Comparison of the Low Contact Stress and Press Fit Condylar Rotating-Platform Mobile-Bearing Prostheses in Total Knee Arthroplasty
A Prospective Randomized Study

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Background: To our knowledge, no study to date has compared the clinical results of posterior cruciate-sacrificing mobile-bearing total knee replacements with those of posterior-stabilized mobile-bearing total knee replacements in the same patients. The purpose of the present study was to compare the clinical and radiographic results of these two designs. We hypothesized that the results would be better for knees treated with the posterior-stabilized mobile-bearing prosthesis.

Methods: The present study consisted of a consecutive series of 107 female patients (mean age, 66.8 years) who underwent bilateral simultaneous total knee arthroplasty at the same surgical setting. All of these patients received a posterior cruciate-sacrificing mobile-bearing prosthesis in one knee and a posterior-stabilized mobile-bearing prosthesis in the contralateral knee. At the time of each follow-up (mean, 7.4 years; range, seven to 7.6 years), the patients were assessed clinically.

Results: The mean postoperative Knee Society knee score (96 compared with 97 points) and Hospital for Special Surgery knee score (93 compared with 94 points) were similar between the two groups. At the time of the latest follow-up, the average range of motion was 127.7° (range, 70° to 150°) in the knees with a posterior cruciate-sacrificing mobile-bearing prosthesis and 132.4° (range, 90° to 150°) in the knees with a posterior-stabilized mobile-bearing prosthesis. With a margin of error of the manual measurement of 5°, this difference was not significant. The estimated survival rate was 97.2% (95% confidence interval, 91% to 99%) at seven years in the posterior-cruciate sacrificing mobile-bearing prosthesis group and 98.1% (95% confidence interval, 92% to 99%) at seven years in the posterior-stabilized mobile-beariing prosthesis group.

Conclusions: After a minimum duration of follow-up of seven years, we found no significant differences between the two groups with regard to the clinical and radiographic results, including knee range of motion.

Level of Evidence: Therapeutic Level I. See Instructions to Authors for a complete description of levels of evidence.

The posterior cruciate-sacrificing mobile-bearing total knee prosthesis (Low Contact Stress Rotating Platform [LCS RP]; DePuy, Johnson & Johnson, Warsaw, Indiana) was introduced by Buechel and Pappas in 1979 to reduce contact stress in the polyethylene and potentially to decrease wear as well as to minimize cement-bone stress at the tibial surface. Compared with the fixed-bearing posterior-stabilized total knee prosthesis introduced by Insall et al. in 1978, the LCS RP device has no so-called post-and-cam mechanism. Instead, stability is provided by the curved design of the tibial insert articulation surface and a balanced flexion gap achieved with exacting surgical technique. Over the past twenty years, good results have been reported in association with the LCS RP device.

The Press Fit Condylar Sigma posterior-stabilized rotating-platform knee (PFC Sigma PS-RP; DePuy, Johnson & Johnson, Warsaw, Indiana) was introduced in 2000. This design was introduced to improve the kinematics of the LCS RP prosthesis by employment of a post-and-cam mechanism. It was anticipated that the post-and-cam mechanism in the PFC
Sigma PS-RP prosthesis would lead to consistent posterior rollback, which, in turn, would lead to better knee range of motion, would reduce polyethylene wear at the articular surface and undersurface, and would provide better stabilization of the tibial insert (Figs. 1-A and 1-B).

Although the design features of the PFC Sigma PS-RP prosthesis have been proposed to improve upon the kinematics of the LCS RP prosthesis, no study, to our knowledge, has compared the clinical results of the PFC Sigma PS-RP prosthesis with those of the LCS RP prosthesis in the same patients. To examine the results associated with the LCS RP and PFC Sigma PS-RP total knee prostheses in patients who had bilateral simultaneous total knee arthroplasty, we sought to determine whether the knee and function scores and the radiographic results for the knees with a PFC Sigma PS-RP prosthesis would be better than those with an LCS RP prosthesis and whether the knees with a PFC Sigma PS-RP prosthesis would have a better range of motion.

**Materials and Methods**

One hundred and twenty-six patients (252 knees) with bilateral knee osteoarthritis (Ahlbäck grade III, IV, or V) underwent simultaneous bilateral sequential total knee arthroplasty. The study protocol and consent forms were approved by the institutional review board. A detailed informed consent form was signed by each patient, and all information was kept confidential. This study was registered in the ClinicalTrials.gov Protocol Registration System (trial number, NCT01075620). Seven patients were excluded because they refused to participate. Seven more patients were excluded because they were male, leaving 112 patients available for participation. Five patients were lost to early follow-up (three
months), leaving 107 patients (214 knees) available for study after a minimum duration of follow-up of seven years (mean, 7.4 years; range, seven to 7.6 years). The study group included 107 women who had a mean age (and standard deviation) of 66.8 ± 5.181 years (range, fifty-four to eighty-one years) at the time of surgery (see Appendix). Twenty-three knees had valgus alignment of 8° to 12°, and the remaining 191 knees had varus alignment of 8° to 20°. Fourteen of 107 patients with an LCS RP total knee prosthesis and twelve of 107 patients with a PFC Sigma PS-RP total knee prosthesis had previous arthroscopic debridement, and the remaining patients had had no previous surgery.

The coronal geometries of both the LCS RP prosthesis and the PFC Sigma PS-RP prosthesis are rounded coronal designs with similar conformity ratios; the contact surface is slightly greater for the PFC Sigma PS-RP prosthesis. The femoral component of the PFC Sigma PS-RP prosthesis has a cam for the tibial post. The sagittal designs of both the LCS RP prosthesis and the PFC Sigma PS-RP prosthesis are multiradiaral. The anterior flange angle is 5° for the LCS RP prosthesis and 0° for the PFC Sigma PS-RP prosthesis. The posterior flange thickness is 8 mm for all sizes of the PFC Sigma PS-RP prosthesis except for size 6 (for which it is 10 mm) and ranges from 6.2 to 9.4 mm in the LCS RP prosthesis.

Randomization to treatment with the LCS RP or PFC Sigma PS-RP total knee prosthesis was accomplished with use of a sealed study number envelope. After the envelope was opened in the operating room before the skin incision was made, the first knee received the prosthesis indicated by the envelope and the contralateral knee received the other prosthesis. There were no cases in which the second procedure was aborted because of intraoperative complications.

All procedures were performed by the senior author (Y.-H.K.). With tourniquet inflation to 250 mm Hg, an anterior midline skin incision (10 to 12 cm in length) was made, followed by a medial parapatellar capsular incision. In the LCS RP group, tibial preparation was performed first, and in the PFC Sigma PS-RP group, femoral preparation was performed first. Ten millimeters of tibial bone was resected, referenced from the least-involved tibial plateau, to achieve a surface perpendicular to the axis of the tibia in the coronal plane. A 7° posterior slope was prepared in the sagittal plane for the knees in the LCS RP group, and a 0° slope was prepared for the knees in the PFC Sigma PS-RP group. Anterior cortical reference was used for the anterior-posterior cut of the distal part of the femur. Femoral component rotation was determined with use of three reference axes: (1) the transepicondylar axis, (2) the midtrochlear posterior slope was prepared in the sagittal plane for the knees in the LCS RP group, (3) the anatomic axis of the limb, the alignment of the components, posterior slope, posterior femoral condylar offset, the level of the joint line, the presence and location of radiolucent lines at the bone-cement or cement-implant interface, and patellar tilt or dislocation by Knee Society scores (Figs. 2-A and 2-B).

All radiographs were made under fluoroscopic guidance to control rotation of the knee.

**Statistical Analysis**

An a priori power calculation was performed with use of a clinically relevant difference in range of motion of 5° and a standard deviation of 9°. For an effect size of 20% in early functional outcome, as measured with a validated instrument such as the linear analog scale assessment for range of motion, with α = 0.05 and β = 0.80, calculation revealed that 104 knees would be needed in each group. In addition to the required number of subjects, ten more patients were recruited to allow for possible attrition. The changes in the Knee Society and Hospital for Special Surgery knee scores were evaluated with use of the paired t test. Pain scores were assessed with use of the chi-square test. Knee motion was compared between the two groups with use of a paired t test. Complication rates and radiographic data were compared between the two groups with a paired t test. The level of significance was set at \( p < 0.05 \).

**Source of Funding**

There was no external funding for this study.

**Results**

The Knee Society and Hospital for Special Surgery knee scores did not differ significantly between the two groups either preoperatively (\( p = 0.612 \) and \( p = 0.291 \), respectively; paired t test) or postoperatively (\( p = 0.167 \) and \( p = 0.087 \), respectively; paired t test). In the LCS RP group, the mean postoperative Knee Society knee score was 96 points (range, 77 to 100 points) and the mean postoperative Hospital for Special Surgery knee score was 93 points (range, 69 to 100 points). In the PFC Sigma PS-RP group, the mean postoperative Knee Society knee score was 97 points (range, 79 to 100 points) and the mean postoperative Hospital for Special Surgery knee score was 94 points (range, 75 to 100 points). In the LCS RP group, ninety knees (84%) had no pain, sixteen (15%) had mild pain, and one (1%) had moderate pain at the time of the latest follow-up. In the PFC Sigma PS-RP group, eighty-eight knees (82%) were pain-free, eighteen (17%) had mild pain, and one (1%) had moderate pain at the time of the latest follow-up.

The mean preoperative range of motion was 127.8° (range, 80° to 150°) in the LCS RP group and 127.2° (range, 85° to 150°) in the PFC Sigma PS-RP group. The mean postoperative range of motion was 127.7° (range, 70° to 150°) in the LCS RP group and 132.4° (range, 90° to 150°) in the
PFC Sigma PS-RP group. This difference was significant ($p < 0.0001$; paired t test). However, if the margin of error of the manual measurement is considered to be $5^\circ$, this difference is not significant ($p = 0.781$; paired t test). The average improvement of the range of motion per patient was $2.5^\circ$ (range, $2^\circ$ to $10^\circ$) in the LCS RP group and $4.2^\circ$ (range, $-15^\circ$ to $12^\circ$) in the PFC Sigma PS-RP group. This difference was not significant ($p = 0.635$; paired t test). Seventeen knees (16%) in
the LCS RP group and fourteen knees (13%) in the PFC Sigma PS-RP group had <110° of motion at the time of the latest follow-up. The maximum flexion in both groups was 150°. The activity level score for the patients was 5 or 6 points at the time of the latest follow-up, indicating participation in strenuous farm work (5 points) or participation in tennis (6 points).

There were no significant differences between the groups in terms of the alignment of the knee (mean, 5.8° of valgus in both groups), the position of the femoral and tibial components in the coronal and sagittal planes, the patellar angle, the posterior slope of the tibia (mean, 4.9° compared with 2.5°), the amount of the tibial surface area that was covered by the implants (tibial capping), the mean level of the joint line, the prevalence of radiolucent lines, or the posterior condylar offset (mean, 24.2 compared with 24.5 mm) (p > 0.05 for all comparisons; paired t test). The prevalence of radiolucent lines measuring <1 mm (on the tibial side only) was 11% (twelve knees) in the LCS RP group and 9% (ten knees) in the PFC Sigma PS-RP group. The prevalence of osteolysis was 1.9% (two knees) in the LCS RP group and 2.8% (three knees) in the PFC Sigma PS-RP group (Table I).

In the LCS RP group, the estimated survival rate according to Kaplan-Meier analysis was 97.2% (95% confidence interval, 91% to 99%) at seven years, with an overall revision rate of 2.8% (three of 107 knees). In the PFC Sigma PS-RP group, the estimated survival rate was 98.1% (95% confidence interval, 92% to 99%) at seven years, with an overall revision rate of 1.9% (two of 107 knees).

Three knees (2.8%) in the LCS RP group and two knees (1.9%) in the PFC Sigma PS-RP group had a deep infection and required a two-stage revision. None of these five knees had had a recurrence of infection at the time of the latest follow-up. One knee (0.9%) in the LCS RP group required open reduction and internal fixation for the treatment of a supracondylar fracture of the femur. Two knees (1.9%) in the PFC Sigma PS-RP group had a patellar clunk syndrome and required arthroscopic debriement, with good results. Instability did not occur in any knee in either group.

### Discussion

Our study investigated whether the PFC Sigma PS-RP prosthesis for total knee arthroplasty provides a greater benefit than the LCS RP prosthesis. Multiple studies have shown that, regardless of the criteria used to measure success or failure, the LCS RP prosthesis achieved essentially equal, or even better, results than the PFC Sigma PS-RP. We found that the intermediate-term clinical outcomes for both prostheses were similar in terms of the Knee Society score, radiographic results, and range of motion. Our survivorship data are agreement with those in other reports on these prosthetic designs.

In previous studies, the mean flexion of the knee has ranged from 102° to 113° for knees with the LCS RP prosthesis and from 101.7° to 130° for knees with the PFC Sigma PS-RP prosthesis. Our patients had comparable range of motion in comparison with the patients in those reports, and we found no clinical difference between the two designs. Because all of our patients had full knee extension, our findings suggest that the PFC Sigma PS-RP design provides no advantage in terms of knee motion.
In the current study, no knee had aseptic loosening or osteolysis that resulted in revision, similar to previous reports on these knee designs. Our findings of a low prevalence of osteolysis in both groups may be related to the inclusion of only female patients, the use of a polished cobalt-chromium tibial baseplate to reduce backside wear of the insert, the use of a polyethylene insert sterilized with gamma irradiation in a vacuum, and the short shelf life of the insert. It is possible that the duration of follow-up was not sufficiently long to reveal osteolysis.

It has been emphasized that an exacting surgical technique, especially balancing of flexion and extension gaps, is mandatory during a mobile-bearing total knee arthroplasty in order to avoid bearing dislocation or instability of the knee. Many surgeons believe that the use of an unconstrained mobile-bearing total knee implant is contraindicated in cases of severe varus and valgus deformity. This idea was challenged by Beverland, who stated that a mobile-bearing total knee implant can be used for virtually every primary total knee arthroplasty, irrespective of deformity. In our series, we were able to use a mobile-bearing total knee implant for every primary total knee arthroplasty selected by the process of randomization, irrespective of the range of deformity, with no postoperative instability.

The present study had limitations. First, it is difficult for a patient who has undergone bilateral total knee arthroplasty to distinguish the independent function of each knee. Although this was a problem when assessing function after the bilateral total knee arthroplasties, the patients were able to grade which knee caused more functional limitation. Second, the duration of follow-up was seven years and long-term variability in outcome cannot be predicted. Third, we performed no interobserver variability testing on the radiographic measurements. Finally, the range of knee motion was not determined under weight-bearing conditions.

The findings of the present intermediate-term, prospective, randomized clinical study suggest that these first and second-generation mobile-bearing knee designs perform well, with no significant differences between the two prostheses that were evaluated.

Appendix
eA A table summarizing the demographic data for the patients is available with the online version of this article at jbs.org.

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