

Eikon[®] LT Adapt Smoke Evacuation Retractor

REF

A90WSE / A90WTSE / A135WSE / A135WTSE / A155WSE / A155WTSE / A175WSE / A175WTSE

Instructions for Use



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Eikon LT Adapt Smoke Evacuation Retractor Instructions for Use

Description

The Eikon LT Adapt Smoke Evacuation Retractors are intended for use in a wide variety of surgical procedures. The retractors and associated slats are non-sterile, reusable instruments comprised of a non-conductive reinforced polymer. The retractors are compatible with the PhotonGuide Adapt Illuminator and the Invuity Single and Bifurcated Fiber Optic Cables.

When assembled, the Eikon LT Adapt Smoke Evacuation Retractors have an integrated channel that enables smoke to be evacuated from the surgical site.

Indications for Use

The Eikon LT Adapt Smoke Evacuation Retractor is intended to provide tissue retraction and surgical site illumination from a high intensity light source. Additionally, the device is intended to remove smoke generated by electrosurgery when used in conjunction with an effective smoke evacuation system.

Contraindications

There are no known contraindications.

For Use With

DESCRIPTION	REF
PhotonGuide Adapt Illuminator	PGA1

Warnings

- <u>Prior to each use, inspect the Retractor and Slat for damage and integrity</u>, to ensure that it is free from debris and defects such as cracks, chips, rough edges, or other general damage. Assess damaged instruments for potential hazard. Do not use damaged instruments.
- Only use Retractors with the PhotonGuide Adapt Illuminator and Invuity Fiber Optic Cable products. Using non-Invuity fiber optic cables may generate excessive heat at the connector and cause thermal injury to the patient or user.
- If the Retractor or Slat is/was used in a patient with, or suspected of having Creutzfeldt-Jakob Disease (Mad Cow Disease), the instrument cannot be reused and must be destroyed due to the inability to reprocess or sterilize to eliminate the risk of cross-contamination. Consult WHO and local regulations for further information.

Precautions

- The Eikon LT Adapt Smoke Evacuation Retractor should be handled and operated by hospital personnel familiar with and trained with their use, assembly, and disassembly
- Use of the Retractor for any purpose or in any manner other than described in this Instructions for Use may cause instrument damage or failure, which could result in patient or user injury.
- The Retractors and associated Slats are provided non-sterile and must be thoroughly cleaned and sterilized prior to use.
- It is recommended to use the appropriate Invuity Sterilization Tray to help protect the Eikon LT Adapt Smoke Evacuation Retractor from damage during transport, decontamination, sterilization, and storage. Space instruments far enough apart to avoid contact with metal instruments or other objects which may dent, ding, scrape or otherwise damage the surface of the instrument
- The Retractors are delicate medical devices and must be handled with care during cleaning and sterilization. If the instruments are not handled appropriately, surface abrasions may occur.
- The Eikon LT Adapt Smoke Evacuation Retractor and Slat must be disassembled prior to cleaning and sterilization.
- If a PhotonGuide Adapt Illuminator is inadvertently left in the retractor, remove the disposable before continuing the cleaning and sterilization process.
- Do not use Sterrad® or other sterilization processes that contain chlorine (bleach) as these are known to cause discoloration of the surface.
- To prevent stains, use distilled or demineralized water and a neutral detergent when cleaning the Eikon LT Adapt Smoke Evacuation Retractor.

- Refer to the Instructions for Use for the Invuity PhotonGuide Adapt Illuminator and Fiber Optic Cables for additional information
- To prevent inadequate suction levels, ensure barb connector of Eikon LT Adapt Smoke Evacuation Retractor is securely connected to the secondary suction hose.
- Ensure that the Instructions for Use provided with the Smoke Evacuation Unit are followed, including all user instructions, warnings, and precautions.

Directions for Use

The following are general directions for the Eikon LT Adapt Smoke Evacuation Retractors:

- 1. Inspect the Retractor and Slat for cleanliness and damage prior to assembling. Do not use if the product is damaged.
- 2. Prior to use, the Eikon LT Smoke Evacuation Retractor must be assembled. First, identify the correct size Slat for the Retractor. Next, fully insert the Slat into the slot in the retractor in the orientation shown below.



- 3. Identify the PhotonGuide Adapt Illuminator and Fiber Optic Cable for the Retractor.
- 4. Securely attach the Fiber Optic Cable and the PhotonGuide Adapt Illuminator to the Retractor. Refer to the PhotonGuide Adapt Illuminator IFU for further details regarding installation in appropriate retractors.
- 5. Connect the Fiber Optic Cable to the ACMI port on the surgical light source. It is recommended to use an LED light source. The lower intensity light output from an LED light source reduces the risk of thermal injury to patient or user or to the PhotonGuide Adapt Illuminator. LED light sources are available from Invuity, Stryker, and Sunoptic.
- 6. Reduce the intensity setting of the light source to the minimum level before turning on the power. Then, turn the surgical light source on.
- 7. Increase the intensity setting of the light source to approximately 60% and assess the illumination transmitted by the PhotonGuide Adapt Illuminator. Avoid increasing the intensity setting of the light source beyond the level of illumination required to perform the procedure. If using a xenon light source, keep the light intensity below 60%.
- 8. Do not use with a xenon light source greater than 300 watts.
- Connect commercially available suction or smoke evacuation tubing to the hub on the bottom of the retractor handle. Then connect the tubing to a compatible smoke evacuation system or suction system.
 NOTE: The secondary suction hose is not provided with the Eikon LT Adapt Smoke Evacuation Retractor. Use a commercially available secondary suction hose compatible with the smoke evacuation unit.
- Position the Retractor as desired. Re-position as needed.
 Note: During the surgical procedure, debris should be removed from the surgical instruments with a sponge, wipe, or gauze and sterile water.
- 11. During the procedure, the PhotonGuide Adapt Illuminator and Fiber Optic Cable may be transferred to another Retractor.
- 12. After the procedure has been completed, turn the surgical light source off and carefully disassemble the Fiber Optic Cable and PhotonGuide Adapt Illuminator from the Retractor and remove the slat from the retractor.
- 13. Disconnect the Fiber Optic Cable from the surgical light source. When removing the Fiber Optic Cable from the light source, grasp the widest area of the black strain relief closest to the ACMI connection on the light source.
- 14. Discard the PhotonGuide Adapt Illuminator according to hospital procedures and in accordance with local, state and federal laws and regulations.

Note: For further instructions, refer to the Fiber Optic Cable and PhotonGuide Adapt Illuminator Instructions for Use.

Preparation for Cleaning

Equipment and Materials

WARNING: Use only Invuity-approved equipment unless otherwise specified. DO NOT modify any system component or accessory.

The following equipment is required to process devices per the instructions in this manual:

- Warm water (See the Water Quality section)
- Prepared, specially formulated cleaning agents (See the Cleaning Agents section)
- Absorbent wipes
- Soft, nonlinting cloth
- Syringe
- Non-abrasive, soft, flexible, synthetic bristle brushes
- Washer-disinfector (complies with ISO 15883 series)
- Cleaning agents as required by the washer-disinfector manufacturer
- Instrument air¹ < 140 kPa [< 20 psi]
- Oven

¹Instrument air is medical gas that falls under the general requirements for medical gases as defined by the National Fire Protection Association (NFPA) Health Care Facilities Code, is not respired, is compliant with the American National Standards Institute (ANSI) / International Society of Automation (ISA) Quality Standard for Instrument Air, and is filtered to 0.01 microns, free of liquids and hydrocarbon vapors, and dry to a dew point of -40 °C [-40 °F].

Cleaning and Sterilization

	1.	Before cleaning and sterilization, ensure the slat has been removed from the retractor.
	2.	Soak surgically used Retractors in an enzymatic solution or a neutral pH detergent according to
		hospital protocol. An alkaline cleaning agent (neat, up to pH 11) is allowable, but not preferred.
		Alkaline cleaning agents may cause cosmetic damage or reduce the life of the product. Pay close
		attention to instructions for correct detergent dilution, temperature, and soak time. Soak for a
		minimum of 1 minute.
	3.	Fully immerse the instrument in solution, and use a small, clean, soft, hand-held brush to remove
Monual		debris from all instrument surfaces. Ensure debris is removed from tips, crevices, and channels.
Ivialiuai		Note: Do not use a stiff bristle brush or metal type brush to clean the instruments. Avoid
		contact with other metal instruments or object which may dent, ding, scrape or otherwise
		damage the surface of the retractor.
	4.	Rinse instruments to remove residual debris and cleaning agents.
	5.	Ensure sterilization trays are properly decontaminated according to hospital procedures.
	6.	If lumens are present, use an appropriately sized lumen brush to clean the lumen, then flush the
		lumen(s) with de-ionized water

	 Before cleaning and sterilization, ensure the slat has been removed from the retractor. Perform the following pre-cleaning steps: Remove all visible soil from the equipment using the prepared cleaning agent, using brushes as necessary. Rinse the equipment to remove all excess cleaning agent. Load the equipment into the washer-disinfector in an appropriate insert tray, a wire basket or other washer-disinfector equipment holder. Avoid contact between components. 					
	Phase	se	Time	Water Temperature	Cleaning Agent	
	Pre-I	Rinse	2 to 4 minutes	< 21 °C [< 70 °F]	Prepared cleaning agent (optional)	
Cleaning: Automated	Was	h ¹	2 to 4 minutes	60 to 82 °C [140 to 180 °F]	Prepared cleaning agent	
	Rins	e	2 to 4 minutes	43 to 82 °C [110 to 180 °F]	-	
	Ther	mal Rinse ²	1 minute	90 °C [194 °F]	-	
	Dry		15 minutes	-	-	
	 ²Thermal rinse is optional. Thermal rinse does not replace sterilization. A thermal rinse phase may be used for up to 5 minutes at a water temperature of not more than 95 °C. 5. Unload the washer-disinfector and visually inspect the equipment for remaining soil or cleaning agent. If soil or cleaning agent remains, repeat the cleaning procedure using freshly prepared cleaning agent. 					
Water Quality	 WARNINGS: Use appropriate water quality for each stage of the cleaning process. Mineral residues from hard water can stain the equipment and/or prevent effective cleaning and decontamination. Use utility water for flushing, washing, and rinsing the equipment. Utility water is water that comes from the tap. Use potable water for diluting cleaning agents. Potable water is water that is treated and delivered in a matter so that it meets United Sates (US) Environmental Protection Agency (EPA) or local guidelines as suitable for drinking. Use critical water for final rinsing of the equipment prior to sterilization. Critical water is water that is extensively treated usually by a multistep treatment process that could include a carbon bed, softening, deionization, and reverse osmosis or distillation to ensure that the microorganisms and the inorganic and organic material are removed from the water. A final submicron filtration could also be part of the treatment process. CAUTION: Poor water quality can adversely affect the life of medical devices. ALWAYS follow the water quality requirements per Association for the Advancement of Medical Instrumentation (AAMI) TIR 34. Warm water with an optimum temperature range of 27 to 44 °C I80 to 110 °E1 is recommended for 					
Dipinfaction	manual clear Disinfection	processes d	o not ensure the	e margin of safety associa	a snould be warm to the tou ated with sterilization proces	ich. sses.
(Optional)	Therefore, disinfection is optional. See the Automated Cleaning section and information related to the thermal rinse phase.					
Packaging	Retractors may be loaded into dedicated instrument trays or general purpose sterilization trays.					

	Retractors (including Invuity Fiber Optic Cables) should be sterilized using steam sterilization by either gravity displacement or pre-vacuum. The following sterilization parameters have been					
	validated for instruments in a tray, wrapped or unwrapped. Note: Invuity endorses the following sterilization parameters per AAMI recommendations for wrapped instruments and container with combined weight of less than 25 lbs. [11.3 kg].					
	Steam Sterilization Parameters					
	De	scription	Gravity Displacement	Pre-Vacuum		
	Ten	nperature	132° C² (270° F)	132-138° C (270-280° F)		
	Сус	cle Time	15 minutes ¹	3-8 minutes ¹		
Sterilization	Di	ry Time	45 minutes	30 minutes		
Otomization	¹ Maximum exposure time may be extended to 18 minutes.					
	Note: Do not use Sterrad® or other sterilization processes that contain chlorine (bleach) or enzymatic solutions as these are known to cause discoloration of the surface.					
	 The final responsibility for verification of sterilization techniques lies directly with the hospital. To ensure the efficacy of hospital processing, all cycles and methods should be verified for different sterilization chambers, wrapping methods, and/or various loading configurations. International sterilization parameters are per the following standards: Australia/New Zealand per AS/NZS 4187 Netherlands per Field Standard for Loaner Instruments, Rev 03.02, April 2008 Europe and the United Kingdom per EN ISO 17664 Canada per CSA ISO 17664 					
Storage	Store instruments in a clean, dry area.					

Warranty

INVUITY warrants, for the earlier of (i) the expiration date, if applicable, or (ii) one (1) year from the date of purchase, that this medical device is free from defects in both material and workmanship when used normally for its intended purposes. Suitability of the medical device for any surgical procedure shall be determined by the user alone (in light of relevant instructions for use) and not by INVUITY, and such user shall be responsible for understanding how to use the device for its intended purpose. In the event that purchaser believes that any INVUITY medical device is defective, purchaser will promptly notify INVUITY. Any INVUITY brand named instrument delivered from INVUITY proving to be defective will, as purchaser's sole and exclusive remedy, be replaced at no charge or the amount paid refunded, at INVUITY's discretion, so long as the defect is discovered and reported to INVUITY within the earlier of thirty (30) days of discovery or one (1) year of purchase, and INVUITY determines that the product is defective and is covered by the warranty. Notwithstanding, INVUITY shall not be responsible for normal wear and tear, including minor cosmetic damage compromise or defect caused by re-sterilization. INVUITY does not repair, sharpen, recoat or otherwise refurbish any used product. The foregoing limited warranty and limited obligation of replacement is void and of no effect in the event that there is any modification made to the medical device or the medical device is used for any purpose other than its intended purpose.

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Customer must obtain a Return Material Authorization to return product to Invuity. Returned product must be thoroughly cleaned and sterilized.

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INSTRUCTIONS FOR USE Eikon LT Adapt Smoke Evacuation, Wide Retractors

REVISION HISTORY

Rev	CO #	Description	Approved Date	Effective Date	Originator
А	19-0328	Initial release	09/16/2019	09/18/2019	J Clark
А	N/A	Admin change: replaced blurry REF symbol on pg 1.	10/17/2019	10/17/2019	S. Mistry

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INSTRUCTIONS FOR USE

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PRINTING INSTRUCTIONS

Print in-house, or at an approved supplier and route to receiving.

Do not print Revision History and Printing Instructions Page (page 8 & 9)

Print in black ink on white paper.

Print subsequent pages double sided, booklet style.

Fold into booklet and staple fold along spine (folded size is 8.5in x 5.5in)