Sensor-based gait training to reduce contact time for runners with exercise-related lower leg pain: a randomised controlled trial

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ABSTRACT

Objectives To assess the effects of a 4-week randomised controlled trial comparing an outdoor gait-training programme to reduce contact time in conjunction with home exercises (contact time gait-training feedback with home exercises (FBHE)) to home exercises (HEs) alone for runners with exercise-related lower leg pain on sensor-derived biomechanics and patient-reported outcomes.

Design Randomised controlled trial.

Setting Laboratory and field-based study.

Participants 20 runners with exercise-related lower leg pain were randomly allocated into FBHE (4 male (M), 6 female (F), 23.4 ± 4 years, 22.0 ± 4.3 kg/m²) or HE groups (3 M, 7 F, 25 ± 5 years, 23.6 ± 3.9 kg/m²).

Interventions Both groups completed eight sessions of HEs over 4 weeks. The FBHE group received vibrotactile feedback through wearable sensors to reduce contact time during outdoor running.

Primary and secondary outcome measures Patient-reported outcome measures (PROMs) and outdoor gait assessments were conducted for both groups at baseline and 4 weeks. PROMs were repeated at 6 weeks, and feedback retention was assessed at 6 weeks for the FBHE group.

Results The FBHE group reported increased function and recovery on PROMs beyond the HE group at 6 weeks (p<0.001). There was a significant group by time interaction for Global Rating of Change (p=0.004) and contact time (p=0.002); the FBHE group reported greater subjective improvement and reduced contact time at 4 and 6 weeks compared with the HE group and compared with baseline. The FBHE group had increased cadence (mean difference: 7 steps/min, p=0.01) at 4 weeks during outdoor running compared with baseline.

Conclusion FBHE was more effective than HE alone for runners with exercise-related lower leg pain, manifested with improved PROMs, reduced contact time and increased cadence.

Trial registration number NCT04270565.

INTRODUCTION

Lower limb injuries constitute up to 50% running-related injuries, and recent literature has advocated using ‘exercise-related lower leg pain’ as the preferred nomenclature when other injuries can be ruled out with clinical examinations. Given the burden exercise-related lower leg pain imposes on runners, recent research has assessed contributing factors to injury development to guide interventions. However, the only recommended care for runners with exercise-related lower leg pain is to perform calf stretching. Rehabilitation is often included in clinical practice and should be considered for exercise-related lower leg pain management, especially as recent work has identified hip and ankle muscle weakness among these patients. Previous research that incorporated a strengthening programme for injured runners successfully reduced patients’ pain and improved patient-reported outcome measures (PROMs). These studies may be used as a framework for developing interventions for runners with exercise-related lower leg pain.

WHAT IS ALREADY KNOWN ON THIS TOPIC

Runners with exercise-related lower leg pain have been found to present with altered gait biomechanics during outdoor running.

WHAT THIS STUDY ADDS

Outdoor gait training with standard of care home exercises (HEs) was more effective than HEs alone on improving self-reported pain and function and on movement patterns during outdoor running.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

Clinicians may consider implementing outdoor-based feedback to improve running biomechanics. Future research is needed to establish the benefit of this modality in a larger, representative sample.
In addition to strength and motion deficits, recent laboratory-based gait analyses have identified increased peak rearfoot eversion during stance,\textsuperscript{8,9} longer stride length,\textsuperscript{10} and slower cadence among runners with lower extremity injuries or active symptoms.\textsuperscript{10} Additionally, runners with a history of lower extremity injuries have been found to present with increased vertical impact peaks and average loading rates.\textsuperscript{11} Although gait-training programmes targeting these factors have been successful,\textsuperscript{12} these interventions have been primarily limited to indoor settings among healthy runners. While several studies have implemented outdoor gait training for healthy runners and runners with tibial stress fractures,\textsuperscript{13,14} more evidence is necessary to support our understanding of treatment success among runners actively experiencing lower limb pain in natural running environments.\textsuperscript{12} Outdoor running assessments implementing wearable technology have identified increased and more variable contact time as the key factor differentiating runners with exercise-related lower leg pain from healthy counterparts.\textsuperscript{6,15} While previous work has not identified a difference in contact time between injured and healthy runners,\textsuperscript{16} this may be attributed to supervised, indoor running that is distinct from typical bouts of outdoor running. Additionally, while longer contact time without significantly different cadence has been found to be associated with lower peak vertical ground reaction forces and higher duty factors among marathoners compared with a control group,\textsuperscript{13} longer contact time with concomitantly slow cadence may be associated with a longer epoch of loading exposure imposed on lower extremity structures.\textsuperscript{18} Based on past outdoor assessments, it is surmised that this longer overall loading contributes to the cumulative stress imposed on the lower limb. As such, longer contact time may be a key contributing factor to exercise-related lower limb pain.

The purpose of this study was to assess the effects of 4 weeks of outdoor gait training to reduce contact time gait-training feedback with home exercise (FBHE) compared with home exercises (HEs) alone for runners with exercise-related lower leg pain. The authors compared groups and timepoints on PROMs and sensor-derived running biomechanics over the 4-week intervention period. PROMs were repeated for both groups at 6 weeks, and feedback retention was assessed for the FBHE group alone at 6 weeks. It was hypothesised that the FBHE group would demonstrate reduced pain and increased function at 4 and 6 weeks compared with baseline and the HE group, and decreased sensor-derived contact time, increased cadence and decreased loading at 4 weeks compared with baseline and the HE group. It was also anticipated that the FBHE group would retain sensor-derived biomechanical changes at 6 weeks.

\textbf{METHODS}

The Consolidated Standards of Reporting Trials flowchart outlining the randomised controlled trial study procedures is presented in figure 1.

\textbf{Participants}

Participants were recruited between February 2020 and May 2021 (end of the academic semester) through our local university and surrounding community. Participants were required to be 18–45 years of age, involved in running at least three times per week for the past 3 months, and report pain between 20 mm and 80 mm on the 100 mm Visual Analogue Scale (VAS) during or following running in the lower leg for ≥1 month, confirmed using a clinical assessment.\textsuperscript{19,21} Participants had to score <90% on the Exercise-Induced Leg Pain Questionnaire, British Version (EILP-Br).\textsuperscript{3,19,20,22} The Exercise-Induced Leg Pain Questionnaire has been found to have excellent internal consistency (intraclass correlation coefficient (ICC): 0.92–0.94) and test–retest reliability (ICC: 0.987–0.995) across patients with exercise-related lower leg pain.\textsuperscript{3,22} Exclusion criteria included pain over the Achilles tendon, popliteal fossa or the superficial posterior compartment of the lower leg, medical diagnoses of compartment syndrome, tibial or fibular stress or full fractures within 3 months.\textsuperscript{3,22} These injuries were exclusionary as these diagnoses would prohibit individuals from completing running due to bone or neurovascular compromise. Participants additionally could not have other pathologies or surgeries, or known pregnancy. The study was approved by our University’s Institutional Review Board for Health Sciences Research (IRB-HSR 22107) and registered as a clinical trial. All participants provided informed consent prior to study procedures.

\textbf{Sample size estimation}

While we originally aimed to recruit 20 participants per group to achieve 80% power and <15% attrition, the global pandemic resulted in resource constraints hindering our sample size.\textsuperscript{23} Based on our available sample and outcomes from our primary variable of interest, the false-positive risk based on the prior probability of 0.5 was 7%.\textsuperscript{24}

\textbf{Patient-reported outcome measures}

In addition to the 100 mm VAS and EILP-Br questionnaires completed during screening, participants completed a running history questionnaire (weekly mileage, number of running days per week, years of experience and pace), Wisconsin Running Injury and Recovery Index\textsuperscript{25} and Lower Extremity Functional Scale (LEFS; table 1) at baseline. The Global Rating of Change (GROC) scale was used to gauge recovery throughout the programme at study timepoints of 2, 4 and 6 weeks.\textsuperscript{26} Each questionnaire included in this study have demonstrated fair to excellent construct validity (LEFS: r range=0.73–0.8,\textsuperscript{27} Wisconsin Index: r range=0.67–0.75,\textsuperscript{28} face validity (GROC: r range=0.72–0.90)\textsuperscript{29} and excellent test–retest
reliability (LEFS: R=0.94, \textsuperscript{27} GROC: ICC=0.90\textsuperscript{29} and Wisconsin Index: ICC=0.934).\textsuperscript{25}

**Patient and/or public involvement**

Four physical therapists (two dual-credentialed as athletic trainers) that had expertise in treating injured runners were involved in designing the functional movement assessments and rehabilitation plans prior to study initiation.

**Procedures**

**Baseline visit: clinical assessments**

Participants reported for a baseline visit at the university research laboratory. Clinical assessments were performed by a blinded athletic trainer with at least 2 years of clinical experience (SLS and XDT) or a trained laboratory assistant with 2 years of laboratory experience (PNF). Lower extremity alignment and range of motion were...
measured using a standard goniometer, and strength was assessed using a hand-held dynamometer. All measurements followed standard patient positioning, anatomical landmarks and equipment based on well-established and published procedures (online supplemental file 1).30–32

Participants completed three functional movements: Y-Balance Test reach distances,33 lateral step-downs from a 15 cm stair,34 and single-leg squats to 45° knee flexion.35 The functional movements were classified by the same blinded assessor into one of the following profiles: (1) medial knee displacement and/or ipsilateral hip drop with or without contralateral trunk lean (valgus), (2) lateral patellar displacement and/or contralateral hip hike with or without ipsilateral trunk lean (varus) and (3) neutral. If participants had one or more movement profile in the valgus or varus groups, they were categorised accordingly. If these adaptations were not present, participants were categorised into the ‘neutral’ group.

Clinical measurements and assessment scores were used to delineate specific HE plans, which were developed by an expert panel of physical therapists with expertise in treating injured runners (online supplemental file 2).

Baseline visit: outdoor running

RunScribe Plus sensors (RunScribe Labs, Half Moon Bay, California, USA) consisting of a triaxial accelerometer, magnetometer and gyroscope were used to collect outdoor running biomechanics (contact time, cadence, pace, stride length, shock (composite score combining impact, or vertical, and braking, or horizontal, force vectors), and pronation excursion and velocity) at a 200 Hz sampling rate, with on-board processing and memory. The sensors have demonstrated fair (pronation excursion ICC: 0.57) to excellent (contact time ICC: 0.93) validity against gold standard three-dimensional motion capture systems.36 37

Participants were issued a set of sensors, instructed on proper usage, and downloaded the associated mobile application. Participants mounted the sensors on their shoelaces and ran on a predetermined 2688 m route for calibration and baseline outdoor running assessment. Participants were asked to run with the sensors two times per week during sustained runs (≥2 miles) over 4 weeks and to record their pain level during runs in the mobile application (0–10).

Baseline visit: group allocation

Clinicians blinded to participant group allocation were dismissed. A random-number generator was used by an investigator who was not involved in patient interactions (JH) to determine the randomisation sequence. Group assignments were placed in sealed opaque envelopes and opened following baseline measures by the clinician administering the feedback. Participants allocated to the HE group were provided a list of their specific HEs with video demonstrations to be completed two times per week over the study period (online supplemental file 3).

| Table 1 | Participant demographics at baseline for FBHE and HE groups |
|-----------------------------------------------|
| | FBHE | HE |
| | n=9; 3 M, 6 F | n=10; 3 M, 7 F |
| | (mean±SD) | (mean±SD) | P value |
| Age (years) | 23±4 | 25±5 | 0.48 |
| Height (cm) | 168±12 | 167±8 | 0.84 |
| BMI (kg/m²) | 22.0±4.3 | 23.6±3.9 | 0.42 |
| Running experience (years) | 6±5 | 5±3 | 0.68 |
| Weekly mileage (km) | 24±18 | 24±19 | 0.97 |
| Average running pace (min/km) | 5:57±1:07 | 5:29±0:39 | 0.75 |
| Shoe mileage | 160±135 | 145±129 | 0.81 |

Note: the participant who dropped out of the study was excluded from the FBHE group.

Heat maps generated based on where patients indicated they experienced pain at baseline. Areas with warmer colours indicate higher density of selected problem areas, while cooler colours indicate lower density of selected problem areas.

*Significant at p≤0.05.

BMI, body mass index; F, female; FBHE, contact time gait-training feedback with home exercise; HE, home exercise; M, male.
Participants were instructed to record exercise compliance in their RunScribe mobile applications.

Participants allocated to the FBHE group received the same HE instructions and were additionally issued a Garmin Forerunner 235 wristwatch (FR235; Garmin Corporation, Olathe, Kansas, USA). The wristwatches were solely used to facilitate gait-training feedback and display the contact time metric in real time and were not used for monitoring purposes. The unblinded researcher manually set a 5% reduction of each participant’s baseline outdoor run average contact time onto the Garmin wristwatch using custom code. The 5% threshold was determined from previous findings among runners with exercise-related lower leg pain. FBHE participants were oriented to the gait-training procedures on the indoor treadmill; they received a vibration of three quick, successive pulses that were intermittently delivered every 125 ms from the watch if the contact time on the RunScribe sensors exceeded the threshold and were sequentially repeated until the contact time fell below the threshold. FBHE runners were instructed to shorten their contact time to reduce the vibration; however, they were not provided further cues. Once participants indicated they were comfortable with the feedback procedures to be completed two times per week, they were dismissed.

Weekly check-ins
Participants completed virtual weekly check-ins due to the COVID-19 pandemic to determine HE compliance and adjust exercises as needed (see online supplemental file 3 for specific criteria-based progressions). At the 2-week timepoint, all participants completed the Wisconsin Running Injury and Recovery Index and the GROC scale, and repeated the 100 mm VAS. The feedback programme was faded for the FBHE group by using the feedback for 50% of their runs for the final 2 weeks (ie, 15 min of a 30 min run).

Follow-up procedures
Participants returned to the laboratory at 4 weeks to repeat outdoor gait assessments, which were completed without feedback for the FBHE group. No further instructions were provided to participants to avoid any potential influence on retention outcomes.

Participants were contacted 2 weeks later (6-week timepoint) to repeat all PROM questionnaires. The FBHE group also repeated the outdoor gait assessment without feedback on the calibration route to assess gait-training retention.

Data processing
Outdoor running biomechanics
Sensor-derived biomechanics were calculated on-board through a proprietary software into the specific spatio-temporal (contact time, cadence, stride pace and stride length), kinetic (shock) and kinematic variables (pronation excursion and maximum pronation velocity). Operational definitions of all sensor-derived outcomes have been published elsewhere. Step-by-step data from each run were extracted from the manufacturer’s dashboard, and averages were taken per limb for each recorded run. Walking and standing events were visually identified in the datasets from when the flight ratio variable fell to 0 and were removed from analyses.

Statistical analyses
Descriptive analyses were conducted using independent samples t-tests to compare baseline age, height, body mass index, questionnaire scores and running experience between groups. We additionally compared running volume at baseline and cumulative distance accrued during the study time frame. Separate 2×4 repeated measures analyses of variance (RMANOVAs) were used to assess the influence of groups (FBHE, and HE) and timepoints (baseline and 4 and 6 weeks) for PROMs. Additionally, separate discrete measures 2×2 RMANOVAs were used to assess the influence of group (FBHE and HE) and timepoints (baseline and 4 weeks) for sensor-derived outdoor running biomechanics. A one-factor RMANOVA was used to assess gait-training retention for FBHE group across three timepoints (baseline and 4 and 6 weeks). All RMANOVA assessments were conducted in using RStudio V.1.2.1335. Alpha was set a priori to .05 for all analyses, and Tukey’s post hoc analyses were used for statistically significant findings.

RESULTS
Groups did not significantly differ at baseline for demographic factors or for cumulative distance accrued across the intervention programme (table 1). One FBHE participant was lost to follow-up due to an unrelated shoulder injury, and there were no adverse events pertaining to this study. Intention-to-treat analyses were not possible as there were no follow-up data. As such, 19 participants (FBHE: 9 and HE: 10) were included in PROM and outdoor gait assessment analyses (figure 1). Compliance with the HE programmes was excellent for both groups (FBHE: 96% and HE: 97%).

PROM results
There were significant time main effects for VAS pain scores. Both groups significantly decreased pain measures at timepoints of 4 and 6 weeks compared with baseline and 2 weeks (table 2). The FBHE group maximum pain change score at 6 weeks was clinically meaningful at −36 mm (CI −55 to −11 mm, d=1.75; 66% of patients reached a minimally clinically important difference of 30 mm), while the HE group improved by only −10 mm (CI −46 to 6 mm, d=0.89; 40% of patients met minimally clinically important difference; table 2).

There were significant group and time main effects for the Wisconsin Running Injury and Recovery Index and EILP-Br and a significant time main effect for the LEFS questionnaire (table 2). There was a significant group by time interaction for the GROC scores (table 2). While both groups reported increased function at follow-ups
of 4 and 6 weeks, the FBHE group reported significantly increased function and recovery than the HE group at 6 weeks and compared with baseline (table 2).

**DISCUSSION**

We determined an added benefit of outdoor gait training to reduce contact time for runners with exercise-related lower leg pain in conjunction with standard care exercises to improve exercise-related lower leg pain patient management. Our findings support the usage of this gait-training approach in conjunction with standard care exercises to improve exercise-related lower leg pain patient management.

**Patient-reported outcome measures**

We support not only that increased contact time is a consequence of exercise-related lower leg pain, but also that contributing to the overall exercise-related lower leg pain disability model, given that the FBHE intervention led to greater improvements in pain and function over time. This information is important for clinicians at baseline compared with 2 weeks after treatment. 40

While patients reported minimal residual pain at 6 weeks, 67% of FBHE patients compared with 40% of HE patients fell below 20 mm on the VAS, which would no longer classify many runners as patients with exercise-related lower leg pain. Our findings support the usage of this gait-training approach in conjunction with standard care exercises to improve exercise-related lower leg pain patient management.

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schema slightly differed from previous study designs as we provided 100% feedback over consistent run times and distances, whereas previous studies gradually built the time of interventions up over the first half of the intervention.43 44 However, we pre-emptively decided on this schedule due to the field-based nature of the intervention. Our intervention still implemented concepts of motor learning theory in which a stimulus is introduced and then gradually removed over time, and we found this overall model to lead to desirable biomechanical changes. The FBHE group surpassed the prescribed contact time feedback at 4 weeks (8.22%) and retained this change at 6 weeks (−0.69% change from 4 weeks), suggesting that patients were able to effectively incorporate the biomechanical adjustment into their motor learning framework to eliminate the feedback stimulus.45 FBHE patients concomitantly increased cadence at 4 and 6 weeks; given that cadence has been identified as a risk factor for RRI in laboratory analyses,9 42 targeting contact time may have a desired effect across affected spatiotemporal parameters for RRI treatment. Given that we only assessed retention for the FBHE group, the long-term effects of the gait-intervention training should be interpreted with some caution as we were unable to repeat these measures and compare against the standard of care patients in the HE group.

Clinical implications and future directions
Our findings support the use of a data-driven, ecological approach to gait training. While clinicians may not have access to extensive gait analysis equipment, commercially-available sensors to prescribe interventions alleviate cost, time, and resource burdens. Future work should seek to replicate this gait-training approach in natural running settings among patients with exercise-related lower leg pain in larger sample sizes. Additionally, future work should consider adopting a specific, evidence-based outdoor gait training programme for patients with other running-related injuries due to mounting accessibility of wearable sensors and growing importance of biometrics.

Table 3 Sensor-derived biomechanical measures between FBHE and HE groups across study timepoints

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>4 weeks</th>
<th>6 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean±SD</td>
<td>Between-group P value</td>
<td>Mean±SD</td>
</tr>
<tr>
<td>Contact time (ms)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>292±34</td>
<td>0.75</td>
<td>268±18*</td>
</tr>
<tr>
<td>HE</td>
<td>288±24</td>
<td></td>
<td>286±19</td>
</tr>
<tr>
<td>Cadence (steps/min)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>170±9</td>
<td>0.24</td>
<td>182±10*</td>
</tr>
<tr>
<td>HE</td>
<td>169±10</td>
<td></td>
<td>170±9</td>
</tr>
<tr>
<td>Pace (m/s)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>3.19±0.43</td>
<td>0.76</td>
<td>3.39±0.49</td>
</tr>
<tr>
<td>HE</td>
<td>3.25±0.41</td>
<td></td>
<td>3.35±0.45</td>
</tr>
<tr>
<td>Stride length (m)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>2.20±0.34</td>
<td>0.49</td>
<td>2.24±0.36</td>
</tr>
<tr>
<td>HE</td>
<td>2.30±0.31</td>
<td></td>
<td>2.37±0.32</td>
</tr>
<tr>
<td>Shock (g)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>13.2±2.1</td>
<td>0.44</td>
<td>13.0±1.7</td>
</tr>
<tr>
<td>HE</td>
<td>14.0±2.5</td>
<td></td>
<td>13.6±2.5</td>
</tr>
<tr>
<td>Pronation excursion (°)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>11.7±4.7</td>
<td>0.54</td>
<td>12.7±5.6</td>
</tr>
<tr>
<td>HE</td>
<td>10.5±4.3</td>
<td></td>
<td>13.3±5.0</td>
</tr>
<tr>
<td>Maximum pronation velocity (°/s)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FBHE</td>
<td>842±288</td>
<td>0.49</td>
<td>855±252</td>
</tr>
<tr>
<td>HE</td>
<td>757±245</td>
<td></td>
<td>746±264</td>
</tr>
<tr>
<td>Foot strike type (1–16)</td>
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<td></td>
<td></td>
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<tr>
<td>FBHE</td>
<td>7±2</td>
<td>0.93</td>
<td>8±2</td>
</tr>
<tr>
<td>HE</td>
<td>7±3</td>
<td></td>
<td>8±4</td>
</tr>
</tbody>
</table>

*Statistically significant compared with baseline at p≤0.05.
†Statistically significant differences between groups at p≤0.05.
FBHE, contact time gait-training feedback with home exercise; HE, home exercise.
in patient care. Finally, there is a need to compare the efficacy of clinic-based supervised gait-training interventions, field-based supervised gait-training interventions and unsupervised gait-training interventions to determine what the optimal implementation to elicit desired patient-centred outcomes and biomechanical changes. We hope that this study framework will set the precedent of specific patient care by meeting runners in their training environment and addressing runners’ specific RRI deficits.

Limitations

Our overall sample size was small due to limitations of human subjects research during the COVID-19 pandemic. As such, our results should be interpreted with some caution, and future research should seek to replicate these methods in a larger sample. Outdoor gait training limited control over external running factors (ie, running surface and environment); however, external validity of the intervention was increased due to this decision. The feedback intervention is currently not commercially available and requires some technical expertise. HEs were individualised, meaning exercise prescriptions varied by patient. This approach was designed with clinicians currently treating injured runners, which strengthens this decision. There was a relatively short follow-up period, and longer-term follow-ups are needed to assess retention length. Furthermore, we did not compare the FBHE group to the HE group for outdoor biomechanics at 6 weeks, and it is unlikely but theoretically possible that there was a time effect that drove the biomechanical findings. Future work comparing groups for long-term outcomes is warranted.

CONCLUSION

Outdoor gait training along with HEs was more effective than HEs alone for runners with exercise-related lower leg pain by improving PROMs, and influencing contact time and cadence at 4 weeks, and with lasting effects at 6 weeks. Clinicians may consider implementing this gait-training approach for runners with exercise-related lower leg pain to improve clinical management.

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Acknowledgements We thank Han Yu Wang and Grady Roberts for their assistance in designing the feedback protocol, and Drs Eric Magrum, Laura Hodges-Long, Kevin Cross and Michael Higgins for their assistance in creating the rehabilitation protocols.

Contributors AFDL: conceptualisation, methodology, validation, formal analysis, investigation, resources, data curation, writing (original draft), visualisation, project administration, and guarantor. SLS, PNF and XT: investigation, data curation and writing (review and editing). JMHE and DJH: conceptualisation, methodology, writing (review and editing) and supervision. JSR: conceptualisation, data curation, formal analysis, writing (review and editing) and supervision. JH: conceptualisation, methodology, formal analysis, resources, writing (review and editing) and supervision.

Funding This study was funded by the Mid-Atlantic Athletic Trainer’s Association Graduate Student Grant, the National Athletic Trainers’ Association Doctoral Dissertation Grant, and a university doctoral dissertation grant award.

Competing interests We received grant funding for this project from the following sources: (1) Mid-Atlantic Athletic Trainer’s Association Graduate Student Grant (grant number not available), (2) National Athletic Trainers’ Association Doctoral Dissertation Grant (grant number 1920DGP01) and (3) University of Virginia School of Education and Human Development IDEAs Grant (grant number not available).

Patient and public involvement Patients and/or the public were involved in the design, conduct, reporting or dissemination plans of this research. Refer to the Methods section for further details.

Patient consent for publication Consent obtained directly from patient(s).

Ethics approval This study involves human participants and was approved by University of Virginia Institutional Review Board for Human Subjects Research (#22107). The participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement No data are available. No data are available per stipulations by the University of Virginia Institutional Review Board.

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