Design-related risk factors for revision of primary cemented stems
Analysis of 3 common stems in the Swedish Hip Arthroplasty Register

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Submitted 09-09-01. Accepted 10-03-23

Background and purpose Even small design variables of the femoral stem may influence the outcome of a hip arthroplasty. We investigated whether design-related factors play any role in the risk of non-aseptic revision of the 3 most frequently used primary cemented stem designs in the Swedish Hip Arthroplasty Register.

Patients and methods We studied 71,184 primary cemented femoral stem implants (21,008 Exeter polished stems, 43,036 Lubinus SPII stems, and 7,140 Spectron EF Primary stems) that were inserted from 1999 through 2006. Design-specific characteristics were analyzed using separate Cox regression models that were adjusted for sex, age, diagnosis, incision, and number of operations (first vs. second).

Results The crude revision rate varied between 0.8% (Lubinus SPII) and 1.4% (Spectron Primary). For the Exeter stem, the smallest femoral head diameter (22 mm) was associated with a higher risk of revision. No other design-specific parameters influenced the risk of revision of the Exeter stem. The smallest Lubinus stem size, a stem with extended neck length combined with a femoral head with increasing neck length, or the use of a cobalt-chromium head had a negative influence on the outcome. For the Spectron stem, the risk of revision was elevated for the smallest stem and for increasing offset calculated as the combined effect of high offset design and increasing neck length.

Interpretation Overall revision rates were low, but for two of the stems studied design factors such as size and neck length or offset influenced the risk of non-aseptic revision.

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Materials and methods From 1999 through 2006, 100,786 femoral implants were inserted. Of these, 1,702 underwent a first-time aseptic revision during the same period. We used only revision of the femoral component with or without any simultaneous revision of the acetabular component as endpoint.

The 3 stem designs studied were Exeter polished (n = 22,577), Lubinus SPII (n = 44,904), and Spectron EF Primary (n = 7,361). Specially designed implants (e.g. some dysplasia...
or extra-long stems) were excluded, as were femoral components used in small numbers (n < 50). All THR revised due to infection, those performed because of tumors, and cases with missing data were excluded. These exclusions left 71,184 cemented femoral stem implants (21,008 Exeter polished stems, 43,036 Lubinus SPII stems, and 7,140 Spectron EF Primary stems).

The article number of each component was recoded into classified variables. Stem size, neck length, and offset were recorded for all designs. An offset variable including the presence of offset version and neck length was constructed (Lubinus SP2, Spectron EF Primary). Head material could be studied in 2 designs (Exeter, Lubinus SP2) and femoral head size (Exeter), type of taper (Exeter), and CCD angle (Lubinus SP2) in one design each (Tables 1–3).

### Femoral stem components

**Exeter polished.** The highly polished tapered collarless Exeter prosthesis (Stryker/Howmedica/Osteonics, Allendale, NJ) is made of low-corrosion stainless steel (Orthinox) and is equipped with a hollow centralizer that is made of pre-polymerized acrylic cement and applied to the tip of the stem to allow subsidence into the centralizer. Its Ra is about 0.1–0.3 mm. The polished Exeter stem has so far been analyzed as a uniform design in the Swedish Hip Arthroplasty Register. However, according to the product lists from the manufacturer, the stem designs used in Sweden between 1999 and 2006 consisted of 2 slightly different prosthesis systems, the older Exeter and the newer Exeter V40. We investigated these designs both individually and together as one group.

The shape of the part of the stem in contact with the cement is equal for both stem designs, but the V40 design has a spigot at the head-neck junction. The stem is available in 7 sizes, with 6 increasing offsets. Different neck lengths with the modular heads provide further offset options.

**Lubinus SPIII.** The Lubinus SPIII stem prosthesis (Lubinus eccentric; Waldemar Link, Hamburg, Germany) without a centralizer is made of cobalt-chromium alloy, is double curved with anterior and posterior ridges, and has an anatomic shape and a matte surface (Ra = 1.5 µm). The stem is available in 7 sizes, with 4 increasing offsets. Different neck lengths with the modular heads provide further offset options.

**Spectron EF Primary.** The Spectron EF Primary stem prosthesis (Stryker) is made of cobalt-chromium alloy, is double curved with anterior and posterior ridges, and has an anatomic shape and a matte surface (Ra = 1.5 µm). The stem is available in 7 sizes, with 4 increasing offsets. Different neck lengths with the modular heads provide further offset options.
increasing sizes, each with a standard or extended neck length, and with 3 neck angles (117°, 126°, and 135°). The analysis was restricted to the most commonly used stem length (150 mm).

Spectron EF Primary. The straight-stem Spectron EF Primary prosthesis (Spectron Primary; Smith and Nephew, Memphis, TN) is made of cobalt-chromium alloy (CoCr). The proximal one-third of the stem is grit-blasted with a surface roughness (Ra) of 2.8 µm and the distal part is smoother with Ra = 0.7 µm. The stem has a centralizer and is available in 5 sizes with increasing length and thickness with normal or high offset. All sizes have the same neck angle (131°). In the high-offset design, the neck is displaced medially and the neck becomes longer for one and the same head taper.

Follow-up
The average follow-up period for all the femoral stems analyzed was 3.5 (SD 2.2) years: 3.3 (SD 2.1) years for the Exeter, 3.3 (SD 2.2) years for the Lubinus SPII, and 3.6 (SD 2.2) years for the Spectron EF Primary.

Statistics
SPSS version 16.0 for Windows was used. Design-specific characteristics were analyzed using separate Cox regression models that were adjusted for sex, age, diagnosis, incision, and number of operation (first or second). The operation was classified as the second operation when the patient had been operated earlier with a total hip arthroplasty on the contralateral side. The assumption of proportional hazards was verified by hazard function plots that showed a reasonable proportionality over time. For the time intervals chosen, no crossing or clearly deviating survival of the design parameters was observed. In the Exeter group, the V40 taper had only been in use during the later part of the period (mean follow-up: 2.4 years; mean follow-up for old taper: 5.4 years). In addition, 22-mm heads for the Exeter stem were mainly used during the later part of the observation period. We therefore decided to restrict the Cox regression analysis of the Exeter stem and only include aseptic revisions up to 3 years. In the Lubinus SPII and Spectron Primary groups, most stems were inserted with 28-mm heads. Thus, implants with other head sizes were excluded. All design parameters were classified using the group with highest number of observations as reference (Tables 1–3). Adjusted risk ratios, 95% confidence intervals (CIs), and p-values are reported. A p-value < 0.05 was considered to be significant.

Results
The Exeter stem (Table 1)
Although the Exeter stem is available in 7 sizes with 6 increasing offsets, the largest stem sizes and offsets were used in insufficient numbers (n < 50) and we therefore analyzed them together as one class for the 2 largest stem sizes and as one class for the 2 largest offsets, resulting in 4 classes of offset and 5 classes of size. For the older version of the Exeter stem and the newer Exeter V40 stem, the average follow-up was 5.4 (SD 1.9) years and 2.4 (SD 1.5) years, respectively. When the stem design factor (“old” or V40) was disregarded, the risk of revision was only higher for 22-mm femoral heads compared to 28-mm heads. None of the other implant-related variables influenced the risk of stem revision (Table 4). However, the “old” Exeter type was revised most frequently because of aseptic loosening and the V40 type was revised most frequently due to dislocation (Table 5).

The Lubinus SPII stem (Table 2)
The largest 2 stem sizes were used in less than 50 cases and they were therefore analyzed together as one class, resulting in 6 classes of stem size. Only head diameters of 28 mm were included because too few other head sizes were used.

Table 4. Relative risks of aseptic stem revision during entire observation period (Lubinus SP, Spectron EF Primary) or within 3 years (Exeter) and 95% CI for implant-related parameters for each stem type. Data from Cox regression adjusted for age, sex, diagnosis, bilaterality, side, and implant-specific parameters (see text). Implant-related parameters with statistically significant influence are shown
Only the smallest stem size (extra-small) was associated with an increased risk of stem revision compared to all other sizes (Table 4). The risk of revision in the short term was almost half for ceramic heads when compared to the group with cobalt-chromium heads (RR = 0.52; p = 0.001) (Table 4).

The reasons for revision of both the ceramic and the cobalt-chromium heads are shown in Table 6. The risk of aseptic revision was reduced in cases with a constructed offset variable corresponding to a short neck length (RR = 0.73; p = 0.3) and increased with use of the longest neck length (RR = 1.72; p = 0.02) (Table 4).

The Spectron implant (Table 3)

Only cobalt-chromium heads with a head diameter of 28 mm were included because the heads made of other materials and with other diameters were used in insufficient numbers (n < 50). The risk of revision was higher for the smallest stem (RR = 2.2; p = 0.006) (Table 4). The risk of aseptic revision was almost 7 times higher in the group with the longest assembled neck lengths based on stem offset and neck length (RR = 6.9; p < 0.001) (Table 4).

Table 5. Revisions for all the femoral stems analyzed (infections excluded)

| Primary                  | Exeter Polished (All) | Exeter (old) | Exeter V40 | Lubinus SPII | Spectron EF |
|--------------------------|-----------------------|--------------|------------|--------------|-------------|
|                          | % (n)                 | % (n)        | % (n)      | % (n)        | % (n)       |
| Aseptic loosening        | 0.4 (96)              | 1.1 (71)     | 0.2 (25)   | 0.3 (114)    | 0.8 (57)    |
| Dislocation              | 0.5 (116)             | 0.6 (39)     | 0.5 (77)   | 0.5 (244)    | 0.5 (35)    |
| Fracture                 | 0.2 (43)              | 0.3 (18)     | 0.2 (25)   | 0.0 (17)     | 0.0 (3)     |
| Implant failure          | 0.0 (3)               | 0.0 (0)      | 0.0 (3)    | 0.0 (8)      | 0.0 (0)     |
| Technical reason         | 0.1 (12)              | 0.1 (7)      | 0.0 (5)    | 0.0 (12)     | 0.1 (4)     |
| Pain only                | 0.0 (3)               | 0.0 (0)      | 0.0 (3)    | 0.0 (5)      | 0.0 (2)     |
| Total                    | 1.2 (273)             | 2.1 (135)    | 1.1 (138)  | 0.8 (400)    | 1.4 (101)   |

Overall, the early revision rate for the 3 most frequently used cemented THRs in Sweden is low (Table 5). Thus, extensive material is required for reliable analysis. The advantage of this study is that it is representative of a wide spectrum of orthopedic surgeons with variable clinical experience, and that it covers a whole nation with complete coverage of all hospitals and very few dropouts (about 1–2% of individual operations annually) (Kärrholm et al. 2008).

We have already performed an analysis similar to the present one (Kärrholm and Herberts 2006). At that time, the follow-up was shorter and all types of aseptic revisions were included—also isolated revisions of the cup. The present analysis is based only on stem revision as the primary outcome variable.

One problem with our analysis is the short follow-up, which is related to the start of recording of article numbers in 1999. The registry is continuously updated with modifications of implants, and we can therefore directly relate any change in the survival of an implant to any modification of that implant design. However, changes in survival in the short term must be interpreted with care, especially as the increased risk of revision turned out to be due to recurrent dislocation, as dislocation is often an early complication (Woolson and Rahimtoola 1999, Sanchez-Sotelo and Berry 2001) and might have nothing to do with the specific modification of the implant design. Sometimes a longer follow-up time is necessary in order to identify substantial changes in survival outcome.

In addition, the definition of failure in the registry—revision with exchange or extraction of at least one part of the THR—is a problem and a limitation in this study. The cohort of unrevised patients with problems is probably as large as the revised cohort (Söderman et al. 2001). Also, any “wait-and-see” approach results in underestimation of the number of revisions because those patients are not registered in the database. Even so, we believe that essential information is obtained from the assessment of implant survival with stem revision as endpoint, because this outcome can be regarded as an underestimation rather than overestimation of the problem.

It has been shown that restoration of the normal anatomical offset during THR is theoretically of advantage (Davey et al. 2009).
1993, McGrory et al. 1995). Unfortunately, we have no information on whether or not the normal anatomy after THR was restored with any particular offset in our analysis. We found that the Spectron stem and the Lubinus stem were associated with an increased risk of revision with increased offset and the use of a long neck. Interestingly, no such tendency was observed for the Exeter design. The reason for a seemingly increased sensitivity to high offset with use of a rough stem is unclear. It may be that a high offset facilitates debonding of the stem due to increased lever arm, resulting in particle production caused by abrasive wear between the stem and the cement mantle. In both the Lubinus SP2 and the Spectron EF Primary design, the head used to obtain maximum neck length has a skirt that might cause impingement and increased risk of dislocation.

It should be emphasized that some implant-related parameters may be biased by factors that are not known to us. The choice of level for neck resection may influence implant fixation. It would definitely influence the flexibility to change the amount of offset during surgery without causing pronounced changes in leg length and stem positioning, which would also influence the true offset. It may be that hips operated with maximum neck length represent technically difficult cases, and to some extent cases where the preoperative planning has been suboptimal.

It appears that there was a difference in outcome for the older version of the Exeter compared to the newer V40 design, as an increased incidence (although not statistically significant), was seen for aseptic loosening in the older version. On the other hand, a higher incidence of revision due to dislocation was seen for the V40 design (Table 5). However, this higher incidence of aseptic loosening for the older version of the Exeter stem must be interpreted with respect to the difference in follow-up time between the two designs, as revisions due to dislocation occur early and revision caused by aseptic loosening in general peaks even later than the mean follow-up period. It would definitely influence the flexibility to change the level for neck resection may influence implant fixation. With the evolution of new and more wear-resistant articulations, there is a trend to use larger heads, which can be expected to reduce the frequency of dislocation (Kelley et al. 1998, Berry et al. 2005, Conroy et al. 2008). There is a similar trend in Sweden, but the follow-up of the few patients is short. We therefore chose to exclude these cases since no relevant comparison could be made.

We found that the smallest sizes of Spectron and Lubinus implants represent an increased risk of stem loosening. The reason for the problems associated with small, unpolished cemented stems is not clear. These stems probably offer poor resistance against torsion, which means that they debond more easily and may then be a source of abrasive wear at the stem/cement interface. A polished stem will most likely cause less abrasion. Thus, if the femoral canal is narrow, polished stems may be preferred.

The overall performance of 3 of the most used stem designs during the past 8 years has been good. In 2006, these designs constituted about 71% of all stems inserted and their 10-year survival rate based on all reasons for revision varied between 93% and 96% (Kärrholm et al. 2008). In spite of this, 2 of the designs had implant variations associated with an increased risk of mechanical failure, which could probably have been avoided if the introduction of these variations had been done using more sophisticated techniques. Our findings underscore previous experiences from other implant designs, where changes in the surface finish or the addition of a new coating did not improve the clinical results (Rockborn et al. 1993, Middleton et al. 1998, Della Valle et al. 2005).

TH and JK: planning of the study, statistical analysis, and preparation of the manuscript. TT: collection and analysis of data.

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