HIP

Vitamin E-doped total hip arthroplasty liners show similar head penetration to highly cross-linked polyethylene at five years: a multi-arm randomized controlled trial

Aims
The most frequent indication for revision surgery in total hip arthroplasty (THA) is aseptic loosening. Aseptic loosening is associated with polyethylene liner wear, and wear may be reduced by using vitamin E-doped liners. The primary objective of this study was to compare proximal femoral head penetration into the liner between a) two cross-linked polyethylene (XLPE) liners (vitamin E-doped (vE-PE)) versus standard XLPE liners, and b) two modular femoral head diameters (32 mm and 36 mm).

Methods
Patients scheduled for a THA were randomized to receive a vE-PE or XLPE liner with a 32 mm or 36 mm metal head (four intervention groups in a 2 × 2 factorial design). Head penetration and acetabular component migration were measured using radiostereometric analysis at baseline, three, 12, 24, and 60 months postoperatively. The Harris Hip Score, University of California, Los Angeles (UCLA) Activity Score, EuroQol five-dimension questionnaire (EQ-5D), and 36-Item Short-Form Health Survey questionnaire (SF-36) were assessed at baseline, three, 12, 36, and 60 months.

Results
Of 220 screened patients, 127 were included in this study. In all, 116 received the allocated intervention, and 94 had their results analyzed at five years. Head penetration was similar between liner materials and head sizes at five years, vE-PE versus XLPE was -0.084 mm (95% confidence interval (CI) -0.173 to 0.005; p = 0.064), and 32 mm versus 36 mm was -0.020 mm (95% CI -0.110 to 0.071; p = 0.671), respectively. No differences were found in acetabular component migration or in the patient-reported outcome measures.

Conclusion
No significant difference in head penetration was found at five years between vE-PE and XLPE liners, nor between 32 mm and 36 mm heads.

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Introduction
The long-term survival of total hip arthroplasties (THAs) is around 80% to 90% after 20 years depending on diagnosis and age.1-4 Implant failure is primarily caused by aseptic loosening, which is associated with polyethylene liner wear.1,5,6 A significant reduction in wear and risk of osteolysis can be achieved by using cross-linked polyethylene (XLPE) liners instead of ultra high molecular weight polyethylene (UHMWPE) liners for metal-on-polyethylene THAs.7,9 To achieve cross-linking, the polyethylene is subjected to radiation. This process produces unwanted free radicals. These are then reduced using either annealing or remelting, but a compromise must be made between the remaining free radicals and reduced wear resistance.10-12 Alternatively, the polyethylene is first cross-linked and then doped with vitamin E (α-tocopherol), an antioxidant that can scavenge free radicals, rendering annealing or remelting redundant.13 This material has shown increased resistance to oxidation and
fatigue crack propagation, as well as a low wear rate in experimental settings and has been investigated in vivo.\textsuperscript{14-20} One advantage of lower wear rates may be to allow the use of larger heads to reduce the risk of dislocation.\textsuperscript{21} Over the past decade we have seen a decreased use of 28 mm diameter modular heads and increased use of 36 mm.\textsuperscript{22} Larger heads have shown increased wear in UHMWPE liners,\textsuperscript{23} but little evidence is available on the effect of head size on vitamin E-doped highly cross-linked polyethylene (vE-PE) liner wear.\textsuperscript{24}

The clinically relevant endpoint is revision rate, but due to low overall revision rate and high longevity, this is often not a feasible outcome measure. As revision rates and polyethylene wear have been associated, polyethylene wear can be measured using RSA and PROMs.\textsuperscript{6,24}

### Methods

#### Trial design

This was a 2 × 2 factorial randomized, single-blinded, controlled trial reported in accordance with the CONSORT statement for multi-arm trials.\textsuperscript{25}

#### Participants

Patients aged 40 to 70 years who were eligible for an un cemented THA due to idiopathic osteoarthritis and could fit at least a 54 mm component were included at Odense University Hospital and Middelfart Hospital between May 2009 and April 2011. Exclusion criteria were severe anteversion of the femoral neck that required non-standard components, acetabular dysplasia (centre edge angle < 20°), malignancy, previous radiotherapy, inability to participate in the rehabilitation programme, or received screws inserted into the acetabular shell screws or femoral cerclage during surgery.

#### Intervention

The patients were allocated to one of four intervention groups: vE-PE liner (E1, Biomet, Warsaw, Indiana, USA) with a 32 mm head (vE-PE, 32 mm); vE-PE liner with a 36 mm head (vE-PE, 36 mm); XLPE liner (ArComXL, Biomet, Warsaw, Indiana, USA) with a 32 mm head, (XLPE, 32 mm); or XLPE liner with a 36 mm head (XLPE, 36 mm).

All patients received plasma-sprayed porous-coated acetal- lular shells (Exceed ABT, Biomet, Warsaw, Indiana, USA) and un cemented porous-coated components (Bi-Metric, Biomet, Warsaw, Indiana, USA) with cobalt-chromium (CoCr) alloy modular femoral heads (Biomet) according to manufacturer’s instructions using the posterior approach. Ten tantalum beads (diameter 0.8 mm) were inserted in the periacetabular bone. Only shell sizes from 54 mm and upward were used to ensure at least 5 mm liner-thickness. The patients received tranexamic acid and antibiotics during surgery. Rehabilitation, pain management, and discharge were standardized.

#### Outcomes

The primary outcome was proximal femoral head penetration into the liner measured using RSA. The secondary outcomes were proximal acetabular component migration measured using RSA and PROMs.

#### RSA

Patients were placed in a supine position above a unipla- nar calibration cage 43 (RSA Biomedical AB, Umeå, Sweden). A ceiling-mounted and a mobile radiography tube exposed two digital radiographs analyzed using UmRSA 7.0 (RSA Biomedical, Umeå, Sweden). The tantalum beads formed the

### Table I. Baseline demographics for allocated patients.

| Characteristic | Individual groups | vE-PE, 32 mm | vE-PE, 36 mm | XLPE, 32 mm | XLPE, 36 mm |
|---------------|------------------|-------------|-------------|------------|------------|
| Total, n      | 24               | 29          | 30          | 33         |
| Male sex, n (%) | 19 (79)          | 20 (69)     | 19 (63)     | 19 (58)    |
| Mean BMI, kg/m² (range) | 28 (22 to 40) | 29 (22 to 36) | 28 (20 to 41) | 27 (21 to 38) |
| ASA class, n (%) | 1               | 7 (29)      | 9 (31)      | 8 (27)     | 13 (39)    |
| 2            | 13 (54)          | 14 (48)     | 15 (50)     | 8 (24)     |
| 3            | 1 (4)            | 1 (3)       | 1 (3)       | 1 (3)      |
| Data not available | 3 (13)         | 5 (17)      | 6 (20)      | 11 (33)    |
| Median blood loss, ml (IQR) | 450 (225 to 550) | 300 (200 to 600) | 350 (300 to 400) | 350 (225 to 400) |
| Median duration of anaesthesia, mins (IQR) | 90 (75 to 105) | 80 (66 to 95) | 75 (65 to 84) | 83 (71 to 99) |
| Median days hospitalized (IQR) | 3 (3 to 3) | 3 (2 to 3) | 3 (2 to 4) | 3 (3 to 5) |
| Mean Harris Hip Score (range) | 47 (20 to 80) | 49 (18 to 81) | 50 (13 to 74) | 44 (21 to 74) |
| Median UCLA Activity Score (IQR) | 6 (4 to 7) | 6 (4 to 8) | 6 (5 to 7) | 5 (4 to 7) |
| Median EQ-5D Health State (IQR) | 0.72 (0.66 to 0.72) | 0.66 (0.59 to 0.72) | 0.72 (0.57 to 0.72) | 0.72 (0.59 to 0.72) |
| Median EQ-5D VAS (IQR) | 64 (46 to 83) | 60 (40 to 70) | 70 (50 to 79) | 75 (60 to 90) |
| Mean SF-36 PCS (range) | 37 (30 to 48) | 37 (32 to 42) | 37 (26 to 44) | 36 (27 to 45) |
| Mean SF-36 MCS (range) | 41 (34 to 53) | 41 (36 to 47) | 42 (32 to 51) | 41 (31 to 54) |

ASA, American Society of Anesthesiologists Physical Status classification system; BMI, body mass index; EQ-5D Health State, EuroQol five-dimensional three-level questionnaire; EQ-5D VAS, EuroQol visual analogue scale; IQR, interquartile range; MCS, mental component score; PCS, physical component score; SF-36, 36-Item, Health Survey questionnaire; UCLA, University of California, Los Angeles; vE-PE, vitamin E-doped cross-linked polyethylene liners; XLPE, cross-linked polyethylene liners.
periacetabular bone segment, and the centre of the acetabular component and head were formed using spherical modelling.

RSA was performed at baseline (seven days after surgery), three months, and one, two, and five years. Head penetration is reported as the total proximal head penetration and as penetration rate at each follow-up. Double examinations were performed at the five-year follow-up, and 95% confidence intervals (CIs) for repeatability in the x- (transverse), y- (vertical), and z-axes (anteroposterior) were 0.246 mm, 0.226 mm, and 0.371 mm, respectively. Analyses where rigid body fitting error > 0.350 mm or condition number > 135 were rejected.

PROMs. The Harris Hip Score is a scale from 0 to 100 and measures pain, function, deformity, and movement related to the hip. The University of California, Los Angeles (UCLA) Activity Score is a ten-point Likert-like scale measuring physical activity and is recommended and used in similar populations. The EuroQol five-dimension questionnaire (EQ-5D) is a utility index in health economics based on five descriptive domains-mobility, self-care, usual activities, pain/discomfort, and anxiety/depression-used with the Danish Time Trade-Off (TTO) value set. The 36-Item Short-Form Health Survey questionnaire (SF-36) is a generic health questionnaire in eight dimensions that can be summarized to a physical and a mental health score. PROMs were assessed at baseline (two weeks prior to surgery), three months, one, three, and five years. Function was evaluated with the HHS and UCLA score, and general health with the EQ-5D and SF-36.
Adverse events. We queried the Danish Hip Arthroplasty Register (DHR) for revision surgery on participating patients. The DHR is a national registry of surgeon-reported parameters including indication for primary and revision hip arthroplasty and has been shown to have high levels of data capture. Sample size. This study was powered as a parallel-group trial to show that vE-PE liners had lower wear than XLPE liners. Power was set to 80%, and the risk of type-I error was set to 5%. Wear was expected to drop from 0.05 mm/year in XLPE to 0.0005 mm/year in vE-PE. The minimal clinically relevant difference was set to 0.05 mm/year. This resulted in a requirement for 15 hips in each group. To account for dropouts and secondary exclusion, study recruitment was planned at 25 patients for each arm. Randomization and blinding. The allocation was computer-generated in two blocks (one block with 100 lots of 25 per group, another block with 28 lots of seven per group) in random order and lots were placed in sealed envelopes. The envelope was opened just before the liner was inserted. The study was blinded for the patients. Registration and ethics. This trial was approved by The Regional Committees on Health Research Ethics for Southern Denmark (S-20080151) and The Danish Data Protection Agency (14/35949 and 18/31287) and complied with the Declaration of Helsinki. The trial was registered at ClinicalTrials.gov (NCT02196792). Statistical analysis. A statistical analysis plan was previously published at ClinicalTrials.gov (NCT02196792). Patients allocated to an intervention group were included in the intention-to-treat (ITT) analysis, and patients who were in the ITT population and present at all RSA follow-ups were included in the per-protocol (PP) analysis. Descriptive statistics were reported using mean and SD or median and interquartile range (IQR) as appropriate. Data were analyzed using mixed-effect analysis using the restricted maximum likelihood approach. Patients were considered as random effects and time, liner material, and head size as fixed effects. Interactions between time and liner material, and time and head size were included. PROM analyses were adjusted for baseline values. Data were analyzed using
In total, 220 patients were assessed for eligibility (Figure 1), and 93 were excluded: 26 patients received an acetabular component size smaller than 54 mm, five patients received other components, four patients received screws, one was excluded from the trial (Supplementary Table ii). No between-group differences were found for any of the secondary outcomes, except for liner materials where vE-PE had significantly lower penetration (vE-PE vs XLPE was -0.117 (95% CI -0.231 to -0.002; p = 0.046)).

The between-group wear rates were similar for liner materials and head sizes at all follow-ups. The effect of liner material and head size did not interact in these analyses. Adverse events. No between-group differences were found for any of the PROMs (Table II). Within-group improvements were found for Harris Hip Score (HHS) for both groups, while only 32 mm improved in UCLA, EQ-5D Health State, EQ-5D Visual Analogue Scale (VAS), and SF-36 Physical Component Score (PCS).

**Results**

**Recruitment.** In total, 220 patients were assessed for eligibility (Figure 1), and 93 were excluded: 26 patients received an acetabular component size smaller than 54 mm, five patients received other components, four patients received screws, one was excluded due to other surgical complications, and 57 patients due to other reasons. The remaining 127 patients were randomized, and 116 patients received the allocated intervention, with a total of 11 patients being excluded post-randomization: four patients received shell screws, one received other components, one suffered a dislocation during surgery and received other components and was excluded, and five were excluded due to screening failure. A total of 22 patients were lost to follow-up, and 94 patients were still enrolled after five years.

**Head penetration.** We found no between-group difference in mean total head penetration for liner materials or head sizes at five years (Figure 2, Table II): vE-PE versus XLPE was -0.084 mm (95% CI -0.173 to 0.005; p = 0.064, mixed-effect analysis), and 32 mm versus 36 mm was -0.020 mm (95% CI -0.110 to 0.071; p = 0.671, mixed-effect analysis). The PP analysis of total head penetration (Supplementary Table i) gave similar results except for liner materials where vE-PE had significantly lower penetration (vE-PE vs XLPE was -0.117 (95% CI -0.231 to -0.002; p = 0.046)).

The between-group wear rates were similar for liner materials and head sizes at all follow-ups. The effect of liner material and head size did not interact in these analyses.
Discussion
This study compared THA head penetration into the liner between two types of liners (vE-PE and XLPE) and between two head sizes (32 mm and 36 mm). It showed that both total head penetration and wear rates were independent of liner type and femoral head size. No patients experienced aseptic loosening but given the low wear rate, small sample size, and short duration of the study, this would have been unlikely. These results do not provide compelling evidence to phase out XLPE liners in favour of vE-PE liners, or to choose 32 mm over 36 mm heads.

The well-known creep and wear phases were not found, and there are a few possible explanations for this. The repeatability of our RSA setup was slightly higher than that of similar trials. The accuracy of the model-based approach used in this study is lower than the marker-based approach, though both methods have the necessary accuracy to detect migration at the osteolysis threshold. Reduced accuracy does not affect the ability to determine true migration, but it increases variance and the necessary sample size for a given power. Lastly, the in-hospital location where RSA was performed changed during this trial. The same calibration box and software version was used for all RSA measurements, but this location change could have imposed a systematic error, although comparison of mean square error (MSE) and condition number (CN) between each follow-up does not suggest this (one-way analysis of variance (ANOVA); p = 0.095 for MSE; p = 0.758 for CN). These reasons may, however, explain the lack of distinct creep and wear phases.

Five other RCT shave compared vE-PE and XLPE liner wear. Two trials report no difference at two or seven years, respectively, while three trials report lower wear in vE-PE liners at five, five, and seven years, respectively. All these studies use the same RSA setup as this study. The disagreement between studies may arise from some of the reasons explained above, differences in sample size, or simply because the true difference in head penetration is low relative to the repeatability.

We have reported linear wear, but volumetric wear may be a more suitable metric for different head sizes. A 32 mm head with a linear wear of 0.100 mm/year has the same volumetric wear as a 36 mm head with a linear wear of 0.079 mm/year, and both head sizes in this study were well below these rates. A recent study by Lindal et al supports the finding that 32 mm heads are not superior to 36 mm in terms of wear.

We did not find a difference in PROMs between 32 mm and 36 mm heads, and this trial was not powered to detect differences in PROMs. The RSA and PROM measurements do not align, as frequent RSA is important in the beginning of follow-up.

A large number of patients were randomized. Due to decentralized and insufficiently strict trial management, excess patients were screened and scheduled for randomization, and the second randomization block was made. This was unintentional, and this cohort has similar baseline demographics to a more recent cohort in a double-blinded RCT from the same department.

The sample size calculation was based on optimistic expectations of the treatment effect, but sample size calculations are inherently subjective and similar RCTs have equal or lower sample sizes.

We conducted both an ITT and a PP analysis to mitigate the risk of bias due to differential dropouts. The ITT and PP analyses showed very similar effect sizes for head penetration. Both were in favour of vE-PE, but the PP was statistically significant while the ITT was not. With repeatability and model-based approach of the RSA setup and the significance discrepancy between ITT and PP for head penetration in mind, this trial may suffer from a Type II error. However, we wonder if the true head penetration of vE-PE and XLPE groups will diverge and become detectably different at a later follow-up.

The intervention groups only differ in terms of liner material and head size, all other parameters were similar between intervention groups, and the primary outcome was measured using a standardized protocol. Among THA patients, 80% are generally diagnosed with osteoarthritis, and we believe the patients in this trial are representative for the majority of THA patients.

In conclusion, in our investigation we identified no differences in head penetration and acetabular component migration between vE-PE and XLPE groups, or between 32 mm and 36 mm heads. In addition, we found no difference in PROMs between 32 mm and 36 mm heads.

Take home message
- Vitamin E-doped cross-linked total hip arthroplasty liners do not show lower wear rates than conventional cross-linked liners.
- Both liners show very low wear rates overall.

Supplementary material
Graphs and estimates from the per-protocol analysis as well as a summary table of dropouts.

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This trial was approved by The Regional Committees on Health Research Ethics for Southern Denmark (S-20080151) and The Danish Data Protection Agency (14/35949 and 18/31287) and complied with the Declaration of Helsinki.

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**Trial registration number**
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