The relationship between foot and ankle symptoms and risk of developing knee osteoarthritis: data from the osteoarthritis initiative

Kade L. Paterson, PhD, Jessica Kasza, PhD, David J. Hunter, PhD, Rana S. Hinman, PhD, Hylton B. Menz, PhD, George Peat, PhD, Kim L. Bennell, PhD

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The relationship between foot and ankle symptoms and risk of developing knee osteoarthritis: data from the osteoarthritis initiative

AUTHORS: Kade L Paterson PhD (kade.paterson@unimelb.edu.au)\textsuperscript{1}, Jessica Kasza PhD (jessica.kasza@monash.edu)\textsuperscript{2}, David J Hunter PhD (david.hunter@sydney.edu.au)\textsuperscript{3}, Rana S Hinman PhD (ranash@unimelb.edu.au)\textsuperscript{1}, Hylton B Menz PhD (H.Menz@latrobe.edu.au)\textsuperscript{4}, George Peat PhD (g.m.peat@keele.ac.uk)\textsuperscript{5}, & Kim L Bennell PhD (k.bennell@unimelb.edu.au)\textsuperscript{1}

AFFILIATIONS: \textsuperscript{1} Centre for Health, Exercise and Sports Medicine, The University of Melbourne, Melbourne, Australia, \textsuperscript{2} Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia, \textsuperscript{3} Institute of Bone and Joint Research, Kolling Institute, University of Sydney, and Rheumatology Department, Royal North Shore Hospital Australia, Sydney, Australia, \textsuperscript{4} School of Allied Health, La Trobe University, Melbourne, Australia, \textsuperscript{5} Arthritis Research UK Primary Care Centre, Keele University, Keele, United Kingdom.

CORRESPONDING AUTHOR: Kade Paterson, Centre for Health Exercise and Sports Medicine, Department of Physiotherapy, School of Health Sciences, University of Melbourne, Parkville, Victoria, Australia 3010. ph: +61 3 8344 0425, fax: +61 3 8344 4188, kade.paterson@unimelb.edu.au
Running headline Foot symptoms and knee OA
ABSTRACT

Objective To investigate whether foot and/or ankle symptoms increase the risk of developing (i) knee symptoms and (ii) symptomatic radiographic knee osteoarthritis (OA).

Design 1020 Osteoarthritis Initiative participants who were at-risk of knee OA, but were without knee symptoms or radiographic knee OA, were investigated. Participants indicated the presence and laterality of foot/ankle symptoms at baseline. The main outcome was development of knee symptoms (pain, aching or stiffness in and around the knee on most days of the month for at least one month in the past year). A secondary outcome was development of symptomatic radiographic knee OA (symptoms plus Kellgren and Lawrence [KL] grade ≥2), over the subsequent four years. Associations between foot/ankle symptoms and study outcomes were assessed by logistic regression models.

Results Foot/ankle symptoms in either or both feet significantly increased the odds of developing knee symptoms (adjusted odds ratio (OR) 1.55, 95% confidence interval (CI) 1.10 to 2.19), and developing symptomatic radiographic knee OA (adjusted OR 3.28, 95% CI 1.69 to 6.37). Based on laterality, contralateral foot/ankle symptoms were associated with developing both knee symptoms (adjusted OR 1.68, 95% CI 1.05 to 2.68) and symptomatic radiographic knee OA (adjusted OR 3.08, 95% CI 1.06 to 8.98), whilst bilateral foot/ankle symptoms were associated with developing symptomatic radiographic knee OA (adjusted OR 4.02, 95% CI 1.76 to 9.17).

Conclusion In individuals at-risk of knee OA, the presence of contralateral foot/ankle symptoms in particular increases risk of developing both knee symptoms and symptomatic radiographic knee OA.
Foot symptoms and knee OA

Key words: Osteoarthritis, Knee Osteoarthritis, Arthritis, Epidemiology
INTRODUCTION

Knee osteoarthritis (OA) is a leading cause of joint pain and disability in middle- and older-aged individuals, and is one of the most commonly managed conditions in primary care. Recent incidence rates suggest around 6% of people aged over 45 years develop knee symptoms each year, whilst 2% develop symptomatic radiographic knee OA. Knee OA symptoms and radiographic change that worsen over time can lead to costly surgical intervention. Thus understanding risk factors associated with the onset of knee symptoms alone or in combination with structural change is a major research focus.

Symptoms in the foot and/or ankle is a potential risk factor for knee pain and OA that has received limited attention to date. Like knee OA, foot/ankle symptoms are very common in middle- and older-aged adults. They affect approximately 24% of people aged over 45 years, and account for a substantial number of primary care consultations in this population. Foot pain is highly disabling, reduces quality of life, adversely affects walking and other daily functional abilities and increases the risk of falls. To date, the majority of studies investigating symptoms at the foot/ankle and knee have examined these problems in isolation. However, isolated joint pain is rare, and concurrent symptoms at the foot/ankle and knee is the most common multi-joint presentation, occurring far greater than expected by chance alone. In a recent cross-sectional study using data from the Osteoarthritis Initiative (OAI), we found that people with both symptomatic radiographic knee OA and foot/ankle symptoms reported significantly worse general and knee OA specific health outcomes, and poorer physical function, than those with
Foot symptoms and knee OA

knee OA but without foot/ankle symptoms\textsuperscript{11}. Despite the strong association between problems at these two sites, their temporal sequence has not yet been evaluated.

Investigating foot/ankle symptoms as a candidate risk factor for knee OA is attractive as it is simple to assess, and there is some evidence of potential modifiability using simple low-cost interventions such as off-the-shelf footwear\textsuperscript{12}. Furthermore, there are a number of plausible biological mechanisms linking foot/ankle symptoms to knee OA development. For example, there may be shared biomechanical risk factors for the two problems, such as a pronated foot type\textsuperscript{13} or inappropriate footwear\textsuperscript{14}. Alternatively, people with foot/ankle symptoms may walk differently to offload their painful foot\textsuperscript{15-17}, altering knee function and increasing the risk of knee OA development. Finally, symptoms at these two sites may represent a widespread pain phenotype or an oligo- or polyarticular form of OA\textsuperscript{18}.

The primary aim of this study was to use longitudinal data from the OAI to examine whether foot/ankle symptoms predict the development of knee symptoms over four years in people without knee symptoms or radiographic knee OA, but at-risk of knee OA, at baseline. A secondary aim was to examine whether foot/ankle symptoms also predict the development of symptomatic radiographic knee OA over four years. It was hypothesized that foot/ankle symptoms would increase the odds of developing knee symptoms and symptomatic radiographic knee OA in people at risk of knee OA.

**METHODS**
Foot symptoms and knee OA

Study population

The OAI is an ongoing prospective multicentre cohort study designed to evaluate and identify biomarkers for the onset and/or progression of knee OA in people aged between 45-79 years. The study enrolled 4796 men and women from four sites in the United States, including Baltimore, Maryland; Columbus, Ohio; Pittsburgh, Pennsylvania; and Pawtucket, Rhode Island. All protocols and procedures were approved by the institutional review board at each site and all participants provided informed consent. Details regarding general exclusion criteria and the wider study protocols are available online for public access (http://www.oai.ucsf.edu/). In the current study, we analyzed OAI participants who were at risk of knee OA, defined as the presence of two or more established characteristics including: overweight, identified using age- and sex-specific criteria; a history of knee injury causing walking difficulties; any knee surgery; an immediate family history of a total knee replacement for OA; Heberden’s nodes; repetitive knee bending during occupational or recreation activities; or aged between 70-79 years. From this subcohort, we only included people who did not have frequent knee symptoms (defined as pain, aching or stiffness in and around the knee on most days of the month for at least one month in the past year) or radiographic evidence of knee OA (Kellgren and Lawrence [KL] grade ≥2) in either knee at baseline. We excluded people (rather than knees) with these outcomes because the presence of symptomatic knee OA in one knee greatly increases the risk of developing contralateral knee OA which may confound results.20-22 Demographic, clinical and radiographic characteristics of both knees for all participants were evaluated at baseline and at 12, 24, 36 and 48-month follow-up visits.
Demographic characteristics and covariates

Demographic data collected included age, sex and race (White, Black/African American or Asian/other non-white). Covariates included body mass index (BMI), comorbidities and depression. As well as recording BMI values, we also classified participants as obese (>30 kg/m²), overweight (≥25 and ≤30 kg/m²) or normal weight (<25 kg/m²). Comorbidities were assessed using the questionnaire version of the Charlson comorbidity index (CCI), and we dichotomized the cohort into those with ‘no comorbidities’ and those with ‘one or more comorbidities’. Depression was measured using the Centre for Epidemiological Studies Depression Scale (CES-D). Scores were summed and a score of ≥ 16 was used to indicate significant depressive symptoms.

Risk factor

The primary risk factor was self-reported foot/ankle symptoms at baseline, defined as pain, aching or stiffness in the foot and/or ankle on more than half of the days during the past 30 days, consistent with definitions used in previous studies. In addition to classifying participants based on the presence or absence of symptoms in either foot/ankle, we further stratified foot/ankle symptoms as ipsilateral, contralateral or bilateral relative to each knee.

Incidence outcomes
Foot symptoms and knee OA

Knee symptoms

Participants were asked about the presence of knee symptoms at baseline, and at the 12, 24, 36 and 48 month follow-up visits for each knee. Incident knee symptoms was defined as development of pain, aching or stiffness in and around the knee on most days of the month for at least one month in the previous year, reported at any of the follow up visits, consistent with the OAI definition and based on American College of Rheumatology criteria for clinical knee OA\textsuperscript{25}.

Symptomatic radiographic knee osteoarthritis

Weightbearing fixed-flexion posteroanterior radiographs of both knees were taken at baseline and at the 12, 24, 36 and 48 month follow-up visits. Radiographs were evaluated using the KL grading system (grades range 0-4) by two central OAI senior musculoskeletal experts blinded to all other participant data and to each other’s readings. Incident symptomatic radiographic knee OA was defined as knee symptoms (as per definition above) and the presence of KL grade $\geq 2$ based on the central OAI reading, at any of the follow up visits.

Statistical analysis

Baseline characteristics of participants with and without foot/ankle symptoms were summarised as number (%) for categorical variables and as mean (SD) or median (interquartile range) for continuous variables, as appropriate. Groups were compared using $\chi$-squared tests, analysis of variance, Wilcoxon rank-sum or Kruskal-Wallis rank tests respectively.
Foot symptoms and knee OA

To investigate the primary aim (development of knee symptoms), we analysed the association between any foot/ankle symptoms (i.e. symptoms in either or both feet/ankle) at baseline and the development of knee symptoms at any point within the four year follow-up period. For both aims, analyses were knee-specific (i.e. conducted at the knee level rather than at the participant level). Since most participants contributed two knees (8 participants with missing data contributed one knee only for the primary aim, and 3 participants with missing data contributed one knee only for the secondary aim), logistic regression models were fitted using generalized estimating equations to account for the correlation between left and right knees within participants. Two models were fitted, adjusting for sets of baseline covariates determined \textit{a priori}. In the first model, only baseline foot/ankle symptoms were included to obtain unadjusted associations between baseline foot/ankle symptoms and the development of outcomes. The second model also included age, sex, race, BMI, Charlson Comorbidity index (dichotomised) and depression to adjust for variables known to be associated with both foot pain \textsuperscript{26} and knee OA \textsuperscript{27}.

In addition to considering whether any foot/ankle symptoms were associated with the outcome (i.e. at the participant level), we also investigated the association with ipsilateral, contralateral or bilateral foot/ankle symptoms (i.e. at the limb level) to see if the association differed by laterality. Logistic regression models were again fitted using generalized estimating equations to adjust for clustering of knees within participants. Covariates were adjusted for in the same way as in the primary analysis. Similar analyses were conducted to address the secondary aim (the development of symptomatic radiographic knee OA), and the set of baseline variables was adjusted as per for the primary aim.
To assess the potential influence of confounders (both measured and unmeasured by the OAI) that were not accounted for in our analyses, we performed sensitivity analyses. More specifically, a causal inference-based approach adapted from Kasza et al\textsuperscript{28} was used. This approach varies a sensitivity parameter that quantifies the differences between participants with and without foot/ankle symptoms, had those without foot/ankle symptoms instead had symptoms. The sensitivity parameter compares the outcomes between two groups with the same exposure (where the exposure is hypothetical in those without foot/ankle symptoms), but with the possibility of differences that were unaccounted for in our analyses leading to differences in the development of knee symptoms and/or symptomatic radiographic knee OA. Values of the sensitivity parameter greater than 1 suggest that unaccounted confounders in those participants who actually had foot/ankle symptoms, such as widespread pain or generalised OA, contributed to the greater likelihood of those participants developing the outcome. Values of the sensitivity parameter equal to 1 suggest that there is no impact of unaccounted-for confounding on the results. Statistical significance was ascribed at p-value $\leq 0.05$. Stata v12 (Stata Corporation, College Station, TX, USA) was used for all analyses.

RESULTS

Sample characteristics

This study used OAI participants who did not have symptomatic radiographic knee OA (n=3306). Patients with knee symptoms (as defined previously) or radiographic knee OA (KL$\geq$2)
in one or both knees at baseline were excluded (n=2286) (Figure 1). Demographic data are presented in Table 1. Of the 1020 participants at baseline, 13% (n=133) reported symptoms in at least one foot/ankle. Those with foot/ankle symptoms were more likely to be female (p=0.014), younger (p=0.029), Black/African American (p<0.001) and have a higher BMI (p=0.003) at baseline. There were no differences in baseline measures of worst KL grade, comorbidities, depressive symptoms, shoulder pain, Heberden’s nodes, or previous knee injury or surgery between those with and without foot/ankle symptoms.

**Development of knee symptoms**

Table 2 shows the odds of developing knee symptoms according to the presence and laterality of foot/ankle symptoms. After excluding knees with missing data, there were 1990 knees from 999 participants available for analysis. Baseline symptoms in any foot/ankle was associated with a significantly increased risk of developing knee symptoms in the subsequent four years (adjusted OR 1.55, 95% CI 1.10 to 2.19). Additional analyses of foot/ankle and knee symptom laterality showed that contralateral foot/ankle symptoms also increased the odds for developing knee symptoms (adjusted OR 1.68, 95% CI 1.05 to 2.68).
Development of symptomatic radiographic knee OA

Table 3 shows the odds of developing symptomatic radiographic knee OA according to the presence and laterality of foot/ankle symptoms. After excluding knees with missing data, there were 1983 knees from 993 people available for analysis. Baseline symptoms in any foot/ankle was associated with a significantly increased risk of developing symptomatic radiographic knee OA at any time in the follow up period (adjusted OR 3.28, 95% CI 1.69 to 6.37). Subgroup analyses based on foot/ankle symptom laterality suggested bilateral foot/ankle symptoms had the highest odds for developing symptomatic radiographic knee OA (adjusted OR 4.02, 95% CI 1.76 to 9.17), and that foot/ankle symptoms that were contralateral to the affected knee also increased the risk of this outcome (adjusted OR 3.08, 95% CI 1.60 to 8.98).

Sensitivity analyses

The results of our sensitivity analyses suggest that it is highly unlikely that any confounder not included in our analyses would have explained the observed association between foot/ankle symptoms and the development of knee symptoms and symptomatic radiographic knee OA. Specifically, the sensitivity analysis for developing knee symptoms (Figure 2) indicates that when the sensitivity parameter is about 1.3, the odds ratio reduces to 1. Hence, for the association to be entirely explained by unaccounted-for confounding, those with foot/ankle
Foot symptoms and knee OA

symptoms would need to be 30% more likely to develop the outcome than those without symptoms would be had they also had foot/ankle symptoms. The sensitivity parameter required to explain the association between foot/ankle symptoms and symptomatic radiographic knee OA is even greater (Figure 3): those with foot/ankle symptoms need to be more than twice as likely to develop the outcome than those without symptoms would be had they also had foot/ankle symptoms. Figures 2 and 3 indicate as the value of the sensitivity parameter gets greater (corresponding to the greater tendency to develop the outcome among those with foot/ankle symptoms), the sensitivity parameter-adjusted OR is further reduced.

DISCUSSION

This is the first study to investigate whether foot/ankle symptoms are a risk factor for the development of knee symptoms and symptomatic radiographic knee OA in people at-risk of the disease. Foot/ankle symptoms in either or both sides were found to increase the risk of developing knee symptoms over the subsequent four years, with contralateral foot/ankle symptoms the only side to show an association with knee symptom development in the laterality analysis. Foot/ankle symptoms in either or both sides were also found to increase the risk of developing symptomatic radiographic knee OA, with bilateral and contralateral foot/ankle symptoms both associated, however there were few cases who developed this outcome and confidence intervals were wide. These findings add to previous cross-sectional studies demonstrating strong associations between symptoms at the foot/ankle and knee\textsuperscript{10,11}, and they
Foot symptoms and knee OA

provide the first longitudinal evidence that foot/ankle symptoms are a risk factor for the development of knee symptoms, and symptomatic radiographic knee OA.

Few studies have investigated risk factors for the onset of knee symptoms. A large prospective cohort study previously identified previous knee injury as the strongest predictor of onset of future knee pain with similar odds ratios to ours (1.59 compared to 1.60). Furthermore, although a number of other risk factors for the development of symptomatic radiographic knee OA have been previously reported, such as age and ethnicity, few are modifiable. Currently, the strongest known modifiable risk factors for developing knee OA are obesity and previous knee injury. Our odds of around 3.3 for developing symptomatic radiographic knee OA are also comparable to these other potentially modifiable factors (pooled OR 2.6 for BMI and 3.9 for knee injury). However, some caution should be used when interpreting the outcomes of our symptomatic radiographic model. Firstly, despite our large cohort with several years' follow-up and our use of knee-level data, few cases developed symptomatic radiographic knee OA. This reduces the precision of the odds ratio for this model, as seen by the wide confidence intervals. With so few cases, and adjustment for six covariates, there is also some risk of over-fitting our regression models. However, our number of events per variable in the model (including covariates) was within recommendations. Finally, our four year follow up may be too short to appropriately evaluate symptomatic radiographic outcomes. However, the OAI only has biennial radiographic data available after four years, and we felt that it would overly complicate our outcome definition to have annual outcomes up to four years and biennial data thereafter. Notwithstanding these points, the results for all of our models were broadly consistent which suggests that it is likely that there is some association between foot/ankle symptoms and
Foot symptoms and knee OA

Symptomatic radiographic knee OA. The findings are also reasonably robust given our sensitivity analyses showed that it is unlikely that our conclusions would be changed had we adjusted for other confounders not included in our analyses.

There are several plausible mechanisms by which foot pain could be linked to the subsequent onset of knee symptoms in people at-risk of knee OA. First, people with foot/ankle symptoms alter their walking pattern and these biomechanical changes may increase the risk of developing knee OA. To date, the effects of foot/ankle symptoms on biomechanics relevant to knee OA have not been explored, however our findings of an association between contralateral but not ipsilateral foot symptoms suggest people with foot/ankle symptoms may shift weight away from the painful foot and increase load on the contralateral knee. Second, it has been suggested that a more pronated or “flatter” foot, which is associated with many painful foot conditions, may increase rotational stress on the tibiofemoral joint, due in part to the tight coupling between movement at the rearfoot and tibia. Over time, this abnormal stress may damage the load-bearing tissues in the knee joint leading to pain and structural damage. However, whilst some cross-sectional studies show increased foot pronation in people with knee OA, and that a more pronated foot is associated with an increased prevalence of knee pain and medial tibiofemoral cartilage damage, other research suggests increased pronation may instead be a compensatory mechanism designed to reduce knee load and pain. Third, footwear may be a shared risk factor for both foot/ankle symptoms and knee OA. For example, inappropriate footwear is a risk factor for foot/ankle symptoms, and some types of footwear such as high heels may also alter knee biomechanics in a detrimental manner. Other researchers have suggested that pain in multiple joints in people with knee OA may reflect a more generalized (e.g. oligo- or
Foot symptoms and knee OA

polyarticular) OA presentation\textsuperscript{39} or a widespread pain phenotype\textsuperscript{18}, partly due to changes in central pain processing\textsuperscript{40, 41}. These central changes may lead to a generalized hypersensitivity to pain and therefore a greater likelihood of developing pain at multiple sites such as the knee and foot. However this does not appear to be explanatory in our findings given our conclusions remained unchanged after we performed sensitivity analyses to account for unaccounted-for confounders.

There is evidence that foot/ankle symptoms may be modifiable given studies have shown simple and relatively inexpensive conservative interventions are effective at treating common causes of foot/ankle symptoms. For example, off-the-shelf footwear was reported to improve general foot pain in older people\textsuperscript{12} and foot pain due to gout\textsuperscript{42}, whilst foot orthoses have been shown to improve pain and function in people with plantar fasciitis\textsuperscript{43}, pes cavus\textsuperscript{44} and rheumatoid arthritis\textsuperscript{45}, amongst others. If the mechanism underpinning the association between foot/ankle symptoms and the development of knee symptoms is due to shifting weight to the contralateral limb to unload the painful foot/ankle, then simple analgesic interventions may also be helpful in reducing the need for this avoidance strategy. Further studies are now required to determine whether treating foot/ankle symptoms using conservative interventions also helps to reduce the incidence of knee pain and symptomatic radiographic knee OA in people at risk of the disease.

There are some limitations to our study. Firstly, although we found a relationship between foot/ankle symptoms and the development of symptomatic and radiographic knee OA, it cannot be determined whether this is an independent relationship to structural or radiographic knee OA.
as these participants are a subset of those who developed knee symptoms. The relationship between foot/ankle symptoms and the development of radiographic knee OA alone was not explored given that radiographic OA without symptoms is not clinically relevant. Second, participants were required to have reported knee symptoms at only one of the follow up visits similar to previous research\(^46\), thus it is possible our analyses included people whose knee symptoms were not sustained over time. We feel that this was appropriate given OAI data have shown knee pain profiles are stable over 6 years\(^47\). However, future studies may wish to examine whether foot/ankle symptoms are associated with more sustained knee pain. Third, we dichotomised BMI and the Charlson comorbidity index which can leave residual confounding\(^48\). However, we found no strong evidence of this when we re-ran the analyses using fractional polynomials to model the continuous scores for these covariates (see Table 1 in the supplementary analyses). It is also possible that our results were biased due to the exclusion of participants because of missing x-rays. However when we compared demographic characteristics and covariates between those with missing and non-missing x-rays, our results showed those with missing x-rays were more similar to OAI participants with KL≥2 at baseline than to those with KL grade 0 and 1 (see Table 2 in the supplementary analyses). Since those with KL≥2 were excluded from the study, it is possible that the participants with missing data would have been excluded regardless. Thus the impact of missing data on our outcomes is likely to be minimal. Finally, we tested a cohort who was already at an increased risk of developing knee OA and thus our results should not be generalised to the wider population. Further research is needed to determine whether foot/ankle symptoms also increase the risk of developing knee symptoms in a population that does not possess other knee OA risk factors.
In conclusion, our study showed that people with foot/ankle symptoms were at an increased risk of developing knee OA symptoms and symptomatic radiographic knee OA compared to those without foot/ankle symptoms. These findings have important clinical and research implications. Although it is unclear whether foot/ankle symptoms directly causes knee symptoms and radiographic changes, or whether its presence is an indirect clinical marker for another variable, our results have identified a potentially modifiable risk factor for knee OA in people at-risk of the disease. Future studies should now determine whether addressing foot/ankle symptoms using conservative interventions reduces the incidence of knee pain and symptomatic radiographic knee OA.

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AUTHOR CONTRIBUTIONS

All authors were involved in conception and design of the study, or in acquisition analysis and interpretation of data, and in revising it critically for important intellectual content. All authors approved the final version to be published. Dr. Paterson takes responsibility for the integrity of the work as a whole, from inception to finished article.

Conception and design. Paterson, Kasza, Hinman, Hunter, Bennell.

Analysis and interpretation of data. Paterson, Kasza, Hunter, Hinman, Menz, Peat, Bennell.
Foot symptoms and knee OA

Drafting of the article. Paterson, Bennell.

Critical revision of the article for important intellectual content. Paterson, Kasza, Hunter, Hinman, Menz, Peat, Bennell.

Final approval of the article. Paterson, Kasza, Hunter, Hinman, Menz, Peat, Bennell.

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CONFLICT OF INTEREST

No authors report competing interests.

RSH and KLB, and the University of Melbourne, received royalties from sales of Gel Melbourne OA shoes from 2012-2014. The manufacturer of the shoes played no role in the study design nor had any input into the analysis and interpretation of data from this study.
Foot symptoms and knee OA

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Foot symptoms and knee OA


Foot symptoms and knee OA


Foot symptoms and knee OA


Foot symptoms and knee OA


Foot symptoms and knee OA


FIGURE LEGENDS

Figure 1. Participants from the Osteoarthritis Initiative included in analysis.

Figure 2. Results of the sensitivity analysis for developing knee symptoms.

Figure 3. Results of the sensitivity analysis for developing symptomatic radiographic knee OA.
Foot symptoms and knee OA

TABLES

Table 1. Baseline characteristics of Osteoarthritis Initiative participants without knee pain classified based on the presence and side of foot/ankle symptoms. One participant had missing foot/ankle symptom status at baseline. Values are N (%) unless otherwise indicated.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Missing (n)</th>
<th>No foot/ankle symptoms (n=887)</th>
<th>Any foot/ankle symptoms (n=133)</th>
<th>P value&lt;sup&gt;†&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) age (years)</td>
<td>0</td>
<td>60.9 (9.1)</td>
<td>59.0 (9.3)</td>
<td>0.029</td>
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<td></td>
<td></td>
<td>0.014</td>
</tr>
<tr>
<td>Male</td>
<td></td>
<td>387 (43.6)</td>
<td>43 (32.3)</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td></td>
<td>500 (56.4)</td>
<td>90 (67.7)</td>
<td></td>
</tr>
<tr>
<td>Race:</td>
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<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Asian and other non-white</td>
<td></td>
<td>14 (1.6)</td>
<td>10 (7.5)</td>
<td></td>
</tr>
<tr>
<td>White/Caucasian</td>
<td></td>
<td>793 (89.4)</td>
<td>107 (80.5)</td>
<td></td>
</tr>
<tr>
<td>Black/African American</td>
<td></td>
<td>80 (9.0)</td>
<td>16 (12.0)</td>
<td></td>
</tr>
<tr>
<td>Median (IQR) BMI kg/m&lt;sup&gt;2&lt;/sup&gt;</td>
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<td>26.4 (23.7, 30.0)</td>
<td>27.7 (24.8, 32.0)</td>
<td>0.003</td>
</tr>
<tr>
<td>BMI categories:</td>
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<td></td>
<td></td>
<td>0.018</td>
</tr>
<tr>
<td>Normal (BMI &lt;25 kg/m&lt;sup&gt;2&lt;/sup&gt;)</td>
<td></td>
<td>317 (35.7)</td>
<td>34 (25.6)</td>
<td></td>
</tr>
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</table>
## Foot symptoms and knee OA

<table>
<thead>
<tr>
<th>Comorbidity</th>
<th>Overweight (BMI 25-30 kg/m²)</th>
<th>Obese (BMI &gt;30 kg/m²)</th>
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<tr>
<td>Overweight</td>
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<td>52 (39.1)</td>
</tr>
<tr>
<td>Obese (BMI &gt;30 kg/m²)</td>
<td>223 (25.1)</td>
<td>47 (35.3)</td>
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<tr>
<td>Worst KL grade *</td>
<td>0</td>
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<tr>
<td>0</td>
<td>568 (64.0)</td>
<td>84 (63.2)</td>
</tr>
<tr>
<td>1</td>
<td>319 (36.0)</td>
<td>49 (36.8)</td>
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<tr>
<td>4</td>
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<td>0 (0)</td>
</tr>
<tr>
<td>Comorbidities:</td>
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</tr>
<tr>
<td>0</td>
<td>692 (78.0)</td>
<td>96 (72.2)</td>
</tr>
<tr>
<td>≥1</td>
<td>195 (22.0)</td>
<td>37 (27.8)</td>
</tr>
<tr>
<td>Depression</td>
<td>0</td>
<td>8 (0.351)</td>
</tr>
<tr>
<td>No</td>
<td>827 (93.7)</td>
<td>118 (91.5)</td>
</tr>
<tr>
<td>Yes</td>
<td>56 (6.3)</td>
<td>11 (8.5)</td>
</tr>
</tbody>
</table>
Foot symptoms and knee OA

SD, standard deviation; IQR, interquartile range; BMI, body mass index; KL, Kellgren Lawrence.

* Baseline values

† P-values from chi-squared test for binary and categorical variables, Wilcoxon rank-sum or Kruskal-Wallis rank tests for variables presented as median (IQR), and analysis of variance tests for variables presented as mean (SD).
Table 2. Logistic regression analyses for the risk of developing knee symptoms during the four year follow up period. GEEs fit to account for the clustering of knees within participants. 50 knees from 29 participants were excluded due to missing data.

<table>
<thead>
<tr>
<th>Laterality of foot/ankle symptoms</th>
<th>Total number of knees (participants)</th>
<th>No knee symptoms</th>
<th>Knee symptoms</th>
<th>Risk for knee symptoms</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>N (%)</td>
<td>N (%)</td>
<td>Unadjusted OR (95% CI)</td>
</tr>
<tr>
<td>No symptoms (ref)</td>
<td>1742 (874)</td>
<td>1135 (89.7)</td>
<td>607 (83.8)</td>
<td>1</td>
</tr>
<tr>
<td>Any side</td>
<td>248 (125)</td>
<td>131 (10.3)</td>
<td>117 (16.2)</td>
<td>1.63 (1.16 to 2.27)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.55 (1.10 to 2.19)</td>
</tr>
<tr>
<td>Ipsilateral</td>
<td>70 (70)</td>
<td>40 (3.2)</td>
<td>30 (4.1)</td>
<td>1.34 (0.83 to 2.17)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.30 (0.80 to 2.12)</td>
</tr>
<tr>
<td>Contralateral</td>
<td>72 (72)</td>
<td>37 (2.9)</td>
<td>35 (4.8)</td>
<td>1.77 (1.11 to 2.84)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.68 (1.05 to 2.68)</td>
</tr>
<tr>
<td>Bilateral</td>
<td>106 (53)</td>
<td>54 (4.3)</td>
<td>52 (7.2)</td>
<td>1.74 (1.06 to 2.86)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>1.65 (0.98 to 2.78)</td>
</tr>
</tbody>
</table>

OR, odds ratios; CI, confidence intervals.

† Adjusted for age, sex, race, BMI, Charlson Comorbidity index (dichotomised) and depression.
Table 3. Logistic regression analyses for the risk of developing symptomatic and radiographic knee OA, during the four year follow-up period. GEEs fit to account for the clustering of knees within participants. 57 knees from 30 participants were excluded due to missing data.

<table>
<thead>
<tr>
<th>Laterality of foot/ankle symptoms</th>
<th>Total number of knees (participants)</th>
<th>No symptomatic knee ROA N (%)</th>
<th>Symptomatic Knee ROA N (%)</th>
<th>Risk for symptomatic knee ROA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Unadjusted OR (95% CI)</td>
</tr>
<tr>
<td>No symptoms (ref)</td>
<td>1736 (869)</td>
<td>1707 (88.1)</td>
<td>29 (64.4)</td>
<td>1</td>
</tr>
<tr>
<td>Any side</td>
<td>247 (124)</td>
<td>231 (11.9)</td>
<td>16 (35.6)</td>
<td>4.26 (2.23 to 8.12)</td>
</tr>
<tr>
<td>Ipsilateral</td>
<td>70 (70)</td>
<td>67 (3.5)</td>
<td>3 (6.7)</td>
<td>2.57 (0.76 to 8.74)</td>
</tr>
<tr>
<td>Contralateral</td>
<td>71 (71)</td>
<td>67 (3.5)</td>
<td>4 (8.9)</td>
<td>4.35 (1.61 to 11.74)</td>
</tr>
<tr>
<td>Bilateral</td>
<td>106 (53)</td>
<td>97 (5.0)</td>
<td>9 (20.0)</td>
<td>5.38 (2.50 to 11.55)</td>
</tr>
</tbody>
</table>

OA, osteoarthritis; ROA, radiographic osteoarthritis; OR, odds ratios; CI, confidence intervals.

† Adjusted for age, sex, race, BMI, Charlson Comorbidity index (dichotomised) and depression.
Osteoarthritis Initiative (OAI) participants at baseline (n=4796)

Excluded:
- Healthy control participants without risk factors for knee OA (n=122)
- Participants with symptomatic radiographic knee OA (n=1368)

OAI participants without symptomatic knee OA (n=3306 participants)

Excluded:
- Participants who reported knee symptoms or had missing knee symptoms at baseline (or both) (n=1061 participants)
- Participants with KL ≥ 2 or missing KL grade at baseline (n=1225)

Eligible sample = 1020 participants (2040 knees)

Aim 1: Development of knee symptoms

Aim 2: Development of symptomatic radiographic knee OA