# **RESEARCH ARTICLE Example 2014 12:30 The SEAR CH ACCESS**



# Does cartilage volume measurement or radiographic osteoarthritis at baseline independently predict ten-year cartilage volume loss?

Andrew McBride<sup>1</sup>, Hussain Ijaz Khan<sup>1\*</sup>, Dawn Aitken<sup>1</sup>, Louisa Chou<sup>1</sup>, Changhai Ding<sup>1</sup>, Leigh Blizzard<sup>1</sup> , Jean-Pierre Pelletier<sup>3</sup>, Johanne Martel-Pelletier<sup>3</sup>, Flavia Cicuttini<sup>2</sup> and Graeme Jones<sup>1</sup>

# Abstract

**Background:** The aim of this study was to examine whether cartilage volume as measured by MRI and radiographic osteoarthritis (OA) at baseline predict cartilage volume loss over ten years independent of each other and other structural co-pathologies.

Methods: 219 participants [mean-age 45(26–61); 57 % female] were studied at baseline and ten years. Approximately half were the adult offspring of subjects who underwent knee replacement for OA and the remainder were randomly selected controls. Joint space narrowing (JSN) and osteophytes were assessed on radiographs and cartilage volume (tibiofemoral), cartilage defects, bone marrow lesions and meniscal tears/ extrusion were assessed on MRI.

**Results:** Mean absolute and percentage per annum cartilage volume loss was 1284 mm<sup>3</sup> and 1.91 % respectively in the medial compartment and 1007 mm<sup>3</sup> and 1.38 % respectively in the lateral compartment. Higher baseline tibiofemoral cartilage volume was independently associated with greater absolute cartilage volume loss in both medial (β(95 % CI) = -300 (-399,-200)) and lateral (β = -338 (-443,-233)) compartments and percentage per annum loss in the lateral compartment( $\beta = -0.15$  (-0.29, -0.01)). Baseline JSN and osteophytes were associated with cartilage volume loss in the univariable analysis, however these associations did not persist after adjustment for other structural co-pathologies.

**Conclusion:** Cross-sectional cartilage volume measurement independently predicts cartilage volume loss over 10 years and can be used to identify fast progressors in clinical trials. Radiographic JSN and osteophytes on the other hand are a reflection of other co-pathologies assessed on MRI and do not independently predict cartilage volume loss over 10 years.

Keywords: Knee, Osteoarthritis, Cartilage volume, Magnetic resonance imaging, Radiographs

\* Correspondence: Hussain.Khan@utas.edu.au

Andrew McBride and Hussain Ijaz Khan are co-first authors <sup>1</sup>Menzies Institute for Medical Research, University of Tasmania, Medical

Science 1 Building, Private Bag 23 17-Liverpool Street, Hobart 7000, Australia Full list of author information is available at the end of the article



© 2016 McBride et al. **Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

#### Background

Osteoarthritis (OA) is characterised by whole joint abnormalities including gradual cartilage volume loss [1]. Recent studies suggest that a cascade of structural changes occur in OA that involve sub-chondral bone expansion, bone marrow lesions (BMLs), meniscal tears, extrusion and eventually gradual loss of articular cartilage [1–7]. Loss of cartilage volume starts around the age of 40 years when radiographic changes are uncommon. [8] In early OA cartilage swelling appears to precede volume loss [6, 9]. This is supported by longitudinal evidence that higher baseline cartilage volume is associated with greater volume loss over a two-year period in early OA [2]. In patients with established OA, lower baseline cartilage volume appears to predict loss over a similar period [3].

Radiographic osteoarthritis (ROA) score has also been found to predict cartilage volume loss [10]. Whether this association is due to the presence of osteophytes or joint space narrowing (JSN) remains controversial. One study found that both JSN and osteophytes act as independent predictors of volume loss in a cohort of randomly selected older adults from community over a two-year period [10]. Other studies have shown that knees with definite osteophytes but without JSN do not show significantly greater rates of cartilage volume loss compared to healthy knees over a one-year period [11]. Similarly studies have shown that both presence [12, 13] and severity of ROA [11] is associated with cartilage thickness loss as well. To our knowledge no papers have looked at the association between ROA scores in early disease and volume loss over a ten-year period.

The aim of this study, therefore, was to examine whether cartilage volume as measured by MRI and ROA at baseline predict cartilage volume loss over ten years independent of each other and other structural co-pathologies.

#### Methods

#### Study subjects

This study was conducted as part of the Offspring study, which is an ongoing population-based study. The Offspring study began in southern Tasmania (primarily in the city of Hobart) in June 2000. Matched sampling was used to recruit the study participants (mean age 45 (26–61) years; 58 % females). Half of the participants were the adult offspring of patients who had a knee replacement performed for idiopathic knee OA at any Hobart hospital from 1996 to 2000 [5]. The diagnosis was confirmed by reference to the medical records of the orthopaedic surgeon and the original radiographs when possible. The other half were age and sex matched controls, randomly selected from the population with no history of knee OA in either parent. Controls were randomly selected from the electoral roll in southern Tasmania (population 229,000), a comprehensive population listing. This study includes data from the baseline visit, 2 year and 10 year follow up.

The Southern Tasmanian Health and Medical Human Research Ethics Committee approved the protocol, and written informed consent was obtained from all participants. Participants were excluded if they had a contraindication to MRI (including metal sutures, presence of shrapnel, iron filing in eye, or claustrophobia). Participants were also excluded if they had undergone a knee replacement surgery or did so after the commencement of the study. Knee pain and knee injury were not a basis for exclusion.

#### Anthropometrics

Weight was measured to the nearest 0.1 kg (with the subject's shoes, socks, and bulky clothing removed), with a single pair of electronic scales (Delta Model 707; Seca, Munich, Germany) that were calibrated using a known weight at the beginning of each clinic session. Height was measured to the nearest 0.1 cm (with shoes and socks removed) using a stadiometer. Body mass index (BMI) was calculated as weight  $(kg)/height$   $(m<sup>2</sup>)$ .

#### Magnetic resonance imaging

MRI of the right knee was performed as described previously [14–16]. All knees were imaged in the sagittal plane on a 1.5-T whole-body magnetic resonance unit (Picker International, USA) using a commercial transmit-receive extremity coil. The following image sequence was used: (i) a T1-weighted fat-suppressed 3D gradient-recalled acquisition in the steady state, flip angle 55°, repetition time 58 msec, echo time 12 msec, field of view 16 cm, 60 partitions,  $512 \times 512$ –pixel matrix, slice thickness of 1.5 mm without an interslice-gap; and (ii) a T2-weighted fat saturation 2D fast spin echo, flip angle 90°, repetition time 3067 ms, echo time 112 ms, field of view 16 cm, 15 partitions,  $256 \times 256$  matrix, slice thickness of 4 mm with an interslice gap of 0.5–1.0 mm.

#### Cartilage volume assessment

Knee cartilage volume was evaluated at baseline and 10 years by a trained observer on T1-weighted gradient echo MR images. Knee cartilage volume was determined by means of image processing on an independent workstation at baseline and follow up. The volumes of individual cartilage plates (medial tibia and femora, and lateral tibia and femora) were isolated from the total volume by manually drawing dis-articulation contours around the cartilage boundaries on a section by section basis. These data were then resampled by means of bilinear and cubic interpolation (area of  $312 \times 312$  µm by 1.5 mm thickness, continuous sections) for the final three-dimensional rendering to calculate the cartilage volume.

Tibial cartilage volume was assessed using Osiris (University of Geneva, Switzerland) software as previously described  $[14, 17]$ . The coefficient of variation $(CV)$  ranged from 2.1 to 2.2 % for intra-observer repeatability [18]. Femoral cartilage volume was determined using Cartiscope (ArthroLab, Montreal, Canada), as previously described [19, 20]. The CV was approximately 2 % for intra-observer and inter-scan repeatability [20]. Total cartilage volume was calculated as: tibial + femoral cartilage volume.

Absolute cartilage volume loss was calculated as: follow-up total cartilage volume - baseline total cartilage volume. Percentage per annum cartilage volume loss was calculated as: ((absolute cartilage volume loss/baseline cartilage volume)/time period between MRI acquisition at baseline and visit-3)  $\times$  100.

#### Cartilage defects

Cartilage defects were assessed at baseline and 10 years on T1-weighted gradient echo MR images at the medial tibial, medial femoral, lateral tibial, and lateral femoral sites on a 0–4 scale, as previously described [16]: grade  $0 =$  normal cartilage; grade  $1 =$  focal blistering and intracartilaginous low-signal intensity area with an intact surface and base; grade  $2$  = irregularities on the surface or base and loss of thickness  $<50$  %; grade  $3 =$  deep ulceration with loss of thickness  $>50$  %; and grade  $4 = \text{full}$ thickness chondral wear with exposure of subchondral bone. Intraobserver reliability (expressed as intraclass correlation coefficient (ICC)) ranged from 0.89 to 0.90. Interobserver reliability was assessed in 50 MR images and yielded an ICC of 0.85–0.90 [16].

#### Meniscal tears

Meniscal tears were assessed by a trained observer on T1-weighted gradient echo and T2-weighted (side by side) MR images at visit-2 and 3 of the study as previously described [19]. The proportion of the menisci affected by a tear was scored separately  $(0-2 \text{ scale}; 0=$ absence of a tear,  $1 =$  simple tear of different types: longitudinal, oblique, radial or horizontal,  $2 =$  complex tear signifying loss > 50 % area of meniscal tissue) at the anterior, middle, and posterior horns (medially/laterally). Anterior, middle and posterior scores were summed to get medial and lateral meniscal tear scores. The intraand inter-observer correlation coefficient ranged from 0.86 to 0.96 [20]. Meniscal tears were scored at visit-2 of the Offspring study, 2 years after the baseline visit.

#### Meniscal extrusion

Meniscal extrusion was assessed by a trained observer on T1-weighted gradient echo MR images at baseline and 10 years as previously described [19]. The proportion of the menisci affected by a partial or full extrusion was scored separately (yes/no) at the anterior, middle, and

posterior horns (medially/laterally). Anterior, middle and posterior scores were summed to get medial and lateral meniscal tear/extrusion scores. The intra- and interobserver correlation coefficient ranged from 0.85 to 0.92 for meniscal extrusion [20].

#### Bone marrow lesions

Bone marrow lesions (BMLs) were assessed on fat suppressed T2-weighted MR images as described previously [21]. BMLs were defined as areas of increased signal intensity in the sub-chondral bone at the medial tibial, medial femoral, lateral tibial, lateral femoral, superior patellar and inferior patellar sites. One trained observer scored the BMLs by measuring the maximum area of the lesion in a specific compartment. The observer manually selected the MRI slice with the greatest BML size. The BML with the highest score was used if more than one lesion was present at the same site. The ICC was 0.97. BMLs were scored at visit-2 of the Offspring study, 2 years after the baseline visit.

#### Radiography

A standing anteroposterior semiflexed x-ray of the right knee was taken in all subjects at baseline and 10 years. The angle was kept to 10–15° by a purpose built goniometer. The tube to film and tube to tibial plateau angle was 90°. Daily quality assurance was performed on the equipment. Radiographs were scored individually for osteophytes and joint space narrowing (JSN), as described previously [22]. Each of the following four features was scored on a scale from 0 to 3 ( $0 =$  normal and  $3 =$  severe): medial JSN, lateral JSN, medial osteophytes (femoral and tibial combined) and lateral osteophytes (femoral and tibial combined). Each score was arrived at by consensus with two readers (LC, AM) simultaneously assessing the radiograph with immediate reference to the Osteoarthritis Research Society International (OARSI) atlas [23]. ROA score was calculated by adding JSN (medial and lateral sites) and osteophytes (medial and lateral tibial and femoral sites) scores. A non-zero score in either JSN or osteophytosis was regarded as evidence of any ROA. Total ROA score had a possible range of 0–18. Reproducibility was assessed in 50 radiographs, two weeks apart, and yielded an agreement (linear weighted kappa value) of 0.87–1.00 for osteophytes and 0.94–1.00 for JSN (p-value < 0.001).

Readers for all the scans were either musculoskeletal radiologists with several years of experience in OA research or health professionals trained by musculoskeletal radiologists. Readers were not blinded to the chronological sequence of the radiographs and MRI scans.

#### Statistical analysis

T-tests and chi-square tests were used to compare differences in means and proportions as appropriate when

examining demographic, cartilage volume and radiographic data. Baseline characteristics of the participants were split into two groups for comparison using mean total cartilage volume loss (absolute) over 10 years: (i) less than mean total cartilage volume loss; (ii) greater than or equal to mean total cartilage volume loss.

Linear regression analysis was used to examine the association between baseline radiographic structures/cartilage and cartilage volume loss (absolute and percentage) over 10 years. β-coefficients were standardised to describe the association between baseline radiographic structures/cartilage volume and cartilage volume loss, so that cartilage volume loss was expressed as loss per standard deviation change in the predictor variables [21, 24, 25]. Multivariable analysis was adjusted for age, sex, BMI, offspringcontrol status, radiographic structures/cartilage volume at baseline and MRI structures which had a higher prevalence (or showed a similar trend) in participants with greater than or equal to mean total cartilage volume loss. All the associations between baseline JSN and cartilage loss were adjusted for baseline osteophytes and vice versa. Interactions terms were calculated to examine significant differences between the offspring and control groups.

Further sub-analyses looking at the association between baseline cartilage volume and/or ROA and absolute cartilage volume loss stratified by mean age was performed to look at the effect of advancing age on the associations described in the study.

To counter the effect of regression to the mean/tracking, when describing the association between baseline cartilage volume and cartilage volume loss, further subgroup analysis was done with baseline cartilage volume stratified by the mean value.

A P-value of less than 0.05 (two-tailed) was considered statistically significant. All statistical analyses were performed on Intercooled Stata 12.0 for windows (Stata-Corp LP).

#### Results

Of the 371 participants included in the Offspring study, 219 between the ages of 26 and 61 years were followed up after 10 years. The characteristics of participants who were followed up compared to participants who were lost to follow up were as follows, respectively: age: 45.25  $(\pm 6.67)$  vs 45.07  $(\pm 7.15)$  years,  $p = 0.806$ ; female sex: 57 % vs 59 %,  $p = 0.749$ ; BMI: 27.2 ( $\pm 4.96$ ) vs 26.8  $(\pm 4.31)$ ,  $p = 0.499$ ; offspring 52 % vs 47 %,  $p = 0.891$ ; knee ROA: 18 % vs 15 %,  $p = 0.486$  and total tibiofemoral cartilage volume at baseline  $(mm^3)$ : 14199 (±3463) vs 14113  $(\pm 3410)$ ,  $p = 0.611$ .

Table 1 describes the baseline characteristics of the group stratified by the mean total cartilage volume loss over 10 years. The average age of the cohort was 45 years. Participants with greater than the mean absolute volume loss were significantly older, had a significantly lower percentage of male participants, a significantly higher prevalence of medial JSN, medial osteophytes and any meniscal tear, and a higher medial and lateral tibiofemoral cartilage volume at baseline visit.

Both absolute and percentage per annum cartilage volume loss were higher in the medial tibiofemoral compartment compared to the lateral tibiofemoral compartment. Mean absolute and percentage per annum cartilage



Table 1 Baseline characteristics of participants split by total (tibiofemoral) cartilage loss (absolute) over 10 years <sup>a</sup>

<sup>a</sup>Mean (SD) except for percentages. P-values determined by t-test or  $x^2$  test (where appropriate)

Bold font signifies statistically significant results

volume loss was 1284 mm<sup>3</sup> and 1.91 % respectively in the medial compartment and 1007 mm<sup>3</sup> and 1.38 % respectively in the lateral compartment.

Figure 1a describes the association between baseline tibiofemoral cartilage volume and absolute cartilage volume loss. A higher baseline cartilage volume was associated with higher absolute cartilage volume loss over 10 years. Figure 1b describes the association between baseline ROA score and absolute cartilage volume loss. ROA score ranged from 0 to 6 (possible range 0–18) in the study population at the baseline visit. A higher baseline ROA score was associated with higher absolute cartilage volume loss on average over 10 years.

Table 2 describes the association between the baseline tibiofemoral cartilage volume and cartilage volume loss over 10 years. Baseline cartilage volume was significantly associated with absolute cartilage volume loss over 10 years in both compartments in the multivariable analysis. Further adjustment for cartilage defects and BMLs did not change the effect size considerably. There was a similar trend for the association between the baseline tibiofemoral cartilage volume and percentage per annum cartilage volume loss but the association reached statistical significance in the lateral compartment



Table 2 Association between baseline cartilage volume and cartilage volume loss over 10 years

<sup>a</sup> Adjusted for age, sex, BMI, offspring-control status, ROA at baseline, meniscal tears at visit-2 and meniscal extrusion at baseline where appropriate		
Lateral tibiofemoral (per SD)	$-226 (-303,-148) -338 (-443,-233)$	
Lateral (tibiofemoral) cartilage volume loss (absolute)		
Medial tibiofemoral (per SD)	$-265$ ( $-337, -194$ ) $-300$ ( $-399, -200$ )	
Medial (tibiofemoral) cartilage volume loss (absolute)		
Baseline cartilage volume (site) $\beta$ (95 % CI)		$\beta$ (95 % CI)
	Unadjusted	Adjusted <sup>a</sup>

SD Standard deviation Bold denotes significant results

only (β (95 % CI) =  $-0.15$  ( $-0.29$ ,  $-0.02$ ),  $p = 0.01$ ) in the fully adjusted model.

Sub-group analysis was done to describe the association between baseline cartilage volume stratified by mean volume and absolute cartilage volume loss over 10 years. Both greater than or equal to the mean total tibiofemoral cartilage volume (β (95 % CI) =  $-744$  ( $-1162$ ,  $-325$ )) and less than mean cartilage volume (β (95 % CI) =  $-423$ (−837, −9)) significantly predicted cartilage volume loss in the fully adjusted model.

Table 3 describes the association between the baseline radiographic measures and cartilage volume loss over 10 years. There were significant associations between medial JSN, lateral JSN and osteophyte scores at baseline and compartment specific absolute cartilage volume loss in the unadjusted analysis. However, none of these associations persisted in the multivariable analysis. Similarly there were no significant associations between the baseline radiographic measures and percentage per annum cartilage volume loss in either the unadjusted or the fully adjusted models.

Further sub-analyses looking at the association between baseline cartilage volume and/or ROA and absolute cartilage volume loss stratified by mean age showed no significant differences between the two age groups





<sup>a</sup>Adjusted for age, sex, BMI, offspring-control status, cartilage volume at baseline meniscal tears at visit-2 and meniscal extrusion at baseline and/or JSN and osteophytes at baseline where appropriate SD Standard deviation Bold denotes significant results

except for a significantly stronger association between baseline cartilage volume and cartilage volume loss over 10 years in the lateral compartment only in the older participants. Participants with mean age ≥45 years showed a significant association ( $\beta$  = −405 (−558, −252)) between the baseline lateral tibiofemoral cartilage volume and cartilage volume loss, whereas participants with mean age <45 years showed no significant association  $(\beta = -173 (-387, +41)).$ 

Analyses to explore interactions between the offspring and control groups found no statistically significant difference between the two groups for any of the associations described above. The association between baseline cartilage volume and cartilage volume loss were statistically significant in both the offspring and controls groups when analysed separately (data not shown).

#### **Discussion**

This longitudinal study documents the associations between baseline cartilage volume/ROA and cartilage volume loss over 10 years. Mean absolute and percentage per annum cartilage volume loss was substantial (19.1 % and 13.8 % in the medial and lateral compartments over 10 years) but less than that seen in older populations [8]. Higher baseline tibiofemoral cartilage volume independently predicted greater absolute cartilage volume loss in both compartments and percentage per annum loss in the lateral compartment only. Baseline JSN and osteophytes did not independently predict absolute or percentage per annum cartilage volume loss in either compartment.

This is the first study to describe an independent association between baseline cartilage volume and absolute cartilage volume loss over 10 years. Some recent studies have shown similar associations between baseline cartilage volume and cartilage volume loss over shorter timeframes but none of these studies accounted for knee structural abnormalities such as meniscal tears, meniscal extrusion and BMLs [2, 26, 27]. All of these structures have been shown to predict cartilage volume loss [28] and are potential confounders for the associations described in this study. The association was independent of these factors in the current study. Furthermore, none of the studies mentioned above described the association between the baseline cartilage volume and cartilage volume loss for both tibial and femoral sites.

A criticism of identifying people who will lose more cartilage using the baseline cartilage volume is that association could be due to regression to the mean/tracking. Regression to the mean is a statistical phenomenon that can make natural variation in repeated data look like real change [29]. It happens when unusually large or small

measurements tend to be followed by measurements that are closer to the mean. Unusually high or low cartilage volume to begin with could be due to a number of factors such as cartilage random variation due to body size, sex and co-pathologies. To counter the effect of regression to the mean, further sub-group analysis was done with baseline cartilage volume stratified by the mean value. Both group showed a significant association between the baseline cartilage volume and cartilage volume loss, albeit with a greater effect size in participants with a higher baseline cartilage volume. This suggests that the significant association we described is not solely due to regression to the mean. Similar independent association in the lateral compartment for percentage per annum loss, which also takes into account the cartilage volume to begin with, also suggests that this association is real. However, we did not see any independent associations for medial compartment percentage per annum loss suggesting there is an increase in cartilage volume, due to cartilage swelling, that precedes cartilage volume loss in early OA [7]. Early OA is characterised by matrix changes including a reduction in cellular and proteoglycan content and subsequent water retention and proteoglycan dilution [30]. This depletion of proteoglycan matrix has been closely related to the progression of OA [9]. The swelling of cartilage, in the form of increased volume [9], detected by MRI in early OA has been shown to correlate with depletion proteoglycan matrix and cartilage volume loss, and would explain the associations described in this study.

Few longitudinal studies have looked at the association between baseline ROA and cartilage volume loss and to date, they have shown mixed results. Preliminary crosssectional findings published from this cohort showed that JSN but not osteophytes were associated with a decreased tibial cartilage volume [18]. Similarly Saunders et al. [10] examined the relationship in a randomly selected older cohort over three years and found that JSN and osteophytes both predicted volume loss in a dose response manner but did not adjust for potential confounders such as meniscal tears/extrusion and BMLs. Furthermore, studies looking at the association between JSN and cartilage thickness loss have shown mixed results as well [20, 31], possibly due to different study populations and shorter follow-up periods. Univariable analysis looking at the association between baseline ROA and cartilage volume loss from our 10 year data showed similar results to Saunders et al. [10] but none of these associations persisted once adjusted for MRI assessed co-pathologies. JSN is a composite of structures that are not visible on radiographs. Variation in JSN and longitudinal changes are a reflection of changes in cartilage and meniscus [32]. Hence when adjusted for abnormalities in these structures, JSN did not independently predict cartilage volume loss. Osteophytes are considered an instigating factor in OA causal pathway and studies have shown that presence of osteophytes is associated with a higher prevalence of cartilage defects and decreased cartilage volume. However, once adjusted for co-pathologies, osteophytes failed to independently predict cartilage volume loss in either compartment. Recent studies have suggested that loss of meniscal function is associated with both cartilage volume loss and presence of osteophytes due to increased bio-mechanical stress on the underlying cartilage and the bone. These results and the data from the present study suggest that osteophytes may be on the OA causal pathway or an attempt at repair and are probably not an independent instigating factor for early cartilage volume loss.

Rate of cartilage volume loss and OA progression varies from patient to patient. Cartilage volume loss is often the end-point in chondro-protective drug trials and has been shown to predict total knee replacement surgery [22]. It is imperative for chondro-protective trials to identify fast progressors to make these trials more responsive and sensitive to change. Previous studies have suggested that degree of JSN can be used to identify the sub-groups, which will lose cartilage faster in chondroprotective drug trials [31]. However, data from this study shows that JSN is not an independent predictor of cartilage volume loss. On the other hand, our results suggest that cartilage volume to begin with can be used to identify fast progressors especially if we can differentiate between swollen and non-swollen cartilage.

The key strength of this study is the long follow up period. To our knowledge this study has the longest follow up period using MRI to monitor disease progression in OA. Another strength is that we examined both femoral and tibial cartilage volume loss whereas previous studies have often only reported on one or the other. Lastly, adjustment for other MRI structural copathologies points towards the mediating mechanisms involved in cartilage volume loss. This study has a number of limitations as well. First, around 40 % of participants were lost to follow up at ten years. Those lost to follow-up however were found to be similar in terms of baseline characteristics compared to the participants who were followed-up. Secondly we examined a specific middle-aged group and therefore the results cannot be generalised to the entire population especially people with advanced OA. We believe our results are generalisable to a middle-aged population as we did not see any significant differences between the offspring and control groups for any of the associations described in this study. Third, meniscal tears and BMLs were scored at two years and not at the baseline visit. However changes in these structures over 8 years was small suggesting that these are unlikely to change the effect size considerably.

#### Conclusion

Cross-sectional cartilage volume measurement independently predicts cartilage volume loss over 10 years and can be used to identify fast progressors in clinical trials. Radiographic JSN and osteophytes on the other hand are a reflection of other co-pathologies assessed on MRI and do not independently predict cartilage volume loss over 10 years.

#### Competing interests

Jean-Pierre Pelletier and Johanne Martel Pelletier are shareholders in ArthroLab Inc; the other authors declare no completing interests. All authors have completed the Unified Competing Interest form (available on request from the corresponding author).

#### Authors' contributions

AM was responsible for data collection, preparation of initial manuscript and revisions of the manuscript. HIK was responsible for the analysis and interpretation of data, preparation of initial manuscript and revisions of the manuscript. DA was responsible for data cleaning and management, data interpretation and drafting of the manuscript. LC was responsible for data collection and drafting of the manuscript. LB was responsible for data analysis and drafting of the manuscript. JPP and JMP were responsible for the measurements of femoral cartilage volume, meniscal tears and meniscal extrusion, and drafting of the manuscript. CD, FC, and GJ were responsible for protocol development, data acquisition and drafting of the manuscript. All authors read and approved the final manuscript.

#### Acknowledgements

We thank the subjects who made this study possible, Catrina Boon and Pip Boon for their role in collecting the data and André Pelletier and Josée Thériault for their expertise in MRI reading.

The National Health and Medical Research Council of Australia and Masonic Centenary Medical Research Foundation supported this work. The study sponsor had no role in the design of the study; the collection, analysis, and interpretation of the data; or the writing of the article and the decision to submit it for publication. The researchers work independently of their funder.

#### Author details

<sup>1</sup>Menzies Institute for Medical Research, University of Tasmania, Medical Science 1 Building, Private Bag 23 17-Liverpool Street, Hobart 7000, Australia. <sup>2</sup> Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia. <sup>3</sup>Osteoarthritis Research Unit, University of Montreal Hospital Research Centre (CRCHUM), Montreal, QC, Canada.

#### Received: 28 July 2015 Accepted: 23 January 2016 Published online: 02 February 2016

#### References

- 1. Ding C, Jones G, Wluka AE, Cicuttini F. What can we learn about osteoarthritis by studying a healthy person against a person with early onset of disease? Curr Opin Rheumatol. 2010;22:520–7.
- 2. Antony B, Ding C, Stannus O, Cicuttini F, Jones G. Association of baseline knee bone size, cartilage volume, and body mass index with knee cartilage loss over time: a longitudinal study in younger or middle-aged adults. J Rheumatol. 2011;38:1973–80.
- 3. Jones G, Ding C, Scott F, Cicuttini F. Genetic mechanisms of knee osteoarthritis: a population based case-control study. Ann Rheum Dis. 2004; 63:1255-9.
- Tessier JJ, Bowyer J, Brownrigg NJ, Peers IS, Westwood FR, Waterton JC et al. Characterisation of the guinea pig model of osteoarthritis by in vivo three-dimensional magnetic resonance imaging. Osteoarthritis Cartilage. 2003;11:845-53.
- 5. Wluka AE, Forbes A, Wang Y, Hanna F, Jones G, Cicuttini FM. Knee cartilage loss in symptomatic knee osteoarthritis over 4.5 years. Arthritis Res Ther. 2006;8:R90.
- 6. Pelletier JP, Raynauld JP, Berthiaume MJ, Abram F, Choquette D, Haraoui B et al. Risk factors associated with the loss of cartilage volume on weightbearing areas in knee osteoarthritis patients assessed by quantitative

magnetic resonance imaging: a longitudinal study. Arthritis Res Ther. 2007;9: R74.

- 7. Ding C, Cicuttini F, Blizzard L, Scott F, Jones G. A longitudinal study of the effect of sex and age on rate of change in knee cartilage volume in adults. Rheumatology (Oxford). 2007;46:273-9.
- 8. Calvo E, Palacios I, Delgado E, Sanchez-Pernaute O, Largo R, Egido J et al. Histopathological correlation of cartilage swelling detected by magnetic resonance imaging in early experimental osteoarthritis. Osteoarthritis Cartilage. 2004;12:878-86.
- 9. Eckstein F, Le Graverand MP, Charles HC, Hunter DJ, Kraus VB, Sunyer T et al. Clinical, radiographic, molecular and MRI-based predictors of cartilage loss in knee osteoarthritis. Ann Rheum Dis. 2011;70:1223-30.
- 10. Saunders J, Ding C, Cicuttini F, Jones G. Radiographic osteoarthritis and pain are independent predictors of knee cartilage loss: a prospective study. Intern Med J. 2012;42:274-80.
- 11. Eckstein F, Nevitt M, Gimona A, Picha K, Lee JH, Davies RY et al. Rates of change and sensitivity to change in cartilage morphology in healthy knees and in knees with mild, moderate, and end-stage radiographic osteoarthritis: results from 831 participants from the Osteoarthritis Initiative. Arthritis Care Res (Hoboken). 2011;63:311-9.
- 12. Eckstein F, Wirth W, Hudelmaier MI, Maschek S, Hitzl W, Wyman BT et al. Relationship of compartment-specific structural knee status at baseline with change in cartilage morphology: a prospective observational study using data from the osteoarthritis initiative. Arthritis Res Ther. 2009;11:R90.
- 13. Jones G, Glisson M, Hynes K, Cicuttini F. Sex and site differences in cartilage development: a possible explanation for variations in knee osteoarthritis in later life. Arthritis Rheum. 2000;43:2543-9.
- 14. Ding C, Cicuttini F, Scott F, Boon C, Jones G. Association of prevalent and incident knee cartilage defects with loss of tibial and patellar cartilage: a longitudinal study. Arthritis Rheum. 2005;52:3918-27.
- 15. Ding C, Cicuttini F, Scott F, Cooley H, Boon C, Jones G. Natural history of knee cartilage defects and factors affecting change. Arch Intern Med. 2006; 166:651-8.
- 16. Ding C, Cicuttini F, Blizzard L, Jones G. Smoking interacts with family history with regard to change in knee cartilage volume and cartilage defect development. Arthritis Rheum. 2007;56:1521-8.
- 17. Jones G, Ding C, Scott F, Glisson M, Cicuttini F. Early radiographic osteoarthritis is associated with substantial changes in cartilage volume and tibial bone surface area in both males and females. Osteoarthritis Cartilage. 2004;12:169-74.
- 18. Berthiaume MJ, Raynauld JP, Martel-Pelletier J, Labonté F, Beaudoin G, Bloch DA et al. Meniscal tear and extrusion are strongly associated with progression of symptomatic knee osteoarthritis as assessed by quantitative magnetic resonance imaging. Ann Rheum Dis 2005;64:556-63.
- 19. Raynauld JP, Martel-Pelletier J, Berthiaume MJ, Beaudoin G, Choquette D, Haraoui B et al. Long term evaluation of disease progression through the quantitative magnetic resonance imaging of symptomatic knee osteoarthritis patients: correlation with clinical symptoms and radiographic changes. Arthritis Res Ther. 2006;8:R21.
- 20. Dore D, Quinn S, Ding C, Winzenberg T, Zhai G, Cicuttini F et al. Natural history and clinical significance of MRI-detected bone marrow lesions at the knee: a prospective study in community dwelling older adults. Arthritis Res Ther. 2010;12:R223.
- 21. Pelletier JP, Cooper C, Peterfy C, Reginster JY, Brandi ML, Bruyere O et al. What is the predictive value of MRI for the occurrence of knee replacement surgery in knee osteoarthritis? Ann Rheum Dis. 2013;72:1594-604.
- 22. Altman RD, Hochberg M, Murphy WA, Jr., Wolfe F, Lequesne M. Atlas of individual radiographic features in osteoarthritis. Osteoarthritis Cartilage. 1995;3(Suppl A):3-70.
- 23. Aitken D, Ding C, Pelletier JP, Martel-Pelletier J, Cicuttini F, Jones G. Responsiveness of magnetic resonance imaging-derived measures over 2.7 years. J Rheumatol. 2014;41:2060-7.
- 24. Dore D, de Hoog J, Giles G, Ding C, Cicuttini F, Jones G. A longitudinal study of the association between dietary factors, serum lipids, and bone marrow lesions of the knee. Arthritis Res Ther. 2012;14:R13.
- 25. Ding C, Martel-Pelletier J, Pelletier JP, Abram F, Raynauld JP, Cicuttini F et al. Two-year prospective longitudinal study exploring the factors associated with change in femoral cartilage volume in a cohort largely without knee radiographic osteoarthritis. Osteoarthritis Cartilage. 2008;16:443-9.
- 26. Wluka AE, Stuckey S, Snaddon J, Cicuttini FM. The determinants of change in tibial cartilage volume in osteoarthritic knees. Arthritis Rheum. 2002;46: 2065-72.
- 27. Chuang-Stein C, Tong DM. The impact and implication of regression to the mean on the design and analysis of medical investigations. Stat Methods Med Res. 1997;6:115-28.
- 28. Ding C, Cicuttini F, Jones G. How important is MRI for detecting early osteoarthritis? Nat Clin Pract Rheumatol. 2008;4:4-5.
- 29. Eckstein F, Benichou O, Wirth W, Nelson DR, Maschek S, Hudelmaier M et al. Magnetic resonance imaging-based cartilage loss in painful contralateral knees with and without radiographic joint space narrowing: Data from the Osteoarthritis Initiative. Arthritis Rheum. 2009;61:1218-25.
- 30. Hunter DJ, Zhang YQ, Tu X, Lavalley M, Niu JB, Amin S et al. Change in joint space width: hyaline articular cartilage loss or alteration in meniscus? Arthritis Rheum. 2006;54:2488-95.
- 31. Ding C, Garnero P, Cicuttini F, Scott F, Cooley H, Jones G. Knee cartilage defects: association with early radiographic osteoarthritis, decreased cartilage volume, increased joint surface area and type II collagen breakdown. Osteoarthritis Cartilage. 2005; 13:198-205.
- 32. Beattie KA, Boulos P, Pui M, O'Neill J, Inglis D, Webber CE et al. Abnormalities identified in the knees of asymptomatic volunteers using peripheral magnetic resonance imaging. Osteoarthritis Cartilage. 2005;13: 181-6.

# Submit your next manuscript to BioMed Central and we will help you at every step:

- **•** We accept pre-submission inquiries
- **•** Our selector tool helps you to find the most relevant journal
- We provide round the clock customer support
- Convenient online submission
- **•** Thorough peer review
- Inclusion in PubMed and all major indexing services
- **•** Maximum visibility for your research

Submit your manuscript at www.biomedcentral.com/submit



# Content courtesy of Springer Nature, terms of use apply. Rights reserved.

### Terms and Conditions Ĩ

Springer Nature journal content, brought to you courtesy of Springer Nature Customer Service Center GmbH ("Springer Nature").

Springer Nature supports a reasonable amount of sharing of research papers by authors, subscribers and authorised users ("Users"), for smallscale personal, non-commercial use provided that all copyright, trade and service marks and other proprietary notices are maintained. By accessing, sharing, receiving or otherwise using the Springer Nature journal content you agree to these terms of use ("Terms"). For these purposes, Springer Nature considers academic use (by researchers and students) to be non-commercial.

These Terms are supplementary and will apply in addition to any applicable website terms and conditions, a relevant site licence or a personal subscription. These Terms will prevail over any conflict or ambiguity with regards to the relevant terms, a site licence or a personal subscription (to the extent of the conflict or ambiguity only). For Creative Commons-licensed articles, the terms of the Creative Commons license used will apply.

We collect and use personal data to provide access to the Springer Nature journal content. We may also use these personal data internally within ResearchGate and Springer Nature and as agreed share it, in an anonymised way, for purposes of tracking, analysis and reporting. We will not otherwise disclose your personal data outside the ResearchGate or the Springer Nature group of companies unless we have your permission as detailed in the Privacy Policy.

While Users may use the Springer Nature journal content for small scale, personal non-commercial use, it is important to note that Users may not:

- 1. use such content for the purpose of providing other users with access on a regular or large scale basis or as a means to circumvent access control;
- 2. use such content where to do so would be considered a criminal or statutory offence in any jurisdiction, or gives rise to civil liability, or is otherwise unlawful;
- 3. falsely or misleadingly imply or suggest endorsement, approval , sponsorship, or association unless explicitly agreed to by Springer Nature in writing;
- 4. use bots or other automated methods to access the content or redirect messages
- 5. override any security feature or exclusionary protocol; or
- 6. share the content in order to create substitute for Springer Nature products or services or a systematic database of Springer Nature journal content.

In line with the restriction against commercial use, Springer Nature does not permit the creation of a product or service that creates revenue, royalties, rent or income from our content or its inclusion as part of a paid for service or for other commercial gain. Springer Nature journal content cannot be used for inter-library loans and librarians may not upload Springer Nature journal content on a large scale into their, or any other, institutional repository.

These terms of use are reviewed regularly and may be amended at any time. Springer Nature is not obligated to publish any information or content on this website and may remove it or features or functionality at our sole discretion, at any time with or without notice. Springer Nature may revoke this licence to you at any time and remove access to any copies of the Springer Nature journal content which have been saved.

To the fullest extent permitted by law, Springer Nature makes no warranties, representations or guarantees to Users, either express or implied with respect to the Springer nature journal content and all parties disclaim and waive any implied warranties or warranties imposed by law, including merchantability or fitness for any particular purpose.

Please note that these rights do not automatically extend to content, data or other material published by Springer Nature that may be licensed from third parties.

If you would like to use or distribute our Springer Nature journal content to a wider audience or on a regular basis or in any other manner not expressly permitted by these Terms, please contact Springer Nature at

[onlineservice@springernature.com](mailto:onlineservice@springernature.com)