



**Summary of safety and clinical
performance
(SSCP)**

Intended for Patients

[United Hip System – UCP Stem and UCP Stem,
Centralizer with Cement Restrictor]

[English]

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The SSCP is not intended to give general advice on the treatment of a medical condition. Please contact your healthcare professional in case you have questions about your medical condition or about the use of the device in your situation. This SSCP is not intended to replace an Implant card or the Instructions For Use to provide information on the safe use of the device.

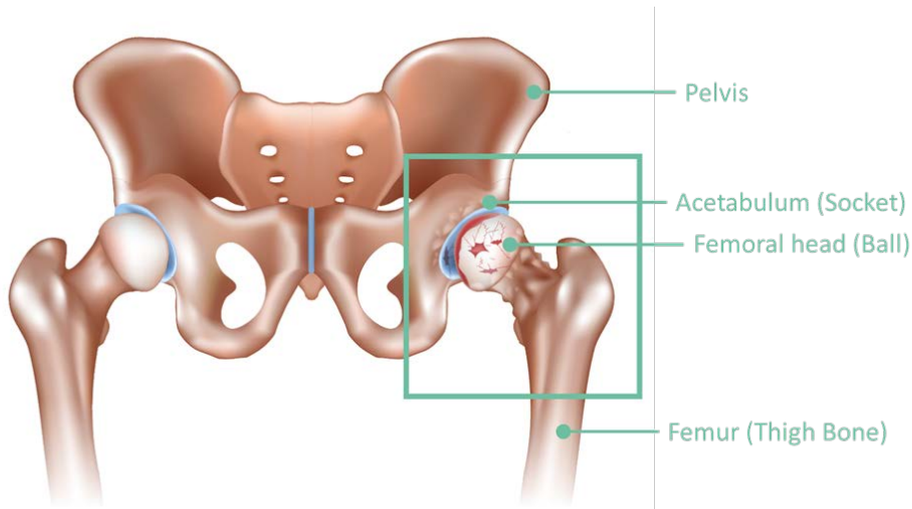
1. Device identification and general information

Device Trade Name	United Hip System – UCP Stem (1107-3000~3050, 1107-3200~3250, 1107-3421~3433) – UCP Stem, Centralizer with Cement Restrictor (1907-3008~3018)
Basic UDI-DI	471987216TD-III-009-2T9 (UCP Stem) 471987216TD-IIIb-018-466 (UCP Stem, Centralizer with Cement Restrictor)
Manufacturer	United Orthopedic Corporation
Address	No. 16, Luke 1st Rd. Luzhu Dist. Kaohsiung City, 82151, Taiwan
First Year of CE Certificate	2014 (UCP Stem) 2001 (UCP Stem, Centralizer with Cement Restrictor)

2. Intended use of the device

2.1. What is the device used for? (Intended purpose)

This document is to help you know more about the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”. If your hip joint is damaged by arthritis, the cartilage and bone of the hip joint become worn out. It may get hard to move and cause pain over time. Total hip arthroplasty is one of the treatment methods. Hip arthroplasty is intended to enhance function and mobility and reduce pain in daily activities. In this surgery, the doctor removes the damaged areas and replaces them with artificial implants. If you want to know more about your hip joint, please check the picture below. The hip consists of ball, thigh bone and pelvis.



2.2. For what conditions do doctor use the device? (Indications and intended patient groups)

United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” is indicated for use in primary or revision total hip arthroplasty in skeletally mature patients with the following conditions:

- A loss of bone and/or cartilage in the hip joint. It may cause by non-inflammatory disease or

autoimmune, and inflammatory disease attack the joints.

- Bone tissue in the thigh bone near the hip joint dies because of a lack of blood supply.
- Revision procedures when other treatments or devices have failed.
- Treatment of bone nonunion, such as femoral neck and trochanteric fractures, involve the proximal femur with the head. It is impossible to deal with by other techniques.

2.3. Who should not have the device? (Contraindications)

Your doctor may decide that total hip arthroplasty is not suitable for you if:

- You currently have hip joint infections.
- Your bone is not strong enough.
- Your joint tissue is not stable.
- Your skeleton is not mature.

3. Device description

3.1. What is the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”? (Device description)

The United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” is used for total hip arthroplasty. Figure 1 shows an overview of it. The United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” includes:

- UCP Stem, Cemented, Standard
- UCP Stem, Cemented, High Offset
- UCP Long Stem, Cemented, High Offset
- UCP Stem, Centralizer with Cement Restrictor.

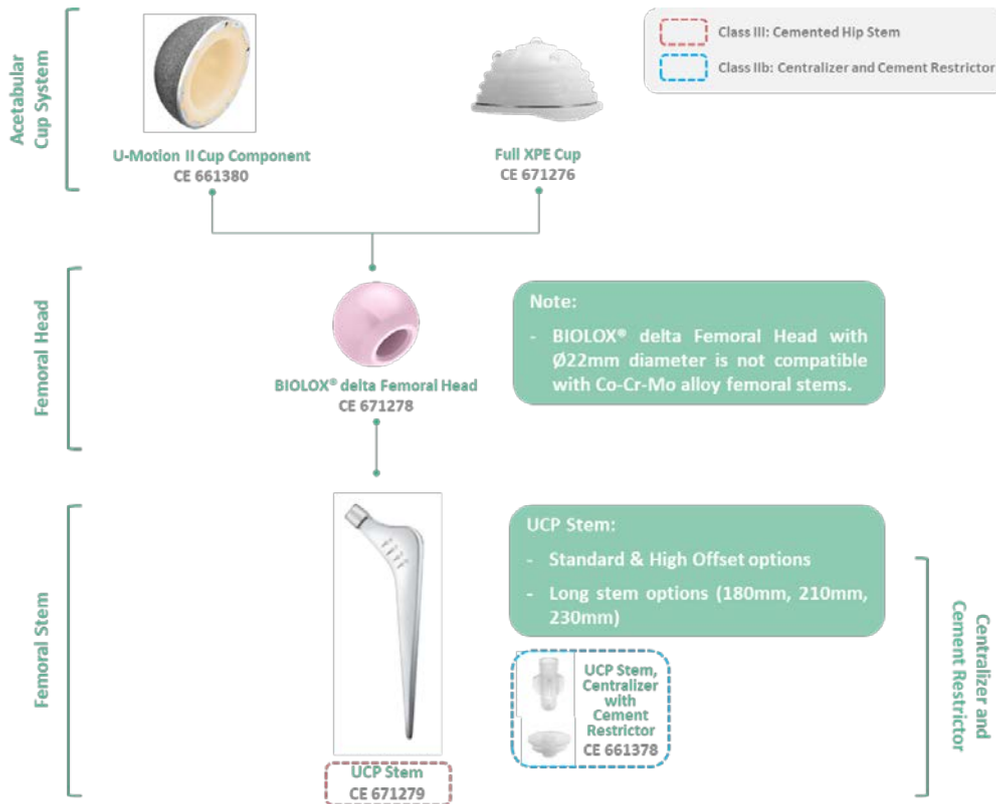
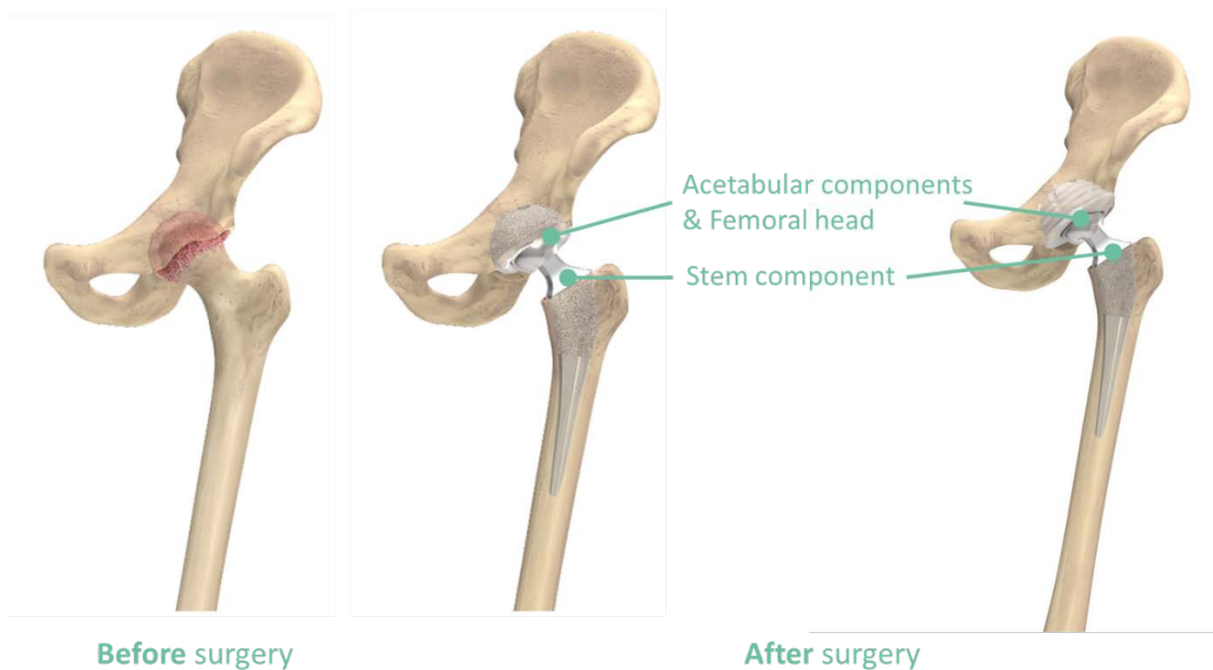


Figure 1. The compatible components for the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”.

If you received the Total Hip Arthroplasty, the expected consequence will be the picture shown below:



3.2. Do the device contain medicinal? (Information about medicinal substances in the device)

None. None of medicinal substances contain in the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”.

3.3. Description of how the device is achieving its intended mode of action

The United Hip System is intended to achieve what your hip work. The hip is a ball-and-socket joint. The ball is on the top of the thigh bone (femur) named the femoral head. The socket is a hollow part of the pelvis called the acetabulum. The ball is controlled by muscles and stabilized by tendons¹ and ligaments². The structures that surround the hip joint allow for hip movement and rotation.

When the hip is diseased or injured, it causes disrupted of natural balance and function. It potentially causes pain, limits mobility, or creates other medical challenges.

3.4. Description of accessories

Accessories will smooth your surgery proceed. The hip instruments will used to complete your surgery.

4. Risks and warnings

4.1. How potential risks have been controlled or managed

All the potential risks have been controlled. To reduce risk as low as possible by following the risk management standards ISO 14971³. This standard is the risk management specified in medical device industries.

4.2. What are the risks of having this surgery? (Remaining risks and undesirable effects)

The following risks have been associated with total hip arthroplasty. Included but are not limited to:

Item	Possible risks and side effects	How often each risk occurs (%) ^a			
		UCP Stem ^b		Benchmark device ^c	
		General problems	Additional surgery to replace the implant	General problems	Additional surgery to replace the implant
1	Negative tissue reaction (Adverse local tissue reaction)	-	-	No incidence rate reported ^d	0.67
2	Metal rust (Corrosion)	-	-	-	No incidence rate reported ^d
3	Broken implant (Implant breakage)	-	-	-	No incidence rate reported ^d
4	Loose implant (Loosening)	-	-	0.04-2.44	0.15-3.42
5	Bone break near implant (Periprosthetic fracture)	-	1.9	0.32-3.65	0.03-3.6
6	Moving downwards (Subsidence)	-	-	13.57	No incidence rate reported ^d
7	Broken bone (Fracture)	-	-	0.23-0.75	No incidence rate reported ^d

¹ Tendons: connect the muscle to the bone.

² Ligaments: connect bones to bones

³ ISO 14971: Medical devices – Application of risk management to medical devices.

Item	Possible risks and side effects	How often each risk occurs (%) ^a			
		UCP Stem ^b		Benchmark device ^c	
		General problems	Additional surgery to replace the implant	General problems	Additional surgery to replace the implant
8	Bruise (Hematoma)	-	-	0.47-6	0.24
9	Infection	1.9	3.45	0.38-7.33	0.7-3.33
10	Mental health condition	-	-	No incidence rate reported ^d	-
11	Pain	-	-	-	No incidence rate reported ^d
12	Muscle, tendon, or ligament problem (Soft tissue complication)	-	-	No incidence rate reported ^d	0.18
13	Surgery-related issue (Surgical complication)	-	-	0.5	-
14	Blood clot (Thrombosis)	-	-	0.27-2.94	-
15	Injury (Trauma)	-	-	-	No incidence rate reported ^d
16	Wound problem (Wound complication)	-	-	0.76-6	3.33
17	Other problems	3.6-3.7	3.45	0.49-2	0.08-1.44

^a Many things can affect the chance of problems, including but not limited to product design, how the surgery is performed, and the patient's health condition.

^b This information is based on patient follow-up studies (PMCF), medical articles, doctor experience, and safety monitoring after the product is on the market.

^c This information is based on medical articles and doctor experience.

^d This side effect is identified from safety monitoring without frequency reported.

4.3. What are the things you must do to avoid potentially serious harm if implanted with the device? (Warnings and Precautions)

- To keeping your quality of life, do not have a large amount of activities, such as sports (running, or muscle strain, etc.) that will place too much stress on the hip joint. Obesity, whose weight is heavier than 75 kg or BMI equal to or more than 30 may produce loads on the hip joint. Your artificial hip joint may fail and not work if placed too much stress on it.
- For those not recommended activities and work after surgery, please follow your doctor's recommendations and physical therapy schedule. Your doctor will give you instructions based on you and your hip implant.
- Inform your doctor if you have signs of infection such as joint pain close to the surgical site, fever, chills, redness, etc.
- Regular X-rays shall be taken. To evaluate if the implant moves, loose, bend, fracture or the cement or bone loss. If one of these conditions occur, please pay attention to it and consider the advantage of revision.
- "UCP Stem" is "MR Conditional," which means you should only receive an Magnetic Resonance

Imaging scan under certain conditions. Metal implants may interact with an Magnetic Resonance Imaging scanner. It may also cause heating or damage to the tissue around the implant. The metal can distort the image taken by the Magnetic Resonance Imaging scanner. You should let your doctor know you have an implant prior to receiving an Magnetic Resonance Imaging scan.

4.4. Have any adverse events ever occurred to the device? (Summary of any field safety corrective action and field safety notice)

A field safety corrective action is any action taken to reduce a risk of death or serious deterioration in the state of health associated with the use of a medical device. It is required when it becomes necessary for the medical device owner to take action to eliminate, or reduce the risk of the identified hazards. The field safety corrective action may include but not limited to:

- The return of the device to the product owner.
- Inspection or examination of the device by user.
- Modification of the device. For example, advice for a change in the way the device is used.
- Exchange of device.
- Damage of device.
- Retrofit by purchaser.
- Advice on the use of the device.
- Advice follow-up of patients, users or others.
- Recall of device.

Until now (the date issued of this document), numerous sets of the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” have been sold worldwide. The volume of sales of the products are summarized below. No field safety corrective action (recall) reported was released.

Product Name	The Volume of Sales (piece)	
UCP Stem, Cemented, Standard	7,516	11,527
UCP Stem, Cemented, High Offset	2,563	
UCP Long Stem, Cemented, High Offset	1,448	
UCP Stem, Centralizer with Cement Restrictor	10,633	

5. Summary of clinical evaluation and post-market clinical follow-up

The Effective Group

Post-market clinical follow-up has been conducted for the United Hip System – “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”.

Post-market clinical follow-up has been conducted for the UCP Stem, Cemented, Standard and UCP Stem, Centralizer with Cement Restrictor. The post-market clinical follow-up results are summarized in the table below. No relevant information in post-market clinical follow-up studies are currently available in the revision surgery of UCP Stem, Cemented, Standard and UCP Stem, Centralizer with Cement Restrictor. United is conducting the post-market clinical follow-up study at multiple clinical study sites. Furthermore, the relevant clinical results are collected.

Device	UCP Stem, Cemented, Standard collocated with UCP Stem, Centralizer with Cement Restrictor	
Diagnosis	Primary Total Hip Arthroplasty	
Cases	54 hips*	
Follow-up	1.3 years (range from 0.3 to 2 years)	
Survival Outcomes	95.5% at 2 years follow-up*	
Clinical Outcomes	Harris Hip Score	90.4 points
	Oxford Hip Score	42 points
Complication	<ul style="list-style-type: none"> - Cellulitis (1.9%) - Mortality not related to implant (3.7%) 	
Revision	- Periprosthetic fracture (1.9%)	

* The usage of UCP Stem, Centralizer with Cement Restrictor was unknown in 2 hips.

* The post-market clinical follow-up is continued to achieve maximum follow-up of 10 years based on our post-market clinical follow-up plan to confirm its safety and performance.

Post-market clinical follow-up has been conducted for the UCP Stem, Cemented, High Offset and UCP Stem, Centralizer with Cement Restrictor. The post-market clinical follow-up results are summarized in the table below. No relevant information in post-market clinical follow-up studies are currently available in the revision surgery of UCP Stem, Cemented, High Offset and UCP Stem, Centralizer with Cement Restrictor. United is conducting the post-market clinical follow-up study at multiple clinical study sites. Furthermore, the relevant clinical results are collected.

Device	UCP Stem, Cemented, High Offset collocated with UCP Stem, Centralizer with Cement Restrictor	
Diagnosis	Primary Total Hip Arthroplasty	
Cases	56 hips*	
Follow-up	1.2 years (range from 0.3 to 2 years)	
Survival Outcomes	100% at 2 years follow-up*	
Clinical	Harris Hip Score	93.1 points

Outcomes	Oxford Hip Score	43.7 points
Complication		- Mortality not related to implant (3.6%)
Revision		- NA

* The usage of UCP Stem, Centralizer with Cement Restrictor was unknown in 4 hips.

* The post-market clinical follow-up is continued to achieve maximum follow-up of 10 years based on our post-market clinical follow-up plan to confirm its safety and performance.

Clinical performance conclusions

In the clinical study to support the device, the scores for clinical outcomes are described below:

Clinical outcomes	Scores
Harris Hip Score	90 to 100: excellent 80 to 89: good 70 to 79: fair <70: poor
Oxford Hip Score	42 to 48: excellent 34 to 41: good 27 to 33: fair <27: poor

The performance of the device was evaluated and summarized below.

The declared lifetime of the “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” is 10 years, where both average Oxford Hip Score and Harris Hip Score are good to excellent and the expected Survival rate for the primary total hip arthroplasty is over 95%, respectively to meet the state-of-the-art.

Total Hip Arthroplasty				
	Items to evaluate		What we expected	Clinical results
Primary	Performance	Harris Hip Score	good to excellent (average \geq 80 points)	excellent (91.8 points)
		Oxford Hip Score	good to excellent (average \geq 34 points)	excellent (42.9 points)
		Survival rate 10 years	\geq 95%	97.9% (2 years) *
Revision	Items to evaluate		What we expected	Clinical results
	Performance	Harris Hip Score	fair to excellent (average \geq 70 points)	No relevant information*
		Oxford Hip Score	fair to excellent	No relevant information*

			(average ≥ 27 points)	
		Survival rate 10 years	$\geq 85\%$	No relevant information*
*The post-market clinical follow-up is continued to achieve maximum follow-up of 10 years based on our post-market clinical follow-up plan to confirm its safety and performance.				

Clinical safety conclusions

In the clinical study to support the device, the safety of the device was evaluated based upon adverse event. After surgery (postoperatively), the complications or adverse events were reported and summarized as table below.

Complications or adverse events	Number of knees with complications or adverse events out of the total number of knees in the study	Percent (%) of patients who had this adverse event
Cellulitis	1 out of 110	0.9%
Mortality not related to implant	4 out of 110	3.6%
Periprosthetic fracture	1 out of 110	0.9%

In conclusion, the study data indicates the United Hip System - “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor” has achieved the clinical performance, which used in skeletally mature patients requiring Total Hip Arthroplasty.

6. What other treatments? (Possible diagnostic or therapeutic alternatives)

If you are considering alternative treatments, please consult your doctor. Your doctor can make suggestions based on your age, general health, and the condition of your hip. The alternative treatments may include but not limited to:

- Nonsurgical treatments:
 - non-pharmacological treatments;
 - pharmacological treatments;
 - cell therapy.
- Other surgical treatments:
 - joint aspiration, joint distraction, joint lavage
 - arthroscopic debridement;
 - cartilage repair technique;
 - internal fixation;
 - different type joint replacement.

7. Who are qualified to use the device? (Suggested training for users)

If you are advised to treat with total hip arthroplasty and implant the United Hip System - “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”, your surgery will be performed by certified Orthopedist.

8. More information

This document gives you information about your treatment choices. It is not intended to replace advice from a doctor. If you have any further questions about the United Hip System - “UCP Stem” and “UCP Stem, Centralizer with Cement Restrictor”, please discuss with your doctor.

9. History of revisions

SSCP revision No.	Date issued (DD-MM-YYYY)	Change description	Revision validated by the Notified Body
0	23-05-2025	First issue.	<input checked="" type="checkbox"/> Yes Validation language: English Note: The SSCP has been approved for the restrictors and centralizers only. <input type="checkbox"/> No
1	23-12-2025	Annually update.	<input checked="" type="checkbox"/> Yes Validation language: English Note: The SSCP has been approved for the restrictors and centralizers only. <input type="checkbox"/> No