INSTRUCTIONS FOR USE

ALL INSTRUCTIONS, PRECAUTIONS AND WARNINGS SHOULD BE CAREFULLY READ AND UNDERSTOOD BEFORE USE. FAILURE TO DO SO MAY RESULT IN COMPLICATIONS.

Caution – Federal (USA) law restricts this device to sale by or on the order of a physician.

Indication for Use
To access and treat the maxillary ostia/ethmoid infundibula in patients 2 years and older, and frontal ostia/recesses and sphenoid sinus ostia in patients 12 years and older using a trans-nasal approach. The bony sinus outflow tracts are remodeled by balloon displacement of adjacent bone and paranasal sinus structures.

Description
The XprESS Multi-Sinus Dilation System is intended to remodel or recreate the sinus outflow tract via trans-nasal balloon dilation. The XprESS device combines features of a curved suction tip and a frontal ostium seeker with the tissue expansion effect of balloon dilation. The familiar features of this device enable a physician to track the device to the sinus ostium. Since the distal end of the device is re-shapeable, one balloon can be modified to work on multiple sinuses within the same patient.

The XprESS device curved suction tip has an atraumatic ball tip. A suction tube may be connected to the proximal barbed fitting to provide active suction by covering the suction vent. An Extension Line connected to a syringe may be connected to the proximal barbed fitting to provide irrigation. The device was designed to prevent fluid from exiting the suction vent during irrigation. The XprESS Multi-Sinus Dilation System is provided sterile and for single use only.

The XprESS Multi-Sinus Dilation System includes the XprESS device, Inflation Syringe, Bending Tool and two Extension Lines. The XprESS LoProfile and Ultra Multi-Sinus Dilation Systems also include the PathAssist LED Light Fiber. The XprESS Pro Multi-Sinus Dilation System also includes a Tuchoy Adapter.

XprESS is available in the following suction tip sizes and balloon sizes. All suction tips and balloon lengths are appropriate for treating all sinuses; selection is based on physician preference.

<table>
<thead>
<tr>
<th>XprESS Pro</th>
<th>XprESS LoProfile</th>
<th>XprESS Ultra</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard Suction Tip (2mm ball tip, 1mm ID, 1.5mm OD)</td>
<td>LoProfile Suction Tip (1.75mm ball tip, 0.7mm ID, 1.2mm OD)</td>
<td>Ultra Suction Tip (1.5mm ball tip, 0.5mm ID, 1.0mm OD)</td>
</tr>
<tr>
<td>Balloon Diameter x Length (mm)</td>
<td>Balloon Diameter x Length (mm)</td>
<td>Balloon Diameter x Length (mm)</td>
</tr>
<tr>
<td>NA</td>
<td>5 x 8</td>
<td>5 x 8</td>
</tr>
<tr>
<td>NA</td>
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<td>6 x 20</td>
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</tr>
<tr>
<td>7 x 18</td>
<td>7 x 20</td>
<td>NA</td>
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</tbody>
</table>

The XprESS Multi-Sinus Dilation System has been tested to withstand multiple inflations and device tip manipulations in a surgical case wherein all 6 sinus ostia are being dilated.

Contraindications
• None known

Warnings
• Never advance or withdraw the XprESS device against any resistance. Do not use excessive force or torque to advance the XprESS device or balloon / slide assembly when positioned in any paranasal space. Such actions could lead to tissue trauma, bleeding, or device damage.
• Do not use breached or damaged packages, since the sterility and functionality of the device may be compromised.
• The XprESS Multi-Sinus Dilation System is provided sterile and intended for single procedure use only. Do not re-sterilize and/or re-use, as it may result in compromised device performance and risk improper sterilization and cross contamination.
• Do not use XprESS device in patients with known allergies to barium sulfate.
• Due to the variability of sinus anatomy, review appropriate radiographic imaging (CT scan) prior to treatment. Do not use the XprESS device to treat a hypoplastic/atelectatic maxillary sinus or atelectatic ethmoid infundibulum.
• Due to the variability of sinus development in pediatric patients, review CT scan to assess each sinus’s development and appropriateness for balloon dilation. Pneumatization may occur as early as 1-2 years of age and continues to develop throughout childhood. Do not use XprESS in a sinus that is not adequately developed.
• Do not exceed the maximum recommended balloon inflation pressure of 12 atm. Over-inflation of sinus balloons can result in serious adverse events.
• Do not use ionic or non-ionic fluoroscopic contrast solution to inflate the balloon in patients with known allergies to contrast media.
• If suction through the XprESS device lumen is used during the procedure, temporarily discontinue suction (remove finger from suction vent, disconnect suction hose from device, or clamp suction hose) at the time of balloon inflation. Suction can resume subsequent to balloon deflation. Using the XprESS device in suction mode while balloon is inflated may result in barometric trauma to sinus tissue which may lead to increased bleeding.
• As in any upper airway procedure or sinus surgery, do not use CPAP until the physician has confirmed that the tissue is adequately healed. CPAP usage prior to soft tissue healing may result in facial and/or neck swelling due to subcutaneous emphysema.
• Do not clean the XprESS device with anti-microbial agents as the compatibility of the XprESS device with these agents has not been tested.
• The XprESS device has been tested only with the iagion Navigation System. Do not attach the XprESS device to other image guidance systems. This may result in inaccurate device positioning. Refer to System Operation 1.b for instructions on how to connect XprESS to the iagion system.
• The XprESS device has been tested only with the Entellus Inflation Syringe. Do not use other inflation devices. Use of other inflation devices may result in serious patient injury.
Precautions
- Store the XprESS device components in a cool and dry place. Never use a device that is beyond its expiration date.
- Handle the XprESS device with care. Prior to use, and during the procedure, inspect the packaging and components for bends, kinks, or other damage. Discontinue the use of the XprESS device if it may have been damaged.
- Select a balloon diameter that will result in expansion of the tissue post-dilation. Do not select a balloon diameter that is larger than the bony margins of the outflow tract as this may damage the balloon.
- Pay special attention when advancing or withdrawing the balloon and slide assembly. If resistance is encountered, use endoscopy or direct visualization to help guide device out of the paranasal space and then attempt to alleviate the resistance. If the cause of resistance cannot be determined, do not use the XprESS device.
- Use direct endoscope visualization with or without PathAssist LED Light Fiber or Light Fiber to ensure accurate placement of the balloon prior to dilation. If balloon location cannot be verified image guidance or fluoroscopy can be used. If balloon location still cannot be verified, the balloon should not be inflated.
- Consider using self-limiting radiation exposure equipment when employing fluoroscopy to confirm device placement. Ensure the equipment is calibrated and maintained per the equipment manufacturer’s user manual.
- Use techniques for reducing fluoroscopic exposure when using fluoroscopy. Examples are applying pulsed beam settings, increasing target-to-panel distance, utilizing posterior-anterior projection, or using appropriate lead shield protection. Total fluoroscopy time should be limited to 30 minutes.
- When fluoroscopy is used, especially in children, minimize radiation dose to the lens of the eye and other proliferating tissues due to the potential of cataract formation or injury to the surrounding tissue.
- Do not advance or withdraw a guidewire through the XprESS Pro or LoProfile suction/irrigation lumen against resistance. This could lead to device damage.
- Be aware that guidewires (including flagon GuideWires) do not track through the XprESS Pro or LoProfile when they are bent in the recommended maxillary configuration, nor the XprESS Ultra in any configuration. Other methods can be used to obtain confirmation of the treatment area such as use of the PathAssist Light Fiber, direct visualization of the XprESS device with an aid of an endoscope or fluoroscopic imaging of the XprESS tip.
- Use standard larger suction tubes for removal of thick secretions or other materials. XprESS Pro has a 1mm ID comparable to that of a 5F suction tube. XprESS LoProfile has a 0.7mm ID comparable to that of a 4F suction tube. XprESS Ultra has a 0.5mm ID comparable to that of a 2.5F suction tube. All are capable of removing blood and thin mucous.
- Fully deflate the balloon and retract the balloon slide assembly before withdrawing the XprESS device from the paranasal sinus space.
- Use only liquid contrast or saline solution for inflation. Do not inflate with air.
- Consider using a new balloon if cross-contamination between sinuses is a concern.

Adverse Effects
Possible adverse effects include, but are not limited to, the following:
- Complication from anesthesia
- Damage to the lamina papyracea
- Damage of the orbital wall or other structures of the eye
- Cerebrospinal fluid leak
- Loss of vision or diplopia (double vision)
- Temporary or permanent facial / nasal pain
- Epistaxis
- cavernous sinus syndrome
- Damage to the lacrimal sac affecting tearing
- Pneumocephalus
- Facial bruising and swelling
- Tissue inflammation
- Fever and infection
- Continued or worsening sinus symptoms
- Revision surgery

Supplies
The following supplies are not provided with the XprESS Multi-Sinus Dilation System and should be available and prepped prior to use of the device.
- Appropriate endoscopes and compatible camera system
- ≥ 50 cc of sterile saline solution or sterile fluoroscopic contrast solution or sterile water
- Needles and syringes as required for injections
- 20-30 cc syringe and Extension Line (if irrigation is to be performed)
- Suction system
- Other supplies or medication as per established laboratory protocol
- If the use of a sterile guidewire is desired (compatible with the XprESS Pro), the recommended guide wire should be sterile and ≤ 0.035 inches in diameter with a minimum length of 50 cm. Example of a guidewire that meets these requirements is the Entellus Medical Sinus Guidewire.
- If desired, Entellus Medical PathAssist™ LED Light Fiber or Light Fiber™ or Light Seeker

Optional Equipment
- flagon Navigation System and GuideWires (GuideWire and GuideWire 0.6 are compatible with XprESS Pro; GuideWire 0.6 is compatible with XprESS LoProfile)

Instructions for Use
System Preparation
1. Prepare the Inflation Syringe and Extension Line
   a. Remove the Inflation Syringe and Extension Line from its sterile package.

Note the 3 referenced Inflation Syringe plunger positions:

**Figure 2 - Plunger all the way in**
- b. Begin with the Inflation Syringe plunger all the way in (Figure 2).
- c. Then submerge tip in sterile saline solution.

**Figure 3 - First Click position**
- d. Fill Inflation Syringe by slowly drawing plunger back to second click position (all the way out) (Figure 4).

**Figure 4 -Second Click position (all the way out)**
- e. Attach an Extension Line to the filled Inflation Syringe.
1. Locate the sinus structure using one of the following confirmation methods:

- Direct Visualization with or without Light Confirmation: Locate the treatment area or sinus structure using XprESS with LED Light Fiber or Light Fiber. Light Seeker, a standard sinus ostium seeker and/or guidewire with the aid of an endoscope. Observe the location of the treatment area relative to the anatomical landmarks through the endoscope. Remove the Light Seeker, sinus ostium seeker or guidewire after locating treatment area.

  Note: If using the PathAssist LED Light Fiber or Light Fiber, refer to the Instructions for Use (IFU) for complete instructions.

- CT Image Guidance: If further confirmation of the treatment area location is desired, CT Image guidance using the fiagon Navigation System and GuideWire or GuideWire 0.6 with XprESS Pro may be used. The fiagon Navigation System and GuideWire 0.6 with XprESS LoProfile may also be used.

  i. If using the GuideWire with XprESS Pro, attach the Tuohy Adapter to the XprESS proximal barbed fitting.

  ii. Load the fiagon GuideWire through the Tuohy Adapter and working lumen of XprESS until the tip of GuideWire aligns with the tip of XprESS.

  iii. Secure the GuideWire in place by tightening the Tuohy Adapter.

  iv. If using GuideWire 0.6 with XprESS Pro or LoProfile, load the GuideWire through the working lumen of XprESS until the luer lock connector meets the proximal barbed fitting of XprESS.

  v. Secure the luer lock connector on the proximal barbed fitting.

  vi. Refer to fiagon Navigation System Instructions for use.

  Note: Neither fiagon GuideWire should be used with any XprESS device in the maxillary bend configuration.

  Note: Do not attach the XprESS device to other image guidance systems.

- Fluoroscopy: If further confirmation of the treatment area is desired, fluoroscopy may be used. Take two orthogonal views (AP and lateral) of the sinus. The XprESS device suction tip is stainless steel and is visible under fluoroscopy. The balloon will be proximal to the tip of the device.

  Notes: Reference marks are located 1 and 2 cm from the tip of the device.

  The XprESS suction tip may be re-shaped to aid in device positioning. Use device as a suction tool to maintain a clear visual field during device positioning. Cover suction vent with finger to allow suction.

2. Under endoscopic visualization, track the XprESS device to the same treatment area identified above.

  Notes: Reference marks are located 1 and 2 cm from the tip of the device.

  The XprESS suction tip may be re-shaped to aid in device positioning. Use device as a suction tool to maintain a clear visual field during device positioning. Cover suction vent with finger to allow suction.

2. Prepare XprESS Multi-Sinus Dilation System.

a. Remove the XprESS device from its sterile package.

b. Remove and discard the balloon protector.

c. Connect the free end of the prepped Extension Line to the XprESS balloon inflation luer.

  Note: Inspect the syringe barrel to ensure there is minimal air in the system. If excessive air remains in the system, repeat the prepping process.

  Figure 5: Alignment between Distal Seal and the Distal Mark Corresponds to 12atm

- Small adjustments to the above bends may be considered to accommodate different patient anatomy.

Using Bending Tool

- The Bending Tool should be used to achieve the proper maxillary bend. The tool also provides a frontal and sphenoid bend configuration if needed.

- Maxillary Bending with Bending Tool: Before shaping the maxillary bend, the device should be close to straight as shown for a Sphenoid Bend. With the bending tool in one hand, position the ball tip into the ball holder in the bending tool (Figure 9). Place a finger at about the 2cm mark on the suction tip and use this finger to form the Maxillary Bend (Figure 10).

Patient Preparation

1. Patient preparation should be consistent with standard practice.

2. Anesthesia should be administered appropriately to allow patient tolerance.

System Operation

1. Locate the sinus structure using one of the following confirmation methods:

a. Direct Visualization with or without Light Confirmation: Locate the treatment area or sinus structure using XprESS with LED Light Fiber or Light Fiber. Light Seeker, a standard sinus ostium seeker and/or guidewire with the aid of an endoscope. Observe the location of the treatment area relative to the anatomical landmarks through the endoscope. Remove the Light Seeker, sinus ostium seeker or guidewire after locating treatment area.

  Note: If using the PathAssist LED Light Fiber or Light Fiber, refer to the Instructions for Use (IFU) for complete instructions.

b. CT Image Guidance: If further confirmation of the treatment area location is desired, CT Image guidance using the fiagon Navigation System and GuideWire or GuideWire 0.6 with XprESS Pro may be used. The fiagon Navigation System and GuideWire 0.6 with XprESS LoProfile may also be used.

  i. If using the GuideWire with XprESS Pro, attach the Tuohy Adapter to the XprESS proximal barbed fitting.

  ii. Load the fiagon GuideWire through the Tuohy Adapter and working lumen of XprESS until the tip of GuideWire aligns with the tip of XprESS.

  iii. Secure the GuideWire in place by tightening the Tuohy Adapter.

  iv. If using GuideWire 0.6 with XprESS Pro or LoProfile, load the GuideWire through the working lumen of XprESS until the luer lock connector meets the proximal barbed fitting of XprESS.

  v. Secure the luer lock connector on the proximal barbed fitting.

  vi. Refer to fiagon Navigation System Instructions for use.

  Note: Neither fiagon GuideWire should be used with any XprESS device in the maxillary bend configuration.

  Note: Do not attach the XprESS device to other image guidance systems.

- Fluoroscopy: If further confirmation of the treatment area is desired, fluoroscopy may be used. Take two orthogonal views (AP and lateral) of the sinus. The XprESS device suction tip is stainless steel and is visible under fluoroscopy. The balloon will be proximal to the tip of the device.

  Notes: Reference marks are located 1 and 2 cm from the tip of the device.

  The XprESS suction tip may be re-shaped to aid in device positioning. Use device as a suction tool to maintain a clear visual field during device positioning. Cover suction vent with finger to allow suction.

2. Under endoscopic visualization, track the XprESS device to the same treatment area identified above.

  Notes: Reference marks are located 1 and 2 cm from the tip of the device.

  The XprESS suction tip may be re-shaped to aid in device positioning. Use device as a suction tool to maintain a clear visual field during device positioning. Cover suction vent with finger to allow suction.
3. Advancing the balloon to the treatment site.
   Under endoscopic visualization, fully advance the balloon slide mechanism forward to position the balloon within the sinus opening.

4. Prior to inflating balloon, discontinue the use of suction (remove finger from suction vent, disconnect suction hose from device, or clamp suction hose) to decrease the risk of sinus barotrauma.

5. Balloon dilation of the treatment site.
   a. Slowly depress the plunger rod to inflate the balloon. The pressure should be increased slowly (3-5 seconds) until the orange piston bottoms out (distal black seal of the piston reaches the distal black mark on the Inflation Syringe – see Figure 5). If these do not align, deflate the balloon and remove the XprESS device and perform a test inflation (as described in steps 2.d and 2.e of the System Preparation section). Alignment of the distal mark and distal seal will ensure that 12 atm of pressure is reached.
   Note: Do not use air or any gaseous medium to inflate the balloon.

   b. When using the 8mm length balloon in the frontal recess, multiple inflations may be needed in order to achieve the desired result. Partially retract the balloon slide mechanism between inflations using the 5mm handle reference marks to ensure full length treatment. Typically 2-3 inflations are needed to treat the frontal recess with an 8mm length balloon. See Figure 11.

   Figure 11: Handle Marks for 8mm Length Balloon

   c. Inflate the balloon until the desired result is achieved or until it reaches 12 atm. Inflate the balloon for up to 20 seconds (less than or equal to 20 second), observe that the balloon is inflated endoscopically.
   Note: Do not exceed 12 atm.

   Warning: To avoid barometric trauma to sinus tissue, do not use device in suction mode (remove finger from suction vent, disconnect suction hose from device, or clamp suction hose) while balloon is inflated.

   d. Deflate the balloon by retracting the Inflation Syringe plunger rod to the second click position and retracting the XprESS balloon slide mechanism. Observe the results endoscopically.

   e. Perform additional inflations if needed until desired result is achieved. Typically 1-2 inflations are performed per sinus with an 18mm length balloon and 1-3 inflations per sinus with an 8mm length balloon.
   Note: To irrigate the sinus, fill a 20-30cc syringe with sterile saline. Connect the syringe to a flexible extension line and purge air. Connect Extension Line to proximal barbed fitting and flush through suction/irrigation lumen as desired. The suction vent does not need to be covered during irrigation.

6. Remove device from treatment site.
   When the sinus outflow tract has been adequately dilated, deflate the balloon (by retracting the Inflation Syringe plunger rod to the stop position and retracting the XprESS balloon slide mechanism) and remove the XprESS device from the treated sinus.

7. If necessary, clean up the ostium site by cutting or removing flaps of tissue, fragments of exposed bone, or any other bone and mucosa that may obstruct or otherwise prevent re-establishment of ventilation and drainage of the sinus.

8. Prepare balloon for dilation of additional sinuses (if desired).
   a. Gently advance the plunger rod into the syringe barrel to slightly expand the balloon to remove any wrinkles using minimal pressure.

   b. Clean the balloon prior to introduction into another sinus. This may be done by wiping the balloon with sterile wet gauze or dipping the balloon in sterile saline or sterile water.

   c. Pull back on the plunger rod to the second click position to apply vacuum to the balloon.

9. Repeat the same procedure to treat additional sinuses if desired.
   Note: The XprESS suction tip may be re-shaped to treat additional sinuses (see Figures 6-8).

10. After completing the entire procedure, dispose of the devices and all waste products according to appropriate environmental health safety guidelines.

How Supplied
The XprESS Multi-Sinus Dilation System is provided sterile and is intended for single-use only. Do not re-sterilize and/or re-use, as it may result in compromised device performance and risk improper sterilization and cross contamination. Do not use breached or damaged packages, since the sterility and functionality of the device may be compromised.

Limited Warranty
Entellus Medical, Inc. warrants that reasonable care has been used in the design and manufacture of this device. Entellus Medical excludes all other warranties, whether expressed or implied, by operation of law or otherwise including, but not limited to, any implied warranties of merchantability or fitness since handling and storage as well as other factors relating to the patient, diagnosis, treatment, medical procedures, and other matters beyond Entellus Medical’s control, directly affect the device and the results obtained from its use. Entellus Medical shall not be liable for any incidental or consequential loss, damage or expense, directly or indirectly arising from the use of this device. Entellus Medical neither assumes, nor authorizes any other person to assume for it, any other or additional liability or responsibility in connection with this device. Refer to Entellus Medical, Inc. Standard Terms and Conditions.

Symbols

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<td>Prescription Use Only</td>
<td>Authorized Representative: MedPass International Ltd.</td>
<td>Australian Sponsor: Compliance Management Solutions</td>
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<tr>
<td>Manufacturer</td>
<td></td>
<td>MedPass International Ltd.</td>
<td>19 Jack William Way</td>
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<td>Sterilization with Ethylene Oxide Gas</td>
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<td>Worcs. WR11 7JJ</td>
<td>BERWICK, VIC, 3806</td>
</tr>
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<td></td>
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</tbody>
</table>

Not made with natural rubber latex.

XPRESS, PATHASSIST and LIGHT FIBER are trademarks of Entellus Medical. patent http://www.entellusmedical.com/patents

Manufactured by: Entellus Medical Inc. 3600 Holly Lane North, Suite 40 Plymouth, MN 55447 USA +1 866-620-7615 (f) +1 866-620-7616 www.entellusmedical.com

Authorized Representative: MedPass International Ltd. 19 Jack William Way BERWICK, VIC, 3806

Australian Sponsor: Compliance Management Solutions

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INSTRUCTIONS FOR USE
PathAssist™ LED Light Fiber™
Read all Instructions prior to use

Caution: Federal (USA) law restricts this device to sale by or on the order of a physician.
Sterility: Provided Sterile, Ethylene Oxide (EO) Sterilization
Single Use: Disposable, For Single Patient Use Only, Do Not Resterilize and/or Reuse
Storage: Store in a cool, dry place. Do not expose to high temperatures above 50°C (122°F).

Indication For Use
To locate, illuminate within, and transilluminate across nasal and sinus structures.

Description
The PathAssist LED Light Fiber is a single use, disposable, flexible instrument that emits light from the distal end. The device consists of a flexible illumination fiber, a protective sheath and an integrated battery powered LED light source. When the LED Light Fiber is activated the fiber will emit red light from the distal tip for over 60 minutes. It has a fiber nominal working length of 27.6cm with an outer diameter of 0.375mm (0.015”).

Figure 1 LED Light Fiber

The LED Light Fiber is packaged alone or may also be packaged with XprESS (LoProfile or Ultra Suction Tips).

Contraindications
None known

Warnings
• Do not use breached or damaged packages, since the sterility and functionality of the device may be compromised.
• Single use only. Do not re-sterilize or re-use, as it may result in compromised device performance and risk improper sterilization and cross contamination.
• Due to the variability of sinus development in pediatric patients, review CT scan to assess each sinus’s development and appropriateness for balloon dilation. Pneumatization may occur as early as 1-2 years of age and continues to develop throughout childhood. Do not use LED Light Fiber in a sinus that is not adequately developed.
• Never advance or withdraw the device against unknown resistances as this can cause tissue trauma or device damage.
• Do not rest the device on the patient during surgery while it is activated, as this could result in burns to the patient.
• No modification of this device is allowed.

Precautions
- Due to the variability of sinus anatomy, review radiographic imaging (CT scan) prior to the procedure.
- Do not kink the LED Light Fiber as this may damage the device.
- Be sure to pre-load the fiber into the XprESS device prior to shaping it into a maxillary bend configuration (i.e., approximately 135° bend) as the fiber will not load when XprESS is pre-shaped in a maxillary configuration.
- Wait to activate the LED Light Fiber just prior to use as once activated the fiber will emit continuous light for over 60 minutes. There is no on/off switch.
- Do not stare directly at LED Light Fiber tip, or point it directly at anyone’s eyes while illumination is active.
- Do not use the device for external transillumination of maxillary sinus by applying the device to the hard palate, as this use has not been tested.
- Do not incinerate the device except for disposal in a controlled incinerator.
### Adverse Effects
Possible adverse effects include, but are not limited to, the following:

- Cerebrospinal fluid leak
- Damage of the orbital wall or other structures of the eye
- Tissue inflammation or trauma

### Compatibility
The device is compatible with the XprESS Multi-Sinus Dilation System *(all suction tip sizes)*

*Please refer to the XprESS Multi-Sinus Dilation System Instructions for Use for detailed information and instructions on the use of XprESS.*

### Instructions for Use

**NOTE:** Steps 1-3 are only necessary if LED Light Fiber is packaged alone. If LED Light Fiber is packaged with XprESS device, go to STEP 4.

1. Remove the LED Light Fiber from the protective packaging.
2. Load the fiber into the working lumen of XprESS (Figure 2).
3. Attach the LED Light Fiber housing to the barbed fitting of the XprESS device (Figure 3). Align the distal tip of the fiber with the distal end of XprESS (Figure 4).
4. Shape loaded XprESS to desired bend configuration for targeted sinus.
5. Activate the LED Light Fiber by removing the pull tab. Confirm that light is being transmitted through the LED Light Fiber.
6. Under endoscopic visualization, place the loaded XprESS device into the target location to illuminate within and transilluminate across nasal and sinus structures.
   - Projected illumination can be enhanced by slightly advancing tip of the LED Light Fiber distal from the XprESS device.
7. After procedure, dispose of device according to Federal, state, and local regulations, and appropriate environmental health safety guidelines. Do not incinerate except for disposal in a controlled incinerator.

### Specifications

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<th>Item</th>
<th>Specification</th>
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<tr>
<td>Weight</td>
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<tr>
<td>Nominal working length of fiber</td>
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<tr>
<td>Fiber outer diameter</td>
<td>0.375mm (0.015”)</td>
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<td>Light source (red LED)</td>
<td>625nm wavelength</td>
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<td>Activation time</td>
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<td>Maximum LED output power for treatment</td>
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<td>Safe operating ambient temperature range</td>
<td>15 - 33°C (59 - 91°F)</td>
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<td>Safe storage and transport temperature range</td>
<td>-10 - 50°C (14 - 122°F)</td>
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<td>Safe operating, storage, &amp; transport relative humidity range</td>
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Complies with medical safety standards: IEC 60601-1:2005; CAN/CSA-C22.2 No. 60601-1-08

Complies with medical EMC standard: IEC 60601-1-2:2007; Type BF applied part
Electromagnetic Compatibility (EMC)

Medical Electrical Equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided in this section. Portable and Mobile RF communications equipment can affect Medical Electrical Equipment.

### Guidance and Manufacturer's Declaration - Emissions

The LED Light Fiber is intended for use in the electromagnetic environments specified below.

The customer or the user of the LED Light Fiber should assure that it is used in such an environment.

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<tr>
<th>Emission Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment - Guidance</th>
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<tr>
<td>RF Emissions CISPR 11</td>
<td>Group 1</td>
<td>The LED Light Fiber uses RF energy only for its internal function. Therefore, its emissions are very low and are not likely to cause any interference in nearby electrical equipment.</td>
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<tr>
<td>RF Emissions CISPR 11</td>
<td>Class B</td>
<td>The LED Light Fiber is suitable for use in all establishments, including domestic, and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonics IEC 61000-3-2</td>
<td>N/A</td>
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<td>N/A</td>
<td></td>
</tr>
</tbody>
</table>

### Guidance and Manufacturer’s Declaration – Immunity

The LED Light Fiber is intended for use in the electromagnetic environments specified below.

The customer or the user of the LED Light Fiber should assure that it is used in such an environment.

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>EN/IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment - Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESD EN/IEC 61000-4-2</td>
<td>±6kV Contact, ±8kV Air</td>
<td>±6kV Contact, ±8kV Air</td>
<td>Floors should be wood, concrete, or ceramic tile. If floors are synthetic, the RH should be at least 30%.</td>
</tr>
<tr>
<td>EFT EN/IEC 61000-4-4</td>
<td>±2kV Mains, ±1kV I/Os</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Surge EN/IEC 61000-4-5</td>
<td>±1kV Differential, ±2kV Common</td>
<td>(LED Light Fiber is powered by internal battery)</td>
<td></td>
</tr>
<tr>
<td>Voltage Dips/Dropout EN/IEC 61000-4-11</td>
<td>&gt;95% Dip for 0.5 Cycle, 60% Dip for 5 Cycles, 30% Dip for 25 Cycles, &gt;95% Dip for 5 Seconds</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Power Frequency EN/IEC 61000-4-8</td>
<td>50/60Hz, Magnetic Field</td>
<td>3 A/m</td>
<td>Power frequency magnetic fields should be that of a typical commercial or hospital environment.</td>
</tr>
</tbody>
</table>

### Guidance and Manufacturer’s Declaration – Immunity

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<tbody>
<tr>
<td>Conducted RF EN/IEC 61000-4-6</td>
<td>3Vrms, 150kHz to 800MHz</td>
<td>N/A (LED Light Fiber is powered by internal battery)</td>
<td>Portable and mobile RF communications equipment should be used no closer to the LED Light Fiber than the distances calculated or listed below.</td>
</tr>
<tr>
<td>Radiated RF EN/IEC 61000-4-3</td>
<td>3Vms, 80MHz to 2.5GHz</td>
<td>3V/m (E1)</td>
<td>Recommended Separation Distance</td>
</tr>
</tbody>
</table>

\[
d = (1.2)(P) \quad 800MHz to 8000MHz
d = (2.3)(P) \quad 2.5MHz
\]

Where \( P \) is the maximum output power rating of the transmitter in watts and \( d \) is the recommended separation distance in meters.

Field strengths from fixed transmitters, as determined by an electromagnetic site survey, should be less than the compliance level (E1). Interference may occur in the vicinity of equipment containing a transmitter.

### Recommended Separation Distances between portable and mobile RF communications equipment and the LED Light Fiber

The LED Light Fiber is intended for use in the electromagnetic environment in which radiated RF disturbances are controlled. The customer or user of the LED Light Fiber can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF Communications Equipment (transmitters) and the LED Light Fiber as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Max Output Power of Transmitter (Watts)</th>
<th>Separation distance according to frequency of transmitter (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150kHz to 800MHz</td>
</tr>
<tr>
<td>0.01</td>
<td>N/A (LED Light Fiber is powered by an internal battery)</td>
</tr>
<tr>
<td>0.1</td>
<td>N/A (LED Light Fiber is powered by an internal battery)</td>
</tr>
<tr>
<td>1</td>
<td>Conducted RF Immunity testing does not apply, resulting in no separation data from 150kHz to 80MHz.</td>
</tr>
<tr>
<td>10</td>
<td></td>
</tr>
<tr>
<td>100</td>
<td></td>
</tr>
</tbody>
</table>
**Limited Warranty**

Entellus Medical, Inc. warrants that reasonable care has been used in the design and manufacture of this device. Entellus Medical excludes all other warranties, whether expressed or implied, by operation of law or otherwise including, but not limited to, any implied warranties of merchantability or fitness since handling and storage as well as other factors relating to the patient, diagnosis, treatment, medical procedures, and other matters beyond Entellus Medical's control, directly affect the device and the results obtained from its use. Entellus Medical shall not be liable for any incidental or consequential loss, damage or expense, directly or indirectly arising from the use of this device. Entellus Medical neither assumes, nor authorizes any other person to assume for it, any other or additional liability or responsibility in connection with this device. Refer to *Entellus Medical, Inc. Standard Terms and Conditions*.

*Graphic Symbols Contained on Device labeling*

<table>
<thead>
<tr>
<th><strong>Consult Instructions for use</strong></th>
<th><strong>LOT</strong></th>
<th><strong>MODEL</strong></th>
<th><strong>#</strong></th>
<th><strong>EC REP</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lot Number</td>
<td>Model Number</td>
<td>Quantity</td>
<td>Authorized Representative in the European Community</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>STERILE EO</strong></th>
<th><strong>Manufacturer</strong></th>
<th><strong>Do Not Use</strong></th>
<th><strong>Rx Only</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sterilization with Ethylene Oxide Gas</td>
<td></td>
<td></td>
<td>Prescription Use Only</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Type BF applied part</strong></th>
<th><strong>REF</strong></th>
<th><strong>Use By</strong></th>
<th><strong>CE Mark</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Reorder Number</td>
<td>Use By</td>
<td>0086 CE Mark</td>
</tr>
</tbody>
</table>

Not made with natural rubber latex.

PathAssist, Light Fiber and XprESS are trademarks of Entellus Medical.

patent  http://www.entellusmedical.com/patents

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