Uncemented computer-assisted design-computer-assisted manufacture femoral components in revision total hip replacement

A MINIMUM FOLLOW-UP OF TEN YEARS

We prospectively evaluated the long-term outcome of 158 consecutive patients who underwent revision total hip replacement using uncemented computer-assisted design-computer-assisted manufacture femoral components. There were 97 men and 61 women. Their mean age was 63.1 years (34.6 to 85.9). The mean follow-up was 10.8 years (10 to 12).

The mean Oxford, Harris and Western Ontario and McMaster hip scores improved from 41.1, 44.2 and 52.4 pre-operatively to 18.2, 89.3 and 12.3, respectively (p < 0.0001, for each). Six patients required further surgery. The overall survival of the femoral component was 97% (95% confidence interval 94.5 to 99.7). These results are comparable to those of previously published reports for revision total hip replacement using either cemented or uncemented components.

The United Kingdom National Joint Registry has reported an increase in the number of revision total hip replacements (THR) of over 250% between 2003 and 2006 (from 2325 to 6145 cases).1 The first generation of cemented revision THR produced some unsatisfactory results.2,3 Technical advances in implant design and surgical techniques have improved survival but significant rates of loosening and bone loss have persisted.4-10 Complications have also been associated with the use of uncemented femoral components, including subsidence9,11 and proximal femoral fractures.12

The variability of the endosteal geometry after removal of the femoral component can make proximal fit and fill difficult to achieve with an ‘off the shelf’ prosthesis. Whatever the anatomy of the proximal femur, it is important to achieve immediate stability, preserve bone stock and protect the femur from cortical defects which can lead to subsequent fracture.13 In revision THR this requires a large inventory of modular femoral components. The use of a computer-assisted design-computer-assisted manufacture (CAD-CAM) prosthesis obviates this need. Encouraging short-term results have been reported with the use of custom femoral components in revision cases with extensive loss of femoral bone stock.14 However, little has been published on the use of custom-made components in revision THR. We report the results of a large cohort of patients who underwent revision THR using CAD-CAM femoral components.

Patients and Methods

A total of 158 consecutive patients, with no exclusions were included in this prospective study. All patients who underwent revision THR between November 1991 and November 1998 were treated using CAD-CAM femoral components. There were 97 men and 61 women. Their mean age was 63.1 years (34.6 to 85.9). The indications for revision are shown in Table I. Age distribution is shown in Table II.

All operations were performed by the senior author (SMA) via a posterior approach without an extended trochanteric osteotomy. A Harris-Galante I (Zimmer, Warsaw, Indiana) was used in 68 patients (43%) and a Trilogy (Zimmer) component in the remainder.

Uncemented acetabular components were used in all but one case when revision of the acetabular component was indicated. In the case where a cemented component was used, the acetabular defect required impaction allografting, followed by insertion of a reconstruction cage into which a highly cross-linked polyethylene liner was cemented.

The custom-made revision femoral components were manufactured by the Biomedical Engineering Unit (Stanmore Implants Worldwide, Stanmore, United Kingdom). The scanning protocol included a true anteroposterior (AP) radiograph of the pelvis and a lateral view of the affected hip. In order to standardise for magnification, a radio-opaque ruler was placed adjacent to the pelvis and the radiographs were...
sent to the manufacturer to create a computer-assisted design which, once approved by the surgeon, formed the template for the final product. This was delivered to the hospital with a tapered reamer and a custom-made rasp. Each femoral component was manufactured from titanium alloy, Ti6Al4V, with a proximal segment which produced the closest possible line-to-line fit with the endosteal border of the femur. This segment was grit-blasted and coated with an 80 μm thick layer of hydroxyapatite (HA) with high crystallinity. All the components had a collar and a lateral flare in order to minimise axial displacement, theoretically improving stability. The design of the distal aspect of the component was based on the degree of femoral bone loss as classified by Pak et al.15 For type 1 defects components of 120 mm to 140 mm in length were used (Fig. 1). For type 2 defects components of 140 mm to 200 mm with a HA coating on the proximal third and a smooth distal aspect to facilitate their entry and centralisation in the femoral canal were used (Fig. 2). For type 3 defects, components of 140 mm to 200 mm were used (Fig. 3). These were fully coated to achieve maximal osseo-integration and with longitudinal cutting flutes to achieve maximal torsional stability.16-18 Where necessary, the components had an anterior bow to fit the anatomy of the femur.

All patients were reviewed clinically and radiologically at six weeks, three months, and annually thereafter. Function was assessed using the Oxford,19 Harris20 and Western Ontario and McMaster (WOMAC)21 hip scores, and any complications were recorded. Assessment was undertaken by two independent observers (NAS, CK), who also reviewed the radiographs for radiolucent lines, and for signs of stable fixation of the femoral component, including advancement of bone trabeculae to the stem-cement interface, spot welding at the bone-stem implant interface, and any evidence of migration.22 Radiolucent lines were defined as linear lucencies at the bone-prosthesis interface > 2 mm wide and occupying > 30% of any Gruen zone. Lytic lesions were defined as balloon shaped lesions without sclerosis.23 Proximal femoral defects were classified by the Paprosky system.15 Loosening of the acetabular component was classified according to the system described by DeLee and Charnley.24 Statistical analysis. Analysis was performed using the unpaired Student’s t-test (GraphPad, La Jolla, California). The level of significance was set at p = 0.05. Kaplan-Meier survival analysis was performed and failure was defined as revision of the femoral component for any reason.

Results

The mean follow-up was 10.8 years (10 to 12). No patient was lost to follow-up. All patients had relief of pain. The mean flexion of the hip was 95° (90° to 110°).

The mean Oxford, Harris and WOMAC scores improved from 41.1 (15 to 60), 44.2 (7 to 91) and 52.4 (10 to 96) pre-operatively to 18.2 (12 to 49), 89.3 (35 to 100) and 12.3 (0 to 62), respectively (p < 0.0001 for each; Fig. 4). Three patients had intermittent anterior thigh pain with activity: each had Paprosky type 3 femoral bone defects and were treated with long-stemmed components.

**Table I.** Patient characteristics and indications for revision

<table>
<thead>
<tr>
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<th>Number of patients</th>
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<tr>
<td>Total</td>
<td>158</td>
</tr>
<tr>
<td>Men</td>
<td>97</td>
</tr>
<tr>
<td>Women</td>
<td>61</td>
</tr>
<tr>
<td>Mean age in years (range)</td>
<td></td>
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<tr>
<td>Male</td>
<td>63.7 (36.0 to 83.4)</td>
</tr>
<tr>
<td>Female</td>
<td>61.6 (34.6 to 80)</td>
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<tr>
<td>Indications for revision</td>
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<tr>
<td>Aseptic loosening</td>
<td>136</td>
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<tr>
<td>Infection</td>
<td>12</td>
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<tr>
<td>Periprosthetic fracture</td>
<td>6</td>
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<tr>
<td>Recurrent dislocation</td>
<td>3</td>
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<td>Liner wear</td>
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**Table II.** Age distribution of patients

<table>
<thead>
<tr>
<th>Age (yrs)</th>
<th>Number of hips</th>
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<tbody>
<tr>
<td>30 to 39</td>
<td>7</td>
</tr>
<tr>
<td>40 to 49</td>
<td>19</td>
</tr>
<tr>
<td>50 to 59</td>
<td>29</td>
</tr>
<tr>
<td>60 to 69</td>
<td>46</td>
</tr>
<tr>
<td>70 to 79</td>
<td>49</td>
</tr>
<tr>
<td>80 to 89</td>
<td>8</td>
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Fig. 1

Radiograph showing the type 1 design.
All femoral components were well fixed radiologically at the time of the final follow-up. Two (1.3%) had subsided by 2 mm and 3 mm, respectively, on AP radiographs taken three years post-operatively, but stabilised thereafter. Radiolucent lines > 2 mm wide were noted in five patients (1.9%). These were in Gruen zones 1, 2 and 7. These patients were asymptomatic (Table III).

Evidence of stress shielding was noted in two hips (1.3%), and these patients were also asymptomatic. Heterotopic ossification was noted in 22 hips (13.9%). These were all stage 1 to 2 according to the classification of Brooker et al.25

Complications occurred in six hips (3.8%). There were two posterior dislocations, one four days and one five years post-operatively. The first required revision
for acetabular malalignment. Two patients had a deep-vein thrombosis and one of these a pulmonary embolus. One patient suffered an undisplaced oblique periprosthetic fracture one month post-operatively. This was treated using stainless steel cables and eventually united after two years.

There were six revisions, one for instability four days post-operatively, one for peri-prosthetic fracture, three for aseptic loosening of the femoral component (at 5, 10 and 11 years post-operatively) and one for liner wear 12 years post-operatively. No hip developed infection. The ten-year survival, with failure defined as revision for any reason, was 97% (95% confidence interval, 94 to 99.7, Fig. 5).

Discussion

Most patients requiring revision THR have proximal femoral bone loss, which can be compounded by further iatrogenic bone loss that can occur during extraction of the femoral component. There is also considerable natural variation in the shape of the proximal femur. The goals of revision surgery include resolution of pain, as close as possible restoration of biomechanics and preservation of bone stock, all with minimal risk to the patient.

The philosophy of the CAD-CAM design is to optimise fit and fill of the femur, which allows maximal contact of bone to the surfaces available for ingrowth, and optimises the initial stability of the femoral component. Failure to achieve initial stability is a major cause of early failure in revision THR. Proximally HA-coated press-fit femoral components have been shown to achieve early fixation and have superior bone ingrowth in the revision setting compared to non-HA-coated components. They also optimise stress transfer to the proximal femur, with minimal micromovement.

All femoral components used in this study had a collar with a smooth superior surface and an undersurface coated with HA. Our rationale for using a collar was to reduce the risk of subsidence because collars have been shown to limit migration of the femoral component in patients with poor bone stock and improve loading of the proximal femur when bony contact is achieved. A lateral flare was also added to all components to minimise axial migration, further enhancing stability. The benefit conferred by collars is controversial, particularly in the revision setting, where bone stock might be deficient. The individual design of each component, and the fact that these patients were having first-time revisions, meant that we were able to achieve collar-bone contact in most cases. Iatrogenic factors such as bone loss during extraction of the component and variation from the planned neck osteotomy were usually responsible when this contact was not achieved (Fig. 2). In these situations the flared design of the body contributed to obtaining stability.

These femoral components were designed to attain immediate stability as well as minimise the risk of fractures to the proximal femur. Pre-operative planning focused on...
achieving the best line-to-line fit in the metaphyseal region of the femur. Attempts to achieve optimal fill of the metaphysis could lead to a bulky prosthesis, which reduces the chances of bone regeneration and increases the risk of fracture. In cases where metaphyseal bone was deficient, the best diaphyseal fit was achieved. In patients with Paprosky type 3 femoral bone defects, long fully HA-coated components with distal cutting flutes were used. Similar designs with flutes have been shown, in experimental studies to maximise torsional stability. These components require a minimum of 6 cm of intimate bone-implant contact below the distal extent of an osteolytic lesion to achieve satisfactory stability. Stem design in the revision setting therefore requires some flexibility. One factor which facilitated the use of this design was that these patients had relatively well preserved proximal femoral bone stock.

Sotereanos et al reported improved hip scores with no revisions at a mean of 5.3 years using custom-made femoral components. The design of their prosthesis was markedly different and based on a different philosophy to ours. Their series was smaller and the patients had different pathologies, each having had a mean of three previous procedures and having a Paprosky grade IIIb or IV proximal femoral defect. Their patients were therefore in a higher risk group, so it is difficult to compare their results with ours.

Although there have been few studies examining the use of custom-made femoral components for revision THR, authors have used several un cemented off the shelf designs with varying success. Mulliken et al reported a 40% periprosthetic fracture rate, relatively modest improvement in the Harris hip score and inadequate fixation using proximally porous-coated femoral components in the short term. Results of extensively porous-coated components have been more encouraging, with excellent clinical and radiographic results at a mean of 6.3 years follow-up. This study had a smaller cohort than that of Mulliken et al, and their patients had better-preserved femoral bone stock. Excellent long-term results have been reported with fully HA-coated components, with only three failures in 40 complex revision cases at follow-up of a mean of 10.2 years. Garbuz et al reported decreased pain and improved quality of life scores following revision THR using tapered, fluted modular titanium femoral components.

The National Institute of Health consensus group have stated that satisfactory implants are those with ten-year survival figures of at least 95%. Our results suggest that CAD-CAM femoral components can be successfully used as revision prostheses. Reporting the outcome of a single component has been shown to be valid.

Despite the encouraging results achieved with these components, some important points need to be considered. The high degree of anatomical conformity of these uncemented components means that their optimal fit and ultimate success depend on highly accurate, reproducible surgical technique, which needs to be acquired prior to undertaking this type of surgery.

Cost is also an issue. Each revision CAD-CAM component costs about £1800. Optional off the shelf systems are available with adjucnts such as cables and plates. A direct comparison of costs with contemporary prostheses is difficult because of the wide variety of systems in common use, as reported by the United Kingdom National Joint Registry, as well as other adjuncts used in revision procedures, including cables, plates, bone grafts and extraction equipment.

Since the advent of the custom-made components described in this study, encouraging medium-term results have been reported with the use of extensively porous-coated components, the ZMR (Zimmer) porous-coated modular stem and fully HA-coated long-stemmed components. Many of these contemporary prostheses are versatile and can be applied to most revision situations, including those with deficient proximal bone stock where the design described in this study would be unsuitable. It is therefore difficult to propose the routine use of custom-made CAD-CAM components for revision THR in the current environment.

These minimum ten-year follow-up results are encouraging and compare favourably with revision prostheses using other means of fixation. They suggest that these custom-made implants provide a viable option in revision THR in suitable cases, particularly where the anatomy of the proximal femur makes the use of a monobloc or modular off the shelf system unsuitable.

No benefits in any form have been received or will be received from a commercial party related directly or indirectly to the subject of this article.

References

1. No authors listed. www.njcentre.org.uk (date last accessed 16 August 2010).


UNCEMENTED CAD-CAM FEMORAL COMPONENTS IN REVISION TOTAL HIP REPLACEMENT


