Footwear for self-managing knee osteoarthritis symptoms: protocol for the Footstep randomised controlled trial

Kade L. Paterson1*, Kim L. Bennell1, Tim V. Wrigley1, Ben R. Metcalf1, Penny K. Campbell1, Jessica Kazsa2 and Rana S. Hinman1

Abstract

Background: Knee osteoarthritis (OA) is a leading cause of musculoskeletal pain and disability globally, and abnormal knee loading is central to disease pathogenesis. Clinical guidelines recommend clinicians provide advice regarding appropriate footwear for people with knee OA, yet there is little research comparing the effects of different footwear on knee OA symptoms. Research suggests that wearing flat flexible shoes is associated with lower knee joint loads compared to stable supportive shoe styles. This two-arm pragmatic, comparative effectiveness randomised controlled trial will compare the effects of daily use of flat flexible shoes and stable supportive shoes on knee OA clinical outcomes, over 6 months.

Methods: 164 people with symptomatic medial tibiofemoral OA of moderate to severe radiographic severity (Kellgren and Lawrence Grade 3 & 4) will be recruited from the community. Following baseline assessment, participants will be randomly allocated to receive either i) flat flexible shoes; or ii) stable supportive shoes. Participants will choose two different pairs of shoes from a selection that fulfil the criteria in their allocated shoe class. Limited disclosure will blind participants to group allocation. Participants will be instructed to wear their allocated shoes daily for 6 months (minimum of 6 h/day), after which participants will be reassessed. The primary outcomes are knee pain severity on walking (measured by numerical rating scale) and self-reported physical function (measured by the Western Ontario and McMaster Universities Osteoarthritis Index), assessed at baseline and 6 months. Secondary outcomes include additional measures of knee pain, function, sport and recreation participation and quality-of-life (measured using subscales of the Knee Osteoarthritis Outcome Score), as well as pain at other sites (measured by numerical rating scale), self-reported global ratings of change in pain and physical function (measured by 7-point rating scale), and physical activity levels (measured by Physical Activity Scale for the Elderly).

Discussion: This study will determine whether daily wear of flat flexible shoes improves clinical outcomes in the management of knee OA, compared to stable supportive shoes. Findings will assist clinicians in providing evidence-based advice regarding appropriate footwear for people with knee OA to self-manage symptoms.

Trial registration: Australian New Zealand Clinical Trials Registry reference: ACTRN12617001098325. Registered 28/07/2017.

Keywords: Osteoarthritis, Knee, Footwear, Shoes, Clinical trial, RCT, Biomechanics, Clinical trial

* Correspondence: kade.paterson@unimelb.edu.au
1Centre for Health, Exercise and Sports Medicine, Department of Physiotherapy, School of Health Sciences, Faculty of Medicine Dentistry & Health Sciences, The University of Melbourne, Melbourne, VIC, Australia
Full list of author information is available at the end of the article

© The Author(s). 2018 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 is the 11th highest contributor to global disability [1]. In 2012, 1.9 million Australians had OA [2], and modelling data estimate a 58% increase in OA by 2032 [2]. The knee joint is commonly affected, and knee OA is extremely debilitating. Pain is dominant, becoming persistent and more limiting as OA progresses. Health expenditure on OA in Australia in 2012 was $3.75 billion AUD [2], with most costs related to conservative and surgical treatments, lost productivity and substantial loss of quality of life. Globally, the annual total direct and indirect cost is estimated to vary between €300 to €19,5000 EUD per patient per year [3].

As OA is a chronic disease with no cure, people with OA have little choice but to self-manage their condition. Accordingly, advice about self-management is the cornerstone of conservative treatment, along with exercise and weight control [4, 5]. As abnormal biomechanics have been implicated in knee OA onset and progression [6, 7], clinical guidelines advocate that clinicians provide advice on “appropriate” footwear to help people with knee OA self-manage their symptoms [4, 8]. However, there is scant evidence from clinical trials to guide footwear choice for this patient group, or indeed, any population with chronic musculoskeletal pain. Guidelines suggest that shoes with thick shock-absorbing soles and arch supports are best for people with OA [8], based largely on expert opinion. Due to the lack of robust clinical trials in this area, footwear trials have been identified as an OA research priority by both the European League Against Rheumatism [8] and the National Institute for Health and Care Excellence (UK) [4].

Abnormal knee joint loading is central to OA pathogenesis [6, 7]. Knee load is typically inferred from three-dimensional gait analysis, usually by parameters of the external peak knee adduction moment. There is some evidence that greater knee load may increase symptoms and risk of structural progression in people with knee OA. For example, an increased knee adduction moment is associated with increased prevalence of medial bone marrow lesions [9] (an important source of pain in knee OA [10]) and medial cartilage defects [11]. The knee adduction moment is also one of the few modifiable factors that predict OA structural progression [7, 12–14]. Reducing knee loads via non-surgical biomechanical treatment strategies is thus an appropriate treatment aim for people with knee OA. Footwear is a promising avenue for self-management, given that foot position and motion influence knee loads.

Biomechanical research shows that wearing shoes significantly increases the knee adduction moment in people with knee OA, compared to barefoot walking [15, 16]. Biomechanical studies also show that some types of footwear increase knee loads more than others [17]. We, and others, have shown that stable supportive shoe styles increase knee loads significantly more compared to flat flexible shoe styles [18–20]. However it is stable supportive shoe styles that are typically recommended by clinicians for people with knee OA, and that are typically worn by people with the condition [21].

There are no high quality clinical trials comparing the effects of flat flexible shoes to stable supportive shoes on knee OA clinical outcomes. Uncontrolled data from the USA show flat flexible shoes reduce knee pain by 36% when worn for 6 h/day over 6 months in 16 people with medial knee OA [22]. In the only randomised controlled trial (RCT) to date [23], a small Brazilian study in 56 women with knee OA compared a standardized flat flexible “Moleca” shoe, worn ≥6 h/day for 6 months, to a control group who wore their own neutral tennis shoes “without characteristics of minimalist footwear”. Flat flexible shoes led to greater pain relief compared to the control group, and significant improvements in function were also observed. Although promising, participants in this study were not blinded, which introduces a high risk of bias given that treatment benefits are often over-estimated when using subjective outcomes with unblinded participants [24]. Thus, further research comparing the effects of flat flexible shoes to stable supportive shoes, using robust RCT designs, is warranted.

The primary aim of this study is to determine if flat flexible shoes lead to significantly greater reductions in knee pain with walking, and improvements in physical function, compared to stable supportive shoes when worn daily over six months. We hypothesise that flat flexible shoes will reduce pain and improve self-reported physical function more than stable supportive shoes.

Methods/design
Trial design
This protocol is described according to SPIRIT guidelines for clinical trials ([25]). The trial is a two-arm pragmatic, comparative effectiveness RCT comparing flat flexible shoes to stable supportive shoes. It will be conducted at The University of Melbourne over 3 years. The primary end-point for analysis of outcomes is after 6 months of shoe wear.

Participants
We will recruit 164 participants with medial tibiofemoral OA from the community via advertisements in print/radio/social media, clinicians and our volunteer database. Knee OA will be classified according to the American College of Rheumatology clinical and radiographic criteria for knee OA [26]. Participants will be included if they:

1) are aged ≥50 years;
ii) report knee pain on most days of the past month;

iii) report average pain during walking (over the previous week) of at least 4 on an 11-point numerical rating scale (NRS, with terminal descriptors of 'no pain' and 'worst pain possible');

iv) demonstrate moderate-severe (Grade 3–4) tibiofemoral OA on x-ray as determined by the Kellgren & Lawrence grading system [27], as recent research suggests these patients may represent a subgroup to which biomechanical interventions designed to reduce the KAM may be more effectively targeted; and

v) demonstrate tibiofemoral osteophytes on x-ray.

Participants will be excluded if they:

i) have lateral joint space narrowing greater than or equal to medial joint space narrowing on x-ray;

ii) have had knee pain for < 3 months;

iii) have had recent knee surgery (past 6 months) or planning surgery in next 6 months;

iv) currently use shoe orthoses, customized shoes or ankle braces;

v) primarily wear high heels, thongs or work boots that would restrict ability to wear allocated shoes 6 h/day;

vi) have had a hip or knee joint replacement on either side;

vii) have had a high tibial osteotomy on either leg;

viii) have had any knee injections in the past 3 months or planning injections in next 6 months;

ix) self-report any other muscular, joint or neurological condition affecting lower limb function;

x) self-report any systemic or inflammatory joint disease (e.g. rheumatoid arthritis);

xi) currently use, or plan to use, a gait aid in the next 6 months;

xii) do not understand written/spoken English; or

xiii) are unable to commit to study requirements (e.g. wearing shoes, attending appointments, completing outcomes, do not have foot size in the range of 8 to 13US for men, and 6 to 11US for women).

Fig. 1 Flow diagram of study phases
Procedure
Figure 1 outlines the trial phases and Table 1 describes the schedule for enrolment, intervention and assessments. Volunteers will be screened by an online form, then over the telephone by the Trial Coordinator. Potentially eligible participants will undergo standardised posteroanterior standing x-rays. Participants who have undergone standing x-ray in the prior 12 months, and can provide the images to research staff for screening, will not undergo new x-rays due to ethical concerns of exposing them to additional radiation. For participants with bilaterally eligible knees, the most symptomatic knee will be deemed the study knee with respect to outcome measurement.

Baseline assessments will be carried out at the Department of Physiotherapy, the University of Melbourne. At the 6-month follow-up, participants will complete questionnaires either on paper or electronically at home. Participants will also complete a short log book to record adherence with allocated footwear for one week of each month during the intervention period. If questionnaires or log books are not returned, the participant will be contacted by email and or phone to prompt their return, or as a last resort, to obtain primary outcome data. Ethical approval has been obtained from the University of Melbourne Human Research Ethics Committee (HREC No. 1748784). All participants will provide written informed consent.

Randomisation and allocation concealment
Eligible participants will be randomised to receive either flat flexible or stable supportive shoes following baseline data collection of primary and secondary outcomes. Randomisation (generated by the study biostatistician) will be by random permuted blocks (size 6 to 12) and stratified by radiographic disease severity (Kellgren & Lawrence grades 3 and 4 [27]). The schedule will be stored on a password-protected website (REDCap) maintained by a researcher not involved in either participant recruitment or administration of outcome measures. Group allocation will be revealed by this same researcher after baseline outcomes have been completed.

Participants will be blinded to group allocation by the process of limited disclosure. Participants will be

| TIMEPOINT | STUDY PERIOD |
|-----------|--------------|
|            | Enrolment | Allocation | Post-allocation | Close-out |
| ENROLMENT: | -1 | 0 | 1m | 2m | 3m | 4m | 5m | 6m |
| Eligibility screen | X | | | | | | |
| Informed consent | X | | | | | | |
| Allocation | X | | | | | | |
| INTERVENTIONS: | | | | | | |
| Flat flexible shoes | | | | | | | |
| Stable supportive shoes | | | | | | | |
| ASSESSMENTS: | | | | | | |
| Primary outcomes | | | | | | |
| Knee pain on walking (11-point numerical scale) | X | X | | |
| Physical function (WOMAC physical function subscale) | X | X | | |
| Secondary outcomes | | | | | | |
| Knee pain (KOOS pain subscale) | X | | X | |
| Sport and recreation activities (KOOS sport and recreation subscale) | X | | X | |
| Quality of life (KOOS quality of life subscale) | X | | X | |
| Patellofemoral pain and OA (KOOS PFJ pain and OA subscale) | X | | X | |
| Other joint pain (11-point numerical scale) | X | | X | |
| Health-related quality of life (AQoL-6D) | X | | X | |
| Physical activity (PASE) | X | | X | |
| Global improvement in pain | | | | | |
| Global improvement in physical function | X | | | |
| Other outcomes | | | | | | |
| Co-interventions | X | | | X | |
| Adverse events | | | X | X | |
| Adherence to wearing shoes | X | X | X | X | X | X |
informed that the trial is evaluating two different undisclosed “classes” of footwear to compare effects on knee OA symptoms. Participants will not be told about the classes of footwear under investigation, nor the specific shoe make/models that will be evaluated. Participants will not be informed about the study hypotheses, or which group they are allocated to until the study is completed, at which time they will be provided a lay summary of study purpose, hypotheses and findings. As the primary and secondary outcomes are participant-reported, and participants are blinded, this study is also considered assessor-blinded. Research staff administering and entering the participant-reported data will be blinded. Statistical analyses will be performed by a blinded biostatistician. Descriptive baseline characteristics that are rese archer-measured using objective methods in the laboratory (e.g. height, weight, foot posture and pressure) will be measured before randomisation by the unblinded staff member who will allocate participants to footwear group, and fit participants to allocated shoes.

Interventions

We previously identified five criteria (Table 2) that distinguish flat flexible shoes from stable supportive shoes and showed that flat flexible shoes were associated with lower knee joint loads compared to stable supportive shoes [18]. Using our established criteria, we have selected a range of commercially-available shoes for each intervention arm. Once allocated to an intervention group, participants will choose two different pairs of shoes from the range of options in order to maximize adherence.

Consumers with chronic knee pain (68 women and 43 men) were consulted in the trial planning phase to help refine shoe selection for the study to ensure that the chosen shoes would be acceptable and likely to be worn as instructed (minimum of 6 h/day for 6 months). In a survey format, consumers were presented with a wide range of commercially available shoes (that fulfilled the criteria for each shoe class) in different styles and colours. Survey respondents mostly indicated that black was the preferred shoe colour, where black was an option. Where possible, the most popular shoe styles in each class were selected for the trial. These are shown in Fig. 2, and include:

**Flat flexible shoes**

Merrell Bare Access (Men and Women), Vivobarefoot Primus Lite (Men and Women), Vivobarefoot Mata Canvas (Men) and Converse Dainty Low (Women).

**Stable supportive shoes**

ASICS Kayano (Men and Women), Merrell Jungle Moc (Men), Nike Air Max 90 Ultra (Women), Rockport Edge Hill (Men) and New Balance 624 (Women).

As footwear manufacturers frequently change shoe models from season to season and year to year, we expect that shoes readily available at trial commencement may cease to be available during the course of the trial or may be rebranded with a different shoe name. In such instances, shoes will be replaced by another pair of commercially available shoes that fulfil the listed criteria in Table 2.

Participants will be instructed to wear either pair of their allocated shoes as much as possible every day for 6 months, and to avoid wearing their own usual footwear as much as possible. At a minimum, participants will be asked to wear the shoes for at least 6 h every day.

Outcome measures

Table 1 summarises the outcome measures that are being collected for this study. Our primary outcomes are validated measures of pain and physical function that have been recommended for use in knee OA clinical trials [28]. Conclusions regarding treatment efficacy will be based on the 6-month changes in our primary outcome measures. Our two primary outcomes, measured at baseline and 6 months, are:

- Severity of knee pain during walking, scored on an 11-point NRS. Average overall pain on walking in the last week will be self-reported using an 11-point NRS. Scores range from 0 to 10, where 0 = no pain and 10 = worst pain possible. This outcome measure has demonstrated reliability in OA [29].

| Table 2 Criteria for selecting shoes |
|-------------------------------------|
| **Flat flexible shoes**              | **Stable supportive shoes**          |
| Heel height/thickness                | > 15 mm                              |
| Shoe pitch                           | < 15 mm                              |
| Arch support/motion control          | Absent                               |
| Sole flexibility                     | “Minimal” rigidity (Footwear Assessment Tool [45]) |
| Weight                               | ≤200 g                               |
| Weight                               | > 300 g                               |

Measurements are based on a size 9 US Men’s and size 9 US Women’s sized shoe

* A tolerance of +/− 10% was used for shoe weight
| Flat flexible shoes | Stable supportive shoes |
|---------------------|------------------------|
| **Men's shoes**     |                        |
| 1) Merrell Bare Access | 1) ASICS Kayano        |
| 2) Vivobarefoot Primus Lite | 2) Merrell Jungle Moc  |
| 3) Vivobarefoot Mata Canvas | 3) Rockport Edge Hill |
| **Women's shoes**   |                        |
| 1) Merrell Bare Access | 1) ASICS Kayano        |
| 2) Vivobarefoot Primus Lite | 2) Nike Air Max 90 Ultras |
| 3) Converse Dainty Low | 3) New Balance 624     |

*Fig. 2 Shoes selected at trial commencement*
Secondary outcome measures will be administered at baseline and at 6 months unless otherwise indicated. These include:

- **Physical function subscale of the WOMAC.** Limitations with physical functioning will be measured by the Western Ontario and McMaster Universities (WOMAC) Osteoarthritis Index (Likert version 3.1) [30]. The WOMAC is a self-report, disease-specific instrument which has established validity, reliability and responsiveness in an extensive range of OA studies [31]. The WOMAC physical function subscale contains 17 questions regarding knee function over the past week, with Likert response options ranging from 0 (no dysfunction) to 4 (extreme dysfunction). WOMAC scores will be extracted from the Knee Injury and Osteoarthritis Outcome Score (KOOS) [32] questionnaire, which contains the WOMAC questions. Total score ranges from 0 to 68, with higher scores indicating worse function.

- **Pain subscale of the KOOS.** The subscale is scored using 9 questions about knee pain experienced in the last week, with Likert response options from None to Extreme. Scores range from 0 to 100, with lower scores indicating worse pain [32].

- **Sport and recreation subscale of the KOOS.** The subscale is scored using five questions regarding function with sport and recreational activities in the last week, with Likert response options ranging from None to Extreme [32]. Scores range from 0 to 100, with lower scores indicating worse function [32].

- **Quality of life subscale of the KOOS.** The subscale consists of four questions about knee-related quality of life experienced in the last week, with five Likert response options for each question. Scores range from 0 to 100, with lower scores indicating worse quality of life [32].

- **Patellofemoral pain and OA subscale of the KOOS.** The subscale is comprised of 11 questions about knee pain and function experienced in the last week, with five Likert response options for each question. Scores range from 0 to 100, with lower scores indicating worse patellofemoral symptoms [33].

- **Lower limb pain severity.** Average overall pain in the last week will be recorded using an 11-point NRS (0 = no pain and 10 = worst pain possible) at each of the following sites: (i) study knee, (ii) contralateral knee, (iii) ipsilateral hip, (iv) contralateral hip, (v) ipsilateral foot/ankle, (vi) contralateral foot/ankle, and (vii) back.

- **Participant-perceived global change.** At 6 months, participants will rate their overall global change in a) pain and b) physical function since enrolling in the study. Response to treatment will be scored using a 7-point global rating of change Likert scale, with terminal descriptors of ‘much worse’ to ‘much better’ [34]. Participants who report ‘moderately better’ or ‘much better’ will be classified as improved. All other respondents will be classified as not improved.

- **Health-related quality of life.** The Assessment of Quality of Life (AQL) [35] (version AQL-6D) will be used to measure health-related quality of life. It consists of 20 items that assess independent living, mental health, relationships, pain, coping and senses. Scores range from −0.04 to 1.00 with higher scores indicating better quality of life.

- **Physical activity levels.** The Physical Activity Scale for the Elderly (PASE) will be used to measure physical activity over the previous week [36]. Scores range from 0 to over 400, with higher scores indicating greater physical activity.

### Treatment adherence
Participants will rate their perceived overall level of adherence over the past 6 months with the instruction to wear their allocated shoes for a minimum of 6 h per day on an 11-point NRS (with terminal descriptors of ‘shoes not worn at all’ and ‘shoes worn completely as instructed’). At the 6-month follow up assessment, participants will also indicate whether they stopped wearing either pair of study shoes during the 6 months on a categorical scale (Yes or No). Participants who score Yes will describe when and why they ceased wearing their allocated shoes, and this will be reported descriptively. Finally, participants will record in log books how many hours each day they wear their allocated shoes, for 7 consecutive days every month, over the 6-month intervention period.

### Adverse effects of treatment and co-interventions
Adverse events will be defined as any problem experienced in the study knee or elsewhere in the body because of wearing the study shoes. Adverse events will be self-reported by participants using a custom-developed table, and by open-ended questioning at 6 months. Participants will also be asked to contact the researchers at any time by telephone or email to report adverse events. Use of co-interventions (medications for knee pain and any other treatments for knee OA) will also be recorded at baseline and 6-months. Participants will complete a custom-developed table to indicate the frequency of use (over the past 6 months) of a range of pain and arthritis medications and co-interventions.

### Descriptive measures
A range of other measures will be collected to describe demographic characteristics, treatment expectations,
self-efficacy, foot biomechanics and current footwear characteristics. These measures will not be used to determine treatment efficacy but will be used for descriptive purposes. They include height, weight and body mass index; radiographic disease severity using the Kellgren & Lawrence scale [27]; current employment status; a rating of participant expectation of treatment outcome, measured on a 5-point ordinal scale; self-efficacy, determined using the Arthritis Self Efficacy Scale [37]; objective measures of foot posture, mobility and function including the Foot Posture Index (FPI [38]), Foot Mobility Magnitude [39], navicular drop [40] and in-shoe regional foot pressure patterns (Novel Pedar, Munich, Germany) during walking; and characteristics of each participant’s three most commonly worn shoes, including shoe weight, heel height, pitch, motion control properties and flexibility [18].

Sample size calculations
We aim to detect the minimal clinically important difference (MCID) on primary outcomes between groups (1.8 (out of 10) [41] for NRS pain and 6 (out of 68) for WOMAC function) [42]. We assume between-participant standard deviations of 2.7 and 11.4, and baseline to 6-month correlations of 0.21 and 0.39 for pain and function respectively (data from our footwear trial in a similar sample) [43]. Using analysis of covariance adjusted for baseline score, we need 46 per arm to achieve 90% power to detect MCID in pain and 65 per arm for function. Allowing for 20% attrition, we will recruit 82 people per arm (n = 164).

Statistical analyses
A biostatistician will analyse data in a blinded manner, with p values less than 0.05 considered significant. Main comparative analyses between groups will be performed using intention-to-treat. If more than 5% of primary outcomes are missing, multiple imputation will be applied. For the primary hypothesis, differences in mean change in pain and function (baseline minus follow-up) will be compared between groups using linear regression modelling adjusted for baseline values and the stratifying variable of Kellgren & Lawrence grade. Similar analyses will be conducted for continuous secondary outcomes. Improvement based on global change will be compared across groups using risk differences, calculated from fitted logistic regression models. The effect of patient characteristics on outcomes will be explored by including relevant terms as covariates in models. Standard diagnostic plots will be used to check model assumptions.

To assess whether the effect of shoe class on the primary outcomes is moderated by any of Kellgren & Lawrence grade, Foot Posture Index score, body mass index or baseline score on the KOOS patellofemoral pain and OA subscale, appropriate interaction terms between randomised group and each of these variables will be included in regression models for the primary outcomes, and for each potential effect modifier separately. For the continuous moderators, a fractional polynomial approach will be applied [44].

Timelines
The application for project funding was successful in October 2016 and funding commenced in August 2017. Ethics approval was obtained from the Human Research Ethics Committee of the University of Melbourne in May 2017. Recruitment commenced in August 2017 and will be completed in December 2019. The trial is due for completion in July 2020 when all participants will have completed 6-month follow-up.

Discussion
This paper has outlined the theoretical foundation, and presented the protocol, for a two-arm pragmatic, comparative effectiveness RCT comparing flat flexible to stable supportive shoes for people with knee OA. Clinical OA guidelines recommend appropriate footwear for knee OA self-management [4, 8], with some guidelines specifying shoes with supportive features, based solely on expert opinion [8]. Stable supportive footwear are also recommended by clinicians for people with knee OA, and are typically worn by people with the disease [21]. Yet these styles increase knee loads compared to flat flexible shoes [18–20], while limited clinical data suggests flat flexible shoes may lead to symptomatic benefits [23]. Given the lack of high quality RCTs comparing footwear styles, leading international organisations [4, 8] have highlighted footwear clinical trials as an important OA research priority. Findings from the Footstep trial will assist clinicians and patients with knee OA in selecting the most appropriate footwear to help self-manage symptoms associated with the disease.

Abbreviations
AQLQ: Assessment of Quality of Life; CONSORT: Consolidated Standards of Reporting Trials; HREC: Human Research Ethics Committee; KOOS: Knee Injury and Osteoarthritis Outcome Score; MCID: Minimal clinically important difference; NRS: Numerical rating scale; OA: Osteoarthritis; PASE: Physical Activity Scale for the Elderly; RCT: Randomised controlled trial; WOMAC: Western Ontario and McMaster Universities

Funding
This study is funded by the National Health and Medical Research Council (Project Grant #1124418). RSH is supported by an Australian Research Council Future Fellowship (FT130100175) and KLB by a National Health and Medical Research Council Principal Research Fellowship (APP1058440).

Availability of data and materials
The datasets used and/or analysed during the current study will be available from the corresponding author on reasonable request once the study has been completed.
Authors’ contributions
KLP, KLB, TVW and RSH conceived the idea for the study and RSH is leading the trial. KLP, TVW, JK and RSH obtained funding for the study. KLP, KLB, TVW and RSH designed the trial protocol with input from BRM, PKC, and JK. KLP drafted the manuscript with input from KLB, TVW, BRM, PKC, JK and RSH. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Ethical approval has been obtained from the University of Melbourne Human Research Ethics Committee (HREC No. 1748784). All participants will be required to provide written informed consent to participate.

Consent for publication
Not applicable.

Competing interests
KLP is a member of the Editorial Board of BMC Musculoskeletal Disorders. The other authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia.
2Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, VIC, Australia.

Received: 7 June 2018 Accepted: 19 June 2018
Published online: 18 July 2018

References
1. Cross M, Smith E, Hoy D, Noile S, Ackerman I, Fransen M, Bridgett L, Williams S, Gullenlin F, Hill CL, et al. The global burden of hip and knee osteoarthritis: estimates from the global burden of disease 2010 study. Ann Rheum Dis. 2014;73:323–30.
2. Arthritis & Osteoporosis Victoria. A Problem Worth Solving. The rising cost of musculoskeletal conditions in Australia. In: Arthritis & Osteoporosis Victoria; 2013.
3. Salmon JH, Rat AC, Sellam J, Michel M, Eschard JP, Guillemin F, Jolly D, Fauret B. Economic impact of lower-limb osteoarthritis worldwide: a systematic review of cost-of-illness studies. Osteoarthr Cartil. 2016;24:1500–8.
4. National Clinical Guideline Centre: Osteoarthritis. Care and management in adults. In: Clinical guideline CG177. Methods, evidence and recommendations. London: National Institute for Health and Care Excellence; 2014.
5. McAlindon TE, Bannuru RR, Sullivan MC, Arden NK, Berenbaum F, Bierma-Zeinstra SM, Hawkew GA, Henrotin Y, Hunter DJ, Kawaguchi H, et al. OARSI guidelines for the non-surgical management of knee osteoarthritis. Osteoarthr Cartil. 2014;22:363–88.
6. Andriacchi TP, Mundermann A. The role of ambulatory mechanics in the initiation and progression of knee osteoarthritis. Curr Opin Rheumatol. 2006;18:514–8.
7. Chehab EF, Favr J, Erhart-Hledik JC, Andriacchi TP. Baseline knee adduction and flexion moments during walking are both associated with 5 year cartilage changes in patients with medial knee osteoarthritis. Osteoarthr Cartil. 2011;19:1833–9.
8. Fernandes L, Hagen KB, Bijsma JW, Andresen O, Christensen P, Conaghan PG, Doherty M, Geenen R, Hammond A, Kjeken I, et al. EULAR recommendations for the non-pharmacological core management of hip and knee osteoarthritis. Ann Rheum Dis. 2013;72:1125–35.
9. Bennett KL, Creaby MW, Wrigley TV, Bowles KA, Hinman RS, Cicuttini F, Hunter DJ. Bone marrow lesions are related to dynamic knee loading in medial knee osteoarthritis. Ann Rheum Dis. 2010;69:1151–4.
10. Lo GH, McAlindon T, Niu J, Zhang Y, Beals C, Dabrowski C, Le Graverand MH, Hunter D. Bone marrow lesions and joint effusion are strongly and independently associated with weight-bearing pain in knee osteoarthritis: data from the osteoarthritis initiative. Osteoarthr Cartil. 2009;17:1562–9.
11. Creaby MW, Wang Y, Bennett KL, Hinman RS, Metcalf BR, Bowles KA, Cicuttini FM. Dynamic knee loading is related to cartilage defects and tibial plateau bone area in medial knee osteoarthritis. Osteoarthr Cartil. 2010;18:380–5.
12. Bennett KL, Bowles KA, Wang Y, Cicuttini F, Davies-Tuck M, Hinman RS. Higher dynamic medial knee load predicts greater cartilage loss over 12 months in medial knee osteoarthritis. Ann Rheum Dis. 2011;70:1770–4.
13. Miyazaki T, Wada M, Kawahara H, Sato M, Baba H, Shimada S. Dynamic load at baseline can predict radiographic disease progression in medial compartment knee osteoarthritis. Ann Rheum Dis. 2002;61:617–22.
14. Chang AH, Mosio KC, Chmiel JS, Eckstein F, Guermazi A, Prasad PV, Zhang Y, Almagor O, Bilelia L, Hayes K, et al. External knee adduction and flexion moments during gait and medial tibiofemoral disease progression in knee osteoarthritis. Osteoarthr Cartil. 2015;23:1099–106.
15. Kemp G, Crossley K, Wrigley T, Metcalf B, Hinman R. Reducing joint loading in medial knee osteoarthritis: shoes and canes. Arthritis Care Res. 2008;59:609–14.
16. Shakoor N, Block J. Walking barefoot decreases loading on the over extremity joints in knee osteoarthritis. Arthritis Rheum. 2006;54:2923–7.
17. Radzimirski AQ, Mundermann A, Sole G. Effect of footwear on the external knee adduction moment — a systematic review. Knee. 2012;19:163–75.
18. Paterson KS, Bennell KL, Wrigley TV, Metcalf BR, Kasza J, Hinman RS. Effects of footwear on the knee adduction moment in medial knee osteoarthritis: classification criteria for flat flexible vs stable supportive shoes. Osteoarthr Cartil. 2017;25:333–8.
19. Shakoor N, Sengupta M, Foucher KC, Wimmer MA, Fogg LF, Block JA. Effects of common footwear on joint loading in osteoarthritis of the knee. Arthritis Care Res. 2010;62:2917–23.
20. Shakoor N, Lidtke RH, Sengupta M, Fogg LF, Block JA. Effects of specialized footwear on joint loads in osteoarthritis of the knee. Arthritis Care Res. 2008;59:1214–20.
21. Paterson KS, Wrigley TV, Bennell KL, Hinman RS. A survey of footwear advice, beliefs and wear habits in people with knee osteoarthritis. J Foot Ankle Res. 2014;7:43.
22. Shakoor N, Lidtke RH, Wimmer MA, Mikolaitis RA, Foucher KC, Thorp LE, Fogg LF, Block JA. Improvement in knee loading after use of specialized footwear for knee osteoarthritis: results of a six-month pilot investigation. Arthritis Rheum. 2013;65:1282–9.
23. Trombini-Souza F, Matias AB, Yokota M, Butugan MK, Goldenstein-Schainer C, Fuller R, Sacco K. Long-term use of minimal footwear on pain, self-reported function, analgesic intake, and joint loading in elderly women with knee osteoarthritis: a randomized controlled trial. Clin Biomech. 2015;30:1194–201.
24. Wood L, Egger M, Gluud LL, Schulz KF, Juni P, Altman DG, Gluud C, Martin RM, Wood AJ, Sterne JA. Empirical evidence of bias in treatment effect estimates in controlled trials with different interventions and outcomes: meta-epidemiological study. BMJ. 2008;336:601–5.
25. Chan A, Tetzlaff JM, Altman DG, et al. Spirit 2013 statement: defining standard protocol items for clinical trials. Ann Intern Med. 2013;158:200–7.
26. Altman ASch, Bloch D, Bole G, Borenstein D, Brandt K, Christie W, Cooke TD, Greenwald R, Hochberg M, et al. Development of criteria for the classification and reporting of osteoarthritis. Classification of osteoarthritis of the knee. Arthritis Rheum. 1988;31:1039–49.
27. Kellgren JH, Jeffrey MR, Ball J. The epidemiology of chronic rheumatism: atlas of standard radiographs. Vol. volume 2. Oxford: Blackwell Scientific; 1963.
28. Bellamy N. Outcome measurement in osteoarthritis clinical trials. J Rheumatol. 1989;16:424–9.
29. Bellamy N. Osteoarthritis clinical trials: candidate variables and clinimetric properties. J Rheumatol. 1997;24:768–78.
30. 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;15:1833–40.
31. McConnell S, Kolopack P, Davis AM. The western Ontario and McMaster universities osteoarthritis index (WOMAC): a review of its utility and measurement properties. Arthritis Rheum. 2001;45:631–4.
32. Roos EM, Roos HP, Lohmander LS, Ekdahl C, Beynon BD. Knee injury and osteoarthritis outcome score (KOOS) - development of a self-administered outcome measure. J Orthop Sports Phys Ther. 1998;28:88–96.
33. Crossley KM, Macri EM, Cowan SM, et al. The patellofemoral pain and osteoarthritis subscale of the KOOS (KOOS-PF): development and validation using the COSMIN checklist. Br J Sports Med. 2017. doi: https://doi.org/10.1136/bjsports-2016-096776
34. ten Klooster PM, Drossaers-Bakker KW, Taal E, van de Laar MA. Patient-perceived satisfactory improvement (PPSI): interpreting meaningful change in pain from the patient’s perspective. Pain. 2006;121:151–7.
35. Osborne RH, Hawthorne G, Lew EA, Gray LC. Quality of life assessment in the community-dwelling elderly: validation of the assessment of quality of life (AQoL) instrument and comparison with the SF-36. J Clin Epidemiol. 2003;56:138–47.
36. Martin K, Rejeski W, Miller M, James M, Ettinger W Jr, Messier S. Validation of the PASE in older adults with knee pain and physical disability. Med Sci Sport Exer. 1999;31:627–33.
37. Lorig K, Chastain RL, Unger E, Shoor S, Holman HR. Development and evaluation of a scale to measure perceived self-efficacy in people with arthritis. Arthritis Rheum. 1989;32:37–44.
38. Redmond AC, Crosbie J, Ouvrier RA. Development and validation of a novel rating system for scoring standing foot posture: the foot posture index. Clin Biomech. 2006;21:89–98.
39. McPoil T, Vicenzino B, Cornwall M, Collins N, Warren M. Reliability and normative values for the foot mobility magnitude: a composite measure of vertical and medial-lateral mobility of the midfoot. J Foot Ankle Res. 2009;2:6.
40. Brody DM. Techniques in the evaluation and treatment of the injured runner. Orthop Clin N Am. 1982;13:541.
41. Bellamy N, Carette S, Ford P, Kean W, Le Riche N, Lussier A, Wells G, Campbell J. Osteoarthritis antirheumatic drug trials. III. Setting the delta for clinical trials- results of a consensus development (Delphi) exercise. J Rheumatol. 1992;19:451–7.
42. Tubach F, Ravaud P, Baron G, Falissard B, Logeart I, Bellamy N, Bombardier C, Felson D, Hochberg M, van der Heijde D, et al. Evaluation of clinically relevant changes in patient reported outcomes in knee and hip osteoarthritis: the minimal clinically important improvement. Ann Rheum Dis. 2005;64:29–33.
43. Hinman RS, Wrigley TV, Metcalf BR, Hunter DJ, Campbell P, Paterson K, Staples MP, Bennell KL. Unloading shoes for osteoarthritis of the knee: protocol for the SHARK randomised controlled trial. BMC Musculoskeletal Disorders. 2014;15:48.
44. Royston P, Sauerbrei W. Two techniques for investigating interactions between treatment and continuous covariates in clinical trials. Stata J. 2009;9:230.
45. Barton CJ, Bonanno D, Menz HB. Development and evaluation of a tool for the assessment of footwear characteristics. J Foot Ankle Res. 2009;2:10.
Author/s:
Paterson, KL; Bennell, KL; Wrigley, TV; Metcalf, BR; Campbell, PK; Kazsa, J; Hinman, RS

Title:
Footwear for self-managing knee osteoarthritis symptoms: protocol for the Footstep randomised controlled trial

Date:
2018-07-18

Citation:
Paterson, K. L., Bennell, K. L., Wrigley, T. V., Metcalf, B. R., Campbell, P. K., Kazsa, J. & Hinman, R. S. (2018). Footwear for self-managing knee osteoarthritis symptoms: protocol for the Footstep randomised controlled trial. BMC MUSCULOSKELETAL DISORDERS, 19 (1), https://doi.org/10.1186/s12891-018-2144-1.

Persistent Link:
http://hdl.handle.net/11343/220315

File Description:
Published version

License:
CC BY