Zimmer Biomet
CoCr and Ceramic
Femoral Heads
Surgical Technique
**Introduction**

Femoral heads are intended for mating with titanium alloy, cobalt-chromium molybdenum alloy, cobalt-chromium-nickel-molybdenum alloy and stainless steel femoral stems.

For specific product compatibility, reference the following compatibility website: [https://IFU.zimmer.com](https://IFU.zimmer.com)

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**CoCr Femoral Heads**

Zimmer Biomet CoCr Femoral Heads and Freedom® Constrained Heads are made from cobalt-chromium molybdenum alloy. They are manufactured to include a 12/14 taper or Type I taper and are offered in a variety of head diameters and neck configurations. See the applicable Ordering Information document for detailed product descriptions.
Ceramic Femoral Heads

Zimmer Biomet offers ceramic heads that are made from an aluminum oxide matrix composite ceramic in accordance with ISO 6474-2.

Ceramic femoral heads are offered in non-sleeved and sleeved designs. Non-sleeved heads are manufactured to include a 12/14 taper or Type I taper. Sleeved options include taper adaptors compatible with 12/14, Type 1 and 6° femoral stem tapers. See the applicable Ordering Information document for detailed product descriptions.

Non-sleeved ceramic femoral heads should only be implanted on pristine/non-damaged stems. In revision cases, a sleeved ceramic femoral head with a taper adaptor should be implanted.

The Zimmer Biomet Ceramic Heads are intended for use as a component of a total hip or hemi-hip prosthesis in primary and revision patients. Refer to the Zimmer Biomet Ceramic Heads package insert for the indications for use.

Do not couple 22.2 mm ceramic head sizes with cobalt-chrome or stainless steel femoral stems. Generally, these materials impose higher stresses on the ceramic head.
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Preoperative Planning

The objectives of pre-operative planning are to define:

- Pre-operative leg length
- Acetabular component size and position
- Femoral component size
- Femoral offset and center of rotation

X-ray templates and digital templates, for some brands, are provided with acetabular and femoral systems.

When performing a revision of the acetabular component only, it is important to identify the stem which will remain in situ during the pre-operative planning. This will help select the appropriate CoCr femoral head type or the right taper adaptor for a ceramic head. The inner taper of the head adapter must fit the stem taper.
Spherical and Freedom Constrained CoCr Femoral Heads

Femoral Head Trial
Select the correct provisional head that matches the stem taper (Figure 2). Visually inspect the trial heads for damage prior to use. Select the appropriate head diameter and offset to create equal leg length and needed lateralization.

Trial Reduction
Perform a trial to assess leg length, range of motion, stability and abductor tension (Figure 3). Repeat as necessary until optimal offset and leg length are established making any necessary adjustments to restore joint mechanics, range of motion and stability.

Make certain that prominent impinging bone and/or osteophytes are removed from the periphery of the acetabulum to maximize range of motion and stability.

Note for Freedom Constrained Heads: Only Freedom provisional heads are compatible for trialing with Freedom implant liners. Never mix the implant components and the provisional components to perform a trial reduction.
Femoral Head Impaction

Thoroughly clean and dry the implant trunnion taper. Assemble the appropriate mating CoCr femoral head onto the stem taper. Impact the femoral head with at least one hard mallet strike using the femoral head impactor axially aligned on the femoral head to the stem taper (Figure 4). Test the security of the head fixation by trying to remove it by hand.

Note for Freedom Constrained Heads: Make sure to position the Freedom head on the stem so that the etch marking on the head is located in the most superior position prior to impaction (Figure 5).

Final Reduction

Once all final implants have been placed, reduce the hip and assess leg length, range of motion, stability, and abductor tension (Figure 6).
Non-Sleeved Ceramic Femoral Head

Femoral Head Trial
Select the correct provisional that matches the stem taper (Figure 7). Visually inspect the provisional heads for damage prior to use. Select the appropriate head diameter and offset to create equal leg length and needed lateralization.

Trial Reduction
Perform a trial reduction to assess leg length, range of motion, stability, and abductor tension (Figure 8). Repeat as necessary until optimal offset and leg length are established making any necessary adjustments to restore joint mechanics, range of motion and stability.

Make certain that prominent impinging bone and/or osteophytes are removed from the periphery of the acetabulum to help maximize range of motion and stability.
Femoral Head Impaction

Thoroughly clean and dry the implant trunnion taper.

Place the non-sleeved ceramic femoral head onto the stem taper by gently turning it (Figure 9). Impact the femoral head with at least one light mallet strike using the femoral head impactor axially aligned on the femoral head to the stem taper. Impact the head with a plastic femoral head impactor only (Figure 10). Test the security of the head fixation by trying to remove it by hand.

Do not couple 22.2 mm ceramic head sizes with cobalt-chrome or stainless steel femoral stems. Generally, these materials impose higher stresses on the ceramic head.

Note: The use of metal impactors or any other metallic objects may scratch or crack the modular head bearing surface, compromising the integrity of the component. If the modular ceramic head becomes scratched or cracked, the head and neck sleeve (if using a sleeved component) must be replaced.

Final Reduction

Once all final implants have been placed, reduce the hip and assess leg length, range of motion, stability, and abductor tension (Figure 11).
Sleeved Ceramic Femoral Head

Stem Trunnion Inspection - Acetabular/Head revision

The sleeved ceramic femoral head can be used on either a new/pristine stem or a previously implanted stem. If the ceramic femoral head is being used with a previously implanted stem, use the Goldberg scale to determine if the condition of the trunnion is tolerable or intolerable.

Tolerable Condition

Used stem tapers displaying fine marks from head/stem disassembly and/or up to, and including a score of 3 as described by Goldberg:

- Taper surface discolored or dull
- <10% of taper surface containing black debris, pits, or etch marks

Intolerable Condition

Used stem tapers with a score of 4 as described by Goldberg:

- >10% of taper surface containing black debris, pits, or etch marks
- Several bands of fretting scars involving several adjacent machine lines, or flattened areas with nearby fretting scars
- A taper with a scratch/defect at a height of 0.25 mm, taper with a broad truncation, slanted taper or a crushed taper are all intolerable taper conditions.

Note: Sleeved ceramic femoral heads must not be used with tapers that exhibit these intolerable condition characteristics (Figure 12).
**Femoral Head Trial**
Verify the taper type on the existing stem if performing an acetabular only revision, or the stem to be inserted, and select the correct provisional that matches the stem taper (Figure 13).

Using the appropriate femoral head provisionals, perform a trial reduction as detailed in previous sections (Figure 14).

**Assembly Instructions**
Verify the correct selection of both the ceramic femoral head and the neck sleeve as predetermined during the trialing process.

Assemble the modular head components prior to positioning them onto the stem. Align the head onto the neck sleeve axially and apply pressure (Figure 15 and 16). A slight resistance will be felt once the taper is engaged.

⚠️ **Note:** Heads and neck sleeves may be packaged either together or separately. See ordering information for detailed packaging notes.
Femoral Head Impaction

Thoroughly clean and dry the implant trunnion taper. Place the ceramic femoral head with sleeve assembled onto the stem taper by gently turning it (Figure 17). Impact the femoral head with at least one firm mallet strike using the femoral head impactor axially aligned on the femoral head to the stem taper. Impact the head with a plastic femoral head impactor only (Figure 18). Test the security of the head fixation by trying to remove it by hand.

Note: The use of metal impactors or any other metallic objects may scratch or crack the modular head bearing surface, compromising the integrity of the component. If the modular ceramic head becomes scratched or cracked, the head and neck sleeve must be replaced.

Final Reduction

Once all final implants have been placed, reduce the hip and assess leg length, range of motion, stability, and abductor tension (Figure 19).
References


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Products manufactured by Zimmer Biomet are not designed to be compatible with products from other manufacturers. There is no assurance that products from different companies may be safely used in combination with each other.

The product compatibility website was developed to address the potential compatibility of product families across Zimmer Biomet companies. In order to establish compatibility, testing and/or engineering evaluations are being completed to assure our global customers that paired products can safely be used in combination. www.zimmerbiomet.com

For product information, including indications, contraindications, warnings, precautions, potential adverse effects and patient counseling information, see the package insert and www.zimmerbiomet.com. Check for country product clearances and reference product specific instructions for use.

Zimmer Biomet does not practice medicine. This technique was developed in conjunction with health care professionals. This document is intended for surgeons and is not intended for laypersons. Each surgeon should exercise his or her own independent judgment in the diagnosis and treatment of an individual patient, and this information does not purport to replace the comprehensive training surgeons have received.

As with all surgical procedures, the technique used in each case will depend on the surgeon’s medical judgment as to the best treatment for each patient. Results will vary based on health, weight, activity and other variables. Not all patients are candidates for this product and/or procedure. Caution: Federal (USA) law restricts this device to sale by or on the order of a surgeon. Rx only.

This surgical technique is applicable to part numbers included within ordering information ‘2307.X-GLBL-en-Ceramic Heads Ordering Information’ and ‘1839.X-GLBL-en-CoCr Heads Ordering Information’.

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