No superior performance of hydroxyapatite-coated acetabular cups over porous-coated cups
An 8-year follow-up of 81 patients from a previously reported controlled trial

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Background and purpose Hydroxyapatite (HA) coating is believed to improve bone-implant ingrowth and long-term survival of prostheses. Recent studies, however, have challenged this view. Furthermore, HA particles may produce third-body wear and initiate aseptic loosening of implants. We report the performance of HA- and porous-coated acetabular cups in a prospective randomized trial.

Methods This was an 8-year follow-up study of our previously published prospective randomized study to compare clinical outcomes, survival, periprosthetic bone mineral density, migration, and wear rates of HA- and porous-coated acetabular cups. Dual X-ray absorptiometry (DXA) and Ein Bild Roentgen Analyse (EBRA) measurements were used. 100 patients who underwent unilateral cementless total hip arthroplasty were randomized to either porous-coated cups or HA-coated cups. Patients were examined preoperatively and at 3, 6, and 9 months, and also 1, 3, and 8 years after surgery. 81 patients were available for 8-year follow-up, 40 with porous-coated cups and 41 with HA-coated cups.

Results Age, sex, bone mineral density, and clinical results (Harris hip score) were similar in the 2 groups. The survival, wear, and migration patterns of the cups were also similar in both groups. The results of periprosthetic bone mineral density scans in region of interest 2 was in favor of the porous-coated cups, but there were no differences between the 2 groups in all the remaining regions of interest.

Interpretation HA coating had no statistically significant effect on clinical results, survival, wear, or migration at the 8-year follow-up.

Loosening and polyethylene (PE) limit the longevity of joint replacements. One of the methods to overcome loosening is osteoconductive coatings of hydroxyapatite (HA), which is believed to enhance component fixation by providing initial stability and by improving bone ingrowth (Soballe et al. 1990, Furlong and Osborn. 1991, Soballe et al. 1992, Rahbek et al. 2005). The increasing use of HA-coated components has given rise to controversy regarding HA debris-induced osteolysis and PE wear (Morscher et al. 1998). Recently, a retrospective study on Mallory-Head cups confirmed that PE wear is greater in HA-coated cups, but without this having a negative influence on revision rate (Gottliebsen et al. 2012). The finding of increased wear in HA-coated cups partially confirms the mechanism of failure for these cups: third-body wear, osteolysis, and loosening. A previous Danish Hip Arthroplasty Register study compared HA-coated implants with non-HA-coated implants (Paulsen et al. 2007) and showed that HA coating did not reduce the overall risk of revision. The same conclusions were reached from a study in the Swedish Hip Arthroplasty Register regarding uncemented femoral stems (Lazarinis et al. 2011). On the other hand, another Swedish registry study found inferior results for acetabular cups coated with hydroxyapatite. The authors questioned the routine use of HA-coated cups in primary total hip arthroplasty (THA) and they were able to show an even higher risk of loosening of HA-coated cups that led to revisions (Lazarinis et al. 2010).

In our previously reported randomized study comparing HA- and porous-coated acetabular cups (Laursen et al. 2007), we found similar outcome, survival, and changes in bone mineral density (BMD) around cups at 3-year follow-up. Here we present data from the 8-year follow-up, adding results on migration and on analysis of PE wear.

Patients and methods

166 consecutive patients were invited to participate in the trial; 60 patients refused. 6 others were excluded because screws had been used for cup fixation. 100 patients who underwent unilateral cementless THA were enrolled in a controlled ran-
domized study from October 1998 to May 2000. Inclusion criteria were primary hip osteoarthrosis, no previous hip surgery in the hip planned to be operated, and no competing medical conditions (neurological disorders or metabolic diseases) that could influence postoperative medication, hindering bone formation or an impending standard mobilization program. Patients were scheduled for unilateral cementless arthroplasty with press-fit fixated cups, and signed an informed consent document. For details of the surgery, the randomization procedure, and the cups used, see Laursen et al. (2007).

The patients were examined preoperatively, at 3, 6, and 9 months, and also 1, 3, and 8 years after surgery. Data regarding clinical results, periprosthetic bone mineral density (BMD) measurements, complications during the first 3-year follow-up, were given in our previous paper (Laursen et al. 2007). At the latest follow-up, the following data were collected: (1) Demographics, complications, prosthesis survival, and clinical results assessed according to the Harris hip score (HHS); (2) Dual X-ray absorptiometry (DXA) for measurement of BMD around the THA cups. Scans of the acetabular region were performed using the same Norland X-ray bone densitometer (Norland Corporation) as in the original study, but with upgraded software (Illuminatus DXA 4.2.0), allowing examination of 4 regions of interest (ROIs) according to Wilkinson et al. (2001) (Figure 1); (3) Cup migration and polyethylene liner wear using Ein Bild Roentgen Analyse (EBRA) software. The EBRA measurement system is based on measurement of 2D migrations from digitalized plain radiographs using a grid of transverse and longitudinal tangents on prominent landmarks of the pelvic contour. Based on the measurement error of the EBRA software, migration values less than 1 mm were not considered significant (Ichmann et al. 1995, Wilkinson et al. 2002).

We used records from the Danish Hip Arthroplasty Registry and The Danish National Patient Registry to obtain information on patients who were lost to follow-up.

**Statistics**

The null hypotheses tested were: (1) There is no difference between the 2 treatment groups concerning migration of the cups, as measured by EBRA; and (2) There is no difference between the 2 treatment groups in any of the 4 ROIs concerning BMD change during the evaluation period. In both cases, a significance level of p < 0.05 was selected.

Sample size was chosen to find differences exceeding BMD of 0.25 g/cm² (least relevant difference: δ = 0.25; SD 0.4), with a risk of type-1 error: $\alpha = 0.05$ (5%); $\beta = 0.2$ (20%); statistical power (risk of type-2 error) = $1 - \beta = 80\%$. The statistical table sample size had 43 patients in each group. In our previous study, 50 patients were chosen in each group to allow for dropouts. Differences between groups were evaluated by calculating the 95% confidence limits of the changes in BMD.

Survival data were calculated as Kaplan-Meier survival estimates, and p-value was calculated from a log-rank test using R statistical software. BMD and EBRA results were calculated as repeated-measures ANOVA, and displayed as mean curves with 1.96 SD error bars. If the mean fell outside the error bar of the other curve, there was a statistically significant difference.

**Ethics**

The Ethics Committee of North Denmark Region approved the study (N-20080005). It was performed in compliance with the Helsinki Declaration. All patients signed an informed consent document. The protocol was registered at ClinicalTrials.gov with the identifier NCT00159497.

**Results**

The patients who were invited and accepted to attend a follow-up had a mean follow-up period of 8.8 (7.9–9.6) years. 4 patients (2 from each group) refused participation in the 8-year follow-up; 13 had died of unrelated causes before the latest follow-up (5 in the standard (porous-coated) group and 7 in the study (HA/TCP-coated) group). Records from Danish national registries revealed that 1 of these patients had received an HA-coated cup and had undergone revision at another hospital due to aseptic loosening of the femoral stem. The patient was excluded from our study population because of the revision, but was included in the survival analysis. In 2 patients, both with non-HA-coated cups, revision surgery had been performed (due to aseptic loosening of the femoral stem in one case and recurrent luxation in other), leaving a study population of 81 patients (Figure 2). Similar numbers of patients were included in the 2 groups, and age, sex, BMI, and clinical results by HHS were similar (Table). Also, Kaplan-Meyer survival analysis showed a similar distribution of patients in the 2 groups (Figure 3).

Changes in periprosthetic BMD were measured in 4 ROIs (Figure 4). There was a decrease in periprosthetic bone density during the first postoperative years in all ROIs except ROI 4. In ROI 1, neither group reached early postoperative BMD levels (measured within 1 week after surgery), whereas in ROI...
4, BMD levels rose during all 8 years of follow-up. We found a small but statistically significant difference in BMD in ROI 2 in favor of the standard (non-HA) cups. Results of peri-prosthetic BMD in ROIs 1, 3, and 4 revealed no differences between the 2 groups after 8 years.

80 patients had radiographs that were adequate for the EBRA analysis. We measured migration of cups and wear rate (Figure 5). Results were calculated as repeated-measures ANOVA, and we found greater migration and wear rates in the standard porous-coated group, but the differences between groups were not statistically significant.

Discussion

In the late 1980s, de Groot et al. (1987) described a technique for direct covering of prosthetic surfaces with hydroxyapatite. This triggered a plethora of studies seeking an explanation of processes in the bone-coating/prostheses interface and looking for data which would support a superior performance for HA-coated components (Furlong and Osborn. 1991, Hardy et al. 1991, Soballe 1993, Overgaard et al. 1998, Rahbek et al. 2005). As the use of HA-coated implants increased, hydroxyapatite granules were suspected of becoming disintegrated from implants and migrating into the joint space, producing third-body wear, osteolysis, and loosening of components (Morscher et al. 1998, Stilling et al. 2009). Despite the fact that there have been many studies, definitive conclusions regarding the performance of HA-coated implants have not been reached because coating properties, implant roughness, and overall design all influence study results. Superior performance in long-term clinical or registry studies should be the most reliable predictor of the benefits of coating (Dumbleton and Manley. 2004). But register and long-term clinical studies also reveal discrepancies regarding the performance of HA-coated hip prostheses. Gottliebsen et al. (2012) reported superior survival of HA-coated Mallory-Head cups in their 11-year follow-up clinical study, but a higher annual PE wear rate in the HA group than in the non-HA group. This was a retrospective non-randomized study and the mean age in the HA group was 6 years younger than that in the non-HA group, indicating that a higher activity level could contribute to the PE wear. Gottliebsen’s findings certainly highlight the risk of HA particles from the implant surface entering the bearing and causing excessive wear, as also shown by Stilling et al. (2009) at 15-year follow-up in an RCT where the HA-coated cups had a higher revision rate than in the Ti-group (57% vs. 17%). An increased risk of need for revision of HA-coated acetabular cups was noted in a Swedish Hip Arthroplasty Register study (Lazarinis et al. 2010), but this could not be confirmed by data based on the Danish Hip Arthroplasty Register (Paulsen et al. 2007).

To come to conclusions based on reliable data, an impeccable study design and long-term follow-up are crucial. On this basis, the present study is of interest concerning the effectiveness of HA coating of acetabular components. The study has several limitations, however. It was statistically underpowered, the findings apply to one specific prosthesis, and there was

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**Table 1: Demographic data and Harris hip score (HHS)**

|                  | Porous-coated Trilogy cup | HA/TCP-coated Trilogy cup |
|------------------|---------------------------|---------------------------|
| **n**            | 41                        | 40                        |
| **Mean age (range) at 8-year follow-up** | 68 (47–80) | 67 (41–81) |
| **Gender, F/M**  | 17/24                     | 16/24                     |
| **Mean BMI (SD)**| 30 (5)                    | 29 (5)                    |
| **Mean HHS (SD) before arthroplasty** | 48 (11) | 53 (11) |
| **Mean HHS (SD) at 8-year follow-up** | 95 (10) | 96 (9) |

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**Figure 2. Flow chart of the patients.**

- 106 patients agreed to participate
- Surgery and randomisation
- Standard, porous-coated cup: 53 patients
  - 3 received acetabular screws
  - 2 refused participation
  - 5 died
  - 2 were revised
- Study, HA/TCP-coated cup: 53 patients
  - 3 received acetabular screws
  - 2 refused participation
  - 7 died
  - 1 was revised
- 8-year follow-up: 41 patients
- 8-year follow-up: 40 patients

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**Figure 3. Kaplan-Meier survival estimates with revision as end-point.**

“Standard” refers to porous-coated cups and “Study” refers to HA/TCP-coated cups. The colored areas represent 95% confidence intervals. Patients who died before the last follow-up appear as censored observations and are marked with crosses. The p-value, calculated from a log-rank test, was 0.5.
a lack of patient-reported outcome measures. The strengths of the study were the similar-sized patient groups, blinded prospective randomization, identical prostheses, and 8-year follow-up. Our previous publication (Laursen et al. 2007) provided data obtained after a 3-year follow-up. These results showed similar BMD changes and clinical outcome between HA and non-HA (standard porous-coated) cups. We assumed that HA coating would show benefit after a longer follow-up. At the 8-year follow-up, both patient groups gave affirmative results with similar clinical outcome. Overall changes in peri-prosthetic BMD had no statistically significant differences and had no negative influence on clinical outcome. All our patients showed major loss of BMD in ROI 1, and the level of BMD afterwards did not reach the preoperative level in either group.

Figure 4. BMD changes in 4 regions of interest (ROIs). Measurements in ROI 2 were significantly favorable for the porous-coated cup.

This is surprising, because ROI 1 represents the most loaded pelvic area, and the body’s response to the load should induce most significant bone strengthening. Schmidt et al. (2012) published a prospective 7-year follow-up, computed tomography-assisted study on cortical and cancellous BMD loss after press-fit HA-coated cup fixation. The authors observed a loss of BMD in the cranial periacetabular area, comparable to our ROI 1, during the follow-up period. Their patients showed no adverse effects on outcome due to BMD loss.

A Swedish registry study (Lazarinis et al. 2010) showed increased revision rates in patients with HA-coated cups. Our
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