Zimmer Biomet Hip Preservation - a new opportunity to treat the hip joint.

All conditions leading up to osteoarthritis present an opportunity to preserve the native anatomy, including:

- Femoroacetabular Impingement
- Dysplasia
- Subchondral Bone Defects
- Avascular Necrosis
- Femoral Neck-Shaft Abnormalities
- Femoral Version Deformities
- Cartilage Damage
- Labral Damage

Using a combination of different procedures, technologies, and products, we are able to help surgeons treat a spectrum of clinical conditions and give patients more treatment options than they would have been afforded in the past.
Zimmer Biomet offers solutions across the continuum of hip preservation procedures to make it simple for surgeons to restore a healthy hip.

**Patient Specific Analysis**

3D Motion Hip Analysis Service, powered by Clinical Graphics
- Delivers next-generation visualization by converting CT and MRI scans into interactive reports

**Hip Arthroscopy**

- JuggerKnot® Long Soft Anchor System
- Quattro® GL Suture Anchor (US Only)¹
- SureLock® All-Suture Anchor (US Only)¹
- SpeedSnare® Suture Passer System
- Dragon Tongue Suture Passing Device for Capsular Closure

¹ Available in the United States only.
Hip Subchondroplasty® Procedure

The Subchondroplasty Procedure is a technique designed to target subchondral bone defects for filling with bone substitute material.²

Surgical Head Dislocation (SHD)

DeNovo® NT (Natural Tissue) Graft

JuggerKnot® Soft Anchor-2.9 mm/2.9mm with Needles
Core Decompression

BioCUE® Blood and Bone Marrow Aspirate (BBMA) Concentration System

PerFuse™ Percutaneous Decompression System

Products are not available in all markets. Please contact your local sales representative for product availability.
Osteotomy

12 Large Titanium Cannulated Screw System (5.0 mm and larger screws)
13 Zimmer® Natural Nail® System (Antegrade and Cephalomedullary options)
14 Versa-Fx® II Femoral Fixation System
15 Dynamic Hip Screw Plate System
16 Intramedullary Bone Saw
17 Zimmer® Plates and Screw System (ZPS)

Products are not available in all markets. Please contact your local sales representative for product availability.
1. Quattro & SureLock Suture Anchors have not been approved for Hip procedures in the European Union. Please refer to the IFU for indications and contraindications.

2. AccuFill® Bone Substitute material is indicated to fill bone voids or defects of the skeletal system that are not intrinsic to the stability of the bony structure, and has not been evaluated for any clinical indications.

3. The BioCUE System is indicated to prepare autologous platelet-rich plasma (PRP) to mix with autograft and/or allograft bone prior to application to an orthopedic site. The PRP has not been evaluated for any clinical indications.

4. Core decompression is an avascular necrosis treatment option. The PerFuse System is designed to access the femoral head for core decompression; and to facilitate mixing/pre-mixing of bone graft material with I.V. fluids, blood, platelet-rich plasma, bone marrow, or other specified blood components.

5. Screws 5 mm and larger are intended for use in the femur; but only screws that are 6.5 mm or larger in diameter are indicated for both the pelvis and femur.

6. The DHS and BP (sterile) are intended for temporary fixation of fractures or osteotomies in the proximal or distal femur, and are not available in the US.

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