ABSTRACT

Introduction Low back pain (LBP) and knee osteoarthritis (OA) are highly prevalent and disabling conditions that cause societal and economic impact worldwide. Two randomised controlled trials (RCTs) will evaluate the effectiveness of a multicomponent lifestyle intervention for patients with LBP and knee OA who are overweight or obese. The key targets of this intervention are to improve physical activity, modify diet and correct pain beliefs. These factors may explain how a lifestyle intervention exerts its effects on key patient-relevant outcomes: pain, disability and quality of life. The aim of this protocol is to describe a planned analysis of a mechanism evaluation for a lifestyle intervention for overweight or obese patients with LBP and knee OA.

Methods and analysis Causal mediation analyses of 2 two-armed RCTs. Both trials are part of a cohort–multiple RCT, embedded in routine health service delivery. In each respective trial, 160 patients with LBP and 120 patients with knee OA waiting for orthopaedic consultation will be randomised to a lifestyle intervention, or to remain part of the original cohort. The intervention consists of education and advice about the benefits of weight loss and physical activity, and the Australian New South Wales Get Healthy Service. All outcome measures including patient characteristics, primary and alternative mediators, outcomes, and potential confounders will be measured at baseline (T0). The primary mediator, weight, will be measured at 6 months post randomisation; alternative mediators including diet, physical activity and pain beliefs will be measured at 6 weeks post randomisation. All outcomes (pain, disability and quality of life) will be measured at 6 months post randomisation. Data will be analysed using causal mediation analysis with sensitivity analyses for sequential ignorability. All mediation models will be randomised to a lifestyle intervention, or to remain randomised to a lifestyle intervention, or to remain part of the original cohort. The intervention consists of education and advice about the benefits of weight loss and physical activity, and the Australian New South Wales Get Healthy Service. All outcome measures including patient characteristics, primary and alternative mediators, outcomes, and potential confounders will be measured at baseline (T0). The primary mediator, weight, will be measured at 6 months post randomisation; alternative mediators including diet, physical activity and pain beliefs will be measured at 6 weeks post randomisation. All outcomes (pain, disability and quality of life) will be measured at 6 months post randomisation. Data will be analysed using causal mediation analysis with sensitivity analyses for sequential ignorability. All mediation models were specified a priori before completing data collection and without prior knowledge about the effectiveness of the intervention.

Ethics and dissemination The study is approved by the Hunter New England Health Human Research Ethics Committee (13/12/11/5.18) and the University of Newcastle Human Research Ethics Committee (H-2015–0043). The results will be disseminated in peer-reviewed journals and at scientific conferences.

Trial registration number ACTRN12615000478516; Pre-results.

Strengths and limitations of this study

► Understanding the underlying causal mechanisms of a lifestyle intervention will explain how the intervention works, or why the intervention failed. These findings will have important clinical and policy implications and could guide implementation strategies.

► We propose to use contemporary methods for causal mediation analysis with sensitivity analyses to evaluate the robustness of the estimated mediation effects to violation of sequential ignorability—a critical assumption required for causal inference in mechanism evaluations.

► The primary mediator (weight) and the outcomes will be captured at the same time point. Thus, it will be challenging to attest the possibility of reverse causation of the mediator–outcome effect.

► Putative mediators including diet and physical activity are measured using self-reported questionnaires.

Newcastle Human Research Ethics Committee (H-2015–0043). The results will be disseminated in peer-reviewed journals and at scientific conferences.

Trial registration number ACTRN12615000490572 and ACTRN12615000478516; Pre-results.

BACKGROUND

Low back pain (LBP) and knee osteoarthritis (OA) are highly prevalent1 2 and disabling musculoskeletal conditions3 4 that cause societal5–7 and economic8 9 impact worldwide. The lifetime prevalence of LBP is 84%,2 and 40%–47% for knee OA.10 Of all health conditions, LBP is ranked first and OA ranked 11th as contributors to global disability.4 11 Direct costs for the management of LBP are estimated at $A4.7 billion in Australia (2012),7 £2.8 billion
in the UK (2013)\textsuperscript{12} and US$90 billion in the USA (1998)\textsuperscript{8}; and the cost of OA accounts for up to 2.5% of the gross national product in Australia, UK and USA.\textsuperscript{9}

A range of risk factors contribute to the development and persistence of LBP and OA. A large proportion of patients with LBP or OA are physically inactive,\textsuperscript{13–14} have poor diet\textsuperscript{14–15} and are overweight or obese.\textsuperscript{16–19} Targeting factors such as diet and physical activity as part of routine management is a plausible strategy to improve outcomes for these patients.\textsuperscript{20–22} Two randomised controlled trials (RCTs) will test the effectiveness of a multicomponent lifestyle intervention for patients with LBP\textsuperscript{23} and knee OA\textsuperscript{24} who are overweight or obese. However, merely evaluating the effectiveness of these interventions is insufficient\textsuperscript{25}; it is important to understand the underlying causal mechanisms that explain how the intervention works, or why the intervention doesn’t work.\textsuperscript{26–27}

**Explaining underlying mechanisms**

Complex interventions for patients with LBP and knee OA are usually evaluated by their effects on patient-relevant outcomes such as pain, disability and quality of life (QoL).\textsuperscript{23–24,25,26,27,28} However, pragmatic interventions such as a lifestyle intervention do not directly target patient-relevant outcomes; they target intermediate factors (often called mediators), such as diet or physical activity, that are then hypothesised to have a causal effect on patient-relevant outcome(s).\textsuperscript{26} Therefore, merely evaluating the effect of the intervention leaves a black-box that conceals the underlying mechanism(s) of the intervention. The aim of a mechanism evaluation is to unpack the black box by decomposing the entire intervention effect into indirect and direct effects. The indirect effect is the effect of the intervention on an outcome that is carried through a selected mediator, and the direct effect is the remaining effect of the intervention that is not explained via the selected mediator. For example, the entire effect of the lifestyle intervention on QoL could be decomposed into an effect carried through changes in diet (indirect effect) and remaining unexplained mechanisms (direct effect).

One way of quantifying causal mechanisms is by conducting causal mediation analysis.\textsuperscript{25–27} This approach can produce important information about the underlying mechanisms of an intervention. If the intervention is effective, causal mediation analysis informs whether the hypothesised mechanisms actually occurred.\textsuperscript{27} Conversely, if the intervention is ineffective, causal mediation analysis can identify where the hypothesised mechanism breaks down.\textsuperscript{27} By using this information, interventions can be refined on the basis of empirical evidence about the underlying mechanism.\textsuperscript{25–27} Elements of the intervention that aim to target proposed mediators that do not affect the outcome can be eliminated; and elements that influence a mediator that actually affects outcome can be retained and optimised.

**Mechanisms of a lifestyle intervention**

Causal mechanisms of lifestyle interventions are unknown. However, there is evidence suggesting that weight loss, inactivity and poor diet are important risk factors that should be considered treatment targets for patients with LBP and OA (ie, mediators). For knee OA, being overweight or obese is a modifiable risk factor.\textsuperscript{18–19,31–32} Further, meta-analyses show that weight loss interventions result in moderate improvements in pain and function for overweight or obese patients with knee OA.\textsuperscript{33} Similarly for LBP, meta-analyses show significant associations between overweight or obesity and a number of LBP outcomes.\textsuperscript{16,34} This suggests that weight might be an appropriate treatment target for both of these conditions to improve patient-related outcomes.

It is also apparent that physical activity and diet may play a role in this mechanism for both conditions because of their effects on weight.\textsuperscript{11,13–15,35–37} Inaccurate beliefs about pain are also associated with poor LBP and OA outcomes.\textsuperscript{38–39} Despite evidence for the relationship between weight, physical activity, and pain beliefs and patient-relevant outcomes, these risk factors have not been tested as underlying mechanisms of lifestyle interventions for patients with LBP and knee OA.

To test these underlying mechanisms, we have embedded a priori mechanism evaluations into two RCTs that will test the effectiveness of a lifestyle intervention for patients with LBP\textsuperscript{23} and knee OA\textsuperscript{24} who are overweight or obese. Our primary hypothesis is that in patients with either LBP or knee OA who are overweight or obese, a lifestyle intervention will have a causal effect on outcomes (pain, disability and QoL) via a primary mechanism through weight. Our secondary hypothesis is that the causal effect of a lifestyle intervention will also be explained via alternative mechanisms including changes in diet, physical activity and pain beliefs.

**Objectives**

The objective of this study is to test the underlying mechanisms of a lifestyle intervention for patients with LBP or OA who are obese or overweight. The specific objectives of this study vary according to whether the lifestyle intervention is effective (unknown at the time of writing this protocol):

- If the intervention is effective, our primary objective is to estimate the extent to which weight mediates this effect. Our secondary objective will be to further refine this mechanism via three serial multiple mediator paths (changes in diet, physical activity and pain beliefs) that then cause changes in weight.
- If the intervention is ineffective, our primary objective is to determine where the causal path breaks down. All potential mediators (weight, diet, physical activity and pain beliefs) will be tested independently.

**METHOD**

**Design**

We will conduct a combined causal mediation analysis of 2 two-armed RCTs.\textsuperscript{23–24} Both trials are part of a cohort multiple RCT,\textsuperscript{10} embedded in routine health service delivery. In both trials, participants are recruited from an existing cohort of patients waiting for orthopaedic consultation; then randomised to receive a lifestyle intervention (intervention group), or to receive usual care by
remaining in the original cohort (control group). The key differences between Williams et al.23 and O’Brien et al.24 are the clinical populations (LBP25 and knee OA),24 and the additional physiotherapy consultations exclusively delivered in the LBP trial.23 Thus, it is plausible that the two different clinical populations may respond differentially to their respective interventions. To accommodate this hypothesis, we will use moderated causal mediation analysis to estimate trial-specific effects, and averaged effects across both trials. If trial assignment (LBP trial vs OA trial) is a significant moderator, we will interpret trial-specific mediation effects in separation; however, if trial assignment is not a significant moderator, we will interpret the averaged mediation effects across both trials.

The trials began recruiting on 11 May 2015 and we expect to close the trial by June 2017. Data collection is still ongoing and all investigators were blind to group allocation at the time of planning and writing this study protocol. Further details of each trial have been outlined by Williams et al.23 (ACTRN12615000478516) and O’Brien et al.24 (ACTRN12615000490572).

Participants and recruitment

One RCT involves 120 patients with OA of the knee,24 and the other, 160 patients with non-specific LBP.23 Patients in both RCTs are those waiting for outpatient orthopaedic consultation at a tertiary referral public hospital in New South Wales (NSW), Australia.

Randomisation

In both trials, eligible patients from the cohort are randomised to an intervention or control group (1:1 ratio). The randomisation schedule was a priori generated by an independent statistician using the SURVEYSELECT procedure (SAS V.9.3). Allocation is concealed and all outcome assessors, patients and investigators are blind to group allocation. Patients are blind to group allocation by nature of the cohort multiple design. This design offers the intervention and control as part of a routine clinical service, where patients consent to routine data collection. Patients randomised to the intervention group are not aware of the offer of the control arm. Likewise, patients randomised to the control group are not aware of the offer of the intervention arm. Thus, patients are not able to discriminate whether the intervention or control are being offered as part of a clinical trial. This reduces the risk of performance bias (how well the participants engage with the intervention). Service providers delivering the intervention are blind to treatment status as they are not aware that patients were being referred from a clinical trial. The outcome assessors do not have access to the randomisation schedule, thus blind to group allocation. This reduces the risk of detection bias (differential outcome measurement between groups).

Intervention groups

Participants in both RCTs23 24 will receive advice and education about the benefits of weight loss and physical activity for their conditions by trained interviewers. Participants are then referred to the NSW Get Healthy Information and Coaching Service (GHS; www.gethealthynsw.com.au).41 The GHS is a free, population-wide, telephone-based health coaching service provided by the NSW Government to support adults in NSW to make sustained healthy lifestyle improvements including diet, physical activity and achieving or maintaining a healthy weight. This service consists of 10 individually tailored coaching calls delivered by university-qualified health coaches, including dieticians, exercise physiologists and psychologists, over a 26-week period. All coaches undergo standardised training before delivering the GHS, thus reducing the potential for differential between coach effects. Coaching is provided on a tapered schedule. Six calls are made in the first 12 weeks to guide, monitor and improve uptake; and four calls are dispersed over the remaining 12 weeks to maintain adherence and avoid relapse.42 This tapered schedule will be kept consistent across all participants, reducing the potential for bias.

Participants with LBP25 will receive an additional clinical consultation with the study physiotherapist before beginning the NSW GHS programme. The consultation aims to correct erroneous pain beliefs, highlight the consequences of unhealthy lifestyle factors, and to provide general encouragement and examples of how improving lifestyle factors can influence pain outcomes and QoL. The consultation also involves behaviour change techniques, informed by self-determination theory43 44 that aims to develop autonomous motivation by increasing perceived competence and self-regulation.41

Control groups

Participants allocated to the control group will remain in usual care. The health service does not provide any active management for patients with knee OA or LBP during the orthopaedic consultation waiting period.

Assessment time points

Patient characteristics, outcome measures, primary and alternative mediators, and potential confounders are measured at baseline (T0) prior to randomisation. The primary putative mediator (weight) will be measured 6 months after randomisation. All putative alternative mediators (diet, physical activity and pain beliefs) will be measured 6 weeks after randomisation. Outcomes will be measured 6 months after randomisation. The intervention and assessment time points are outlined in table 1.

Primary outcome measures

Average pain intensity over 7 days will be measured using an 11-point pain Numeric Rating Scale (NRS; 0=no pain, 10=pain as bad as it could be).45 We will measure self-perceived disability using the 24-item Roland-Morris Disability Questionnaire in patients with LBP46; and the Western Ontario and McMaster Universities Osteoarthritis Index47 in patients with knee OA. We will measure QoL using the Short Form Health Survey V.2.48
Putative mediators

The primary mediator, weight, will be measured to the nearest 0.1 kg by a trained research assistant using the International Society for the Advancement of Kinanthropometry procedures. Physical activity will be measured using the Active Australia Survey, which has moderate reliability (Cohen’s kappa=0.52) and good face and criterion validity. Dietary intake will be measured using a Short Food Frequency Questionnaire, which has moderate reliability (weighted kappa range=0.37–0.85) and criterion validity. Pain-related attitudes and beliefs will be measured using the Survey of Pain Attitudes Questionnaire. All putative mediators are measured in both control and intervention groups in both trials. These mediators are measured using self-reported questionnaires with known limitations.

Potential confounders

We will control for the following pretreatment confounders: pain duration, baseline pain, disability and QoL. These variables were selected on the basis of their theorised causal relationships with the mediator and outcome variables. We will include baseline measures of the mediators and outcomes in the regression models.
as covariates. Directed acyclic graphs specific to each model are presented in figure 1.

Causal mediation analysis
We plan to construct single and multiple mediator models based on current recommendations for causal mediation analysis. The details of each model are illustrated in figure 1 and table 2; and the overall analysis plan is outlined in figure 2.

Justification for primary and alternative mechanisms
Our hypothesised mechanisms are based on theory and evidence. We selected weight at 6-month follow-up as our primary mediator because the key component of the lifestyle intervention was targeted to reduce weight, and because the target population was overweight or obese. Evidence suggests that weight might have direct causal effects on patient-related outcomes (pain, disability and QoL). This is because initial consultations in the LBP trial aimed to reassure patients and reframe erroneous beliefs about pain. Although patients with OA did not receive a clinical consultation that directly targeted pain beliefs, the GHS may have inadvertently changed pain beliefs through the promotion of physical activity. The physical activity component could enable the patients to realise that pain does not need to be a barrier to keeping a physically active lifestyle. This theory is informed by Albert Bandura’s techniques of verbal persuasion, modelling and mastery. These refined mechanisms will be tested in serial multiple mediator models (figure 1B).

Sample size
Both trials are sufficiently powered (90%) to detect clinically meaningful between-group changes in pain (1.5-point reduction on NRS) and weight (6% reduction). To gain a general appreciation for the required sample size to detect an indirect effect through the primary mediator (weight), we used the sample size estimator for joint indirect effects developed by Vittinghoff and Neiands. With a two-sided alpha of 0.05, exposure-mediator error term correlation coefficient of 0, and mediator-outcome error term correlation coefficient of 0.2, a sample of 71 per group provides 80% power to detect a proportion mediated of 50%, with clinically meaningful treatment-mediator (r=0.5) and

<table>
<thead>
<tr>
<th>Model</th>
<th>Treatment (X)</th>
<th>Alternative mediator (M₂) at 6 weeks</th>
<th>Primary mediator (M₁) at 6 months</th>
<th>Outcome (Y) at 6 months</th>
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<tr>
<td>1.0</td>
<td>Rx</td>
<td>Weight</td>
<td></td>
<td>Pain/Disability/QoL</td>
</tr>
<tr>
<td>1.1*</td>
<td>Rx</td>
<td>Diet</td>
<td>Weight</td>
<td>Pain/Disability/QoL</td>
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<tr>
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<td>Rx</td>
<td>Physical activity</td>
<td>Weight</td>
<td>Pain/Disability/QoL</td>
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<tr>
<td>1.3*</td>
<td>Rx</td>
<td>Pain beliefs</td>
<td>Weight</td>
<td>Pain/Disability/QoL</td>
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<tr>
<td>1.4</td>
<td>Rx</td>
<td>Diet</td>
<td></td>
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<tr>
<td>1.5</td>
<td>Rx</td>
<td>Physical activity</td>
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<td>Pain/Disability/QoL</td>
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<tr>
<td>1.6</td>
<td>Rx</td>
<td>Pain beliefs</td>
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<td>Pain/Disability/QoL</td>
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<tr>
<td>2.0</td>
<td>Rx</td>
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<td>2.1*</td>
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<td>2.3</td>
<td>Rx</td>
<td>Pain beliefs</td>
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<td>Pain/Disability/QoL</td>
</tr>
</tbody>
</table>

*Multiple mediator models will only be tested if there is a significant relationship between Mₙ and M₁. If the relationship is non-significant, then the alternative mediators will be tested in separate single mediator models with the mediator measured at week 6. Significance levels are set a priori at p<0.05.

QoL, quality of life.
Figure 2 Overall analysis plan. Note: ‘pain’ is interchangeable with disability and QoL. PA, physical activity; PB, pain beliefs; QoL, quality of life.
mediator-outcome \((r=0.3)\) effects. The sample sizes for both trials were primarily estimated to detect the main effect of the intervention on pain and weight. Therefore, this post hoc power calculation provides indication that both trials would be powered to detect an indirect effect that consists of moderate treatment-mediator and mediator-outcome effects. Moderate effects would be considered clinically meaningful effects based on previous work.\(^{67,68}\) Sample size estimators for multiple mediator models are currently unavailable.\(^{69}\) O’Rourke and Mackinnon provide evidence that multiple mediator models have more power than single mediator models.\(^{70}\) Thus we expect this study to have sufficient power for multiple mediator models.

**Methodological considerations**

**No-confounding assumption (sequential ignorability)**

Estimating indirect effects that have causal meaning relies on satisfying the ‘no-confounding’ assumption, often termed ‘sequential ignorability.’\(^{65}\) It is critical that the treatment-mediator effect and the mediator-outcome effect are not confounded.\(^{65}\) In mediation analyses of standard RCTs, this assumption only holds for the treatment-mediator and treatment-outcome effects (due to randomisation). However, since the mediators cannot be randomised, this assumption does not hold for the mediator-outcome relationship.\(^{66}\) There may be unknown or unmeasured confounders that might induce a spurious relationship between the mediator and outcome. Recent advances in causal mediation analysis have developed sensitivity analysis techniques that can estimate the impact of violating this assumption, which we will employ in this study.\(^{71}\) These methods are an extension of the traditional methods (Baron and Kenny)\(^{72}\) and reflect contemporary advances in causal mediation analysis.\(^{61}\)

**Alternative mediator as a post-randomisation confounder in multiple mediator models**

In mediation analyses, post-randomisation confounders are variables that are affected by the treatment that then simultaneously influence the mediator and outcome. The presence of a post-randomisation confounder effectively induces bias for indirect and direct effects.\(^{73}\) By construction of the multiple mediator model, an alternative mediator \((M_2)\) is a post-randomisation confounder for the primary mediator-outcome relationship (ie, the alternative mediator that is affected by the treatment might causally affect both the primary mediator and the outcome and induce a spurious relationship). For example, changes in diet caused by the treatment can subsequently have a causal effect on weight and QoL, thereby inducing a spurious relationship between weight and QoL. To overcome this problem, we will assess the dependence between the alternative mediators (diet, physical activity, pain beliefs) and the primary mediator (weight). If an alternative mediator and a primary mediator are significantly correlated, we will build serial multiple mediator models, as recommended by Imai et al.\(^{79}\) If the alternative and primary mediators are not related, then we will not treat the alternative mediator as a post-randomisation confounder, and test the alternative mediators in independent single mediator models.

**Data analysis**

Analyses will be performed in R (The R Foundation for Statistical Computing) using the mediation package.\(^{74}\)

**Single mediator models**

A model-based inference approach will be used to estimate the average causal mediation effect (ACME), average direct effect (ADE) and the average total effect.\(^{74}\) First, we will fit two regression models: the mediator model and the outcome model. The mediator model is constructed with the treatment status as the independent variable and the mediator as the dependent variable. The outcome model is constructed with the treatment status and the mediator as independent variables, the outcome as the dependent variable, and the set of observed pretreatment confounders as covariates. Continuous mediators and outcomes that are normally distributed will be modelled using linear models \((l m)\); but if skewed, they will be modelled using generalised linear models \((g l m)\) with appropriate family and link functions.\(^{75}\) The ordinal mediator (diet) will be modelled using the proportional odds logistic model \((p o l r)\).\(^{74}\)

Because it is plausible that the indirect and direct effect sizes might depend on treatment allocation (treated and non-treated), we will include a treatment-mediator interaction term in the outcome model. We will calculate two separate ACMEs that are conditional on treatment status \((x=1 and x=0)\) and their marginal effects. We will interpret both conditional effects to generalise to their respective treatment group (treated and non-treated) and the marginal effect to generalise to the overall population. Not accounting for small non-significant interaction effects can dramatically influence the indirect and direct effect estimates.\(^{69}\)

The mediates function will use the mediator and outcome models to estimate the potential values of the mediator and outcome. The simulated potential values of the mediator and the outcome will be used to compute the ACME, ADE and average total effects. We will use 1000 bootstrap stimulations to generate 95% CIs. We will interpret the unstandardised point estimate of ACME and its 95% CIs.

Trial assignment (OA trial vs LBP trial) could moderate indirect and direct effects. Therefore, we will test the moderating effect of trial assignment by using the \texttt{test\_modmed} function. This function directly tests the difference in the ACME and ADE between two levels of the hypothesised moderator (OA trial vs LBP trial). If the ACME and ADE are statistically different, we will analyse the two trials separately to estimate the ACME and ADE that are specific to each trial. However, if they are not different, we will estimate an averaged ACME and ADE across both trials.
A sensitivity analysis will be conducted to determine the robustness of the ACME to the influence of violating the no-confounding assumption (sequential ignorability). The level of confounding due to unknown confounders is represented by the correlation between the residuals (error terms) from the mediator and outcome models, denoted $\rho$ ($\rho$ho). If $\rho=0$ (ie, no correlation between residuals), then this can be hypothetically interpreted as no unmeasured confounding. We will use the medsens function to explore how varying levels of $\rho$ (between the extremes of $-1$ and $+1$) influence the ACME. The output will provide the values of $\rho$ at which the CIs for the ACME include 0 (a non-significant ACME). That is, how strong the effect of unmeasured confounding would need to be to invalidate the estimated ACME.

Multiple mediator models

For multiple mediator models, we will use an expanded mathematical framework. Multiple mediator models will only be constructed if the alternative mediator (diet, physical activity and pain beliefs) and primary mediator (weight) are related. We will use the multimed function from the mediation package to estimate the ACME and ADE, and the sensitivity parameters. We will use 1000 bootstrap stimulations to generate 95% CIs.

Conclusion

We present an analysis plan for a mechanism evaluation of a lifestyle intervention for patients with knee OA and LBP who are overweight or obese. In the event that the intervention is effective, this investigation will provide evidence for hypothesised causal mechanisms through changes in weight, diet, physical activity and pain beliefs. If the intervention is ineffective it will provide explanations for why the intervention did not work. These results will help refine the intervention and guide intervention strategies.

Author affiliations

1School of Medicine and Public Health, Hunter Medical Research Institute, University of Newcastle, Newcastle, Australia
2Centre for Rehabilitation Research, Centre for Statistics in Medicine, Nuffield Department of Orthopaedics, Rheumatology and Musculoskeletal Sciences, University of Oxford, Oxford, United Kingdom
3Centre for Pain, Health and Lifestyle
4Neuroscience Research Australia, University of New South Wales, Sydney, Australia
5Hunter New England Population Health, Wallsend, Australia
6Musculoskeletal Division, The George Institute for Global Health, University of Sydney, Sydney, Australia
7Ambulatory Care Centre, John Hunter Hospital, Newcastle, Australia

Contributors HL, JW, SK, AW, KMO, RHJ, LW, SLY, EC, RH, EKR, JHM and CMW were responsible for the design of the study. CMW and JW procured funding. All authors contributed to developing the intervention and data collection protocols and materials, and reviewing, editing, and approving the final version of the paper. HL drafted the manuscript, and all authors subsequently contributed to the manuscript. All authors have read and approved the final manuscript.

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Mechanism evaluation of a lifestyle intervention for patients with musculoskeletal pain who are overweight or obese: protocol for a causal mediation analysis

Hopin Lee, John Wiggers, Steven J Kamper, Amanda Williams, Kate M O’Brien, Rebecca K Hodder, Luke Wolfenden, Sze Lin Yoong, Elizabeth Campbell, Robin Haskins, Emma K Robson, James H McAuley and Christopher M Williams

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