DUR®-Digital Invisio® Flexible Ureteroscope/Choledochoscope

Operations and Technical Manual
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Repair/Return/Warranty

Repair/Return
All returns for credit must have prior written authorization. All medical devices returned to ACMI for any reason must be 1) disassembled, 2) cleaned, and 3) high-level disinfected or sterilized in accordance with the product’s instructions for use or labeling, and shipped in accordance with ACMI’s return procedures (available upon request) and all applicable regulations. To obtain a return authorization number, return addresses, and instructions, please call toll-free (888) 524-7266.

Limited Express Warranty
SHOULD THE PRODUCT BECOME INOPERABLE, DURING NORMAL AND PROPER USE IN ACCORDANCE WITH APPLICABLE INSTRUCTIONS, WITHIN THE TIME FRAME SPECIFIED BELOW FROM THE DATE OF SHIPMENT, ACMI WILL REPAIR OR REPLACE THE PRODUCT, AT ITS SOLE OPTION, AT NO CHARGE. ACMI MAKES NO OTHER WARRANTIES WITH RESPECT TO THE PRODUCTS AND EXPRESSLY DISCLAIMS ALL OTHER WARRANTIES, EXPRESS OR IMPLIED, AS TO MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE OR ANY OTHER MATTER. IN NO EVENT SHALL ACMI BE LIABLE FOR ANY CONSEQUENTIAL DAMAGES. IN NO EVENT SHALL ACMI BE LIABLE FOR ANY BREACH OF WARRANTY IN ANY AMOUNT EXCEEDING THE PURCHASE PRICE OF THE PRODUCT.

This Warranty runs only to the original user and may be voided if the product(s) are serviced or repaired by anyone other than ACMI or an organization duly authorized by ACMI for such purpose.

<table>
<thead>
<tr>
<th>Product</th>
<th>Warranty Period</th>
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<tbody>
<tr>
<td>DUR-Digital Invisio Flexible Ureteroscope (excluding cable)</td>
<td>1 year</td>
</tr>
<tr>
<td>DUR-Digital Invisio Flexible Ureteroscope cable only</td>
<td>1 year</td>
</tr>
<tr>
<td>Accessories</td>
<td>90 days</td>
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Warnings and Cautions

The following warnings and cautions apply to the use, care, and/or maintenance of the ACMI® DUR-Digital Invisio Flexible Ureteroscope System. Failure to comply with, or abide by, any warning or caution set forth in this manual will void the video system’s Limited Express Warranty.

Warnings
(indicate that a danger to life or health can result from misusing the equipment)

1. SERVICE WARNING AND DISCLAIMER NOTICE: It is recommended that the medical equipment for which these documents apply be returned to ACMI for servicing or repair. Service or repair provided by any party other than ACMI or an ACMI-authorized repair facility may result in the user and/or repair facility being liable and responsible for damages, including patient or user injuries, arising from such servicing or repair, and any such servicing or repair shall void the manufacturer’s Limited Express Warranty, if any, applicable to such medical equipment.

2. To prevent electric shock and breakage of seals, do not remove covers on the instrument or IDC Invisio Digital Controller as a potential electric shock hazard exists. Refer all repairs to ACMI-authorized service personnel.

3. Study this manual and other labeling thoroughly for safe handling and storage. Misuse of the equipment can cause injury to the patient and could have an adverse effect on the procedure being performed. Do not drop equipment, or allow it to be struck by other objects.

4. Procedures should be performed only by persons with adequate training and preparation. Consult the medical literature regarding techniques, complications, and hazards prior to any procedure.

5. Do not use electromedical energy sources in the presence of flammable detergents, anesthetics, nitrous oxide (N2O), or oxygen.

6. Consult the operating manuals of all electromedical energy sources used with endoscopic instruments for appropriate instructions, warnings, and cautions prior to use. Such sources of energy include electrical, electrohydraulic, electrosurgical, heat, hydraulic, laser, light, pressure, sound, ultrasound, and vacuum.

7. Make no repairs to any instrument or IDC Invisio Digital Controller. Personal injury or damage to units may result. Refer units to ACMI-authorized service personnel for repair.

8. Never look directly into the light emitted from the instrument. Damage to the eyes can result.

9. Keep the distal tip of any electrode, probe, laser fiber, or other ancillary device in the field of view at all times when active.

10. Use only those lubricants specified in the labeling.

11. Prior to use, examine each electrode and its insulation for damage; do not use if damaged.

12. Follow the disinfectant or sterilizer manufacturer’s recommended procedures and cautions.

13. Do not use disinfectant solutions that contain long-life surfactants; such solutions can leave conductive residues.

14. Do not use an instrument that fails to meet the criteria stated in the labeling or that has been damaged.

15. When instruments from different manufacturers are used together, verify that any isolation or grounding is not violated.

16. Follow the labeling instructions regarding the disposal or reuse of accessories. Reuse of disposable accessories could compromise patient safety.

Cautions
(indicate that equipment or other property may be damaged or may malfunction by misuse)

1. Pay close attention to the care, cleaning, disinfection, and sterilization instructions in this manual. Any deviation can cause damage and void ACMI’s Limited Express Warranty. Do not steam sterilize the instrument or IDC Invisio Digital Controller.

2. To ensure continued satisfactory performance, perform the prescribed inspections and operational tests as recommended.

3. Thoroughly check all electrical cables and plugs before each use and replace any which are damaged or excessively worn. Do not use if damage is suspected or discovered.
4. Check the outer surface of the flexible shaft prior to each use to make certain that the instrument is free of any cuts, holes, rough surfaces, sharp edges, or protrusions. Do not use if damage is suspected or discovered.

5. Do not insert a wet connector into the IDC Invisio Digital Controller receptacle, as poor video performance and/or damage to the system may result.

6. These instruments contain no user-serviceable parts. Refer all repairs to ACMI-authorized service personnel.

7. Connect electromedical energy source power cables only to properly wired grounding receptacles.

8. The instrument is listed by CSA as meeting Medical and Dental Equipment Standard UL 2601 and CSA Standard 22.2 No. 601 and IEC-60601.

9. Prior to use of a cardiac defibrillator, remove endoscopic instruments from the patient. Failure to remove an endoscopic instrument from a patient during use of a cardiac defibrillator could result in damage to the instrument due to the discharge of the cardiac defibrillator.

10. Testing to ensure that chassis leakage does not exceed the allowable levels per the appropriate standard should be performed at least once each year.

11. Do not adjust electronic circuitry. Electronic circuitry is set at the factory and no further adjustment is necessary.

12. Never remove the instrument from the IDC Invisio Digital Controller by pulling on the cable. Always pull the insertion (connector) body to remove the cable from the IDC Digital Controller.

13. U.S. Federal law restricts this device to sale by or on the order of a physician.

Section 1.0 Unpacking and Initial Inspection

Upon receipt, examine the shipping carton and its contents for signs of damage. Check for rattling or loose material inside. Examine the instrument, IDC Invisio Digital Controller (sold separately), and accessories for damage. Do not use a damaged product—contact ACMI Customer Service at (888) 524-7266.

It is very important that upon unpacking the equipment the proper video format selection is made: PAL, VGA (Progressive Scan), or NTSC.


Section 2.0 General Description and Intended Use

The instrument system consists of a Digital Flexible Ureteroscope with an integrated cable, accessories, an IDC Invisio Digital Controller, and all required video connection cables.

The ACMI instrument system (which includes the DUR-Digital Invisio Flexible Ureteroscope, Choledochoscope, and IDC Invisio Digital Controller) is intended for use to examine body cavities, hollow organs and canals in the body, in the urinary tract, and can be used percutaneously to examine the interior of the kidney; and using additional accessories, can be used to perform various diagnostic and therapeutic procedures.

The DUR-D System is also indicated for the examination of bile ducts, and using additional accessories, to perform various diagnostic and therapeutic procedures during cholecystectomy.
Section 3.0 Description of Components

**DUR-D Digital Invisio Flexible Ureteroscope**

The distal tip of the instrument contains a digital imaging sensor that converts light energy into an electrical signal for the production of an intra-operative video image that is displayed on a monitor. The three remote push buttons on the Ureteroscope allow fingertip control of DUR-D System functions, as well as control of external devices and accessories.

**Figure Identifications**

1. Deflection Control Levers UP and DOWN
2. Programmable Remote Control Buttons
3. Working Channel
4. Irrigation Port
5. Leak Test Valve
6. Strain Reliefs
7. Flexible Shaft
8. Deflection Section
9. Insertion (Connector)
10. Card Edge Printed Circuit Board Plug

### 3.1 The Instrument & Accessories

**DUR-D** Invisio Flexible Ureteroscope, 3.1mm, 65cm, 3.6Fr channel

**DUR-DBA** Invisio Flexible Ureteroscope, backward articulating, 3.1mm, 65cm, 3.6Fr channel

**DUR-D and DUR-DBA are compatible with IDC-1000 and IDC-1500 controllers. For appropriate controller software upgrades, contact ACMI Customer Service or your local ACMI representative.**

**Accessories**

- Remote Printer Cable, Sony Printers: MV-10350
- Remote Printer Cable, Toshiba Printers: MV-10351
- BNC Cables, 4 ft. (2/pkg): MV-14024
- BNC Cables, 12 ft. (2/pkg): MV-9144
- BNC Cables, 40 ft. (2/pkg): MV-9144-40
- Y/C (S-Video) Cables, 4 ft. (2/pkg): 103017-1
- Y/C (S-Video) Cables, 12 ft. (2/pkg): MV-9162
- Y/C (S-Video) Cables, 30 ft. (2/pkg): MV-9162-30
- RGB (Component) Cable, 6 ft.: IDC-0110
- VGA (Progressive Scan) Cable, 6 ft.: IDC-0100
- Power Cables, 4 ft., 120 V (2/pkg): MV-10428
- Connector Maintenance Kit, small: MV-10454
- Connector Maintenance Kit, large: MV-10455
3.2 Technical Specifications
(for reference only—typical measurements/performance)

**DUR-D and DUR-DBA Invisio Flexible Ureteroscopes (typical)**

- **Working Length:** 65cm
- **Distal Tip:** 2.7mm (8.7Fr)
- **Shaft:** 3.1mm (9.3Fr)
- **Working Channel:** 1.3mm (3.6Fr)
- **Active Deflection - Primary:** 250° up, 250° down
- **Leak Test Valve:** Yes
- **AUTOSEAL**
  (no vent valve for sterilization): Yes
- **Minimum Bend Radius:** 8.0 mm
- **Field of View**: 80°
- **Focus:** Automatic 2–40 mm
- **Direction of View:** 9° (towards working channel)

* The DUR-D objective head features a removable lens cell design. A high-magnification objective head may be special ordered.

3.2.1 Safety Specifications

This equipment meets the Safety Requirements of—

- IEC 60601:1995
- IEC 60601-1-2:2004
- UL 60601-1:2003
- CSA C22.2.125:1999

**Type Construction:** Portable Equipment

**Protection Against Electric Shock Hazards—**
- **Class of Protection:** Class 1
- **Degree of Protection:** Type BF

**Protection Against the Ingress of Water:** Ordinary

**Rated Operation:** Continuous Operation

**Other Safety Information:**

Equipment is not for use in presence of flammable anesthetics (gases) or detergents.

**NOTE:** Use of electrically active surgical devices in the working channel may increase the patient leakage currents.

It is important that the user perform an operational and functional test of the instrument(s) prior to each use. If an irregularity or substandard function is suspected, the instrument(s) should not be used. Notify ACMI Customer Service immediately.
Section 4.0  Operation

4.1  Inspection of the Instrument

Perform the following steps before using the instrument—do not use if damage is found.

1. Visually inspect, and feel with fingertips, the entire surface of the shaft (including the distal tip) for dents, protrusions, holes, or other irregularities. Examine the instrument in all deflection modes: non-deflected (straight), fully deflected up, and fully deflected down.

   CAUTION: Never force the flexible tip into a straight or flexed position while holding the deflection control—this can damage the control mechanisms.

2. Ensure that irrigant flows freely and that accessories pass easily through the instrument working channel.

4.2  Optical Field Inspection

Clean all lens surfaces, including the optical window, with a gauze pad or cotton swab moistened with 70% isopropyl alcohol. Do not use if image on the monitor remains distorted or cloudy after cleaning. While supporting the distal tip, position the instrument on an object 8–10 mm away. The image on the monitor should be crisp and clear.

Refer to ACMI document Quality Check of Flexible and Semi-Rigid Endoscopes (PI136) for additional information.

4.3  Deflection Controls

Hold the instrument with the shaft extended downward with the irrigation port pointing upward; place the thumb on the deflection control lever. When the deflection lever is in the middle (or neutral) position (1) the tip will be straight. Logical primary deflection moves the tip up 250° when the lever is moved up or pushed back (2) and down 250° when the lever is moved down or pushed forward (3).

   CAUTION: Never bend the deflection sections of the shaft by the fingers as this can damage the deflecting mechanism.

NOTE: The DUR-DBA’s deflection control is international-style: pushing the deflection lever forward articulates the distal tip “up” and pushing the lever back articulates the distal tip “down” (opposite to the logical DUR-D deflection).
4.4 Leakage Tester

The ACMI Leakage Tester enables the surgical staff to easily verify the integrity of the flexible endoscope’s working channel.

Note: The leakage tester connector and air vent must both be completely dry.

1. Secure the leakage tester connector fitting (1) to the special port (2) located on the proximal end of the endoscope. Rotate the leakage tester connector so as to lock in place. Proper connection will require alignment of the leakage tester connector with the pins located on the air vent and clockwise rotation of the leakage tester connector.

2. Pressurize the scope by pumping the hand bulb (3) until the indicator on the gauge is in the range of 140–200 mmHg. DO NOT overpressurize as this may cause serious damage to the scope. Several squeezes are required to achieve the test pressure range. Achieving the test pressure with a single squeeze may indicate a valve problem.

3. Monitor the gauge on the leakage tester to determine if the indicator remains in the range of 140–200 mmHg. If the indicator drops from this region rapidly, then a major leak has been detected.

4. Release the pressure release valve (4) for a minimum of 20 seconds on the leakage tester. Engage the pressure release valve to verify the gauge indicates ‘0’ (or below 40 mmHg). Disconnect the leakage tester from the scope.

If a leak is detected, or the leak tester does not pressurize and behave as though it is pressurizing the device properly, disconnect the leakage tester connector and verify that it and the air vent are completely clear and dry. Reattach and repeat steps 1, 2, and 3 above to confirm whether a leak is present.

If scope leakage or problems are confirmed, DO NOT use endoscope. Contact ACMI customer service immediately.

4.5 AUTOSEAL™ Feature

The instrument is equipped with AUTOSEAL technology. The AUTOSEAL feature eliminates the need to attach a venting valve, which allows for ventilation of ethylene oxide gas (EO) from the instrument after sterilization.

4.6 Rotating Working Channel/Irrigation Port

The working channel and irrigation port each rotate ±45°; both have Luer lock fittings.

An adjustable working channel seal (ref ABP) can be attached by turning clockwise and pushing down, and detached by turning counterclockwise and pulling up. The ABP will accept working instruments up to 3 Fr in diameter.

A standard, floppy tip .038” guidewire may be used to check the integrity of the channel. If an accessory does not pass easily even though the guidewire passes freely, the accessory may be too large. If the guidewire does not pass freely, contact ACMI Customer Service.
4.7 Strain Relief
The external strain relief covers the proximal portion of the shaft. The strain relief protects the shaft during use and storage. Do not bend the shaft sharply as light fiber breakage and/or shaft damage can result.

4.8 Cable Compensation
The instrument is equipped with a cable compensating mechanism to guard the pull cables against excessive loads.

Section 5.0 Procedure

1. Attach the Irri-Flo™ Tubing (ref 5920000) (or similar tubing) to the irrigation port. Attach a 30 cc sterile syringe to the three-way stopcock to provide controlled irrigation, or attach a self-sealing biopsy seal to the accessory port (see Section 4.6 Rotating Working Channel/Irrigation Port) for insertion of appropriate-size accessories.

2. When transporting the instrument, support the tip of the shaft with one hand and the proximal housing with the other to prevent damage to the shaft.

3. Accessories cannot be inserted when the tip is deflected; attempting to do so may perforate the working channel.

4. When the endoscope is straight, less resistance is noted. When the tip is deflected, resistance increases as the angle of deflection increases. Attempts to forcibly insert or withdraw accessory devices can damage the instrument channel or accessory.

5. If the scope is deflected, it may be necessary to withdraw it slightly to straighten the tip, advance the accessory to the distal tip, reposition the scope, and then slowly advance the accessory.

IMPORTANT: To introduce an accessory (1) into the working channel with seal (2), grasp the accessory shaft close to the working channel entry and advance slowly, using repeated short strokes. If resistance is encountered, slowly withdraw the accessory several millimeters, then continue introduction. DO NOT FORCE ACCESSORY—see Section 6.1 for more information.

CAUTION: Before attempting to pass instruments or devices through the working channel of a flexible endoscope, the entire working channel must be as straight as possible, with no tip deflection.

The 9º direction of view will allow the device to enter the field of view close to the tip. It is recommended that the device then be pulled just out of the field of view before further endoscope manipulation. This will assure easy advancement of the device once the endoscope has arrived at the pathology. Attempting to pass instruments or devices through a deflected flexible endoscope will cause damage to the working channel. This damage can result in leakage, requiring repair of your flexible endoscope. ACMI’s Limited Express Warranty does not cover such damage.

ACMI recommends a 270 micron laser fiber for maximum deflection and access to the kidney. The superior deflection capability of ACMI flexible endoscopes can exceed the minimum bend radius of larger fibers. A larger fiber may break and be accidentally fired inside the working channel, damaging the endoscope.

CAUTION: Sharp accessories, such as laser fibers and EHL probes, can cause damage to the working channel if passed when the scope is deflected—pass these instruments only when the scope is not deflected (straight).

NOTE: As irrigation flow diminishes with an accessory in the channel, pressurized irrigation may be necessary.

WARNING: Before removing the endoscope from the patient, be sure the shaft is straight (not deflected). Patient injury and possible damage to the instrument could result.

Immediately flush the working channel with sterile distilled water using a syringe to remove any blood or mucus.
5.1 Working Channel

IMPORTANT: When the endoscope is straight, less resistance is noted. When the tip is deflected, resistance increases as the angle of deflection increases. Attempts to force entry or withdrawal of accessory devices—especially sharp devices such as laser fibers and electrohydraulic lithotripsy (EHL) probes—can cause damage to the working channel or accessory.

Pass all accessories all the way through the endoscope before deflecting the tip whenever possible.

5.2 White Balancing

The DUR-Digital Invisio Flexible Ureteroscope does not require white balancing as it is set at the factory.

5.3 Focusing

The DUR-Digital Invisio Flexible Ureteroscope requires no focusing.

5.4 Remote Head Button Setup

The DUR-Digital Invisio Flexible Ureteroscope is equipped with three remote head buttons on the top of the hand piece to control DUR-D System features and peripheral devices. The middle button (2) is dedicated to brightness, while the distal and proximal buttons (1, 3) are programmable through the On-Screen-Display (OSD) system (see Sections 5.5 and 5.6).

5.5 On-Screen-Display (OSD) System

The OSD enables the configuration of system functions, as well as the customization of the remote button functions to a specific preference.

To access the OSD (available anytime during the procedure when the DUR-D is in Active Mode):
- Depress and hold the distal and proximal remote buttons for two seconds.

To configure the remote buttons and system settings:
1. Depress the proximal button to display options, cycling through the various choices.
2. Depress the distal button to select a specific option.
3. No other buttons are active during programming.
4. The DUR-D Invisio System will remember the last confirmed remote button settings when the power is disconnected from the IDC Invisio Digital Controller.
5.6 OSD Options and Navigation

1. Remote Button (a) – proximally located
   a. Control (select one)—the system will display the available choices for the distally located remote button to control: Enhancement, Accessory 1, Accessory 2, or Zoom.

2. Remote Button (b) – middle button
   a. This button is dedicated to Brightness.
      - Allows the surgeon to increase the maximum overall brightness of the image.
      - The control will increase the image brightness if there is sufficient reserve.
      - The image brightness will increase from level 1 to 5 and then back to 1 in this mode.

3. Remote Button (c) – distally located
   a. Control (select one)—the system will display the available choices for the proximally located remote button to control: Enhancement, Accessory 1, Accessory 2, or Zoom.

4. Remote Button Choices (Options)
   a. Enhancement
      - Allows the surgeon to increase the enhancement of the image.
      - The image will be enhanced from level 1 to 6 and then back to 1 in this mode.
      - Level 6 is the Laser Mode and is recommended for use with a Holmium YAG laser.

5. Dual Accessory Control
   a. A VCR, digital capture device, or video printer can be controlled when a remote button is assigned to Accessory 1 or Accessory 2.
   b. VCRs will RECORD/STOP with each button press.
   c. Video Printers/Digital Capture Devices will CAPTURE/PRINT with each button press. These devices must be set correctly according to their manufacturers instructions.
   d. Connect the accessory cable (#1) from the rear of the IDC Invisio Digital Controller to a device.
   e. Connect the accessory cable (#2) from the rear of the IDC Invisio Digital Controller to a device.
   f. The remote buttons, if configured, will now control or trigger a peripheral device.

6. DUR-D, DUR-DBA Pre-Set Buttons and Levels
   a. Enhancement – Top Button (end of scope)
      The IDC Invisio Digital Controller utilizes proprietary digital image processing to increase image details. The digital enhance setting allows the user to see fine details that were previously indistinguishable from their surroundings.
      - Enables the base level of enhancement to be set from 1 to 6 and back to 1.
      - Level 6 is the Laser Mode and is recommended for use with a Holmium YAG laser. (Note: on some lasers, best results are with aiming beam turned to a lower setting.)
      - Factory setting from ACMI is Laser Mode.
      - Enhancement level can be adjusted from the remote buttons, if configured, or via the front panel of the IDC Invisio Digital Controller.
   b. Brightness – Square, Center Button
      The IDC Invisio Digital Controller utilizes proprietary digital image processing to provide a bright image across the entire visual field.
      - Enables the base level of brightness to be set from 1 to 5 and back to 1.
      - Factory setting from ACMI is level 3.
      - Brightness level can be adjusted from the remote buttons, if configured, or via the front panel of the IDC Invisio Digital Controller.
   c. Zoom – Bottom Button (closest to deflection trigger)
      The IDC Invisio Digital Controller utilizes proprietary image processing to digitally zoom or enlarge the image size on the monitor. This makes it easier to see details during a procedure.
      - Enables base level of zoom to be set from 65% super sharp to 135% super large in 5 steps.
      - Factory setting is 100%, Zoom 3.
      - Zoom level can be adjusted from the remote buttons, if configured.
5.7 Inspection
Before using your instrument, inspect the endoscope, control unit, and cabling (including connectors). A malfunction of any of these components can degrade the video image.

DUR-Digital Invisio Flexible Ureteroscope Assembly
1. Thoroughly dry the connector prior to insertion into the IDC Invisio Digital Controller.
2. Confirm that the optical window on the distal end of the instrument is clean and free of condensation.
3. Check the cable connector for signs of moisture, contamination, or physical damage (e.g., broken housing or chipped circuit board).

System
1. Plug the instrument into the receptacle on the IDC Invisio Digital Controller.
2. Connect video output signal(s) to the monitor (see IDC Invisio Digital Controller Operations and Technical Manual).
3. Point the instrument at a nearby object to verify that you are obtaining a clear, high quality, color image on the video monitor.
4. Verify that the instrument will focus on objects at close range (5 mm) as well as further away (50 mm) from the endoscope. Check that the image brightness on the monitor remains reasonably constant. If not, see Section 6.2 Troubleshooting: Video Image.

Monitors
1. The instrument and monitor should be electrically isolated from electrocautery and other electrical medical equipment.
2. Verify that the electrical equipment is safely grounded. Test your electrical sockets for proper grounding and check that all plugs contain a ground prong.
3. Check that the monitor termination switch, if present, is set to 75 ohm ON. (If no number is shown, switch the monitor termination switch back and forth until you obtain the best picture.) If two or more monitors are being used, only terminate the last connected monitor.
4. Make sure all of the cable connections are correct (see IDC Invisio Digital Controller Operations and Technical Manual).

Section 6.0 Troubleshooting Guide
6.1 Troubleshooting the Instrument
1. Accessory does not pass smoothly.
   a) Accessory is bent or kinked; tip is sharp.
      - Use new accessory.
   b) Working channel is obstructed.
      - Thoroughly clean channel of debris.
   c) Instrument shaft is damaged.
      - Inspect entire shaft for indentations, kinks, or other unusual wear that may indicate internal damage. If damaged, return to ACMI for repair.

2. Accessory cannot be inserted.
   a) Accessory is too large.
      - Be sure accessory is appropriate size for channel. See Section 3.2.
   b) Accessory is “hung up” on working channel diaphragm.
      - Push through seal gently with short strokes.
   c) Accessory jaws are not closed completely.
      - Close accessory jaws before insertion.
3. Accessory cannot be passed while instrument is deflected.
   a) Be sure tip is not deflected.
      - Gradually withdraw scope to straighten tip until accessory passes.

4. Accessories do not operate smoothly.
   a) Inspect accessory.
      - Thoroughly clean; or, if damaged, discard and replace.

5. Articulation is not smooth or does not operate at all.
   a) Listen for grinding noise when activating the deflection mechanism(s). If grinding noise is heard, contact ACMI Customer Service for a replacement or to arrange for service.

6. Remote buttons are not activating a peripheral device.
   a) Check OSD configuration (see Sections 5.4, 5.5, and 5.6 for more information).
   b) Check remote accessory cable connections from the IDC Invisio Digital Controller to the peripheral device.
   c) Check that the peripheral device is configured correctly to handle the signal from the IDC Invisio Digital Controller (see manufacturer’s instructions).

7. Remote buttons are not activating a system setting (i.e., Brightness or Enhancement)
   a) Check OSD configuration (see Sections 5.4, 5.5, and 5.6 for more information).

6.2 Troubleshooting: Video Image

1. Picture is grainy; image “whites out” easily when scope is close to tissue. Colors are slightly off.
   a) Un-terminated monitor.
      - Set termination switch at back of monitor to 75 ohm ON or termination position.

2. Periphery of image is cloudy or hazy. Colors are washed out. Spots appear on screen.
   a) Dirty optics.
      - Clean the distal tip of the DUR-D Invisio with a cotton swab and alcohol.
      - Apply anti-fog agent if desired.
   b) Leaking, nicked, or damaged scope.
      - Inspect distal end of scope optics.

3. Picture is dark, grainy.
   a) Insufficient light reaching subject.
      - Inspect the distal tip of the instrument for dirt or other foreign materials on the outer lens (see Section 4.2).
   b) Improperly adjusted or double-terminated monitor or video printer. - See monitor’s instruction manual and/or Section 5.7.
   c) Increase brightness on front of controller or from the remote buttons on the instrument.

4. No picture generated.
   a) Broken video cable.
      - Check that connecting ends of all cables are intact. Look for slices or other visual damage to the cable. Replace cable as appropriate.
      - Direct connect a cable from the IDC Invisio Digital Controller to the input of the monitor.
   b) Lack of power to a system component.
      - Ensure that all systems are plugged in and all accessories are switched to ON.
   c) Off-line VCR or other peripheral device.
      - Set VCR input switch to “line” position and/or refer to your VCR’s instruction manual.
   d) Other components not in proper channel (e.g., the monitor may be in line “B” while hookup is in line “K”).
      - Connect components correctly.
5. Picture color is incorrect.
   a) Improperly adjusted color.
      - Check the monitor’s chroma and phase settings.
      - Check the printer’s color and phase settings.

6. Picture is too dark.
   a) Insufficient brightness.
      - Increase brightness on front of controller or from the remote buttons on the instrument.
   b) Damaged instrument.
      - Check condition of instrument.
   c) Incorrect contrast or brightness settings on monitor or printer.
      - Adjust contrast and/or brightness on monitor and/or printer.

7. Picture is too bright.
   a) Excessive light intensity.
      - Decrease brightness on front of controller or from the remote buttons on the instrument.
   b) Un-terminated monitor or secondary monitor cable.
      - Set termination switch at back of monitor to 75 ohm ON or termination position.

8. Noise or “snow” appears on screen when no electrocauter y is in use.
   a) Excessive enhancement.
      - Decrease enhancement level to low or off.
   b) Faulty video cables.
      - Check video cables and replace as necessary.

9. Noise or “snow” appears on screen when using electrocauter y.
   a) Dirty contacts in instrument receptacle (front panel).
      - Clean contacts with ACMI Connector Maintenance Kit (ref MV-10454 or MV-10455).
   b) Interference from electrocautery unit.
      - Plug electrocautery equipment into separate electrical outlet.
   c) Interference from electrocautery cable.
      - Separate video cable from electrocautery cable.
   d) Poor video cable connection(s).
      - Check all video cables for good condition and secure connection.

10. Picture is intermittent.
    a) Wet instrument connector.
        - Check cable-to-IDC Invisio Digital Controller connection.
        - Shake and air-dry connection.
    b) Dirty instrument connector contacts
        - Clean contacts with ACMI Connector Maintenance Kit (REF MV-10454 or MV-10455).
    c) Failing instrument cable.
        - Contact ACMI Customer Service.

11. Intermittent picture accompanied by green or unusual coloring.
    a) Wet instrument connector.
        - Check cable-to-IDC Invisio Digital Controller connection. Shake and air-dry connection.
    b) Bad instrument cable.
        - Contact ACMI Customer Service.

• For all other problems or when a recommended corrective action does not correct the identified problem, contact ACMI Customer Service at (888) 524-7266.
Section 7.0 General Care

Like many precision surgical devices, the instrument and IDC Invisio Digital Controller are delicate and should be handled with care. Follow the suggestions below to prolong your system’s life and avoid repairs:

- Never remove the instrument cable from the IDC Invisio Digital Controller by pulling on the cable itself—always pull from the connector body to remove.
- Coil the instrument cable loosely—never sharply bend, kink, strain, or crush the cable. Exercise care when handling the connector at the end of the instrument cable. Damage may occur if the connector is dropped or forced into the receptacle.
- Keep unprotected instruments, cables, and IDC Invisio Digital Controllers separate from one another. Do not stack system components.
- Pick up and carry instruments or system components separately, not in groups.
- Never insert a wet cable connector into the IDC Invisio Digital Controller receptacle.
- Use ACMI Connector Maintenance Kits (REF MV-10454, MV-10455) for proper care of connector and cable receptacle.

Section 8.0 Maintenance

The following cleaning, disinfection, and sterilization procedures are recommended by ACMI. Use of any procedures not expressly recommended by ACMI may adversely affect or damage ACMI devices and may void ACMI’s Limited Express Warranty.

Follow all applicable bloodborne pathogen procedures as indicated by OSHA and/or your hospital requirements when cleaning, disinfecting, and sterilizing instruments and accessories.

WARNING: Always turn off the power on the IDC Invisio Digital Controller and disconnect the instrument before cleaning.

8.1 Cleaning

Clean the instrument and optics after each use with an enzymatic cleaner formulated to dissolve proteinaceous material.

1. Separate any assembled components.
2. Rinse the scope with warm (38–49°C) water on all exterior surfaces.
3. Flush the working channel with warm (38–49°C) water using a syringe.
4. Prepare enzymatic detergent solution per the manufacturer’s written instructions.
5. Immerse the scope in the enzymatic solution. Wash the exterior of the scope with the enzymatic detergent solution and a soft brush.
6. Allow the scope to soak in the enzymatic solution for 10 minutes.
7. Brush the exterior of the scope thoroughly in the enzymatic solution. Pay special attention to the area around the deflection handle and the function buttons.
8. Clean the interior working channel with enzymatic detergent solution using a working channel cleaning brush (ACMI REF CB-3115: 3 Fr, 115 cm long). Ensure both working channel and inlets are cleaned through to the distal end of the scope.
9. Flush each working channel inlet with a minimum of 50 cc of enzymatic detergent solution using a syringe.
10. Rinse the exterior of the scope by immersing it in warm water.
11. Rinse each inlet with 50 mL minimum of warm water.
12. Purge air through each inlet using a 50 cc syringe.
13. Repeat steps 6–11 at least two times.
14. Clean the lens with a cotton swab moistened with 70% isopropyl alcohol. Avoid scratching the lens with a hard object.
15. Dry the exterior of scope using a sterile, lint-free towel or air jet.
8.2 Cleaning the Optics on the Instrument
CAUTION: Do not attempt to open the DUR-Digital Invisio Flexible Ureteroscope. Opening these sealed assemblies affects the DUR-D’s moisture seals and will void your warranty.

8.3 Cleaning the Instrument Connector and Receptacle
ACMI recommends the use of the Connector Maintenance Kits (REF MV-10454, MV-10455). Follow the cleaning instructions provided with these kits.

8.4 Cleaning the IDC Invisio Digital Controller
CAUTION: Never immerse the IDC Invisio Digital Controller in any solution.
If dirt or liquid gets on the IDC Invisio Digital Controller, wipe with a damp cloth or sponge. Dry with a towel after cleaning it. Depending on use, clean at least once a week.

8.5 Disinfection
1. Keep all components disassembled.
2. After cleaning and inspection, completely immerse the instrument and cable in a 2.4% activated alkaline glutaraldehyde solution in a plastic or enamel basin while keeping the shaft in a soft curl.
3. Fill the working channel with the solution, using a syringe.
4. Soak for a minimum of 45 minutes (not to exceed 60 minutes).
5. Thoroughly rinse the outside of the instrument and the cable and flush the working channel three times with sterile water, using a syringe.
6. Wipe the instrument including cable and cable connector paddle dry using a sterile towel. The cable connector paddle must be completely dry before connecting to the controller.
7. DO NOT DISINFECT THE IDC DIGITAL CONTROLLER.

8.6 Autoclave Sterilization
CAUTION: Never steam sterilize the instrument or IDC Digital Controller. Exposing the instrument or IDC Digital Controller to temperatures greater than 145°F (63°C) will cause irreparable damage.

8.7 Ethylene Oxide Sterilization
The instrument can be sterilized in an ethylene oxide (EO) sterilizer.
1. Thoroughly clean, rinse, and dry prior to sterilization.
2. The instrument can be placed in an ethylene oxide (EO) sterilizer controlled within the following parameters:

<table>
<thead>
<tr>
<th>EO (100%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp °F (°C): 131 (55)</td>
</tr>
<tr>
<td>Rel. Humidity: &gt; 35%</td>
</tr>
<tr>
<td>Vacuum: 2–3 psia</td>
</tr>
<tr>
<td>Preconditioning Time: ≥ 60 minutes</td>
</tr>
<tr>
<td>Gas Conc.: 725–750 mg/l</td>
</tr>
<tr>
<td>Exposure: ≥ 60 minutes</td>
</tr>
</tbody>
</table>
3. After EO exposure, aerate for a minimum of 12 hours at 122°F (50°C) in a mechanical aerator.
8.8 STERIS SYSTEM 1™ Sterilization
The instrument is compatible with the STERIS SYSTEM 1 when used according to the STERIS written instructions. It is important to use the C1140 Covered Flexible Processing Tray/Container and Quick Connect QCC1671 to process the instrument.

8.9 STERRAD® Sterilization
In countries where STERRAD is approved for use with flexible endoscopes, the DUR-Digital Invisio Flexible Ureteroscope has been validated for both sterilization and material compatibility. Sterrad NX is an approved and validated process for use with the DUR-D.

Section 9.0 Storage
To pack the instrument system components for storage, disconnect the IDC Invisio Digital Controller from the instrument and the monitor.
Separate the power cable from the IDC Invisio Digital Controller, coil it loosely, and place it into the storage container. Store the system and components in a clean, dry, well-ventilated environment.
Take special care to provide the instrument assembly with an extra level of protection during storage; the distal tip, shaft, cable, and connector are fragile. Never strain, crush, kink, or sharply bend the power cable, the instrument cable, or shaft. Exercise care when handling the connector at the end of the instrument cable—damage may occur if the connector is dropped.
Store the instrument in a protected vented tray; do not store in shipping case. Be certain instruments are completely dry prior to storage. Keep instruments separate from one another to avoid damage. Keep the shaft in the non-deflected (straight) position when storing.

Section 10.0 Theory of Operation
The system consists of two major components: the instrument and the IDC Invisio Digital Controller. The DUR-Digital Invisio Flexible Ureteroscope contains a single image sensor placed within the distal tip of the endoscope upon which a scene is imaged. The image sensor consists of an active area that converts light into electrical charges that form the basis for generating an image. The parts of the active area are often referred to as "pixels," short for "picture elements."
Refer to IDC Invisio Digital Controller Operations and Technical Manual for more information.

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