**MEDLINEUNITE® Calcaneal Fracture Plating SYSTEM**

**INSTRUCTIONS FOR USE**
Attention Operating Surgeon

**DEVICE DESCRIPTION**
The Medline UNITE® Calcaneal Fracture Plating System consists of 3 styles of Sinus Tarsi plates (Sinus Tarsi, Sinus Tarsi Offset, and Sinus Tarsi Extension) made from Titanium Alloy (Ti6Al4V ELI) and Perimeter plates made from Commercially Pure Titanium. The Sinus Tarsi, Sinus Tarsi Offset, and Sinus Tarsi Extension, and Perimeter plates are each offered in small, medium, and large sizes in left and right configurations. Previously cleared 3.5mm locking and non-locking screws (K151235) that range from 10-60mm in length will be included in the tray with the Medline UNITE® Calcaneal Fracture Plates. The system also includes reusable instrumentation necessary to implant the plates and screws, e.g. targeting guide, drill guides, drill bits.

**INDICATIONS**
Medline UNITE® Calcaneal Fracture Plating System are intended for use in stabilization of fresh fractures, revision procedures, joint fusion, and reconstruction of bones in the feet and ankles including extra-articular, intra-articular, joint depression, tongue-type, and severely comminuted fracture of the calcaneus. The system can be used in both adult and pediatric (adolescent and child) patients.

**INFORMATION FOR USE**
The surgeon must select the type of and size implant that best meets the patient’s surgical needs.

**CONTRAINDICATIONS**
1. Any previous or active infection or blood supply limitations.
2. Insufficient quality of bone or soft tissue.
3. Patients who are unwilling or incapable of following postoperative care instructions.
4. Material sensitivity. If suspected, tests should be conducted prior to implantation.
5. This device is not intended for screw attachment or fixation to the posterior elements (pedicles) of the cervical, thoracic, or lumbar spine.

**ADVERSE EFFECTS**
- Fracture of implant due to excessive activity, prolonged loading upon the device, incomplete or inadequate healing, or excessive force exerted on the implant during insertion.
- Implant migration and/or loosening requiring revision surgery.
- Bone resorption or over-production.
- Metal sensitivity or histological or allergic reaction resulting from implant material.
- Infection or painful, swollen or inflamed implant site.
- Unexpected histological response possibly involving macrophages and/or fibroblasts.
- Migration of particle wear debris possibly resulting in bodily response.
- Embolism.
- Pain, discomfort, or abnormal sensations due to the presence of an implant.
- Necrosis of the bone.
- Necrosis of the tissue.
- Nerve damage resulting from surgical trauma.

**PRE-OPERATIVE PRECAUTIONS**
It is the surgeon’s responsibility to determine the best course of action for each patient. The surgeon should carefully consider the following when selecting a patient for surgery:
- More conservative treatment options.
- Patient’s weight, occupation, and activity level.
- Patient’s expectations of the device.
- Patient’s willingness to follow post-operative instructions.
- Patient conditions that may limit their ability to recognize limitations of the device that may lead to device failure, such as senility, mental illness, or alcoholism.
- Known or suspect material allergies.
- Comorbidities, eg. diabetes, smoking.

The patient should be warned of surgical risks and be made aware of possible adverse effects. For safe and effective use, the surgeon must be thoroughly familiar with this type of implant, the method of application, instrumentation, and the recommended surgical technique and indications for this type of device. Improper implantation of the device can increase the possibility of loosening or migration. No metal implants can be expected to withstand loads at the same level as healthy bone.

**INTRA-OPERATIVE PRECAUTIONS**
- The surgeon must select the type and size implant that best meets the patient’s surgical needs.
- Inspect all implants for scratching and nicking prior to use as such stress concentrations can lead to failure. Avoid flawing the implant surface during insertion to minimize the potential for failure.
- An implant shall never be reused. Previous stresses may have created imperfections, which can lead to a device failure.
- Instruments, particularly drills, and drivers shall be inspected for wear or damage prior to use.
- Medline UNITE® Calcaneal Fracture Plates are designed specifically for use with Medline instrumentation. The use of other instrumentation is not recommended.

**POST-OPERATIVE PRECAUTIONS**
The postoperative care instructions provided by the surgeon should be strictly followed to avoid adverse stresses applied to the device. Failure to follow postoperative care instructions can cause implant and/or treatment failure. Periodic follow-up, including x-rays for comparison to early post-op conditions, is recommended to monitor the position and state of the implant, the condition of the bone, and any signs of implant migration, loosening, bending, or cracking.

**RECOMMENDATIONS REGARDING DEVICE FRAGMENTS**
1. In the event of implant fracture, carefully consider and discuss with the patient, if possible, the risk/benefit of removing the hardware versus leaving the fragment in the patient. Things to consider and discuss with the patient include; the material composition of the fragment (if known), the size and location of the fragment, and the potential mechanisms of injury if the fragment is not removed, including MRI exams.
2. Any decision to remove the device should take into consideration the potential risk to the patient of a second surgical procedure.
3. If removal is determined to be the best course of action, inspect the device immediately after removal from patient for signs of damage. If the device is damaged, retain the device to assist the manufacturer’s investigation of the event. Device removal should be followed by adequate postoperative care.
CONCERNING MAGNETIC RESONANCE ENVIRONMENT
Medline UNITE® Calcaneal Fracture Plates have not been evaluated for safety and compatibility in the MR environment. It has not been tested for heating, migration, or image artifact in the MR environment. The safety of the Medline UNITE® Cannulated Screws in the MR environment is unknown. Scanning a patient who has this device may result in patient injury.

CLEANING & STERILIZATION
All implants and instruments are provided non-sterile in trays or individually packaged and must be adequately cleaned and sterilized prior to use or re-use. A manual cleaning method is provided and has been developed and tested consistent with TIR 30 (manual). The provided sterilization recommendations have been developed and tested consistent with ANSI/AAMI ST79. An implant should never be re-sterilized after contact with human body tissues or fluids. Devices labeled for single use only should never be reused. Reuse of these devices may result in, but are not limited to: decreased performance of the product, cross-infection, and contamination. Trays should be thoroughly cleaned and inspected prior to use. Visually inspect the tray for cleanliness. Repeat cleaning process, as necessary, until tray is visually clean. All instruments, where applicable, must be disassembled prior to cleaning.

Manual Cleaning
Do not allow excessive debris and soft tissue to dry after use. Begin cleaning process as soon after use as possible. Additional scrubbing may be required if debris and soft tissue dry.

1. Rinse under cool running tap water to remove visual soil.
2. Prepare enzymatic detergent per manufacturer’s recommendation using lukewarm tap water.
3. Allow the device to soak in the prepared enzymatic detergent for 1 minute.
4. Using a soft bristled brush, thoroughly brush the device to remove soil.
5. Run a stylet through the lumens a minimum of 3 times to remove soil.
6. Using a syringe, aggressively flush the lumens with the prepared enzymatic detergent.
7. Remove the device from the detergent and rinse under cool running tap water to remove detergent residuals.
8. Using a syringe, aggressively flush the lumens.
9. Prepare neutral detergent per manufacturer’s recommendation using warm tap water.
10. Allow the device to soak in the prepared neutral detergent for 3 minutes.
11. Using a soft bristled brush, thoroughly brush the device to remove soil.
12. Run a stylet through the lumens a minimum of 3 times to remove soil.
13. Using a syringe, aggressively flush the lumens with the prepared neutral detergent.
14. Remove the device from the detergent and rinse in running RO/DI water to remove detergent residuals.
15. Prepare enzymatic detergent per manufacturer’s recommendation in a sonication unit.
16. Allow the device to sonicate for 10 minutes.
17. Remove the device from the sonicator and thoroughly rinse under running critical water.
18. Using a syringe, aggressively flush the lumens.
19. Dry the device with a disposable, lint-free cloth.
20. Visually inspect the device for cleanliness.

Instruments should be inspected for any damage or wear prior to use. Instruments that have corrosion, pitting, and/or discoloration should not be used.

Sterilization
Sterilization should be performed in the provided tray double-wrapped in FDA cleared sterilization wrap using the following method:

<table>
<thead>
<tr>
<th>Cycle Type</th>
<th>Parameter</th>
<th>Minimum Set Point</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-vacuum</td>
<td>Exposure Temperature</td>
<td>270°F (132°C)</td>
</tr>
<tr>
<td></td>
<td>Exposure Time</td>
<td>4 Minutes</td>
</tr>
<tr>
<td></td>
<td>Dry Time</td>
<td>40 Minutes</td>
</tr>
</tbody>
</table>

Do not stack trays during sterilization.

Ensure that the implants and instruments are at room temperature prior to use.

STORAGE: Store all devices in a clean and dry environment. The devices are manufactured from non-degradable materials. When stored under the recommended conditions, the shelf life of this product is not limited.

CAUTION: Federal Law (USA) restricts this device to sale by or on the order of a physician.

For additional product information, please visit www.medline.com or contact customer service at 1-800-MEDLINE.
Manufactured for and distributed by Medline Industries, Inc. Three Lakes Dr, Northfield, IL 60093 USA 1-800-MEDLINE

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