IMPLANTS For Total Knee Replacement



IMPORTANT PATIENT LEAFLET





READ CAREFULLY THIS LEAFLET BEFORE SURGERY.

IT CONTAINS IMPORTANT INFORMATION FOR YOU.

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AMPLITUDE

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KEEP THIS LEAFLET

You may need to read it again. If you have any further questions, ask your surgeon.

If you experience any side effect, including **possible side effects not listed** in this leaflet, talk to your doctor or pharmacist. See page 4.

This leaflet has been written assuming the person receiving the implants is reading it .

INTENDED USE

The Total Knee system is intended to replace the natural knee joint (knee arthroplasty) through an artificial system.

Patients with a mature skeleton may be implanted with a total knee prosthesis in cases of severe joint pathology (cartilage wear, inflammation like arthritis or after an injury such as a bone fracture after a fall) and where other devices or treatments have failed.

Please ask your surgeon for details. The decision to implant a total knee prosthesis is left to the surgeon after evaluation of the risk/benefit balance and after discussion with you.

WARNING AND PRECAUTIONS

You must be informed the factors that could compromise the success of the surgery and post operative results:

- Overweight;
- Case history detailing infections and/or falls;
- Metabolic disorders that reduce your resistance or induce progressive bone deterioration;
- Local bone tumours;
- Severe bone deformities;
- Severe osteoporosis;
- Playing sports intensively;

- Playing risky sports or engaging in risky activities;
- Addictive behaviour.

SPECIFIC WARNINGS & MRI INFORMATION

The materials used in your implant may trigger security gates/scanners.

PLEASE INFORM PERSONNEL ABOUT YOUR IMPLANTED DEVICE.

Non-clinical testing demonstrated that the implants are **MR Conditional**. A patient with one of these devices can be scanned safely in a MR system under the following conditions:

- Static magnetic field of 1.5T or 3T for a 15min-procedure;
- Maximum spatial gradient field of 1,500 Gauss/cm;
- Normal operation mode only;
- The image artefact caused by the implants extends approximately 99mm from these implants. MR images may be blurred next to the implants.

	8.6°C orimetry WB-SAR of 3.27W/kg)	Maximum measured temperature rise with 1.5T MRI, for a 15min procedure	A
with 3T MRI, for a 15min procedure (calorimetry WB-SAR of 2.39W/kg)	9.9°C orimetry WB-SAR of 2.39W/kg)	Maximum measured temperature rise with 3T MRI, for a 15min procedure	



REPORTING ADVERSE EVENTS

Any incident in relation to the device shall be reported to your surgeon, healthcare facilities or family doctor.

YOU HAVE BEEN IMPLANTED WITH:

The surgeon needs to tick boxes corresponding to the devices you have been implanted with.

Associations with other devices have to be validated by Amplitude.

FEIVIORAL COIVIPONENTS	
SCORE II Femoral component - cementless	
SCORE II Femoral component - cemented	
TIBIAL COMPONENTS	
SCORE Tibial baseplate - cementless	
SCORE Tibial baseplate - cemented	
SCORE II Tibial mobile bearing insert	
Tibial augment for Total Knee Prosthesis - cemented	
Offset adapter for Total Knee Prosthesis	
Extension stem for Total Knee Prosthesis - cemented	
PATELLAR COMPONENTS	
Resurfacing patellar implant - cemented	
Resurfacing patellar implant NM - cemented	
Inset patellar implant - cemented	
Inset patellar implant - cementless	
MATERIALS	
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Cobalt-Chromium alloy Cr: 26.50% - 30.00% / Mo: 4.50% - 7.00% / Ni ≤ 1.00% / Fe ≤ 1.00% / C ≤ 0.35% / Mn ≤ 1.00% / Si ≤ 1.00% / Co: Balance	SCORE II Femoral component - cemented SCORE Tibial baseplate - cemented
Cobalt-Chromium alloy Cr: 26.50% - 30.00% / Mo: 4.50% - 7.00% / Ni ≤ 1.00% / Fe ≤ 1.00% / C ≤ 0.35% / Mn ≤ 1.00% / Si ≤ 1.00% / Co: Balance Hydroxyapatite coating (pentacalcium hydroxide tris(orthophosphate) Ca5(PO4)3OH: 100%) & titanium undercoating (C ≤ 0.10% / H ≤ 0.30% / Fe ≤ 0.60% / N ≤ 5.00% / O ≤ 10.00% / Ti: Balance)	SCORE II Femoral component - cementless SCORE Tibial baseplate - cementless
Polyethylene	SCORE II Tibial mobile bearing insert
PE: 100%. Possible traces of Ti, Al, Ca, Cl	Patellar components
	Standard keel (provided with tibial baseplates - cemented or cementless)
Stainless steel C ≤ 0.08% / Si ≤ 0.75% / Mn : 2.00% - 4.25% / Ni : 9.00% - 11.00% /	Tibial augment for Total Knee Prosthesis - cemented
Cr : 19.50% - 22.00% / Mo : 2.00% - 3.00% / Nb : 0.25% - 0.80% / S ≤ 0.01% / P ≤ 0.025% / Cu ≤ 0.25% / N : 0.25% - 0.50% / Fe: Balance	Offset adapter for Total Knee Prosthesis
	Extension stem for Total Knee Prosthesis cemented

DEVICES EXPECTED LIFETIME

The SCORE II total knee prosthesis expected survivorship is at least 95% at 10 years for primary cases and 80% at 10 years for revision cases. This can be reduced or increased depending on your activity level or events that could compromise implants integrity (falls, accidents...).

Please attend regular check-ups according to your surgeon's advice.

CONTRAINDICATIONS

At the time of the surgery, these conditions are not recommended:

- Pregnancy and breastfeeding;
- Allergy to the implant materials;
- Infections;
- Severe mental, muscular, neurological or vascular deficiencies affecting the limb in question;
- Destruction of bone or poor bone quality which may affect stability of the implant;
- Highly localized arthrosis requiring osteotomy or unicompartmental arthroplasty;
- Anatomic disorder requiring a constrained or hinge prosthesis.

These conditions should be discussed with your surgeon. Any pathology (even if not listed above) must be mentioned to your surgeon beforehand as well.

POSSIBLE SIDE EFFECTS

You must be informed about the side effects in relation with the device that might occur. Those recorded by Amplitude are all rare (lower than 1 per 1000 cases) or occasional* side effects (between 1 for 1000 cases and 1 for 100 cases). Please consult your surgeon if any doubt.

- Difficult recovery after major surgery
- Injury to a major artery or nerve in the lower limb
- Disruption or rupture of the thigh tendons
- Tear of a thigh muscle
- Infection
- Stiffness*
- Instability
- Pain
- Patellar problems
- Allergy to implanted materials

- Bone degeneration
- Tissue reactions to prosthetic debris and wear particles
- Bone fracture
- Prosthesis dislocation
- Prosthesis loosening
- Premature wear on joint surfaces
- Prosthesis breakage
- Prosthesis displacement
- Friction/conflict between the prosthesis and the surrounding tissue

Other events linked to the surgery or current knowledge can also occur:

- Phlebitis, fat or pulmonary embolism
- Bruises
- Complex regional pain syndrome (Algoneurodystrophy)

- Bone ossification abnormalities
- Poor bone healing

You must also be aware of the risks and possible side effects of surgery - please ask your surgeon for further information.

After a joint replacement, you should pay attention during your daily life: any blow, fall or accident may **jeopardise** your recovery or your implant lifetime. Please ask your surgeon for further information.

