FI	TH113



(i) The IMAGE1 S HX-FI (TH112) and IMAGE1 S HX-P-FI (TH113) camera heads do not support the CLARA mode.

Compatible camera heads for IMAGE1 S X-LINK without S-Technologies

Item	Order no.
IMAGE1 S HX	TH110
IMAGE1 S HX-P	TH111

Compatible videoendoscopes for IMAGE1 S X-LINK with S-Technologies

Item	Order no.
Video bronchoscope	11900BP
Video bronchoscope	11900BN
Video bronchoscope HD	11910T
Video bronchoscope HD	11910D
Video bronchoscope	11910P
Video bronchoscope	11910S
Video Mediastinoscope	10973HD
Video rhino-laryngoscope	11101VP
Video rhino-laryngoscope	11101VN
Video rhino-laryngoscope	11101VPS
Video rhino-laryngoscope	11101VNS
HD Video rhino-laryngoscope	11101HD
Video-gastroscope 9.3/2.8/1,100	13821PKS
Video-gastroscope 9.3/2.8/1,100	13821NKS
Slim gastroscope 5.9/2.0/1,100	13820PKS
Slim gastroscope 5.9/2.0/1,100	13820NKS
Video-gastroscope 12/4.2/1,100	13807PKS
Video-gastroscope 12/4.2/1,100	13807NKS
Video-gastroscope 12/2.8-3.4/1,100	13806PKS
Video-gastroscope 12/2.8-3.4/1,100	13806NKS
Video-gastroscope 9.7/2.8/1,100	13801PKS
Video-gastroscope 9.7/2.8/1,100	13801NKS
Video duodenoscope 12.6/4.2/1,260	13885PKS
Video duodenoscope 12.6/4.2/1,260	13885NKS
FRIMBERGER Video duodenoscope	13883PKS
Video duodenoscope 12.6/4.2/1,250	13882PKS
Video duodenoscope 12.6/4.2/1,250	13882NKS
Video colonoscope 12.9/3.8/1,600	13925HPKS



Accessories and spare parts

Item	Order no.
Video colonoscope 12.9/3.8/1,600	13925HNKS
Video colonoscope 12.9/3.8/1,600	13925PKS
Video colonoscope 12.9/3.8/1,600	13925NKS
Video colonoscope 12.9/3.8/1,400	13924HPKS
Video colonoscope 12.9/3.8/1,400	13924HNKS
Video colonoscope 12.9/3.8/1,400	13924PKS
Video colonoscope 12.9/3.8/1,400	13924NKS
Video colonoscope 10.6/2.8/1,600	13907PKS
Video colonoscope 10.6/2.8/1,600	13907NKS
Video colonoscope 10.6/2.8/1,400	13906PKS
Video colonoscope 10.6/2.8/1,400	13906NKS
Video colonoscope 13/3.8/1,600	13905PKS
Video colonoscope 13/3.8/1,600	13905NKS
Video colonoscope 13/3.8/1,400	13904PKS
Video colonoscope 13/3.8/1,400	13904NKS
Medium gastroscope	13823PKS
Medium gastroscope	13823NKS
Sigmoidoscope	13920PKS
Sigmoidoscope	13920NKS
Slim colonoscope	13926PKS
Slim colonoscope	13926NKS
Slim colonoscope	13927PKS
Slim colonoscope	13927NKS
Interventional gastroscope	13826PKS
Interventional gastroscope	13826NKS
Video uretero-renoscope FLEX-X C	11278VSU
Video uretero-renoscope FLEX-X C	11278VSUA
Video uretero-renoscope FLEX-X C	11278VSUE
Video uretero-renoscope FLEX-X C	11278VS
Video uretero-renoscope FLEX-X C	11278VSA
Video uretero-renoscope FLEX-X C	11278VSE
Flexible video urethro-cystoscope	11272VP
Flexible video urethro-cystoscope	11272VPU
Flexible video urethro-cystoscope	11272VN
Flexible video urethro-cystoscope	11272VNU
Flexible HD video cystoscope	11272VH



Accessories and spare parts

Item	Order no.
Flexible HD video cystoscope	11272VHU
TROIDL flexible rectoscope	13912PKS
TROIDL flexible rectoscope	13912NKS
Video choledochoscope	11292VP
Video choledochoscope	11292VPU
Video choledochoscope	11292VS
Video choledochoscope	11292VSU
Flexible video choledochoscope	11292VSA
Flexible video choledochoscope	11292VSUA
Flexible video choledochoscope	11292VH
Video hysteroscope	11264V

Compatible single-use videoendoscopes for IMAGE1 S X-LINK with E-Box

Item	Order no.
FIVE S 3.5x65	091361-06
FIVE S 5.3x65	0915612-06
Video rhino-laryngoscope	091330-06
Video esophagoscope	091370-06
Video-uretero-renoscope FLEX-X ^{C1}	091271-06
Video-uretero-renoscope FLEX-X ^{C1}	091279-06

Compatible videoendoscopes and camera heads for IMAGE1 S D3-LINK with S-Technologies

Item	Order no.
TIPCAM1 S 3D, 0°, dia. 10 mm,	26605AA
TIPCAM1 S 3D, 30°, dia. 10 mm	26605BA
TIPCAM1 S 3D, 0°, dia. 4 mm	7240AA3D
TIPCAM1 S 3D, 30°, dia. 4 mm	7240BA3D
TIPCAM1 S 3D, 0°, dia. 4 mm	28164AA3D
TIPCAM1 S 3D, 30°, dia. 4 mm	28164BA3D
TIPCAM1 S 3D, 45°, dia. 4 mm	7240FA3D
TIPCAM1 S 3D, 45°, dia. 4 mm	28164FA3D
VITOM 3D, dia. 4 mm	TH200



Compatible camera heads and videoendoscopes for IMAGE1 S 4U-LINK with S-Technologies in combination with IMAGE1 S CONNECT II

Item	Order no.
IMAGE1 S 4U	TH120
IMAGE1 S 4U Rubina	TH121
TIPCAM1 Rubina, 0°	26606ACA
TIPCAM1 Rubina, 30°	26606BCA

The IMAGE1 S 4U camera head can also be used together with the IMAGE1 S CONNECT.

Operable SCB light sources

Item	Order no.
XENON 300 SCB	20133120-1
XENON 175 SCB	20132120-1
POWER LED 175 SCB	20161420-1
XENON 100 SCB	20132620-1
Cold light fountain CO2mbi LED SCB	TL100
Cold light fountain power LED 300	TL300
D-LIGHT C	20133620-1
D-LIGHT C/AF	20133620-133
D-LIGHT P	20133720-1

Operable KS HIVE light sources

Item	Order no.
Power LED Rubina	TL400
Power LED Saphira	TL410

Operable SCB insufflators

Item	Order no.
THERMOFLATOR SCB	26432020-1
ENDOFLATOR SCB	26430520-1
ENDOFLATOR 40	UI400
ENDOFLATOR 50	UI500



3D monitors

Item	Order no.
3D monitor (FHD), 26"	9826NB-3D
3D monitor (FHD), 32"	9832NB-3D
3D monitor (FHD), 32"	TM323
3D monitor (FHD), 32"	TM330
3D monitor (FHD), 26"	TM263
4K 3D monitor, 32"	TM350
4K/3D monitor, 55"	TM450

2D monitors

Item	Order no.
2D monitor (FHD), 19"	9619NB
2D monitor (FHD), 26"	9626NB
2D monitor (FHD), 27"	9627NB
2D monitor (FHD), 24"	WM100
2D monitor (FHD), 27"	TM220
4K 2D monitor, 32"	TM324
4K/2D monitor, 32"	TM340
4K 2D monitor, 31"	TM341
4K 2D monitor, 31"	TM342
4K 2D monitor, 58"	TM440
4K 2D monitor, 32"	TM343
4K 2D Monitor, 43"	TM441

KS HIVE compatible motorized support arms

Item	Order no.
ARTIP CRUISE	UR500B
ARTIP CRUISE	UR500C

Compatible veterinary videoendoscopes for IMAGE1 S X-LINK with S-Technologies

Item	Order no.
Veterinary videoendoscope	60914NKS
Veterinary videoendoscope	60914PKS
Veterinary videoendoscope	60714NKS
Veterinary videoendoscope	60714PKS
Veterinary videoendoscope	60511NKS
Veterinary videoendoscope	60511PKS



Accessories and spare parts

Item	Order no.
Veterinary videoendoscope	60332NKS
Veterinary videoendoscope	60332PKS
Veterinary videoendoscope	60130NKS
Veterinary videoendoscope	60130PKS
Veterinary videoendoscope	60125NKS
Veterinary videoendoscope	60125PKS
Veterinary videoendoscope	60118NKS
Veterinary videoendoscope	60118PKS
Veterinary videoendoscope	60278VS
Veterinary videoendoscope	60278VSU



9 Electromagnetic compatibility

9.1 General notes on the operating environment

The product is suitable for use in professional healthcare settings. Professional healthcare facilities include physician offices, dental offices, limited care facilities, freestanding surgical centers, freestanding birth centers, multiple treatment facilities, hospitals (emergency rooms, patient rooms, intensive care, surgical rooms, outside the HF-shielded room of an ME system for MRT).

(i) The emission characteristics of this product make it suitable for use in industrial areas as well as in hospitals (CISPR 11 Class A) and other professional healthcare environments. If it is used in a residential environment (for which CISPR 11 Class B is normally required), the product may not offer sufficient protection for radio transmission operation. The user might need to take mitigation measures, such as relocating or re-orienting the product.

MR unsafe!

This product is MR unsafe.

▶ Keep the product away from magnetic resonance imaging scanner room.

A WARNING

Electromagnetic interferences! Malfunction!

Use of this equipment adjacent to or stacked with other equipment could result in improper operation.

- Avoid this situation.
- If such use is necessary: Ensure that this equipment and the other equipment are operating normally.

Reduced immunity or increased emissions! Malfunction!

Use of the product with accessories, transducers and cables other than those specified in this manual may result in increased emissions or decreased immunity.

• Only use the accessories specified in the manual.

9.2 Accessories and cables

Degradation of performance! Malfunction!

Portable RF communications equipment (including peripherals such as antenna cables and external antennas) could result in degradation of the performance of the product.

• Do not use portable communications equipment closer than 30 cm (12 inches) to any part of the product, including cables.



System cables and maximum lengths used for EMC compliance.				
Type Shield Längth [m] Ferrite Us		Use		
LINK cable (IML ca- ble)	Yes	2	No	Interconnection of intermodule LINK signals
BNC to BNC (12G- SDI cable)	Yes	10	No	Connection of 3G/12G-SDI video signals
DVI	Yes	3	Yes	Connection of DVI signals
4-pin Mini DIN	Yes	1	No	Connection of SCB signals
Power cord	No	2	No	Connection of device to AC Mains
Ethernet - CAT6A	Yes	2	No	Connection of HIVE signals

9.3 Table 1 – Compliance level for immunity tests

Guidelines and manufacturer's declaration - electromagnetic immunity

The product is intended for use in the electromagnetic environment specified below. The user of the product should make sure that it is used in such an environment.

Immunity tests	EN/IEC 60601 test level	Compliance level	Electromagnetic envi- ronment – guidelines	
Electrostatic dis- charge (ESD) acc. to IEC 61000-4-2	± 8 kV contact discharge ± 15 kV air discharge	± 8 kV contact discharge ± 15 kV air discharge	Floors should be made of wood, concrete, or cov- ered with ceramic tiles. If floors are covered with synthetic material, the rel- ative humidity must be at least 30%.	
Electrical fast tran- sients/bursts acc. to IEC 61000-4-4	 ± 2 kV for power lines ± 1 kV for input and output lines 100 kHz repetition 	± 2 kV for power lines ± 1 kV for input and out- put lines 100 kHz repetition	The power supply quality should be that of a typical commercial or hospital environment.	
Surges acc. to IEC 61000-4-5	±1 kV voltage outer con- ductor – outer conductor ±2 kV voltage outer con- ductor – ground	±1 kV voltage outer con- ductor – outer conductor ±2 kV voltage outer con- ductor – ground	The power supply quality should be that of a typical commercial or hospital environment.	
Voltage dips, short interruptions, and voltage variations acc. to IEC 61000-4-11	Voltage dip: Dip to 0% for 1 cycle at 0° phase angle Dip to 70% for 25/30 cy- cles at 0° phase angle Dropout to 0% for 0.5 cycles @ 0°, 45°, 90°, 135°, 180°, 225°, 270°, and 315° phase angles Voltage interruption: 100% for 250/300 cy- cles	Voltage dip: Dip to 0% for 1 cycle at 0° phase angle Dip to 70% for 25/30 cy- cles at 0° phase angle Dropout to 0% for 0.5 cycles @ 0°, 45°, 90°, 135°, 180°, 225°, 270°, and 315° phase angles Voltage interruption: 100% for 250/300 cy- cles	The power supply quality should be that of a typical commercial or hospital environment. If the user of the product requires continued operation in the event of interruptions to the power supply net- work, it is recommended that the product be oper- ated with an uninterrupt- ible power supply or a battery.	

Electromagnetic compatibility



Immunity tests	EN/IEC 60601 test level	Compliance level	Electromagnetic envi- ronment – guidelines
Magnetic field at the power fre- quency (50/60 Hz) acc. to IEC 61000-4-8	30 A/m at 50 Hz / 60 Hz	30 A/m at 50 Hz / 60 Hz	If image distortion occurs, it may be necessary to in- stall the product further from sources of electro- magnetic fields or to in- stall magnetic shielding. Before the product is in- stalled, the electromag- netic field should be mea- sured to ensure that it is sufficiently low.
Immunity test acc. to IEC 61000-4–3	3 V/m 80 MHz to 2.7 GHz	3 V/m 80 MHz to 2.7 GHz	
for radiated, radio- frequency electro- magnetic fields	* Refer to Table 2 for wireless proximity RF field test levels		
Immunity to con- ducted distur-	3 V _{rms} on 150 kHz to 80 MHz	3 V _{rms} on 150 kHz to 80 MHz	
bances, induced by radio-frequency	1 kHz 80% AM modula- tion	1 kHz 80% AM modula- tion	
fields acc. to IEC 61000-4-6	$6 V_{ms}$ in ISM band	$6 V_{ms}$ in ISM band	

9.4 Table 2 – Test levels for near fields from HF wireless communications equipment

Test frequency MHz	Frequency band MHz	Radio service	Modulation	Immunity test level V/m	Compliance level V/m
385	380 – 390	TETRA 400	Pulse modula- tion 18 Hz	27	27
450	430 – 470	GMRS 460, FRS 460	FM ± 5 kHz de- viation 1 kHz sine wave	28	28
710	704 – 787	LTE band 13	Pulse modula-	9	9
745		and 17	and 17	tion 217 Hz	
780					
810	800 – 960	GSM 800/900,	Pulse modula-	28	28
870		TETRA 800, iDEN 820,	tion 18 Hz		
930		CDMA 850, LTE band 5			
1720	1700 – 1990	GSM 1800,	Pulse modula-	28	28
1845		CDMA 1900, GSM 1900, DECT,	tion 217 Hz		



Test frequency MHz	Frequency band MHz	Radio service	Modulation	Immunity test level V/m	Compliance level V/m
1970		LTE band 1, 3, 4, 25, UMTS			
2450	2400 – 2570	Bluetooth, WLAN 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modula- tion 217 Hz	28	28
5240	5100 – 5800	WLAN 802.11	Pulse modula-	9	9
5500		a/n	tion 217 Hz		
5785					

9.5 Table 3 – Test levels for radiated and conducted immunity tests

Guidance and manufacturer's declaration - Electromagnetic immunity

The product is intended for use in the electromagnetic environment specified below. The user should make sure that it is used in such an environment.

Immunity tests	EN/IEC 60601 test level	Compliance level	Electromagnetic environ- ment – Guidance	
Conducted RF distur- bances acc. to IEC 61000-4-6	3 V _{rms} 150 kHz to 80 MHz 6 V _{eff} in ISM ands be- tween 0.15 MHz and 80 MHz	3 V _{rms} 150 kHz to 80 MHz 6 V _{eff} in ISM ands between 0.15 MHz and 80 MHz	Portable and mobile RF communications equipment should be used no closer to any part of the product, in- cluding cables, than the rec ommended safety distance calculated from the equation applicable to the frequency	
Radiated RF disturbances acc. to IEC 61000-4-3	3 V/m 80 MHz to 2.7 GHz	3 V/m 80 MHz to 2.7 GHz	of the transmitter. Recommended safety dis- tances: $d = 1.2 \sqrt{P}$ Where P is the rated power of the transmitter in watts [W] according to the infor- mation provided by the transmitter manufacturer and d is the recommended safety distance in meters [m]. The field strengths of fixed transmitters as determined by an electromagnetic site survey ^a should be less than the compliance level ^b in all frequency ranges.	



Immunity tests	EN/IEC 60601 test level	Compliance level	Electromagnetic environ- ment – Guidance
			<i>d</i> = 1.2 √P 80 MHz to 800 MHz
			<i>d</i> = 2.3 √ <i>P</i> 800 MHz to 2.7 GHz
			Interference may occur in the vicinity of devices marked with the following symbol:
			(((••))

Note: At 80 MHz and 800 MHz, the higher frequency range applies.

Note: These guidelines may not apply in all situations. The propagation of electromagnetic waves is affected by absorptions and reflections of buildings, objects, and people.

^a Field strengths from fixed transmitters, e.g., 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 with regard to fixed transmitters, an electromagnetic site survey should be considered. If the measured field strength at the location where the product is being used exceeds the compliance levels above, the product should be monitored to ensure that it is functioning as intended. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the product.

^b Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

9.6 Table 4 – Emission class and group

Guidelines and manufacturer's declaration - Electromagnetic emissions

The product is intended for use in such an environment as specified below. The customer or user of the product should ensure that it is used in such an environment.

Interference emission measure- ments	Compliance	Electromagnetic environment – guidelines	
RF emissions acc. to CISPR 11	Group 1	The device 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 acc. to CISPR 11	Class A	The device is suitable for use in all es- tablishments other than domestic and those directly connected to the public low voltage power supply network that supplies buildings used for domestic purposes.	
Emission of harmonic oscillations acc. to IEC 61000-3-2	Class A		
Voltage fluctuations/flicker emissions acc. to IEC 61000-3-3	Complies		



9.7 Table 5 – Recommended separation distances between portable and mobile HF communications devices and the product

The product is intended for use in an electromagnetic environment in which HF disturbances are controlled. The customer or user of the product can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile HF communications equipment (transmitters) and the product as recommended below, according to the output energy of the communications equipment.

Rated maximum out-	Separation distance d [m] according to frequency of transmitter			
put power of the transmitter [W]	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.7 GHz	
	$d = 1.2 \sqrt{P}$	$d = 1.2 \sqrt{P}$	d = 2.3 √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 whose maximum rated power is not listed in the table above, the recommended separation distance d in meters (m) can be estimated using the equation from the respective column, whereby P is the maximum rated power of the transmitter in watts (W) according to the transmitter manufacturer.

Note: At 80 MHz and 800 MHz, the separation distance for higher frequency range applies.

Note: These guidelines may not apply in all situations. The propagation of electromagnetic waves is affected by absorptions and reflections of buildings, objects, and people.

(i) The product was tested for compatibility with HF surgical devices in accordance with IEC 60601-2-2 Appendix BB.



Subsidiaries

10 Subsidiaries

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LZA705 • EN • V3.0 • 10-2022 • IFU • CE-MDR

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