Long-term outcome of a cementless, hemispherical, press-fit acetabular component

SURVIVORSHIP ANALYSIS AND DOSE-RESPONSE RELATIONSHIP TO LINEAR POLYETHYLENE WEAR

Between 1988 and 1998 we implanted 318 total hip replacements (THRs) in 287 patients using the Plasmacup (B. Braun Ltd, Sheffield, United Kingdom) and a conventional metal-on-polyethylene articulation. The main indications for THR were primary or secondary osteoarthritis.

At follow-up after a mean 11.6 years (7.6 to 18.4) 17 patients had died and 20 could not be traced leaving a final series of 280 THRs in 250 patients. There were 62 revisions (22.1%) in 59 patients. A total of 43 acetabular shells (15.4%) had been revised and 13 (4.6%) had undergone exchange of the liner. The most frequent indications for revision were osteolysis and aseptic loosening, followed by polyethylene wear. The mean Kaplan-Meier survival of the Plasmacup was 91% at ten years and 58% at 14 years. Osteolysis was found around 36 (17.1%) of the 211 surviving shells. The median annual rate of linear wear in the surviving shells was 0.12 mm/year and 0.25 mm/year in those which had been revised (p < 0.001).

Polyethylene wear was a strong independent risk factor for osteolysis and aseptic loosening. The percentage of patients with osteolysis increased proportionately with each quintile of wear-rate.

There is a high late rate of failure of the Plasmacup. Patients with the combination of this prostesis and bearing should be closely monitored after ten years.

Cementless modular acetabular prostheses were introduced in the 1980s as a response to the perceived poor results of cemented, all-polyethylene, acetabular implants. The intention was to eliminate cemented fixation, which was thought at the time to be the principal cause of osteolysis, and to make the transfer of load from the bearing surface to the pelvis more even. Since then it has become apparent that polyethylene wear, rather than ‘cement disease’, is the major cause of pelvic osteolysis. McCombe and Williams have shown in a randomised, clinical trial that a conventional polyethylene liner mounted in a metal shell has approximately twice the rate of annual linear wear of a cemented all-polyethylene component and would therefore be expected to have a higher rate of osteolysis.

The Plasmacup (B Braun Ltd, Sheffield, United Kingdom) consists of a cementless, hemispherical, modular, titanium acetabular shell and a press-fit ultra-high-molecular-weight polyethylene (UHMWPE) liner. Its outer surface has a porous plasma-sprayed titanium coating 0.35 mm thick. Primary stability is achieved by interference fit, and secondary stability by ongrowth of bone. Between 1988 and 2002 this prosthesis was used at our hospital for younger patients, generally under 60 years of age, undergoing primary total hip replacement (THR). Throughout this period the polyethylene liners were made of conventional UHMWPE and sterilised by gamma irradiation in an inert gas. In 2002, our short- to medium-term studies on the use of this prosthesis showed good results, with a survival of 97.7% at five years in 128 hips and a prevalence of radiological osteolysis of 3.0%. However, over the last five years we have seen an increasing incidence of acetabular osteolysis and high rates of annual linear wear (Fig. 1). We therefore undertook a review of the surviving patients who had undergone THR using this prosthesis between 1988 and 1997 in order to establish its long-term survivorship.

Patients and Methods

Between 1988 and 1997, 318 primary THRs were performed using the Plasmacup in 287 patients by a variety of surgeons of both consultant and training grade. In 2007, patients with a minimum follow-up of ten years were invited to attend for clinical and radiological review. By this stage 20 patients...
(21 hips) had been lost to follow-up and 17 (17 hips) had died from unrelated causes, leaving a final series of 280 hips in 250 patients. There were 125 men and 125 women in the group of whom 219 (245 hips) underwent clinical and radiological review. The remaining 31 (35 hips) were unable to attend but agreed to a telephone interview. This review was approved by the Audit Department of the Sheffield Teaching Hospitals NHS Foundation Trust.

The median age of the patients at surgery was 54 years (Interquartile range (IQR) 46 to 59, minimum 19 to maximum 77). The indication for THR was primary osteoarthritis in 204 hips, developmental dysplasia in 32, fracture of the hip in 16, inflammatory arthritis in seven, previous Perthes’ disease in seven, avascular necrosis in six, as a conversion from an arthrodesis in four, previous slipped capital femoral epiphysis in three, and a neuromuscular disorder in one. All the procedures were performed in a laminar-flow theatre with the patient in the lateral decubitus position. The surgical approach was anterolateral in 243 (86.8%) hips, posterior in 28 (10.0%), and lateral with a trochanteric osteotomy in nine (3.2%). The technique for insertion of the shell included under-reaming by 2 mm and pulsed lavage of the reamed acetabulum. The median diameter of the shell was 52 mm (interquartile range (IQR) 48 to 54). The median thickness of the liner was 13 mm (minimum 8; maximum 20; IQR 12 to 15). The femoral components used were the Exeter (Stryker Ltd, Staines, United Kingdom) in 194 hips, the Charnley (DePuy Ltd, Leeds, United Kingdom) in 53, the BiContact (B Braun Ltd) in 14, the TPS (DePuy Ltd) in 12, the Stanmore (Biomet, Swindon, United Kingdom) in 5, the BiContact (B Braun Ltd) in 5, Huckstep (B Braun Ltd) in one and Kent (Biomet) in one. All the cemented femoral components were inserted after pulsed lavage of the prepared femur with saline, and using vacuum-mixed Palacos R cement containing gentamicin (Schering Plough Ltd, Welwyn Garden City, United Kingdom). This was introduced retrograde using a cement gun over a cement restrictor, and pressurised before insertion of the component. Cementless femoral component were inserted according to the manufacturers’ instructions. A conventional UHMWPE acetabular...
lar liner, gamma irradiated in an inert gas, was used in each hip. A 22 mm metal head was used in 142 hips and a 28 mm metal head in 138.

At clinical review a standardised anteroposterior radiograph of the pelvis was taken with the beam centred on the symphysis pubis and with a focus-to-film distance of 100 cm. A lateral radiograph was also taken. Linear wear of the liner was measured by a uni-radiological technique on digitised radiographs using EBRA digital software (EBRA-Cup version 2003; University of Innsbruck, Austria) according to a previously described protocol. In patients who had had their acetabular component revised, polyethylene wear was measured from the pre-revision radiograph by the same method. Loosening of the acetabular component was defined radiologically according to the criteria of Hodgkinson, Shelley and Wroblewski. Linear and expansile osteolytic lesions were recorded in the three zones described by DeLee and Charnley. Loosening of the femoral component was defined according to the criteria of Harris and McGann. Femoral osteolysis and radiolucent lines were recorded in the zones described by Gruen, McNeice and Amstutz.

Statistical analysis. All analyses were two-tailed using SPSS statistical software version 15 (SPSS Inc., Chicago, Illinois). A p-value ≤ 0.05 was deemed to be significant. The Kaplan-Meier survivorship analyses were calculated using Graphpad Prism version 5 (Graphpad Software Inc, La Jolla, California). Revision was used as the endpoint and was defined as "an operation that involved the removal and/or replacement of one or more components of a joint replacement". Continuous data were analysed using the Mann-Whitney U test and categorical data were analysed using the chi-squared test. Acetabular revision was further subdivided into ‘major’ involving removal and/or replacement of the acetabular shell and ‘minor’ which did not involve the removal and/or replacement of the shell, such as an exchange of liner. Risk factors for revision of the Plasmacup were analysed using the Cox proportional hazards model.

Results
Survivorship of the prosthesis. The revision-free survival for any indication was 90.2% (95% confidence interval (CI) 86.1 to 93.2) at ten years, but only 56.3% (95% CI 43.2 to 67.6) at 14 years after operation (Fig. 1). The ten-year survival of the Plasmacup was 91.0% (95% CI 86.9 to 93.8), but fell rapidly thereafter to 57.8% (95% CI 44.4 to 69.0) at 14 years. When minor revisions (liner exchange) were excluded, the survival of the Plasmacup shell at ten and 14 years was 93.5% (95% CI 89.8 to 95.8) and 71.1% (95% CI 59.2 to 80.0), respectively. The survival of the femoral component was 95.6% (95% CI 92.4 to 97.5) and 80.2% (95% CI 72.1 to 92.8) at ten and 14 years, respectively.

We have revised 62 hips (22.1%) in 59 patients (23.6%). Of these, 43 (15.4%) required revision of the acetabular shell. In 24 of these 43 hips (55.8%) only the shell was revised, in 18 (41.8%) the femoral component and acetabular shell were revised, and in one hip (2.3%) a liner exchange was followed by an exchange of both the shell and the femoral component. In 13 hips (21.0%) only the liner was revised. Three hips had revision of both the liner and the femoral component, and in three only the femoral component was revised.

The indication for revision of the acetabular shell was osteolysis in 28 hips, infection in five, aseptic loosening in four, excessive wear with shell damage but without radiological osteolysis in four, and recurrent dislocation in two. The indication for liner exchange was wear in 13 hips and dislocation in four. The indication for revision of the femoral component was osteolysis in ten hips, aseptic loosening in five, fracture of the component secondary to osteolysis in three, and wear damage of the femoral head in a well-fixed monobloc component in two.

Radiological complications in the surviving prostheses. In the 245 hips which had undergone radiological review, there were 211 surviving primary acetabular shells and 241 femoral components. Osteolysis was present around 36 (17.1%) of the surviving shells. This was expansile in 28, involving a single DeLee and Charnley zone in 17 hips and multiple zones in 11, and in eight the pattern of osteolysis was linear involving a single DeLee and Charnley zone. Osteolysis was present around 37 (15.0%) of the surviving femoral components. In 14 hips this was expansile, involving a single Gruen zone in four hips and multiple zones in ten, and in 23 the pattern of osteolysis was linear involving a single Gruen zone in 12 and multiple zones in 11. One further femoral component was loose and migrated.

Polyethylene wear and other risk factors for revision. It was possible to measure the rate of polyethylene wear in 237 (84.6%) hips. In the remainder, radiographs suitable for EBRA analysis were unavailable. The median rate of annual linear wear in those revised for any indication was 0.25 mm/year (IQR 0.18 to 0.32) compared with 0.12 mm/year (IQR 0.08 to 0.18) in those which remained in situ (Mann-Whitney U test, p < 0.001). The median rate of annual linear wear in surviving Plasmacups with osteolysis was 0.16 mm/year (IQR 0.11 to 0.23) compared with 0.12 mm/year (IQR 0.07 to 0.17) for those without signs of osteolysis (Mann-Whitney U test, p = 0.009). The median rate of wear in patients who had a 22 mm diameter femoral head was 0.15 mm/year (IQR 0.08 to 0.20) and 0.15 mm/year (IQR 0.09 to 0.22) in those with a head of 28 mm diameter (p = 0.477).

The rate of polyethylene wear was an independent risk factor for revision of the acetabular shell (Cox proportional hazards model, B = 3.89, SEM 0.77, p < 0.001). The age at surgery, gender, diagnosis, surgical approach, diameter of the shell, thickness of the liner, size and material of the femoral head (22 mm vs 28 mm; stainless steel/cobalt-chrome), and the design of the femoral component were not risk factors for survival of the acetabular shell (Cox univariate analyses, all comparisons p > 0.05). These covariates were also not significant predictors of the rate of polyethylene wear (multiple linear regression, all comparisons p > 0.05).
The body mass index (BMI) and level of activity, as measured by the Oxford hip score,\(^1\) were assessed for patients who attended for clinical review with a surviving prosthesis. The mean BMI in this group was 27.8 (95% CI 27.0 to 28.7) and the mean Oxford hip score 22.1 (20.4 to 23.9). Neither of these variables was a significant predictor of polyethylene wear in this group of survivors (linear regression p > 0.05).

The patients were divided into quintiles (q) based on the rate of annual linear wear, and the odds ratio was calculated using a method described previously.\(^4\) The lower and upper boundaries for each quintile (units = mm/yr) were 0.01 to 0.08 (q1), 0.08 to 0.12 (q2), 0.12 to 0.17 (q3), 0.17 to 0.24 (q4), and 0.25 to 0.74 (q5). The proportion of patients who had revision for osteolysis, loosening or radiological evidence of osteolysis increased with each quintile from 12.5% for quintile 1 to 52.2% for quintile 5 (chi-squared test, p < 0.001; Fig. 2a). The odds of osteolysis, loosening, or radiological evidence of osteolysis also increased with each quintile (Fig. 2b). Modelling of the effect of each quintile was also examined using survivorship analysis by the Cox proportional hazards model. This also showed a decline in osteolysis-free survival with increasing quintile (Fig. 3). This analysis was performed including age at surgery, gender, diagnosis, material and diameter of the femoral head, and thickness of the polyethylene liner as analysis covariates, and the resultant survivorship curves for each quintile are presented after adjustment for any potential confounding effects of these other variables.

**Discussion**

We previously reported a favourable early outcome with this prosthesis in 128 THRs from the current series.\(^1\) The survival rate was 98% at 59 months, with few radiological signs of osteolysis. These results were similar to those of Badhe, Quinnell and Howard,\(^1\) who showed a survival of the Plasmacup of 97% at 82 months in 153 hips. These findings were also similar to the mid-term results of studies on many other cementless, modular, acetabular shells using conventional polyethylene liners, including the Duraloc 300,\(^1\) the Anatomic,\(^1\) and the Harris-Galante I and II.\(^1\) However, the survival of this type of bearing couple rapidly
declined in our series after ten years. This was consistent with the findings of the Scandinavian joint registries, and suggested that the reporting of 'mid-term' survivorship is a poor guide to the long-term survival of this type of bearing couple. For example, according to the 2007 report of the Swedish hip register, the ten-year survival of components implants in patients between the ages of 50 and 59 years at surgery was 90%, but by 16 years had fallen to 78% in men and 68% in women. Similarly, the Finnish hip register showed that the ten-year survival of this type of component and bearing in patients aged between 55 and 64 years at surgery was 95% (95% CI 94.0 to 96.0) at ten years, but 81% (95% CI 77.0 to 85.0) at 15 years.  

Osteolysis and aseptic loosening are thought to result from an inflammatory osteolytic response to wear debris from the implants. The rate of wear and osteolysis are closely associated. The odds of revision or osteolysis in our series had a wear rate of 0.12 mm/year or less. Furthermore, this dose-response relationship resisted the potential confounding effects of age at surgery, gender, diagnosis, the size and material of the femoral head and the thickness of the polyethylene liner which were used as analysis covariates in the Cox regression model.

The wear rates which we observed in our series were similar to those previously reported for conventional polyethylene liners articulating with a metal head. McCombe and Williams, 1 in a randomised trial of 162 patients, found a mean rate of wear over six years of 0.15 mm/year. Previous studies which have reported the annual wear of modular polyethylene liners have reported the annual wear of 0.15 mm/year. These are approximately twice as great as those reported in cemented monobloc polyethylene acetabular components articulating against metal heads, which have wear rates in the range 0.05 to 0.11 mm/year.  2,20,28  

Although osteolysis and revision of the acetabular shell were strongly associated with the rate of wear of the bearing surface, we found no association with other factors such as age at surgery, gender, indication for primary surgery, surgical approach, diameter of the shell, thickness of the liner, size and material of the femoral head or the design of the femoral component. Our finding that the size of the femoral head (22 mm/28 mm) had no effect on the rate of osteolysis contradicts that of Livermore et al, 28 but agrees with that of Hallan, Lie and Havelin. 29 The likely reason for this lack of association is that, in our series, only 1% of liners were ≤ 8 mm thick and the median thickness was 13 mm. Thus, our data did not suffer from the potential confounding effect of the use of liners of inadequate thickness in which the bulk mechanical properties of the material rather than the surface wear rate become the limiting factors.

Our data support the findings of others that conventional polyethylene liners in cementless shells have a high rate of wear and poor long-term survivorship. They also confirm a dose-response relationship between polyethylene wear and osteolysis which does not support the concept of a ‘safe’ threshold for the wear rate. Furthermore, the rapid decline in the survivorship of the prosthesis after ten years suggests limitations in the ten-year benchmarking classification system used by the National Institute of Clinical Excellence (NICE) in the United Kingdom.  

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


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