Efficacy of heel lifts versus calf muscle eccentric exercise for mid-portion Achilles tendinopathy (HEALTHY): a randomised trial

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ABSTRACT
Objective(s) To compare the efficacy of in-shoe heel lifts to calf muscle eccentric exercise in reducing pain and improving function in mid-portion Achilles tendinopathy.

Methods This was a parallel-group randomised superiority trial at a single centre (La Trobe University Health Sciences Clinic, Discipline of Podiatry, Melbourne, Victoria, Australia). One hundred participants (52 women and 48 men, mean age 45.9, SD 9.4 years) with clinically diagnosed and ultrasonographically confirmed mid-portion Achilles tendinopathy were randomly allocated to either a (1) heel lifts (n=50) or (2) eccentric exercise (n=50) group. The primary outcome measure was the Victorian Institute of Sport Assessment-Achilles (VISA-A) questionnaire at 12 weeks. Differences between groups were analysed using intention to treat with analysis of covariance.

Results There was 80% follow-up of participants (n=40 per group) at 12 weeks. The mean VISA-A score improved by 26.0 points (95% CI 19.6 to 32.4) in the heel lifts group and by 17.4 points (95% CI 9.5 to 25.3) in the eccentric exercise group. On average, there was a between-group difference in favour of the heel lifts for the VISA-A (adjusted mean difference 9.6, 95% CI 1.8 to 17.4, p=0.016), which approximated, but did not meet our predetermined minimum important difference of 10 points.

Conclusion In adults with mid-portion Achilles tendinopathy, heel lifts were more effective than calf muscle eccentric exercise in reducing pain and improving function at 12 weeks. However, there is uncertainty in the estimate of effect for this outcome and patients may not experience a clinically worthwhile difference between interventions.

Trial registration number ACTRN1261701225303.

BACKGROUND
Mid-portion Achilles tendinopathy is a common musculoskeletal condition, presenting as pain and stiffness between 2 and 6 cm above the posterior calcaneus.1–3 The pathological tendon shows abnormal changes on ultrasound imaging, such as altered tendon structure, disorganisation of fibres and thickening.4,5 Individuals with Achilles tendinopathy often report reduced participation in physical activity,6 impaired performance,7 and deficits in ankle joint plantarflexion strength and endurance.8 Achilles tendinopathy can also impact health-related quality of life.9 Accordingly, effective interventions are essential for this common and disabling condition.

Calf muscle eccentric exercise has been extensively researched, frequently administered,10–13 and is the recommended intervention for the management of mid-portion Achilles tendinopathy.14 Numerous eccentric exercise protocols exist, varying in load, frequency and speed. Nevertheless, similar treatment effectiveness has been reported, suggesting there is currently no optimum protocol.15–17 Although eccentric exercise continues to be the ‘gold standard’ for mid-portion Achilles tendinopathy,10 success rates are varied, with up to 44% of those affected reporting the exercise to be ineffective.15,16 One reason for the varied success could be heterogeneous exercise adherence.17

Heel lifts are inserts placed inside footwear, and are also commonly prescribed for mid-portion Achilles tendinopathy.18,19 Heel lifts do not require active engagement to exert their effects—adherence is usually high. Heel lifts reduce ankle joint dorsiflexion during running, and reduce gastrocnemius muscle length and Achilles tendon strain during walking.20 While these biomechanical findings suggest heel lifts may be effective for mid-portion Achilles tendinopathy, there have been no high quality randomised trials of this intervention. The aim of this study was, therefore, to compare heel lifts to calf muscle eccentric exercise for reducing pain and improving function in individuals with mid-portion Achilles tendinopathy.

METHODS
Study design
The Heel lifts vs calf muscle eccentric Exercise for Achilles Tendinopathy (HEALTHY) study was a parallel-group randomised superiority trial. The description of the methods of this trial have been summarised from the published protocol.21 Findings are reported according to the Consolidated Standards of Reporting Trials 2010 Statement22 and the TIDieR guide.23

Protocol deviations
In the trial registration and study protocol,21 we specified that we would randomise 92 participants. However, we had more eligible volunteers for study participation and therefore enrolled 100 participants into the study.
Study population and setting
Between September 2017 and December 2017, participants were recruited from advertisements placed in local newspapers and on social media platforms. Full inclusion and exclusion criteria have been detailed previously; a summary of the eligibility criteria is below. Participants aged at least 18 years of age, with clinically diagnosed and ultrasonographically confirmed mid-portion Achilles tendinopathy, with pain present for at least 2 months and reporting a minimum level of pain of at least a 3 out of 10 on a Numerical Rating Scale-11 in the last week were included. Participants were excluded if they had previous Achilles tendon rupture or surgery in the most symptomatic limb(s), chronic ankle instability, treatment with in-shoe heel lifts or a calf muscle eccentric exercise programme in the previous 3 months, or a medical condition (eg, inflammatory arthritis) that deemed the participant unsuitable for participation. All assessments were conducted at a single centre (La Trobe University Health Sciences Clinic, Discipline of Podiatry, Melbourne, Victoria, Australia).

Study interventions
Participants allocated to the heel lifts group received three pairs for bilateral use (ie, irrespective of unilateral/bilateral mid-portion AT) of Clearly Adjustable 12 mm heel lifts (A. Algeo, Australia) made from firm (Shore A 90) multilayered clear vinyl. Heel lifts were placed in up to three pairs of each participant’s most frequently worn footwear. According to each participant’s shoe size, small, medium or large heel lifts were dispensed. A 3.2 mm PPT Ultralux top cover was adhered to the top surface to maximise comfort (figure 1A). Existing insoles were removed from participant’s footwear to minimise heel slippage.

The participants allocated to the eccentric exercise group received a 12-week calf muscle eccentric exercise programme, based on the Alfredson method.11 The technique for performing this exercise programme has been detailed previously.21 The program involved performing the exercises twice daily, 7 days per week for 12 weeks. Participants with bilateral symptoms performed the programme on both sides. The programme involved two exercises: the first with the knee extended (figure 1B,C), and the second, with the knee flexed to increase activation of soleus (figure 1D,E). Both exercises involved three sets of 15 repetitions, performed twice daily on the symptomatic limb(s). Initially, resistance consisted of bodyweight, and the participants stood with all their bodyweight on the injured leg. Participants were advised to continue the exercise even if they experienced pain. However, they were advised to stop and resume the following day if the pain became disabling. Participants were advised to increase the resistance component of the programme once they could perform both exercises with no pain or discomfort on the injured side. Participants were told to use a weighted backpack to increase resistance, beginning with the addition of 5 kg of mass to the backpack, and to increase the load by 5 kg increments.

Both groups received a handout explaining their allocated intervention and a modified physical activity programme based on the pain-monitoring model,13 as detailed previously;21

Treatment preference, credibility and expectancy
Participants were asked prior to randomisation to indicate their preferred treatment.28 Given that participants were not blinded to their allocated intervention, their beliefs about treatment were considered important, so participants completed the Credibility/Expectancy Questionnaire (CEQ)27 after being allocated to an intervention group.

Outcome measures
Outcome measures were collected at 2, 6 and 12 weeks unless stated otherwise. The primary end-point was at 12 weeks. To minimise participant burden, a postal questionnaire was used for the assessment at 6 weeks.

Primary outcome measure
The primary outcome was the Victorian Institute of Sport Assessment-Achilles (VISA-A) questionnaire. The VISA-A is a reliable self-administered patient-reported outcome measure containing eight questions evaluating pain, function and activity.26 The maximum obtainable score is 100, with a lower score indicating greater clinical severity. Participants with bilateral symptoms completed the questionnaire on the most painful side (or right side, if Achilles tendons were equally painful).

Secondary outcome measures
Secondary outcome measures included the following.
1. Achilles tendon structure using ultrasound tissue characterisation (UTC) and anterior–posterior thickness at baseline and 12 weeks, as described previously.21 Tendon structure was quantified by the percentage of echo-types: aligned fibrillar structure (AFS; echo types I and II) and disorganised structure (DIS; echo types III and IV).29 30
2. Severity of pain (at worst in the previous week) measured on a 100 mm Visual Analogue Scale (VAS).31
3. Health status (using the EuroQol 5 Dimension 5 Level (EQ-5D-5L) questionnaire) was expressed for both of its components, the index value (scores range from <0 (worse than dead) to 0 (dead) to 1 (full health)) and the VAS value (scores range from 0 (worst imaginable health state) to 100 (best imaginable health state)).32
4. Magnitude of physical activity in the previous week (using the 7-day Recall Physical Activity Questionnaire).33
5. Calf muscle function at baseline and 12 weeks (using the standing heel rise test, assessed on flat ground).34 This test recorded the maximum number of repetitive unilateral weight-bearing raises (concentric-eccentric muscle action) that could be performed until fatigue, and was conducted to the sound of a metronome (one heel rise every 2 s with a metronome set at 60 beats per minute).
6. Self-reported treatment effectiveness for pain and function (using the Patient Impression of Change (PIC) questionnaire).31 This variable was then dichotomised into the categories of ‘effective’ (‘much improved’ or ‘very much im-
proven) and ‘ineffective’ (‘minimally improved’ to ‘very much worse’).35

7. Use of paracetamol rescue medication and cointerventions to relieve Achilles pain.

**Intervention adherence**
Adherence to the interventions was measured at 2, 6 and 12 weeks via questionnaires. Participants in the heel lifts group were asked to report the number of days and hours per day they had worn the heel lifts in the preceding period. Similarly, participants in the eccentric exercise group were asked to report the number of days per week they had performed the exercise programme in each week of the preceding period.

**Adverse events**
Adverse events were measured at 2, 6 and 12 weeks via questionnaires. Participants were asked to document the type of adverse event, the body location, the frequency and/or severity of the effect.36

**Randomisation and blinding**
To ensure allocation concealment, randomisation was provided by an independent computer (online) generated service (Griffith University, Queensland, Australia). The allocation for each participant was provided once eligibility was confirmed and baseline measures were completed. To ensure balance of participants in both groups, the random allocation sequence was generated with a 1:1 allocation using permuted blocks of random sizes (2, 4, 6). Due to the nature of the interventions, neither the participants, nor research staff administering the interventions (CLR and JMG), assessing outcomes (CLR), or entering data (CLR) were blinded to group allocation. However, data entry was independently validated, and research staff analysing data (CLR, HBM and JMG) were blinded to group allocation (masking used).

**Sample size**
An a priori sample size calculation estimated that a total of 92 participants (46 per group) would be needed to detect a difference between two groups, based on a power of 80%, minimal important difference (MID) of 10 points in the VISA-A.16 SD of 16.9,26 standardised effect size of 0.59 and an alpha of 0.05. We did not allow for participant loss to follow-up in our calculation as we used multiple imputation for missing data.38

**Data analysis**
SPSS V26 (IBM) was used for statistical analysis using the intention-to-treat principle. All data were explored for normality, and non-normally distributed data were transformed where possible. Data unable to be transformed were analysed using non-parametric tests. To avoid overtesting and reduce the risk of type I-associated errors, the end-point was the completion of the study (12 weeks). Multiple imputation was used to replace missing data using five iterations, with sex, age, baseline scores and group allocation used as predictors,38 for all primary and secondary outcome measures. The exception was for the variables; self-reported treatment effectiveness, use of cointerventions, adherence and adverse events, where data substitution was not applied. Continuously scaled measures were analysed using analysis of covariance (ANCOVA) with baseline scores and the intervention group as the independent variable.41 For dichotomous-scaled outcome measures, relative risk, risk difference and number needed to treat (NNT) were used for analysis.

Where appropriate, 95% CIs and p values were calculated to complement point estimates. The participants’ expectations (how much they expected they would benefit from the treatment) and credibility (beliefs about the logic of an intervention), measured using the CEQ at baseline, were compared between groups using the Mann-Whitney U (for non-normally distributed data) and independent t-tests (for normally distributed data).

**RESULTS**

**Participants**
Between September 2017 and December 2017, 368 participants contacted the researcher (CLR) for information regarding the trial. One hundred participants were enrolled into the study. Fifty participants were allocated to the heel lifts group and 50 participants were allocated to the eccentric exercise group. There were no clinically relevant differences in baseline characteristics between groups (table 1). Overall, 83, 73 and 80 participants completed the 2, 6 and 12 weeks follow-up assessments, respectively (figure 2).

**Primary outcome measure**
There was 80% follow-up of participants (n=40 per group) at 12 weeks. The mean VISA-A score improved by 26.0 points (95% CI 19.6 to 32.4) in the heel lifts group and by 17.4 points (95% CI 9.5 to 25.3) in the eccentric exercise group. On average, there was a between-group difference in favour of the heel lifts for the VISA-A (adjusted mean difference 9.6, 95% CI 1.8 to 17.4, p=0.016).

**Secondary outcome measures**
At 12 weeks, there were no statistically significant between-group differences in Achilles anterior–posterior diameter or Achilles tendon integrity (AFS and DIS) assessed using UTC (table 2). However, there was a statistically significant between-group difference for severity of pain (VAS) (adjusted mean difference −19.2, 95% CI −31.0 to −7.4, p=0.002) in favour of the heel lifts. There was also a statistically significant between-group difference for the EQ-5D-5L index score (adjusted mean difference 0.1, 95% CI 0.0 to 0.2, p=0.044) in favour of the heel lifts. There were no statistically significant between-group differences for EQ-5D-5L VAS, level of physical activity or calf muscle function. Participants in the heel lifts group were more likely to report an improvement in pain (RR 1.4, 95% CI 1.0 to 2.0, NNT 4) and function (RR 1.6, 95% CI 1.0 to 2.3, NNT 4), as measured using the PIC (online supplemental file 1).

**Treatment preference, credibility and expectancy**
Fifty-two per cent of participants in the heel lifts group and 64% of participants in the eccentric exercise group had a preference for the eccentric exercise programme (table 1). Participants in both groups considered their interventions to be credible and they expected to benefit from them (online supplemental file 2). There were no statistically significant between-group differences regarding the credibility and expectancy of treatment.

**Intervention adherence**
Table 3 shows the adherence at each time point for both treatment groups. Adherence was greater in the heel lifts group at all time periods.

**Use of rescue medication and cointerventions**
Overall, seven participants reported using cointerventions or rescue medication (paracetamol) during the study (online...
Table 1  Participant characteristics at baseline

<table>
<thead>
<tr>
<th></th>
<th>Heel lifts group (n=50)</th>
<th>Eccentric exercise group (n=50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demographics and anthropometrics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age, years</td>
<td>46.1 (9.0)</td>
<td>45.6 (9.8)</td>
</tr>
<tr>
<td>Female*</td>
<td>26 (52)</td>
<td>26 (52)</td>
</tr>
<tr>
<td>Height, cm</td>
<td>171.9 (9.1)</td>
<td>172.6 (10.7)</td>
</tr>
<tr>
<td>Weight, kg</td>
<td>85.6 (17.7)</td>
<td>89.9 (22.0)</td>
</tr>
<tr>
<td>Body mass index, kg/m²†</td>
<td>28.0 (24.3 to 31.8)</td>
<td>28.8 (26.3 to 31.1)</td>
</tr>
<tr>
<td>Waist to hip ratio</td>
<td>0.9 (0.1)</td>
<td>0.9 (0.1)</td>
</tr>
<tr>
<td>FPI score†</td>
<td>3 (2 to 4)</td>
<td>3 (0 to 4)</td>
</tr>
<tr>
<td>Supinated (FPI -12 to 0)*</td>
<td>10 (20)</td>
<td>15 (30)</td>
</tr>
<tr>
<td>Normal (FPI 1 to 7)*</td>
<td>40 (80)</td>
<td>34 (68)</td>
</tr>
<tr>
<td>Pronated (FPI 8 to 12)*</td>
<td>0 (0)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Ankle joint dorsiflexion range of motion, degrees‡</td>
<td>38.5 (6.5)</td>
<td>39.1 (7.8)</td>
</tr>
</tbody>
</table>

Self-reported education*
- Primary school completed 1 (2) 3 (6)
- High school (or equivalent) completed 14 (28) 7 (14)
- Vocational training 11 (22) 12 (24)
- College/university completed 15 (30) 15 (30)
- Postgraduate degree completed 9 (18) 12 (24)
- Don’t know 0 (0) 1 (2)

Self-reported medical conditions*
- Asthma 4 (8) 0 (0)
- Hypercholesterolaemia 0 (0) 2 (4)
- Hypertension 10 (20) 8 (16)
- Leg cramps 10 (20) 4 (8)
- Osteoarthritis 4 (8) 6 (12)
- Hands 1 (2) 2 (2)
- Feet 2 (4) 0 (0)
- Other 4 (8) 9 (18)

Self-reported medications*
- Uses at least 1 medication 20 (40) 28 (56)

Physical activity
- Partakes in physical activity* 43 (86) 41 (82)
- Average hours of physical activity per week, hour* 4.5 (1.9 to 7.7) 5.1 (0.9 to 8.2)

Footwear
- Women’s shoe size, USA† 8.0 (7.5 to 9.0) 8.0 (8.0 to 10.0)
- Men’s shoe size, USA† 10.0 (9.0 to 11.0) 10.5 (10.0 to 11.0)
- Athletic shoe worn most frequently* 20 (40) 21 (42)
- Rearfoot height, mm† 31.6 (26.0 to 37.0) 33.1 (26.8 to 36.9)
- Forefoot height, mm† 19.1 (13.0 to 23.4) 20.6 (12.9 to 24.1)
- Pitch, mm† 12.3 (8.4 to 15.8) 12.5 (9.7 to 16.7)
- Current use of foot orthoses* 6 (12) 12 (24)
- Current use of ankle brace* 1 (2) 1 (2)

Achilles tendinopathy characteristics
- Left/right/bilateral* 11 (22)/14 (28)/25 (50) 10 (20)/7 (14)/33 (66)
- Duration of symptoms, month† 18.0 (6.0 to 36.0) 22.5 (8.0 to 36.0)

Outcome measures
- VISA-A 57.0 (15.4) 53.3 (17.1)
- Achilles tendon AP diameter, mm 8.61 (1.88) 9.83 (2.12)
- Achilles tendon integrity: AFS, % 76.3 (17.6) 71.7 (19.6)
- Achilles tendon integrity: DIS, % 23.7 (17.6) 28.3 (19.6)
- Severity of pain (VAS), mm 56.1 (24.6) 60.1 (21.9)
- EQ-SD-5L, Index 0.76 (0.11) 0.73 (0.11)
- EQ-SD-5L, VAS 76.1 (12.7) 77.4 (15.4)

Self-reported Physical Activity Recall (PAR), kilocalories per day† 3493 (2916 to 4118) 3688 (3197 to 4497)

Table 1  Continued

<table>
<thead>
<tr>
<th></th>
<th>Heel lifts group (n=50)</th>
<th>Eccentric exercise group (n=50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calf muscle function, repetitions</td>
<td>18.9 (10.6)</td>
<td>17.9 (11.0)</td>
</tr>
</tbody>
</table>

Preferred treatment*
- Preferred HL/EE/impartial 19 (38)/26 (52)/5 (10)
- Received preferred intervention 19 (38) 32 (64)

EQ-SD-5L, EuroQol 5 dimension 5 level questionnaire was expressed for both of its components, the index value (scores range from <0 (worse than dead) to 1 (full health)) and the VAS value (scores range from 0 (worst imaginable health state) to 100 (best imaginable health state)). PAR questionnaire. Values are mean (SD) unless otherwise noted.

* Median (IQR), where IQR is expressed as Q1–Q3 (non-normally distributed data).
† Measured using the weight-bearing lunge technique. Details of the measurements can be found in the study protocol.21
* n (%).
§ Preferred treatment: one participant reported using a cointervention or rescue medication.
Achilles tendon integrity: AFS, anterior-posterior; DIS, disorganised structure; EE, eccentric exercise; FPI, Foot Posture Index; HL, heel lifts; VAS, Visual Analogue Scale; VISA-A, Victorian Institute of Sports Assessment-Achilles.

Figure 2  Flow of participants through the study. AT, Achilles tendon; NRS, Numerical Rating Scale.
exercises from a physiotherapist, one reported applying tape and one reported using a topical analgesic/non-steroidal anti-inflammatory gel.

Adverse events
There were no serious adverse events. The most common adverse events were pain in the knee, lower leg and foot and ankle. There were no statistically significant between-group differences in the number of participants reporting adverse events (table 4).

Sensitivity analysis
Sensitivity analysis was performed to assess the robustness of the findings for the primary outcome measure where multiple imputation was used to handle missing data. We used a series of five alternate approaches to handle missing data (online supplemental file 4). Results demonstrated that the magnitude of difference between groups for the VISA-A ranged from 6.5 to 10.0 points in favour of the heel lifts, with one of the five alternate approaches to handling missing data displaying a statistically significant between-group difference (10.0 points) that could be considered clinically meaningful (online supplemental file 4).

DISCUSSION
This randomised trial found heel lifts to be more effective in reducing pain and improving function in adults with mid-portion

Table 2 Primary and secondary outcome measures

<table>
<thead>
<tr>
<th>Outcome measure</th>
<th>Group</th>
<th>Adjusted mean difference*</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>VISA-A</td>
<td>Heel lifts*</td>
<td>57.0 (15.4)</td>
<td>53.3 (17.1)</td>
</tr>
<tr>
<td>Baseline</td>
<td>Eccentric exercise*</td>
<td>69.3 (14.1)</td>
<td>58.9 (19.0)</td>
</tr>
<tr>
<td>2 weeks</td>
<td>71.9 (16.8)</td>
<td>64.2 (20.3)</td>
<td>–</td>
</tr>
<tr>
<td>6 weeks</td>
<td>83.0 (16.9)</td>
<td>70.7 (22.2)</td>
<td>9.61 (1.83 to 17.40)</td>
</tr>
<tr>
<td>Achilles tendon AP diameter, mm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>8.61 (1.88)</td>
<td>9.83 (2.12)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>8.31 (2.23)</td>
<td>8.98 (2.32)</td>
<td>3.82 (−1.43 to 9.08)</td>
</tr>
<tr>
<td>Achilles tendon integrity: AFS, %†</td>
<td></td>
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<tr>
<td>Baseline</td>
<td>76.3 (17.6)</td>
<td>71.7 (19.6)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>74.1 (18.2)</td>
<td>69.7 (21.0)</td>
<td>3.41 (−6.34 to 13.14)</td>
</tr>
<tr>
<td>Achilles tendon integrity: DIS, %§</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>23.7 (17.6)</td>
<td>28.3 (19.6)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>26.9 (16.6)</td>
<td>30.0 (28.5)</td>
<td>−2.18 (−14.15 to 9.78)</td>
</tr>
<tr>
<td>Severity of pain (VAS), mm</td>
<td></td>
<td></td>
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<tr>
<td>Baseline</td>
<td>76.1 (24.6)</td>
<td>60.1 (21.9)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>40.5 (26.2)</td>
<td>49.3 (25.6)</td>
<td>–</td>
</tr>
<tr>
<td>EQ-5D-5L, Index</td>
<td></td>
<td></td>
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<tr>
<td>Baseline</td>
<td>0.76 (0.11)</td>
<td>0.73 (0.11)</td>
<td>–</td>
</tr>
<tr>
<td>2 weeks</td>
<td>0.81 (0.10)</td>
<td>0.77 (0.15)</td>
<td>–</td>
</tr>
<tr>
<td>6 weeks</td>
<td>0.79 (0.15)</td>
<td>0.76 (0.21)</td>
<td>–</td>
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<tr>
<td>EQ-5D-5L, VAS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>76.1 (12.7)</td>
<td>77.4 (15.4)</td>
<td>–</td>
</tr>
<tr>
<td>2 weeks</td>
<td>82.5 (10.8)</td>
<td>79.9 (14.8)</td>
<td>–</td>
</tr>
<tr>
<td>6 weeks</td>
<td>80.0 (14.3)</td>
<td>80.0 (17.9)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>84.9 (11.4)</td>
<td>82.8 (13.2)</td>
<td>−0.89 (−2.21 to 0.44)</td>
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<tr>
<td>Self-reported physical activity (PAR), kilocalories per day††</td>
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<td></td>
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<tr>
<td>Baseline</td>
<td>3440 (2916 to 4118)</td>
<td>3650 (3197 to 4497)</td>
<td>–</td>
</tr>
<tr>
<td>2 weeks</td>
<td>3230 (2662.9 to 4065)</td>
<td>3377 (3007 to 4526.2)</td>
<td>–</td>
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<tr>
<td>6 weeks</td>
<td>1846 (1669.1 to 2287)</td>
<td>1855 (1653.4 to 2348.1)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>1856.5 (1568 to 2232)</td>
<td>1908 (1558 to 2417)</td>
<td>–</td>
</tr>
<tr>
<td>Calf muscle function, repetitions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>18.9 (10.6)</td>
<td>17.9 (11.2)</td>
<td>–</td>
</tr>
<tr>
<td>12 weeks</td>
<td>20.1 (8.8)</td>
<td>21.6 (10.3)</td>
<td>−1.76 (−5.65 to 2.13)</td>
</tr>
</tbody>
</table>

EQ-5D-5L, EuroQol 5 dimension 5 level version questionnaire, (scores on the EQ-5D-5L index range from <0 (worse than dead) to 0 (dead) to 1 (full health); scores on the EQ-5D-5L VAS range from 0 (worst imaginable health state) to 100 (best imaginable health state)). PAR, Values are expressed as mean (SD) and postmultiple imputation unless stated otherwise. Bold values denote statistical significance at the p < 0.05 level.

*Values are presented prior to data transformation.
†Values in parentheses represent the 95% CI.
‡Data transformation was not applied.
§Values expressed postdata transformation (square root transformation applied to non-normally distributed data).
¶Values expressed postdata transformation (participant 16 removed (outlier) from non-normally distributed data).
**Values expressed postdata transformation (participant 83 removed (outlier) from non-normally distributed data).
††Values expressed as median (IQR), where IQR is expressed as Q1–Q3.
‡‡Mann-Whitney U test performed as non-normally distributed data was unable to be transformed.
AFS, aligned fibrillar structure; AP, anterior–posterior; DIS, disorganised structure; VAS, Visual Analogue Scale; VISA-A, Victorian Institute of Sports Assessment–Achilles.
Achilles tendinopathy. This is supported by findings for the primary outcome measure, the VISA-A questionnaire, where a 9.6 point between-group difference was observed in favour of the heel lifts intervention. However, it is unclear if the observed magnitude of difference in the VISA-A is large enough to be clinically meaningful, based on a priori MID of 10 points. The estimated MID for the VISA-A ranges between 6.5 and 20 points, with 10 points the most commonly reported.61,62

There were also several important findings for the secondary outcome measures at 12 weeks that favoured the heel lifts intervention, but the clinical importance of the observed differences remains uncertain. First, the severity of pain (measured using aVAS) was significantly less in the heel lifts group, although it is unclear if the magnitude of difference (19 points) is clinically meaningful for mid-portion Achilles tendinopathy.46 Second, self-reported treatment effectiveness for both pain and function, measured using the PIC, was significantly greater in participants allocated the heel lifts, with the NNT indicating that four participants would need to be treated with heel lifts to achieve one additional successful outcome. However, the CIs around these values were wide. Third, health-related quality of life (measured using the EQ-5D-5L Index) favoured the heel lifts, but the clinical importance of the observed difference (0.10) is not clear.47

**Comparison of findings with previous research**

This is the first randomised trial to compare the efficacy of heel lifts to calf muscle eccentric exercise for mid-portion Achilles tendinopathy. Consequently, there are no previous trials with which to compare these findings to. In our study, the mean within-group improvement in VISA-A at 12 weeks was 17.4 points for the eccentric exercise group. The magnitude of improvement is lower than what was reported in a recent systematic review with meta-analysis, where the mean within-group improvement at 12 weeks in VISA-A following an eccentric exercise programme was reported to be 21 points.49 However, there was variability in response across included studies, and the authors concluded that the improvement was likely to have been overestimated. Therefore, we believe that our findings regarding the effect of the eccentric exercise programme are valid.

**Strengths and limitations**

This randomised trial was pragmatically designed, incorporated rigorous methodology and included two groups with similar baseline characteristics.

Potential confounding factors were controlled for with the use of appropriate participant randomisation and use of ANCOVA statistics for data analysis. Bias was minimised with the use of allocation concealment, double checking of data entry and blinded data analysis. An appropriate sample size to ensure adequate statistical power and the use of diagnostic imaging to confirm mid-portion Achilles tendinopathy of study participants strengthened this trial.

The participants of this study were representative of individuals with Achilles tendinopathy that present to healthcare providers, based on age and sex distribution, although the duration of symptoms of participants of this study (mean 34.4 months) was far greater than previously reported (mean 2.6 and 7.4 months).49,50 Further studies are required to test these findings in sub-groups such as younger or highly active individuals. We also used interventions that are readily available and frequently prescribed in clinical practice.

Our results should be interpreted in context of inherent study design limitations. Due to the nature of the interventions, participants and research staff administering the intervention could not be blinded. However, the eccentric exercise programme was the preferred intervention in both groups, and there were no between-group differences in treatment credibility or expectations. Adherence was also self-reported and may not truly reflect actual adherence. Finally, the study duration was 12 weeks, so it remains unknown if the observed effects persist beyond this point.

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**Table 3** Self-reported adherence

<table>
<thead>
<tr>
<th>Time period</th>
<th>Heel lifts group</th>
<th>Eccentric exercise group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Days†</td>
<td>%</td>
</tr>
<tr>
<td>1–2</td>
<td>7 (6 to 7)</td>
<td>94</td>
</tr>
<tr>
<td>3–6</td>
<td>7 (7 to 7)</td>
<td>87</td>
</tr>
<tr>
<td>7–12</td>
<td>7 (6 to 7)</td>
<td>91</td>
</tr>
</tbody>
</table>

Days refers to the number of days per week the intervention was reported to be used over the time period.

*Percentage adherence was calculated as follows. For the heel lifts group, the number of days the heel lifts were worn was divided by 7 (ie, the maximum possible number of days per week the heel lifts could be worn) and multiplied by 100. For the eccentric exercise group, the number of days the eccentric exercise programme was performed was divided by 7 (ie, the maximum possible number of days per week the eccentric exercise programme could be performed) and multiplied by 100.

†Values are expressed as median (IQR), where IQR is expressed as Q1–Q3.

**Table 4** Self-reported adverse events

<table>
<thead>
<tr>
<th>Adverse event*</th>
<th>Heel lifts group (n=50)†</th>
<th>Eccentric exercise group (n=50)†</th>
<th>RR†</th>
<th>P value</th>
<th>ARI (%)</th>
<th>NNT-H‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Developed new pain</td>
<td>23 (46)</td>
<td>22 (44)</td>
<td>1.05 (0.68 to 1.61)</td>
<td>0.841</td>
<td>2.0 (−16.9 to 20.7)</td>
<td>50 (−6 to 5)</td>
</tr>
<tr>
<td>Upper body</td>
<td>0 (0)</td>
<td>3 (6)</td>
<td>UTD</td>
<td>UTD</td>
<td>−6.0 (−16.2 to 2.2)</td>
<td>−17 (−6 to 47)</td>
</tr>
<tr>
<td>Lower back</td>
<td>6 (12)</td>
<td>3 (6)</td>
<td>2.00 (0.53 to 7.56)</td>
<td>0.307</td>
<td>6.0 (−6.1 to 18.4)</td>
<td>17 (−17 to 5)</td>
</tr>
<tr>
<td>Hip</td>
<td>3 (6)</td>
<td>2 (4)</td>
<td>1.50 (0.26 to 8.60)</td>
<td>0.659</td>
<td>2.0 (−8.3 to 12.6)</td>
<td>50 (−12 to 8)</td>
</tr>
<tr>
<td>Knee</td>
<td>6 (12)</td>
<td>7 (14)</td>
<td>0.86 (0.31 to 2.37)</td>
<td>0.767</td>
<td>−2.0 (−15.8 to 11.8)</td>
<td>−50 (−6 to 9)</td>
</tr>
<tr>
<td>Lower leg</td>
<td>8 (16)</td>
<td>9 (18)</td>
<td>0.89 (0.37 to 2.11)</td>
<td>0.790</td>
<td>−2.0 (−16.9 to 13.0)</td>
<td>−50 (−6 to 8)</td>
</tr>
<tr>
<td>Foot or ankle§</td>
<td>7 (14)</td>
<td>9 (18)</td>
<td>0.78 (0.31 to 1.93)</td>
<td>0.587</td>
<td>−4.0 (−18.6 to 10.7)</td>
<td>−25 (−5 to 9)</td>
</tr>
<tr>
<td>Blister/skin condition</td>
<td>2 (4)</td>
<td>1 (2)</td>
<td>2.00 (0.19 to 21.36)</td>
<td>0.566</td>
<td>2.0 (−7.0 to 11.6)</td>
<td>50 (−14 to 9)</td>
</tr>
</tbody>
</table>

Some participants reported more than one adverse event. The heel lifts group was considered the experimental group for all calculations.

*Values are the number of participants who reported at least one adverse event.

†Values are expressed as n (%) for each group.

‡Values in parentheses represent the 95% CI.

§Excludes skin conditions.

ARI, absolute risk increase; NNT-H, number needed to harm; RR, relative risk; UTD, unable to determine.
Benefits. Potential disadvantages of heel lifts include costs, ability to use across multiple shoe styles and supply availability.

CONCLUSION
In this study, heel lifts were more effective than eccentric exercise in reducing pain and improving function at 12 weeks in adults with mid-portion Achilles tendinopathy. Further studies are required to determine if this difference is large enough to be clinically meaningful. Healthcare providers can consider heel lifts as a first-line treatment for mid-portion Achilles tendinopathy.

What are the findings?

► In-shoe heel lifts are used for mid-portion Achilles tendinopathy, yet evidence from high-quality randomised controlled trials has been lacking.

► We found that heel lifts were effective, yet it remains unknown if heel lifts result in an improvement considered important by patients above the current first-line treatment, eccentric exercise.

How might it impact on clinical practice in the future?

► Healthcare providers can consider heel lifts as a first-line treatment for mid-portion Achilles tendinopathy.

REFERENCES
Original research


