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
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RESEARCH ARTICLE

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Increased development of radiographic hip osteoarthritis in individuals with high bone mass: a prospective cohort study

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Abstract

Background: Individuals with high bone mass (HBM) have a greater odds of prevalent radiographic hip osteoarthritis (OA), reflecting an association with bone-forming OA sub-phenotypes (e.g. osteophytosis, subchondral sclerosis). As the role of bone mineral density (BMD) in hip OA progression is unclear, we aimed to determine if individuals with HBM have increased incidence and/or progression of bone-forming OA sub-phenotypes.

Methods: We analysed an adult cohort with and without HBM (L1 and/or total hip BMD Z-score $\geq +3.2$) with pelvic radiographs collected at baseline and 8-year follow-up. Sub-phenotypes were graded using the OARSI atlas. Superior/inferior acetabular/femoral osteophyte and medial/superior joint space narrowing (JSN) grades were summed and Δ osteophyte and Δ JSN derived. Pain and functional limitations were quantified using the WOMAC questionnaire. Associations between HBM status and change in OA sub-phenotypes were determined using multivariable linear/logistic regression, adjusting for age, sex, height, total body fat mass, follow-up time and baseline sub-phenotype grade. Generalised estimating equations accounted for individual-level clustering.

Results: Of 136 individuals, 62% had HBM at baseline, 72% were female and mean (SD) age was 59 (10) years. HBM was positively associated with both Δ osteophytes and Δ JSN (adjusted mean grade differences between individuals with and without HBM $\beta_{\text{osteophyte}} = 0.30$ [0.01, 0.58], $p = 0.019$ and $\beta_{\text{JSN}} = 0.10$ [0.01, 0.18], $p = 0.019$). Incident subchondral sclerosis was rare. HBM individuals had higher WOMAC hip functional limitation scores ($\beta = 8.3$ [0.7, 15.98], $p = 0.032$).

Conclusions: HBM is associated with the worsening of hip osteophytes and JSN over an average of 8 years, as well as increased hip pain and functional limitation.

Keywords: Hip osteoarthritis, Progression, High bone mass, BMD, WOMAC

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Introduction

Osteoarthritis (OA) of the hip is highly prevalent, affecting approximately 1% of the worldwide population, significantly contributing to global disability [1]. Currently, no disease-modifying medications are available; therapy consists of pain management until severity warrants a total hip replacement (THR). Detection of risk factors for hip OA progression offers an opportunity to identify potential targets for the development of therapeutic interventions.

Higher bone mineral density (BMD) has been associated with *prevalent* hip OA in several case-control [2, 3] and population-based studies [4–7]. However, such analyses are complicated as BMD is often measured at the hip and therefore it is hard to determine whether increased BMD is a cause, or feature, of hip OA [4–6, 8]. In men with discordant hip OA, Arokoski et al. found femoral neck (FN)-BMD to be 4% higher in the more severely affected hip, reflecting increased FN volume (measured by MRI) [9]. This may reflect a process known as buttressing, whereby osteophytes extend across the FN to artefactually increase measured BMD [10]. However, Chaganti et al. identified a relationship between total hip (TH) cortical volumetric BMD (vBMD, measurement of which it not artefactually increased by bone size) and hip OA in 3886 men in the Study of Osteoporotic Fractures in Men (MrOS) [5]. Moreover, lumbar spine (LS)-BMD can be artefactually elevated by the presence of spinal osteophytes, a feature of spinal OA [4–6]. However, Nevitt et al. found that the relationship between LS-BMD and severe hip OA persisted despite adjustment for spinal osteophytes [4]. Furthermore, they found a relationship between calcaneal BMD and hip OA in over 4000 women from the Study of Osteoporotic Fractures (SOF), although of lower magnitude than seen for TH-BMD [4].

More recently, in a unique population of individuals with high bone mass (HBM), Hardcastle et al. reported those with HBM to have an increased odds of hip OA, reflecting a greater odds of osteophytosis but not joint space narrowing (JSN) [10]. HBM in index cases was defined as a TH or LS-BMD Z-score of at least +3.2, with a Z-score of at least +1.2 at the other site, identifying a generalised high BMD phenotype [11]. Genetic analysis of HBM individuals suggests that the HBM phenotype is at least in part determined by the polygenic inheritance of multiple BMD-associated loci [12]; thus, the temporal relationship is suggestive of a causal pathway between generalised high BMD and prevalent hip OA.

Fewer longitudinal analyses have addressed the relationship between BMD and the *incidence* and/or *progression* of hip OA. In the Johnston County OA project (JoCo) studying 928 older adults over median 6.5 years, although BMD did not predict incident radiographic hip

OA, it was inversely associated with incident *symptomatic* radiographic hip OA [13]. Furthermore, Bergink et al. identified an increased odds of both hip OA incidence and progression in those in the highest quartile of FN-BMD compared to the lowest quartile [14], whilst Hochberg et al. identified a dose-response relationship between both forearm and FN-BMD and the incidence of hip OA in SOF [15].

We aimed to determine the role of high BMD in hip OA by examining whether HBM individuals also have an increased odds of hip OA *incidence* and/or *progression*, using 8-year follow-up data collected in this unique cohort. We further aimed to determine the relationship between HBM and clinical features of OA, namely pain and functional limitations.

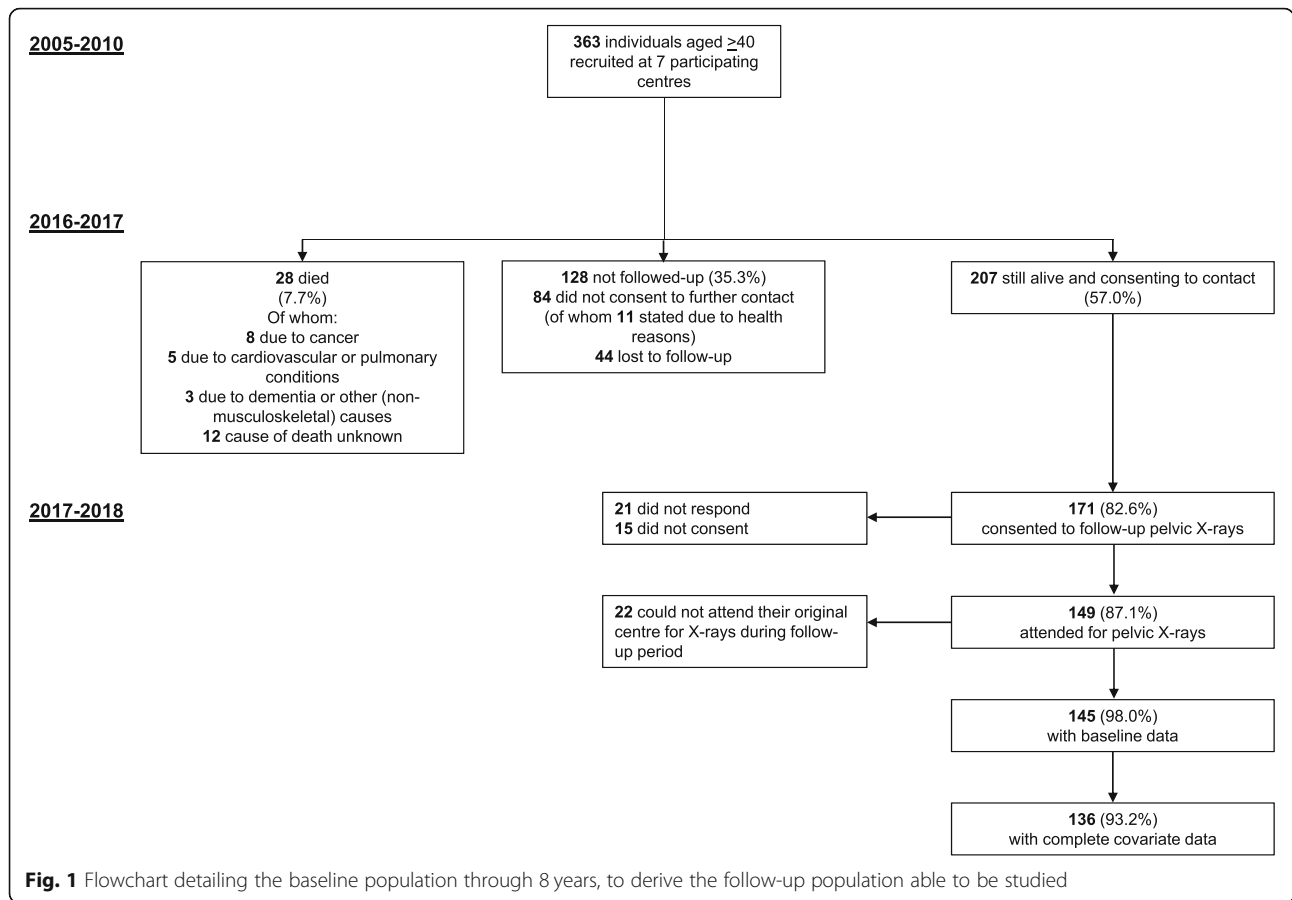
Methods

The high bone mass study

Participants were recruited as part of the UK-based HBM study. Index cases were initially identified by screening routine clinical National Health Service (NHS) dual-energy X-ray absorptiometry (DXA) databases (254, 736 DXA scans from seven UK hospitals) for individuals who had had *T* and/or *Z*-scores $\geq +4$. All 1290 DXA images were inspected by trained clinicians to exclude scans with artefactual elevations of DXA BMD (e.g. degenerative disease, OA, surgical/malignant artefacts). Full details of DXA database screening and participant recruitment have been published [11]. Generalised HBM was defined as a L1 or TH-BMD *Z*-score $\geq +3.2$ with a *Z*-score $\geq +1.2$ at the other skeletal site. A +3.2 threshold was consistent with the only published precedent for identifying HBM using DXA [16] and most appropriately differentiated generalised HBM from artefact [11]. The use of *Z*-score, rather than *T*-score, limited age bias [11]. Index cases passed on invitations to first-degree relatives and spouses/partners who underwent the same assessments. HBM in spouses was defined as per index cases. In first-degree relatives, HBM was defined as summed L1 plus TH *Z*-score $\geq +3.2$, as this identified relatives with BMD overlapping the index case BMD distribution [11]. Participants who were aged < 18, pregnant or unable to give written informed consent were excluded. Baseline recruitment of 363 adults (237 [65%] with HBM) ran between 2005 and 2010 across seven NHS centres (which participated in follow-up). Two hundred seven (57%) were alive and contactable in 2016; 149 (72%) of whom completed a postal questionnaire and attended for follow-up hip radiographs between 2017 and 2018 (Fig. 1).

Assessment of BMD

DXA scans were performed of the TH and LS at baseline and, after 8 years follow-up, of the TH, LS and total



body (TB) using standard protocols at each assessment centre. All but five (97%) participants re-attended their original centre, limiting measurement error due to differential procedures. DXA scans were performed on Hologic scanners in Bath, Bristol, Sheffield and St George’s London and GE Lunar scanners in Cambridge and Hull. Known differences in calibration exist between Hologic and Lunar [17, 18]. We limited systematic bias by converting TH and LS-BMD measures to standardised BMD (sBMD) [18, 19]. All images were visually

inspected for positioning and metal artefacts (e.g. hip prosthesis).

Assessment of radiographic OA

Standing anteroposterior (AP) pelvic X-rays were performed at baseline and follow-up using standard protocols at each centre. To limit observer bias, all radiographs were pooled for analysis, with the reader blinded to HBM status, demographics and timepoint. Radiographs were graded for semi-quantitative OA sub-

Table 1 Variables generated by Croft scoring and the OARSI atlas, with additional derived variables

Variable	Grading	Variable used in the analysis
Osteoarthritis (Croft score)	0–5	Progressive OA: Croft score ≥ 3 at baseline and an increase in score at follow-up Incident OA: Croft score < 3 at baseline and ≥ 3 at follow-up
Osteophytes		Change in osteophyte score: sum of all semi-quantitative osteophyte grades at follow-up minus their sum at baseline
Superior femoral	0–3	
inferior femoral	0–3	
Superior acetabular	0–3	
inferior acetabular	0, 1	
JSN		Change in JSN score: sum of both superior and medial semi-quantitative JSN grades at follow-up minus this sum at baseline
Superior	0–3	
Medial	0–3	

Abbreviations: OARSI Osteoarthritis Research Society International, JSN joint space narrowing, mJSW medial minimal joint space width

phenotypes (osteophytes and JSN, graded 0–3) and subchondral sclerosis (graded as present or absent) using the OARSI atlas [20]. The presence or absence of subchondral cysts was also evaluated. Overall OA was graded using Croft scoring [21]. Generated and derived progression variables are summarised in Table 1. Radiographs, viewed in open source ImageJ software [22], were inspected for poor image quality, rotation and/or tilt. All readings were performed by one assessor (AH) after focussed radiological training with a musculoskeletal radiologist (MW) and rheumatologist (SAH). A random selection of 72 hips (20%) were regraded to determine intra-rater reliability and graded by a second reader (SAH) to determine inter-rater reliability.

Assessment of clinical OA

Hip pain and limitation of function were assessed by postal questionnaire at 8-year follow-up. To limit non-response bias, the questionnaire was resent if not returned within 3 weeks. If still unreturned after a further 2 weeks, a reminder telephone call was made. The postal questionnaire included the short version WOMAC function scale [23, 24], which limited participant burden. The pain subscale (five questions relating to pain walking on a flat surface, ascending/descending stairs, at night, sitting or lying and standing upright) and function (seven questions relating to difficulty ascending stairs, rising from sitting, walking on flat, getting in/out of a car, putting on socks/stockings, rising from bed and sitting) each had five possible responses (none, mild, moderate, severe, extreme) scored 0–4, respectively. Missing values for pain or function questions were mean-imputed if a participant was missing one question on the pain scale and ≤ 3 on the function scale. Average scores were calculated for each subscale and scaled to give a score ranging from 0 to 100, with 0 representing no pain or functional limitation [25].

Covariate data

At baseline, structured interviews and clinical examination determined participant characteristics including age and standing height. Total body fat mass (TBFM) was assessed by TB DXA scan. Baseline menopausal status, alcohol consumption and history of hormone replacement therapy (HRT) use and smoking were determined by researcher-administered questionnaires. Baseline physical activity levels were determined using the International Physical Activity Questionnaire sent by post [26–28]. Menopausal status, history of smoking and highest educational status were determined by postal questionnaire at follow-up.

Statistical analysis

Associations between HBM status and OA incidence were determined by multivariable logistic regression. We included all hips to increase sample size and thus statistical power, using generalised estimating equations (GEE), which account for correlation between hips from the same individual and produce unbiased estimates in analyses of clustered data [29]. This analysis was restricted to hips with a Croft score < 3 at baseline.

Associations concerning change in osteophytes and JSN (continuous variables) were determined by multivariable GEE linear regression with robust standard errors to account for any non-normal distributions in outcome variables [30, 31]. Betas from analysis of continuous variables represent the difference in mean outcome between those with and without HBM (e.g. a beta of 1 for change in osteophyte score represents a 1-point greater increase in osteophyte score in HBM individuals). Osteophyte and/or JSN scores of 0 at baseline were included in analyses of change in osteophytes and JSN, optimising sample size. Analyses were initially performed unadjusted (model 1) and then adjusted for age, sex and time between radiographs (and baseline sub-phenotype score for continuous outcomes) (model 2). Our previous analyses found HBM to be associated with increased TBFM, with evidence suggesting this is a consequence rather than a cause of HBM [32]. Therefore, adiposity, hypothesised to be on the causal pathway in these analyses, was adjusted for as TBFM in model 3 along with height, to investigate a possible mediating effect of adiposity. Analyses were restricted to individuals with complete data for model 3. Statistical analysis was performed in Stata version 15 (Statacorp, USA) and R version 3.5.1.

Sensitivity analyses

Joints with THR were excluded from the main analyses; however, as THR may have been performed due to severe OA, those with a baseline Croft score < 3 and THR at follow-up were coded as incident OA cases, if they had stated that their THR was performed due to 'arthritis' ($n = 4$). Two individuals without OA at baseline, who had a THR at follow-up due to fracture, were coded as having no incident OA. A person-level analysis, using the sum of the osteophyte and the sum of the JSN scores for the two hips, used GEE to account for correlation within families. Incident OA in person-level analyses represents incident OA in either hip. A model adjusting for metal artefacts on DXA images, analyses removing individuals with DXA positioning errors potentially leading to under-measurement of TBFM (10 hips) and analyses removing individuals who visited a different study site for follow-up (10 hips) were all performed. To check that associations between HBM and change in OA sub-

phenotypes were not explained by bone size, we performed an additional analysis adjusting for the FN area (measured at follow-up). Finally, to check if conclusions were valid despite skewed continuous outcomes, all linear analyses were repeated using a Poisson model.

Results

Characteristics of the study population

Follow-up radiographic and covariate data were available for 136 individuals, with 62% having HBM (index cases or relatives with HBM). The proportion of individuals with HBM did not differ between the populations with and without follow-up data. Those with follow-up data were younger, were less likely to have had hip OA at baseline, to have ever smoked, to be postmenopausal, but were more physically active (Supplementary Table 1). Mean follow-up time for those with complete data was 8.2 (SD 1.0) years and did not differ between those with and without HBM (Table 2). HBM cases were more commonly female (85 vs 50%), with a trend towards a higher proportion of postmenopausal women. HBM individuals had greater baseline BMD (mean TH-BMD

1.24 vs 0.98 g/cm²), BMI (29.8 vs 27.5 kg/m²) and TBFM (33.0 vs 29.1 kg) than individuals without HBM (Table 2), consistent with previous observations in this population [11, 32]. Physical activity levels did not differ between HBM individuals and those without HBM.

Repeatability of radiographic OA variables

Weighted intra-rater kappa statistics for the Croft score and all osteophyte (except inferior acetabular) were > 0.7. The intra-rater reliability kappa for inferior acetabular osteophytes was 0.49, for medial JSN was 0.66 and for superior JSN was 0.49. AH observed no acetabular sclerosis or subchondral cysts. Intra-rater reliability for femoral sclerosis was perfect. Inter-rater weighted kappas for the Croft score and all osteophyte grades (except inferior acetabular) were > 0.6, representing substantial agreement [33]. The inter-rater kappa for inferior acetabular osteophytes was 0.38, with kappas of 0.48 for medial JSN and 0.39 for superior JSN. There was disagreement on the one observed case of femoral sclerosis and the one case of subchondral cysts, so these variables were excluded from analyses.

Table 2 Characteristics of the study population, constituting individuals with and without HBM, who were followed up at 8 years

	All, N = 136	HBM, N = 86	Relatives without HBM, N = 50	p value for difference
	N (%)			
Female gender	98 (72.1)	73 (84.9)	25 (50.0)	< 0.001
Postmenopausal ^b	75 (76.5)	59 (80.8)	16 (64.0)	0.087
Menopause transition during the follow-up period	11 (11.2)	6 (8.2)	5 (20.0)	0.177
History of HRT use ^f	49 (50.0)	39 (53.4)	10 (40.0)	0.508
History of smoking ^f	66 (48.9)	42 (49.4)	24 (48.0)	0.874
Physical activity category ^b				
Low	14 (10.7)	9 (11.0)	5 (10.2)	
Medium	46 (35.1)	26 (31.7)	20 (40.8)	0.567
High	71 (54.2)	47 (57.3)	24 (49.0)	
Education category ^f				
Up to GCSE/O level	55 (42.0)	42 (50.0)	13 (27.7)	
A level or equivalent	26 (19.9)	17 (20.2)	9 (19.2)	0.019
Degree or equivalent	50 (38.2)	25 (29.8)	25 (53.2)	
			Mean (SD)	
Age, years ^b	59.2 (10.2)	60.2 (9.9)	57.5 (10.6)	0.136
Height, cm ^b	167.8 (9.6)	166.1 (8.4)	170.8 (10.8)	0.005
Weight, kg ^b	81.5 (17.0)	82.1 (16.0)	80.6 (18.7)	0.619
BMI (kg/m ²) ^b	28.9 (5.5)	29.8 (5.6)	27.5 (5.1)	0.017
TBFM (kg) ^f	31.6 (10.6)	33.0 (10.9)	29.1 (9.5)	0.035
TH-BMD, g/cm ² ^b	1.143 (0.182)	1.242 (0.129)	0.976 (0.131)	< 0.001
L1-BMD, g/cm ² ^b	1.255 (0.215)	1.377 (0.149)	1.049 (0.141)	< 0.001
Follow-up time, years	8.2 (1.0)	8.2 (0.7)	8.2 (1.4)	0.817

Abbreviations: HBM high bone mass, HRT hormone replacement therapy, BMI body mass index, TBFM total body fat mass, TH-BMD total hip bone mineral density

^bAssessed at baseline

^fAssessed as follow-up

Table 3 Prevalence of radiographic and clinical sub-phenotypes of OA in the study population, stratified by HBM status

	All hips		HBM hips		Non-HBM hips	
	Total N	N (%) with sub-phenotype	Total N	N (%) with sub-phenotype	Total N	N (%) with sub-phenotype
OA (Croft ≥ 3)						
Baseline	285	22 (7.7)	179	13 (7.3)	106	9 (8.5)
Follow-up	275	33 (12.0)	173	24 (13.9)	102	9 (8.8)
Incident	257	18 (7.0)	162	15 (9.3)	95	3 (3.2)
Progressive	18	5 (27.8)	11	2 (18.2)	7	3 (42.9)
Hip replacement (identified on radiograph)						
Baseline	290	5 (1.7)	184	5 (2.7)	106	0
Follow-up	290	15 (5.2)	184	11 (6.0)	106	4 (3.8)
Incident	285	10 (3.5)	179	6 (3.4)	106	4 (3.8)
Osteophyte score						
Baseline	285		179		106	
0		203 (71.2)		126 (70.4)		77 (72.6)
1–4		75 (26.3)		50 (27.9)		25 (23.6)
≥ 5		7 (2.5)		5 (2.8)		4 (3.8)
Follow-up	275		173		102	
0		161 (58.6)		94 (54.3)		67 (65.7)
1–4		105 (38.2)		73 (42.2)		32 (31.4)
≥ 5		9 (3.3)		6 (3.5)		3 (2.9)
Delta	275		173		102	
< 1		201 (73.1)		121 (69.9)		80 (78.4)
1		48 (17.5)		32 (18.5)		16 (15.7)
> 1		26 (9.5)		20 (11.6)		6 (5.9)
JSN score						
Baseline	285		179		106	
0		253 (88.8)		160 (89.4)		93 (87.7)
1–2		27 (9.5)		16 (8.9)		11 (10.4)
≥ 3		5 (1.8)		3 (1.7)		2 (1.9)
Follow-up	275		173		102	
0		241 (87.6)		149 (86.1)		92 (90.2)
1–2		28 (10.2)		20 (11.6)		8 (7.8)
≥ 3		6 (2.2)		4 (2.3)		2 (2.0)
Delta	275		173		102	
< 1		261 (94.9)		161 (93.1)		100 (98.0)
1		12 (4.4)		10 (5.8)		2 (2.0)
> 1		2 (0.7)		2 (1.2)		0 (0.0)
	All individuals		HBM individuals		Relatives without HBM	
	Total N	Median (IQR)	Total N	Median (IQR)	Total N	Median (IQR)
WOMAC at follow-up						
Pain	145	0 (0, 25)	92	10 (0, 35)	53	0 (0, 15)
Function	145	3.6 (0, 25)	92	10.7 (0, 30.4)	53	0 (0, 14.3)
	Total N	N (%)	Total N	N (%)	Total N	N (%)
Hip replacement (self-reported)	145	16 (11.0)	92	13 (14.1)	53	3 (5.7)

Abbreviations: HBM high bone mass, OA osteoarthritis, JSN joint space narrowing, WOMAC Western Ontario and McMaster Universities Osteoarthritis Index

HBM and the incidence and progression of overall hip OA

Radiographic hip OA was observed in 7.7% of all 290 hips at baseline and 12.0% at follow-up (Table 3). Of the 257 OA-free hips at baseline, 7.0% developed OA. There was no clear evidence that HBM was associated with an increased risk of overall incident OA measured by Croft score, before (model 1, OR = 2.54 [95%CI 0.66, 9.71], Fig. 2) or after adjustment for age, sex and follow-up time (model 2, 1.65 [0.41, 6.70]). Due to the low baseline prevalence of overall OA defined as Croft score ≥ 3 (i.e. the presence of osteophytes and JSN), we were unable to analyse OA progression. Using Croft score ≥ 1 to define OA at baseline, 82 hips had potential to progress, of which 16 had a higher Croft score at follow-up than baseline (12 with HBM). However, no clear association between HBM and overall OA progression was observed (model 3, OR 4.14 [0.81, 21.3], Fig. 2). When combining incident and progressive OA to generate a variable for any incident or progressive hip OA, HBM was still not clearly associated with the overall change in hip OA severity (model 3, OR 1.72 [0.58, 5.11]).

Combined incidence and progression of radiographic hip OA sub-phenotypes

Of the total population, 28.8% hips had at least one osteophyte at baseline, rising to 41.5% at follow-up (Table 3). JSN was much less prevalent at baseline and follow-up (11.3% and 12.4%, respectively). In unadjusted analyses, we found evidence that individuals with HBM experienced greater changes in both osteophyte and JSN scores than individuals without HBM ($\beta_{\text{osteophyte}} = 0.30$ [0.05, 0.54], $p = 0.019$ and $\beta_{\text{JSN}} = 0.09$ [0.01, 0.16], $p = 0.019$, β reflects the difference in the mean change in osteophyte/JSN score between those with and without HBM). These associations persisted after adjustment for age, sex, follow-up time, baseline score, height and TBFM (model 3) (Fig. 2).

HBM and clinical features of hip OA

HBM was associated with 12-point [95% CI 5, 18] higher WOMAC pain scores and 13-point [7, 19] higher function scores in unadjusted analyses. Adjustment for age, sex, height and TBFM attenuated these relationships by

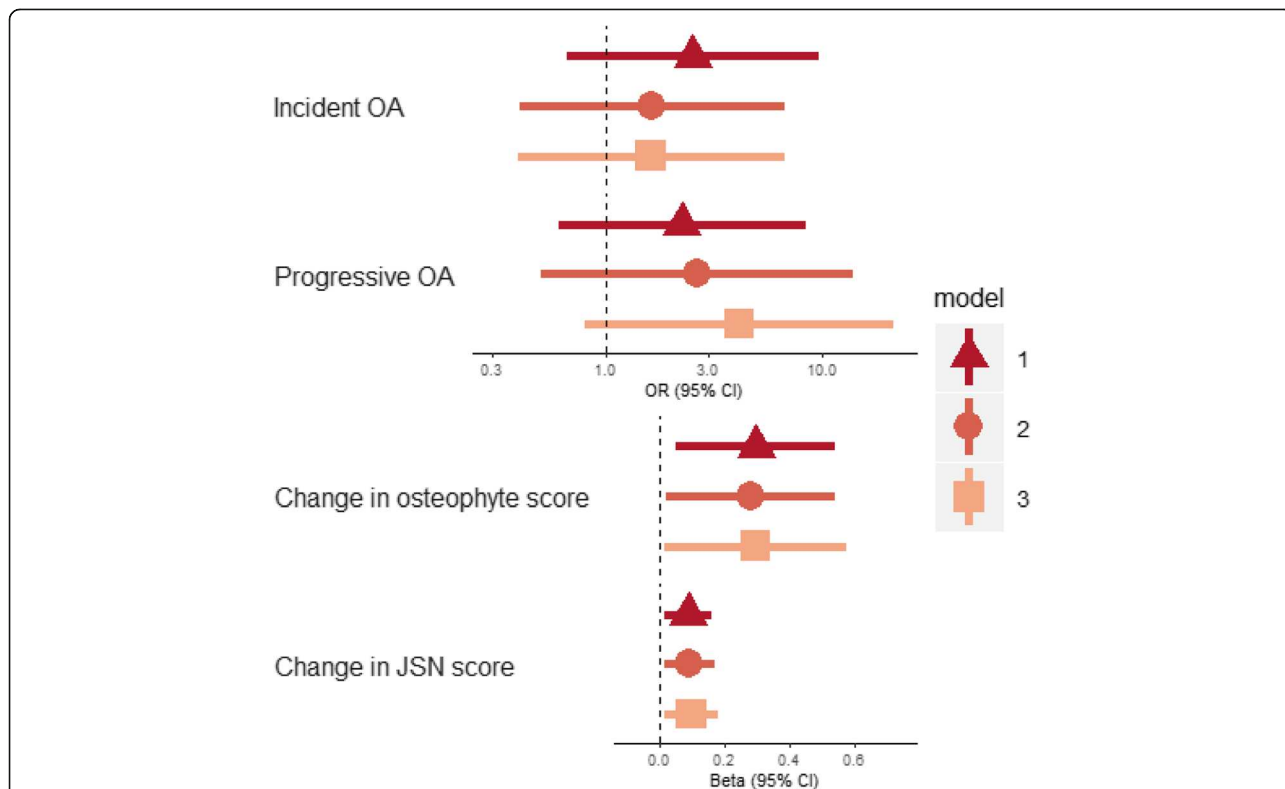


Fig. 2 Associations between HBM and incident and progressive OA and change in OA sub-phenotypes. Points for continuous outcomes represent the difference in mean outcome between individuals with and without HBM (for example, a beta of 1 for change in osteophyte score over 8 years or the increase in the size of an osteophyte already present). Points for binary outcomes represent the odds ratio for individuals with HBM compared to their relatives with normal BMD. Model 1: unadjusted; model 2: adjusted for age, sex and follow-up time (plus baseline score for continuous outcomes); model 3: adjusted for age, sex, follow-up time, height and TBFM (plus baseline score for continuous outcomes). $N_{\text{incident OA}} = 248$; $N_{\text{continuous outcomes}} = 263$. Abbreviation: JSN joint space narrowing

approximately one-third to one-half ($\beta_{\text{pain}} = 6.4 [-1.4, 14.2]$, $p = 0.105$ and $\beta_{\text{function}} = 8.3 [0.7, 15.8]$, $p = 0.032$, β represents the difference in mean WOMAC score between those with and without HBM). Further adjustment for osteophyte or JSN score at follow-up did not appear to explain these relationships (Fig. 3). There was some weak evidence supporting an increased odds of self-reported hip replacement in individuals with HBM who completed the follow-up questionnaire, compared to those without HBM (age, sex, height and TBFM-adjusted OR = 4.27 [0.94, 19.5], $p = 0.061$, $N = 148$).

Sensitivity analyses

Including six individuals with an incident THR and a Croft score < 3 at baseline in the analysis of incident OA did not alter conclusions drawn. Neither did removing 10 hips from individuals who visited a different study site for follow-up radiographs, removing hips from individuals with DXA positioning errors, nor additional adjustment for TB DXA artefact. Conclusions were unchanged when performing a person-level analysis accounting for within-family clustering, although CIs were wider due to the reduced sample size (Supplementary Figure 1). Additional adjustment for the FN area (as a measure of bone size) marginally attenuated effect estimates ($\beta_{\text{osteophyte}} = 0.26 [0.01, 0.52]$ to $0.21 [-0.02, 0.44]$ and $\beta_{\text{JSN}} = 0.08 [0.01, 0.16]$ to $0.07 [4.74 \times 10^{-3}, 0.13]$). Conclusions were unchanged when using a Poisson

model, although the association between HBM and change in JSN score was stronger than the association with change in osteophyte score; however, it should be noted that these analyses may be biased by the need to recode negative values as zero.

Discussion

We have found evidence for increased osteophyte development (i.e. incidence and/or progression) and JSN at the hip, over an average of 8 years in individuals with HBM, compared to their relatives without HBM. Radiographic JSN is thought to indirectly reflect cartilage loss [34]. Furthermore, individuals with HBM have more hip pain and limitation of function in their daily activities, which adds further evidence for increased OA severity in this population. Low statistical power limited our ability to draw strong conclusions about the relationship between HBM and overall incident OA, based on the Croft score.

Few studies have determined the association between BMD and hip OA incidence or progression. Bergink et al. observed a relationship between FN-BMD and both hip OA incidence and progression in the Rotterdam study population [14]. We have extended these findings by determining the relationship between high BMD and the incidence and/or progression of individual radiographic sub-phenotypes. Barbour et al. identified weak evidence for worsening osteophytes with increasing BMD in JoCo, but no evidence for a relationship with

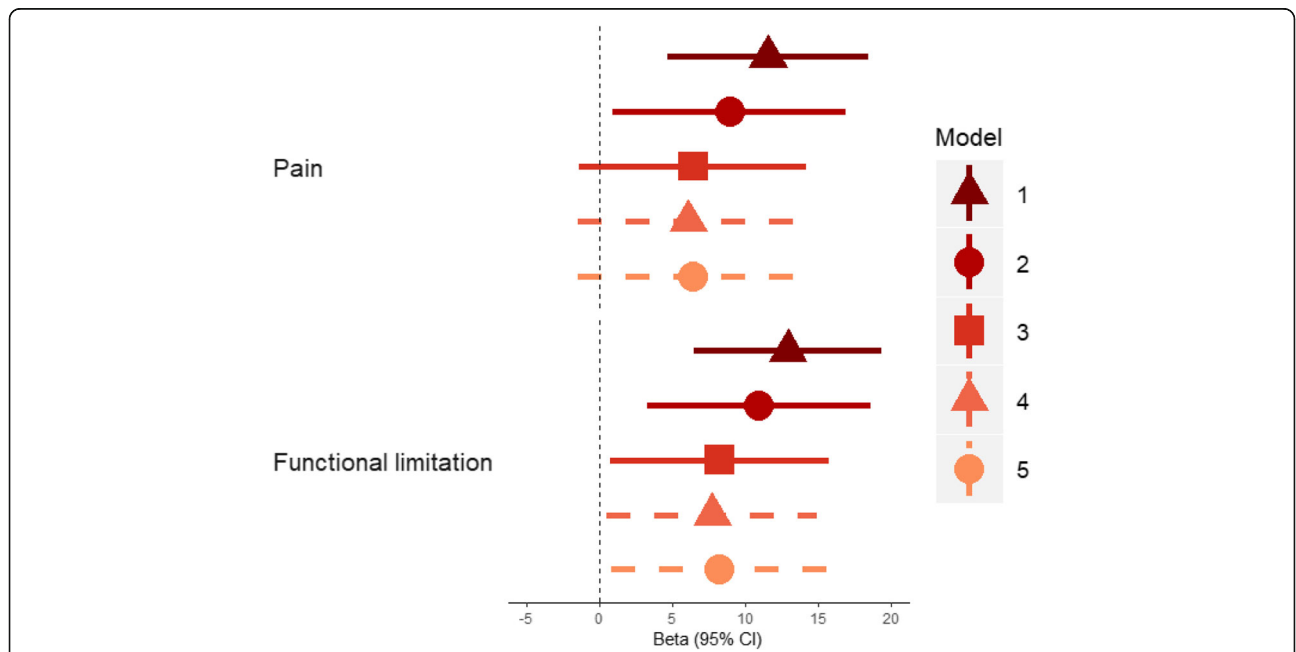


Fig. 3 Associations between HBM status and WOMAC pain and function sub-scale scores. Points represent the mean difference in WOMAC scores between individuals with HBM and relatives/spouses without HBM. Person-level analysis, accounting for clustering in families. Follow-up osteophyte and JSN score is the highest of the two hips. Model 1: unadjusted; model 2: adjusted for age and sex; model 3: adjusted for age, sex, height and total body fat mass; model 4: model 3 plus osteophyte severity at follow-up, model 5: model 3 plus JSN severity at follow-up. $N = 127$

JSN progression [13], which is inconsistent with our observed (albeit weak) relationship between HBM and change in JSN score. Hochberg et al. identified a dose-response relationship between BMD and subsequent incidence of OA in SOF [15]. However, this relationship was no longer present when defining incidence based on JSN alone. In our analyses, whilst we did not observe strong evidence for an association between HBM and incident hip OA, possibly due to low numbers, the direction of effect was consistent with previous findings.

Our observed relationship between HBM and hip pain is consistent with studies of population-based cohorts, which have identified an increased BMD in those reporting hip pain [4]. The severity of OA sub-phenotypes did not appear to explain the relationship between HBM and hip pain or functional limitations, suggesting that HBM individuals have an increased risk of clinical OA independent of radiographic severity. The WOMAC questionnaire measures pain over the past 48 h, which may explain why radiographic OA severity did not explain current pain, as pain could increase during stages of rapid OA progression not captured by radiographs [35]. An analysis of the Framingham and OA Initiative populations found that fewer than 25% of individuals with radiographic hip OA reported hip pain, and fewer than 20% reporting hip pain had radiographic hip OA [36]. It is possible that increased pain and functional limitation in the HBM population could reflect other conditions of the hip, such as bursitis [37] or features of a mild skeletal dysplasia, or inflammation not detected on the radiograph.

Increased TBFM in the HBM population [32] did not appear to explain the relationship between HBM and change in radiographic OA sub-phenotypes. Adjustment for the FN area, as a measure of bone size, only explained a small proportion of the relationship. Unfortunately, we do not have measures of FN width, a reported risk factor for hip OA progression [38]. It is plausible that HBM individuals would have greater FN width due to greater bone mass meaning measures of FN area may not equate to FN width in this population. Another factor which may mediate the relationship between HBM and development of hip OA sub-phenotypes is differences in hip shape. HBM individuals more commonly have features of cam-type deformity (i.e. larger femoral head size and reduced sphericity) compared to their relatives without HBM [39]. Evidence suggests that cam-type deformities are a risk factor for end-stage hip OA and hence potentially for hip OA progression [40, 41].

Although HBM is likely to be caused by the polygenic inheritance of multiple BMD loci [12], or the monogenic inheritance of rare variants [42], indicating that HBM precedes OA development, we cannot rule out the possibility that biological pleiotropy, rather than a causal

effect, explains our results. We have previously identified an increased prevalence of pelvic enthesophytes in the HBM population, leading to the hypothesis that HBM individuals may have a genetic predisposition to form extra bone [43]. We observed a stronger effect size for the relationship between HBM and change in hip osteophyte score, than we did for that between HBM and change in hip JSN score, which further suggests a 'bone-forming' phenotype in this population. Further evidence for pleiotropy was provided by Hackinger et al. who found weak evidence for a genetic correlation between hip OA and LS-BMD, but not hip OA and FN-BMD [44]. By performing a cross-phenotype meta-analysis between overall OA and LS-BMD, the authors identified novel loci in the *SMAD3* gene [44]. *SMAD3* is part of the transforming growth factor β (TGF β) signalling pathway, regulating osteoblast differentiation and thus bone formation. The first discovered hip OA locus, growth differentiation factor-5 (*GDF5*), is a ligand for this pathway [45], suggesting that this pathway contributes to both BMD and hip OA.

Strengths and limitations

The HBM study constitutes the largest population of individuals with extreme, unexplained, generalised HBM [11]. We analysed change in OA sub-phenotypes separately, which allowed us to detect the stronger relationship with osteophyte development compared to change in JSN. We analysed change in osteophytes and JSN as continuous measures, increasing statistical power to detect associations and reducing the possibility of a ceiling effect by increasing the range of possible values from 0 to 6 for JSN and 0 to 10 for osteophytes and eliminating the possibility of selection bias in a case-only analysis.

The method of identifying individuals from NHS DXA databases ascertained a predominantly female and older population such that a relatively large proportion were unable to be followed up after 8 years, due to death or poor health. Hence, there was a lower baseline prevalence of radiographic hip OA in the population able to be followed up, meaning we had limited power to assess hip OA incidence and progression based on the overall Croft score. The baseline cross-sectional study was powered to determine if the odds of OA differed between HBM individuals and their relatives with an expected recruitment of 200 cases and 200 controls [10]. However, loss-to-follow-up over 8 years reduced our sample size and a retrospective power calculation for the analyses presented here showed that we had approximately 65% power to detect the change in osteophyte and JSN scores reported here and lower power to detect a difference in proportion of incident hip OA between HBM individuals and their non-HBM relatives. Radiographs and DXA scans were performed using standard protocols at each

centre but were not standardised across centres. However, as 97% of individuals re-attended the same centre for follow-up, this is unlikely to affect our measures for change in radiographic features. Furthermore, measuring change in sub-phenotype variables did not separate hip OA sub-phenotype progression from incidence since these results had to be pooled to optimise sample size. As baseline and follow-up radiographs were not read as pairs, we did observe a few negative scores for change in osteophytes (8%) and change in JSN (1.5%), which were included in analyses, because removing these values as ‘measurement error’ would have biased results as there was likely to have been the same proportion of measurement error overinflating change, for which we would not have been able to account (hence the reasoning for not basing conclusions on the Poisson analysis). Radiographic grading of OA sub-phenotypes is subjective, which we limited using an established atlas [20], although our intra-rater and inter-rater reliability were low for a few variables, attenuating the conclusions we can draw from this analysis. As the reader was blinded to timepoint, it is unlikely that radiographic features were systematically under-graded at baseline and over-graded at follow-up, meaning measurement error is unlikely to explain our results. WOMAC scores were only collected at follow-up, and therefore, we cannot draw conclusions about the relationship between HBM and symptomatic OA progression. Finally, as HBM individuals represent a rare and extreme tail of the BMD distribution, findings may not be generalisable to the wider population.

Conclusions

We have found evidence for associations between HBM and worsening of radiographic sub-phenotypes of hip OA over 8 years. We further provide evidence for greater symptoms of OA in HBM individuals. These associations are independent of the elevated fat mass observed in HBM individuals. Further genetic analyses are planned to determine the BMI-independent causal role of BMD in hip OA progression and to identify the underlying biological pathways explaining these associations.

Supplementary Information

The online version contains supplementary material available at <https://doi.org/10.1186/s13075-020-02371-0>.

Additional file 1: Supplementary Table 1. Baseline characteristics for those with and without follow-up data. **Supplementary Figure 1.** associations between HBM and incident OA and incident and progressive OA sub-phenotypes in person-level analyses.

Abbreviations

AP: Anteroposterior; BMD: Bone mineral density; BMI: Body mass index; CI: Confidence intervals; DXA: Dual-energy X-ray absorptiometry; FN: Femoral neck; GEE: Generalised estimating equations; HBM: High bone mass; JoCo: Johnston County Osteoarthritis Project; JSN: Joint space

narrowing; LS: Lumbar spine; MrOS: Study of Osteoporotic Fractures in Men; OA: Osteoarthritis; REC: Research Ethics Committee; sBMD: Standardised bone mineral density; SOF: Study of Osteoporotic Fractures; TB: Total body; TBFM: Total body fat mass; TH: Total hip; THR: Total hip replacement; vBMD: Volumetric bone mineral density

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Authors’ contributions

Conceptualization: JHT and CLG. Methodology: AH, SAH, JHT and CLG. Software: JP. Formal analysis: AH, JHT and CLG. Investigation: AH, SAH, MF, EM, KESP, MKJ, MA, KM, MW, JHT and CLG. Writing of the original draft: AH, JT and CLG. Writing, review and editing: all authors. The authors read and approved the final manuscript.

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Availability of data and materials

The datasets during and/or analysed during the current study available from the corresponding author on reasonable request.

Ethics approval and consent to participate

Written informed consent was obtained in line with the Declaration of Helsinki [46]. The study was approved by the Bath Multi-centre Research Ethics Committee (REC reference 05/Q2001/78) and each local NHS REC. Follow-up data collection was approved by the Central Bristol REC and NHS Health Research Authority.

Consent for publication

Not applicable.

Competing interests

The authors have no competing interests to disclose.

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References

- Cross M, Smith E, Hoy D, Nolte S, Ackerman I, Fransen M, et al. The global burden of hip and knee osteoarthritis: estimates from the global burden of disease 2010 study. *Ann Rheum Dis*. 2014;73(7):1323–30.
- Stewart A, Black A, Robins SP, Reid DM. Bone density and bone turnover in patients with osteoarthritis and osteoporosis. *J Rheumatol*. 1999;26(3):622–6.
- Glowacki J, Tuteja M, Hurwitz S, Thornhill TS, Leboff MS. Discordance in femoral neck bone density in subjects with unilateral hip osteoarthritis. *J Clin Densitom*. 2010;13(1):24–8.
- Nevitt MC, Lane NE, Scott JC, Hochberg MC, Pressman AR, Genant HK, et al. Radiographic osteoarthritis of the hip and bone mineral density. The Study of Osteoporotic Fractures Research Group. *Arthritis Rheum*. 1995;38(7):907–16.
- Chaganti RK, Parimi N, Lang T, Orwoll E, Stefanick ML, Nevitt M, et al. Bone mineral density and prevalent osteoarthritis of the hip in older men for the Osteoporotic Fractures in Men (MrOS) Study Group. *Osteoporos Int*. 2010; 21(8):1307–16.
- Antoniades L, MacGregor AJ, Matson M, Spector TD. A cotwin control study of the relationship between hip osteoarthritis and bone mineral density. *Arthritis Rheum*. 2000;43(7):1450–5.
- Edwards MH, Paccou J, Ward KA, Jameson KA, Moss C, Woolston J, et al. The relationship of bone properties using high resolution peripheral quantitative computed tomography to radiographic components of hip osteoarthritis. *Osteoarthritis Cartilage*. 2017;25(9):1478–83.
- Burger H, van Daele PL, Odding E, Valkenburg HA, Hofman A, Grobbee DE, et al. Association of radiographically evident osteoarthritis with higher bone mineral density and increased bone loss with age. *Rotterdam Stud Arthritis Rheum*. 1996;39(1):81–6.
- Arokoski JP, Arokoski MH, Jurvelin JS, Helminen HJ, Niemitukia LH, Kroger H. Increased bone mineral content and bone size in the femoral neck of men with hip osteoarthritis. *Ann Rheum Dis*. 2002;61(2):145–50.
- Hardcastle SA, Dieppe P, Gregson CL, Hunter D, Thomas GE, Arden NK, et al. Prevalence of radiographic hip osteoarthritis is increased in high bone mass. *Osteoarthritis Cartilage*. 2014;22(8):1120–8.
- Gregson CL, Steel SA, O'Rourke KP, Allan K, Ayuk J, Bhalla A, et al. 'Sink or swim': an evaluation of the clinical characteristics of individuals with high bone mass. *Osteoporosis Int*. 2012;23(2):643–54.
- Gregson CL, Newell F, Leo PJ, Clark GR, Paternoster L, Marshall M, et al. Genome-wide association study of extreme high bone mass: contribution of common genetic variation to extreme BMD phenotypes and potential novel BMD-associated genes. *Bone*. 2018;114:62–71.
- Barbour KE, Murphy LB, Helmick CG, Hootman JM, Renner JB, Jordan JM. Bone mineral density and the risk of hip and knee osteoarthritis: the Johnston county osteoarthritis project. *Arthritis Care Res*. 2017;69(12):1863–70.
- Bergink AP, Rivadeneira F, Bierma-Zeinstra SM, Carola Zillikens M, Arfan Ikram M, Uitterlinden AG, et al. Are bone mineral density and fractures related to the incidence and progression of radiographic osteoarthritis of the knee, hip and hand in elderly men and women? The Rotterdam study. *Arthritis Rheumatol*. 2018;71(3):361–9.
- Hochberg MC. Do risk factors for incident hip osteoarthritis (OA) differ from those for progression of hip OA? *J Rheumatol Suppl*. 2004;70:6–9.
- Little RD, Carulli JP, Del Mastro RG, Dupuis J, Osborne M, Folz C, et al. A mutation in the LDL receptor-related protein 5 gene results in the autosomal dominant high-bone-mass trait. *Am J Hum Genet*. 2002;70(1):11–9.
- Shepherd JA, Fan B, Lu Y, Wu XP, Wacker WK, Ergun DL, et al. A multinational study to develop universal standardization of whole-body bone density and composition using GE Healthcare Lunar and Hologic DXA systems. *J Bone Miner Res*. 2012;27(10):2208–16.
- Fan B, Lu Y, Genant H, Fuerst T, Shepherd J. Does standardized BMD still remove differences between Hologic and GE-Lunar state-of-the-art DXA systems? *Osteoporos Int*. 2010;21(7):1227–36.
- Hanson J. Standardization of femur BMD. *J Bone Miner Res*. 1997;12(8): 1316–7.
- Altman RD, Gold GE. Atlas of individual radiographic features in osteoarthritis, revised. *Osteoarthritis Cartilage*. 2007;15(Suppl A):A1–56.
- Croft P, Cooper C, Wickham C, Coggon D. Defining osteoarthritis of the hip for epidemiologic studies. *Am J Epidemiol*. 1990;132(3):514–22.
- Schneider CA, Rasband WS, Eliceiri KW. NIH Image to ImageJ: 25 years of image analysis. *Nat Methods*. 2012;9(7):671–5.
- Whitehouse SL, Lingard EA, Katz JN, Learmonth ID. Development and testing of a reduced WOMAC function scale. *J Bone Joint Surg Br*. 2003; 85(5):706–11.
- Bellamy N, Buchanan WW, Goldsmith CH, Campbell J, Stitt LW. Validation study of WOMAC: a health status instrument for measuring clinically important patient relevant outcomes to antirheumatic drug therapy in patients with osteoarthritis of the hip or knee. *J Rheumatol*. 1988;15(12): 1833–40.
- Wylde V, Lenguerrand E, Gooberman-Hill R, Beswick AD, Marques E, Noble S, et al. Effect of local anaesthetic infiltration on chronic postsurgical pain after total hip and knee replacement: the APEX randomised controlled trials. *Pain*. 2015;156(6):1161–70.
- Craig CL, Marshall AL, Sjostrom M, Bauman AE, Booth ML, Ainsworth BE, et al. International physical activity questionnaire: 12-country reliability and validity. *Med Sci Sports Exerc*. 2003;35(8):1381–95.
- Hagstromer M, Oja P, Sjostrom M. The International Physical Activity Questionnaire (IPAQ): a study of concurrent and construct validity. *Public Health Nutr*. 2006;9(6):755–62.
- Hallal PC, Victora CG. Reliability and validity of the International Physical Activity Questionnaire (IPAQ). *Med Sci Sports Exerc*. 2004;36(3):556.
- Ballinger GA. Using generalized estimating equations for longitudinal data analysis. *Organ Res Methods*. 2004;7(2):127–50.
- Huber PJ, editor. The behavior of maximum likelihood estimates under nonstandard conditions. *Proceedings of the Fifth Berkeley Symposium on Mathematical Statistics and Probability, volume 1: statistics*; 1967 1967; Berkeley, Calif: University of California Press.
- White H. A heteroskedasticity-consistent covariance matrix estimator and a direct test for heteroskedasticity. *Econometrica*. 1980;48(4):817–38.
- Gregson CL, Paggiosi MA, Crabtree N, Steel SA, McCloskey E, Duncan EL, et al. Analysis of body composition in individuals with high bone mass reveals a marked increase in fat mass in women but not men. *J Clin Endocrinol Metab*. 2013;98(2):818–28.
- Landis JR, Koch GG. The measurement of observer agreement for categorical data. *Biometrics*. 1977;33(1):159–74.
- Guermazi A, Hunter DJ, Roemer FW. Plain radiography and magnetic resonance imaging diagnostics in osteoarthritis: validated staging and scoring. *J Bone Joint Surg Am*. 2009;91(Suppl 1):54–62.
- Wesseling J, Bierma-Zeinstra SMA, Kloppenburg M, Meijer R, Bijlsma JWJ. Worsening of pain and function over 5 years in individuals with 'early' OA is related to structural damage: data from the Osteoarthritis Initiative and CHECK (Cohort Hip & Cohort Knee) study. *Ann Rheum Dis*. 2015;74(2):347.
- Kim C, Nevitt MC, Niu J, Clancy MM, Lane NE, Link TM, et al. Association of hip pain with radiographic evidence of hip osteoarthritis: diagnostic test study. *BMJ*. 2015;351:h5983.
- Nieuwenhuijse MJ, Nelissen RG. Hip pain and radiographic signs of osteoarthritis. *BMJ*. 2015;351:h6262.
- Javadi MK, Lane NE, Mackey DC, Lui LY, Arden NK, Beck TJ, et al. Changes in proximal femoral mineral geometry precede the onset of radiographic hip osteoarthritis: the study of osteoporotic fractures. *Arthritis Rheum*. 2009; 60(7):2028–36.
- Patel A, Baird D, Hardcastle S, Gregory J, Aspenden R, Faber B, et al. Do alterations in hip shape explain the increased risk of hip osteoarthritis in individuals with high bone mass? [Conference abstract]. *Bone research society annual meeting 2016*, Liverpool, UK. 2016.
- Nicholls AS, Kiran A, Pollard TCB, Hart DJ, Arden CPA, Spector T, et al. The association between hip morphology parameters and nineteen-year risk of end-stage osteoarthritis of the hip: a nested case-control study. *Arthritis Rheumatism*. 2011;63(11):3392–400.
- Agricola R, Heijboer MP, Bierma-Zeinstra SM, Verhaar JA, 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.
- Gregson CL, Bergen DJM, Leo P, Sessions RB, Wheeler L, Hartley A, et al. A rare mutation in SMAD9 associated with high bone mass identifies the SMAD-dependent BMP signaling pathway as a potential anabolic target for osteoporosis. *J Bone Miner Res*. 2020;35(1):92–105.
- Hardcastle SA, Dieppe P, Gregson CL, Arden NK, Spector TD, Hart DJ, et al. Osteophytes, enthesophytes, and high bone mass a bone-forming triad with potential relevance in osteoarthritis. *Arthritis Rheumatol*. 2014;66(9): 2429–39.

44. Hackinger S, Trajanoska K, Styrkarsdottir U, Zengini E, Steinberg J, Ritchie GRS, et al. Evaluation of shared genetic aetiology between osteoarthritis and bone mineral density identifies SMAD3 as a novel osteoarthritis risk locus. *Hum Mol Genet.* 2017;26(19):3850–8.
45. Cibrian Uhalte E, Wilkinson JM, Southam L, Zeggini E. Pathways to understanding the genomic aetiology of osteoarthritis. *Hum Mol Genet.* 2017;26(R2):R193–r201.
46. World Medical Association Declaration of Helsinki ethical principles for medical research involving human subjects. *J Am Med Assoc.* 2013;310(20): 2191–4.

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