LINK SLED Prosthesis
with MITUS Instrument Set

Surgical Technique
## Explanation of Pictograms

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<th>Meaning</th>
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<tr>
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</tr>
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<td>Material (number)</td>
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<td><img src="image" alt="Rx only" /></td>
<td>Rx only</td>
</tr>
<tr>
<td><img src="image" alt="Caution" /></td>
<td>Caution: Federal law restricts this device to sale by or on the order of a physician</td>
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</tbody>
</table>
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LINK SLED Prosthesis
with MITUS Instrument Set

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Important Information
The successful design of LINK’s Unicondylar Sled Prosthesis which was originated in 1969 has remained unchanged since its last modification in 1981.

Further advantages:
- high joint mobility
- short recovery period

The design of the femoral component preserves bone substance and permits femoral resurfacing. This treatment is therefore available as a fall-back option.

The instruments and the surgical technique are regularly optimized to ensure ease of use and reliable implantation.

The LINK Unicondylar Sled Prosthesis is available in four sizes.
Femoral Components
The large radii of the femoral surface distributes the contact stress more homogenously. The globular structure of the concave inner surface of the sled provides optimal bonding between implant and cement. The design incorporates two posts whose shape and alignment aid in positioning the sled. The implant is easy to remove should revision become necessary.

Tibial Plateaus
The tibial plateaus can be used medially as well as laterally owing to their symmetrical shape. The sizing is adapted to the anatomical shape of the head of the tibia. Two designs are available:

- **Type all-polyethylene** (non metal-backed)
  This design is available in four heights and four diameters. The structured underside allows a very good interface between implant and bone cement.

- **Type metal-backed**
  In this design, the tibial plateaus are available in three heights and three diameters. The globular structure on the underside of the plateau offers optimum bonding between the implant and bone cement.

**LINK PorEx (TiNbN = Titanium Niobium Nitride)**
Surface Modification
The LINK PorEx surface modification results in a ceramic-like surface, which significantly reduces chrome and nickel ions release.\(^1\)

Thanks to its outstanding hardness, abrasion properties similar to ceramics and larger wetting angle, in contact with liquids the LINK PorEx surface against UHMWPE has a lower coefficient of friction compared to CoCrMo surfaces.\(^1\)

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\(^1\) Internal technical report: Study of the influence of TiNbN-coating on the ion release of CoCrMo-alloys in SBF buffer simulator testing
Rünow Minimally Invasive Surgical Technique

For implantation of a sled protheses it is essential to select the correct indication. The concept is based on the fact that in early stages of knee osteoarthritis (OA) the cartilage damage is limited to a single compartment within the knee joint.

The design of the LINK Sled Prosthesis ensures that only minimal bone resection is required when preparing the bone to receive the femoral and tibial components. This preserves high-quality bone, particularly the hard sub-chondral bone, which is important for secure long-term fixation of the implant.

The Tibial Saw Guide supports resection according to anatomical conditions and ensures precise, reproducible bone cuts.

The MITUS Instrument Set offers distinct advantages to the surgeon:

- minimal bone resection
- full control over the level of tibial resection
- opportunity to try out different sizes using trial implants
- option to perform the surgery using either conventional or minimally invasive surgical techniques
- medial or lateral use of instruments possible
Rünow Minimally Invasive Surgical Technique

Two different forms of surgical approach can be used

Conventional Approach: through a midline or a medial parapatellar skin incision. The joint cavity is reached via a medial parapatellar incision and splitting of the quadriceps tendon. The patella is everted laterally.

Minimally Invasive Approach: through a short parapatellar skin incision. The capsular incision is also parapatellar allowing access to the joint with minimal disturbance of the extensor mechanism and without dislocating the patella.

The minimally invasive technique reduces complications and can be performed with great precision provided the LINK instruments are used correctly.
Patient Positioning

The limb is placed in a thigh support with 45° flexion of the hip. The leg is hanging down. It should be possible to flex the knee at least 120°. When using a medial incision a lateral thigh support is needed.

The operation is performed with the surgeon sitting in front of the flexed knee. The other leg is placed in a leg support leaving plenty of space for the surgeon and the assistant. The operation is performed in a bloodless field.
Patient Positioning

With the knee flexed 90°, a medial parapatellar incision is made starting at the margin of the vastus medialis 2–3 cm medial to the patella and extending distally and diagonally to the tibial tuberosity.

A medial parapatellar capsule incision is made. For better visualization the incision is angulated in its proximal part. The vastus medialis is detached. The capsule is released from the tibia almost to the front of the medial collateral ligament. The meniscus is removed. Partial excision of the retropatellar fat pad is necessary to gain better exposure of the intercondylar notch.

A retractor is placed in the lateral recess, allowing inspection of this compartment. To examine the patellar articulation, the knee is extended. If there are any doubts preoperatively about the condition of the other compartments diagnostic arthroscopy or MRI can be performed prior to the operation. After inspection, the retractor is placed in the intercondylar notch and the curved retractor behind the femoral condyle, to get a full view of the medial compartment.
The purpose of the LINK Unicondylar Sled Prosthesis is to restore the damaged joint surfaces and the mechanical axis; a slight under-correction is desirable.

The Tibial Saw Guide allows the surgeon to determine and achieve the desired cutting depth precisely and to control the cutting in the frontal and the sagittal planes. The Saw Guide can be used with either a minimally invasive technique or the traditional exposure.
In knee replacement surgery by the traditional technique, the deepest point and the most damaged area of the tibial plateau are taken as the basis for determining the depth of the tibial resection. The depth of the resection is then highly dependent on the surgeon’s experience. Often further resection is needed or the height of the Tibial Plateau must be changed to obtain the desired alignment and stability of the knee. The best aid to determining the depth of the horizontal cut is weight-bearing radiographs of the knee and pre-operative observations of the degree of cartilage damage. These allow a slight undercorrection of only a few degrees of varus to be achieved. The analysis of the weight-bearing radiographs is based on the classification of Ahlbäck.

The proposed resection depths are based on the use of a 9-mm high Tibial Plateau.

**Grade I**  The joint space is reduced by one-half. The cartilage of the tibial condyle is preserved but reduced in height. The Cutting Platform should be adjusted to 11 mm depth. The stylus is placed at the deepest point of the remaining cartilage of the tibial condyle.

**Grade II**  Total loss of the cartilage on both the femoral and the tibial condyles. The Cutting Platform should be adjusted to 9 mm depth and the stylus placed at the deepest point of the exposed bone of the tibial condyle.

**Grade III**  Half a centimeter bone attrition of the femoral and tibial condyles on the frontal view weightbearing radiograph. The Cutting Platform should be adjusted to 7 mm and the stylus placed at the border between the exposed and the eroded bone.

The stylus is not placed at the level of the planned surface of the Tibial Plateau. In Grade I the surface of the Tibial Plateau will be lower than the surface of the tibial condyle, and correspondingly in Grade III the surface of the Tibial Plateau will be higher than the surface of the damaged tibial condyle.
Tibial Resection

<table>
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<tr>
<th>Height of Tibial Plateaus</th>
<th>Resection Height</th>
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<td>9 mm</td>
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</tr>
<tr>
<td>11 mm</td>
<td>13</td>
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Table 1:
Depth of the tibial resection (mm) in relation to the chosen height of the Tibial Plateau.

<table>
<thead>
<tr>
<th>Resection Height</th>
<th>Height of Tibial Plateaus</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Grade I</td>
</tr>
<tr>
<td>7 mm</td>
<td>–</td>
</tr>
<tr>
<td>9 mm</td>
<td>7</td>
</tr>
<tr>
<td>11 mm</td>
<td>9</td>
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</table>

Table 2:
Height of the Tibial Plateau (mm) in relation to the depth of the tibial cutting.

Table 1
When a 7-mm high Tibial Plateau is used, the tibial resection depth should be 9 mm in Grade I knees and 7 mm in Grade II knees. Because the construction of the Tibial Saw Guide does not permit less than 7 mm resection depth between the tip of the stylus and the Cutting Platform a 7-mm Tibial Plateau cannot be used in Grade III osteoarthritis, and such knees must therefore be undercorrected. According to suggestions given above, the depth of the resection when using an 11-mm Tibial Plateau will be 13 mm in Grade I, 11 mm in Grade II, and 9 mm in Grade III. These resection depths will unnecessarily be too deep and will remove more bone than necessary.

Table 2
It is convenient to use the same resection depth independent of the degree of cartilage and bone damage. This means that a resection depth of 9 mm in relation to the tibial surface is needed in order to use a 7-mm Tibial Plateau in Grade I, a 9-mm Tibial Plateau in Grade II and an 11-mm Tibial Plateau in Grade III to achieve the same degree of alignment.
The clamp of the Tibial Saw Guide is placed at the level of the ankle directly proximal to the malleoli.

**The posterior slope of the Tibial Component**

Note that the Cutting Platform has a posterior slope of 6° in relation to the long axis of the Guide. The Tibial Saw Guide should be adjusted in the vertical plane parallel to the long axis of tibia by moving the vertical rod ventrally. In most cases the Guide needs to be moved 20–25 mm anteriorly to obtain the required posterior angle of a 6°. **Lock Screw A.** Orient the posterior slope to the natural preoperative situation, so that the biomechanics of the individual patient are not changed. The resection can be checked for control purposes. It should have the same thickness ventral-dorsal parallel.

**Attention:** Kinematic results suggest that 5° to 7° of posterior slope is preferable, and that excessive posterior slope (> 7°) should be avoided.

**The varus-valgus alignment of the Tibial Component**

The varus-valgus alignment of the Tibial Component can be adjusted by placing the distal fixation of the long rod beneath the actual tibia condyle. In women the rod is moved approximately 20–25 mm and in men 25–30 mm from the center to achieve a cutting surface perpendicular to the long axis of tibia. The horizontal slope is controlled with the Alignment Rod. **Lock Screw B.**

**Warning:** Overcorrection in valgus should be avoided under all circumstances. Position the tibia in 0° to 3° varus. Place the Alignment Rod through the Tibial Cutting Block.
Tibial Resection

The Eminentia Saw Guide (E) is placed close and parallel to the eminentia along the planned sagittal cut.

Attention: The eminentia intercondylaris and in particular the insertion of the anterior cruciate ligament serve for orientation. The sagittal cut should be made just medial to the ACL attachment point on the tibial spine in order to maximize the size of the tibial base.

There are Cutting Platforms (P) for the medial as well as the lateral compartments. The cutting depth can be set between 7 and 13 mm by using a Screwdriver in the adjustment Hole (B). The Cutting Platform is secured and locked with Screw (A).

The Tibial Saw Guide is fixed with a Fixation Pin in the central hole of the platform. The Pin is angulated centrally towards the eminentia. A second Fixation Pin is placed in the Tibial Saw Guide to secure the position.
Tibial Resection

Bone Cuts

The **vertical cut** is performed along the Eminentia Saw Guide. The **horizontal cut** is guided by the Cutting Platform.

The resected Tibial Plateau and remaining parts of the meniscus are then removed.
Tibial Resection

Depending on implant selection a **Template** is used for the sizing of the Tibial Plateau. Both are available in three sizes (45, 50 and 55 mm).

The size of the Tibial Plateau in the sagittal plane is determined by placing the hook of the Template behind the tibial condyle. If the anterior part of the Template is in alignment with the anterior border of the tibia, that is the right size.

The size must be checked medially to ensure there is no medial overhang.

**Attention:** It is important to achieve maximal coverage of the tibial plateau. Determine the tibial component as large as possible. However, an overhang, especially anteriorly, should be avoided.
Femoral Resection

Do not remove the Tibial Saw Guide during the preparation for the Femoral Component.

Begin the preparation of the femoral condyle by cutting 3–5 mm of its posterior aspect to remove undamaged cartilage.

Resect central and medial osteophytes, with attention to osteophytes behind the medial collateral ligament.
Femoral Resection

There are four sizes of the Femoral Components (40, 46, 52 and 60 mm) and corresponding Drill Guides to determine the correct size. The selected femoral Drill Guide is placed centrally on the femoral condyle and fixed with two short Fixation Pins.
Surgical Technique

Trial Reduction

Drill the anchoring holes. If it is difficult to drill the lower hole at 100-110° of flexion of the knee, the Femoral Drill Guide is either too large or has been placed too far dorsally. Either change its position or chose a smaller Drill Guide.

Mark the borders of the Drill Guide. Remove any cartilage inside the area marked for the Femoral Component.
Corresponding to the Femoral Drill Guides are four **Femoral Trial Sled Prostheses**. Before trialing the chosen size, use a chisel or a saw to prepare a groove between the two anchoring holes. Place the Femoral Trial Sled Prosthesis using the **Inserting Forceps**.  
**Attention:** Orient the pegs as shown in Figures A & B.

Test knee flexion and extension to make certain that the Femoral Trial Sled Prosthesis does not make contact with the patella at any point during the movement. If it does, remove any patellar bone that contacts the Femoral Trial Sled Prosthesis.
The **Tibial Plateaus** (all-polyethylene) are available in 4 heights (7, 9, 11 and 13 mm) and the Tibial Plateaus (metal-backed) in 3 heights (9, 11 and 13 mm).

With the Femoral Trial Sled Prosthesis in place, a 9-mm **Tibial Trial Plateau** is positioned. This is easiest when the knee is flexed at least 90°. Some valgus load may be needed. If the Tibial Component has a tendency to tilt anteriorly, the posterior angle of slope is too small. This can be corrected with a rasp.

Move the knee through its entire range of motion to check joint stability. Select the Tibial Component height that most closely restores the natural tension of the ligaments. With valgus loading of the knee joint, it should be possible to open the medial joint space 1-2 mm.

**Warning:** Overcorrection in valgus should be avoided under all circumstances. Position the tibia in 0° to 3° varus. It is important to ensure a slight under correction of the limb alignment and have appropriate ligamentous tension restored (2-3 mm of laxity) in flexion and extension.
Surgical Technique

Trial Reduction

If the knee is too tight, remove the Tibial Trial Component and the Fixation Pin in the Cutting Platform and loosen Screw (A). Deepen the resection by lowering the platform to the appropriate level by turning Screw (B) using a Screwdriver. As a rule 1-mm increase in resection depth increases varus angulation by 2 degrees.

Secure the Cutting Platform by tightening Screw (A) and stabilize it with a Fixation Pin through one of the unused holes in the Cutting Platform.

Perform the cut and repeat the trial by using the same height of the Tibial Trial Component.
Cementation

Prepare the space for the keel of the Tibial Plateau (metal-backed), place the head of the **Cancellous Bone Compressor** into the recess of the tibial Template and impact it using the **Impactor**.

The keel of the Tibial Plateau (all-polyethylene) is larger. To prevent fractures of the tibial condyle remove some bone with a chisel before impacting the Bone Compressor.

Whichever Tibial Plateau is being used, the tibial surface needs to be protected during the compression of the bone with the tibial Template, which is laid on the sawing platform. Test that the final choice of Tibial Plateau fits and can be placed easily. Some valgus stress will be needed. The keel slot may be extended anteriorly if necessary.
Cementation

Preparation
Warning:
Good fixation of the implant is a prerequisite to achieve long-term success. Cementing technique is one factor that plays an important role in this respect. Therefore the following instructions should be carefully considered.

In sclerotic bone, multiple holes should be drilled with a small drill (Max diam 3.0 mm-drill pins can be used as an alternative) to ensure better bone cement interdigitation. Due to the preparation technique, this is particularly important for the femoral condyle. Cleanse all cement-receiving bone surfaces thoroughly using pulse lavage and dry with a clean, dry lap sponge.

General remarks
The tibial and femoral components should be cemented separately from each other in two stages. This ensures that there is sufficient time to position the component, remove excess bone cement, and allow it to harden without inadvertently manipulating the implant-bone cement-bone interface.
Prepare the bone cement according to the manufacturer’s instructions. Apply the bone cement to both the back of the implant and to the bone. Beginning with the tibial component, carefully apply the bone cement evenly to ensure a homogenous cement mantle.

Attention: Consider applying bone cement at the vertical plane too.
Maintain a steady pressure with the tibial impactor during curing. Then proceed to cement the femoral component.

Option: Start the implantation with the femoral component.

Cementation of Tibial Component
Apply a thin layer of cement over the entire underside of the tibial component. The cement should just overfill the bead structure on the underside of the tray, up to 1 mm proud posteriorly and 2 mm proud anteriorly. Apply cement to the tibia and pressurize the cement, striving for penetration of 3-4 mm.
Use of a cement gun/cartridge equipped with a pressurizing nozzle is recommended to deliver and pressurize cement into the prepared holes and across the flat surface. Alternatively, cement may be applied manually and pressurized into the bone using a flat osteotome.

Cementation of Femoral Component
Apply bone cement to the back of the Femoral Component. In addition, fill both drill holes for the fixation pegs with bone cement. Position the femoral component using Inserting Forceps, and insert both pegs into the prepared drill holes. Seat the Femoral Component with the Femoral Impactor, maintaining a steady pressure on the Femoral Impactor during curing.
**Implants**

**Femoral Components**

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<th>REF CoCrMo</th>
<th>REF CoCrMo/LINK PorEx*</th>
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* LINK PorEx: TiNbN = Titanium Niobium Nitride; surface modification (gold color).

**Tibial Plateaus – metal-backed**

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<th>REF CoCrMo/LINK PorEx*</th>
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* LINK PorEx: TiNbN = Titanium Niobium Nitride; surface modification (gold color).
Important information:
Tibial Components of 7 mm height offer the advantage of greater bone preservation and allow a good range of motion. The suitability of these particular components have to be medically indicated. They are not suitable for obese or very active patients.

Tibial Plateaus – All-polyethylene

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**MITUS Instrument Set**

Minimally Invasive Surgical Technique for LINK Unicondylar Sled Prosthesis

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<th>Description</th>
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<td>Set, complete in 2 standard containers, on 3 trays with storage inserts,</td>
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*Fittings: How to order: 317-649/08B = Hudson fitting*

- B Hudson
- C Harris
- D A-O
- E Jacobs
- H Zimmer
Instruments

Lower Tray, Container 1

1 15-2200/02 Lower Tray (Container 1), empty, 550 x 265 x 50 m
Drill and Saw Guide (templates) for tibial plateaus all-polyethylene

2 15-2201/55 55 mm

3 15-2201/50 50 mm

4 15-2201/45 45 mm

5 15-2040/05 Sled Impactor, for sled prosthesis metal-backed, 170 mm

6 15-2201/70 Curette, to remove excess cement

7 15-2201/71 Spatula, double end, to remove excess cement

8 15-2040/03E* Twist Drill with stop, Ø 5.5 mm, 160 mm, fittings optional (see page 25)*
Drill and Saw Guide (templates) for tibial plateaus metal-backed

9 15-2202/55 55 mm

10 15-2202/50 50 mm

11 15-2202/45 45 mm

12 15-2040/06 Plateau Impactor, 250 mm

13 15-2105 Chip Chisel, 15 mm wide, 240 mm

14 15-2201/16 Lambotte Osteotome, width 9 mm

15 15-2201/17 Lambotte Osteotome, width 11 mm

16 15-2102/03 Lambotte Osteotome, width 15 mm

17 15-2040/02E* Twist Drill, Ø 3.0 mm, 160 mm, fittings optional (see page 25)*
1  15-2200/03  Upper Tray (Container 1), empty, 550 x 265 x 50 mm

Drill Guides for sled prostheses

2  15-2201/60  large

3  15-2201/52  medium

4  15-2201/46  medium-small

5  15-2201/40  small

6  15-2042  Inserting Forceps, for tibial plateaus all-polyethylene and trial plateaus, 215 mm

Cancellous Bone Compressors, for tibial plateaus all-polyethylene

7  15-2201/14  Ø 50-55 mm

8  15-2201/15  Ø 45 mm

9  15-2201/19  Cancellous Bone Compressor, for tibial plateaus metal-backed, Ø 45-55 mm

Trial Sled Prostheses

10  15-2021/05  large

11  15-2021/04  medium

12  15-2021/03  medium-small

13  15-2021/02  small

14  15-2201/12  Fixation Pins, for drill guides, Ø 2 mm, 60 mm

15  15-2201/53  Fixation Pin, to stabilize the drill guide, Ø 5.4 mm, 50 mm (4 ea.)

16  15-2201/13  Holding and Inserting Forceps, for drill guides

17  15-2040/09  Inserting Forceps, for tibial plateaus metal-backed

18  15-2040/08  Set of Trial Plateaus, on storage tray, Ø 45, 50, 55 mm, heights: 7, 9, 11, 13 mm (12 ea.)
# Instruments

## Tray, Container 2

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>15-2200/01 Tray (Container 2), empty, 550 x 265 x 50 m</td>
</tr>
<tr>
<td>2</td>
<td>317-586 Inserter/Extraction Forceps, for fixation pins, 210 mm</td>
</tr>
<tr>
<td>3</td>
<td>15-2201/18 Extractor, for fixation pins, to be used with 317-586</td>
</tr>
<tr>
<td>4</td>
<td>317-585/95 Fixation Pins, Ø 3 mm, 95 mm (6 ea.)</td>
</tr>
<tr>
<td>5</td>
<td>15-2201/32 Eminentia Saw Guides, left, height A</td>
</tr>
<tr>
<td>6</td>
<td>15-2201/37 Eminentia Saw Guides, left, height B</td>
</tr>
<tr>
<td>7</td>
<td>15-2201/33 Eminentia Saw Guides, right, height A</td>
</tr>
<tr>
<td>8</td>
<td>15-2201/38 Eminentia Saw Guides, right, height B</td>
</tr>
<tr>
<td>9</td>
<td>15-2201/34 Tibial Alignment Device, extramedullary</td>
</tr>
<tr>
<td>10</td>
<td>15-2201/35 Stylus</td>
</tr>
<tr>
<td>11</td>
<td>15-2201/39 Spacer Bolt, to 15-2201/31</td>
</tr>
<tr>
<td>12</td>
<td>15-2201/11 Retractor</td>
</tr>
<tr>
<td>13</td>
<td>15-2201/10 Inserting Forceps, for trial sled prostheses</td>
</tr>
<tr>
<td>14</td>
<td>317-538/01 Flexible Belt, 495 mm</td>
</tr>
<tr>
<td>15</td>
<td>15-2201/31 Tibial Saw Guide Base, adjustable</td>
</tr>
<tr>
<td>16</td>
<td>15-2201/36 Alignment Rod, transversal, 200 mm</td>
</tr>
<tr>
<td>17</td>
<td>10-5373 Hex Screwdriver, hex 2.5 mm, 180 mm</td>
</tr>
<tr>
<td>18</td>
<td>317-648 Universal Wrench, hex 6.0 mm, 140 mm</td>
</tr>
<tr>
<td>19</td>
<td>130-611 Impactor, 280 mm</td>
</tr>
</tbody>
</table>
Instruments

Additional Instruments (not included in Instrument Set, complete)

Trial Tibial Plateaus, Ø 58 mm, suitable for tibial plateaus all-polyethylene (without metal-backed)

<table>
<thead>
<tr>
<th>REF</th>
<th>Height mm</th>
<th>Width mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>15-2047/13</td>
<td>7</td>
<td>31</td>
</tr>
<tr>
<td>15-2047/14</td>
<td>9</td>
<td>31</td>
</tr>
<tr>
<td>15-2047/15</td>
<td>11</td>
<td>31</td>
</tr>
<tr>
<td>15-2047/16</td>
<td>13</td>
<td>31</td>
</tr>
</tbody>
</table>

Trial Tibial Plateaus Height 8 mm, suitable for tibial plateaus with metal-backed

<table>
<thead>
<tr>
<th>REF</th>
<th>Height mm</th>
<th>Width mm</th>
<th>Ø mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>15-2040/33</td>
<td>8</td>
<td>22.5</td>
<td>45</td>
</tr>
<tr>
<td>15-2040/34</td>
<td>8</td>
<td>25.0</td>
<td>50</td>
</tr>
<tr>
<td>15-2040/35</td>
<td>8</td>
<td>27.5</td>
<td>55</td>
</tr>
</tbody>
</table>

15-2048/04
Storage Tray, separate
for all-polyethylene trial tibial plateaus, Ø 58 mm

15-2048/05
Storage Tray, separate
for trial tibial plateaus height 8 mm

15-2201/58
Drill and Saw Guide (templates)
for tibial plateaus all-polyethylene, Ø 58 mm
The tibial saw guide consists of a base with one cutting platform for medial resection and another for lateral resection. The stylus is inserted into a hole on the cutting platform. The spacer bolt is fixed on the opposite side. To protect the intercondylar eminence, eminentia saw guides are available.

The adjustable extramedullary tibial saw guide base determines the correct axial alignment. A transversal alignment rod helps to align the saw guide horizontally. The saw guide is fixed distally using a plastic connector. The saw guide is secured proximally with fixation pins.
Important Information for X-ray Investigations

X-ray investigations
X-ray images can be used to evaluate implant positioning post-operatively. Images taken from certain angles can create the impression that the implant has broken.

Note:
The LINK tibial plateau metal-backed is delivered as one piece, i.e. the polyethylene component and the metal component are pre-assembled as a single unit. The manufacture of the components has remained unchanged since today. For secure connection the polyethylene engages with a mechanical coupling device.

These technical specifications can lead X-ray images taken from certain angles to appear distorted, which may give the impression that the tibial plateau is broken. Examples of such distorted images are shown below:

As a broken tibial plateau is most unlikely, the diagnosis must be verified with additional X-ray images. Verification: Rotation of the tibia ensuring strictly lateral alignment for the follow-up X-ray.
## X-ray Templates, 110% actual size, one sheet

<table>
<thead>
<tr>
<th>REF</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>15-2021/10</td>
<td>for Unicondylar Sled Prosthesis 15-2020/40 to 15-2020/60</td>
</tr>
<tr>
<td>15-2021/14</td>
<td>for Tibial Plateaus, metal-backed 15-2030/02 to 15-2030/13 and 15-2230/02 to 15-2230/13</td>
</tr>
<tr>
<td>15-2021/13</td>
<td>for Tibial Plateaus, all-polyethylene 15-2028/01 to 15-2028/16</td>
</tr>
</tbody>
</table>
The **LINK SLED Knee System** is indicated for patients with severe joint diseases with limitation of mobility due to degenerative, rheumatoid or post-traumatic arthrosis or arthritis. Joint fractures which disallow an osteosynthetic reconstruction. These devices are intended for cemented use only.

**The LINK SLED Knee System is indicated for the following conditions:**

1) Unicondylar arthrosis by intact ligaments including both cruciate ligaments
2) Valgus/Varus deformities <10°

**Contraindications:**

1) Acute or chronic infections, local and systemic
2) Distinctive muscular, nerve, vascular or other diseases which put the affected limb at risk
3) Insufficient / inadequate bone mass- or quality which prevents a stable anchor of the prosthesis

**Relative Contraindications:**

1) Adiposity
2) Insufficient collateral ligaments
3) Insufficient musculature
4) Lacking or foreseeable not assured compliance
5) Foreseeable overload of joint prosthesis
Please note the following regarding the use of our implants:

1. **Choosing the right implant is very important.**
   The size and shape of the human bone determines the size and shape of the implant and also limits the load capacity. Implants are not designed to withstand unlimited physical stress. Demands should not exceed normal functional loads.

2. **Correct handling of the implant is very important.**
   Under no circumstances should the shape of a finished implant be altered, as this shortens its life span. Our implants must not be combined with implants from other manufacturers. The instruments indicated in the Surgical Technique must be used to ensure safe implantation of the components.

3. **Implants must not be reused.**
   Implants are supplied sterile and are intended for single use only. Used implants must not be used again.

4. **After-treatment is also very important.**
   The patient must be informed of the limitations of the implant. The load capacity of an implant cannot compare with that of healthy bone!

5. **Unless otherwise indicated, implants are supplied in sterile packaging.**
   Note the following conditions for storage of packaged implants:
   - Avoid extreme or sudden changes in temperature.
   - Sterile implants in their original, intact protective packaging may be stored in permanent buildings up until the “Use by” date indicated on the packaging.
   - They must not be exposed to frost, dampness or direct sunlight, or mechanical damage.
   - Implants may be stored in their original packaging for up to 5 years after the date of manufacture. The “Use by” date is indicated on the product label.
   - Do not use an implant if the packaging is damaged.

6. **Traceability is important.**
   Please use the documentation stickers provided to ensure traceability.

7. **Further information** on the material composition is available on request from the manufacturer.

Follow the instructions for use!
This publication is intended for distribution in the U.S. only.