DISTAL RADIUS

Instructions for Use
1. **INTENDED USE**

The Conventus CAGE™ – Distal Radius is intended for the fixation of distal radius fractures.

2. **DEVICE DESCRIPTION**

The Conventus CAGE™ – Distal Radius (DR CAGE) is an expandable implant which is deployed into the intramedullary canal of the distal radius and provides a scaffold to which bone fragments can be attached using fragment screws. The DR CAGE is flexible during device placement, but once deployed creates a stable bone-implant construct at the completion of the surgical implantation procedure.

The DR CAGE is comprised of an Expandable Scaffold (Inner Scaffold, Outer Scaffold, Central Member), Fragment Screws, Proximal Plate and Fragment Screws in a proximal location. All components of the system are made of known biocompatible materials as listed in **Table 1**.

<table>
<thead>
<tr>
<th>Component</th>
<th>Material and Standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expandable Scaffold</td>
<td>Scaffolds: Nitinol (ASTM F2063)</td>
</tr>
<tr>
<td></td>
<td>Central Member: (ASTM F136)</td>
</tr>
<tr>
<td>Fragment Screws</td>
<td>Titanium alloy (ASTM F136)</td>
</tr>
<tr>
<td>Proximal Plate</td>
<td>Titanium alloy (ASTM F136)</td>
</tr>
</tbody>
</table>

3. **METHOD OF IMPLANTATION**

Please see the DR CAGE Surgical Technique Manual for a more-detailed description of the implantation procedures. The surgical technique includes the following general steps:

3.1. Reduce and stabilize fracture.
3.2. Template and size implant location.
3.3. Insert guide pin at entry point.
3.4. Prepare cavity for DR CAGE.
3.5. Insert and lock Expandable Scaffold.
3.6. Secure bone fragments with Fragment Screws into Expandable Scaffold.
3.7. Attach Proximal Plate.

4. **CONTRAINDICATIONS**

The DR CAGE should not be implanted in patients with:

- active infections at the operative site or other active systemic infection
- suspected or known sensitivity or allergies to Nickel or Titanium
- mental conditions that preclude cooperation with the rehabilitation regimen

5. **WARNINGS AND PRECAUTIONS**

**Warnings:**

5.1. Fracture fixation devices are neither intended to carry the full load of the patient, nor intended to carry a significant portion of the load for extended periods of time.

5.2. Do not use the DR CAGE with components from other manufacturers. Use only Conventus Orthopaedics devices.

**Precautions:**

5.3. Use care in handling and storage of implant components. Cutting, sharply bending or scratching the surface can significantly reduce the strength and fatigue resistance of the implant system. This, in turn, could induce cracks and/or stresses that could lead to fracture of the implants.
5.4. Additional surgical technique information is available upon request. The surgeon should be familiar with the devices, instruments and surgical technique before surgery.

5.5. Please refer to the surgical technique manual for proper implantation procedures.

5.6. The surgeon should take care with patient selection and avoid patients with insufficient quality of bone, an obliterated medullary canal or conditions which tend to retard healing, including but not limited to, blood supply limitations or previous infections.

5.7. Postoperative instructions to patients and appropriate nursing care are critical. Early weight bearing substantially increases implant loading and increases the risk of loosening, bending or breaking the device. Early weight bearing should only be considered where there are stable fractures with good bone-to-bone contact. Patients who are obese and/or noncompliant, as well as patients who could be pre-disposed to delayed union or non-union, must have auxiliary support.

5.8. Even after full healing, the patient should be cautioned that refracture is more likely with the implant in place and soon after its removal, rather than later when voids in the bone left by implant removal have been filled in completely.

5.9. Patients should be cautioned against unassisted weight-bearing activity without physician direction or medical release.

5.10. Postoperative care and physical therapy should be structured to prevent loading of the radius until stability is evident.

Potential adverse events may be related to surgery in general or the device. These may include, but are not limited to the following:

Potential adverse events related to any surgery: reactions to anesthesia, the anesthetic or other medications, bleeding, infection (both deep and superficial), ileus, blood vessel damage, nerve or soft tissue damage, atelectasis, pneumonia, hematoma, seroma, wound dehiscence or incisional hernia, urologic problems, embolism, anemia, colitis, thrombophlebitis, heart attack, stroke or death.

Potential adverse events related to the device: bending, cracking or fracture of the implant components, failure to achieve fracture healing, or loosening of the implant, limb shortening or loss of anatomic position with nonunion, malunion with rotation or angulation, irritational injury of soft tissues, including impingement syndrome, or tissue reactions which include macrophage and foreign body reactions adjacent to implants. Metal sensitivity or histological or allergic reaction resulting from implantation of a foreign material may occur. Additional surgery may be necessary for implant removal, repositioning or replacement.

The Expandable Scaffold and Delivery System of the DR CAGE are provided sterile. Each package should be inspected prior to use to ensure package integrity prior to use. Do not resterilize implants or single-use accessories if the package has been opened but the implant is not used.

Fragment Screws and the Proximal Plate are provided NON-Sterile in an autoclavable tray and must be sterilized prior to use according to the one of the following cycles:

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Minimum Exposure Time</th>
<th>Minimum Temperature</th>
<th>Minimum Drying Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gravity Autoclave</td>
<td>30 Minutes</td>
<td>132°C</td>
<td>20 Minutes</td>
</tr>
<tr>
<td>Pre-vacuum Autoclave</td>
<td>4 Minutes</td>
<td>132°C</td>
<td>20 Minutes</td>
</tr>
<tr>
<td>Pre-vacuum Autoclave</td>
<td>3 Minutes</td>
<td>135°C</td>
<td>20 Minutes</td>
</tr>
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All implants should be inspected prior to use for possible damage or defects. Any damaged or defective component should not be used and should be returned to the manufacturer. If an implant has been contaminated by blood or secretions, clean and steam sterilize the implant prior to returning it to the manufacturer.

The manufacturer is not responsible for any complications arising from incorrect diagnosis, incorrect choice of implant, incorrect operating techniques, the limitations of treatment methods or inadequate asepsis.

During the post operative phase, in addition to mobility and muscle training, it is of particular importance that the physician keeps the patient well informed about post-surgical activity requirements.
6. OPERATIVE PRECAUTIONS

The surgeon is to be thoroughly familiar with the DR CAGE, method of implantation, instruments and surgical technique.

Proper surgical procedures and techniques are the responsibility of the medical professional. Each surgeon must evaluate the appropriateness of the procedures based on his or her personal medical training and experience.

The implants are for single-implant use only. An explanted device must never be re-implanted.

7. MRI SAFETY INFORMATION

MRI Safety Information

MR Conditional
Non-clinical testing demonstrated that the DR CAGE is MR Conditional. A patient with this device can be scanned safely, immediately after placement 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 2.0-W/kg for 15 minutes of scanning (i.e., per pulse sequence)
- Normal Operating Mode of operation for the MR system

MRI-Related Heating

In non-clinical testing, the DR CAGE produced a temperature rise of less than 3°C under the scan conditions defined above in a 1.5-Tesla/64-MHz (Magnetom, Siemens Medical Solutions, Malvern, PA. Software Numaris/4, Version Syngo MR 2002B DHHS Active-shielded, horizontal field scanner) MR Scanner.

In non-clinical testing, the DR CAGE produced a temperature rise of 2°C under the scan conditions defined above in a 3-Tesla/128-MHz (Excite, HDx, Software 14X.M5, General Electric Healthcare, Milwaukee, WI) MR Scanner.

Artifact Information

MR image quality may be compromised if the area of interest is in the exact same area or relatively close to the position of the DR CAGE. Therefore, optimization of MR imaging parameters to compensate for the presence of this device may be necessary.

The image artifact extends approximately 10mm from the device, when scanned in nonclinical testing using a gradient echo pulse sequence at 3-Tesla (Excite, HDx, Software 14X.M5, General Electric Healthcare, Milwaukee, WI).
8. **PRODUCT COMPLAINTS**

Any healthcare professional (e.g., a surgeon using the product) who has a complaint or who has experienced any dissatisfaction in the quality, identity, reliability, safety, efficacy, and/or performance of any Conventus product should notify Conventus Orthopaedics, or, where applicable, their distributor. In the event of serious incident, or risk of serious incident, having resulted in, or may potentially result in, the death or severe deterioration in the state of health of a patient or user, Conventus Orthopaedics or the distributor must be notified as soon as possible. When filing a complaint, please provide the component(s) reference number (model number), manufacturing lot number(s), your name and address, and the nature of the complaint in full detail, as well as notification of whether a written report is requested.

9. **STORAGE INSTRUCTIONS**

Store in a cool dry place and keep away from direct sunlight. Prior to use, inspect instruments and tray for signs of tampering.

10. **WARRANTY**

THE COMPANY REPRESENTS AND WARRANTS THAT THE DR CAGE WILL CONFORM TO THE COMPANY’S SPECIFICATIONS AND COMPLY WITH ALL APPLICABLE FDA STANDARDS, AS SUCH STANDARDS MAY BE AMENDED FROM TIME TO TIME. THE COMPANY MAKES NO OTHER EXPRESS OR IMPLIED WARRANTIES REGARDING THE DR CAGE.

11. **FURTHER INFORMATION**

For further Information, please contact:

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Maple Grove, MN 55369
Phone: 763-515-5000
Fax: 763-315-4980
www.conventusortho.com
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Caution: Federal Law (USA) restricts this device to sale by or on the order of a physician.

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<thead>
<tr>
<th>REF</th>
<th>Model /Part Number</th>
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<td>Material</td>
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<td>🔄️</td>
<td>Contents are Non-Sterile</td>
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<tr>
<td>🛠️</td>
<td>Manufacturer</td>
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<tr>
<td>MR</td>
<td>MR Conditional</td>
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