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Cam morphology but neither acetabular dysplasia nor pincer morphology is associated with osteophytosis throughout the hip: findings from a cross-sectional study in UK Biobank

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Summary
Objectives: To examine whether acetabular dysplasia (AD), cam and/or pincer morphology are associated with radiographic hip osteoarthritis (rHOA) and hip pain in UK Biobank (UKB) and, if so, what distribution of osteophytes is observed.
Design: Participants from UKB with a left hip dual-energy X-ray absorptiometry (DXA) scan had alpha angle (AA), lateral centre-edge angle (LCEA) and joint space narrowing (JSN) derived automatically. Cam and pincer morphology, and AD were defined using AA and LCEA. Osteophytes were measured manually and rHOA grades were calculated from JSN and osteophyte measures. Logistic regression was used to examine the relationships between these hip morphologies and rHOA, osteophytes, JSN, and hip pain.
Results: 6,807 individuals were selected (mean age: 62.7; 3382/3425 males/females). Cam morphology was more prevalent in males than females (15.4% and 1.8% respectively). In males, cam morphology was associated with rHOA [OR 3.20 (95% CI 2.41–4.25)], JSN [1.53 (1.24–1.88)], and acetabular [1.87 (1.48–2.36)], superior [1.94 (1.45–2.57)] and inferior [4.75 (3.44–6.57)] femoral osteophytes, and hip pain [1.48 (1.05–2.09)]. Broadly similar associations were seen in females, but with weaker statistical evidence. Neither pincer morphology nor AD showed any associations with rHOA or hip pain.
Conclusions: Cam morphology was predominantly seen in males in whom it was associated with rHOA and hip pain. In males and females, cam morphology was associated with inferior femoral head osteophytes more strongly than those at the superior femoral head and acetabulum. Further studies are justified to characterise the biomechanical disturbances associated with cam morphology, underlying the observed osteophyte distribution.

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Introduction
Hip osteoarthritis (OA) is a common condition that causes considerable morbidity often leading to costly total hip replacements (THR) 1,2. Differences in hip morphology have long been postulated as risk factors, including acetabular dysplasia (AD), and cam and pincer morphologies 3. AD is associated with under-coverage of the acetabulum over the femoral head and is considered a consequence of milder forms of developmental dysplasia of the hip (DDH) 4,5. Severe DDH is strongly associated with hip OA whereas AD shows inconsistent associations 5,6. Cam morphology, which represents bulging of the lateral femoral head leading to an aspherical appearance, and pincer morphology, comprising increased coverage of the acetabulum over the femoral head, both have been suggested to cause OA via femoro-acetabular impingement (FAI). The biomechanical concept of aberrant forces due to impingement of
the superolateral femoral head on the lateral acetabulum during hip movement in particular flexion, abduction and internal rotation14,15. An individual’s hip morphology develops through gestation, childhood and adolescence well before the onset of OA3,10. Genetic loci have been associated with different hip morphologies including DDH indicating a genetic predisposition1,12. Observational studies suggest cam morphology forms in adolescence when the metaphysis fuses, with increased physical activity implicated as a risk factor1,14. FAI syndrome is recognised as a cause of hip pain in younger individuals, diagnosis of which is supported by relevant examination findings and either cam and/or pincer morphologies in the absence of OA3,12. Several studies suggest that surgery to correct the hip morphologies implicated in FAI improves symptoms such as pain14–18. Conceivably, surgery to correct these hip morphologies and prevent FAI might also prove useful in reducing the risk of developing OA. However, whether FAI is a risk factor for hip OA in the general population remains unclear. Whereas cam morphology is associated with an increased risk of radiographic hip OA (rHOA) and THR, pincer morphology does not appear to be a risk factor for hip OA20,21. FAI has been proposed to cause hip OA in patients with cam and/or pincer morphologies secondary to impingement19 but as yet the precise mechanism remains unclear. A systematic review showed labral deformities are associated with cam morphology but the authors concluded causality could not be inferred from the studies21. No population studies have explored the distribution of osteophytes in individuals with these shape morphologies, which might give some indication as to any underlying biomechanical disturbance.

In the present study, we sought to establish the importance of hip morphology as a risk factor for OA by examining whether AD, cam and/or pincer morphology are related to rHOA and/or hip pain. In particular, we aimed to determine what distributions of osteophytes, if any, are associated with these hip morphologies. We used high resolution dual-energy X-ray absorptiometry (DXA) scans of the hip (previously validated for the use of detecting rHOA25) from a sub-sample of UK Biobank (UKB), and applied a novel automated method for ascertaining hip morphology to address these questions.

Materials and methods

Population

UKB is a mixed sex cohort, based in the UK, which prospectively recruited 500,000 adults aged 40–69 years old between 2006 and 2010. The UKB Ethics Advisory Committee oversees the maintenance, development and use of UKB data and its approval covers this study. The participants underwent extensive genetic and physical assessment, development and use of UKB data and its approval covers this study. The participants underwent extensive genetic and physical

A detailed description of the DXA mark up and derivation of parameters related to rHOA is available25. In brief, a machine learning algorithm placed 85 outline points around the left femoral head and acetabulum26,27. The points were manually checked and corrected where necessary. All osteophytes were marked up using a custom tool (University of Manchester) which allows the user to shade/identify pixels where an osteophyte is visible (Fig. 1), at the lateral acetabulum, superolateral femoral head, and interomedial femoral head. Femoral head osteophytes are referred to as superior and inferior femoral head osteophytes for simplicity. Outline points were moved to the internal boundary of an osteophyte if present (Fig. 1). Osteophyte area was used to derive osteophyte grade, based on thresholds identified from receiver operating characteristic (ROC) analyses comparing osteophyte area with osteophyte grade assessed semi-quantitatively in a subset of images. Superior minimum joint space width (mJSW) in millimetres (mm) was automatically measured between lines drawn through points 78–84 on the acetabulum and points 22–31 on the femoral head (Fig. 1). From mJSW semi-quantitative joint space narrowing (JSN) was calculated by applying ROC-derived thresholds to height adjusted mJSW measures, as these were more accurate (greater area under the curve) than using mJSW alone1.3. Repeatability for the presence of osteophytes intra-reader kappa of 0.80–0.91 was obtained with repeat readings of 300 images more than 2 months after initial grading and JSN on 100 images giving a kappa of 0.93. RHOA was defined as the presence of both grade ≥1 JSN and a grade ≥1 osteophyte at any location28,29. In addition, we employed a more stringent threshold, termed rHOA grade ≥2, requiring the presence of a grade ≥2 osteophyte and grade ≥2 JSN. Subchondral sclerosis and cysts were not examined as part of this study due to their relative infrequency30. A binary hip pain variable was derived from the following question: “Have you had hip pains for more than 3 months?” The question was not side-specific and the cause of hip pain is not identified.

Alpha angle

To automatically derive alpha angle (AA), a custom Python script was developed that fits a circle of best fit using the outline points 15–28 around the femoral head31. The script calculates the angle between a line passing through the centre of the femoral head and neck, and a line passing through the centre of the femoral head and the point at which the femoral head–neck junction leaves the circle of best fit (Fig. 1). An in-depth description of these methods including validation experiments has previously been published32. Cam morphology was defined as AA ≥60°33,34. For repeatability, 100 images were reassessed more than 2 months after initial reading with the same methods. The AA from each assessment was compared giving a concordance correlation coefficient 0.84, and cam morphology comparison gave a kappa 0.81 (97% agreement).

Lateral centre-edge angle

To automatically derive the lateral centre-edge angle (LCEA), a custom Python script was developed that calculates the angle between a line passing through the lateral edge of the acetabulum (defined by outline point 78) and the centre of the femoral head (defined by the circle of best fit as described above), and a line which passes perpendicular to the image x-axis through the centre of the femoral head (Fig. 1)35. Pincer morphology was defined as a LCEA of ≥45° and AD as a LCEA <25°36,37. 100 images were reassessed for repeatability 2 months after initial reading. The LCEA from each assessment was compared giving a concordance correlation coefficient 0.98, pincer morphology comparison gave a kappa 0.94 (99% agreement), and AD gave a kappa 1 (100% agreement).
Patient and public involvement

A patient and public involvement group made up of OA patients (University of Bristol), reviewed the plans for this analysis at an early stage. They supported the overall research aim and they emphasised the importance to focus on hip pain. The results of this work will be shared with the same group as well as the wider public and patient communities via social media and our university press teams.

Statistical analysis

The demographic data are given as mean and range for continuous variables and binary variables are given as counts and frequency. Due to the clear differences in cam prevalence between the sexes, sex stratified analyses were conducted alongside combined sex models. We examined associations between hip morphologies and the presence of rHOA and its constituent features (osteophytes and JSN), using logistic regression. The results are presented as odds ratios (OR) with 95% confidence intervals (CI), comparing those having each morphology with the remainder. A sensitivity analysis was done comparing pincer morphology and AD with all rHOA based outcomes using logistic regression with a reference group including those with a LCEA ≥25° & <45° as both ends of the LCEA spectrum have been associated with rHOA (Supplementary Results). Logistic regression was also used to examine relationships between morphology and hip pain. Directed acyclic graphs informed the a priori selection of covariates for the adjusted model, namely age, height, weight and ethnicity, with sex also added to the adjusted combined sex models. Sensitivity analyses were performed with rHOA grade ≥2 as the outcome. All statistical analyses used Stata version 15 (StataCorp, College Station, TX, USA).

Results

Population characteristics

7,000 UKB participants with a left hip DXA were initially selected, 193 were excluded (due to poor image quality or removal of consent) leaving 6,807 individuals (mean age: 62.7 years) in the final analysis. The sample comprised 3425 [50.3%] females and 3382 [49.7%] males. 1489 [21.9%] participants, 581 [17.2%] males and 908 [26.5%] females, had a self-reported diagnosis of OA (no joint locations were specified in the question) and 594 [8.7%] participants, 219 [6.5%] males and 375 [11.0%] females, reported hip pain for more than 3 months.

DXA-derived hip shape characteristics

AA was greater in males [mean: 51.6° (range: 35.8–106.2)] than females [44.2° (33.2–115.0)] and cam morphology, defined as AA ≥50°, was more frequently found in males [519 (15.4%)] than
rHOA and its constituent features

Prevalent rHOA, defined as the presence of a grade ≥1 osteophyte combined with grade ≥1 JSN, was more frequent in males [245 (7.2%)] than females [108 (3.2%)] (Table I). JSN was more common in males [35.5 (7.9%)] with pincer morphology, defined as the presence of a grade ≥1 osteophyte throughout the hip: findings from a cross-sectional study in UK Biobank, Osteoarthritis and Cartilage, https://doi.org/10.1016/j.joca.2021.08.002

Cam vs rHOA and its constituent features

Cam morphology was associated with increased risk of rHOA in males [OR: 3.24 (95% CI 2.44–4.30); Table II], females [2.37 (1.07–6.94; Table III]), and males and females combined [4.08 (3.15–5.27; Supplementary Table 1)]. Similar associations were seen after adjustment for demographic covariates, namely age, height, weight, and ethnicity, with sex added to the combined sex model. In addition, cam morphology was associated with JSN in unadjusted and adjusted analyses in males [1.53 (1.25–1.88) & 1.75 (1.57–1.96) respectively] (Fig. 2 & Table I). In females, there was little evidence of association between pincer morphology and grade ≥2 inferior femoral osteophytes in females. Pincer and AD vs rHOA and its constituent features

There was little evidence of association between pincer morphology and rHOA, in males, females, or males and females combined (Table II and III, Supplementary Table 1). In contrast, pincer morphology showed strong associations with JSN in males [4.03 (3.16–5.130), females [4.03 (3.10–5.24)], and males and females combined [4.00 (3.36–4.77)], with equivalent findings after adjustment. Pincer morphology was unrelated to the presence of osteophytes. AD was unrelated to rHOA or osteophytes in males, females, or males and females combined (Table II and 3, Supplementary Table 1).
that cam morphology was associated with an increased risk of

Discussion

between pincer or AD and hip pain, in males, females, or males and

sensitivity analysis was conducted for pincer morphology and AD, comparing their associations with rHOA based outcomes with those of a reference group which included those without AD and pincer morphology, yielding similar results (Supplementary Table 4).

Morphological measures vs hip pain

Cam morphology was associated with hip pain in males, in both unadjusted and adjusted analyses [1.51 (1.08–2.12) and 1.48 (1.05–2.09) respectively] (Table IV). In further analyses, this association was partially attenuated by additional adjustment for the presence of osteophytes [adjusted OR for the presence of acetabular 1.43 (1.01–2.01), superior 1.42 (1.01–2.00), inferior 1.30 (0.91–1.85) osteophytes and all osteophytes combined 1.27 (0.89–1.81)]. In contrast, cam morphology was unrelated to hip pain in females, or males and females combined apart from in the adjusted model (Supplementary Table 5). There was no evidence of association between pincer or AD and hip pain, in males, females, or males and females combined (Table IV and Supplementary Tables 4 and 5).

Discussion

In a large cross-sectional study of 6,807 individuals, we found that cam morphology was associated with an increased risk of prevalent hip OA, as reflected by rHOA and self-reported hip pain. In contrast, neither pincer morphology nor AD were related to either rHOA or hip pain, although they were associated with a greater and lower risk of JSN respectively. To further understand the relationship between cam morphology and hip OA, we explored the relationship between cam morphology and osteophyte distribution. Cam morphology was associated most strongly with inferior femoral head osteophytes, rather than those at the superior-lateral femoral head and acetabulum. In addition, the association between cam morphology and hip pain was partially attenuated by adjusting for the presence of inferior femoral osteophytes. This suggests that a mechanism involving the inferior femoral head contributes to the relationship between cam morphology and hip pain.

This is the first study to use DXA scans to define FAI-related morphologies with AA and LCEA. Comparison between DXA-derived AA [males: mean 51.6° (range 35.8–106.2); females: 44.2° (33.2–115.0)] and LCEA [males: 35.5° (7.9–61.8); females: 35.2° (8.4–59.7)] from our study with comparative studies which used x-rays to derive AA [males: 52.6° (30–108); females: 45° (26–92)] and LCEA [males: 34.4° (8–62); females: 35.3° (6–67)] show similar population level statistics. Our findings are also consistent with previous population studies showing that cam morphology is associated with rHOA. However, in contrast to the presented results, previous large population studies found no relationship between cam and hip pain. In our study, cam morphology was predominantly a male characteristic, and although cam was associated with hip pain in males, a similar relationship was not seen in females, possibly due to a lack of power. These findings are consistent with previous work suggesting that cam is much less likely to occur in females and therefore cannot explain the majority of female hip OA or hip pain. It may be that different thresholds for cam morphology based on AA are required in males and females, to account for sex differences in hip shape but further research is needed.

Further, our findings are consistent with previous studies which found that pincer morphology is not associated with rHOA or hip pain, and provide further evidence against an important role of pincer-type FAI in the development of hip OA. Though pincer morphology was unrelated to rHOA or osteophytes, it was associated with an increased risk of JSN. This could be a true relationship, but we are cautious of this conclusion as analysis of the site of maximal JSN showed this tended to be more lateral. This might represent an artefact related to 2-dimensional imaging creating the appearance of a narrowed joint space in the presence of acetabular

| Male & Females | rHOA | Acetabular OP | Superior Femoral OP | Inferior Femoral OP | JSN |
|-------------|------|--------------|---------------------|-------------------|-----|
| **Unadjusted analysis** | | | | | |
| Cam | 3.24 | 3.47 × 10⁻¹⁶ | 1.89 | 1.04 × 10⁻⁰⁷ | 1.94 | 4.61 × 10⁻⁰⁶ | 4.77 | 1.47 × 10⁻²¹ | 1.53 | 4.88 × 10⁻⁰⁵ |
| | [2.44–4.30] | | [1.50–2.39] | | [1.46–2.58] | | [3.46–6.57] | | [1.25–1.88] | | |
| Pincer | 1.30 | 0.22 | 0.88 | 0.49 | 0.62 | 0.06 | 0.86 | 0.60 | 3.16–5.13 | 0.28 [0.07–0.47] | 1.30 × 10⁻⁰⁶ |
| | [0.85–1.97] | | [0.62–1.25] | | [0.37–1.02] | | [0.48–1.53] | | | | |
| AD | 0.87 | 0.64 | 1.34 | 0.13 | 1.06 | 0.83 | 1.86 | 0.02 | 0.81 | 0.41 |
| | [0.48–1.58] | | [0.91–1.97] | | | | | | | |
| **Adjusted analysis** | | | | | |
| Cam | 3.20 | 9.24 × 10⁻¹⁸ | 1.87 | 2.02 × 10⁻⁰⁷ | 1.94 | 5.74 × 10⁻⁰⁵ | 4.75 | 3.13 × 10⁻²¹ | 1.53 | 6.02 × 10⁻⁰⁵ |
| | [2.41–4.25] | | [1.48–2.36] | | [1.45–2.57] | | [3.44–6.57] | | [1.24–1.88] | | |
| Pincer | 1.30 | 0.22 | 0.86 | 0.41 | 0.63 | 0.08 | 0.81 | 0.47 | 3.25–5.30 | 0.28 |
| | [0.85–1.98] | | [0.61–1.23] | | [0.38–1.05] | | [0.45–1.45] | | | | |
| AD | 0.89 | 0.70 | 1.41 | 0.08 | 1.07 | 0.79 | 1.95 | 0.02 | 0.28 | 1.30 × 10⁻⁰⁶ |
| | [0.49–1.62] | | [0.96–2.08] | | [0.64–1.79] | | [1.13–3.35] | | [0.16–0.47] | | |

Table II: Results from logistic regressions examining the relationships between different hip morphologies, and rHOA, as well as grade ≥1 osteophytes and JSN in males. Unadjusted and adjusted results are shown in the form of odds ratios (OR), 95% confidence intervals (CI) and p-values (P). Adjusted models include age, height, weight and ethnicity. rHOA, radiographic hip osteoarthritis; OP, osteophyte; JSN, joint space narrowing.
### Table III

Results from logistic regression examining the relationships between different hip morphologies, and rHOA, as well as grade ≥1 osteophytes and JSN in females. Unadjusted and adjusted results are shown in the form of odds ratios (OR), 95% confidence intervals (CI) and p-values (P). Adjusted models include age, height, weight and ethnicity. rHOA, radiographic hip osteoarthritis; OP, osteophyte; JSN, joint space narrowing.

#### Females

|                  | Acetabular OP | Superior Femoral OP | Inferior Femoral OP | JSN |
|------------------|---------------|---------------------|---------------------|-----|
|                  | OR [95% CI]   | P                   | OR [95% CI]         | P   | OR [95% CI] | P   | OR [95% CI] | P   |
| **Unadjusted analysis** |
| Cam             | 2.73 [1.07–6.94] | 0.04 | 1.12 [0.51–2.47] | 0.78 | 2.01 [0.80–5.10] | 0.14 | 10.97 [4.93–24.39] | 4.24 × 10⁻⁶⁰ | 1.83 [1.03–3.25] | 0.04 |
| Pincer          | 1.30 [0.69–2.45] | 0.43 | 0.91 [0.60–1.39] | 0.68 | 1.24 [0.70–2.18] | 0.45 | 2.09 [0.97–4.48] | 0.06 | 4.03 [3.10–5.24] | 1.31 × 10⁻²⁵ |
| AD              | 0.64 [0.26–1.59] | 0.34 | 1.15 [0.76–1.75] | 0.50 | 0.68 [0.31–1.47] | 0.33 | 1.12 [0.40–3.13] | 0.83 | 0.31 [0.18–0.54] | 3.43 × 10⁻⁰⁵ |
| **Adjusted analysis** |
| Cam             | 2.47 [0.96–6.36] | 0.06 | 0.99 [0.45–2.21] | 0.99 | 1.83 [0.72–4.67] | 0.20 | 10.07 [4.49–22.61] | 2.13 × 10⁻⁰⁶ | 1.75 [0.97–3.14] | 0.06 |
| Pincer          | 1.23 [0.65–2.33] | 0.53 | 0.83 [0.54–1.26] | 0.38 | 1.15 [0.65–2.03] | 0.64 | 1.96 [0.91–4.23] | 0.09 | 4.05 [3.10–5.3] | 1.52 × 10⁻²⁴ |
| AD              | 0.72 [0.29–1.79] | 0.48 | 1.37 [0.90–2.09] | 0.15 | 0.75 [0.35–1.64] | 0.48 | 1.28 [0.46–3.62] | 0.64 | 0.34 [0.19–0.58] | 1.10 × 10⁻⁰⁴ |

**Fig. 2**

Logistic regression results are shown for the associations between cam morphology and osteophyte presence at three locations: acetabular, superior femoral, and inferior femoral head. Odds ratios are plotted with 95% confidence intervals either side. Results are presented as different models, diamonds represent the male only model (n = 3382), circles represent the female only model (n = 3425) and squares represent the combined sex model (n = 6807). Unadjusted results are shown by hollow shapes and results adjusted for age, height, weight and ethnicity are shown by filled shapes. The adjusted combined sex model also has sex as an additional covariate. Y-axis is natural log based.

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over coverage which could represent a limitation when examining this outcome against an acetabulum-based hip morphology.

The lack of association between AD and hip OA in our study is in keeping with a previous study by Gosvig et al., but contrary to other previous studies, in particular a systematic review which reported that longitudinal studies found acetabular under coverage associated with OA progression. This may because acetabular coverage can mimic osteophytes and vice versa, despite high resolution images being inspected individually it can still be difficult to discriminate the two features thus potentially preventing cross-sectional studies from detecting associations between AD and rHOA. Direct comparisons between studies are difficult because of the different LCEA cut-offs used to define AD, along with differences in the imaging modalities used and outcomes employed. For example, Saberi Hosnijeh et al. used a more stringent threshold of LCEA (<20°) (compared to <25° in the present study) and reported associations between AD and total hip replacement (THR) as opposed to rHOA or hip pain.

Whilst any mechanistic links cannot be reliably determined in the context of this cross-sectional analysis, it is possible that the relationship between cam morphology and rHOA is causal, such that pre-existing cam morphology causes aberrant biomechanical forces which in turn lead to osteophyte formation. Since the strongest associations were observed between cam morphology and inferior femoral osteophytes, as opposed to superior femoral and acetabular osteophytes, this suggest aberrant biomechanical forces are present throughout the joint. Our study did not show a predisposition for osteophytes at the site of impingement, i.e., acetabular or superior femoral head osteophytes. This aligns with a previous study that found cam-type hip shape modes obtained from statistical shape modelling derived from DXA scans were associated with osteophytes both superiorly and inferiorly on the acetabulum and femoral head measured on x-rays taken 5 years later. Other authors have suggested inferior femoral head osteophytes to be a marker of hip instability but further work is needed to understand how cam morphology might contribute to this.

The association between cam morphology and hip pain which we observed may partly be mediated by osteophyte formation, particularly superior osteophytes, adjustment for which led to partial attenuation of this relationship. Although not a formal mediation analysis this indicates that osteophyte formation may mediate the relationship between cam morphology and hip pain. This is consistent with findings from our recent study based on the same DXA images, where we found osteophytes at different locations to be independently associated with hip pain. This view is also in agreement with several other emerging lines of evidence that osteophytes are an important source of pain in hip OA.

This represents the largest population study to date of relationships between hip morphology and hip OA, which was made feasible by the development of automated means of deriving AA and LCEA on hip DXA scans. However, although well suited for derivation of hip morphology and rHOA, use of DXA scans has some inherent limitations. For example, when deriving LCEA, since only one hip is visualised per scan, it was not possible to adjust for pelvic tilt as performed when deriving equivalent measures from radiographs. Another limitation arises from examining only left hips when the hip pain measure used in our study was not side specific. The latter reduces precision, although this would likely bias our results towards the null rather than inducing false associations. Another limitation is the cross-sectional nature of our study. For example, it is possible that spurious associations may be introduced between hip morphology and rHOA, if measures such as AA and LCEA incorporate osteophytes because it is difficult to identify the true contour of the bone and as already mentioned we cannot comment on causality of any observations seen. Unfortunately, our study does not include measures of subchondral sclerosis or cysts which are well recognised constituents of rHOA again decreasing the precision of our measurement of rHOA. Additionally, DXA scans are done supine rather than weight bearing which could theoretically increase mJSW. However, a comparison between JSW on weight bearing and non-weight bearing hip x-rays found only a minimal change in JSW (0.1 mm mean difference) in those who already had JSN and OARSI clinical trial guidance suggests supine hip x-rays are acceptable for assessing rHOA. Finally, our study is based on 2-dimensional imaging which limits our ability to detect differences in hip morphology in planes better visualised on 3-dimensional imaging. Of note is that a recent study comparing x-rays with CT scans showed similar sensitivity and specificity between the two modalities when defining cam and pincer morphology.

In conclusion, using novel methods developed and applied to high resolution DXA images from a large cross-sectional study, we found that cam morphology is associated with hip OA, as reflected by rHOA and self-reported hip pain. These associations were strongest in men, in whom cam morphology was much more common than in women. We found associations between cam morphology and osteophytes to be located throughout the joint with the strongest relationship with those at the inferior femoral head. Further work is needed to understand the biomechanical consequences of cam morphology underlying the pattern of osteophytes with which this is associated, as a prelude to developing tailored strategies for reducing OA progression.

### Author contributions
All authors have made significant contributions to the conception and design of this study, the acquisition of data, its analysis and interpretation. All authors helped draft the article before approving...
the final version of this manuscript. Dr B Faber (ben.faber@bristol.ac.uk) takes responsibility for the integrity of the work in its entirety.

Conflict of interest
TC & CL have a patent Image processing apparatus and method for fitting a deformable shape model to an image using random forest regression voting. This is licensed with royalties to Audax, and to Optasia Medical. NH reports consultancy fees and honoraria from UCB, Amgen, Kyowa Kirin, Thornton Ross, Consilient.

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Supplementary data

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References

1. Hunter DJ, Bierma-Zeinstra S. Osteoarthritis. Lancet. 2019;393(10182):1745–59.
2. Registry NJ. report National Joint Registry Annual Report 2017 2017.
3. Faber BG, Frysz M, Tobias JH. Unpicking observational relationships between hip shape and osteoarthritis: hype or hope? Curr Opin Rheumatol 2020;32(1):110–8.
4. Lloyd-Roberts GC. Osteoarthritis. Postgrad Med 1955;31(362): 618–22.
5. Saberi Hosnjieh F, Zuiderwijk ME, Versteeg M, Smeete HT, Hofman A, Uitterlinden AG, et al. Cam deformity and acetabular dysplasia as risk factors for hip osteoarthritis. Arthritis Rheum 2017;69(1):86–93.
6. Thomas GE, Palmer AJ, Batra RN, Kiran A, Hart D, Spector T, et al. Subclinical deformities of the hip are significant predictors of radiographic osteoarthritis and joint replacement in women. A 20 year longitudinal cohort study. Osteoarthritis Cartilage 2014;22(10):1504–10.
7. Gosvig KK, Jacobsen S, Sonne-Holm S, Palm H, Troelsen A. Prevalence of malformations of the hip joint and their relationship to sex, groin pain, and risk of osteoarthritis: a population-based survey. J Bone Joint Surg Am 2010;92(5):1162–9.
8. Ganz R, Parvizi J, Beck M, Leunig M, Nötzli H, Siebenrock KA. Femoroacetabular impingement: a cause for osteoarthritis of the hip. Clin Orthop Relat Res 2003;417:112–20.
9. Murphy NJ, Eyles JP, Hunter DJ. Hip osteoarthritis: etiopathogenesis and implications for management. Adv Ther 2016;33(11):1921–46.
10. Frysz M, Gregory J, Aspden RM, Paternoster L, Tobias JH. Sex differences in proximal femur shape: findings from a population-based study in adolescents. Sci Rep 2020;10(1):4612.
11. Baird DA, Evans DS, Kamanu FK, Gregory JS, Saunders FR, Giuraniuc CV, et al. Identification of novel loci associated with hip shape: a meta-analysis of genomewide association studies. J Bone Miner Res 2019;34(2):241–51.
12. Hatzikoutoula K, Roposch A, Consortium DDHC, Shah KM, Clark MJ, Bratherton S, et al. Genome-wide association study of development dysplasia of the hip identifies an association with GDF5. Commun Biol 2018;1:56.
13. Agricola R, Weinans H. What causes cam deformity and femoroacetabular impingement: still too many questions to provide clear answers. Br J Sports Med 2016;50(5):263–4.
14. van Klij P, Heijboer MP, Gini AZ, Verhaar JAN, Waarsing JH, Agricola R. Cam morphology in young male football players mostly develops before proximal femoral growth plate closure: a prospective study with 5-yearfollow-up. Br J Sports Med 2019;53(9):532–8.
15. Griffin DR, Dickinson EJ, O’Donnell J, Agricola R, Awan T, Beck M, et al. The Warwick Agreement on femoroacetabular impingement syndrome (FAI syndrome): an international consensus statement. Br J Sports Med 2016;50(19):1169–76.
16. Palmer AJR, Ayyar Gupta V, Fernquest S, Rombach I, Dutton SJ, Mansour R, et al. Arthroscopic hip surgery compared with physiotherapy and activity modification for the treatment of symptomatic femoroacetabular impingement: multicentre randomised controlled trial. BMJ 2019;364:k1185.
17. Mansell NS, Rhon DI, Meyer J, Slevin JM, Marchant BG. Arthroscopic surgery or physical therapy for patients with femoroacetabular impingement syndrome: a randomized controlled trial with 2-year follow-up. Am J Sports Med 2018;46(6):1306–14.
18. Griffin DR, Dickinson EJ, Wall PDH, Achana F, Donovan JL, Griffin J, et al. Hip arthroscopy versus best conservative care for the treatment of femoroacetabular impingement syndrome (UK FASHIoN): a multicentre randomised controlled trial. Lancet 2018;391(10136):2225–35.
19. Agricola R, Heijboer MP, Roze RH, Reijman M, Bierma-Zeinstra SMA, Verhaar JAN, et al. Pincer deformity does not lead to osteoarthritis of the hip whereas acetabular dysplasia does: acetabular coverage and development of osteoarthritis in a nationwide prospective cohort study (CHECK). Osteoarthritis Cartilage 2013;21(10):1514–21.
20. Leunig M, Beaule PE, Ganz R. The concept of femoroacetabular impingement: current status and future perspectives. Clin Orthop Relat Res 2009;467(3):616–22.
21. Frank JM, Harris JD, Erickson BJ, Slikker 3rd W, Bush-Joseph CA, Salata MJ, et al. Prevalence of femoroacetabular impingement imaging findings in asymptomatic volunteers: a systematic review. Arthroscopy 2015;31(6):1199–204.
22. Yoshida K, Barr RJ, Galea-Soler S, Aspden RM, Reid DM, Gregory JS. Reproducibility and diagnostic accuracy of kellgren–lawrence grading for osteoarthritis using radiographs and dual-energy X-ray absorptiometry images. J Clin Densitom 2015;18(2):239–44.
23. Bycroft C, Freeman C, Petkova D, Band G, Elliott LT, Sharp K, et al. UK Biobank resource with deep phenotyping and genomic data. Nature 2018;562(7726):203–9.
24. Harvey NC, Matthews P, Collins R, Cooper C, Ukbma Group. Osteoporosis epidemiology in UK Biobank: a unique oppor-
tunity for international researchers. Osteoporos Int 2013;24(12):2903–5.
25. Faber BG, Ebsim R, Saunders FR, Frysz M, Lindner C, Gregory JS, et al. Osteophyte size and location on hip DXA scans are associated with hip pain: findings from a cross sectional study in UK Biobank. medRxiv 2021. 2021.04.26.21255905.
26. Ebsim R, Lindner C, Faber B, Frysz M, Saunders FR, Gregory JS, et al. Development of a machine learning-based fully automated hip annotation system for DXA scans. Proceedings of the Bone Research Society Annual Meeting 2020:2020.
27. Lindner C, Thiagarajah S, Wilkinson JM, arc OC, Wallis GA.
28. Kellgren JH, Lawrence JS. Radiological assessment of osteoarthritis of the hip throughout the hip: a survey of 4151 subjects of the Copenhagen Osteoarthritis Study. Acta Radiol 2008;49(4):436–41.
29. Faber BG, Baird D, Gregson CL, Gregory JS, Barr RJ, Aspden RM, et al. DXA-derived hip shape is related to osteoarthritis: findings from in the MrOS cohort. Osteoarthritis Cartilage 2017;25(12):2031–8.
30. Kijima H, Yamada S, Konishi N, Kubota H, Tazawa H, Tani T, et al. The differences in imaging findings between painless and painful osteoarthritis of the hip. Clin Med Insights Arthritis Musculoskelet Disord 2020;13. 1179544120946747.
31. Hunter DJ, McDougall JJ, Keefe FJ. The symptoms of osteoarthritis and the genesis of pain. Rheum Dis Clin N Am 2008;34(3):623–43.
32. Hardcastle SA, Dieppe P, Gregson CL, Hunter D, Thomas GE, Arden NK, et al. Prevalence of radiograph hip osteoarthritis is increased in high bone mass. Osteoarthritis Cartilage 2014;22(8):1120–8.
33. A Kanatani K, A Rangarajan P. Hyper least squares fitting of circles and ellipses. Comput Stat Data Anal 2011;55:2197–208.
34. Faber BG, Ebsim R, Saunders FR, Frysz M, Davey Smith G, Cootes T, et al. Deriving Alpha Angle from Anterior-Posterior Dual-Energy X-Ray Absorptiometry Scans: An Automated and Validated Approach. Wellcome Open Research, 2021, https://wellcomeopenresearch.org/articles/6-60/v1.
35. Agricola R, Heijboer MP, Bierma-Zeinstra SMA, Verhaar JAN, Weinans H, Waarsing JH. Cam impingement causes osteoarthritis of the hip: a nationwide prospective cohort study (CHECK). Ann Rheum Dis 2013;72(6):918–23.
36. Gosvig KK, Jacobsen S, Sonne-Holm S, Gebuhr P. The prevalence of cam-type deformity of the hip joint: a survey of 4151 subjects of the Copenhagen Osteoarthritis Study. Acta Radiol 2008;49(4):436–41.
37. van Klij P, Reiman MP, Waarsing JH, Reijman M, Bramer WM, Verhaar JAN, et al. Classifying cam morphology by the alpha angle: a systematic review on threshold values. Orthop J Sports Med 2020;8(8): 2325967120938312.
38. Roling MA, Mathijsen NMC, Bloem RM. Diagnostic sensitivity and specificity of dynamic three-dimensional CT analysis in detection of cam and pincer type femoroacetabular impingement. BMC Musculoskel Disord 2020;21(1):37.