**COR-KNOT MINI® EU TECHNOLOGY GUIDE**

**FIG. 1**

**COR-KNOT® QUICK LOAD® UNIT DESCRIPTION**

Each COR-KNOT® QUICK LOAD® UNIT provides one sterile COR-KNOT® FASTENER ① held in a customized loading unit consisting of a purple target ②, a wire snare ③, and a blunt curved handle ④. Made from medical grade titanium, a COR-KNOT® FASTENER is a mushroom-shaped hollow sleeve, which is crimped by the COR-KNOT MINI® DEVICE to fasten together segments of suture ⑤.

**COR-KNOT MINI® DEVICE DESCRIPTION**

Each sterile package (kit) contains two 17cm long SINGLE PATIENT USE COR-KNOT MINI® DEVICES and twelve COR-KNOT® QUICK LOAD® UNITS (COMBO KIT) or two COR-KNOT MINI® DEVICES only (DEVICE KIT). A COR-KNOT® FASTENER is loaded into the distal tip of the 4mm diameter shaft ⑥. A rotational knob ⑦ with an indicator fin ⑧, a white handle ⑨ and purple lever ⑩ are located at the proximal end of the device. By squeezing the purple lever, the COR-KNOT MINI® DEVICE crimps the COR-KNOT® FASTENER at the closure site and can trim away excess suture.

**INDICATIONS**

The COR-KNOT MINI® DEVICE used in conjunction with LSI SOLUTIONS® specified 2-0 polyester suture and a COR-KNOT® titanium fastener is indicated for use to fasten and trim suture in general and cardiovascular surgical applications.

**FIG. 2**

Rotating Shaft

**COR-KNOT MINI® DEVICE**

4mm Diameter, 17cm Length Shaft

Courtesy of Peter A. Knight, M.D.
AT SCRUB TABLE

LOADING WITH A COR-KNOT® QUICK LOAD® UNIT
Use proper operating room technique to pass the sterile COR-KNOT® QUICK LOAD® UNIT from its packaging. While maintaining appropriate sterile technique, follow the steps indicated in the illustrations.

1. INSERT the blunt tip of the curved handle into the distal slot at the end of the COR-KNOT MINI® DEVICE shaft. ROTATE the curved handle through the distal slot and out of the suture slot until the COR-KNOT® FASTENER occupies the shaft’s distal slot. Fully ENGAGE the COR-KNOT® FASTENER within the tip of the COR-KNOT MINI® DEVICE by pushing on the purple target or by pulling on the curved handle.

2. PUSH-OUT and remove the purple target.

3. RELEASE the curved handle from the distal slot at the end of the COR-KNOT MINI® DEVICE shaft.

4A. INSPECT to ensure that the COR-KNOT® FASTENER is well loaded and fully seated.

ACTIONS
When the COR-KNOT MINI® DEVICE is loaded with a COR-KNOT® FASTENER and appropriately positioned at a suture closure site, squeezing the purple lever can instantly secure and trim the suture. The surgical titanium used in a COR-KNOT® FASTENER is not absorbed by the body and is generally not associated with significant inflammatory reactions.

CONTRAINdications
• Endoscopic procedures should only be performed by physicians having adequate training and familiarity with endoscopic techniques. Medical literature should be consulted relative to techniques, complications and hazards prior to the performance of endoscopic procedures.
• The COR-KNOT® QUICK LOAD® UNIT is not intended to be used with any device other than the COR-KNOT MINI® DEVICE or the COR-KNOT® DEVICE. The COR-KNOT MINI® DEVICE is not intended to be loaded with anything other than a COR-KNOT® QUICK LOAD® UNIT.
• The COR-KNOT® FASTENER is NOT intended for placement into circulating blood unless used with compatible suture under conditions judged by the surgeon to be clinically appropriate.
• Use only with LSI SOLUTIONS® specified 2-0 polyester suture.
• Each COR-KNOT MINI® DEVICE is not intended to be fired more than 12 times.

WARNINGS
• Users should be familiar with standard procedures and techniques involving surgical suture and titanium usage before employing the COR-KNOT MINI® DEVICE with a COR-KNOT® QUICK LOAD® UNIT for fastening and trimming suture.
• Adequate COR-KNOT® FASTENER security requires reasonable clinical judgment and appropriate surgical techniques as warranted by surgical circumstances and the experience of the surgeon.
• When securing suture with a COR-KNOT MINI® DEVICE, ensure any needle caps or needles are removed from the suture ends to be loaded prior to loading the suture through the COR-KNOT MINI® DEVICE.
• Excessive suture tensioning can cause suture breakage.
• Single patient use only. Do not reclean or resterilize. Adequate cleaning or removal of blood and other foreign materials from used COR-KNOT® products cannot be guaranteed. Validation of resterilization is not established. Failure to eliminate inflammatory or infectious agents may cause patient harm. Product functional performance may be compromised in reprocessed devices or COR-KNOT® FASTENERS.
• Discard any open (unsealed), unused, expired or damaged COR-KNOT® product.
• COR-KNOT® QUICK LOAD® UNIT components and each COR-KNOT MINI® DEVICE, along with packaging, must be inspected, handled and disposed of consistent with standard, accepted medical device disposal procedures.
• Direct contact between sensitive tissue structures (e.g., pulsatile arteries, cardiac valve leaflets, valve chordae, etc.) and foreign materials can lead to tissue injury or damage, such as tissue erosion. Always orient COR-KNOT® FASTENERS and remnant suture tails to avoid direct contact between delicate tissue or prosthetic structures.
• As with any foreign body, prolonged contact of any suture with salt solutions, such as urine and bile, may result in calculus formation.
• While the titanium of the COR-KNOT® FASTENER is physiologically very inert, routine surgical precautions must be employed whenever foreign materials are left in a patient.

PRECAUTIONS
• When handling the COR-KNOT® QUICK LOAD® UNIT care should be taken to avoid damage.
• Do not rotate the shaft while the fastener is loaded with suture.
• Do not squeeze the purple lever of COR-KNOT MINI® DEVICE while loading the COR-KNOT® QUICK LOAD® UNIT.
• Irreparable damage to COR-KNOT MINI® DEVICE suture cutting blade will occur if the purple lever is squeezed while the COR-KNOT® QUICK LOAD® UNIT curved handle is in place at the tip of the instrument.
• Ensure that obstructions do not interfere with the firing of COR-KNOT MINI® DEVICE.
• Do not squeeze the purple lever of the loaded COR-KNOT MINI® DEVICE, until the COR-KNOT® FASTENER has been appropriately positioned directly upon the tissue or prosthetic material and the suture accurately tensioned at the targeted site.
• Always squeeze and hold the purple lever and then fully release it before moving the COR-KNOT MINI® DEVICE tip. Failure to appropriately release the purple lever can cause suture breakage. Inspect each COR-KNOT® FASTENER and its suture tails.
• Do not squeeze the purple lever on the same COR-KNOT® FASTENER more than once.
• Cut suture with scissors if the COR-KNOT MINI® DEVICE fails to trim suture or release COR-KNOT® FASTENER.
• Avoid crushing or crimping damage to the COR-KNOT® FASTENER due to inappropriate squeezing of COR-KNOT MINI® DEVICE purple lever, and/or to application of surgical instruments like forceps, needle holders, clamps, etc.
• If COR-KNOT® FASTENER falls out of tip or is not properly loaded, retrieve loose fastener, reload with new fastener and start again.
• If the purple lever of the COR-KNOT MINI® DEVICE does not return completely forward on its own (i.e., without assistance), manually push the lever forward all the way to release the COR-KNOT® FASTENER.
• Check for hemostasis or leakage where appropriate.
• Before endoscopic instruments and accessories from different manufacturers are employed together in a procedure, verify compatibility and ensure that electrical isolation or grounding are not compromised.

ADVERSE REACTIONS
Adverse effects associated with the use of surgical suture and titanium can include, but are not limited to: wound dehiscence, thrombus formation, embolism, calcification formation in urinary and biliary tracts when prolonged contact with salt solutions such as urine and bile occurs, infected wounds, minimal acute inflammatory tissue reaction, and transitory local irritation. Surgical titanium is not absorbed by the body and is generally not associated with inflammatory reactions. Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the country competent authority.
**THREADING SUTURE THROUGH A LOADED COR-KNOT® FASTENER**

Extracorporeally: surgeons typically use their non-dominant hand to hold the device near the end of its shaft and their dominant hand to complete the suture threading technique. Rotate the shaft by turning the indicator knob to orient the suture slot toward the center of the prosthetic.

5. **PASS** both ends of the suture through the open wire snare at the end of the shaft.

6. **PULL** the curved handle with its attached wire snare containing the suture ends towards the purple lever to draw the snared bends in the suture into the COR-KNOT® FASTENER.

7. **THREAD** the suture through the COR-KNOT® FASTENER and out of the suture slot near the end of the shaft by continuing to pull the curved handle until the wire snare and both ends of the suture exit through the suture slot.

8A. **GRASP** both ends of the suture after passing off the curved handle with the wire snare.

8B. **INSPECT** to ensure fastener is fully seated in distal device tip.

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**OUTSIDE OF PATIENT**

5. **PASS**

6. **PULL**

7. **THREAD**

8A. **GRASP**

8B. **INSPECT**

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**INSIDE OF PATIENT**

9. **SLIDE** the COR-KNOT®-MINI DEVICE distal tip gently over the partially tensioned suture down to the targeted site with suture slot and rotational knob’s indicator fin oriented toward the center of the prosthesis.

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**SQUEEZE & HOLD / TUG & RELEASE / REMOVE DEVICE**

With the distal tip on the prosthesis, use one hand to apply sufficient suture tension to hold tissue and prosthesis in appropriate apposition, then the other hand to:

10. **SQUEEZE** the purple lever until it stops, maintain the device tip’s position, **HOLD** lever for one second.

11. **TUG** the suture gently to cut free both suture ends and **RELEASE** the purple lever fully to release the crimped COR-KNOT® FASTENER.

12. **REMOVE DEVICE** and inspect to ensure COR-KNOT® FASTENER and suture tails are oriented away from delicate tissue and prosthesis structures.

**PLEASE NOTE** While abruptly squeezing and releasing of the purple lever provides average suture holding forces above EP and USP standards, this recommended **SQUEEZE & HOLD** technique assures optimized suture holding force. Visually inspect each suture tail and COR-KNOT® FASTENER.

If the crimped COR-KNOT® FASTENER does not readily release from the distal tip, ensure purple lever is released, then gently **PUSH** inward and **ROTATE** the handle 90° about the shaft. If still necessary, **ROTATE** the handle back, then turn 90° in the opposite direction. If COR-KNOT® FASTENER will still not release, cut suture.

**FAILURE TO PROPERLY LOAD SUTURE**

**NOTE:** TO REMOVE A RETAINED COR-KNOT® FASTENER IF THE WIRE SNARE IS INADVERTENTLY REMOVED WITHOUT PROPER SUTURE THREADING, SQUEEZE AND RELEASE THE PURPLE LEVER AND THEN TAP THE DISTAL SHAFT ON A TABLE OR USE A SCALPEL TO PRY OUT THE CRIMPED COR-KNOT® FASTENER.
**COR-KNOT® QUICK LOAD® UNIT**

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**CRIMPED COR-KNOT® FASTENER**

ACTUAL SIZE

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**OVERALL QUALITATIVE AND QUANTITATIVE COMPOSITION**

The only portion of the COR-KNOT® QUICK LOAD® UNIT which is an implant is the COR-KNOT® FASTENER, which is made up of 100% commercially pure titanium.

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**COR-KNOT® QUICK LOAD® UNIT DESCRIPTION**

Each COR-KNOT® QUICK LOAD® UNIT provides one sterile COR-KNOT® FASTENER 1 held in a customized loading unit consisting of a purple target 2, a wire snare 3, and a blunt curved handle 4. Made from medical grade titanium, a COR-KNOT® FASTENER is a mushroom-shaped hollow sleeve, which is crimped by the COR-KNOT® MINI® DEVICE to fasten together segments of suture.

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**COR-KNOT® QUICK LOAD® UNIT**

1. COR-KNOT® FASTENER
2. PURPLE TARGET
3. WIRE SNARE
4. CURVED HANDLE

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ACTUAL SIZE
SUTURE CUTTING DIFFICULTY - THE CAUSES OF SUTURE NOT CUTTING EASILY WHEN USING A COR-KNOT MINI® DEVICE CAN INCLUDE:

• USER ERROR INDUCED DAMAGE DULLING THE SUTURE CUTTING BLADE OR
• DEVICE MALFUNCTION, WHICH MAY ALSO REDUCE FASTENER STRENGTH AND SECURITY

If suture cutting difficulty occurs while using any COR-KNOT MINI® DEVICE, discontinue its intraoperative use and REMOVE DEVICE from the surgical field. Visually INSPECT FASTENER to compare its crimp to other fasteners. Pull or tug on the fastener with a forceps or clamp to TEST FASTENER and suture security.

REMOVE DEVICE–INSPECT FASTENER–TEST FASTENER–RETURN DEVICE

COR-KNOT MINI® DEVICE suture cutting difficulty can be induced by the inadvertent squeezing of the purple lever while the metal loading components are still in the distal device shaft. This user error can lead to irreparable damage to the suture cutting blade by driving the blade into the metal curved handle or metal wire snare.

MRI SAFETY INFORMATION – MR CONDITIONAL

Non-clinical testing demonstrated that the COR-KNOT® Titanium Fastener is MR Conditional. A patient with this device can be scanned safely in an MR system under the following conditions:

• Static magnetic field of 1.5-Tesla and 3-Tesla, only
• Maximum spatial gradient magnetic field of 4,000-Gauss/cm (40-T/m)
• Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2-W/kg for 15 minutes of scanning (i.e., per pulse sequence) in the Normal Operating Mode.

Under the scan conditions defined, the COR-KNOT® Titanium Fastener is expected to produce a maximum temperature rise of 1.5°C after 15-minutes of continuous scanning (i.e., per pulse sequence).

In non-clinical testing, the image artifact caused by the COR-KNOT® Titanium Fastener extends approximately 2-mm from this implant when imaged using a gradient echo pulse sequence and a 3-Tesla MR system.

HARVESTED AT 18 MONTHS

Courtesy of Scott M. Goldman, M.D.
### FIG. 6  **COR-KNOT MINI®** PRODUCT ORDERING

<table>
<thead>
<tr>
<th>REORDER</th>
<th>PRODUCT</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>031400</td>
<td><strong>COR-KNOT MINI® DEVICE KIT</strong></td>
<td>Box of 6 Kits (2 Devices per Kit)</td>
</tr>
<tr>
<td>030950</td>
<td><strong>COR-KNOT® QUICK LOAD® SINGLES</strong></td>
<td>Box of 12 Singles (1 Fastener per Pouch)</td>
</tr>
<tr>
<td>030902</td>
<td><strong>COR-KNOT® QUICK LOAD® 6-POUCH</strong></td>
<td>Box of 12 Pouches (6 Fasteners per Pouch)</td>
</tr>
<tr>
<td>031450</td>
<td><strong>COR-KNOT MINI® COMBO KIT</strong></td>
<td>Box of 6 Kits (2 Devices &amp; 12 Fasteners per Kit)</td>
</tr>
</tbody>
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**DESCRIPTION**

**PRODUCT**

**SUPPLIED:** STERILE

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**INTRA OP**

![Image of surgical procedure](image1)

 nonzeroannorphotof.png

Courtesy of Peter A. Knight, M.D.

**3D ECHO**

![Image of surgical procedure](image2)

 nonzeroannorphotof.png

Courtesy of Peter A. Knight, M.D.

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The Summary of Safety and Clinical Performance (SSCP) Report, with information on device suitability and patient target group, shall be made available in the European database on medical devices (EUDAMED) at http://ec.europa.eu/tools/eudamed once the database is publicly accessible. The Basic UDI-DI for this device is 0850200006COR-KNOT2N.

Patents: www.lsisolutions.com/patents

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Symbol Glossary: www.lsisolutions.com/symbols

This Product Comes with our LSI SOLUTIONS® Perfect Performance Policy®

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