Endo-Model® – M
Modular Knee Prosthesis System with Segmental Bone Replacement Components

Surgical Technique
Endo-Model®—M
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with Segmental Bone Replacement Components

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Important information
The Endo-Model® – M Modular Intracandylar Total Knee Joint Prosthesis is an additional version of the LINK® Endo-Model® Rotational and Hinge Knee System.

Based on the same low friction principle, the rotational movement of this prosthesis passes smoothly through a pivot point in the physiological region. The Endo-Model® – M Rotation Knee Joint Prosthesis permits flexion of the joint up to 165°. Due to the special shape of the tibial contact surfaces and physiological rotation, the kinematics of this prosthesis allow for a cushioned transmission of force. The hinge knee prosthesis permit only flexion of the joint up to 165°, without rotation.

With every step, and especially when falling, torsional stresses are transmitted to the prosthetic anchorage which adversely affects the longevity of the cement interface. The cushioned transmission of forces, made possible by the design features, provides a dampened impact upon the boundary layer of the cement interface. The resection required during implantation of the Endo-Model® – M Knee Joint Prosthesis amounts to only 14 mm in the tibia-femur joint plane. With the medium sized intracandylar component only 30 mm wide, there is usually ample bone mass left in the event a revision is necessary. Normally, the resection is smaller compared with a total knee implant. Design and dimensioning of the rotation knee joint prosthesis significantly simplify the surgical procedure. Mounting of the femoral and tibial components is a simple task, requiring only one special introducer instrument for the UHMWPE Plateau. Both components are linked by the special anti-luxation device of the plateau without reducing the motional and rotational sequences. Implantation is facilitated by a small number of easily manipulated instruments. The hinge knee prosthesis is linked by an axis mechanism.

Flexion and rotation of the knee prosthesis occurs in a cross joint. Hyperextension amounts to 3°. The compromise axis lies in the region of the physiological pivot point. Flexion of up to 165° is possible. Often during endoprosthetic replacement of the knee joint, an advancement of the patella or of the patella bearing surface is observed. By displacing the femoral component dorsally relative to the tibial axis, the natural range of motion is also preserved in the patello-femoral joint. This protects against progression of retropatellar arthrosis.

Rotation of the prosthesis terminates in the extended position by form closure and assures a secure posture.
The rotational option increases continuously with flexion. This rotation is limited primarily by the capsule ligament apparatus. The body weight, bearing on the joint, elastically dampens further rotation. The femoral component of the Endo-Model® – M Total Knee Joint Prosthesis has a normal valgus position of 6°.

Both prosthesis components are broadly supported on the respective knee joint surfaces so that the compression strength of the cancellous bone in the femur and tibia, is not exceeded. The flanges of the femoral component are anatomically shaped. Its ventral depression provides a smooth transition from the implant to the trochlear groove.

The modular prosthesis stems are available in cementable and cementless versions (with smooth surface or longitudinal ribs respectively). To achieve a central position within the medullary canal, the tips of the cemented stems are fitted with star shaped UHMWPE caps. Direct contact of the metal stem with the inner wall of the bone is thereby prevented.

The stems are supplied in lengths of 50 up to 280 mm. Special femoral segments for revision surgery of resurfacing knee implants (reconstruction of condyles) and for tumor cases are also available. It is absolutely necessary to use these segments only in combination with longer stem.


## Assembly: Plateau with anti-luxation device

After cementation of tibial and femoral components the UHMWPE plateau is removed from the tibial tray by loosening the trial screw. With the knee in flexion both components are assembled.

The tibial plateau is attached to the introducer and slid between the femoral and tibial components so that its medial lip grabs over the flange of the femoral bushing. Care must be taken that the dovetailed medial and lateral parts fit into the groove at the posterior rim of the metal tibial tray (fig. B).

In this position the UHMWPE plateau is pressed down into the metal tray and firmly fixed by the self-locking screw.

Implanted Endo-Model® Modular Knee Prosthesis.
The Modular Stems are joined to the femoral and tibial components by a cone assembly. To secure the fixation two opposite tongues at the stems are inserted into the medial and lateral grooves at both the tibial and femoral components.

By tightening the locking screw (2) located in the taper (3) of the tibial respectively femoral component its pointed tip presses the stem (1) firmly onto the taper. A counter screw (4) secures the stem locking screw against loosening. The screw fixation is performed medially.
### Indications / Contraindications

<table>
<thead>
<tr>
<th></th>
<th>Rotational version</th>
<th>Hinged version</th>
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<tbody>
<tr>
<td><strong>General Indications</strong></td>
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<tr>
<td>• Severe joint diseases with limitation of mobility due to degenerative, rheumatoid or post-traumatic arthrosis or arthritis. Joint fractures which disallow an osteosynthetic reconstruction.</td>
<td>X</td>
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<tr>
<td><strong>Indications</strong></td>
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<tr>
<td>• Bone necroses</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Bicondylar arthrosis by partly damaged collateral ligaments</td>
<td>X</td>
<td>-</td>
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<tr>
<td>• Bicondylar arthrosis by completely damaged ligaments and muscular instability.</td>
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<td>X</td>
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<tr>
<td>• Revision surgery after hinge knee or rotational knee joint</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Revision surgery by insufficient / inadequate bone mass</td>
<td>X</td>
<td>X</td>
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<tr>
<td><strong>Differential Indications</strong></td>
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<tr>
<td>• Arthrosis of patella flange</td>
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<td>X</td>
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<tr>
<td>• Valgus/Varus deformities &lt;10°</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Valgus/Varus deformities 10-15°</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Valgus/Varus deformities 15-20°</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>• Valgus/Varus deformities 20-30°</td>
<td>-</td>
<td>X</td>
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<tr>
<td><strong>Contraindications</strong></td>
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<tr>
<td>• Acute or chronic infections, local and systemic</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Allergies due to (implant) materials</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Distinctive muscular, nerve, vascular or other diseases which put the affected limb at risk.</td>
<td>X</td>
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<tr>
<td>• Insufficient / inadequate bone mass- or quality which prevents a stable anchor of the prosthesis.</td>
<td>X</td>
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<tr>
<td><strong>Relative Contraindications</strong></td>
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<tr>
<td>• Adiposity</td>
<td>X</td>
<td>X</td>
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<td>• Insufficient musculature</td>
<td>X</td>
<td>-</td>
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<tr>
<td>• Lacking or foreseeable not assured compliance</td>
<td>X</td>
<td>X</td>
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<tr>
<td>• Foreseeable overload of joint prosthesis</td>
<td>X</td>
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</table>

Please note: These indications/contraindications refer to standard cases. The ultimate decision on whether or not an implant is suitable for a patient must be made by the surgeon based on his/her individual analysis and his/her experience.
Surgical Technique

Opening

01
A rein placed posteriorly around the condyles helps to stabilize the femur when the knee is flexed.

02
Osteophytes are removed to restore the physiologic contour of the condyles.

03
The width of the condyles determines which size of knee prosthesis to select (A).

The red line indicates the axis resulting from the initial opening of the femoral canal. The blue line indicates the axis of the femur and the final position of the stem of the femoral prosthesis components (B).
**Femoral Preparation**

04. To initially open the femoral canal, the bone awl is placed at the lowest point (red) of the trochlear groove in the patellofemoral joint. The point lies at the tip of the intercondylar fossa.

05. The femoral canal is opened with an 8 mm milling drill at the point previously marked.

06. The femoral canal is carefully widened with a ball reamer.
Femoral Preparation

07
The femoral saw guide corresponds in shape to the portion of the femoral component that will lie within the bone (A). The threaded rod with handle is attached to the anterior part of the femoral saw guide. To centrally align the instrument in the femoral canal, the extension stem is screwed into the femoral saw guide. At the proximal end of the extension stem, a metal centralizer adapted to the diameter of the canal is attached (B). The saw guide is advanced until its box is in contact with the condyles. The rotational alignment is correct when the anterior groove of the resection box matches the trochlear groove when viewed from above (C).

08
Alternatively, the femoral saw guide can be aligned with the femur and the handle with the aid of the alignment guide and can be fixed with fixation pins.
Femoral Preparation

09

Anterior and bilateral femoral osteotomies are performed with the oscillating saw along the surface of the box (D).

The resected bone at the junction of the condyles should be carefully osteotomized step by step with the saw. This should allow the box of the femoral saw guide to be inserted up to the level of the condyles after the resected bone has been removed with a lambotte osteotome (E).

10

In order to adapt the anterior femoral canal to the slight groove in the prosthesis and the saw guide between the anterior surface of the intracondylar box and the stem, the cancellous bone retrograde to the trochlear groove is hollowed out with the oscillating saw. This process creates a step in the cancellous bone attached in a proximal direction, so that this can be removed with a ball reamer or broach.
Femoral Preparation

The femoral canal is intentionally opened far in a posterior direction first (Fig. F, red point). The inserted femoral guide must then be shifted anteriorly (Fig. F, blue point) until the stem of the femoral saw guide (locked in extension) is aligned with the distal femur (Fig. G, blue position of femoral saw guide), so as to prevent hyperextension.

If proper axial alignment of the femoral saw guide cannot be achieved after the previous steps have been performed, the femoral saw guide will have to be shifted further in an anterior direction. This requires resection of more bone from the intracondylar portion of the trochlear groove. A lambotte osteotome impacted into the bone can be used to define the extent of the osteotomy with the oscillating saw. The impacted osteotome also guides the saw blade until the saw has cut a groove in the bone. It is important to perform the resection gradually, verifying proper alignment of the axis of the femoral saw guide with the axis of the femur. After this additional resection, the steps of hollowing out the cancellous bone proximal to the trochlear groove and removing cancellous bone from the anterior portion of the femoral canal proximal to the joint must be repeated.
Femoral Preparation

13
The box of the femoral saw guide is advanced into the bone until the contour of the anterior groove of the saw guide is congruent with the trochlear groove in the patellofemoral joint. The femoral saw guide is fixed with two fixation pins. The condyles are then adapted to match the curved surface of the resection box with an oscillating saw.

14
After the trochlear groove resection guide is attached to the saw guide, the anterior surface of the distal femur is resected to fit the shape of the implant.

15
The remaining anterior edge is rounded off with a trochlear groove rasp. Residual cartilage should be removed.
Femoral Preparation

16
In sclerotic bone, it is a good idea to achieve a slight sagittal curvature of the inner edges of the condyles to match the interior contour of the implant between box and flanges. Soft cancellous bone will conform to the interior contour of the flanges; sclerotic bone must be shaped to fit. In cases where the medial femoral condyle is sclerotic, cement retention holes are drilled to improve fixation of the implant.

17
When using cemented modular stems, a cylindrical bone plug is inserted into the proximal region of the femur to seal the femoral canal. The bone cylinder is advanced to a depth corresponding to the overall length of the stem (see Implants and Instruments Catalog, page 12). This plug provides hemostasis higher up in the femoral canal and seals the canal when cement is injected during later implantation of the femoral component.

18
When using cementless conical or cylindrical modular stems, the corresponding attachment plate (femur or tibia) is affixed to the shaft of the broach so that the inscribed side of the plate is flush with the bone on the averted side and the black marking ring on the shaft of the broach is barely visible (see arrow). The attachment plate is fixed with the inbus screw.
Femoral Preparation

19
The broach is secured to a handle for milling and broaching. Machine operation of the broach is not permitted.

20
The awl is subsequently carefully screwed in a clockwise direction until contact of the attachment plate with the bone. In this position, stable cortical anchoring should be achieved.

21
The selected trial stem is screwed onto the femoral trial prosthesis.
Femoral Preparation

22
Fitted trial femoral prosthesis.
Tibial Preparation

23 Next, the tibia is exposed. The hand on the opposite side to the approach is placed in the popliteal fossa with the thumb abducted, while the other hand grasps the ankle anteriorly. Applying traction and shear force with the hands with slight external rotation exposes the proximal tibia.

24 The tibial canal is marked with a broach at the junction between the anterior and middle third of the sagittal diameter of the tibial surface, immediately anterior to the intercondylar eminence. The point described lies above the center of the medullary canal of the tibia. An 8 mm twist drill is used to open the tibial canal.

25 The guide bar is subsequently introduced and the T-shaped handle is unscrewed.
# Surgical Technique

## Tibial Preparation

26
The coupling is pushed over the guide bar and is connected to the tibial saw guide with a caliper. The caliper is aligned to the highest level of healthy plateau surface during initial procedures. The maximum resection level is 10 mm. In the case of revision, precisely enough material should be resected in order to leave a planar osseous surface. In each case, the dorsal slope is 0°. To check the slope angle and to rotationally align the saw guide, the alignment rod can be affixed to the tibial saw guide anteriorly.

27
The saw guide is secured by means of fixation pins (at least four). Of these, at least two angled boreholes (black borders) and two straight boreholes are to be used. Resection is subsequently carried out. The rotational alignment is marked for the later positioning of the tibial implant with a sterile pin or scoring of the anterior cortex.

28
The preparation of the tibial canal is carried out with ball reamers (for cemented modular stems) or broaches (conical or cylindrical for cementless modular stems) as described in femoral preparation. The broaches are joined to the tibial attachment plate.
**Tibial Preparation**

29 The profile of the implant in the corresponding implant size is inserted into the proximal tibia using the proximal tibial impression instrument. The anterior alignment pin of the instrument is then aligned with the rotation marking of the anterior cortex carried out prior to this. The instrument is driven in deep enough so that the upper surfaces are in line with the resection level.

30 The trial tibial prosthesis is attached posteriorly to the selected trial stem. The threaded rod with handle is screwed on in a proximal direction and the trial implant is driven in.

31 The tibial and femoral trial implants are coupled. The trial tibial plateau is fitted with the aid of the introducer.
It is now important to check the sagittal and rotational alignment of the implant before cementing the implant components in place:

- **H** hyperextension
- **J** impaired extension
- **K** external hyperrotation
- **L** internal hyperrotation

must be corrected.

Figures **M** and **N** show correct axial alignment of the implant.
Implantation: Rotational Knee Prosthesis

33
Once it has been verified that the implant is correctly positioned, the components are individually cemented in place. When using cemented modular stems with a length of 120 mm, it is recommended that the femoral components are usually implanted in around 80 g bone cement. The use of longer cemented modular stems requires a correspondingly larger amount of bone cement. Excess cement at the sides is removed.

When cementless modular stems are used, the femoral joint portion must be anchored with bone cement.

34
The plastic strip that protects the joint mechanism from the entry of cement is removed. The strip should be pulled posteriorly.

35
Before the tibial component is cemented with at least 40 g of cement, when cemented modular stems with a length of 135 mm are used, the tibial canal is sealed with a bone plug. The use of longer cemented modular stems requires a correspondingly larger amount of bone cement. Note the marked rotational position when aligning the implant. When cementless modular stems are used, the tibial joint portion must be anchored with bone cement.

Caution!
The tibial component may only be cemented without the polyethylene plateau removed where the trial screw has been inserted to a maximum depth. To remove the polyethylene plateau, the trial screw is removed and the plateau taken off of the metal tibial tray using the tibial plateau introducer. Subsequently, the trial screw is reintroduced until it stops in the metal plateau. This is necessary to prevent cement from entering the threaded hole in the metal plateau.

Excess cement is removed.
### Surgical Technique

#### Implantation: Rotational Knee Prosthesis

36

Before the cemented components are fitted together, the polyethylene plateau still has to be removed from the tibial metal tray. With the knee flexed, the femoral component is inserted onto the pin of the tibial component.

37

To insert the polyethylene tibial plateau, the femoral component is lifted slightly. The tibial plateau is then inserted from an interior direction between the proximal and distal component of the prosthesis (O). Care should be taken to ensure that the chamber of the plastic plateau merges with the flange of the femoral component, and that the dovetail recess on the underside of the plastic plateau snaps into the marginal groove on the metal tibial tray (P).

38

The polyethylene plateau is secured in place on the metal tibial tray with the self-locking fixation screw.

**Caution!**

The self-locking fixation screw may only be used during the final assembly of the plateau. Loosening the fixation screw destroys the screw retention system in the polyethylene plateau, and a new plateau must then be inserted.
Implantation: Rotational Knee Prosthesis

The implanted rotational knee prosthesis should allow up to 90° of flexion, depending on the soft tissue structures. In extension, a resilient extension impairment of approximately 5° is optimal. This extension impairment helps to ensure secure contact between the two prosthesis components.
Surgical Technique

Implantation: Hinge Knee Prosthesis

The preparation of the osseous implant position is exactly the same as for the rotational variant. With regards to instruments, the complementary instrument set for the Endo-Model® Total Hinge Knee Joint Prosthesis is required.

01

After the femur has been prepared to receive the femoral component, the polyethylene bearings (A), which are supplied assembled, are removed from the box of the femoral component and replaced with trial bearings (B). This takes place with special applying forceps (C).

Later, the trial bearings are replaced with the final bearings.

02

The introducer (D) is inserted into the femoral component. The drill guide must lie medially. The cylindrical portion of the shank of the introducer, opposite the drill guide, is first inserted into the medial bearing. Once the moveable spacer on the shaft of the instrument has been inserted into the intracondylar opening, the introducer is locked in the box of the femoral component by tightening the knurled-head screw (E).
Implantation: Hinge Knee Prosthesis

03
A cylinder of bone is placed as a cement restrictor to seal the end of the femoral canal and the femoral component is implanted with at least 80 g of bone cement.

After the bone cement has hardened, a cylinder of bone is harvested from the medial condyle with the trephine and removed from the reamer for use later on in the procedure.

04
The tibial trial component is advanced into the prepared tibial canal with the introducer.
05 The upper and lower prosthetic components are connected. The pin (F) on the coupling of the tibial trial component is introduced into the lateral trial bearing and locked with the coupling jig (G).

06 Correct location of the prosthesis components with respect to axial alignment, rotation, and sufficient extension and flexion should be verified. In certain cases, additional tibial resection may be required or a spacer should be used.
Implantation: Hinge Knee Prosthesis

07
The tibial component is implanted with at least 40 g of bone cement. Correct rotation is maintained with the introducer (H) while distal pressure is applied to the implant.

08
Finally, the trial bearings are replaced with the final bearings using the applying forceps (I).

Caution!
The open bearing must be placed medially, as the prosthesis axle is introduced medially.
The tibial component is introduced into the femoral component and is adjusted with the trial axle (J). A trial run is carried out. The upper and lower parts of the prosthesis are then sealed with the final prosthesis axle (K), which is mounted on the threaded rod (L). A set screw firmly screwed into the screw hole of the axle (M) ensures the axial position.

In order to prevent loosening of the screw, the terminal screw hole must be sealed over the set screw with some bone cement. The milled bone cylinder is introduced into the medial femoral condyle again.
Implantation: Hinge Knee Prosthesis

Completely implanted Endo-Model® Hinge Knee Prosthesis.
15-2599/05

X-ray template for Endo-Model®-M
Modular Knee Prosthesis System, including modular stem extensions,
110% actual size, 1 set of 9 sheets
Further information about:

**Endo-Model® – M, Implants & Instruments**
Modular Knee Prosthesis System with Segmental Bone Replacement Components
(Catalogs: 718dt_Impl.Instr / 718en_Impl.Instr.)

available on request

Further information about:

**Titan Niobium Nitride Coatings** for custom-made orthopaedic implants
– Material and Coatings
(catalog: 914en)

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Important Information

Please note the following regarding the use of our implants:

1. **Choosing the right implant is extremely important.**
   The size and shape of the human bone determine the size and shape of the implant and also limit 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 exceedingly 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 reused.

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!*

WALDEMAR LINK GmbH & Co. KG, Hamburg

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The Surgical Technique described has been written to the best of our knowledge and belief, but it does not relieve the surgeon of his/her responsibility to duly consider the particularities of each individual case.

Unless otherwise indicated, all instruments are made of surgical stainless steel.