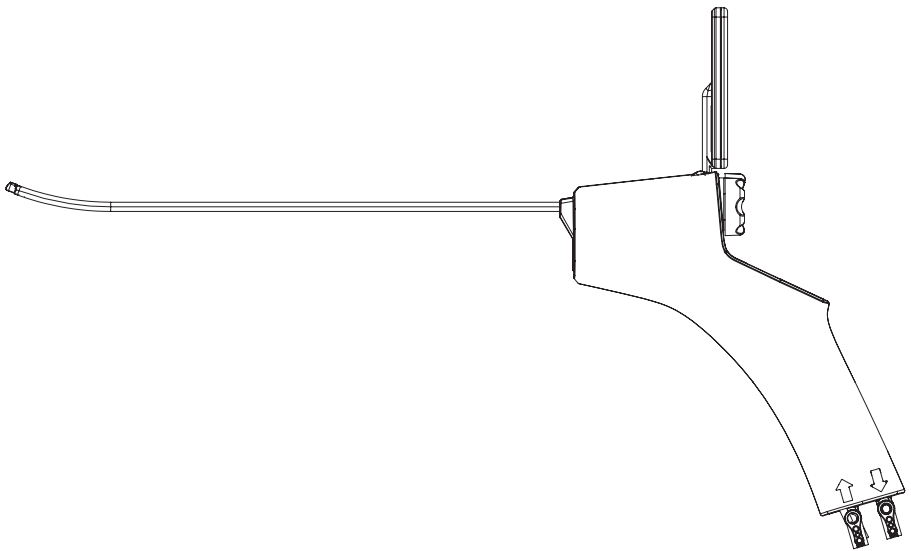




LiNA OperåScope™





LiNA OperaScope™ Instructions for Use

INTENDED USE AND PRODUCT DESCRIPTION

The LiNA OperaScope™ is intended for use in visualization of the cervical canal and uterine cavity during diagnostic and therapeutic gynecological procedures.

The LiNA OperaScope™ is a single-use battery operated hysteroscope with a cannula and LCD display with a 3m HDMI cable attached to the handle (**REF OP-201-1**). The OperaScope is provided sterile via ethylene oxide and the battery pack is provided sterile via radiation.

The 4.2 mm (nominal) diameter cannula contains a miniature camera and LED illumination at the distal tip, and a pre-curved tip section of the cannula is rotated by a turning knob (**Figure 7**) on the handle. The center of this turning knob is the insertion channel for introduction of Hysteroscopic instruments/accessories up to 1.86 mm (5.5 Fr) nominal in diameter and 310mm minimum working length. Distention and flushing of the uterine cavity with distention medium is facilitated with separate in-flow and out-flow luer lock stopcocks at the bottom of the handle. The HDMI cable at the bottom of the handle can be connected to an external monitor with HDMI input, or alternatively connected to the OperaScope™ Recording Module™ for recording of video or pictures to a standard USB drive.

During operation, visualization can be optimized by adjusting the brightness (illumination). The brightness is controlled by brightness button (**Figure 5**) located on the handle.

In addition to the OperaScope, the OperaScope Recording Module (**Figure 4**) is available to capture both still and video images for later viewing, The OperaScope Recording Module is provided nonsterile and designed to be connected to the OperaScope HDMI cable. The OperaScope Recording Module is not intended to come in contact with the sterile field or the patient.

Caution: Federal Law (USA) restricts the device to sale by or on the order of a physician.

The LiNA OperaScope™ is designed with 60 minutes of battery life.

The LiNA OperaScope™ and Recording Module are to be stored under room temperature and dry conditions.

ACCESSORIES

The following accessories can be used with LiNA OperaScope:

- OperaScope Recording Module
- Commercially available fluid management tubing with a male luer lock
- Commercially available devices for diagnostic/therapeutic use

The types of procedures where the OperaScope could offer visualization include:

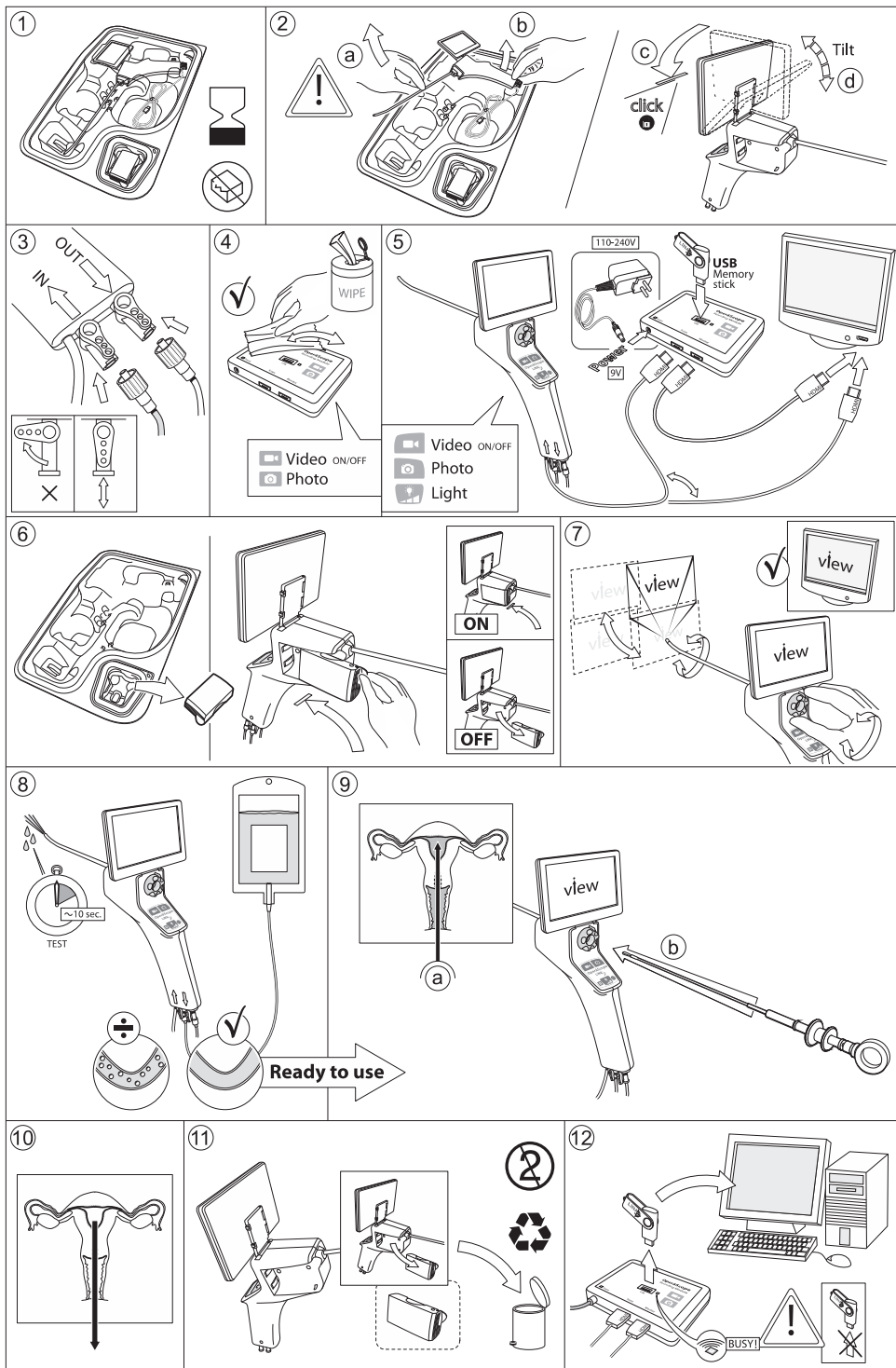
- Assessment of abnormal bleeding, pelvic pain, amenorrhea and abnormal findings from hysterosalpingogram;
- Assessment of infertility and pregnancy wastage;
- Confirmation of the presence of intrauterine foreign body;
- Assist in locating of submucosal fibroids and polyps targeted for removal;
- Provide visual guidance during directed biopsy, submucosal myomectomy, transection of intrauterine adhesions and septa.

CONTRAINDICATIONS:

General

The device is contraindicated for use in:

- Inability to distend the uterus
- Cervical Stenosis
- Cervical/Vaginal infection
- Uterine bleeding or menses





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- Known Pregnancy
- Known carcinoma of the cervix and/or the uterus
- Recent uterine perforation
- Known Pelvic Inflammatory Disease (PID)
- Medical contraindication or intolerance to anesthesia

WARNINGS:

Failure to follow all instructions or any warnings or precautions could result in serious patient injury.

- The LiNA OperaScope™ is provided STERILE via ethylene oxide (OperaScope™) and radiation sterilization (Battery). Carefully inspect the packaging for any damage prior to use. Do NOT attempt to use the device if sterile barrier is damaged. Do NOT use past expiration date. Do NOT use if the device is exposed to non-sterile surfaces before procedure.
- To mitigate the risk of perforation, only advance/manipulate the OperaScope while viewing a live camera image and illumination, allowing observation of the cervical and endometrial cavity.
- Low light intensity at the lowest available setting may limit visualization and be unsuitable for surgical procedures. Device operators should check that the light intensity range is suitable for the planned procedure prior to use.
- Do not advance the OperaScope should resistance be experienced while in the patient.
- Do not attempt to alter the shape of the distal cannula.
- For single use only. Do NOT reuse, reprocess or re-sterilize the LiNA OperaScope™. Any reprocessing may impede the functions of this device. Reusing single use devices may also increase the risk of cross contamination. Attempts to clean the device results in risk of device malfunction and/or erroneous pathology specimen collection due to residual tissue in the LiNA OperaScope™.
- Hysteroscopy procedures using the LiNA OperaScope™ should be performed only by medical professional who have adequate training in hysteroscopy.
- Do not remove the battery during the treatment process.
- Always have a backup device readily available for immediate use.
- For use only with the batteries provided with each LiNA OperaScope™ device – batteries are identified as AA Energizer alkaline batteries in a dedicated battery pack for the LiNA OperaScope™.
- LiNA OperaScope™ is designed with 60 minutes of battery life.
- If any malfunction should occur during use or battery life expires, stop the procedure immediately, and slowly withdraw the LiNA OperaScope™ and replace with a new LiNA OperaScope™.
- A liquid distension medium is used and strict fluid intake and output surveillance should be maintained. Intrauterine instillation exceeding 1 liter should be followed with great care to reduce the possibility of fluid overload.
- Portable RF communication equipment (including peripherals such as antenna cables and external antennas) should not be used within 30 cm (12 inches) of the OperaScope, as degradation of OperaScope performance could result.
- Use of accessories, transducers and cables other than those specified could result in a change in electromagnetic properties resulting in potential improper operation.
- Use of the OperaScope equipment adjacent to other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.
- Avoid submersing the OperaScope handle, battery, HDMI port and/or LCD screen in a fluid bath. Should the OperaScope be submerged, ensure the OperaScope is dried off and remains operational before clinical use. If the OperaScope is nonoperational, replace with a new device.

US specific warnings:

- Federal Law (USA) restricts the device to sale by or on the order of a physician
- The LiNA OperaScope™ is intended only as an adjunct in assessing patient condition. It must be used in conjunction with clinical signs and symptoms.



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PRECAUTIONS:

Potential complications of Continuous Flow Hysteroscopy:

- Hyponatremia
- Hypothermia
- Uterine perforation resulting in possible injury to bowel, bladder, major blood vessels and ureter.
- Pulmonary edema
- Cerebral edema
- Infection
- Bleeding
- Pain

The use of normal saline as a distending medium, and limiting the infused volume to less than 1000 ml is recommended to decrease the risk of the above complications. Intrauterine distension can usually be accomplished with pressures in the range of 35 - 75 mmHg. Unless the systemic blood pressure is excessive, it is seldom necessary to use pressures greater than 75-80 mmHg.

Before each use or after a change of OperaScope settings, check to ensure OperaScope view provides a live image (rather than a stored one) and has the correct image orientation.

If a continuous flush/irrigation during OperaScope operation is not maintained, the distal tip operating temperature may exceed 41°C (105.8°F).

While the OperaScope Recording Module is connected to the OperaScope, ensure that the power supply unit is easily accessible if power should need to be disconnected.

Ensure the OperaScope Recording Module and monitor are outside of the patient's reach.

INSTRUCTION FOR USE:

Read the instruction for use prior to using this device!

1. Carefully inspect the packaging for any damages prior to use. Do NOT attempt to use the device if the sterile barrier is damaged. Do NOT use past expiration date.
2. Using sterile technique, remove the hysteroscope from the sterilized blister (**Figure 2**).
3. Attach the in flow fluid tubing to the inflow port and the out flow tubing to the outflow port (**Figure 3**).
4. OperaScope can be connected via HDMI cable to an external monitor or the Recording Module.

Note: Prior to using the Recording Module, ensure the unit is plugged in and green power light is illuminated.

5. Remove the battery from the sterilized blister and insert the battery into the OperaScope Handle (**Figure 6**).
6. The light source at the distal tip of the cannula will illuminate and the camera image should be visible on the onboard LCD. If illumination or camera image is not functional, discontinue setup and open a new OperaScope (**Figure 7**).
7. Flush the fluid through the cannula until all air bubbles have cleared from the tubing (**Figure 8**).
8. The hysteroscope is now ready for use.
9. With fluid flowing, the cannula is inserted through the cervical canal while viewing the image, as is standard procedure for introduction of a hysteroscope (**Figure 9**).
10. If picture or video recording is desired, insert a USB memory stick into the Recording Module and make sure the HDMI cable is connected from the OperaScope. Both the OperaScope handle and Recording Module have picture and video capture buttons. Press and release the picture button to capture a still image. Press and release video button to start recording; press and release the video button again to stop recording (**Figure 13**).
11. The LiNA OperaScope has central lumen located within the turning knob of the Handle (**Figure 9**) and provides a conduit to delivery diagnostic and therapeutic devices to uterine cavity. Devices within the following dimensions are compatible with the OperaScope central lumen:



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- Maximum outer diameter: 1.86 mm (5.5 Fr)
 - Minimum working length: 310 mm
12. After the procedure is completed, remove the hysteroscope (**Figure 10**).
 13. Remove the battery pack and dispose of the battery. The device may now be safely disposed in accordance with local governing ordinances (**Figure 12**).
 14. Remove USB stick from Recording Module and transfer the recordings to a computer using a USB port.

Technical specification for LiNA OperaScope

Nominal input voltage: 3V DC ±10%
Nominal Current: 330mA ±20%
Battery type: 2pcs AA 1.5V Energizer Alkaline (Non-rechargeable)
Operating time: ≥60 minutes
Camera resolution: 400x400
Diagonal size of LCD display: 4.3 Inch
Minimum usable length: 240mm
Pre-fixed tip angle: 20° ±5°

Shelf Life Conditions

The labeled shelf life was based on test units that were real time aged for 3 months at a nominal temperature of 22oC. The test units were then subjected to simulated shipment conditions per ASTM D4169, Cycle 13, assurance level 1. All units were inspected and subjected to design verification testing and met all acceptance criteria.

OPERASCOPE RECORDING MODULE

Download of Video/Still Imagery Files Post Procedure

Video and still imagery can be downloaded from the OperaScope via the OperaScope Recording Module USB and imported into the patient's medical record. Since there is no patient identifying information on the video/still imagery file, the downloaded file should be uploaded and saved to the patient's medical record immediately post procedure ensuring the proper link between the file and the patient.

Playback of recorded Video/Still images on computer

- Step 1
Insert the USB-stick into the computer USB port
- Step 2
Download and install some supplement codec software for Windows Media Player such as Media Player Codec Pack.
- Step 3
Click "Start" in Windows and select the folder in which your H.264 file is located, such as My Videos or My Downloads.
- Step 4
Right-click on the video and select "Open With." Choose "Windows Media Player" from the available options to begin playing your video/Images.

Of note, the play back on some computers with older software may occur at double speed.

Post Procedure Cleaning

The OperaScope Recording Module is intended to be used outside the sterile field and away from the patient. Where the OperaScope Recording Module is not intended to come in direct contact with patient, the device can be exposed to body fluids and/or blood as it is used in the operating room during patient treatment.

After each patient use, the following cleaning/disinfecting instructions to ensure proper cleaning of the external components



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DO NOT SUBMERGE OPERASCOPE RECORDING MODULE

DURING CLEANING/DISINFECTING, ORIENT THE OPERASCOPE RECORDING MODULE POSITION TO MINIMIZE LIQUID FROM COMING IN CONTACT WITH THE POWER, USB, SCOPE AND MONITOR CONNECTORS

OPERASCOPE RECORDING MODULE CASE SHOULD NOT BE OPENED FOR CLEANING/DISINFECTING

1. Power off and unplug the OperaScope Recording Module
2. Place the OperaScope Recording Module and power cord on a clean, dry surface
3. Using a cleaning/disinfecting wipe, completely wipe the top, bottom and all 4 sides of the OperaScope Recording Module.
4. After wiping, visually inspect the exterior for residual bodily fluids or dirt and repeat wipe.
5. Let the OperaScope Recording Module sit for at least 3 minutes.
6. Using a 70% Isopropyl swab, thoroughly wipe the seams of the Module ensuring the Module is free of bodily fluid.
7. Once the Module is visually clean, place the device in a storage area for future use.

OperaScope Recording Module Battery 2-year Shelf Life:

The OperaScope Recording Module battery has an operating life of 2 years from the date of manufacture and will be replaced prior to the anticipated shelf life expiration.

Nominal input voltage: 100-240V AC ±10%

Nominal input frequency: 50-60Hz

Nominal input current: 0.16-0.08Arms @ max load

Nominal output voltage: Uout: 9V DC +5% / -5%

Nominal output current Iout: 800mA

AC adapter provided: Friwo Fox 9VDC/800mA

Complies with EN 60601-1-2: 2015 for EMC.

- Note: the emissions characteristics of this equipment make it suitable for use in industrial areas and hospitals/clinics (CISPR 11 Class A). The equipment is not indicated for use outside a hospital or clinic setting.
- Immunity according to Professional healthcare facility environment

	Catalog Number		Consult Instructions for Use
	Lot Number		Do Not Use Device If Package Is Damaged
	Manufacturer		Keep Dry
	Authorized Representative in European Community		Sterilized Using Ethylene Oxide
	Use-by Date		Do Not Resterilize
	Prescription		Do Not Reuse
	Consult Operations Manual		Caution, Consult Accompanying Documents



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Open Source Software used in OperaScope Recording Module

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DK-2600 Glostrup
Denmark
www.lina-medical.com

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