Development And Early Feasibility Testing Of A Mind-Body Physical Activity Program For Patients With Heterogeneous Chronic Pain; The GetActive Study

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Background: Increasing physical function is a challenging, yet imperative goal of pain management programs. Physical activity can improve physical function, but uptake is low due to chronic pain misconceptions, poor pain management skills, and doing too much too soon.

Purpose: To increase physical function by 1) adapting an evidence-based, group, mind-body program to address the needs of patients with heterogeneous chronic pain and to facilitate individually tailored quota-based pacing with a Fitbit (GetActive with Fitbit) or without it (GetActive) (phase 1), and 2) assessing preliminary feasibility benchmarks (phase 2).

Methods: We followed evidence based frameworks for developing interventions and for early feasibility testing. In phase 1 we conducted 4 focus groups with 22 patients with heterogeneous chronic pain and adapted the mind-body program. In phase 2 we conducted a nonrandomized pilot trial of the 2 programs (N=7 and 6) with qualitative exit interviews.

Results: Focus groups showed high interest in increasing activity, a preference for walking linked to pleasurable activities, using a Fitbit to track number of steps, and learning skills to manage pain and aid with increased activity. Both programs had good to excellent feasibility markers. Participation in both programs was associated with signal of improvements in physical and emotional function, as well as intervention targets. Exit interviews confirmed high satisfaction and suggested modification.

Conclusion: Results informed subsequent adaptations of the 2 programs and methodology for an ongoing pilot randomized controlled trial (RCT) of the 2 programs, necessary before an efficacy RCT of the 2 programs against an education control.

Keywords: mind-body, chronic pain, fitbit, physical activity, focus groups, feasibility, physical function

Introduction

Chronic pain, defined as persistent pain that lasts more than 3–6 months,1 is prevalent,2 costly,2 and associated with decreased physical and emotional function.3 Over the last decade, mind-body (e.g. meditation and mindful-movement based) programs have been increasingly popular among patients with chronic pain, and have been shown to lead to improvements in anxiety, depression, and acceptance of symptoms.4 However, effect sizes for improvements in self-reported emotional4–6 and physical functioning6–8 have been small to moderate, and tended to dissipate...
over time. Existent mind-body interventions do not incorporate systematic gains in physical activity despite its positive effects on emotional and physical function. Thus, there is a need for novel interventions that incorporate mind-body and physical activity skills in order to sustainably and substantially improve emotional and physical function among patients with chronic pain. The importance of comprehensive assessment of physical function has recently been specifically emphasized. The Recent Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials (IMMPACT) and International Classification of Functioning, Disability and Health (ICF) guidelines, as well as a recent review of mind-body interventions for chronic pain clearly specify that physical function should be comprehensively assessed by combining self-report, performance-based (e.g., walk test), and objective (e.g., accelerometer) measures in order to better understand effects of interventions and also improve overall efficacy and effect sizes. However, no chronic pain clinical trials to date comply. There is therefore a need for comprehensive assessments of physical function among patients with chronic pain.

Physical activity is an important target for patients with chronic pain, who tend to be sedentary, deconditioned, and at risk for further pain conditions and disability. Moreover, such sedentariness negatively impacts patients’ mood, motivation, emotional function, and mental health. Walking – the preferred physical activity modality in chronic pain patients – is efficacious in improving self-reported function, particularly when quota-based (i.e. when activity is dependent on a given quota rather than dependent on pain levels), although uptake and adherence are problematic. Activity monitoring devices such as Fitbit provide the user with feedback about their level activity and progress, and present an opportunity to actively reinforce the patient in incremental quota-based gains tailored to each patient’s ability, thereby facilitating physical activity. Such devices are showing promising use in patients with chronic illness, including initial reports of effectiveness among patients with chronic pain. Our guiding hypothesis is that the most effective and efficient way to sustainably increase emotional and physical function among chronic pain patients is by 1) gradual and quota contingent increases in activity (walking) paired with re-engagement in activities of daily living; 2) monitoring and reinforcing quota contingent activity goals (number of steps) with a digital activity monitoring device (Fitbit); and 3) teaching mind-body skills to manage pain, and target barriers to engaging in physical activity.

The goal of this paper is to set the stage for a future fully powered efficacy RCT to test our guiding hypothesis. Here, we report on a 2-phased study informed by the National Institute of Health (NIH) stage model for behavioral intervention development, and National Institute of Complementary and Integrative Health (NCCIH) model for developing and testing mind-body intervention, and recent recommendations for early feasibility testing. In phase 1, we conducted focus groups to with the aim of informing adaptation of the 8-session mind-body Relaxation Response Resiliency Program (3RP) and developing the GetActive and GetActive with Fitbit programs, including treatment manuals and study procedures. The 3RP program was chosen specifically due to its demonstrated efficacy and its multimodal structure (e.g. including relaxation response, coping, CBT, and positive psychology skills), which has significant benefits over unimodal programs. In phase 2, we conducted a non-randomized pilot trial with exit interviews of the GetActive and GetActive with Fitbit programs with the aim of exploring feasibility markers and within group improvements on physical function and other relevant outcomes, to inform additional modifications before future phases of the study - phase 3 (RCT of GetActive versus GetActive with Fitbit) and phase 4 (efficacy RCT of GetActive versus GetActive with Fitbit versus health education control).

Phase 1: Development Of The GetActive And GetActive With Fitbit Programs

Methods

First, our multidisciplinary team with expertise in chronic pain, mind-body medicine, and physical activity modified the 3RP manual to target the needs of patients with chronic pain. The 3RP teaches relaxation response (e.g., diaphragmatic breathing, mindfulness), stress awareness and coping (e.g., adaptive thinking, acceptance), and positive psychology (e.g., humor, optimism) skills. We also proposed new pain specific skills including a discussion on the rationale and practice of quota-based, non pain contingent increases in activity using time goals (GetActive) and step counts (GetActive with Fitbit). Further we added behavioral activation techniques (e.g. identifying and scheduling physical activities that are achievable for the patient; targeting motivation and paired increased activity with reengagement in activities...
of daily living (e.g. increasing step count via activities such as walking to the grocery store or commuting home). Next, we developed a semi-structured qualitative interview script to gauge participants’ views on the challenges of living with chronic pain, perception of the proposed modifications of the 3RP skills, perception of the additional proposed skills targeting pain, physical activity, emotional and physical functioning, and barriers and facilitators for Fitbit use and program participation.

Participants And Recruitment
Twenty-two adults with chronic pain participated in four focus groups (N=2-9 each). Participants were recruited between September 2017 and January 2018, primarily via the Pain Clinic at the Massachusetts General Hospital, through direct referrals from study staff, and through flyers distributed through a hospital-wide email advertising clinical research. Interested participants were screened over the phone by a trained research assistant. Inclusion criteria were: 1) age ≥ 18; 2) nonmalignant musculoskeletal pain for more than 3 months; 3) able to perform a 6 min walk test; 4) own a smartphone with Bluetooth 4.0 or computer; 5) on a stable dose of psychotropic or pain medication and willing to maintain a stable dose. Exclusion criteria were: 1) medical illness expected to worsen in the next 6 months; 2) serious psychiatric illness (untreated schizophrenia, active suicidality); 3) current substance use disorder; 4) practice of yoga/meditation once per week for 45 mins or more within the last 3 months; and 5) regular physical exercise for >30 min daily. The latter two exclusion criteria were chosen to ensure that participants interviewed would be similar to those who will participate in the intervention, thus eliminating confounds. All screenings were reviewed by a clinical psychologist. Of the 86 participants screened, 39 did not meet study criteria, 16 had scheduling difficulties, and 9 dropped prior to enrollment. All participants provided written informed consent prior to the focus groups. All study procedures were approved by the Institutional Review Board of Massachusetts General Hospital, and all procedures were in compliance with the Declaration of Helsinki. Study procedures are further detailed in Figure 1.

Focus Groups
The interview focused on 4 major areas: 1) challenges of living with chronic pain, 2) perception of modified 3RP specific skills to address needs of patients with chronic pain, 3) perception of additional skills targeting activity, emotional and physical functioning, and 4) barriers/facilitators for Fitbit use and program participation. Interviews lasted 40–60 mins depending on the size of each group and were audio-recorded. Groups were led by clinical health psychologists with expertise in treating individuals with chronic pain. At the end of each group participants completed a demographic questionnaire (Table 1). Each participant was reimbursed $20 for participation.

Analysis Plan
We used guidelines for qualitative data analyses that focused on a framework approach with the goal of minimizing bias. The framework approach is a method of qualitative data analysis which involves the indexing of thematic content into categories, which are then used to compare and contrast the content and draw descriptive
conclusions. Study staff transcribed all recordings which were subsequently independently checked by an additional staff member. Next, we identified themes and developed a coding structure. Two additional staff members not involve in the development of the coding structure independently coded the data using NVivo 11. Whenever the data did not fit an existing code, new ones were added. Discrepancies were resolved through consensus between the coders and feedback from study staff with expertise in qualitative methods. A kappa coefficient was calculated to determine coding precision.

Results

The Kappa index for coding was high (0.76), indicating high consensus between raters. Coding yielded 4 major themes (bolded) and 9 subthemes (italicized). Table 2 provides further details about the themes and subsequent adaptations to the program manual.

<table>
<thead>
<tr>
<th>Table 1 Demographic And Clinical Patient Characteristics Of Participants In Focus Groups And Open Pilot Groups</th>
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<tr>
<td><strong>Focus Groups n=22</strong></td>
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<td><strong>M (SD)</strong></td>
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<tr>
<td>Age</td>
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<tr>
<td>Overall pain (scale 1–10)</td>
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<td>Gender</td>
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<td>Some college/Associates degree (&lt;16 years)</td>
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<td>Completed college (16 years)</td>
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<td>Graduate/professional degree (&gt;16 years)</td>
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<tr>
<td>Theme</td>
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<tr>
<td>Challenges of Living with Chronic Pain</td>
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<td>Perception of Mind-Body 3RP Specific Skills to Address Needs of Patients with Chronic Pain</td>
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### Table 2 (Continued).

<table>
<thead>
<tr>
<th>Theme</th>
<th>Sub-Theme</th>
<th>Quotes</th>
<th>Example Of Subsequent Changes To Manual Based On Quote</th>
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| Perception of Additional Skills Targeting Emotional and Physical Functioning | Pain-specific skills | "I don't have great body awareness… I had injuries first that I didn’t take care of right away because I wasn’t very good at figuring out body awareness. So, it was sort of things like… is this a good sort of pain? Did I overexert myself? And that’s something I still have trouble with… my own activity has sort of slowed down because I can’t really tell if I’m having like beneficial sort of sensations from it.”  
"I just want to get back to being somewhat normal, physically.”  
"It might not make the pain go away at all, but I feel like being able to work through it or walk through it and still be active without feeling like it's going to come back and bite you, I feel like that’s what I would look for or expect.” | Emphasis on body awareness (session 2) and cultivating mindfulness of pain (session 3) |
| | Physical activity skills | "The past year I’ve restricted myself so bad that I’m too scared to do anything, so … I just basically don’t.”  
"Maybe I’ll walk faster to the parking garage, and then I’ll have the worst joint pain when I get home. I don't know how to even manage that.” | Providing more detail on activity pacing (session 2) and activity limitation due to misconceptions such as “hurt always means harm” (session 1) |
| Barriers and Facilitators | Using a Fitbit | "I have enough trouble charging everything else that has to be charged.”  
"Every time it reminds me, ‘Oh, you haven't got enough steps, you have to do this,’ it’s always on the list of your every day schedule… and you always set it less priority. And it stays there for a long time, and you get really frustrated.”"The Fitbit is a reminder, it’s something in your face. We’re … creatures of habit, we need to see something that makes us go.” | Added detail of Fitbit, emphasis on making a specific plan to remember to wear, sync, and change Fitbits (Session 1) |
| | Program participation | "I’m limited because I don’t drive now … and it’s difficult to get a ride.”  
"I would be interested in in-person … because it gives me an excuse to get out of the house.”  
“So to have to come here to attend a session—that would be tough … just ’cause of the travel issues … whereas online you can just do it anywhere.” | Emphasis on motivation factors, link between attendance and potential benefit from program (session 1) |
Theme #1: Challenges Of Living With Chronic Pain
Social functioning. Most participants reported that their chronic pain affects their loved ones and impairs their own social functioning. They felt disconnected and misunderstood by others, who may not be aware of their pain. Participants expressed a wish to re-strengthen and restore their relationships.

Physical functioning. Participants noted inactivity/significant reduction in activities due to pain and associated physical symptoms such as exhaustion. They reported that their chronic pain significantly interferes with their ability to keep a job, engage with family or friends, and impacts their performance at work due to impaired concentration or frequent absenteeism.

Emotional functioning. Many participants reported feeling overwhelmed, anxious, and depressed due to pain. They also felt frustrated about the medical system and noted that their pain management needs are unmet (e.g., insurance companies not covering their preferred treatments, medical providers unable or unwilling to offer useful treatment, being told to “get over it,” and being dismissed from care). Some participants reported feeling that their doctors invalidate their experiences, are unable to understand them, or set unrealistic expectations.

Theme #2: Perception Of The Modified 3RP Specific Skills To Address Needs Of Patients With Chronic Pain
The majority of participants liked the modified 3RP skills. Participants appreciated the provided rationale for using these skills to improve pain management and reported their interest in learning these skills themselves. Several participants noted that they had been or were currently intermittently using some of the 3RP strategies, such as gratitude, humor, and optimism, as well as practices such as deep breathing and meditation. They noted these skills as being helpful and associated with improvement in pain, relaxation, and motivation. However, many noted challenges adhering to these skills (e.g., finding time to practice) and reported their belief that the group setting would help with motivation. They were receptive to pain specific meditations and learning to use these skills consistently to manage pain. Other skills, such as goal setting, were also perceived as helpful means to increase motivation and adherence over time. A minority of participants, however, expressed some reluctance, as they associated mind-body skills with inactivity or laziness. Other participants shared concerns about the logistics of practicing these skills in public.

Theme #3: Perception Of Additional Skills Targeting Emotional And Physical Functioning
Pain specific skills. Participants indicated that understanding the disability spiral, myths about pain (e.g., harm versus hurt), learning quota-based (non-pain contingent) pacing, the use of metaphors such as “the true pain alarm” (i.e., when pain signifies damage) vs the “false pain alarm” (i.e., when it does not) and setting specific goals toward increasing activity can all be useful tools to help manage their pain and become more physically active regardless of pain. Participants reported that they expected these new skills to better enable them to tend to their day-to-day activities despite the pain, rather than eliminate their pain altogether.

Physical activity skills. Generally, participants reported being sedentary, and indicated fear of exacerbating pain as a primary motive for avoiding activity. They reported continuously restricting their activities and the heavy toll associated with these restrictions. Several participants attempted to be active, primarily through walking, but noted immediate discouragement due to increased pain. All participants noted increased activity during the “good days,” and then needing to rest the following day(s) because of activity-related pain. Despite these challenges, most participants preferred walking as a primary mode of physical activity, although a minority of participants reported interest in other activities, such as aquatic exercise. The provided rationale for using pacing and quota-based, non-pain contingent increases in walking paired with re-engagement in daily activities made sense to participants.

Theme #4: Barriers And Facilitators
Using a Fitbit. Participants noted an understanding of the rationale behind using a Fitbit to help them gradually increase their daily steps while learning to pace themselves. They described their desire to meet a predetermined step goal, and to develop a habit of wearing the Fitbit and incorporate it into their normal routines. Participants noted two barriers to engaging with the Fitbit: forgetting to charge it and forgetting to wear it. They noted that having a concrete plan for when to charge their Fitbit (e.g., while taking a shower) as well as reminders to wear it would help adherence. They also agreed that receiving reminder phone calls from study staff and checking their progress on their phones would help increase both adherence and motivation.
Program participation. Participants identified several barriers to participation, including burden of travel to the clinic, challenges finding parking, cost of parking, distance of clinic to the closest train station, and scheduling conflicts. Several participants noted an increased challenge with both attendance and activity during the winter months in New England. Participants also identified several facilitators to program participation, such as the interactive nature of the group setting, the opportunity to leave their houses, and optimism regarding the program’s potential to help them better manage their pain and increase their ability to engage in activities that they enjoyed prior to the onset of their chronic pain.

**Phase 2: Open Pilot Studies**

**Methods**

**Participants And Recruitment**

Recruitment took place between January and April 2018 via the same methods described in Phase 1. Inclusion criteria were identical to those of phase 1, with an additional criterion requiring the ability to participate in 8 in-person group treatment sessions. We also excluded participants who used a Fitbit tracker in the past 3 months. A total of 13 participants enrolled (Figure 2 and Table 1). The first six participated in the GetActive program and the next seven in the GetActive with Fitbit program. All participants provided written informed consent prior to participation. All study procedures were approved by the Institutional Review Board of Massachusetts General Hospital, and all procedures were in compliance with the Declaration of Helsinki. Study procedures are further detailed in Figure 3.

**Procedure For GetActive And GetActive With Fitbit**

A research assistant scheduled the consent and assessments for all participants. A clinician described the study in detail and performed consent procedures. After consent was provided, participants completed study questionnaires in person, as a group. A research assistant was available to answer questions and check that all questions were answered. The research assistant also performed the 6 min walk test.

At the end of the 2 hr assessment session, participants received a wGT3X-BT ActiGraph accelerometer (Pensacola, FL, USA) to wear daily for 1 week, wear instructions (including no changes in routine activities), and were asked to keep a wear time log. The research assistant called participants daily with reminders to wear the ActiGraphs and help problem solve issues in real time. Participants returned the ActiGraphs again for one week, following the same guidelines previously described. They returned for the final assessment session a week later, during which they completed the post intervention questionnaires, the 6 min walk test, and exit interviews.

**Additional Procedural Steps For GetActive With Fitbit**

The research assistant downloaded the ActiGraph data while the study clinician delivered the first group session, and prepared individual Fitbit accounts with individual steps counts. Step goals for each participant were determined based on predefined criteria (see GetActive and GetActive with Fitbit programs section below). At the end of the GetActive with Fitbit group session, the research assistant paired each participant’s phone with the Fitbit. During each of the subsequent group visits, the research assistant manually programmed steps goals directly into each participant’s Fitbit using the same criteria.

**GetActive And GetActive With Fitbit Programs**

The two 8-session weekly programs retained the core components of the 3RP, modified for pain: 1) elicitation of the relaxation response (RR) through general, pain and activity-specific meditations; 2) reducing overall reactivity to stress and pain through adaptive thinking and acceptance-based techniques, and 3) increasing connectedness to self and others generally and when in pain through social support and positive psychology skills, as well as the emphasis on home practice (adaptive goal setting: “SMART goals”; defined as goals that are specific, measurable, attainable, relevant, and time-based”), daily RR practice and appreciations.

We also added 5 components: 1) education regarding myths about chronic pain and activity to help participants challenge their ingrained misconceptions regarding pain and activity; 2) the “true versus false pain alarm skill,” to help participants understand the difference between harmful pain (e.g., acute pain, new injury), and non-harmful pain (e.g., chronic pain which is not indicative of current physical damage); 3) quota-based, non-pain contingent activity pacing, to help them gradually increase activity; 4) behavioral activation (e.g., reengagement in meaningful activities of daily living) paired with quota-based pacing based increases in activity; 5) challenging misconceptions around mindfulness meditation (Table 3).
Figure 2 Participant flow; Phase 2.
For the GetActive with Fitbit group, we used the Fitbit’s step-count to assess weekly physical activity, determine participants’ pacing plans, and set weekly activity SMART goals. Each week, steps were assessed over 5-day blocks. After the ActiGraph-monitored baseline step-count was determined, participants set this count as their initial step goal, which was uploaded onto the Fitbit. Participants had to reach their goal, on average, by the day of the session. We excluded the highest and lowest step-count days. Patients were allowed 2 “rest days” per week. Missed samples or days on which patients forgot to charge or wear their devices were counted as rest days. If participants met the 5-day goal criterion, they were given the option of repeating their step goal the following week or increasing their step goal by 10%. Participants paired their Fitbits with their smartphones, which automatically downloaded data on a daily basis. A staff member calculated patients’ upcoming step goals and provided the study clinician with individual adherence data for each participant, used when discussing homework adherence and to set homework including new step goals. In the GetActive program, participants’ physical activity levels were assessed via activity logs. We calculated their corresponding pacing plans and SMART goals based on time spent being physically active or distance walked, based on individual patient preference.

Figure 3 Phase 2 procedure.
Feasibility And Acceptability Assessments

Assessment milestones were set a priori and were based on guidelines for feasibility studies.\(^40,41\) Feasibility of recruitment was assessed by determining the proportion of contacted patients who agreed to participate. A feasibility rate of 80% was considered excellent and 70% was considered good. Program acceptability was calculated by reporting the proportion of participants who attended 6 out of 8 treatment sessions. Feasibility was considered excellent when at least 70% participants attended at least 6 of the 8 planned treatment sessions (75%). Therapist adherence to the manual was determined via checklists. An adherence rate of 100% was considered excellent and 75% good. Feasibility of quantitative measures was deemed acceptable if no questionnaires were fully missing in more than 25% participants, and if reliability was higher than 0.70. Adherence to homework and activity were assessed via number of logs turned in by participants over the course of the study, and via reaching SMART goals and frequency of reported RR practice and appreciations. Adherence was deemed excellent if data were successfully collected at least 5 out of 6 days a week (>7 wear hours a day; excluding the days participants received and returned the ActiGraphs), from at least 80% of participants during the pre- and post-assessments. Adherence was deemed good if data were collected from at least 70% of participants. Fitbit adherence was assessed via daily wear of a charged device for the GetActive with Fitbit group. Adherence to Fitbit was deemed excellent if data were collected from at least 5 of 7 days from at least 80% of participants and good if collected from 70% of participants. Satisfaction with the program was assessed via the Client Satisfaction Questionnaire (CSQ-3)\(^42\) and was determined by the percentage of participants scoring above the scale midpoint (≥75% excellent; ≥ 70% good).

Physical Function Assessments

Objective Physical Function/Physical Activity

During baseline and post-program testing, all participants wore a wGT3X-BT ActiGraph accelerometer (minimum 7 hrs/day) at baseline and post intervention assessments. Adherence was deemed excellent if data were successfully collected at least 5 out of 6 days a week (>7 wear hours a day; excluding the days participants received and returned the ActiGraphs), from at least 80% of participants during the pre- and post-assessments. Adherence was deemed good if data were collected from at least 70% of participants. Fitbit adherence was assessed via daily wear of a charged device for the GetActive with Fitbit group. Adherence to Fitbit was deemed excellent if data were collected from at least 5 of 7 days from at least 80% of participants and good if collected from 70% of participants. Satisfaction with the program was assessed via the Client Satisfaction Questionnaire (CSQ-3)\(^42\) and was determined by the percentage of participants scoring above the scale midpoint (≥75% excellent; ≥ 70% good).

Performance-Based Physical Function

We used the 6 min walk test,\(^44\) which measures the distance participants walk for 6 mins. The MCID in walking distance is 54 m.\(^44\)

![Table 3 GetActive With Fitbit Session Topics And Skills](image-url)
Patient-Reported Physical Function

We measured self-reported physical function with the Patient-Reported Outcomes Measurement Information System (PROMIS) Physical Function, v.1.2.8h, which assesses one’s ability to perform activities ranging from self-care (activities of daily living) to more complex activities requiring a combination of skills, often within a social context. The MCID for this measure is 8.6 We additionally used the World Health Organization Disability Assessment Schedule (WHODAS) 2.0, a 36-item questionnaire assessing functional difficulties in various domains (e.g., understanding and communicating, getting around, self-care, getting along with people, life activities, and participation in society). Internal reliability in the current sample was good (Cronbach’s alpha=0.81–0.87) for the PROMIS and excellent (0.95–0.96) for the WHODAS. There is no MCID available for WHODAS. We used both measures in order to be able to compare their respective sensitivity in preparation for a future RCT.

Self-Reported Physical Activity Scale For Individuals With Physical Disabilities (PASIPD)48

A 13-item questionnaire which assesses engagement in leisure, household, and work activities of varying intensities for the past 7 days. This measure has established validity, with low-to-moderate Cronbach’s alpha values (0.37–0.65) within each of its five factors due to the small number of items in each factor. There is no MCID available for this measure.

Emotional Function

We assessed emotional function with the PROMIS depression v1.0.8h as well as the PROMIS anxiety, v1.08a. The MCID for PROMIS depression is 5.19 and for PROMIS anxiety is 4.28. Internal reliability for both measures in the current sample was excellent (Cronbach’s alpha=0.87–0.96 for depression, 0.93–0.96 for anxiety).

Other Patient-Reported Assessments

While our main quantitative outcomes, which we will test in the future efficacy RCT are emotional and physical function, at this stage it is important to test other variables that assess constructs that are directly targeted by the program, which may be mediators in a future RCT and will inform about the mechanisms of improvement. We used the following additional measures (MCID indicated when available):

Pain. We used the Numerical Rating Scale (NRS); almost always representing no pain and “10” representing the worst pain possible, during rest and activity. A 1-point change is considered the MCID for patients with musculoskeletal pain.54

Social isolation was assessed via the PROMIS Social Isolation short form (4a), a 4-item questionnaire assessing perceptions of being avoided, excluded, detached, disconnected from, or unknown by, others. Internal reliability (Cronbach’s alpha) for this measure was 0.60–0.82.

Mindfulness was assessed by the Cognitive and Affective Mindfulness Scale- Revised (CAMS-R), a 12-item self-report measure of the frequency with which patients use mindfulness behaviors (e.g., noticing thoughts without judgement) in their daily lives. Items are scored on a 4-point Likert scale, with “0” representing “rarely/not at all” and “3” representing “almost always”. Internal reliability (Cronbach’s alpha) for this measure was 0.63–0.77.

Emotional support was assessed via the PROMIS emotional support (4a), a 4-item questionnaire measuring perceived feelings of being cared for and valued as a person and having close relationships. This measure had excellent internal reliability in the current sample (Cronbach’s alpha=0.94–0.97).

Coping. Coping was assessed using (1) the Pain Catastrophizing Scale (PCS), which assesses hopelessness, helplessness and rumination about pain; (2) the Tampa Kinesiophobia Scale (TSK), which assesses fear of pain and activities that cause pain, and the Measure of Current Status (MOCS), which assesses ability to engage in a series of healthy coping skills (e.g., relaxation, social support, adaptive thinking). Internal reliability was found to be excellent (Cronbach’s alpha=0.93–0.94) for the PCS, good (0.77–0.87) for the TSK, and acceptable (0.71–0.77) for the MOCS.

Perceived Improvement was assessed with the Modified Patient Global Impression of Change (MPGIC), in which participants assessed on a 7-point Likert scale (with 1 representing “very much improved”) and “7” representing “very much worse” the extent to which they perceived that the intervention improved their pain, physical activity, physical function, emotional function, pain resiliency, and the degree to which the Fitbit helped to increase their activity (for the GetActive with Fitbit group). The rationale for using all of these measures concurrently is to help determine which the assessments best capture the effects of the program, and should be included in the future feasibility and efficacy RCTs.
Exit Interviews
Following program completion, participants participated in semi-structured group-based exit interviews to assess their views on program components feasibility. All participants agreed to maintain confidentiality. Interviews lasted ~60 mins and were audio-recorded, with an emphasis on the importance of honest feedback to enable program optimization.

Analysis Plan
Consistent with common guidelines for feasibility studies, we did not test efficacy or perform between-group analyses. Rather, we limited our quantitative analyses to descriptive statistics for each quantitative measure, as well as within-group pre-post comparisons using paired t-tests and Cohen’s D effect sizes (ES) to broadly gauge for potential signals of improvement. To verify results of these comparisons due to the low sample size, we additionally conducted the non-parametric related-samples Wilcoxon signed rank test comparing pre and post values. Qualitative data analysis for the exit interviews followed the methods outlined above in Phase 1.

Results
Feasibility And Acceptability Markers
Feasibility of recruitment, credibility and expectancy, client satisfaction, acceptability of treatment, adherence to ActiGraphs and Fitbit, adherence to homework, therapist adherence to manual, and perception of improvement were good to excellent and similar for both groups (Table 4).

Physical Function (Table 5)
Objective physical function. Overall, both groups, when considered together, showed a significant increase in ActiGraph-measured number of steps from baseline to posttest (Figure 4). Participants in the GetActive with Fitbit group had a relatively high baseline step-count

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Feasibility And Acceptability Of The Open Pilot Studies</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Outcome</strong></td>
<td><strong>GetActive</strong></td>
</tr>
<tr>
<td>Feasibility of recruitment</td>
<td>47 participants out of 52 successfully contacted agreed to complete screening (excellent)</td>
</tr>
<tr>
<td>Credibility and expectancy</td>
<td>5 out of 6 participants (83%) scored over median split (excellent)</td>
</tr>
<tr>
<td>Client satisfaction</td>
<td>5 out of 6 participants (83%) scored over median split (excellent)</td>
</tr>
<tr>
<td>Acceptability of treatment</td>
<td>4 out of 6 participants (66%) attended ≥ 6 out of 8 sessions (acceptable)</td>
</tr>
<tr>
<td>Adherence to ActiGraphs and Fitbit</td>
<td>6 out of 6 participants at baseline (100%) and 5 out of 6 at post-test (83%) recorded ≥ 5 days of ActiGraph data (excellent)</td>
</tr>
<tr>
<td>Adherence to homework</td>
<td>4 out of 6 participants (66%) completed ≥ 5 out of 7 weeks of homework (acceptable)</td>
</tr>
<tr>
<td>Therapist adherence to manual</td>
<td>100% adherence (excellent)</td>
</tr>
<tr>
<td>Feasibility of quantitative measures</td>
<td>No questionnaires missing fully (excellent)</td>
</tr>
<tr>
<td>Patients’ perception of improvement</td>
<td>5 out of 6 participants (83%) reported overall improvement (excellent)</td>
</tr>
<tr>
<td>Analgesics (non-narcotic)</td>
<td>Stable</td>
</tr>
<tr>
<td>Narcotic analgesics</td>
<td>Stable</td>
</tr>
<tr>
<td>Adverse events</td>
<td>None</td>
</tr>
</tbody>
</table>
(>7300 steps, with most (5 out of 7) participants walking more than 5,000 steps a day). Improvements in number of steps in both groups were clinically meaningful, over the MCID of 1000 steps45 (medium effect size). Four out of 7 participants within the GetActive with Fitbit group and 4 out of 6 within the GetActive group showed increases in step-count over the MCID (medium effect size).

Performance-based physical function. As a group, participants in the GetActive group generally increased their distance walked in the 6 min walk test (4 out of 6 participants; 1 above the MCID; medium effect size). Participants in the GetActive with Fitbit showed no change in the 6 min walk test (4 out of 7 participants increased distance but did not reach the MCID).

Self-report physical function. Both the GetActive and GetActive with Fitbit groups showed some improvement in physical function (medium effect size; Table 5).

Emotional Function
Both groups exhibited non-significant improvement in anxiety (small effect size) and depression (small to medium effect size; Table 6).

Other Measures
The GetActive group exhibited clinically meaningful reductions in levels of pain during rest and activity (medium effect size; Table 6), pain resilience (large effect size), and non-significant improvements in kinesiophobia (large effect size), and coping (medium-large effect size; Table 6). Participants in the GetActive with Fitbit group similarly exhibited a significant and clinically meaningful decrease in levels of pain during rest and activity (large effect sizes), as well as improvements in coping (large effect size), and a non-significant improvement in pain resilience (medium-large effect size). These comparisons were repeated using the non-parametric related-samples Wilcoxon signed rank test (see Analysis Plan) and yielded similar results in terms of direction and significance.

Exit Interviews
Participants reported high satisfaction with the learned relaxation, mind-body, and coping skills, and reported an improved ability to manage stress, pain, and an increased quality of life. They indicated that the pacing of the program, particularly in the first sessions, could be more relaxed and spread out over more sessions. Three participants indicated that healthy behaviors topics (e.g., healthy eating, sleep) detracted from the rest of the intervention. Participants in the GetActive

<table>
<thead>
<tr>
<th>Table 5</th>
<th>Objective, Performance-Based And Self-Report Activity/Function Outcomes</th>
<th>GetActive</th>
<th>GetActive With Fitbit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical Activity</td>
<td></td>
<td>Baseline M (SD)</td>
<td>Post-Test M (SD)</td>
</tr>
<tr>
<td>ActiGraph Average Steps</td>
<td>5341.33 (2470.81)</td>
<td>6514.83 (3728.3)</td>
<td>1173.5 (−1136.30, 3483.30)</td>
</tr>
<tr>
<td>6 min Walk Test Distance (m)</td>
<td>405.88 (109.98)</td>
<td>436.95 (41.05)</td>
<td>31.07 (29.26, 32.6)</td>
</tr>
<tr>
<td>Physical function (PROMIS)</td>
<td>37.98 (7.76)</td>
<td>43.28 (4.77)</td>
<td>5.30 (4.75, 5.85)</td>
</tr>
<tr>
<td>Self-reported physical activity</td>
<td>8.60 (8.02)</td>
<td>11.40 (11.94)</td>
<td>2.80 (2.45, 3.15)</td>
</tr>
</tbody>
</table>

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with Fitbit group reported that the Fitbit was helpful in improving sleep and body awareness. However, they also reported feeling guilty and discouraged whenever they did not meet their step-goal or when receiving reminders about lack of activity.

**Discussion**

Despite decades of research, improvements in emotional and physical function following mind-body treatments among chronic pain patients are modest and generally fade over time. Further, no clinical trials to date assess physical function using IMMPACT and ICF criteria. Although there is evidence that physical activity helps patients with chronic pain, uptake and adherence remain problematic. The current study used the NIH stage model and NCCIH model for intervention development and adapted an evidence-based, group, mind-body program for the specific needs of patients with chronic pain. The study targeted comprehensive improvement in emotional and physical function (self-report, performance based and objective) with a Fitbit device (GetActive with Fitbit) or without it (GetActive; phase 1). We also assessed preliminary feasibility markers for the two programs via 2 nonrandomized open pilot studies (phase 2).

Focus groups confirmed the physical, emotional, and social challenges associated with chronic pain. Fear of exacerbating pain, difficulties with pacing, and frustration with current functioning were common complaints from participants. They held largely favorable views of the modified 3RP and the additional pain management skills described, the use of a Fitbit device to increase activity, and the value of learning skills in a supportive group setting. The rationale of combining pain-specific mind-body skills with increased in walking linked to re-engagement in activities of daily living resonated with focus group participants. They identified the group setting and the Fitbit as particularly motivating in regards to meeting their goals. These results confirmed prior reports in this population.

Results are also in line with evidence suggesting that this population is in need of extra support to engage in physical activity, and the essential roles of quota-based pacing and mind-body skills as aids in this endeavor.

**Figure 4** Pre-post changes in performance-based measures in the 2 programs.
Table 6: Quantitative Self-Report Outcomes Of Pain, Emotional Function And Coping

<table>
<thead>
<tr>
<th>Measures</th>
<th>GetActive</th>
<th></th>
<th></th>
<th></th>
<th>GetActive</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline M (SD)</td>
<td>Post-Test M (SD)</td>
<td>Pre-Post Change M (95% CI)</td>
<td>t value</td>
<td>p value</td>
<td>Cohen’s D</td>
<td>Baseline M (SD)</td>
<td>Post-Test M (SD)</td>
</tr>
<tr>
<td>Pain at rest</td>
<td>5.17 (2.23)</td>
<td>3.83 (2.64)</td>
<td>1.33 (0.25, 2.42)</td>
<td>3.16</td>
<td>0.025</td>
<td>1.30</td>
<td>5.29 (2.29)</td>
<td>3.57 (2.64)</td>
</tr>
<tr>
<td>Pain with activity</td>
<td>6.67 (1.51)</td>
<td>4.67 (3.56)</td>
<td>2.00 (-0.89, 4.89)</td>
<td>1.78</td>
<td>0.136</td>
<td>0.73</td>
<td>7.57 (1.81)</td>
<td>6.00 (2.24)</td>
</tr>
<tr>
<td>Pain resilience</td>
<td>28.83 (6.62)</td>
<td>39.17 (6.85)</td>
<td>-10.33 (-20.10, -0.56)</td>
<td>2.72</td>
<td>0.042</td>
<td>1.11</td>
<td>29.57 (11.52)</td>
<td>33.71 (7.50)</td>
</tr>
<tr>
<td>Depression</td>
<td>59.02 (6.65)</td>
<td>54.70 (10.10)</td>
<td>4.32 (-4.42, 13.05)</td>
<td>1.27</td>
<td>0.260</td>
<td>0.52</td>
<td>55.11 (10.86)</td>
<td>53.66 (4.42)</td>
</tr>
<tr>
<td>Anxiety</td>
<td>59.38 (5.55)</td>
<td>56.78 (11.61)</td>
<td>2.60 (-7.19, 12.39)</td>
<td>0.68</td>
<td>0.525</td>
<td>0.28</td>
<td>55.00 (12.02)</td>
<td>51.74 (9.21)</td>
</tr>
<tr>
<td>Social isolation</td>
<td>48.63 (8.16)</td>
<td>46.53 (9.33)</td>
<td>2.10 (-9.70, 13.90)</td>
<td>0.46</td>
<td>0.667</td>
<td>0.19</td>
<td>51.94 (6.46)</td>
<td>50.54 (5.24)</td>
</tr>
<tr>
<td>Emotional support</td>
<td>45.48 (9.73)</td>
<td>45.68 (9.90)</td>
<td>-0.20 (-4.72, 4.32)</td>
<td>0.11</td>
<td>0.914</td>
<td>0.05</td>
<td>49.80 (7.76)</td>
<td>50.64 (10.26)</td>
</tr>
<tr>
<td>Mindfulness</td>
<td>29.83 (4.58)</td>
<td>33.33 (5.35)</td>
<td>-3.50 (-9.89, 2.89)</td>
<td>1.41</td>
<td>0.218</td>
<td>0.58</td>
<td>27.29 (4.54)</td>
<td>29.00 (4.76)</td>
</tr>
<tr>
<td>Pain catastrophizing</td>
<td>44.00 (12.51)</td>
<td>33.17 (14.32)</td>
<td>10.83 (-7.63, 29.29)</td>
<td>1.51</td>
<td>0.192</td>
<td>0.62</td>
<td>34.14 (11.05)</td>
<td>29.86 (8.55)</td>
</tr>
<tr>
<td>Kinesiophobia</td>
<td>43.17 (8.91)</td>
<td>34.83 (8.61)</td>
<td>8.33 (-2.09, 18.76)</td>
<td>2.05</td>
<td>0.095</td>
<td>0.84</td>
<td>40.43 (1.90)</td>
<td>37.14 (7.01)</td>
</tr>
<tr>
<td>Coping</td>
<td>25.67 (8.12)</td>
<td>32.83 (3.87)</td>
<td>-7.17 (-17.22, 2.89)</td>
<td>1.83</td>
<td>0.126</td>
<td>0.75</td>
<td>22.00 (6.76)</td>
<td>30.29 (7.48)</td>
</tr>
</tbody>
</table>
The open pilot studies showed excellent therapist adherence, client satisfaction, perceptions of improvement, feasibility of quantitative measures, and feasibility of recruitment for both programs. One hundred percent of participants with baseline data completed post-testing in both groups, and attrition was lower compared to other studies of chronic pain. Credibility and adherence to ActiGraph and Fitbit were high, and minimal adverse events were reported. Results indicate that the interventions and procedures are feasible and accepted by study participants.

In line with common guidelines for feasibility studies, this study did not test efficacy or perform any between-group analyses. Nevertheless, preliminary exploration of effect sizes of within group post-change suggests that both programs show promise for improvements in most outcome measures. Critically, both groups exhibited step-count increases that are clinically meaningful. Moreover, we found large effect sizes and clinically meaningful decreases in pain following the programs, as well as large and significant effects for improvements in physical function and medium-large effect sizes for improvements in coping, pain resilience, and kinesiophobia.

Qualitative exit interviews indicated high satisfaction with both programs and the mind-body, pain-specific and physical activity skills learned, and suggested additional modifications including: 1) increasing the total number of sessions in both programs from 8 to 10 to allow for a more relaxed pace of teaching the skills; 2) including 3RP specific information on healthy eating and sleep in an appendix to avoid deterring from the flow of the program; 3) debunking the myths regarding mindfulness, meditation, and other mind-body practices which had associations with “laziness” for some participants; 4) incorporating a more thorough discussion on challenges associated with forming new habits, as well as introducing the skill of self-compassion to help patients remain motivated even when not meeting step-count goals; 5) developing a comprehensive assessment of activity (e.g., job activity, seating time, Physical Activity Questionnaire-Short Form IPAQ-SF scores) during screening to ensure that participants are sedentary (e.g., less than 5,000 steps) so that they can fully benefit from the programs.

Conclusions
This study is the first to integrate physical activity with mind-body skills to increase comprehensive physical and emotional functioning in a heterogeneous group of patients with chronic pain, the first to use a comprehensive assessment of physical function consistent with IMMPACT and ICF criteria, and the first to compare increase in activity with and without a Fitbit. Focus groups, open pilot, and exit interview data provided preliminary evidence for feasibility and acceptability of the novel interventions as well as a signal of improvement in physical activity, pain, physical function, and other measures. Consistent with the aims of pilot feasibility studies, our aim was not to examine the efficacy of the programs or generalize findings to the larger pain population. Rather, we collected meaningful information which allowed us to further modify the GetActive with Fitbit and GetActive programs and study procedures which are currently being tested within a definitive feasibility pilot RCT, following the NIH stage model and NCCIH framework. If feasibility benchmarks are met within this feasibility RCT of GetActive versus GetActive with Fitbit, we will next conduct a fully powered RCT of the GetActive with Fitbit versus Get Active versus a health education control in order to determine comparatively their efficacy and sustained improvement in emotional and physical function, as well as elucidate whether the Fitbit is a necessary ingredient. Results of the future efficacy RCT will have important implications for the treatment of chronic pain and may lead to a paradigm shift in the manner in which pain is assessed and treated, support implementation of ICF and IMMPACT criteria across all settings, and potentially provide evidence for the use of digital activity trackers as activity reinforcers within chronic pain treatments.

Acknowledgement
We thank Stephanie Moffat and Christopher Funes for transcribing the interviews.

Funding
This study was funded by an R34 grant from the National Center for Complementary and Integrative Health (1R34AT009356-01A1) to the senior author.

Disclosure
Dr. Schatman serves as a consultant to Kaleo Pharma and Quest Diagnostics and received ad board honorarium from Salix Pharmaceuticals. All other authors declare that they have no conflicts of interest in this work.

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