Patient-specific lateral lumbar interbody with independent *in-situ* adjustment of height & lordosis

**Patient-specific**
- *In-situ* parallel expansion up to 16.1mm
- Independent lordotic adjustment up to 30 degrees*

**Designed for Surgical Efficiency**
- Single instrument for *in-situ* height and lordotic adjustment
- Decreased need for multiple trials

**Streamlined Inventory**
- Five implants to treat a wide range of patients
- Individual sterile pre-packaged implants

**Designed to Improve Clinical Outcomes**
- Patient-specific fit and lateral MIS approach minimizes trauma
- Lateral approach reduces tissue disruption, anesthesia time and post-operative complications

510(k) K181531
Sagittae® is a lateral lumbar interbody with independent height and lordosis adjustment designed to restore sagittal balance.

Sagittae is available in the following sizes:

<table>
<thead>
<tr>
<th>Length (mm)</th>
<th>Width (mm)</th>
<th>Height Range (mm)</th>
<th>Lordosis Range</th>
<th>Bone Graft Volume (cm³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>42.0</td>
<td>21.5</td>
<td>8.4-17.1</td>
<td>0-30</td>
<td>1.4-6.1</td>
</tr>
<tr>
<td>46.0</td>
<td>21.5</td>
<td>8.4-17.1</td>
<td>0-30</td>
<td>1.6-6.6</td>
</tr>
<tr>
<td>50.0</td>
<td>21.5</td>
<td>8.4-17.1</td>
<td>0-30</td>
<td>1.8-7.2</td>
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<tr>
<td>54.0</td>
<td>21.5</td>
<td>8.4-17.1</td>
<td>0-30</td>
<td>2.0-7.9</td>
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<tr>
<td>58.0</td>
<td>21.5</td>
<td>8.4-17.1</td>
<td>0-30</td>
<td>2.2-8.6</td>
</tr>
</tbody>
</table>

Fig 1. Sagittae Inserter

Fig 2. Sagittae interbody placement using lateral MIS approach

Fig 3. Sagittae height and lordosis adjusted in-situ

Fig 4. Fully expanded Sagittae interbody

For more information about Sagittae, please contact your SpineEX representative, or email: info@spineEXinc.com

*These devices are intended to be used with supplemental fixation systems that have been cleared for use in the lumbosacral spine (e.g. posterior pedicle screw and rod systems, anterior plate systems, and anterior screw and rod systems). Hyperlordotic interbody devices (≥20° lordosis) must be used with at least anterior supplemental fixation.

Please refer to package insert for a detailed device description, indications, contraindications, warnings, precautions and other important information.