S	М	I	т	н	&	N	E	Р	н	E	W

SPECTRONEF H		Р	S	Y	S	т	E	М
--------------	--	---	---	---	---	---	---	---

S	U	R	G	С	Α	L I	Т	E	С	н	Ν	Q	U	E

SPECTRON⁺⁺ EF

H I P S Y S T E M

J. Rod Davey, M.D., F.R.C.S.

Acting Head of Orthopaedic Surgery, The Toronto Hospital Assistant Professor of the Department of Surgery, University of Toronto Toronto, Ontario, Canada

Paul Di Cesare, M.D., F.A.C.S.

Hospital for Joint Diseases Orthopaedic Institute Director of Musculoskeletal Research Center Co-Director of Surgical Arthritis Service Assistant Professor of Orthopaedic Surgery, New York University School of Medicine New York, New York

Henrik Malchau, M.D., Ph.D.

Sahlgrenska University Hospital Associate Professor Department of Orthopaedics Göteborg, Sweden

Nota Bene: The technique description herein is made available to the healthcare professional to illustrate the authors' suggested treatment for the uncomplicated procedure. In the final analysis, the preferred treatment is that which addresses the needs of the patient.

Stem Specifications

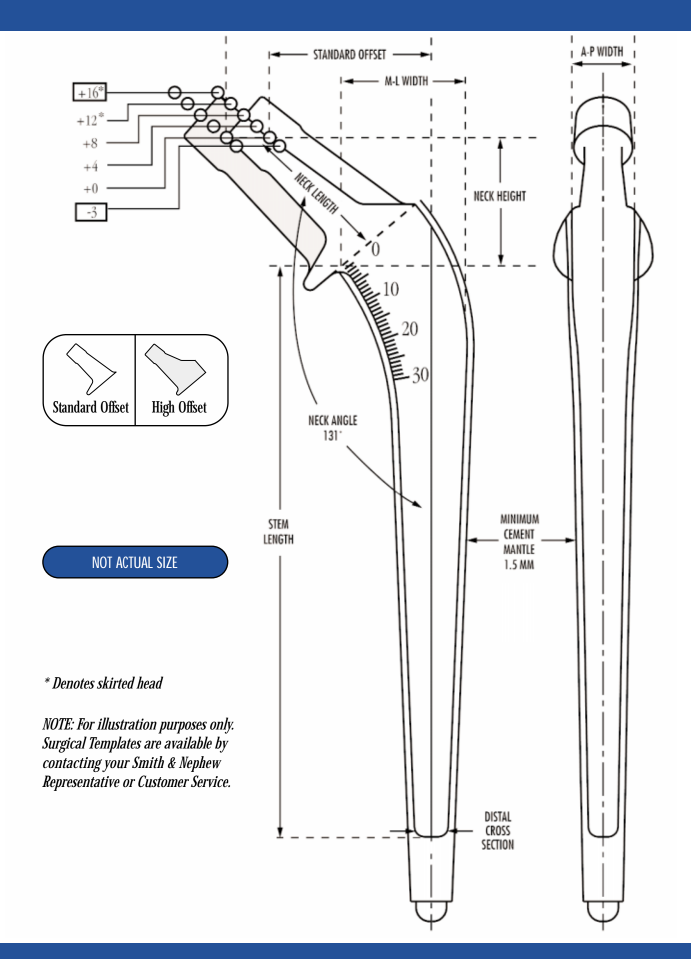
	SPECIFICATIONS									
Size	Neck Angle	Distal Cross Section	Stem Length	A-P Width	M-L Width					
1, 1H	131º	6 mm	115 mm	12.9 mm	25.4 mm					
2, 2H	131º	7 mm	125 mm	13.7 mm	27.2 mm					
3, 3H	131º	8 mm	135 mm	14.5 mm	28.9 mm					
4, 4H	131º	10 mm	135 mm	15.3 mm	30.7 mm					
5, 5H	131º	12 mm	135 mm	16.1 mm	32.5 mm					

NECK HEIGHT MM										
	W	When Femoral Head Component Selected Is:								
Size	-3	+0	+4	+8	+12	+16				
1	24	26	28	31	34	36				
1H	24	26	28	31	34	36				
2	26	28	30	33	36	38				
2H	26	28	30	33	36	38				
3	28	30	32	35	37	40				
3H	28	30	32	35	37	40				
4	30	32	34	37	39	42				
4H	30	32	34	37	39	42				
5	32	34	36	39	41	44				
5H	32	34	36	39	41	44				

NECK OFFSET MM									
	When Femoral Head Component Selected Is:								
Size	-3	+0	+4	+8	+12	+16			
1	32	35	38	41	44	47			
1H	38	41	44	47	50	53			
2	34	36	39	42	45	48			
2H	42	44	47	50	53	56			
3	35	38	41	44	47	50			
3H	45	48	51	54	57	60			
4	37	39	42	45	48	51			
4H	47	49	52	55	58	61			
5	38	41	44	47	50	53			
5H	48	51	54	57	60	63			

	NECK LENGTH MM								
	W	When Femoral Head Component Selected Is:							
Size	-3	+0	+4	+8	+12	+16			
1	27	30	34	38	42	46			
1H	31	34	38	42	46	50			
2	29	32	36	40	44	48			
2H	34	37	41	45	49	53			
3	31	34	38	42	46	50			
3H	37	40	44	48	52	56			
4	33	36	40	44	48	52			
4H	39	42	46	50	54	58			
5	35	38	42	46	50	54			
5H	41	44	48	52	56	60			

For use with Smith & Nephew 12/14 femoral heads only.



Femoral Osteotomy

1. Femoral Osteotomy

Use the osteotomy guide to determine the level of resection, using one of the following techniques:

A. Place the template over the X-ray of the hip. Determine the stem size. Determine length of femoral head to be used. A graduation scale can be found on the medial aspect of the stem on the template. Make note of how many graduations above the lesser trochanter where the osteotomy will take place, as determined by the collar of the stem.

In the O.R., place the osteotomy guide on the femur by referencing the lesser trochanter at the same graduation mark as noted during templating. Osteotomize the neck.

B. Place the template over the X-ray of the hip. Determine the stem size. Determine length of femoral head to be used. Determine the base neck length on the standard offset stem as indicated on the template. Add to this number the length of the femoral head.

In the O.R., place the greater trochanter block on the osteotomy guide at this numerical position. If the number does not match perfectly, use the lower number. Place the guide on the femur by resting the block on the top of the greater trochanter. Osteotomize the neck.

2. Prepare Acetabulum

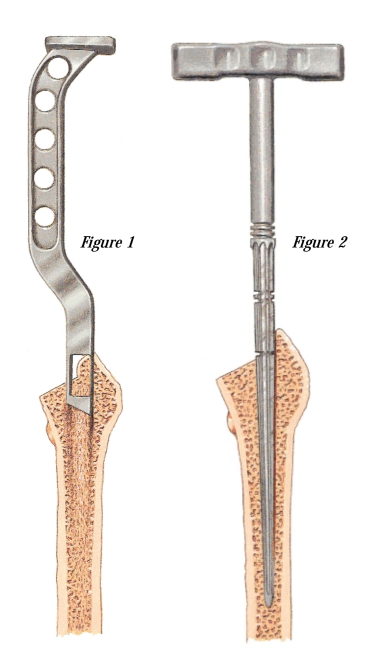
If acetabular reconstruction is required, prepare acetabulum using the technique for the intended acetabular component.



Femoral Canal Preparation

3. Femoral Canal Preparation

Open the medullary canal at the transected neck using the box chisel. Stay posterior and lateral in order to obtain a neutral stem position *(Figure 1).* Open the femoral canal using the blunt medullary reamer *(Figure 2).*



Femoral Broaching

4. FEMORAL BROACHING

Start the broaching procedure along the mid-axis of the femur with the Size 1 broach and progressively broach to the appropriate femoral stem size. Seat the broach slightly below the level of the femoral neck resection to facilitate calcar reaming.

The Spectron EF broach is designed to provide a minimum 1.5 mm cement mantle per side. Additional cement mantle thickness is achieved by pressurizing the cement into the cancellous bone. The broach is slightly longer than the corresponding implant to accommodate the distal centralizer.



Calcar Preparation & Trialing

5. CALCAR PREPARATION

With the final broach fully seated, remove the broach handle and ream the calcar with the appropriate calcar reamer. The smaller calcar reamer is used with broach sizes 1-3, and the larger calcar reamer is used with broach sizes 4 and 5. Plane the calcar until it is level with the broach.

6. TRIALING

Remove the calcar reamer and place the matching standard or high offset trial neck (as determined by templating) onto the broach post. Select the trial femoral head of desired diameter and neck length and reduce the hip to assess stability. Soft tissue tension can be improved by using the high offset trial neck instead of the standard offset trial neck without increasing leg length.

If trialing for the universal Bipolar or Unipolar, trial according to the appropriate technique for the selected device.

Fer	Femoral Head And Neck Length Options									
Trial Color	22 mm	26 mm	28 mm	32 mm						
Green			-3	-3						
Yellow	+0	+0	+0	+0						
Red	+4	+4	+4	+4						
White	+8	+8	+8	+8						
Blue	+12*	+12*	+12*	+12*						
Black	_	_	+16*	+16*						

Spectron EF

*Denotes skirted heads.





Placing The Buck⁺ Cement Restrictor

7. PLACING THE BUCK CEMENT RESTRICTOR

> Attach the broach handle to the broach and remove the broach from the femoral canal. Use the canal sounds to determine the distal canal diameter. The proximal flange of the cement restrictor should always be larger than the distal canal diameter (Table 1). Accurate cement restrictor depth placement is then determined by placing the Spectron EF stem (with attached distal centralizer) next to the inserter tool and adding 20 mm to the length (Table 2).

Remove the vent-occluding membrane by inserting the vent opening tool into the distal end of the restrictor and pushing the pin through the vent hole. Remove and discard the plastic debris.

Thread the cement restrictor onto the inserter using a clockwise motion. Insert the device to the level of the medullary canal that has been predetermined. Once this level is reached, disengage the restrictor from the inserter using a counterclockwise twisting motion. Remove the inserter from the medullary canal. If it is necessary to remove the restrictor prior to cement insertion, it can be reattached to the inserter rod and pulled out of the canal. The surgeon may adjust the restrictor as many times as required.

Buck Plug Size	Canal Size	Catalog Number
18.5 mm	<15 mm	129418
25 mm	16-21 mm	129419
30 mm	22-26 mm	7127-9420
35 mm	27-31 mm	7127-9421

Table 1



Stem Size	BUCK Cement Restrictor Insertion Depth (mm)
1	140
2	150
3	160
4	160
5	160

Table 2

Preparing The Femoral Canal

8. Preparing The Femoral Canal

Irrigate the canal with saline solution and pulsatile lavage to remove all debris. Continue preparing the femur with the femoral canal brush to remove any remaining weak cancellous bone, blood clots, and marrow fats. Repeat lavaging as necessary to remove all remaining debris.



Insert the femoral canal suction absorber into the femoral canal to dry the canal while mixing the cement.

Injecting Cement

10. Loading Cement

Load the cement powder and monomer into the cement mixer. Load the powder first, and then the monomer.

11. INJECTING CEMENT

After mixing the cement, remove the femoral canal suction absorber. Insert the nozzle of the cement gun to the top of the Buck cement restrictor and inject cement into the canal in retrograde fashion. Continue injecting cement until the canal is completely full and the distal tip of the nozzle is clear of the canal.



Pressurizing Cement

12. Pressurizing Cement

Break off the long nozzle and place the femoral pressurizer over the remaining short nozzle. Place the disposable femoral pressurizer into the mouth of the canal. This will occlude the canal and compress the cement. Maintain firm pressure until the cement is in a doughy state and can withstand displacement. This process will allow for proper cement interdigitation into trabecular bone. Withdraw the femoral pressurizer and remove any extruded cement around the periphery of the canal.



13. Selecting Stem & Distal Centralizer

Use the implant which corresponds to the last broach seated in the femur. An optional distal centralizer may be placed on the stem to provide neutral alignment and predictable cement mantle. Each implant has a corresponding distal centralizer which is intended to provide a uniform 1.5 mm distal cement mantle. Note, however, all of the stems will accommodate any of the available distal centralizers to address variations in distal femoral geometries. Using clean gloves, place the round plug of the selected centralizer into the hole at the distal end of the stem and push the centralizer superiorly until snug.

NOTE: If a distal centralizer is not used, place the distal hole plug which is packaged with the implant into the centralizer hole prior to inserting the stem.

Stem Size	Minimum Centralizer Size
Sizes 1, 1H	8 mm
Sizes 2, 2H	9 mm
Sizes 3, 3H	10 mm
Sizes 4, 4H	12 mm
Sizes 5, 5H	13 mm

Stem Insertion

14. STEM INSERTION

Insert the selected femoral stem into the canal. Fit the femoral stem driver into the stem driving platform and push into place. Advance the stem approximately 1 cm per second to avoid air inclusions at the stem/cement interface.

Trim away excess cement with Concise^{\dagger} cement sculps. Remove the stem driver and maintain steady pressure with the thumb on the neck taper until cement is cured.

15. FINAL TRIAL REDUCTION

A final trial reduction may be performed at this time using trial femoral heads.



Femoral Head Assembly

16. Femoral Head Assembly

Clean and dry the neck taper with a clean sterile cloth. Place the prosthetic femoral head on the neck taper and firmly impact several times with a head impactor and a mallet.



SPECTRON EF 12/14 FEMORAL STEM & HEAD COMPONENTS



Spectron EF 12/14 Primary Collared Stems Cobalt Chromium – ASTM F 799

Size	Stem Length	Implant Cat. No.	Broach/Trial Cat. No.	Trial Neck Cat. No.
1	115 mm	7131-2101	7136-5001	7136-5081
2	125 mm	7131-2102	7136-5002	7136-5082
3	135 mm	7131-2103	7136-5003	7136-5083
		7131-2104		7136-5084
5	135 mm	7131-2105	7136-5005	7136-5085



CoCr 12/14 Taper Femoral Heads Cobalt Chromium – ASTM F 799

Neck Length	22 mm	26 mm	
-3	_	_	
+0	7130-2200	7130-2600	
+4	7130-2204	7130-2604	
+8	7130-2208	7130-2608	
+12	7130-2212	7130-2612	
+16		—	
Neck			
Length	28 mm	32 mm	36 mm
-3	7130-2803	7130-3203	7130-3603
+0	7130-2800	7130-3200	7130-3600
+4	7130-2804	7130-3204	7130-3604

+0 7130-2800 7130-3200 7130-3600 +4 7130-2804 7130-3204 7130-3604 +8 7130-2808 7130-3208 7130-3608 +12 7130-2812 7130-3212 7130-3612 +16 7130-2816 7130-3216 —



Spectron EF 12/14 Primary High Offset Stems Cobalt Chromium – ASTM F 799

Size	Stem Length	Implant Cat. No.	Broach/Trial Cat. No.	Trial Neck Cat. No.
	- U			
1H	115 mm	7131-2111	7136-5001	7136-5091
2H	125 mm	7131-2112	7136-5002	7136-5092
3H	135 mm	7131-2113	7136-5003	7136-5093
4H	135 mm	7131-2114	7136-5004	7136-5094
5H	135 mm	7131-2115	7136-5005	7136-5095



Oxinium[†] 12/14 Taper Femoral Heads

28 mm	32 mm
7134-2803	7134-3203
7134-2800	7134-3200
7134-2804	7134-3204
7134-2808	7134-3208



Spectron Invis[†] Distal Contralizors

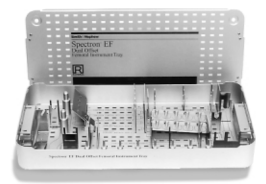
Distal Cellu alizers					
Cat. No. Size O.D.					
7131-3101	1	8 mm			
7131-3102	2	9 mm			
7131-3103	3	10 mm			
7131-3104	4	12 mm			
7131-3105	5	13 mm			

Invis Distal Centralizers

Cat. No.	0.D.	Cat. No.	0.D.
7131-3208	8 mm	7131-3215	15 mm
7131-3209	9 mm	7131-3216	16 mm
7131-3210	10 mm	7131-3217	17 mm
7131-3211	11 mm	7131-3218	18 mm
7131-3212	12 mm	7131-3219	19 mm
7131-3213	13 mm	7131-3220	20 mm
7131-3214	14 mm	7131-3221	21 mm

Neck Length	36 mm
-3	7134-3603
+0	7134-3600
+4	7134-3604
+8	7134-3608

SPECTRON EF 12/14 INSTRUMENTATION



12/14 Dual Offset Sterilization Tray Cat. No. 7136-9112

12/14 Standard Offset Sterilization Tray (Not Shown) Cat. No. 7136-9113

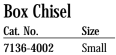


Osteotomy Guide			
Cat. No.	Size		
7136-5036	Sizes 1-5		



Broach Handle Cat. No. 7136-4007







Anteversion Handle Cat. No. 7136-4012



Blunt Medullary Reamer Cat. No. 11-9657



Broac	nes/Trials	
Cat. No.	Size	

JIZC
Size 1
Size 2
Size 3
Size 4
Size 5

SPECTRON EF 12/14 INSTRUMENTATION



Laical Realliers				
Cat. No.	Size			
7136-5023	Size 1–3			
7136-5025	Size 4–5			



Femoral Component Driver Cat. No. 11-9817



12/14 Taper Trial Necks

Size	Primary Collared Cat. No.	Size	Primary High Offset Cat. No.	
1	7136-5081	1H	7136-5091	
2	7136-5082	2H	7136-5092	
3	7136-5083	3H	7136-5093	
4	7136-5084	4H	7136-5094	
5	7136-5085	5H	7136-5095	



Femoral Head Impactor Cat. No. 7136-4009

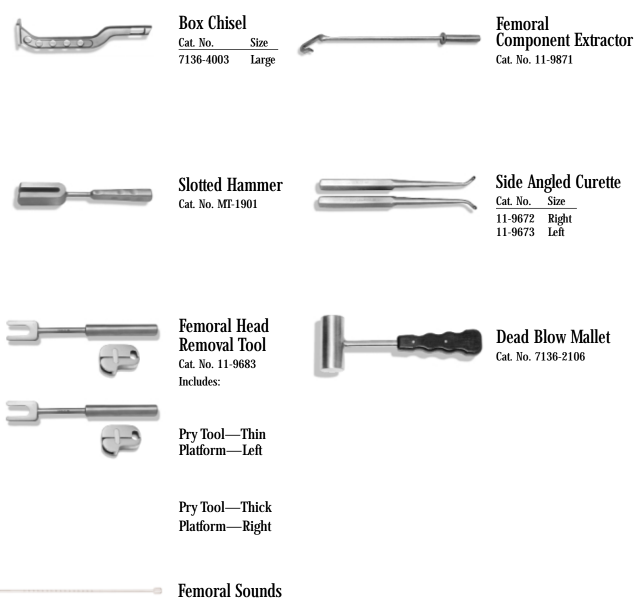


Trial 12/14 Taper Femoral Heads

Neck	Color		I	I	I	
Length	Code	22 mm	*26 mm	*28 mm	32 mm	36 mm
-3	Green			7135-2803	7135-3203	7135-3603
+0	Yellow	7135-2200	7135-2600	7135-2800	7135-3200	7135-3600
+4	Red	7135-2204	7135-2604	7135-2804	7135-3204	7135-3604
+8	White	7135-2208	7135-2608	7135-2808	7135-3208	7135-3608
+12	Blue	7135-2212	7135-2612	7135-2812	7135-3212	7135-3612
+16	Black	—		7135-2816	7135-3216	7135-3616

*Space allowed for 26 mm and 28 mm heads in instrument tray.

ADDITIONAL SPECTRON EF 12/14 INSTRUMENTS



I chiorar sounds			
Cat. No.	Size		
7136-3508	8-9 mm		
7136-3510	10-11 mm		
7136-3512	12-13 mm		
7136-3514	14-15 mm		
7136-3516	16-17 mm		
7136-3518	18-19 mm		



VersaBond^{††} Cat. No. 7127-1140



Femoral Pressurizers

Cat. No. Size 7127-0026 7127-0027 7127-0028

Small Medium Large

 \frown

>



Prep-IM⁺⁺ Total Hip Preparation Kit

Cat. No. 12-1010

- Includes the following:
 - 2 Buck Cement Restrictors
- Femoral Canal Brush 1
- 1 Buck Disposable Inserter
- 1 Femoral Canal Suction Absorber
- 2 Concise Cement Sculps
- 1 Medium Femoral Pressurizer

Buck Cement Restrictors

10		
2	2	
-	-	

Cat. No. **Description** 91-4535 13 mm 12-9418 18.5 mm 25 mm 12-9419 7127-9420 30 mm 35 mm

Vent Opening Tool Cat. No. 11-0028



Buck Cement Restrictor

Cat. No.	Size
12-9418	18.5 mm
12-9419	25 mm
7127-9420	30 mm
7127-9421	35 mm



Vortex[†] Vacuum Mixer

Cat. No. 7127-0070

Vortex Nozzles

Cat. No. 7127-0080 Standard Breakaway Long Tapered Revision Umbrella Re-use Kit (not shown) 7127-0072 Adaptor

Description

7127-0081 7127-0084 7127-0085 7127-0071

7127-9421



Concise Cement Sculps Kit Cat. No. 11-1000 (one of each)



Buck Femoral Cement Restrictor Inserter

Femoral Canal

Cat. No.

11-0037

11-0038

Suction Absorber

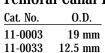
Size

19 mm

25 mm

Cat. No. 11-2428

Femoral Canal Brush



19

CEMENT & ACCESSORIES



Connector, Schraeder Cat. No. 7127-0050



Connector, Drager Cat. No. 7127-0051



MixOR⁺⁺ Vacuum Mixing System with Syringe Cat. No. 7127-0020



Connector, D.I.S.S. Cat. No. 7127-0052



Femoral Cement Compressor Cat. No. 11-1434



Handpiece with Zimmer Coupling Cat. No. 7127-7000



Disposable Femoral Cement Compressor Cap Cat. No. 11-1435



Powerhose with Zimmer Coupling Cat. No. 7127-7001



Hip with Suction Cat. No. 7127-7004

Hip without Suction Cat. No. 7127-7005



MixOR Pump and Hose Kit Cat. No. 7127-0040

MixOR Hose Only (not shown) Cat. No. 7127-0041

MixOR Pump Only (not shown) Cat. No. 7127-0042



InjectOR Gun Cat. No. 7127-2000

IMPORTANT MEDICAL INFORMATION Warnings and Precautions Total Hip System

IMPORTANT NOTE

Total hip replacement (THR) arthroplasty has become a successful procedure for relieving pain and restoring motion in patients who are disabled from hip arthropathy. The goals of total hip replacement are to decrease pain, increase function, and increase mobility. The Constrained Acetabular liner is designed for those THR patients who suffer or are at risk for recurrent dislocations.

MATERIALS

Femoral components are cobalt chromium alloy, titanium 6AI-4V alloy or stainless steel. Femoral heads are cobalt chromium alloy, zirconia ceramic, alumina ceramic, Oxinium' oxidized zirconium or stainless steel. Acetabular liners are ultra-high molecular weight polyethylene or alumina ceramic. All poly acetabular components are ultra-high molecular weight polyethylene. Acetabular shells and the constrained liner adapter are titanium 6AI-4V alloy. The component material is provided on the outside carton label.

Note: Ceramic/ceramic implants are not available in the USA.

Some of the alloys needed to produce orthopedic implants contain some metallic components that may be carcinogenic in tissue cultures or intact organism under very unique circumstances. Ouestions have been raised in the scientific literature as to whether or not these alloys may be carcinogenic in implant recipients. Studies conducted to evaluate this issue have not identified conclusive evidence of such phenomenon, in spite of the millions of implants in use.

DESCRIPTION OF SYSTEM

The Total HIp System consists of femoral components, proximal pads, taper sleeves, distal sleeves, acetabular components, fixation screws and pegs, hole covers, centralizers, and femoral heads. Components may be grit blasted, porous coated, hydroxylapatite (HA) coated, or HA porous coated. All implantable devices are designed for single use only.

Femoral Components

Femoral components are available in a variety of sizes. Porous coated components are coated for biological ingrowth. Proximally and distally modular femoral components accept proximal pads and distal sleeves, respectively. Non-porous femoral components can feature PMMA centralizers that help produce a uniform thickness of cement.

Femoral components are available with a Small (10/12), Large (14/16), or 12/14 global taper.

Small taper femoral components mate and lock directly with a 22 mm metal, oxidized zirconium or ceramic head. The Small taper also mates with a taper sleeve which, in turn, mates with either metal or ceramic heads (26, 28, or 32 mm), bipolar or unipolar components.

Large taper femoral components mate and lock with either metal heads (26, 28, or 32 mm), ceramic heads (28 or 32 mm), bipolars or unipolar components.

Femoral components with a 12/14 taper mate and lock with either metal heads (22, 26, 28, or 32 mm), oxidized zirconium heads (28 or 32 mm), ceramic heads (22, 26, 28, or 32 mm), bipolar or unipolar components.

Small, Large, and 12/14 taper femoral component tapers are machined to mate and lock with ceramic heads, thus preventing rotation of the ceramic head on the stem, which would cause wear of the stem taper.

Taper Sleeves

A taper sleeve is required to be impacted on the Small taper femoral component prior to impacting a femoral head size 26, 28, or 32 mm. A taper sleeve is required to attach a unipolar head. Unipolar taper sleeves are available in Small, Large, and 12/14 tapers. Never place more than one taper sleeve on a femoral component.

Femoral Heads

Cobalt chromium (22, 26, 28, and 32 mm), oxidized zirconium (28 and 32 mm) and ceramic (22, 26, 28, and 32 mm) heads are available in multiple neck lengths for proper anatomic and musculature filt. Heads are highly polished for reduced friction and wear. The following zirconia ceramic heads are available for use only with Small and Large taper femoral components:

Zirconia Ceramic	Head Diameter	Neck Length	
42-7815	32 mm	Standard	0 mm
42-7816	32 mm	Long	+ 4 mm
42-7817	32 mm	X-Long	+ 8 mm
42-7818	28 mm	Standard	0 mm
42-7819	28 mm	Long	+ 4 mm
42-7820	28 mm	X-Long	+ 8 mm

Note: 32 mm heads with a -3 mm neck length are not available for use with the Small taper stems.

In addition to the components listed above, the following components are available for use only with Small taper femoral components:

Zirconia Ceramic	Head Diameter	Neck Length	
7132-0002	22 mm	Long	+ 4 mm
7132-0006	22 mm	X-Long	+ 8 mm

Note: 22 mm Zirconia Ceramic Heads used with Small taper femoral components are not available in the U.S.A.

The following zirconia ceramic heads are available for use only with 12/14 taper femoral components:

1010		,5:011	•	
Zirconia Ceramic	Head Diameter	Neck I	ength	
7132-0028	28 mm	Standard	0 mm	
7132-0428	28 mm	Long	+ 4 mm	
7132-0828	28 mm	X-Long	+ 8 mm	
7132-0026	26 mm	Standard	0 mm	
7132-0426	26 mm	Long	+ 4 mm	
7132-0826	26 mm	X-Long	+ 8 mm	
7132-0422	22 mm	Long	+ 4 mm	
7132-0822	22 mm	X-Long	+ 8 mm	
The following alumina ceramic heads are available for use only with 12/14 taper femoral components:				
Alumina Ceramic Head Diameter		Neck I	ength	
7133-2800	28 mm	Standard	0 mm	
7133-2804	28 mm	Lona	+ 4 mm	

7133-2804 28 mm 7133-2808 28 mm

7133-3200 32 mm 0 mm Standard 7133-3204 32 mm Long + 4 mm 7133-3208 32 mm X-Long + 8 mm Acetabular Components Acetabular components can be one piece all polyethylene, two-piece component consisting of a titanium shell and a polyethylene liner or a titanium shell and an alumina ceramic liner. Constrained liners are two components consisting of a polyethylene liner and a titanium adapter. Please see Warnings and Precautions for specific information on

X-Long

+ 8 mm

screws, pegs and hole covers use. Acetabular reinforcement and reconstruction rings are used with an all polyethylene acetabular component. Note: The metal shell and ceramic liner in the Ceramic/Ceramic Acetabular System are not available in the U.S.A.

Note: 10 Mrad cross-linked polyethylene (UHMWPE) Reflection¹¹ acetabular liners may be used with metal (CoCr) femoral heads or zirconia ceramic femoral heads.

Femoral components and femoral heads are designed for use with any Smith & Nephew polyethylene acetabular component or polyethylenelined, metal-backed acetabular component having an appropriatelysized inside diameter.

INDICATIONS, CONTRAINDICATIONS, AND ADVERSE EFFECTS

Hip components are indicated for individuals undergoing primary and revision surgery where other treatments or devices have failed in rehabilitating hips damaged as a result of trauma or noninflammatory degenerative joint disease (NIDJD) or any of its composite diagnoses of osteoarthritis, avascular necrosis, traumatic arthritis, slipped capital epiphysis, fused hip, fracture of the pelvis, and diastrophic variant.

Hip components are also indicated for inflammatory degenerative joint diseases including rheumatoid arthritis, arthritis secondary to a variety of diseases and anomalies, and congenital dysplasia; old, remote osteomyelitis with an extended drainage-free period, in which case, the patient should be warned of an above normal danger of infection postoperatively; treatments of nonunion, femoral neck fracture and trochanteric fractures of the proximal femur with head involvement that are unmanageable using other techniques; endoprosthesis, femoral osteotomy, or Girdlestone resection; fracture-dislocation of the hip; and correction of deformity.

The hip joint metal/polymer constrained cemented or uncemented prosthesis is intended to replace a hip joint. The constrained liner is intended for primary or revision patients at high risk for hip dislocation due to a history of prior dislocation, bone loss, soft tissue laxity, neuromuscular disease, or intra-operative instability and for whom all other options to constrained acetabular components have been considered.

Acetabular reinforcement and reconstruction rings are intended to be used in primary and revision surgeries where the acetabulum has the deficiencies of the acetabular roof, anterior or posterior pillar, medial wall deficiency, and / or protrusion as a result of the indications listed previously.

Some of the diagnoses listed above and below may also increase the chance of complications and reduce the chance of a satisfactory result. Contraindications

- Conditions that would eliminate or tend to eliminate adequate implant support or prevent the use of an appropriately- sized implant, e.g.:
- a. blood supply limitations;
- insufficient quantity or quality of bone support, e.g., osteoporosis, or metabolic disorders which may impair bone formation, and osteomalacia; and
- c. infections or other conditions which lead to increased bone resorption.
- Mental or neurological conditions which tend to impair the patient's ability or willingness to restrict activities.
- Physical conditions or activities which tend to place extreme loads on implants, e.g., Charcot joints, muscle deficiencies, multiple joint disabilities, etc.
- 4. Skeletal immaturity.
- The zirconia ceramic head is contraindicated for use with any other product than an UHMW polyethylene cup or a metal backed UHMW polyethylene cup.
- 6. The alumina ceramic liner is contraindicated for use with any product other than the metal shell with the correlating inner taper geometry and the appropriate sized alumina ceramic head. The alumina ceramic liner should only be used with the alumina ceramic head.

Contraindications may be relative or absolute and must be carefully weighted against the patient's entire evaluation and the prognosis for possible alternative procedures such as non-operative treatment, arthrodesis, femoral osteotomy, pelvic osteotomy, resection arthroplasty, hemiarthroplasty and others.

Conditions presenting increased risk of failure include: osteoporosis, metabolic disorders which may impair bone formation, and osteomalacia. Possible Adverse Effects

Possible Adverse Effects

- Wear of the polyethylene and ceramic articulating surfaces of acetabular components may occur. Higher rates of wear may be initiated by the presence of particles of cement, metal, or other debris which can develop during or as a result of the surgical procedure and cause abrasion of the articulating surfaces. Higher rates of wear may shorten the useful life of the prosthesis, and lead to early revision surgery to replace the worn prosthetic components.
- 2. With all joint replacements, asymptomatic, localized, progressive bone resorption (osteolysis) may occur around the prosthetic components as a consequence of foreign-body reaction to particulate wear debris. Particles are generated by interaction between components, as well as between the components and bone, primarily through wear mechanisms of adhesion, abrasion, and fatigue. Secondarily, particles may also be generated by third-body particles lodged in the polyethylene or ceramic articular surfaces. Osteolysis can lead to future complications necessitating the removal or replacement of prosthetic components.
- Loosening, bending, cracking, or fracture of implant components may result from failure to observe the Warnings and Precautions below. Fracture of the implant can occur as a result of trauma, strenuous activity, improper alignment, or duration of service.
- 4. Dislocations, subluxation, decreased range of motion, or lengthening or shortening of the femur caused by improper neck selection, positioning, looseness of acetabular or femoral components, extraneous bone, penetration of the femoral prosthesis through the shaft of the femur, fracture of the acetabulum, intrapelvic protrusion of acetabular component, femoral impingement, periarticular calcification, and/or excessive reaming.
- 5. Fracture of the pelvis or femur: post-operative pelvic fractures are usually stress fractures. Femoral fractures are often caused by defects in the femoral cortex due to misdirected rearning, etc. Intraoperative fractures are usually associated with old congenital deformity, improper stem selection, improper broaching, and/or severe osteoporosis.
- 6. Infection, both acute post-operative wound infection and late deep wound sepsis.
- Neuropathies; femoral, sciatic, peroneal nerve, and lateral femoral cutaneous neuropathies have been reported. Temporary or permanent nerve damage resulting in pain or numbness of the affected limb.
- 8. Wound hematoma, thromboembolic disease including venous thrombosis, pulmonary embolus, or myocardial infarction.
- Myositis ossificans, especially in males with hypertrophic arthritis, limited pre-operative range of motion and/or previous myositis. Also, periarticular calcification with or without impediment to joint mobility can cause decreased range of motion.
- Trochanteric nonunion usually associated with early weight bearing and/or improper fixation of the trochanter, when a transtrochanteric surgical approach is used.
- Although rare, metal sensitivity reactions and/or allergic reactions to foreign materials have been reported in patients following joint replacement.
- 12. Damage to blood vessels.
- Traumatic arthrosis of the knee from intraoperative positioning of the extremity.
- 14. Delayed wound healing.
- Aggravated problems of the affected limb or contralateral extremity caused by leg length discrepancy, excess femoral medialization, or muscle deficiency.
- Failure of the porous coating/ substrate interface or hydroxylapatite coating/ porous coating bonding may result in bead separation delamination.
- Stem migration or subsidence has occurred in conjunction with compaction grafting procedures usually resulting from insufficient graft material or improper cement techniques. Varus stem alignment may also be responsible.

WARNINGS AND PRECAUTIONS

The patient should be warned of surgical risks, and made aware of possible adverse effects. The patient should be warned that the device does not replace normal healthy bone, that the implant can break or become damaged as a result of strenuous activity or trauma, and that it has a finite expected service life and may need to be replaced in the future. Do not mix components from different manufacturers. Additional Warnings and Precautions may be included in component literature.

Preoperative

 Use extreme care in handling and storage of implant components. Cutting, bending, or scratching the surface of components can significantly reduce the strength, fatigue resistance, and/or wear characteristics of the implant system. These, in turn, may induce internal stresses that are not obvious to the eye and may lead to fracture of the component. Implants and instruments should be protected from corrosive environments such as salt air during storage. Do not allow the porous surfaces to come in contact with cloth or other fiber-releasing materials.

- Allergies and other reactions to device materials, although infrequent, should be considered, tested for (if appropriate), and ruled out preoperatively.
- 3. Fixation and expected longevity of components expected to be left in place at revision surgery should be thoroughly assessed.
- Surgical technique information is available upon request. The surgeon should be familiar with the technique. Refer to medical or manufacturer literature for specific product information.
- Intraoperative fracture or breaking of instruments can occur. Instruments which have experienced extensive use or excessive force are susceptible to fracture. Instruments should be examined for wear, or damage, prior to surgery.
- Do not cold water quench ceramic components and never sterilize ceramic heads while fixed on the stem taper. (See sterilization section, below.)
- Select components such that the Zirconia ceramic, oxidized zirconium, and alumina heads always articulates with a UHMW polyethylene cup or a metal-backed UHMW polyethylene cup. Zirconia ceramic, oxidized zirconium, and alumina heads should never articulate against metal because severe wear of the metal will occur.
- Select only Smith & Nephew femoral components that indicate their use with ceramic heads. This is important because the taper on the stem is machined to tightly mate and lock with the ceramic head thus preventing rotation of the ceramic head on the stem. Also, an improperly dimensioned taper could result in fracture of the ceramic head.
- 9. The zirconia ceramic head is composed of a new ceramic material with limited clinical history. Although mechanical testing demonstrates that when used with a polyethylene acetabular component, the yttria stabilized zirconia ball produces a relatively low amount of particulates, the total amount of particulate remains undetermined. Because of the limited clinical and preclinical experience, the biological effect of these particulates can not be predicted.
- 10. Alumina ceramic should never articulate against metal because severe wear could occur.

Intraoperative

- The general principles of patient selection and sound surgical judgment apply. The correct selection of the implant is extremely important. The appropriate type and size should be selected for patients with consideration of anatomical and biomechanical factors such as patient age and activity levels, weight, bone and muscle conditions, any prior surgery and anticipated future surgeries, etc. Generally, the largest cross-section component which will allow adequate bone support to be maintained is preferred. Failure to use the optimum-sized component may result in loosening, bending, cracking, or fracture of the component and/or bone.
- 2. Correct selection of the neck length and cup, and stem positioning, are important. Muscle looseness and/or malpositioning of components may result in loosening, subluxation, dislocation, and/or fracture of components. Increased neck length and varus positioning will increase stresses which must be borne by the stem. The component should be firmly seated with the component insertion instruments.
- Care should be taken not to scratch, bend (with the exception of the Reconstruction Rings) or cut implant components during surgery for the reasons stated in Number One of the "Pre-Operative" section of "Warnings and Precautions."
- A +12 mm or +16 mm femoral head should not be used with any Small taper stems.
- Distal sleeves should not be used to bridge cortical defects that lie within 25 mm of the tip of the base stem.
- 6. Matrix¹¹ Small taper stem sizes 8S 10L must have a minimum neck length of +8 mm when used with a bipolar component; and Small taper stem sizes 12S - 16L must have a minimum neck length of +4 mm when used with a bipolar component.
- 7. Modular heads and femoral components should be from the same manufacturer to prevent mismatch of tapers.
- Stainless steel heads and stainless steel stems should only be used together. Neither should be used with other metal components.
- 9. Use only Reflection Liners with Reflection Shells.
- Clean and dry stem taper prior to impacting the femoral head or taper sleeve. The modular femoral head component must be firmly seated on the femoral component to prevent disassociation.
- 11. Take care, when positioning and drilling screw and peg holes, to avoid penetration of the inner cortex of the pelvis, penetration of the sciatic notch, or damage to vital neurovascular structures. Perforation of the pelvis with screws that are too long can rupture blood vessels causing the patient to hemorrhage. Do not place a screw in the center hole of the actabular prosthesis. Placement of drills and screws in the anterior or medial portions of the prosthesis is associated with a high risk of potentially fatal vascular injury. Bone screws must be completely seated in the holes of the shell to allow proper locking for the actabular component liner. If the tapered pegs need to be removed from the shell after impaction of the pegs, do not reuse the pegs or the peg shell holes. Use new pegs and different shell holes, or a new shell if necessary.
- USE ONLY REFLECTION' TITANIUM SPHERICAL HEAD BONE SCREWS, UNIVERSAL CANCELLOUS BONE SCREWS, TAPERED PEGS, AND HOLE COVERS with the Reflection Acetabular Component and USE ONLY OPTI-FIX" TITANIUM BONE SCREWS

AND UNIVERSAL CANCELLOUS BONE SCREWS with the Opti-Fix Acetabular Component. Reflection Spherical Head Screws are only for use with SP3, FSO and InterFit Shells. The Reflection SP3, V, InterFit" and the Reflection For Screws Only (FSO) shells accept Universal Cancellous, Reflection screws, and tapered screw-hole covers, not pegs. Locking Head Pegs and Reflection Screw Hole Covers are only for use with SP3 Shells. Tapered pegs can only be used with Reflection V Shells. The threaded center hole in Reflection Shells only accepts the threaded hole cover, not screws or pegs. The InterFit threaded hole cover is only for use with Reflection InterFit, SP3, Spiked and No Hole Shells. The Reflection threaded hole cover can be used with all Reflection shells. Refer to product literature for proper adjunctive fixation and hole cover usage.

- 13. Prior to seating modular components, surgical debris including tissue must be cleaned from the surfaces. Debris, including bone cement, may inhibit the component locking mechanism. If the shell is to be cemented in place, remove extraneous cement with a plastic sculps tool to ensure proper locking of the liner. During liner insertion, make sure soft tissue does not interfere with the shell/liner interface. Chilling the liner reduces the impaction force required to seat the liner. Modular components must be assembled securely to prevent disassociation. Debris inhibits the proper fit and locking of modular components which may lead to early failure of the procedure. Failure to properly seat the acetabular liner into the shell. can lead to disassociation of the liner from the shell.
- Avoid repeated assembly and disassembly of the modular components which could compromise the critical locking action of the locking mechanism.
- 15. Care is to be taken to assure complete support of all parts of the device embedded in bone cement to prevent stress concentration which may lead to failure of the procedure. During curing of the cement, care should be taken to prevent movement of the implant components.
- 16. If the head is removed from a femoral component that will be left in place at revision surgery, it is recommended that a metal head be used. A ceramic head may fracture from irregularities on the femoral component taper.
- 17. If components are to be left in place at revision surgery, they should first be thoroughly checked for signs of looseness, etc. and replaced if necessary. The head/neck component should be changed only when clinically necessary.
- 18. Once removed from the patient, implants previously implanted should never be reused, since internal stresses which are not visible may lead to early bending or fracture of these components.
- 19. With the congenitally dislocated hip, special care should be taken to prevent sciatic nerve palsy. Also, note that the femoral canal is often very small and straight and may require an extra-small straight femoral prosthesis; however, a regular-sized prosthesis should be used when possible. Note that the true acetabulum is rudimentary and shallow. A false acetabulum should not ordinarily be utilized as a cup placement site for anatomical and biomechanical reasons.
- 20. With rheumatoid arthritis, especially for those patients on steroids, bone may be extremely osteoporotic. Care should be taken to prevent excessive penetration of the acetabular floor or fracture of the medial acetabular wall, femur, or greater trochanter.
- 21 Revision procedures for previous arthroplasty, Girdlestone, etc., are technically demanding and difficult to exercise. Common errors include misplacement of the incision, inadequate exposure or mobilization of the femur, inadequate removal of ectopic bone, or improper positioning of components. Postoperative instability as well as excessive blood loss can result. In summary, increased operative time, blood loss, increased incidence of pulmonary embolus and wound hematoma can be expected with revision procedures.
- 22. Prior to closure, the surgical site should be thoroughly cleaned of cement, bone chips, ectopic bone, etc. Ectopic bone and/or bone spurs may lead to dislocation or painful or restricted motion. Range of motion should be thoroughly checked for early contact or instability.
- Proper positioning of the components is important to minimize impingement which could lead to early failure, premature wear, and/or dislocation.
- 24. In order to minimize the risks of dislocation and loosening of the shell-acetabular bone or shell-bone cement interface that may occur when using a metallic shell intended for biological fixation or cemented use only, surgeons should consider providing immediate resistance to tensile forces between the metallic shell and the acetabular bone or bone cement interface through the use of orthopedic bone fixation devices such as bone screws, spikes, screw threads, pegs, fins, or other bone fixation devices.
- 25. To correctly position the metallic locking ring that is a part of the constrained liner assembly, surgeons should consult the manufacturer's instructions for appropriate device assembly.
- 26. Physicians should consider component malposition, component placement, and the effect on range of motion when using modular heads (with sleeves or skirts) and extended liners.

Postoperative

1. Postoperative directions and warnings to patients by physicians, and patient care, are extremely important. Gradual weight bearing is begun after surgery in ordinary total hip arthroplasty. However, with trochanter osteotomy or certain complex cases, weight-bearing status should be individualized with the non or partial weightbearing period extended.

- Patients should be warned against unassisted activity, particularly use of toilet facilities and other activities requiring excessive motion of the hip.
- 3. Use extreme care in patient handling. Support should be provided to the operative leg when moving the patient. While placing the patient on bedpans, changing dressings, and clothing, and similar activities, precautions should be taken to avoid placing excessive load on the operative part of the body.
- Postoperative therapy should be structured to regain muscle strength around the hip and a gradual increase of activities.
- 5. Periodic X-rays are recommended for close comparison with immediate postoperative conditions to detect long-term evidence of changes in position, loosening, bending and/or cracking of components or bone loss. With evidence of these conditions, patients should be closely observed, the possibilities of further deterioration evaluated, and the benefits of early revision considered.
- Prophylactic antibiotics should be recommended to the patient similar to those suggested by the American Heart Association for conditions or situations that may result in bacteremia.

PACKAGING AND LABELING

Components should only be accepted if received by the hospital or surgeon with the factory packaging and labeling intact. If the sterile barrier has been broken, return the component to Smith & Nephew, Inc.

STERILIZATION/RESTERILIZATION

Most implants are supplied sterile and have been packaged in protective trays. The method of sterilization is noted on the package label. All radiation sterilized components have been exposed to a minimum of 25 kiloGrays of gamma radiation. If not specifically labeled sterile, the implants and instruments are supplied non-sterile and must be sterilized prior to use. Inspect packages for punctures or other damage prior to surgery.

Metal Components

Nonporous or non-HA coated metal components and oxidized zirconium heads may be initially sterilized or resterilized, if necessary, by steam autoclaving in appropriate protective wrapping, after removal of all original packaging and labeling. Protect the devices, particularly mating surfaces, from contact with metal or other hard objects which could damage the product. The following process parameters are recommended for these devices:

- Prevacuum Cycle: 4 pulses (Maximum = 26.0 psig (2.8 bars) & Minimum = 10.0 inHg (339 millibars)) with a minimum dwell time of 4 minutes at 270°F to 275°F (132°C to 135°C), followed by a 1 minute purge and at least 15 minutes of vacuum drying at 10 inHg (339 millibars) minimum.
- For the United Kingdom, cleaning and sterilization should be carried out in accordance with HTM 2030 and HTM 2010. The recommended prevacuum sterilization cycle is: Evacuation to 100mBar for 2-3 minutes, Negative Pressure pulsing (5): 800mBar-100mBar, Positive Pressure pulsing (5): 2.2Bar – 1.1 Bar, Sterilization exposure: 3 minutes at 134°-137°C, Drying vacuum 40mBar for 5-10 minutes. Note: mBar absolute.
- Gravity Cycle: 270°F to 275°F (132°C to 135°C) with a minimum dwell time at temperature of 10 minutes, followed by a 1 minute purge and at least 15 minutes of vacuum drying at 10 inHg (339 millibars) minimum.

Smith & Nephew does not recommend the use of low temperature gravity cycles or flash sterilization on implants.

Do not resterilize femoral prostheses with ceramic heads seated on the stem. Do not steam autoclave femoral prostheses with proximal or distal centralizers attached. If resterilization is required for femoral prostheses with proximal or distal centralizers attached, use ethylene oxide gas.

If porous coated or HA coated implants are inadvertently contaminated, return the unsolied prosthesis to Smith & Nephew for resterilization. DO NOT RESTERILIZE porous coated or HA coated implants. The porous coating requires special cleaning procedures.

Plastic Components

Plastic components may be resterilized by ethylene oxide gas. The following parameters are recommended as starting points for cycle validation by the health care facility:

Sterilant	Temp.	Humidity	Maximum Pressure	Concen- tration	Exposure Time
100% EtO	131'F	40-80%	10 PSIA	725	60-180
	(55°C)	(70% Target)	(689 millibar)	mg/l	minutes

Suggested initial starting point for aeration validation is 12 hours at 120°F (49°C) with power aeration. Consult aerator manufacturer for more specific instructions.

Ceramic Components

Do not resterilize ceramic femoral heads or liners.

INFORMATION

For further information, please contact Customer Service at (800) 238-7538 for calls within the continental USA and (901) 396-2121 for all international calls.

Authorized EC Representative: Smith & Nephew Orthopaedics GmbH, Tuttlingen, Germany.

Caution: Federal Law (USA) restricts this device to sale by or on the order of a physician.

H₂O₂ – hydrogen peroxide sterilization

†Oxinium is a trademark of Smith & Nephew. U.S. Patent #5,037,438 ††InterFit, Opti-Fix and Reflection are trademarks of Smith & Nephew. REG. U.S. PAT. & TM. OFF.

07/02

NOTES



First Choice in Orthopaedics

Smith & Nephew, Inc. • 1450 Brooks Road • Memphis, TN 38116 U.S.A. (901) 396-2121 • For information: 1-800-821-5700 • For orders and order inquiries: 1-800-238-7538

+Buck, Concise, Invis and Vortex are trademarks of Smith & Nephew.
+Oxinium is a trademark of Smith & Nephew. U.S. Patent #5,037,438
++InterFit, Matrix, MixOR, Opti-Fix, Prep-IM, Reflection, Spectron and VersaBond are trademarks of Smith & Nephew. REG. U.S. PAT. & TM. OFE
©2003 Smith & Nephew, Inc.