The Paragon 28® Grappler® Suture Anchor System implants are comprised of suture anchors and suture that are intended for soft tissue repair. The Grappler® Suture Anchor System implants are offered in various configurations and sizes to accommodate patient anatomy.

IMPLANT MATERIALS
For implants of the Grappler® Suture Anchor System are made from polyetheretherketone (PEEK), titanium alloy, polyester, and ultra-high molecular weight polyethylene (UHMWPE). The instrumentation is manufactured from medical grades of stainless steel, nitinol, and polymer.

INDICATIONS FOR USE
The Paragon 28® Grappler® Suture Anchor System is intended for the fixation of soft tissue to bone including:
- Elbow: Biceps Tendon Reattachment, Lateral Epicondylitis Repair, Tennis Elbow Repair
- Shoulder: Rotator Cuff Repair, Bankart Repair, SLAP Lesion Repair, Biceps Tendonosis, Acromio-Clavicular Separation Repair, Deltoit Repair, Capsular Shift or Capsulolabral Repair
- Hand/Wrist: Scapholunate Reconstruction, Ulnar or Radial Collateral Ligament Reconstruction, TFCC
- Foot/Akle: Lateral Stabilization (Bromstrom, Bromstrom-Gould, Chrisman-Snook Repair), Ankle Ligament Repair, Medial Stabilization (Deltoid Repair, Spring Ligament Reconstruction), Achilles Tendon Repair, Metatarsal Ligament Repair, Syndesmosis Repair, Hallux Valgus Reconstruction, Digital Tendon Transfers, Midfoot Reconstruction, LisFranc Repair
- Knee: Medial Collateral Ligament Repair, Lateral Collateral Ligament Repair, Posterior Oblique Ligament Repair, Iliotibial Band Tendonosis, Extra Capsular Reconstruction, Patellar Ligament and Tendon Avulsion Repair
- Hip: Capsular Repair, Acetabular Labral Repair

CONTRAINDICATIONS
The Paragon 28® Grappler® Suture Anchor System implants are not designed or sold for any use except as indicated. Use of the Grappler® Suture Anchor System is contraindicated in the following situations:
- Active, suspected or latent infection in the affected area
- Patients who are physiologically or psychologically inadequate
- Coagulation; an overweight or corpulent patient can strain the implant to such a degree that stabilization or implant failure can occur
- Patients with a known allergy to the implant material(s)
- Patients previously sensitized to titanium
- Insufficient quantity or quality of bone or soft tissue to permit stabilization, conditions that retard healing (not including pathological fractures) and conditions causing poor blood supply

POTENTIAL COMPLICATIONS AND ADVERSE REACTIONS
In any surgical procedure, the potential for complications and adverse reactions exist. The risks and complications with these implants include:
- Loosening, deformation or fracture of the implant
- Acute post-operative infections and late infections with possible sepsis
- Migration, subluxation of the implant
- Wound hematoma and delayed wound healing
- Temporary or protracted functional neurological perturbation
- Tissue reactions as a result of allergy or foreign body reaction to dislodged particles or implant material
- Implant degradation with localized reaction and pain
- Pain, a feeling of malaise or abnormal sensations due to the implant used
- Bone resorption or over-production

All possible complications listed here are not typical of Paragon 28®, Inc. products but are in principle observed with any implant. Promptly inform Paragon 28®, Inc. as soon as complications occur in connection with the implants or surgical instruments used. In the event of premature failure of an implant in which a causal relationship with its geometry, surface quality or mechanical stability is suspected, please provide Paragon 28®, Inc. with the explant(s) in a cleaned, disinfected and sterile condition. Paragon 28®, Inc. cannot accept any other returns of used implants. The surgeon is held liable for complications associated with inadequate asepsis, inadequate preparation of the osseous implant bed in the case of implants, incorrect indication or surgical technique or incorrect patient information and consequent incorrect patient behavior.

WARNINGS AND PRECAUTIONS
- Re-operation to remove or replace implants may be required at any time due to medical reasons or device failure. If corrective action is not taken, complications may occur
- Use of an implant in areas of high functional stresses may lead to implant fracture and failure
- Implants, wires, or other appliances of dissimilar metals should not be used together in or near the implant site
- The implants are intended for single use only.
- Instruments and implants are to be treated as sharps.
- Avoid K-wires through the implant.
- Avoid flawed implant surfaces to minimize the potential for early fatigue failure.
- Do not use other manufacturer’s instruments or implants in conjunction with the Grappler® Suture Anchor System.
- Do not resterilize the Grappler® Suture Anchor System Implants and Instruments.

MR SAFETY INFORMATION
The Grappler® Suture Anchor System has 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 Grappler® Suture Anchor System in the MR environment is unknown. Scanning a patient who has this device may result in patient injury.

MAINTAINING DEVICE EFFECTIVENESS
- The surgeon should have specific training, experience, and thorough familiarity with the use of fixation devices.
- Sutures should be considered in vivo performance when selecting a suture type.
- The surgeon must exercise reasonable judgment when deciding which implant type to use for specific indications.

CONTRAINDICATIONS
- Consideration should be taken in the use of absorbable sutures in tissues with poor blood supply as suture extrusion and delayed absorption may occur.
- The Grappler® Suture Anchor System implants are not intended to endure excessive abnormal functional stresses.
- Failure to use dedicated, unique Grappler® Suture Anchor System instruments for every step of the implantation technique may compromise the integrity of the implanted device, leading to premature device failure and subsequent patient injury. Failed devices may require re-operation and removal.
- Carefully inspect the implants prior to use, inspect the instruments before and after each procedure to assure they are in proper operational condition. Instruments which are faulty, damaged or suspect should not be used.
- Paragon 28®, Inc. recommends the use of Paragon 28®, Inc. products in a sterile environment.

HANDLING AND STERILIZATION
Sterile Product:
Paragon 28® Grappler® Suture Anchor System products are provided sterile using ethylene oxide. Do not re-sterilize. SINGLE USE ONLY. The risk of re-use of device includes potential for patient to develop infection. Do not use product after expiration date. Packages for product should be intact upon receipt.

Product in sterile packaging should be inspected to ensure that the package has not been damaged or previously opened. If the inner package integrity has been compromised, DO NOT USE THE DEVICE. Contact the manufacturer for further instructions. The product should be opened using aseptic technique. Once the seal of the product is broken, the product should not be re-sterilized.

All product should be stored in a clean, dry environment.

INSTRUCTIONS FOR USE
Only surgeons who are fully experienced in the use of such implants and the required specialized surgical techniques should implant the Paragon 28® Grappler® Suture Anchor System. Refer to the Grappler® Suture Anchor System Surgical Technique P44-STG-0001 for complete instructions for use. For product information or to obtain a copy of the surgical technique manual, please contact Paragon 28®, Inc. by phone, (855) 786-2828.

IMPLANT REMOVAL (IF NECESSARY)
- Instrumentation can be provided for implant removal.
- Removal instructions are provided in the Grappler® Suture Anchor System Surgical Technique (P44-STG-0001).

PRODUCT COMPLAINTS
The customer or health care provider should report any dissatisfaction with the product quality, labeling, or performance to Paragon 28®, Inc. immediately. Paragon 28®, Inc. should be notified immediately of any product malfunction by telephone or written correspondence. When filing a complaint, the name, part number and lot number of the part should be provided along with the name and address of the person filing the complaint.

Please contact company for product inquiries, cleaning instructions and surgical techniques, or to report any adverse event.