

USER'S MANUAL

COGENTIX MEDICAL

Flexible Video Cystoscope/Hysteroscope

CST-5000/5000*i*

and Slide-On® EndoSheath® Technology

NOTE: Federal (USA) law restricts this device to sale by, or on the order of, a physician or other appropriately licensed medical professional.

www.cogentixmedical.com



How to Use This Manual

This User's Manual contains the recommended procedures for preparing and using the Cogentix Medical **CST-5000/5000**/ Flexible Video Cystoscope/Hysteroscope and the **Slide-On**® **EndoSheath**® **Technology**. It is intended for physicians and other medical personnel who will come in contact with the equipment before, during, and after any patient procedures performed with them. The manual also contains pertinent information on the proper care and handling of the equipment. Please read and become familiar with this entire manual before using the equipment.

This manual contains the following information:

- Description of the endoscope and Slide-On® EndoSheath® Technology
- The endoscope's intended use
- Components and features of the endoscopes and peripheral equipment used in conjunction with the endoscope
- Complete instructions on endoscope preparation, inspection, operation, reprocessing, and storage
- Warning and Caution statements that must be observed by endoscope users to ensure patient and user safety

If you are a **first time endoscope user**, Cogentix Medical strongly recommends that you read this manual from beginning to end and become intimately familiar with the endoscope and its use.

If you are an **experienced endoscope user**, select specific chapters and/or sections that pertain to features and procedures that you are using.

Organization of this Manual

Following is a list of the chapters included in this User's Manual. Each chapter's title is listed at the top of all pages after the title page, so that you can quickly access the information you need.

Chapter 1, Symbols and Terms – This chapter defines the symbols on the endoscope and peripheral equipment. There is also a brief list of the terms that are commonly used in the manual.

Chapter 2, Important Information – The information in this chapter is a summary of critical Warning and Caution statements in the manual. This information is essential to the safe operation and reprocessing of the endoscope. Cogentix Medical strongly recommends that this chapter be read thoroughly and completely understood by all users before working with the endoscope.

Chapter 3, Endoscope and Accessories – Introduces the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope and compatible peripheral equipment. This chapter includes instrument diagrams, identifies components, and defines their functions.

How to Use This Manual

Chapter 4, Installing and Removing the Slide-On® EndoSheath® Technology – The **Slide-On® EndoSheath® Technology** is a sterile, single-use barrier placed over the endoscope's Insertion Tube before the procedure, and removed and discarded after the procedure is completed. This chapter includes the procedures for installing the sheath prior to the procedure and removing it when the procedure is over.

Chapter 5, Preparation, Inspection and Operation – This chapter describes how to prepare the endoscope and peripheral equipment for use, and how to assemble the equipment into a system. The chapter also leads you through a detailed inspection procedure to confirm that the equipment is undamaged and working properly before it is used in a procedure.

Chapter 6, Reprocessing – This chapter contains important instructions on the proper cleaning, disinfection, and sterilization of the endoscope before its first use and after each subsequent use. Strict adherence to the instructions in this chapter will render the endoscope "patient-ready" for each procedure.

Chapter 7, Care and Storage – If the endoscope will not be used for a prolonged period, refer to this chapter for instructions on safe, secure storage.

Chapter 8, Troubleshooting – Describes possible problems that may be encountered with the endoscopic system, and suggests corrective actions to take towards resolving minor problems.

Chapter 9, Warranty and Service – This chapter contains the terms of the Cogentix Medical warranty on the endoscope, any restrictions that apply and user actions that may void the warranty if taken. This chapter also includes shipping instructions in case the endoscope must be returned to Cogentix Medical for repair.

The *Appendix* contains the technical specifications for the **CST-5000/5000**/ Flexible Video Cystoscope/Hysteroscope and infection control information.

Additional Information

The information in this User's Manual is subject to change without notice. If you have any questions regarding any of the material contained in this manual, or wish to confirm that this is the most-comprehensive information available for this product, please contact your local distributor or Cogentix Medical Customer Service Department at 866 258-2182 (toll free in U.S.) or (+1) 952 426-6189 (international calls).

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1 Symbols and Terms

Symbols

The symbols listed below can be found on the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope and on other components of the endoscopic system.



Type BF applied part (Safety degree specified by IEC 60601-1)



Alerts the user to the presence of important operating, maintenance, and service instructions. Refer to the user's manuals for warnings and safety precautions associated with equipment used in the procedure.



Equipment bearing this mark has been designed, tested, and certified as essentially compliant with all applicable European Union (EU) regulations and recommendations.



Serial number of the endoscope



Up position for the Angulation Lever



Down position for the Angulation Lever



STERIS[®] and STERRAD[®] Reprocessing Compatibility
Refer to Chapter 6, Reprocessing (Endoscope must feature the

S symbol for STERRAD[®] / STERIS[®] compatibility)



Products do not contain natural rubber latex



Consult Instructions for Use



The presence of this symbol on the product or packaging indicates that the device is RoHS compliant.

Symbols and Terms

Terms

The following terms are used throughout this User's Manual:

"Cystoscope/Hysteroscope," "Videoscope" and "Endoscope" refer to the Cogentix Medical CST-5000/5000; Flexible Video Cystoscope/Hysteroscope.

"Slide-On® EndoSheath® Technology" or "Sheath" refers to the disposable CV-2.1 or CV-1.5 Slide-On® EndoSheath® Technology for the CST-5000/5000i Flexible Video Cystoscope/Hysteroscope.

"Processor" refers to the DPU-5000/7000 Series Video Processors.

2 Important Information

The information in this chapter is essential for the correct and safe operation of the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope. Please read and understand this information before preparing or using the endoscope or any peripheral equipment with which it will be used.

Intended Use

Cystoscopy:

The Cogentix Medical **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope with **Slide-On**[®] **EndoSheath**[®] **Technology** is intended to be used for endoscopic access to and examination of the lower urinary tract, including the bladder. When combined with accessory instruments, the endoscopic system allows the user to perform various diagnostic and therapeutic procedures.

Hysteroscopy:

The Cogentix Medical **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope and **Slide-On**[®] **EndoSheath**[®] **Technology** is also intended to be used for direct viewing of the cervical canal and the uterine cavity, for the purpose of performing diagnostic and therapeutic surgical procedures.

Do not use the equipment for any purpose other than these intended uses.

Contraindications for Hysteroscopy

The **CST-5000/5000***i* endoscope and **Slide-On**[®] **EndoSheath**[®] **Technology** should not be used to perform hysteroscopy if any one of the following conditions exists. **Do not perform hysteroscopy using this equipment if**:

- The patient is pregnant or suspected to be pregnant
- The patient's cervix cannot be properly dilated
- The patient's uterus cannot be distended
- The patient has acute pelvic inflammatory disease (PID)
- The patient has invasive carcinoma of the cervix
- The patient has had a recent uterine perforation

User Qualifications

This equipment should only be used in a medical facility by or under the supervision of a physician trained in cystoscopy/hysteroscopy. Use of the system does not require any deviation from standard cystoscopy or hysteroscopy technique. However, the operator should have complete familiarity with the operation of the entire system prior to clinical use.

Important Information

Only practitioners with adequate training in hysteroscopy should perform hysteroscopic procedures. Cogentix Medical strongly recommends a thorough review of all relevant medical literature relative to techniques, complications, and hazards prior to undertaking any hysteroscopic procedure.

For preparation of the endoscope before use, and disassembly and proper cleaning after use, users should be thoroughly trained in the proper procedures. Failure to completely understand these details, such as (but not limited to) **EndoSheath® Technology** installation and authorized disinfection protocols, may pose an infection control risk and/or cause equipment damage.

Only use the endoscope and peripheral equipment according to the instructions and under the operating conditions given in this User's Manual. Failure to do so could result in compromised safety, equipment malfunction and/or instrument damage.

If training assistance is desired from either the manufacturer or local distributor, please contact Cogentix Medical Customer Service at 866 258-2182 (toll free in U.S.) or (+1) 952 426-6189 (international calls).

Reprocessing

The endoscope must be thoroughly cleaned, disinfected, and/or sterilized before its first use and after each subsequent use. This is the <u>only</u> way to ensure that a "patient-ready" endoscope is used in every procedure. See Chapter 6, **Reprocessing**, for information on all reprocessing equipment and procedures.

Maintenance and Repair

The endoscope contains **no** user-serviceable parts; **never** attempt to modify or repair it. Doing so may cause further equipment damage and/or compromise patient safety if the endoscope is subsequently used in a procedure. The endoscope may only be serviced / repaired at an authorized Cogentix Medical facility.

In addition to thoroughly inspecting the endoscope before each procedure, it should be periodically inspected to determine if there is damage or wear that requires attention.

Signal Words

Information included in this manual to warn users of the possibility of patient injury and/or equipment damage is signified by the Warning and Caution symbols shown in this section. Warnings, Cautions and Notes will appear throughout this manual; carefully read and follow all statements.



Alerts the user to situations which, if not avoided, could result in death or serious injury.



Alerts the user to situations which, if not avoided, could result in moderate or minor injury to the user or patient. It is also used to alert the user to conditions and actions that could cause equipment damage.



Note: Indicates additional helpful information.

Important Warnings and Safety Precautions

The following precautions should always be exercised when using the endoscope and all medical equipment to ensure safety for all involved parties – user(s), patient(s), etc. They are summarized here in the order of the stages of the endoscope's use.

Preparation, Inspection, and Assembly



The CST-5000/5000*i* endoscope and the Slide-On[®] Endo-Sheath[®] Technology are designed to operate as an integrated system. Neither component can be used independently of the other.

The **Slide-On**[®] **EndoSheath**[®] **Technology** is designed for a single patient use only. Do not reuse or attempt to re-sterilize the Sheath. For installing the Sheath in a sterile environment, users should wear two pairs of sterile gloves. Refer to Chapter 4, **Installing and Removing the Slide-On**[®] **EndoSheath**[®] **Technology** for complete details.

Carefully inspect all equipment before using it in a procedure, and do not use any equipment that is damaged or excessively worn. Doing so could lead to patient injury and/or further damage to the equipment.

If inspection reveals difficulty in articulation of the endoscope's Distal Bending Section, the endoscope may be damaged. Do not use the endoscope if damage occurs; doing so could cause patient injury, and may result in further damage to the endoscope.

All devices that are connected to the **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope and **DPU-5000/7000 Series** Video Processor must be Classified Medical Equipment. Before using any additional equipment, confirm that it complies with the appropriate end-product safety standard (such as IEC 60950-1) and the Standards for Medical Electrical Equipment (UL 60601-1 or IEC 60601-1).

Important Information

CAUTION

Never drop the equipment or subject it to severe impact, as it could compromise the functionality and/or safety of the equipment or system. Should the equipment be mishandled or dropped, do not use it. Immediately return it to an authorized Cogentix Medical service facility for inspection and repair.

While the Sheath is being installed, the endoscope should move freely without the application of force. Should any resistance be encountered, verify that the Sheath's channel is properly aligned. If the channel of the Sheath is misaligned, straighten out the channel before continuing scope insertion, as shown in Figure 2-1 below. Also refer to Chapter 8, **Troubleshooting**, for further information. If the suggestions given in that chapter do not solve the alignment or insertion problems, contact your local distributor or Cogentix Medical Customer Service Department.



Note: The Cogentix Medical CST-5000/5000*i* Flexible Video Cystoscope/Hysteroscope and Slide-On® EndoSheath® Technology are not made with natural rubber latex.

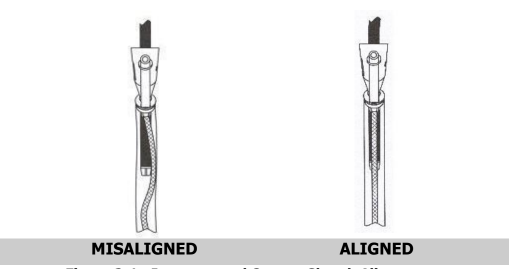


Figure 2-1: Incorrect and Correct Sheath Alignment

During Use — General



Do not use this equipment in the presence of a flammable anesthetic mixture containing air, oxygen or nitrous oxide. There is a possibility of fire or explosion. If any component of the endoscopic system malfunctions during the procedure, or if the endoscopic image is lost or compromised, immediately move the endoscope's Distal Bending Section to the neutral position and slowly withdraw the endoscope from the patient.

WARNING

Using an endoscope that is not functioning properly could cause patient injury and/or further damage to the equipment.

Do not use any accessory instrumentation that is damaged. Doing so may result in instrumentation malfunction during the procedure, which could cause serious patient injury and/or further damage to the equipment.

CAUTION

Always wear appropriate personal protective equipment when using the endoscope and/or sheath, such as a gown, gloves, and face and eye shields.

Avoid excessive bending or twisting of the endoscope's Insertion Tube and Videoscope Cable. Although they are designed to bend, excessive bending can damage the fiber bundles and internal components.

Do not apply excessive pressure to the endoscope's Angulation Lever, as it could damage the endoscope and lead to patient injury.

Do not look directly at the intense light emitted from the endoscope tip to avoid the possibility of eye injury.

If resistance is encountered when inserting an instrument into the Sheath, do not force it, as it could damage the instrument, the endoscope and/or the CV-2.1 Slide-On® EndoSheath® Technology.

The Irrigation/Working Channel of the **CV-2.1 Slide-On® EndoSheath® Technology** accommodates instrumentation indicated by the accessory manufacturer to be compatible with a 2.1 mm or smaller working channel. All instrumentation must be tested for compatibility with the channel prior to clinical use. If assistance is needed to determine compatibility, contact your local distributor or Cogentix Medical Customer Service Department.

Do not attempt to advance an exposed needle or other sharp instruments through the working channel. Doing so could damage the sheath's channel.

A thorough understanding of the principles and techniques involved in laser, electrosurgical, and ultrasonic procedures is essential to avoid shock and burn hazards to both patient and medical personnel, and to avoid damage to the device and other medical instruments. Ensure that insulation or grounding is not compromised.

During Use — Hysteroscopy



Prior to performing a hysteroscopic procedure, it is extremely important that the user review the following information which may be critical to ensuring patient safety.

Continuous CO₂ Flow Hysteroscopy

If CO₂ gas is used as a distention medium, operative hysteroscopy is contraindicated due to the risk of gas embolization, with the exception of non-electrosurgical polypectomy or directed biopsy. CO₂ gas may be used for diagnostic procedures. It is extremely important that a hysteroscopic insufflator is used. Death has been reported when laparoscope CO₂ insufflators were used during hysteroscopy. The CO₂ flow rate should be limited to less than 100 ml/min, and the patient's intrauterine pressure should not exceed 100 mm Hg.

Potential complications of continuous flow hysteroscopy with CO₂ include:

- CO₂ embolization
- Circulatory collapse
- Death

Continuous Fluid Flow Hysteroscopy

If a liquid distention medium is used, strict fluid intake and output surveillance should be maintained to ensure that fluid deficit is known at all times. Depending on whether a non-electrolytic or an electrolytic solution is being used, when excessive fluid deficit occurs, consideration should be given to stopping further infusion and concluding the procedure.

When using a fluid distention medium, closely monitor fluid intake/ output. Excessive intake of distention fluid can lead to fluid overload and complications.

Potential complications of continuous flow hysteroscopy with fluid distention media include:

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

If pregnancy is suspected, perform a pregnancy test prior to performing hysteroscopy. **DO NOT** perform hysteroscopy if the patient is pregnant.

If performing Endometrial Ablation or Hysteroscopic Myomectomy, verify that the patient does not have procedural contraindications.

Vaginal ultrasonography before hysteroscopy may identify clinical conditions that will alter patient management.

> Gravity fed intrauterine fluid distention can usually be accomplished with pressures in the range of 35-75 mm Hg. Hanging the fluid distention medium 42 inches above the patient can generate intrauterine pressure of approximately 80 mm Hq. Unless the systemic blood pressure is excessive, it is seldom necessary to use pressures greater than 75-80 mm Hg.



Note: Hysteroscopes are used as tools to access the uterine cavity and are not, in and of themselves, a method of surgery.

Diagnostic hysteroscopy can be used to evaluate numerous conditions including the following:

- Abnormal uterine bleeding
- Infertility
- Evaluation of abnormal hysterosalpingogram or sonohysterogram
- Intrauterine foreign body
- Amenorrhea
- Pelvic pain

Operative hysteroscopy is used to perform numerous procedures including the following:

- Directed endometrial biopsy
- Polypectomy
- Submucous Myomectomy
- Transection of Intrauterine Adhesions
- Transection of Intrauterine Septa
- **Endometrial Ablation**

Reprocessing



The endoscope must be properly reprocessed, by cleaning, disinfecting and/or sterilizing, before its first use and after each subsequent use. Using an endoscope in a procedure that has not been properly reprocessed presents an acute infection-control risk to both the patient and medical personnel performing or assisting in the procedure.



Always wear appropriate personal protective equipment when reprocessing the endoscope and/or sheath, such as a gown, gloves, and face and eye shields.

Use extreme care when reprocessing the endoscope. Do not forcefully pull, push, or drag wipes, towels, or cloths along the Insertion Tube. The use of excessive force could damage the endoscope.

Important Information

Do Not immerse the endoscope in disinfectant solution for long periods of time (>1 hour). Prolonged immersions may damage the outer coverings of the endoscope and allow fluid infiltration.

Do Not place the endoscope in or near contaminated areas after it has been reprocessed. Doing so can re-contaminate the endoscope and require reprocessing to be repeated.

Do Not place the endoscope in awkward or confining areas between procedures as this could result in equipment damage.

3 Endoscope and Accessories

The Cogentix Medical **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope and **Slide-On**[®] **EndoSheath**[®] **Technology** are designed to perform safe, sterile, and efficient cystoscopic / hysteroscopic procedures.

The two major components of the endoscopic system are:

- The CST-5000/5000i Flexible Video Cystoscope/Hysteroscope, which is shown in Figure 3-1 and Figure 3-2 on pages 13 and 14. The endoscope's Insertion Tube has no working channel; it contains the video camera module and illumination bundles.
- The disposable **Slide-On**[®] **EndoSheath**[®] **Technology**. The sterile, disposable Sheaths are installed over the endoscope's Insertion Tube, acting as a protective barrier to protect patients and users from the spread of potentially pathogenic materials. The Sheaths also contain working channels through which irrigation can be performed and/or accessory equipment can be used. There are two types of Sheaths available for use with the **CST-5000/5000***i*:
 - □ The **CV-2.1 Slide-On**[®] **EndoSheath**[®] **Technology** (Figure 3-3 on page 16). This Sheath contains a 2.1mm (6.3Fr) channel through which irrigation fluids can be instilled into the patient, or accessory instruments can be inserted.
 - □ The **CV-1.5 Slide-On**[®] **EndoSheath**[®] **Technology** (Figure 3-4 on page 18) has a 1.5mm (4.5Fr) channel that supports irrigation, but cannot accommodate the insertion of accessory devices.

Inspect the Standard Set



Do not use any equipment that is observed to be damaged or excessively worn. Doing so could lead to patient injury and/or further damage to the equipment.

When the endoscope is received from Cogentix Medical, immediately confirm that all of the applicable items listed in Table 3-1 have been shipped, and inspect them for damage. If any item is missing or damaged, do not use the endoscope. Contact Cogentix Medical to obtain a replacement part.

CST-5000/5000i Cystoscope/Hysteroscope Standard Set					
COGENTIX MEDICAL CATALOG NO.	DESCRIPTION				
08-5201	CST-5000 Flexible Video Cystoscope/Hysteroscope, NTSC				
08-5202	CST-5000 Flexible Video Cystoscope/Hysteroscope, PAL				
08-5203	CST-5000/ Flexible Video Cystoscope/Hysteroscope, NTSC				
08-5204	CST-5000/ Flexible Video Cystoscope/Hysteroscope, PAL				
07-6180	Cogentix Medical CST-5000/5000/ Carrying Case				
07-6015	Vent Cap				
CST-5000/5000i User's Manual (this document)					
VIDEO PR	VIDEO PROCESSORS (NOT SHIPPED WITH ENDOSCOPE)				
07-5050	DPU-5050 Video Processor with LCD Display *				
07-5051	DPU-5050A Video Processor with LCD Display *				
07-7001	DPU-7000A Video Processor with LCD Display				
OPTIONAL ITEMS	OPTIONAL ITEMS AND ACCESSORIES (NOT SHIPPED WITH ENDOSCOPE)				
07-6160	Installation Stand, Floor Model				
07-6161	Installation Stand, Clamp Model				
07-6162	Installation Stand, Wall Mount				
07-6010	Videoscope Leak Tester				
ENDOSHEATH® TECHNOLOGY					
08-5101	CV-2.1 Slide-On [®] EndoSheath [®] Technology with 2.1mm (6.3 Fr) Working Channel for the CST-5000/ 5000 <i>i</i> Flexible Video Cystoscope/Hysteroscope				
08-5102	CV-1.5 Slide-On® EndoSheath® Technology with 1.5mm (4.5 Fr) Irrigation Channel for the CST-5000/5000i Flexible Video Cystoscope/Hysteroscope				

Table 3-1: CST-5000/5000i Flexible Video Cystoscope/Hysteroscope – Components

^{*}Available in select markets only

Equipment Diagrams

CST-5000/5000/ Flexible Video Cystoscope/Hysteroscope

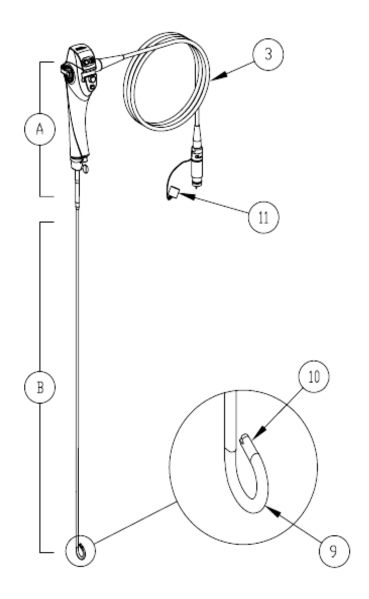


Figure 3-1: CST-5000/5000i Flexible Video Cystoscope/Hysteroscope

A: Control Body

B: Insertion Tube

CST-5000/5000/ Flexible Video Cystoscope/Hysteroscope

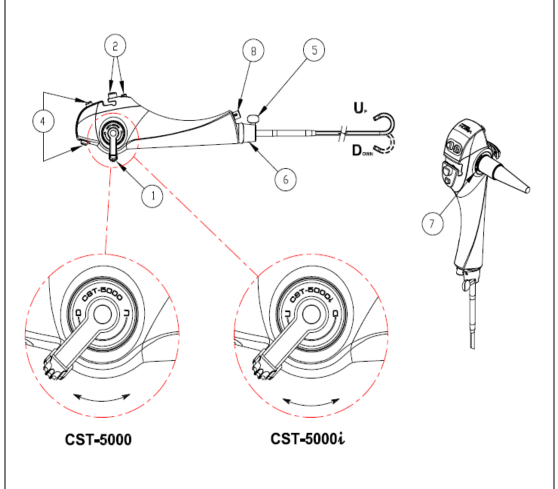


Figure 3-2: CST-5000/5000i Flexible Video Cystoscope/Hysteroscope -Control Body

Instrument Components

1. **Angulation Lever:** Located on the Control Body, this component Controls Distal Bending Section deflection. There is only one difference between the CST-5000 and 5000/ Models: the Up and Down conventions are reversed in the Angulation Lever and its effect on articulation/deflection in the Distal Bending Section.

CST-5000 Model:

The Angulation Lever on the **CST-5000** provides control of the bending section in the "traditional" directions. This indicates that when the Angulation Lever is moved toward the "top" of the Control Body, the Distal Bending Section will articulate in the visual "down" direction. When the Angulation Lever is moved toward the Insertion Tube, the Distal Bending Section will articulate toward the visual "up" direction.

CST-5000/ Model:

The Angulation Lever on the **CST-5000**/ provides control of the bending section in the "intuitive" directions. This indicates that when the Angulation Lever is moved toward the "top" of the Control Body, the Distal Bending Section will articulate in the visual "up" direction. When the Angulation Lever is moved toward the Insertion Tube, the Distal Bending Section will articulate in the visual "down" direction.

- 2. **Flow Control Valve:** Secures the EndoSheath[®] Irrigation Tubing for button-actuated irrigation during use with Open Flow and Close Flow buttons.
- 3. **Videoscope Cable:** The connector at the end of the cable connects to the **DPU-5000/7000 Series** Video Processor.
- 4. Control Buttons: Four programmable function buttons which allow the user to activate different functions of the video system. Consult the DPU-5000/7000 Series Video Processor User's Manual for instructions regarding the control functions and how to program the buttons.
- 5. **Locking Knob:** Mates with the EndoSheath[®] Rigid Connector to secure the **Slide-On**[®] **EndoSheath**[®] **Technology** to the endoscope.
- 6. **EndoSheath**[®] **Interface:** Secures the disposable EndoSheath[®] cover to endoscope body.
- 7. **Identification Ring:** Includes the Serial Number, which is a unique number identifying the endoscope, and the \bigcirc S symbol, which indicates the endoscope can be sterilized using a validated STERRAD® / STERIS® system. The endoscope must feature this symbol on the Identification Ring in order for STERRAD® / STERIS® compatibility to apply.
- 8. **Vent Valve:** When the Vent Cap is connected, this valve allows access to the interior of the endoscope for EtO and STERRAD[®] gas sterilization, and should be connected during transport. The Vent Cap **must** be attached to the valve prior to EtO and STERRAD[®] gas sterilization and prior to shipping. The valve is also used as a Leak Tester Connector for Leak Testing.
- 9. **Distal Bending Section:** Deflects up and down when the Angulation Lever is actuated.
- 10. **Distal Tip:** The terminating point of the video camera module and the light guide fiber bundles [Light Guides].
- 11. **Sealing Cap:** Seals the plug prior to soaking for leak testing or disinfection.
- **A) Control Body:** This component provides physician control over endoscopic functions.
- **B) Insertion Tube:** This segment, along with the Distal Tip, is the part of the endoscope that is inserted into the patient.

CV-2.1 Slide-On® EndoSheath® Technology

The **CV-2.1 Slide-On® EndoSheath® Technology** shown in Figure 3-3 is a sterile, disposable barrier for the Cogentix Medical **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope. The Sheath isolates the endoscope from contact with patient fluid and material during the procedure.

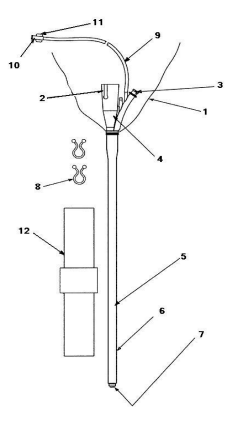


Figure 3-3: CV-2.1 Slide-On® EndoSheath® Technology

- 1. **Control Body Cover:** Covers and protects the endoscope's Control Body from contaminants during the procedure.
- 2. **Slot for Locking Knob:** Slides past the Locking Knob to secure the Sheath to the endoscope.
- 3. **Accessory Port with Removable Seal:** Allows rapid fluid injection and passage of accessory instruments during procedures. A removable Accessory Port Valve Seal allows for secure Luer-Lock syringe attachment for fluid withdrawal.
- 4. **EndoSheath**[®] **Connector:** Secures the Sheath to the endoscope.
- 5. **Working Channel:** A solid, impermeable tubing that runs the length of the Sheath, allowing for maximum fluid irrigation volume and evacuation while also allowing the passage of accessory devices.

Endoscope and Accessories

- 6. **Insertion Tube Barrier:** A flexible elastomeric sleeve that creates a barrier covering for the endoscope's Insertion Tube.
- 7. **Sheath Window:** A patented, optically clear window that covers the Distal Tip of the endoscope.
- 8. **Control Body Cover Clips (2):** Secures the Control Body Cover to the Video Cable and Irrigation Tube.
- 9. **Irrigation Tube:** Provides access for irrigation or evacuation suction to the distal tip during a procedure.
- 10. **Irrigation Tube Connector:** A standard Luer-Lock fitting to allow for coupling to the Irrigation/Suction tube.
- 11. **Luer Cap:** A standard Luer-Lock fitting cap to prevent internal contamination of the Irrigation tube before making a connection.
- 12. **Drape Bag:** Covers the Installation Stand to prevent contamination. The Drape Bag should also be used to conveniently dispose of the Sheath.

CV-1.5 Slide-On® EndoSheath® Technology

The **CV-1.5 Slide-On**[®] **EndoSheath**[®] **Technology** shown in Figure 3-4 is a sterile, disposable barrier for the Cogentix Medical **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope. The Sheath isolates the endoscope from contact with patient fluid and material during the procedure, and supports irrigation, but does not support accessory instrument passage.

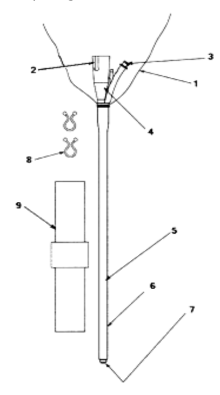


Figure 3-4: CV-1.5 Slide-On® EndoSheath® Technology

- 1. **Control Body Cover:** Covers and protects the endoscope's Control Body from contaminants during the procedure.
- 2. **Slot for Locking Knob:** Slides past the Locking Knob to secure the Sheath to the endoscope.
- 3. **Irrigation Port:** Luer-Lock syringe attachment allows for fluid irrigation and withdrawal.
- 4. **EndoSheath® Connector:** Secures the Sheath to the endoscope.
- 5. **Channel:** A tube that runs the length of the Sheath and allows for fluid irrigation and evacuation.
- 6. **Insertion Tube Barrier:** A flexible, elastomeric sleeve that creates a barrier covering for the endoscope's Insertion Tube.

- 7. **Sheath Window:** A patented, optically clear window that covers the Distal Tip of the endoscope.
- 8. **Control Body Cover Clips (2):** Secures the Control Body Cover to the Video Cable.
- 9. **Drape Bag:** Covers the Installation Stand to prevent contamination. The Drape Bag can also be used to conveniently dispose of the Sheath.

DPU-5000/7000 Series Digital Video Processor



The **CST-5000/5000** Flexible Video Cystoscope/ Hysteroscope is not compatible with any other manufacturers' video processors. Attempting to connect the endoscope to or use it in conjunction with another manufacturer's video processor could cause damage to the endoscope and/or the video processor.

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope **must** be used in conjunction with a **DPU-5000/7000 Series** Video Processor (shown in Figure 3-5 below). Refer to the **DPU-5000/7000 Series** User's Manual for complete instructions on the operation of the unit.

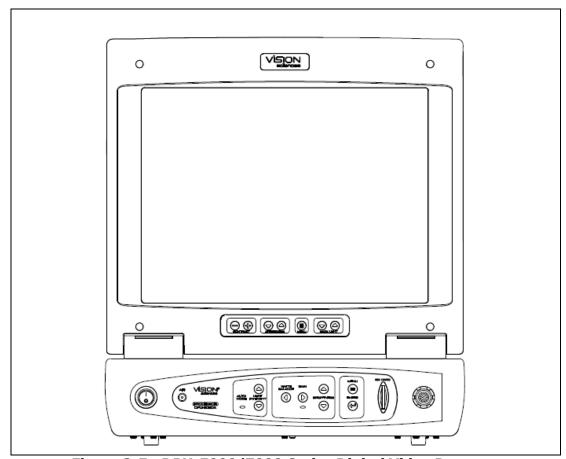


Figure 3-5: DPU-5000/7000 Series Digital Video Processor

Endoscope and Accessories

Installation Stand



Installation and removal of the **Slide-On® EndoSheath® Technology** onto and from the Cystoscope/Hysteroscope should always be performed using the Installation Stand. Failure to do so may cause difficulty in installation and/or equipment damage.

Refer to Figure 3-6 below and to the Installation Stand's Instructions For Use for information on the correct preparation and use of the stand.

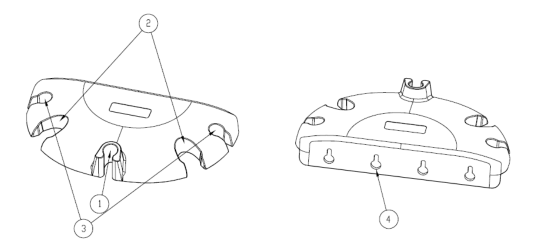


Figure 3-6: Installation Stand

- 1. **EndoSheath**® **Slot/Holder:** The slot that securely holds the EndoSheath® Connector, which is designed to fit with its Accessory Port facing out.
- 2. **Endoscope Storage Slots:** Post-procedure slots for hanging endoscopes. (Not intended for long-term storage unless the Installation Stand is wall-mounted.)
- 3. **Cable Storage Slots:** For securing the Videoscope Cable when the endoscope is not in use or being stored.
- 4. **Wall-Mounting Bracket/Holes:** For mounting on a wall or inside a cabinet for storage purposes.

Accessories



Do not use any accessories that are not in compliance with the equivalent safety requirements of this equipment. Doing so may reduce the operational safety of the system and could cause patient and/or user injury. For all accessories, confirm that safety certifications have been performed in accordance with the appropriate standard (IEC 60601-1 and/or IEC 60601-1-1).

The use of accessories not specified in this manual or sold by Cogentix Medical may result in increased electromagnetic emissions or decreased immunity of the equipment or system.

Video Processor

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope is designed to work with the Cogentix Medical **DPU-5000/7000 Series** Video Processors. The endoscope is not compatible with any other manufacturers' video processors.

Light Sources

The **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope has an integrated, solid-state light source which is controlled by the **DPU-5000/7000 Series** Video Processors. No external light source is required for the **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope.

Leak Testing

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope may only be leak tested with a Cogentix Medical V1 Endoscope Leak Tester.

Reprocessing

The **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope may be reprocessed by a variety of methods. Refer to Chapter 6, **Reprocessing** for the accessories that will be used when reprocessing the endoscope. Contact Cogentix Medical Customer Service for advice on compatibility issues.

Therapeutic Accessories

The **CV-2.1 Slide-On**[®] **EndoSheath**[®] **Technology** features a 2.1mm Working Channel for the passage of therapeutic tools. Cystoscopy/Hysteroscopy tools and accessories that are compatible with a 2.1mm channel should be compatible with this system.



The **CV-1.5 Slide-On**[®] **EndoSheath**[®] **Technology** does not support accessory tool passage. Attempting to insert an accessory or tool could damage this Sheath.

For further information regarding tools and accessories, please contact your local distributor or Cogentix Medical Customer Service Department.

CAUTION

Installation and removal of the Slide-On® EndoSheath® Technology onto and from the Cystoscope/Hysteroscope should always be performed with the Installation Stand. Failure to utilize the installation stand for setup and preparation may lead to difficulty in installation and ultimately equipment damage.

Ensure that the endoscope is clean and dry prior to installing the **Slide-On® EndoSheath® Technology**. Any moisture on the endoscope prior to installation may result in difficulty removing the endoscope from the **Slide-On® EndoSheath® Technology** after the procedure.

Wear appropriate protective gear when using the **Slide-On® EndoSheath® Technology** and Cystoscope/Hysteroscope, including gown, gloves, and face and eye shields. To maintain a sterile field during installation, it is recommended that users wear two (2) pairs of sterile gloves.

The Endoscope and Sheath do not have any user-serviceable parts. Do not attempt any repairs. If malfunction occurs, refer to the Troubleshooting section of this manual or call Cogentix Medical for assistance.

Install the Slide-On® EndoSheath® Technology



Before installing the Sheath, carefully inspect the endoscope's Insertion Tube for any damage or defects. If any irregularities are found, do not use the endoscope. Using a damaged or defective endoscope could damage the Sheath, cause further damage to the endoscope itself, and/or cause patient or user injury.

The **Slide-On**[®] **EndoSheath**[®] **Technology** is supplied sterile, and is intended for a single use only. Do not reuse or attempt to re-sterilize the Sheath, as it could become damaged, which could in turn cause damage to the endoscope and/or present an infection-control risk to the patient and/or user.



Do not expose the interior or exterior of the Sheath to alcohol or other cleaning agents prior to use.



Note: Refer to **Instructions For Use** for **CV-2.1** and **CV-1.5** Sheaths.



Do not use this equipment in the presence of a flammable anesthetic mixture containing air, oxygen or nitrous oxide. There is a possibility of fire or explosion.

Prepare the Endoscope and Sheath

- 1. Before installing the Sheath, clean the endoscope's lens on the Distal Tip with an alcohol prep pad. Do not use abrasive materials to clean any part of the endoscope, particularly the lens at the Distal Tip. Doing so could damage the lens and impair the endoscope's imaging capability.
- 2. Connect the **CST-5000/5000***i* video cable to the **DPU-5000/7000 Series** video processor, and turn the processor on.
- 3. Check the **EndoSheath**[®] disposable packaging for defects or damage.
- 4. Open the Sheath's package carefully; put on two pairs of sterile gloves.
- 5. Remove the Drape Bag from the tray, open, remove the adhesive strip backing and press the adhesive area to the back of the Installation Stand. Secure the Drape Bag with a plastic clip if necessary.
- 6. Carefully remove the Sheath from the package.
- 7. Place the Sheath into the Installation Stand with the Accessory Port facing outward (toward you). NOTE: The Sheath fits in the Stand in only one direction.
- 8. Discard the disposable packaging.
- 9. Fold back the Control Body Cover to expose the top opening of the $\mathsf{EndoSheath}^{\$}\mathsf{Connector}.$

Insert the Endoscope Into the Sheath

- 10. Take the endoscope in hand and verify that the articulation section (Distal Bending Section) of the endoscope is straight.
- 11. Hold the endoscope vertically above the Sheath. Align the endoscope's D-shaped Insertion Tube with the D-shaped opening of the EndoSheath® Connector the flat portion of the Insertion Tube should face toward the Sheath's Accessory Port (toward you).
- 12. Gently slide the endoscope's Insertion Tube into the Sheath. Keep the Insertion Tube as straight as possible. Do not rotate the Insertion Tube during insertion (this will cause resistance).

CAUTION

When inserting the endoscope into the Sheath, avoid excessive bending of the Insertion Tube, as it could damage the Sheath and/or the endoscope.

Do not apply excessive force to install the Sheath onto the endoscope if insertion is found to be difficult. If you experience difficulty installing the Sheath, refer to Chapter 8, **Trouble-shooting**. If the recommended actions given there do not ease endoscope insertion, call your local distributor or Cogentix Medical Customer Service at 866 258-2182 (toll free in U.S.) or (+1) 952 426-6189 (international calls) for further instruction.



Note: It is extremely important to maintain the alignment between the flat portion of the Insertion Tube and the Biopsy Port on the EndoSheath[®] Connector during installation and removal. If there is resistance in loading the endoscope into the Sheath, check this alignment, and straighten the channel before continuing endoscope insertion.

Figure 4-1: Incorrect and Correct Sheath Alignment



13. Make sure the Locking Knob on the endoscope is in a vertical position.

ALIGNED

MISALIGNED

- 14. Align the endoscope's Locking Knob with the vertical slot on the EndoSheath® Connector, and continue sliding the endoscope into the Sheath until the Locking Knob is fully seated at the base of the slot.
- 15. Rotate the Locking Knob to the horizontal position (locked position). This will ensure a secure fit between the EndoSheath® Connector and the endoscope.
- 16. Using the angulation lever, articulate the Bending Section in both directions to ensure proper sheath fit.
- 17. Visually inspect the Sheath and confirm that the endoscope's Distal Tip is flush with the Sheath's window. If the Distal Tip is not flush with the Sheath's

window, articulate the Distal Bending Section until the window is seated properly.

Connect Irrigation Tubing / Complete Sheath Attachment

- 18. When using the **CV-2.1 Slide-On**[®] **EndoSheath**[®] **Technology**: Connect the Irrigation Tubing to endoscope's Flow Control Valve by pressing the Open Flow button and stretching the tubing into the slot of the valve. This allows the Flow Control Valve to secure the tubing in place. Do not pull the tubing too tightly. Ensure that the tubing is placed into slot **completely** before pressing the Close Flow button.
- 19. When using the **CV-2.1 Slide-On**[®] **EndoSheath**[®] **Technology**: Remove the Sterile Cap from the Irrigation Tubing.
- 20. Connect a fluid management accessory, such as saline line, to the Sheath's Irrigation Tube Luer fitting.
- 21. Connect the Irrigation Luer Lock to the Irrigation source.



Note: The Flow Control Valve may be bypassed and a stopcock used for on-off fluid management.

- 22. Remove outer pair of gloves.
- 23. Pull the Sheath's Control Body Cover up over the endoscope's Control Body and the Videoscope Cable to completely cover them.
- 24. Secure the Control Body Cover and Irrigation Tubing to the Videoscope Cable using the Control Body Cover Clips.



Note: Fluid withdrawal and irrigation may also be accomplished by connecting to the standard luer fitting on the Sheath's Accessory Port.

25. Turn the Flow Control Valve on and off and confirm that it operates properly.

Observe the Endoscopic Image

The Sheath's window should now be in direct contact with the Insertion Tube's Distal Tip. To confirm this, view an endoscopic image with illumination on; there should be no glare.

If a gap is observed between the Distal Tip and the Sheath's window, or if glare is observed in the endoscopic image, move the Angulation Lever up and down several times to articulate the Distal Bending Section. This process should properly seat the Sheath. In some cases, it may be necessary to hold the edge of the optical window (while wearing gloves) and move it to fully seat the Sheath.

Remove the Slide-On® EndoSheath® Technology

When the endoscopic procedure is complete, the **Slide-On**[®] **EndoSheath**[®] **Technology** must be removed and disposed of properly to eliminate the possibility of infection-control risks.



Wear appropriate personal protective equipment when removing the Sheath from the endoscope, to prevent the possibility of infection from contact with patient material. Cogentix Medical strongly recommends that a double set of gloves be worn for this procedure.

Always use the Installation Stand to remove the endoscope from the Sheath. Attempting to remove the endoscope without using the Installation Stand could cause damage to the endoscope.

- 1. Put on two pairs of sterile gloves.
- 2. Ensure that the Drape Bag is completely covering the installation stand.
- 3. Slide the sheathed endoscope into the draped Installation Stand. The sheathed Insertion Tube must be contained within the Drape Bag to capture and contain any patient material and/or fluids. The Accessory Port must be facing toward you.
- 4. Disconnect Irrigation tubing from source.
- 5. Remove the Control Body Cover Clips and discard them in the Drape Bag. Keeping your hands on the outside of the contaminated Control Body Cover, pull the Cover forward and down off of the endoscope.
- 6. Remove the outer set of gloves. **Do not** handle the endoscope's Control Body with contaminated gloves.
- 7. Disconnect the Irrigation Tubing from the Flow Control Valve by pressing the Open Flow button, and then stretch the tubing while pulling it away from the base of the slot towards the open slot of the valve. Continue pulling on the tubing until it is clear of the valve.
- 8. Rotate the Locking Knob into the vertical position so that it is aligned with the slot on the EndoSheath[®] Connector.
- 9. Using the Angulation Lever, articulate the Distal Bending Section of the endoscope into the straight/neutral position.
- 10. Hold the endoscope's Control Body in one hand. Using the bottom outside portion of the Drape Bag as a barrier between your fingers and the Sheath, gently grasp the Sheath's Window, and then <u>gently</u> and <u>slowly</u> withdraw the endoscope from the Sheath.



Do not twist the endoscope when removing it from the Sheath. Doing so can damage the Insertion Tube.



If you experience difficulty removing the endoscope from the Sheath, **do not** use excessive force in trying to remove it. Refer to Chapter 8, **Troubleshooting**, for further instructions.

- 11. Place the endoscope in a non-contaminated area. Power off the video processor.
- 12. Inspect the Insertion Tube and Distal Tip, and confirm that these areas are dry. If moisture is observed, there may have been a leak into the Sheath during the procedure, provided that the endoscope was dry when the Sheath was attached. In this case, the endoscope must be high-level disinfected or sterilized following the instructions given in Chapter 6, **Reprocessing**.
- 13. Fold the top end of the contaminated Drape Bag over the EndoSheath[®] Connector and remove it from the Installation Stand. Carefully discard the contaminated Sheath per hospital policy.
- 14. The endoscope should then undergo the recommended cleaning procedure detailed in Chapter 6, **Reprocessing**.



If an abnormality is detected during endoscope preparation, do not use the equipment; refer to the tables in Chapter 8, **Troubleshooting**. If the problem cannot be solved using the information in that chapter, contact your regional distributor or Cogentix Medical Customer Service.

When using the **Slide-On**[®] **EndoSheath**[®] **Technology** with the endoscope, refer to the instructions for use that are shipped with the system. These instructions will provide complete details on preparing, installing and removing the disposable sheath.

During the procedure, the temperature at the distal end of the endoscope may exceed 41°C (106°F) due to the intense endoscope illumination. Surface temperatures over 41°C (106°F) may cause mucosal burns. Always use the minimum level of illumination necessary for adequate viewing. Whenever possible, avoid close stationary viewing and do not leave the distal end of the endoscope in close proximity to mucous membranes for a long time.

The **Slide-On**[®] **EndoSheath**[®] **Technology** is shipped sterile and intended for a single use only; do not reuse it. When the procedure is complete, remove the Sheath from the endoscope and dispose of it. Reusing the Sheath can damage it, and in turn cause endoscope damage. In addition, a reused Sheath presents a marked infection-control risk to the next patient.

A complete review and understanding of the **DPU-5000/7000 Series** User's Manual is recommended before using the **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope.

Preparation and Inspection

Follow the inspection steps listed below before connecting any equipment or using the system. Do not use the equipment if abnormalities are detected:

Select an Installation Site

It is important to select an appropriate location in which to install the Video Processor.

- Place the Video Processor on a stable rigid surface such as a cart, counter-top, or solid stand.
- The location must not contain explosive or flammable gases.

- Place the **CST-5000/5000***i* Videoscope and Video Processor away from radios, televisions, cell phones, or any other devices that emit electromagnetic energy. These can interfere with proper operation. Avoid stacking the Videoscope or the Video Processor on other equipment to avoid possible electromagnetic interference.
- Place the Video Processor in a dry place, and avoid contact with liquids.
- Do not allow the Video Processor's vents to be obstructed; full ventilation is necessary for proper operation. Vents are located on the bottom and back of the unit.
- 1. Check the Insertion Tube for holes, superficial cuts, or abrasions.
- 2. Lightly run your fingertips over the entire length of the Insertion Tube to confirm that it is smooth and does not exhibit looseness or bagging.



Do not apply excessive pressure to the endoscope's Insertion Tube. Doing so can damage the internal components of the insertion tube.

- 3. Check the Distal end of the Sheath's working channel for any sharp edges, being careful to minimize contact with the clear window.
- 4. Check for full Distal Tip deflection by actuating the Angulation Lever up and down.

Avoid applying excessive pressure when using the Angulation Lever. Doing so can damage the angulation mechanism.

5. Clean the Lens on the Distal Tip with an alcohol prep pad.



Do not use abrasive materials to clean the Lens. Doing so could damage the Lens and impair the endoscope's imaging capability.

6. Insert the endoscope's Videoscope Cable Connector into the connector on the front panel of the DPU-5000/7000 Series Video Processor and power on the Video Processor when ready for the procedure.



Turn off the Video Processor's power switch before connecting or disconnecting the Videoscope Cable. Connecting or disconnecting the cable with the power on could damage both the endoscope and the Video Processor.

7. Connect an irrigation source to the Sheath's Irrigation Tube.



Note: When using the **CV-2.1 Sheath** with irrigation tubing, connecting the irrigation source to the Sheath's Irrigation Tube will allow for irrigation while an accessory device is either inserted into or attached to the Accessory Port.

Fluid withdrawal and irrigation may be accomplished by connecting to the standard luer-lock fitting on the Accessory Port by removing the Accessory Port Seal when no accessory device is used.



Note: The Flow Control Valve may be bypassed, and a stopcock used on the Accessory Port, for on-off fluid management.

Endoscope Operation



Avoid excessive bending or twisting of the endoscope's Insertion Tube, particularly at the distal end. While the tube is designed to bend, excessive pressure can damage the fiber bundles and internal components.

Excessive angulation or excessive pressure placed on the Angulation Lever may cause equipment damage. Do not exert force to move the lever beyond its natural limits.

- 1. Hold the endoscope so that the Control Body fits comfortably in your hand, allowing easy manipulation of the Angulation Lever. The other hand is free to manipulate the Insertion Tube.
- 2. The **DPU-5000/7000 Series** Video Processor should be on; adjust the settings to the desired level using the Processor's controls.
- 3. Perform the White Balance function.
- 4. Prepare the patient using normally acceptable clinical practice prior to endoscope insertion.
- 5. Lubricate the outside of the Sheath before inserting the endoscope into the patient. Cogentix Medical recommends that the Sheath be lubricated with water or a water-based lubricant just prior to insertion.
- 6. Introduce the sheathed endoscope into the patient using normally acceptable clinical practice. Operate the Angulation Lever as necessary for advancement and observation.



Note: The four (4) programmable buttons on the endoscope's Control Body may be programmed on the Video Processor to perform designated image-control functions. These functions include:

- Image Freeze
- Image Capture
- □ Image Enhancement
- □ Remote Activation (Copy/Print)
- Gain Control

Refer to the **DPU-5000/7000 Series** Video Processor User's Manual for further information on available control functions and instructions on programming the Control Buttons.

7. When the procedure is completed, withdraw the endoscope under direct visualization without holding the Angulation Lever. This will allow the Distal Bending Section to move freely during withdrawal.

Aspirating Fluids

- 1. Attach a syringe to the Sheath's Accessory Port. When using the **CV-2.1 Slide-On® EndoSheath® Technology**, first remove the Accessory Port Seal.
- 2. Aspirate fluids by withdrawing the syringe's plunger.
- 3. To aspirate fluids via the **CV-2.1 Sheath's** irrigation tubing, confirm that the Accessory Port Seal is in place, and then open the endoscope's Flow Control Valve. Press the Close Flow button to stop aspiration.

Instilling Fluids

To inject anesthetic, saline or other fluids, attach a syringe to the Sheath's Accessory Port. When using the **CV-2.1 Slide-On**[®] **EndoSheath**[®] **Technology**, the tip of the syringe may be inserted directly into the Accessory Port Seal's opening.

If necessary, remove the Accessory Port Seal before withdrawing fluid back into the syringe.

The irrigation tubing may also be used to instill fluids. Open the Flow Control Valve to enable instillation, and close the Valve to stop it.



Make sure that the syringe is seated properly and straight in the Sheath's Accessory Port. Otherwise, the seal may be compromised and fluid may not flow smoothly into the Working Channel.

Inserting Accessories



Accessories may only be inserted when using the CV-2.1 Slide-On® EndoSheath® Technology. The CV-1.5 Slide-On® EndoSheath® Technology does not support accessory instrument passage.

Before inserting an accessory into the Sheath, confirm that it is compatible with a 2.1 mm diameter Working Channel. If the accessory is too large, it will damage the Sheath and may compromise the integrity of the barrier.

Do not continue advancing an accessory if excessive resistance is encountered, and do not use excessive force to try and advance it. The use of excessive force may result in damage to the Sheath and/or the endoscope.

- 1. Before inserting an accessory, straighten the endoscope's Distal Bending Section.
- 2. Insert the tip of the accessory through the Sheath's Accessory Port. Using straight, steady strokes, pass the accessory through the Working Channel until the tip of the accessory is visible on the video monitor.
- 3. If resistance is encountered while inserting the accessory, withdraw the accessory, straighten the endoscope's Distal Bending Section, and attempt to insert the accessory again. If resistance is still felt, confirm again that the accessory's diameter is compatible with the size of the Working Channel. Refer to Chapter 8, **Troubleshooting** for additional suggestions.

Electrosurgical Devices/Accessories

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope and **Slide-On**[®] **EndoSheath**[®] **Technology** may be used with high-frequency (HF) electrosurgical devices. Operators utilizing HF devices and accessories should follow all manufacturer and facility guidelines for proper and safe use. Refer to the user manuals of all HF devices being used in the procedure, and closely follow all indications, instructions, and safety precautions.

WARNING

Before using any electrosurgical devices for high frequency cauterization, users should be thoroughly familiar with all guidelines, safety precautions, and proper use of the equipment. Follow all manufacturer instructions on proper equipment preparation and use. Accessories should be inspected for damage before and after each procedure.

Before electrosurgery, inspect the endoscope for any physical damage to surfaces and components. If damage is discovered, discontinue use and contact Cogentix Medical for repair. Continued use of damaged equipment during electrosurgery may lead to equipment damage and/or patient injury.

A thorough understanding of the principles and techniques involved in electrosurgical procedures is essential to avoid shock and burn hazards to both patient and medical personnel and prevent damage to the device and other medical instruments. Ensure that insulation or grounding is not compromised.

Do not immerse electrosurgical instruments in liquids unless the instruments are specifically designed and labeled to function in liquids.

WARNING

Set the high-frequency (peak) voltage level of the electrosurgical unit no higher than the voltages given below for the respective operating modes:

CUT: 560 V_p
 COAG: 775 V_p
 SPRAY: 1,700 V_p

Always utilize the lowest output setting necessary on the Electrosurgical Unit. This reduces the potential for patient injury or equipment damage.

When the recommended voltages shown in the instructions for electrosurgical accessories differ from normally acceptable limits, always use the lowest recommended voltage.

Always confirm that the electrode section of the electrosurgical accessory is an appropriate distance from the Distal Tip of the endoscope and that the electrode is clearly in view. If the electrode is in close proximity to the Distal Tip or still within the **Slide-On**[®] **EndoSheath**[®] **Technology** during use, the endoscope and/or **Slide-On**[®] **EndoSheath**[®] **Technology** may be damaged and patient injury may occur.

Do not supply oxygen or use in presence of combustible gases during electrosurgery. There is the potential for combustion during cauterization.

Cogentix Medical recommends the use of isolated electrosurgical accessories. Use of non-isolated accessories may result in operator injury.

To best determine the necessary minimum output, operators should conduct basic tests before electrosurgery according to the User's Manual of the Electrosurgical Unit.

Laser Devices/Accessories



Before using any laser devices, users should be thoroughly familiar with all guidelines, safety precautions, and proper use of the equipment. This includes, but is not limited to, proper eye and skin safety guidelines.

Do not use a damaged laser fiber accessory. Utilizing a probe with a damaged cover or distal end may result in patient injury and/or equipment damage.

Always confirm that the distal tip of the laser fiber accessory is an appropriate distance from the Distal Tip of the endoscope and that the laser fiber tip is clearly in view. If the distal tip of the accessory is in close proximity to the endoscope's Distal Tip or still within the Slide-On® EndoSheath® Technology during use, the endoscope and/or Slide-On® EndoSheath® Technology may be damaged and patient injury may occur.

Do not supply oxygen when performing laser surgery; do not perform laser surgery in the presence of combustible gases. There is the potential for combustion during cauterization.

Follow all manufacturer instructions on proper equipment preparation and use. Accessories should be inspected for damage before and after each procedure.



Before inserting or removing the laser fiber accessory, ensure that the endoscope's Distal Bending Section is in the neutral position and straight. If the Distal Bending Section is articulated, there is a risk of damaging the Instrument Channel of the **Slide-On**[®] **EndoSheath**[®] **Technology**.

6 Reprocessing

The CST-5000/5000*i* Flexible Video Cystoscope/Hysteroscope utilizes the Slide-On® Endo-Sheath® Technology. The Slide-On® EndoSheath® Technology is a sterile, disposable protective covering which limits the need for elaborate chemical disinfection or sterilization procedures after every endoscopy procedure. The complete system enables the user to implement a fast and effective method of reprocessing the endoscope and benefits both user and patient by providing an Insertion Tube covered with a sterile Sheath for every procedure. For cleaning/disinfection procedures when using the EndoSheath® Technology, see Cleaning After Slide-On® EndoSheath® Technology Use on page 41.



The endoscope must be properly reprocessed by cleaning, disinfection or sterilization, before its first use and after each subsequent use according to protocols in this section. Using an endoscope in a procedure that has not been properly reprocessed presents an acute infection-control risk to both the patient and medical personnel performing or assisting in the procedure.



Some methods of disinfection and sterilization may be harmful to the endoscope and exposure to them could result in extensive equipment damage. Please contact Cogentix Medical Customer Service to verify the compatibility of a cleaning method not listed in this manual and/or a complete list of functionally compatible agents.

Clean the endoscope immediately after use in a procedure. Failure to do so may allow patient debris to harden on the endoscope's external surfaces, which can become difficult to remove and could inhibit subsequent disinfection/sterilization processes.

Do not use an endoscope that has been determined to have a leak, and do not immerse such an endoscope in any fluids. Fluid entry into the endoscope can cause equipment damage and render the endoscope unfit for patient use.

Always wear appropriate personal protective equipment when reprocessing the endoscope or any of its components. Appropriate protective equipment includes items such as a gown, gloves, and face and eye shields.

Complete and thorough reprocessing of the endoscope is the only way to ensure that a "patient-ready" endoscope is used in all patient procedures. Closely adhere to the reprocessing instructions given in this chapter.

Reprocessing Steps

The endoscope reprocessing procedure is made up of a series of discrete steps, each of which is essential to successful reprocessing. The steps are listed below in their proper order, and the complete instructions for each step are given in greater detail in this chapter.

- **Leak Testing** The reprocessing procedure requires exposing the endoscope's surface to and immersing the endoscope itself in fluids. If there is a leak in any part of the endoscope, the internal components of the endoscope are vulnerable and will likely be damaged by fluid ingress. Before cleaning, disinfecting and/or sterilizing the endoscope, it is essential to perform a leak test to ensure the interior of the endoscope is resistant to fluid invasion.
- **Cleaning** Visible debris is removed from the surface of the endoscope in this procedure, which uses water and an instrument-grade detergent. When the Slide-On® EndoSheath® Technology is used and inspection after the procedure confirms that the Sheath was not compromised, surface cleaning and intermediate level disinfection of the endoscope should be sufficient to prepare it for the next procedure.
- Intermediate-Level Disinfection After use with an EndoSheath® **Technology** and proper cleaning, the endoscope should undergo intermediate-level disinfection. For the complete routine, see Cleaning After Slide-On[®] EndoSheath[®] Technology Use on page 41.
- **High-Level Disinfection** If the user suspects the endoscope has been contaminated, it will be necessary to immerse the endoscope in a high-level disinfectant.
- **Sterilization** In addition to high-level disinfection, the endoscope may be sterilized using ethylene oxide (EtO) gas. It must then be thoroughly aerated to ensure that all residues have been removed. The endoscope may also be sterilized using a validated STERRAD[®] or STERIS[®] system. Refer to the STERRAD®/ STERIS® section in this chapter.

Leak Testing

The CST-5000/5000i Flexible Video Cystoscope/Hysteroscope must be evaluated for possible leaks in the Control Body and/or the Insertion Tube before immersion in any fluids. The Leak Tester accessory should be used for this test. (Cogentix Medical Leak Tester is required.) Follow the steps given below.



It is essential that gloves be worn when performing the leak test procedure, in case the endoscope's Insertion Tube has been contaminated and requires further disinfection or sterilization. An endoscope in this condition can present an infection-control risk to the person(s) reprocessing the endoscope.

Attach the Leak Tester to the Endoscope

Connect the Leak Tester to the endoscope's Vent Valve (see Figure 6-1). Align the slot on the Leak Tester's connector with the pin on the Vent Valve, then push down and rotate the connector clockwise until it locks.

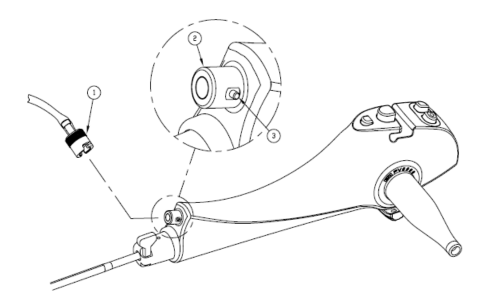


Figure 6-1: Leak Tester Connection

Leak Tester Components

- 1. Connector
- 2. Vent Valve
- 3. Alignment Pin (align with slot in Connector)

Pressurize the Endoscope



The Videoscope Cable's Sealing Cap must be attached to the plug end of the Videoscope Cable Connector prior to leak testing or immersion of the endoscope in water or disinfecting solution.

- 1. Make sure that the Leak Tester's Pressure-Relief Valve is closed by moving the button to the "out" position.
- 2. Pump the hand bulb of the Leak Tester until the pressure gauge's needle reaches **the green zone**. Due to the size of the internal space of the endoscope, 2-3 pumps of the hand bulb may be required to pressurize the entire chamber. After the first pump, the needle may drop out of the green zone and reach a stable position in the white zone. Continue with additional pumps until the needle no longer falls back into the white zone.

CAUTION

Do not over-pressurize the interior of the endoscope (do not allow the needle to go above the green area on the gauge). Overpressurizing the interior of the endoscope can damage the lighttransmission and/or optical system components.

- 3. Maintain the pressure for ten (10) seconds, observing the position of the needle on the pressure gauge. If the pressure decreases, the Sealing Cap of the Videoscope Cable may not be secured onto the plug; or the Leak Tester to endoscope connection may be loose. Check the Pressure-Relief Valve on the Leak Tester, it may still be open and should be closed. Make sure the Sealing Cap is securely placed over the plug; remove and reattach the Leak Tester to the endoscope and repeat Steps 1-3.
 - If the pressure decreases after the connections are restored, the endoscope has a damaged seal. Do not continue to use the endoscope or immerse it in fluids in this condition. Contact your regional distributor or the Cogentix Medical Customer Service Center to arrange for evaluation and/or repair. When returning the endoscope, follow the instructions given in Chapter 9, Warranty and Service.
- 4. If the needle's position remains steady on the Leak Tester, immerse the entire endoscope in water, and observe it for thirty (30) seconds. Angulate the Distal Bending Section up and down while the endoscope is immersed, as holes in the soft covering of the Distal Bending Section may not be evident while it is in a relaxed position.
- 5. A steady stream of air bubbles at a given location indicates a small leak in the endoscope that was not detected by the pressure gauge. If a leak is detected, the air pressure in the endoscope will prevent water from entering through the leak. However, immediately remove the endoscope from the water and do not immerse it in any more fluids.

Do not continue to use an endoscope if leaks are detected. Contact your local distributor or the Cogentix Medical Customer Service Center to arrange for evaluation and/or repair. When returning the endoscope, follow the instructions given in Chapter 9, Warranty and Service.



Note: Do not mistake the release of trapped air from the crevices on the endoscope's outer surface for a leak. Trapped air can be released by tapping the endoscope gently after immersing it in

- 6. The absence of air bubbles confirms that the endoscope is watertight. Remove it from the water and open the Leak Tester's Valve.
- 7. Make sure that the needle on the pressure gauge falls to zero (0), and disconnect the Leak Tester from the endoscope. The endoscope can now be safely immersed in cleaning solutions.



Failure to discharge/depressurize the endoscope after leak testing may place stress on the soft covering of the Insertion Tube, potentially producing a "rolling over" of the covering.

Cleaning/Disinfection/Sterilization

Use of the Vent Cap



Failure to follow the instructions given in this section regarding the use of the Vent Cap may result in damage to the endoscope. Any such damage will void the product warranty.

The Vent Cap is to be **attached** to the endoscope prior to <u>all</u> of the following procedures in order to prevent damage to the endoscope caused by changes in pressure and temperature:

- Gas Sterilization
- Aeration
- Shipping

The Vent Cap is to be **removed** from the endoscope prior to:

- Patient Procedures
- Immersion in Fluids

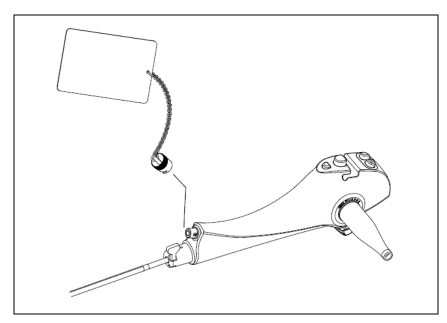


Figure 6-2: Opening the Vent Valve

Cleaning After Slide-On® EndoSheath® Technology Use



The **Slide-On**[®] **EndoSheath**[®] **Technology** is intended for a single use only; **do not reuse it**. When the procedure is complete, remove the Sheath from the endoscope and dispose of it as described in Chapter 4, **Installing and Removing the Slide-On**[®] **Endo-Sheath**[®] **Technology**. Reusing the Sheath can present an acute infection-control risk to the user and the next patient on whom the endoscope is used.

Reprocessing

After a procedure in which the **Slide-On® EndoSheath® Technology** was attached over the endoscope's Insertion Tube, Cogentix Medical recommends performing the following prophylactic cleaning and intermediate disinfection routine between endoscopic procedures:

CAUTION

Because of the possibility that a Sheath could be torn, or that the endoscope or Sheath could come in contact with contaminated surfaces, the user should develop and follow a prophylactic routine when handling the endoscope, sheathed or unsheathed.

- After removing the Sheath, inspect the Insertion Tube and Distal Bending Section, and confirm that these areas are dry. If moisture is observed, there may have been a leak into the Sheath during the procedure, providing the endoscope was dry when the sheath was attached. In this case, the endoscope must be high-level disinfected or sterilized following the instructions given in this chapter.
- 2. **For Cleaning** Gently wash all external surfaces of the endoscope with an appropriate instrument-grade detergent. An ample size basin must be used for cleaning the endoscope. If the basin is too small, the endoscope may inadvertently be kinked or damaged during cleaning.
- 3. After washing, thoroughly rinse the outside of the endoscope with clean, lukewarm water and place it on a clean, dry surface.
- 4. **For Intermediate Level Disinfection** Wipe down the entire endoscope with a soft, lint-free cloth (gauze) soaked in 70% ethyl or isopropyl alcohol.
- 5. Ensure that all external surfaces of the endoscope are **thoroughly dried** prior to attaching another Sheath or storing the endoscope.

High-Level Disinfection and Sterilization

In the event that the endoscope is contaminated, it should be high-level disinfected or sterilized after cleaning. Use caution when cleaning and then high-level disinfecting or sterilizing the endoscope.

Recommended Disinfection and Sterilization Procedures

The following procedures have been determined by Cogentix Medical to be compatible with the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope.

- High-Level Disinfection in Glutaraldehyde: All Cogentix Medical endoscopes are validated for high-level disinfection in 2.4% glutaraldehyde solutions. Perform the High-Level Disinfection Protocol described in this chapter.
- **Sterilization by Ethylene Oxide (EtO) Gas**: The endoscope may be sterilized using a validated EtO protocol. The acceptable processing parameters and procedure are given in the "Ethylene Oxide (EtO) Gas Sterilization" section beginning on page 45.

- **Sterilization by STERRAD**®: The endoscope may be sterilized using a validated **STERRAD**® protocol. Refer to Table 6-1 on page 46 for the approved systems / cycles suitable for use with this endoscope.
- **Sterilization by STERIS**[®]: The endoscope may be sterilized using a validated **STERIS**[®] protocol. Refer to Table 6-1 on page 46 for the approved system suitable for use with this endoscope.



Disinfection and sterilization methods not listed here may be harmful to the endoscope and could cause extensive equipment damage. Please contact Cogentix Medical Customer Service to determine the compatibility of a disinfection or sterilization method not listed in this manual and/or a complete list of functionally compatible agents.

Acceptable Reprocessing Materials

Cleaning	 Soft Material Lint-Free Gauze (4x4) Enzymatic Cleaner Instrument Grade Detergent
Intermediate Level Disinfection	70% Isopropyl Alcohol or70% Ethyl Alcohol
High Level Disinfection	2.4% Glutaraldehyde-based solution
Sterilization	 EtO Gas Sterilization STERRAD[®] 100S, NX, 100NX* STERIS[®] System 1E*

^{*} Endoscope **must** feature the (S) symbol for STERIS[®] / STERRAD[®] compatibility

Incompatible Methods

The high-level disinfection and sterilization chemicals and methods shown below are not compatible with the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope; **DO NOT USE THEM**, as they could cause extensive damage to the endoscope. If you have any questions regarding the compatibility of a given disinfection or sterilization method, please contact your local distributor or the Cogentix Medical Customer Service Center.

Incompatible High Level Disinfection and Sterilization Methods/Chemicals High Level Disinfection DO NOT USE • Chlorines • Formaldehyde

Chemicals	DO NOT USE		Formaldehyde Iodophors
Sterilization Methods	DO NOT USE	•	Autoclave Ultrasonic

High-Level Disinfection Protocol

If the endoscope was determined to be free of leaks, it may be immersed in a glutaraldehyde solution for the amount of time recommended by the disinfectant manufacturer to achieve high-level disinfection.



It is imperative that the endoscope be leak tested and cleaned prior to immersion in high-level disinfectant. Failure to do so may not detect leaks that could allow fluid ingress and damage the endoscope. Failure to clean the endoscope may allow gross debris to remain on external surfaces, which could impair proper disinfection. In the event that the endoscope fails the leak test, do not immerse the endoscope in liquids and do not use it in a procedure. Return the endoscope to the manufacturer for repairs.

Make sure that the sealing cap is securely attached to the Videoscope Cable prior to immersion. Failure to do so will lead to fluid invasion and severe equipment damage.

Pre-Cleaning

- 1. Gently wipe down the Insertion Tube and Distal Bending Section with a soft, lint-free cloth or gauze (4x4) to remove visible debris.
- 2. Perform the Leak Test procedure.
- 3. Gently wash down all external surfaces with an enzymatic cleaning solution and soak the endoscope in the enzymatic cleaning solution for the time recommended by the enzymatic solution's manufacturer.
- 4. Remove the endoscope from the cleaning solution and rinse it thoroughly with clean, lukewarm water.
- 5. Dry all external surfaces of the endoscope.

Disinfection

- 1. Immerse the endoscope in the disinfectant solution at the temperature recommended by the disinfectant manufacturer.
- 2. Allow the endoscope to remain immersed in the disinfectant solution for the period of time recommended by the disinfectant manufacturer.
- 3. Following disinfection, remove the endoscope from the solution.

Rinsing

- 1. Immerse the endoscope in a container of clean, lukewarm water.
- 2. Thoroughly rinse the outside of the endoscope with clean, lukewarm water and place it on a clean, dry surface.
- 3. Wipe all external surfaces of the endoscope with a soft, lint-free cloth or gauze (4x4) until it is completely dry.
- 4. Confirm that the Lens at the endoscope's Distal Tip is free of disinfectant residue.

Ethylene Oxide (EtO) Gas Sterilization

The **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope may be sterilized using a validated ethylene oxide (EtO) gas sterilization protocol, following the processing parameters given below.



If the Vent Valve is not opened during gas sterilization, the increased heat and pressure from the sterilization process will cause pressure to build up inside the endoscope and could rupture the watertight seals and/or softer materials of the endoscope.



The Videoscope Cable's Sealing Cap <u>must be removed</u> from the Videoscope Cable Connector prior to EtO Sterilization.

EtO Gas Sterilization Parameters

Temperature: $125^{\circ} \pm 5^{\circ}F$ ($52^{\circ}C \pm 3^{\circ}C$) **Relative Humidity:** $50\% \pm 10\%$ **EtO Concentration:** 600 mg/liter **Exposure Time:** 3 hours + 1/-0 hour

Post-Sterilization Aeration: 12 hours at 130°F (55°C) or 72 hours

at 75°F (24°C)



NOTE: EtO Gas Sterilization at the above parameters has been validated by Cogentix Medical, and will sterilize the device to a sterility assurance level (SAL) of 10⁻⁶.

Prior to EtO Gas Sterilization, the endoscope must be leak tested, precleaned and dried as described for the High-Level Disinfection Protocol.

CAUTION

Failure to properly pre-clean the endoscope may inhibit the EtO gas sterilization process.

Prior to Gas Sterilization, the Vent Valve must be opened as shown in Figure 6-2 (page 41) to accommodate the heat and pressure changes of the Gas Sterilization process. To open the Valve, press the red Vent Cap onto the Vent Valve, and rotate it clockwise until it is seated and locked.

After EtO Gas Sterilization

Effective aeration must be completed after EtO gas sterilization. Cogentix Medical recommends following the instructions-for-use supplied by the manufacturer of the gas sterilizer, and that a biological indicator be used to confirm sterilization efficacy.

STERRAD® and STERIS® Sterilization

Prior to STERRAD® and STERIS® Sterilization, the endoscope must be leak tested, cleaned and dried as described in this chapter.

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope has been validated for material and functional compatibility with the following sterilization systems/cycles:

STERRAD [®] 100S	STERRAD [®] NX	STERRAD [®] 100NX	STERIS [®] System 1E
✓	✓	✓	✓
Short and Long Cycles	Short and Long Cycles	Flex Cycle Only	

Table 6-1: STERRAD® and STERIS® Validated Systems/Cycles

Refer to the STERRAD® or STERIS® Sterilization System User's Manual for complete details on instructions for use.



Failure to properly clean the videoscope may inhibit the STERRAD® or STERIS® sterilization process.



The Vent Cap **must** be **attached** for the STERRAD® sterilization process. The Vent Cap must be removed for the STERIS® sterilization process.



STERRAD® or STERIS® compatibility applies only to endoscopes which feature the (S) symbol located on the Control Body's Identification Ring.

7 Care and Storage

Follow the instructions in this chapter if you anticipate that the endoscope will not be used for a prolonged period of time. Do not leave the endoscope exposed to the elements in such circumstances.

Storage

Follow the instructions below when storing the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope:

- **DO NOT** store the endoscope with a **Slide-On**[®] **EndoSheath**[®] **Technology** installed on the Insertion Tube. Over time, the Sheath material may adhere to the Insertion Tube and become difficult to remove.
- When storing the endoscope, be sure to keep the Insertion Tube as straight as possible. The Videoscope Cable may be stored either straight or neatly coiled to prevent kinking or bending.
- The equipment should be completely clean and dry before storing.
- This endoscope should be maintained in a clean condition during storage so that it is ready for subsequent use.
- The endoscope should be stored in a dry, well ventilated environment avoid high humidity, direct sunlight, and temperatures below -10°C or above 60°C.
- Do not store the endoscope in its carrying case. This case is only intended for endoscope transport; it is not properly ventilated for storage.
- Avoid storing the endoscope in heavily trafficked areas where there is a chance that it may sustain physical damage.



The endoscope should **NEVER** be stored in areas where it could be exposed to liquids or environmental conditions such as high temperature, humidity, direct sunlight, dust, salt, etc., which could adversely affect its operation.

The endoscope should **NEVER** be stored in the presence of flammable or explosive gases or chemicals.

Care and Storage

Disposal

The equipment should be returned to Cogentix Medical for disposal.

Contact your local Cogentix Medical representative or service facility for further information.

The information in this chapter is intended to help users diagnose problems that may occur during operation of the endoscope. The tables include some of the problems that could arise during operation, possible causes for those problems, and suggested corrective action.



LAUTION If the problem persists even after the corrective action has been taken, or a problem occurs that is not covered in the tables, do not use the endoscope. Contact Cogentix Medical for service or repairs using the information given in Chapter 9, Warranty and Service.

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope requires a video processor to process and display images. In order to identify issues related to image problems, you may also have to refer to the Troubleshooting chapter in the **DPU-5000/7000 Series** Video Processor User's Manual.

PROBLEM	Probable Cause	Action
Angulation feels stiff	Damaged Distal Bending Section causing impaired angulation.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Angulation alignment is no longer up/down	Insertion Tube has become twisted.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Loss of angulation	Angulation wires have been stretched or broken during use.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Cloudy or foggy images or poor image quality when the	Patient debris or other material on the Objective Lens.	⇔ Clean the Objective Lens with an alcohol prep pad to remove material or stain. Excess staining may not be correctable and the lens may require replacement.
endoscope is unsheathed	Fluid incursion into the videoscope's optical system.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
	Improper settings on the Video Processor or Display.	 Adjust the settings on the Video Processor or Display. Perform a White Balance procedure on the Video Processor.
	Damaged optics, sensor or electronics in the endoscope.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
No image	Video Processor is not powered on.	
	Connection between the endoscope and the Video Processor is lost.	⇔ Check the cable connection between the endoscope and the Video Processor.
	No video output signal to a monitor.	⇔ Check the video output cable connections when using an external monitor. Replace the cable if necessary.
		⇒ If the problem cannot be corrected, send the endoscope and Video Processor to Cogentix Medical for repair.

Table 8-1: Troubleshooting

PROBLEM	PROBABLE CAUSE	ACTION
Loss of illumination	Patient material or other substance on the Light Guides.	Clean the Distal Tip with an alcohol prep pad to remove material or stain. Excess staining may not be correctable and the lens may require replacement.
	Light Intensity is set too low.	⇒ Adjust Light Intensity setting.
	Damaged light guide fiber bundles.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
	Internal light source is deteriorating.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Wrinkles and/or folds in the Insertion Tube	These may be a result of excessive force applied to the Insertion Tube during cleaning or Sheath removal, or the long-term effects of repeated immersion in chemical disinfecting solutions, which could stretch and weaken the outer coverings.	⇒ Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Insertion tube is dented	Dents can be caused by physical trauma to the endoscope [e.g., closing the case on the Insertion Tube].	⇒ Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Loss of pressure during leak test	The Leak Tester is not connected properly to the Vent Valve.	⇒ Re-connect the Leak Tester and perform the test again.
	The Leak Tester's Pressure-Relief Valve is open.	
	A hole or crack has broken the endoscope's watertight seal.	Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.

Table 8-1: Troubleshooting (cont'd)

Cannot insert The Sheath's channel is Straighten the channel before continuing scope endoscope into insertion. misaligned. the sheath Make sure the Locking Knob is positioned The endoscope's Locking vertically. Align Locking Knob with slot and Knob is not aligned with fully seat the endoscope in the sheath. Rotate the slot. the Locking Knob clockwise until it is horizontal. The endoscope's Insertion Return the endoscope to Cogentix Medical Tube has been damaged for repair. Refer to Chapter 9, Warranty and its diameter has and Service. increased. Place the Angulation Lever in the neutral The Angulation Lever has position so the Distal Bending Section of the been actuated. endoscope is straight. The Sheath's Window is Articulate the Insertion Tube back and forth not aligned with the several times to fully seat the Sheath. Insertion Tube. Gently manipulate the edges of the Sheath's Window to fully seat it. The D-shaped opening of Align the opening of the EndoSheath® the EndoSheath® Connector with the Insertion Tube. Connector is not aligned with the Insertion Tube. The Sheath is torn or Replace the Sheath. punctured. Cannot remove Locking Knob not rotated. Rotate Locking Knob to the vertical position. the endoscope from the sheath The Sheath is torn or Contact your regional distributor or punctured. Cogentix Medical for removal instructions. Disinfect/sterilize the endoscope. The Sheath has been Use a syringe without a needle to introduce attached to the small amounts of 70% alcohol into the opening at the proximal end of the EndoSheath® endoscope for an extended period of time, Connector. When the liquid reaches the Distal causing it to adhere to the Tip of the endoscope/sheath, carefully attempt Insertion Tube. to remove the endoscope from the Sheath. If necessary, contact your Regional Distributor or Territory Manager for removal instructions. Disinfect/sterilize the endoscope. The flat portion of the Align the flat portion of the Insertion Tube with Insertion Tube is not the Biopsy Port and then remove the properly aligned with the endoscope from the Sheath. Biopsy Port. Gently grasp the Sheath's Optical Window The Distal Head of the using the hanging drape bag as a barrier Insertion Tube is lodged between your fingers and the Sheath. Then in the Sheath. slowly and gently withdraw the endoscope from the Sheath.

Table 8-1: Troubleshooting (cont'd)

			1
Glare evident when viewing through the sheath	The Sheath's window is not in contact with the Distal Tip of the endoscope due to:		
	- Incomplete insertion into the Sheath.	↑ ↑	Move the Distal Bending Section back and forth several times to fully seat the Sheath. Gently manipulate the edges of the Sheath's
	OR		window to properly seat the endoscope.
	 The endoscope's tip is damaged and/or enlarged. 	₽	Remove the endoscope from the Sheath. Return the endoscope to Cogentix Medical for repair. Refer to Chapter 9, Warranty and Service.
Insufficient or no irrigation during procedure	The stopcock is closed.	↔	Open the stopcock.
	The Irrigation Tubing is kinked, crushed, or open to the atmosphere.	⇔	Straighten out or relieve any tension on the tubing.
	 Irrigation Tubing is tightly positioned in the endoscope's Flow Control Valve. 	⇔	Relieve tension on the tubing by repositioning it to provide more slack ahead of the Flow Control Valve.
	The Flow Control Valve is closed.	\Rightarrow	Open the Flow Control Valve.
Cannot inject fluid through the Accessory Port	The Working Channel is clogged.	↔	Remove the endoscope from the patient and inject 10cc of saline through the channel. If the channel remains blocked, remove the Sheath from the endoscope and install a new Sheath.
	 The Syringe is not fully seated into or aligned with the Accessory Port. 	⇔	Insert the syringe tip fully and straight into the Accessory port.

Table 8-1: Troubleshooting (cont'd)

The Accessory will not pass through the Sheath's Working Channel	Accessory is too large.	⇧	Verify that the CV-2.1 Slide-On® Endo- Sheath® is being used. Check the diameter of the accessory. Make sure that the Accessory is compatible with a 2.1 mm Working Channel.
	The Channel is occluded by patient debris	↔	Remove the endoscope from the patient and inject 10cc of saline through the Channel. If, after repeated attempts, the channel remains blocked, remove the Sheath from the endoscope and install a new one. Do not force the Accessory through the working Channel as this may damage the Channel and compromise the integrity of the barrier.
	The endoscope's Distal Bending Section is angulated.	↔	Maneuver the endoscope's Distal Tip to an area of the anatomy in which it can be safely straightened out. Insert the Accessory until its tip can be seen on the endoscopic image. Now angulate the endoscope's Distal Bending Section and proceed to the desired area.

Table 8-1: Troubleshooting (cont'd)

9 Warranty and Service

Warranty Information



NOTE: Alterations or repairs done by persons not authorized by Cogentix Medical will void this warranty.

Cogentix Medical is not liable for any damages to the **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope resulting from misuse, negligence, or improper cleaning or storage. The warranty defined herein shall apply only to the original buyer. In no event shall Cogentix Medical be liable for anticipated profits, consequential damages or loss of time incurred by the buyer with the purchase or use of this equipment.



Note: Cogentix Medical sells many of its products through regional distributors. Before sending equipment to Cogentix Medical, contact your regional distributor for repair/return procedures.

Cogentix Medical warrants that the **CST-5000/5000** Flexible Video Cystoscope/Hysteroscope and its accessories will be free from defects in materials and workmanship for a period of one year from the date of the invoice. Replacement parts are warranted for a period of ninety (90) days from the date of the invoice.

All non-warranty repairs will be warranted to be free from defects in materials and workmanship for a period of ninety (90) days from the date of the invoice.

Upon receipt of a **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope for repair, Cogentix Medical will evaluate the instrument and make the final decision as to the warranty status.

The above warranties are in lieu of all other warranties, either expressed or implied, including warranties of fitness or merchantability.

Cogentix Medical Service Information

CST-5000/5000i Flexible Video Cystoscopes/Hysteroscopes are serviced at authorized Cogentix Medical repair facilities only. Use the following procedure to expedite returned goods for repair or replacement:

1. Telephone your Regional Distributor, Territory Manager, or Cogentix Medical Customer Service (Monday through Friday from 8:00 AM to 7:00 PM EST).

USA customers call **1-866-258-2182** International customers call (+1) 952-426-6189 for Cogentix Medical Customer Service or call your regional distributor

- 2. Provide a detailed description of the problem.
- 3. If troubleshooting cannot solve the problem, a Returned Goods Authorization (RGA) number will be issued.
- 4. Complete an Incident Report Form and send it to Cogentix Medical along with the returned goods. Returned merchandise will only be accepted with an RGA number.

Shipping to Cogentix Medical or Distributor



If the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope has been used in a clinical setting, disinfect all system components before shipping as described in Chapter 6, Shipping contaminated equipment could Reprocessing. present an acute infection-control risk for those handling the endoscope, both during shipping and at Cogentix Medical or authorized repair facility.

If the **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope has been used in a clinical setting but cannot be disinfected before shipping, place a red biohazard label on the shipping container to indicate that the contents are contaminated, in accordance with OSHA standards 29 CFR 1910.1030.

Observe the following precautions before shipping the endoscope:

- 1. Attach the Vent Cap to the endoscope's Vent Valve in preparation for shipping.
- 2. If the endoscope has a leak or tear or fails the leak test, or for some other reason cannot be disinfected properly as described in Chapter 6, **Reprocessing**, wipe the endoscope down with 70% ethyl or isopropyl alcohol to remove debris. Indicate on the outer package that the contents are contaminated.
- 3. Ship the endoscope in its carrying case. Place the carrying case inside a corrugated box containing protective shipping material to prevent damage during shipment.

Warranty and Service

Regardless of warranty status, all shipping charges to and from an authorized Cogentix Medical facility are the responsibility of the customer.



Note: The customer will be contacted and advised of the estimated repair costs. Repairs will not begin on any equipment until authorization or a purchase order has been issued indicating approval of the charges.

Appendix

Specifications CST-5000 / CST-5000i

	1	1
Insertion Tube Diameter	CV-2.1 Sheath (08-5101)	5.6 mm (16.8 Fr) with sheath
(Nominal)	CV-1.5 Sheath (08-5102)	5.2 mm (15.6 Fr) with sheath
Overall Endoscope Length (Dis	tal Tip to	650 mm
Top of Control Body)		
Insertion Tube Working Length	1 w/ Sheath	370 mm
Field of View in Air/Water		110° (air) / 76° (water)
Direction of View		Forward
Depth of Field		3 - 50 mm
Environmental Effects on Optic Performance	cal	None
Angulation		215° Up/140° Down
Working Channel Diameter	CV-2.1 Sheath	2.1 mm (6.3 Fr)
	(08-5101)	
Channel Diameter	CV-1.5 Sheath (08-5102)	1.5 mm (4.5 Fr)
Operating Environment		
Temperature		50° to 104° F (10° to 40° C)
Relative Humidity		30 to 85%
Air Pressure		700 to 1060 hPa
Storage Environment		
Temperature		14° to 140° F (-10° to +60° C)
Relative Humidity		0 to 95%
Air Pressure		700 to 1060 hPa
Mode of Operation		Continuous
Electrical Safety		IEC 60601-1 & IEC 60601-2-18
Thermal Safety		IEC 60601-1 & IEC 60601-2-18
Electromagnetic Compatibility		IEC 60601-1-2
Degree of Protection Against Electrical Shock		Type BF
Degree of Protection Against I	nvasion of	Fully Immersible (as per
Liquids		Reprocessing Instructions)

Table A-1: Specifications

Infection Control Information

The Slide-On® EndoSheath® Technology for the CST-5000/5000i Flexible Video Cystoscope/Hysteroscope is designed to offer practitioners the ability to perform safe, efficient endoscopy. The Slide-On® EndoSheath® Technology has been proven to be an effective barrier to microorganisms as small as 27 nanometers¹. Efficacy testing for barrier qualities has been performed by an independent laboratory per FDA required guidelines². All **Slide-On**® EndoSheath® Technology undergoes a rigorous Quality Assurance process to ensure the utmost in product quality and efficacy.

Please contact Cogentix Medical Customer Service for a detailed information packet regarding Infection Control and the Slide-On® EndoSheath® Technology.

¹ Viral challenge testing performed with bacteriophage 27 nanometers in size, per FDA guidance.

² Per FDA Clearance requirements, barriers must be tested by the guidelines set within the "FDA Guidance for Manufacturers Seeking Marketing Clearance of ... Endoscope Sheaths Used as Protective Barriers"

Electromagnetic Compatibility Declarations



Use of accessories not specified in this manual or sold by Cogentix Medical may result in increased electromagnetic emissions or decreased immunity of the equipment or system.

Guidance and manufacturer's declaration - electromagnetic emissions

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope with **DPU-5000/7000 Series** Digital Video Processor [the "System"] is intended for use in the electromagnetic environments specified below. The customer or the user of the System should ensure that it is always used in such environments.

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

Table A-2: Electromagnetic Emissions Declaration

Guidance and manufacturer's declaration – electromagnetic immunity

The CST-5000/5000*i* Flexible Video Cystoscope/Hysteroscope with DPU-5000/7000 Series Digital Video Processor [the "System"] is intended for use in the electromagnetic environments specified below. The customer or the user of the System should ensure that it is always used in such environments.

IEC 60601 test level	Compliance level	Electromagnetic environment – guidance
±6kV contact ±8kV air	±6kV contact ±8kV air	Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative
		humidity should be at least 30%.
±2kV for power supply lines	±2kV for power supply lines	Mains power quality should be the equivalent of that in a typical commercial or hospital environment.
input/output lines	±1 kV for input/output lines	
±1 kV differential mode ±2 kV common mode	±1 kV differential mode ±2 kV common mode	Mains power quality should be the equivalent of that in a typical commercial or hospital environment.
>95% dip U_T for 0.5 cycle 60% dip U_T for 5 cycles 30% dip U_T for 25 cycles 95% dip U_T for 5 sec	Compliant with all levels of voltage dips for $U_T = 100 \text{ VAC}$ and $U_T = 240 \text{ VAC}$	Mains power quality should be the equivalent of that in a typical commercial or hospital environment. If the user of the System requires continued operation during power mains interrupts, it is recommended that the System be powered from an uninterruptible power supply or a battery.
3 A/m	3 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a standard commercial or hospital environment.
	±6kV contact ±8kV air ±2kV for power supply lines ±1 kV for input/output lines ±1 kV differential mode ±2 kV common mode >95% dip U _T for 0.5 cycle 60% dip U _T for 5 cycles 30% dip U _T for 25 cycles 95% dip U _T for 5 sec	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$

Table A-3: Electromagnetic Immunity Declaration

Guidance and manufacturer's declaration – electromagnetic immunity

The **CST-5000/5000***i* Flexible Video Cystoscope/Hysteroscope with the **DPU-5000/7000 Series** Video Processor [the "System"] is intended for use in the electromagnetic environment specified below. The customer or the user of the System should assure that it is always used in such environments.

Immunity Test	IEC 60601 test level	Compliance level	Electromagnetic environment – guidance
			Portable and mobile RF communications equipment should be used no closer to any part of the System, including cables, than the recommend separation distance calculated from the equation applicable to the frequency of the transmitter.
Conducted RF	3 Vrms	3 V	Recommend separation distance $d = 1.17\sqrt{P}$
IEC 61000-4-6	150 kHz to 80	3 v	d = 1.17√P 80MHz to 800MHz
	MHz		d = 2.33√P 800MHz to 2.5GHz
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m	where <i>P</i> is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and <i>d</i> is the recommended separation distance in meters (m).
			Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, ^a should be less than the compliance level in each frequency range. ^b
			Interference may occur in the vicinity of equipment marked with the following symbol:
			$((\bullet))$

NOTE 1: At 80 MHz and 800 MHz the higher frequency range applies.

NOTE 2: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Table A-4: Electromagnetic Immunity Declaration

Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the System is used exceeds the applicable RF compliance level above, the System should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary such as re-orienting or relocating the System.

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

Recommended separation distances betweenportable and mobile RF communications equipment and the System

The CST-5000/5000/ Flexible Video Cystoscope/Hysteroscope with the DPU-5000/7000 **Series** Video Processor [the "System"] is intended for use in the electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the System can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the System as recommended below according to the maximum output power of the communications equipment.

Radiated maximum output power of	Separation distance according to frequency of transmitter				
transmitter	m				
w	150 KHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.5 GHz		
	$D = [3.5/V_1]\sqrt{P}$	$D = [3.5/E_1]\sqrt{P}$	$D = [7/E_1]\sqrt{P}$		
0.01	D = 0.12	D = 0.12	D = 0.23		
0.1	D = 0.37	D = 0.37	D = 0.74		
1	D = 1.17	D = 1.17	D = 2.33		
10	D = 3.69	D = 3.69	D = 7.38		
100	D = 11.67	D = 11.67	D = 23.33		

For transmitters rated at a maximum output power not listed above, the recommended separation distance, d, in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1: At 80 MHz and 800 MHz the separation distance for the higher frequency range

applies.

NOTE 2: These guidances may not apply in all situations. Electromagnetic propagation is

affected by absorption and reflection from structures, objects and people.

Table A-5: Recommended Separation Distances





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For additional product information or questions pertaining to sales and service, please contact the local distributor or the manufacturer.



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Covered by one or more of the following U.S. Patents: 6,350,231; 6,530,881; 6,733,440. Other U.S. and international patents pending.

7/15 09645-EN Rev G (13215)