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ORIGINAL RESEARCH

Psychosocial Correlates of Objective, Performance-Based, and Patient-Reported Physical Function Among Patients with Heterogeneous Chronic Pain

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Background: Improving all aspects of physical function is an important goal of chronic pain management. Few studies follow recent guidelines to comprehensively assess physical function via patient-reported, performance-based, and objective/ambulatory measures.

Purpose: To test 1) the interrelation between the 3 types of physical function measurement and 2) the association between psychosocial factors and each type of physical function measurement.

Methods: Patients with chronic pain (N=79) completed measures of: 1) physical function (patient-reported disability; performance-

based 6-minute walk-test; objective accelerometer step count); 2) pain and non-adaptive coping (pain during rest and activity, pain-catastrophizing, kinesiophobia); 3) adaptive coping (mindfulness, general coping, pain-resilience); and 4) social-emotional dysfunction (anxiety, depression, social isolation and emotional support). First, we tested the interrelation among the 3 aspects of physical function. Second, we used structural equation modeling to test associations between psychosocial factors (pain and non-adaptive coping, adaptive coping, and social-emotional dysfunction) and each measurement of physical function.

Results: Performance-based and objective physical function were significantly interrelated (r=0.48, p<0.001) but did not correlate with patient-reported disability. Pain and non-adaptive coping ($\beta=0.68$, p<0.001), adaptive coping ($\beta=-0.65$, p<0.001) and social-emotional dysfunction ($\beta=0.65$, p<0.001) were associated with patient-reported disability but not to performance-based or objective physical function (p>0.1).

Conclusion: Results suggest that patient-reported physical function may provide limited information about patients' physical capacity or ambulatory activity. While pain and non-adaptive reactions to it, adaptive coping, and social-emotional dysfunction may potentially improve patient-reported physical function, additional targets may be needed to improve functional capacity and ambulatory activity.

Trial Registration: ClinicalTrials.gov NCT03412916.

Keywords: physical function, chronic pain, six-minute walk test, accelerometer, psychosocial factors

Introduction

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Chronic pain is financially costly, widespread¹ and decreases physical function. Most patients with chronic pain report challenges with physical function and

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© 2020 Greenberg et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/ terms.php and incorporate the Creative Commons Attribution — Non Commercial (unported, v3.0) License (http://creativecommons.org/licenses/by-nc/3.0/). By accessing the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). activity-related increased symptoms.² As a result, chronic pain treatments are increasingly focused on improving physical function.^{3,4}

Most physical function research in chronic pain has been focused on patient-reported outcomes (PROs).^{5–7} However, physical function is a multidimensional construct that encompasses more than an individual's perception of functioning. Consistently, the Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials (IMMPACT)^{8,9} and the International Classification of Functioning, Disability and Health (ICF)¹⁰ recommend comprehensively assessing physical function, with PROs, performance-based (which capture functional capacity during a time-limited standard test), and objective assessments (ambulatory measurement of daily step-count via accelerometers across a specified time period).¹¹

Studies evaluating the relationship between the different aspects of physical function in chronic pain have generally found conflicting results on the association between PROs and accelerometer-measured physical function.^{12–14} They found a small to moderate association between PROs and performanced based physical function,^{15–18} as well as between accelerometer-measured physical function and PROs.^{19,20} Currently, only 2 published papers looked at all 3 physical function measures within 1 study and found support for an association between PROs and performance-based function but little or no support for associations with step-count.^{21,22}

Treatments aimed at improving physical function in chronic pain typically target psychosocial factors such as pain, coping, and social-emotional factors,^{23–25} which are associated with improvement in PROs.^{8,26} However, it is unclear whether targeting these constructs also improves performance-based and objective/ambulatory-measured physical function, as prior studies found mixed results for associations with performance-based physical function.^{12,21,28–30}

The success of chronic pain treatments lies in targeting factors that can improve all facets of physical function.^{31,32} To inform such pain treatments, the association between all facets of physical function, as well as their relationship with psychosocial factors typically targeted in chronic first needs to be established. Consistently, we aimed to test, within 1 model, the association among psychosocial factors typically targeted in chronic pain trials with PROs, performance-based, and objective physical function. We examined a theoretically driven model of latent psychosocial factors (pain and non-adaptive coping,

adaptive pain coping and social-emotional dysfunction) and modeled their relationships to these 3 physical function facets. First, we wanted to understand to which extent PROs, performance-based and objective physical function are interrelated. Next, we explored cross-sectional associations of the 3 latent constructs with the 3 physical function facets.

Method

Participants and Recruitment

We analyzed baseline data from 2 sequential pilot mindbody clinical trials.^{33,34} The trials constituted 2 phases of the same study, tested different iterations of the same mind-body interventions, targeted the same population and used identical inclusion/exclusion criteria, referral sources, and recruitment materials. Participants were recruited through direct referrals from the Center for Pain Medicine at Massachusetts General Hospital, online mailing lists and advertisements within Partners Healthcare. Recruitment occurred between July 2018 and September 2019. Inclusion criteria were: 1) 18 years of age or older, 2) nonmalignant self-reported chronic pain for at least 3 months, consistent with the definition of chronic pain as persistent or recurrent pain lasting longer than 3 months,³⁵ 3) ability to walk for at least 6 minutes, 4) access to a mobile device with Bluetooth 4.0, 5) if on pain or psychotropic medication, being on a stable dose for the past 3 months, and 6) cleared for participation by a medical doctor. Exclusion criteria were: 1) medical illness that was expected to worsen over the following 6 months, 2) active suicidality or untreated and serious psychiatric illness, 3) current untreated substance use disorder, 4) participation in meditation or yoga practice for more than 45 minutes a week during the prior 6 months, 5) using a Fitbit device in the prior 6 months, and 6 >30 min daily of regular physical exercise. The mind-body physical activity programs aimed to improve PRO, performancebased and objective physical function through teaching mind-body skills (eg, breathing meditation, body scan, mindfulness), pain-specific cognitive-behavioral skills (eg, goal setting, behavioral activation, adaptive thinking), and physical restoration skills (eg, quota-based pacing). We tested our research questions using baseline data collected prior to program participation.

A total of 95 participants met inclusion and exclusion criteria across the 2 trials. Participants had a mean age of 50.69 years (SD=14.68), were predominantly female (67%), white (76%), non-Hispanic (85%) (Table 1). The

Table I Demographics

Age (M (SD))	50.69 (14.68)	
	N (%)	
Gender		
Male	31 (32.6%)	
Female	64 (67.4%)	
Ethnicity		
Hispanic or Latino/Latina	12 (12.6%)	
Not Hispanic or Latino/Latina	81 (85.3%)	
Missing	2 (2.1%)	
Race		
White	72 (75.8%)	
American Indian/Alaskan Native	2 (2.1%)	
Asian	6 (6.3%)	
Black/African American	8 (8.4%)	
More Than One Race	7 (7.4%)	
Marital Status		
Single, never married	34 (35.8%)	
Living with significant other	(.6%)	
Married	27 (28.4%)	
Separated/Divorced	19 (20%)	
Widowed	4 (4.2%)	
Employment		
Employed full-time	26 (27.4%)	
Employed part-time	12 (12.6%)	
Going to school full or part time	3 (3.2%)	
Retired	17 (17.9%)	
Unemployed	20 (21%)	
Other	17 (17.9%)	
Education		
High school (12 years)	14 (14.7%)	
Some college/Associates degree (<16 years)	28 (29.5%)	
Completed college (16 years)	20 (21.1%)	
Graduate/professional degree (>16 years)	33 (34.7%)	
Pain Type		
Multi-site	65 (68.4%)	
Back pain	10 (10.5%)	
Neck pain	3 (3.2%)	
Fibromyalgia	2 (2.1%)	
Other	14 (14.7%)	
Missing	1 (1.1%)	
Pain duration (median)	10 years	

most common marital, education and employment statuses were single or never married (36%), holding a graduate or professional degree (35%), and being fully employed (27%), respectively. Participants reported a median pain duration of 10 years (SD=10.48; min=3 months; max=60 years) and the majority (68%) had more than 1 type of chronic pain.

After providing written informed consent, participants completed a battery of PROs assessing physical function, pain and non-adaptive coping, and socialemotional dysfunction (see Measures), and underwent the 6-minute walk test (6MWT³⁶). At the end of the baseline visit, participants received a wGT3X-BT ActiGraph accelerometer (ActiGraph, LLC, Fort Walton Beach, FL, USA) and were instructed to wear it during all waking hours over their right hip. They were also instructed to maintain their regular levels of activity, and to fill out a daily device-wear and activity log. Participants were offered the option to receive text message reminders to wear the ActiGraph devices. All study procedures were approved by the Institutional Review Board at Massachusetts General Hospital and were conducted in accordance with the Declaration of Helsinki.

Measures

We administered reliable and valid measures that captured the constructs of interest and are routinely used in chronic pain clinical trials. In line with our research question and recent IMMPACT criteria, we assessed PROs, performancebased (6MWT) and objective/ambulatory (step-count) physical function. We combined the rest of our psychosocial measures into categories based on conceptual overlap and prior research. The first category included pain (ie, pain during rest and activity) and non-adaptive coping (ie, pain catastrophizing, kinesiophobia). These factors were grouped together because they are all negative responses to an initial nociceptive stimulus.^{37,38} The second category consisted of adaptive coping with pain (ie, general coping strategies, pain resilience, and mindfulness). Positive psychology research indicates that these factors provide a useful framework for assessing strengths in individuals who experience chronic pain, beyond the absence of the non-adaptive reactivity patterns in the first category.^{39,40} We grouped these factors together since cultivating mindfulness and relaxation is an established adaptive coping strategy in the face of pain and difficulty.41-43 The third category consists of socialemotional dysfunction (anxiety, depression social isolation, emotional support), as these constructs are often interrelated and are frequently coupled in relevant literature.^{44–46}

Objective Physical Function

We measured average daily step count over the course of 7 days with a wGT3X-BT ActiGraph accelerometer.⁴⁷

ActiGraph settings are detailed in previous studies.^{33,34} Participants were instructed to wear the ActiGraph during all waking hours over their right hip using an elastic belt for one week, except when in water (eg, bathing or swimming). The mean daily ActiGraph wear time in our sample was 12.2 hours per day. Data were stored, cleaned and analyzed with ActiLife software. The devices recorded counts in epochs of 30 seconds.⁴⁸ We defined non-wear time as 90 or more consecutive minutes having no activity counts.⁴⁹ We allowed 2 minutes or less of activity counts between 0 and 100,⁵⁰ and any wear periods shorter than 10 minutes long were ignored. A staff member verified the validity of ActiGraph wear time data and compared them with self-reported activity logs to ensure consistency.

Performance-Based Physical Function

We used the 6-minute walking test (6MWT³⁶) to measure the total distance in meters that participants walk in the span of 6 minutes while observed in the clinic. This test provides a snapshot of an individual's functional capacity and has demonstrated excellent reliability in patients with chronic pain.⁵¹ Higher distance traveled depicted higher physical function. We used a standardized method to prevent random error or reactivity. The test was performed in a closed, flat, oval indoor space marked in meter increments. Minimal and uniform encouragements were given to all participants. The same study staff performed both the baseline and post-program 6MWT.

Patient-Reported Physical Function (Disability)

We used the World Health Organization Disability Assessment Schedule (WHODAS 2.0) (World Health Organization, 2010) to assess patient reported physical function. This measure had previously been demonstrated to be valid in patient with chronic pain.⁵² The WHODAS 2.0 is a 36-item scale assessing difficulty in performing various daily living activities in domains such as cognitions (understanding and communicating), mobility (eg, getting around), self-care (eg, hygiene, dressing), getting along and interacting with others, life activities (eg, household, leisure, school and work responsibilities), and participation (eg, community activities). Internal reliability for the WHODAS 2.0 in the current sample was excellent (Cronbach's alpha=0.96). High scores depict higher disability and lower physical function. For consistency herein we refer to this construct as "disability".

Pain and Non-Adaptive Coping Strategies The Numerical Rating Scale (NRS)^{53,54} was used to measure pain intensity. This is a 2-item scale measuring pain intensity at rest and pain intensity during activity. High scores depict higher pain intensity.

The Pain Catastrophizing Scale (PCS)⁵⁵ was used to measure participants' tendencies toward rumination, magnification and helplessness when faced with pain. The PCS has 13 items and had excellent internal reliability in our sample (Cronbach's alpha=0.94). High scores depict higher pain catastrophizing.

The Tampa Scale for Kinesiophobia (TSK)⁵⁶ was used to measure fear of movement and physical activity due to pain. The TSK is a 17-item questionnaire and had good internal consistency (Cronbach's alpha=0.86). High scores depict higher levels of kinesiophobia.

Adaptive Coping Strategies

The Measures of Current Status (MOCS-A⁵⁷) was used to measure participants' abilities to use general coping efforts and strategies such relaxation, adaptive thinking, and social support. The MOCS-A is a 13-item questionnaire and had good internal reliability (Cronbach's alpha=0.88). High scores depict better use of coping strategies.

The Pain Resilience Scale (PRS)⁵⁸ was used to capture participants' abilities to maintain behavioral engagement and regulate emotions in the face of pain. This is a 14-item measure and had good internal reliability in the current sample (Cronbach's alpha=0.89). High scores depict greater pain resilience.

The Cognitive and Affective Mindfulness Scale – Revised (CAMS-R)⁵⁹ was used to assess cognitive and affective aspects of mindfulness (ie, participants' abilities to pay attention to their experiences in the present moment in a non-judgmental manner). This is a 12-item questionnaire and had good internal reliability (Cronbach's alpha=0.84). High scores depict greater mindfulness.

Social-Emotional Dysfunction

The PROMIS anxiety $(v1.08a)^{60}$ was used to measure anxiety symptoms. This is an 8-item questionnaire and had excellent internal reliability (Cronbach's alpha=0.95). High scores depict higher levels of anxiety.

The PROMIS depression $(v1.08b)^{61}$ was used to measure symptoms of depression. This is an 8-item questionnaire and had excellent internal reliability (Cronbach's alpha=0.96). High scores depict higher levels of depression.

The PROMIS social isolation short form $(4a)^{62}$ was used to capture patients' perceptions of being left out and disconnected from others. This is a 4-item questionnaire and had good internal reliability (Cronbach's alpha=0.88). High scores are indicative of social isolation.

The PROMIS emotional support $(4a)^{63}$ was used to capture the availability of others with whom to talk to and feeling appreciated by others. This is a 4-item questionnaire and had excellent internal reliability (Cronbach's alpha=0.92). We reversed the scores on this measure such that high scores depict greater dysfunction.

Statistical Analysis

Sample Characteristics and Correlations

Data analyses were conducted in R 3.6.1.⁶⁴ We examined the sample for missing data. Of the entire N=95 sample, one participant had missing data on the NRS pain at rest, and 15 participants had missing ActiGraph data (N=6 dropped out and did not wear the ActiGraph; N=5 had no valid wear days, and 4 had missing data due to a technical error). A total of N=79 participants had full data for all study measures and were included in the main analyses, consistent with listwise requirements for structural equation modeling (SEM).^{65,66} We also verified normality (kurtosis = 0.49-0.53, skewness = 0.25-0.27) and the absence of univariate outliers to ensure the suitability of the data for SEM. Descriptive statistics were used to characterize the sample by study measures. Pearson correlations assessed the bivariate associations between the different measures of physical function. All measures were standardized to z-scores for subsequent latent variable analyses.

Determining Latent Variable Structure

Prior to SEM, we confirmed that our psychosocial measures had an underlying factor structure. Separate constructs for measures of pain and non-adaptive coping, adaptive coping, and social-emotional dysfunction were established using confirmatory factor analysis (CFA) with maximum likelihood estimation. This approach is widely used to evidence the validity of psychosocial constructs.⁶⁷ Factors were retained as latent variables in the SEM if they met commonly accepted criteria:⁶⁸ Bartlett's test of sphericity (p < 0.05), the Kaiser-Meyer-Olkin (KMO) measure of sampling adequacy (>0.60), and loadings ≥ 0.40 .

Test of Measurement and Structural Models

SEM was conducted using the lavaan package in $R^{.69}$ SEM allowed us to build on the CFA by testing the hypothesized relationships between psychosocial latent variables (pain and non-adaptive coping, adaptive coping, and social-emotional dysfunction) and physical function measurements (objective, performance-based, and patient-reported). First, we developed a measurement model by assigning each measure to its respective latent variable from the CFA. Next, we tested the degree to which non-adaptive coping, adaptive coping, and socialemotional dysfunction were separate yet related psychosocial constructs. Finally, we entered physical function variables as single indicators and specified directional paths to each of the psychosocial latent variables. This allowed for examination of differential associations between physical function modality (objective, performance-based, and patient-reported) and latent variables (pain and non-adaptive coping, adaptive coping, socialemotional dysfunction), consistent with our research questions.

Estimation and Model Fit

Maximum likelihood estimation, which is the most commonly used method in SEM,^{67,70} was used to estimate all model parameters simultaneously. Decisions regarding model were based on commonly accepted criteria:71,72 Chi-square test of fit (Cmin/df) value near 2, a comparative fit index (CFI); $^{73} \ge 0.95$, and the root-meansquare error approximation (RMSEA⁷⁴) value between 0.002 and 0.08. We considered each of these indices when interpreting goodness of fit as no single measure is considered a "gold standard".^{74,75} Within the model. correlation coefficients were used to measure the bidirectional strength of association between the psychosocial latent variables. Statistically significant regression coefficients (standardized beta values) suggest a directional relationship between the psychosocial latent variables and the physical function measures.

Results

Preliminary Analyses

Participants were generally sedentary (M daily step count of ~5550 steps),⁷⁶ had lower performances on the 6MWT (~350 meters),⁵¹ and higher levels of disability on the WHODAS 2.0 (~32)^{7,77} compared to previously reported values in other chronic pain samples. Table 2 illustrates descriptive statistics for all study measures.

Table 2 Descriptive Statistics for Study Measures

	M (SD)
ActiGraph average steps	5548.63 (2824.95)
6-minute walk test distance (m)	348.23 (89.96)
Physical function (WHODAS)	31.26 (18.72)
Pain at rest	5.34 (2.30)
Pain with activity	7.05 (2.02)
Pain catastrophizing	21.63 (12.06)
Kinesiophobia	39.45 (8.33)
Coping strategies	26.91 (9.28)
Pain resiliency	34.29 (10.10)
Mindfulness	30.83 (6.54)
Anxiety	56.50 (9.80)
Depression	56.63 (10.35)
Social isolation	50.34 (10.15)
Emotional support	48.99 (8.46)

Performance-based (6MWT) and objective (ActiGraph steps) physical function were significantly and moderately correlated (r = 0.47, p < 0.001). Patient-reported disability (WHODAS) was not significantly correlated with performance-based (r = -0.19, p = 0.09) or objective/ambulatory measured (r = -0.18, p = 0.11) physical function.

Confirmatory Factor Analyses (CFA)

Three CFAs were conducted to determine that the psychosocial measures loaded onto their respective constructs. The NRS (activity and rest), PCS, and TSK loaded onto the first factor for pain and non-adaptive coping (KMO = 0.70; Bartlett's test, p < 0.001, standardized loadings = 0.58-0.76, variance explained = 44%). The MOCS-A, PRS, and CAMS-R loaded onto the second factor for adaptive coping (KMO = 0.61; Bartlett's test, p < 0.001, standardized loadings = 0.66-0.99, variance explained = 58%). The PROMIS subscales of anxiety, depression, social isolation, and emotional support loaded onto the third factor for social-emotional dysfunction (KMO = 0.66; Bartlett's test, p < 0.001, standardized loadings = 0.33-0.90, variance explained = 51%). Pain and nonadaptive coping, adaptive coping, and social-emotional dysfunction were retained as the latent variables in subsequent SEM.

Structural Equation Modeling (SEM)

The measurement model, with the psychosocial measures loaded onto their respective latent variables, demonstrated adequate fit (Cmin/df = 1.57, CFI = 0.944, RMSEA = 0.085). Next, we explored interrelationships between the

psychosocial constructs before adding the physical activity measures. Prior to entering the physical function measures, the pain and non-adaptive coping factor was significantly negatively correlated with adaptive coping (r = -0.57, p = 0.014) and significantly positively correlated with social-emotional dysfunction (r = 0.67, p = 0.001). Higher adaptive coping and social-emotional dysfunction were also significantly and negatively correlated (r = -0.79, p < 0.001).

As illustrated in Figure 1, the 3 physical function measures were added to the model as indicators and directional paths were specified to assess the relationships between psychosocial latent variables and physical function measures. This final structural model retained adequate fit (Cmin/df = 1.39, CFI = 0.946, RMSEA =0.070). Greater pain and non-adaptive coping ($\beta = 0.68$, p < 0.001), social-emotional dysfunction ($\beta = 0.65$, p < 0.001) 0.001), and adaptive coping ($\beta = -0.65$, p < 0.001) were all significantly associated with higher patient-reported disability. The latent variables for pain and non-adaptive coping, adaptive coping, or social-emotional dysfunction were not significantly correlated with the objective (range $\beta = -0.11 - 0.08$, range p = 0.390 - 0.639) nor performancebased (range $\beta = -0.18 - 0.028$, range p = 0.150 - 0.806) measures of physical function.

Discussion

Improving all aspects of physical function (PROs, performance-based and objective/ambulatory measured) is an important goal of chronic pain treatments. Using SEM, we first tested to which extent the 3 types of physical function measurement are interrelated, and second, to what degree common psychological factors targeted within non-pharmacological clinical trials are associated with each of the 3 physical function measures.

Testing our first research question revealed that while the 6MWT and accelerometer-measured daily average step-count were moderately correlated, their correlation with disability was not statistically significant. This finding partially aligns with previous research,^{21,22} which identified little or no association between PRO physical function and accelerometer data. However, these 2 studies also found a significant association between PRO and performance-based physical function. This discrepancy may be due to differences in the PROs used to measure physical function, and use of different types of ActiGraph measures with different levels of accuracy and calibrations across the different studies. Our finding highlights the importance of assessing all facets of physical function (PRO,

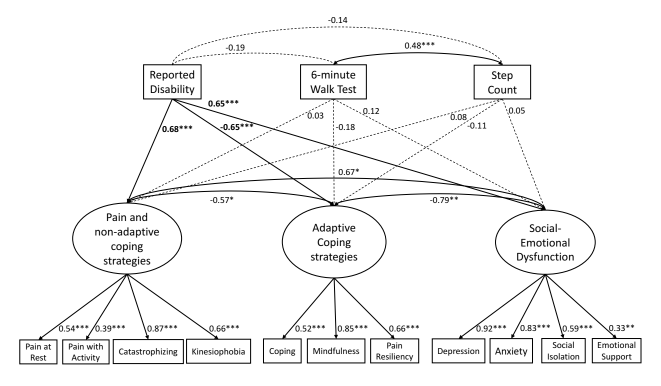


Figure I SEM model.

Notes: Structural model with directional paths specified from the 3 physical function measures to the psychosocial latent variables. Model fit was adequate (Cmin/df = 1.39, CFI = 0.946, RMSEA = 0.070). The bold values represent significant paths for the physical function measurements. The reported significance levels are standard as follows: *p < 0.05, **p < 0.01, **p < 0.001.

functional capacity/performance-based and ambulatory measured step-count) which are not redundant but provide complementary information regarding an individual's overall physical function, and supports recommendations from IMMPACT and ICF.

Testing our second research question revealed that all latent psychosocial factors (ie, pain and non-adaptive coping, adaptive coping, and social-emotional dysfunction) significantly correlated with PRO physical function, while none correlated with 6MWT and objective/ambulatory physical function. This supports previous findings highlighting the multidimensionality of chronic pain²⁵ and the link between psychosocial factors and PRO measured physical function^{8,26} but not objective/ambulatory-measured physical function, 12,21,28 with associations with performance-based physical function being mixed.^{21,27,28,78} This finding further highlights the importance of a comprehensive assessment of all aspects of physical function. Our results confirm the potential of improvement in PRO physical function through commonly targeted psychosocial factors, while pointing to the need for additional targets in order to also improve functional capacity and ambulatory activity, which are important aspects of physical function with strong implications for morbidity and mortality.

Taken together, these findings have several important implications. It is important to include PROs, performance-based and objective measures of physical function, in line with IMMPACT criteria,^{8,9} as each provides important and unique information about particular aspects of physical function.^{21,22} Each measurement has its own advantages and limitations which should be taken into consideration in interventions for chronic pain. For instance, PROs provide information about the patient's perceptions and experiences, which are critical determinants of their well-being,79 but provide limited information regarding their functional capacity or ambulatory activity. Performance-based measures provide information on an individual's functional capacity during a timelimited test that may be influenced by an individual's motivation, without capturing other information regarding activity in the individual's day to day life.⁸ Accelerometers can provide useful and more objective activity estimates of an individual's ambulatory activity across a longer specified time period,⁸⁰ but do not capture important activities of daily living such as dressing, eating, engagement in domestic activities, school or work, which are better captured by PROs. Further, accelerometers may pose a burden to participants and are limited to the degree to which they

are worn, highlighting the importance of carefully designed research protocols to ensure reliable and valid assessment of step-count.

The lack of association between any of the common psychosocial intervention targets in chronic pain programs (eg, pain and non-adaptive coping, adaptive coping, and social-emotional function) and performance-based or objective measures of physical function suggests another implication - in order to improve patients' performance - and accelerometer-based physical function, other components are needed. IMMPACT recommends directly targeting increased activity,⁸ and current skills targeting activity such as setting activity goals, quota-based activity pacing and behavioral activation may be particularly promising.^{33,34} Using a commercially available digital monitoring device such as a Fitbit offers the potential to help individuals monitor their step count and reinforce activity goals.^{33,43} Finally, our findings highlight the extent of associations within measurement modalities and dissociations across measurement modalities. There were significant and mostly moderate-tostrong correlations across all of the patient-reported measures, regardless of the specific construct they were intended to assess. Patients' subjective experiences tend to converge and are evident across PROs. This offers numerous avenues through which interventions may be sensitive to patientreported outcomes but also conveys how little these relate to performance and objective physical function outcomes.

The current study has several strengths. First, it provides a rare comprehensive assessment of physical function among patients with chronic pain, using patientreported, performance-based and objective measures, in accordance with IMMPACT and ICF guidelines.8 Second, we used SEM to test relationships between variables. This approach enables concurrent examination of the relationships between a multitude of factors, as well as consideration of latent variables that cannot be directly observed. Third, we used step-count as our accelerometer output variable. Recent data suggest that step-count is better correlated with measures such as physical activity energy expenditure,⁸¹ compared to accelerometer activity intensity measures such as moderate-to-vigorous activity (MVPA⁸¹). Importantly, step-count provides information regarding movement regardless of intensity, which may optimally capture the construct of physical function and performance of activities of daily living. Finally, to objectively measure step-count, we used hip-mounted ActiGraph accelerometers, which maximize accuracy and bypass the overestimation of steps in daily living associated with wrist-worn devices such as Fitbit.^{82,83}

Some limitations of the study should also be considered. The sample size is small, and our model may have overfit our data. Our SEM approach may be seen as justified, nevertheless, given the relatively strong factor loadings and overall fit indices. Additionally, the latent variables were highly correlated with each other, raising some doubt about whether they represent distinct constructs. This is consistent with the significant overlap found between psychological measures in pain samples which may reflect patients' overall underlying emotional distress.⁸⁴ The cross-sectional study design precludes the ability to determine causality in this study. Finally, generalizability may be limited by the fact that participants in this study had all signed up to participate in an intervention and were recruited from a single urban center. While testing the subsection of the population with chronic pain who are willing and able to participate in interventions may help inform treatment programs directed specifically at these patients, future studies may strengthen the current findings by testing participants from multiple sites and using larger samples with longitudinal assessments.

Conclusion

This study was the first to examine patient-reported, performance-based, and objective physical function along with commonly used psychosocial targets in a single model among patients with chronic pain. Findings confirm that patient-reported, performance-based, and objective assess different aspects of function. Targeting psychosocial variables such as pain and non-adaptive coping, adaptive coping, and social-emotional dysfunction has the potential to improve patient-reported physical function, but additional targets may be needed to improve patients' functional capacities and ambulatory activities. Future research can extend our findings by examining the longitudinal association between the 3 physical function assessments and the PROs during chronic pain interventions.

Abbreviations

PF, physical function; IMMPACT, Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials; ICF, International Classification of Functioning, Disability and Health; PRO, patient-reported outcomes; 6MWT, 6-minute walk test; WHODAS, World Health Organization Disability Assessment Schedule; NRS, Numerical Rating Scale; PCS, Pain Catastrophizing Scale; TSK, The Tampa Scale for Kinesiophobia; MOCS-A, The Measures of Current Status; PRS, Pain Resilience Scale; CAMS-R, Cognitive and Affective Mindfulness Scale – Revised; SEM, structural equation modeling; CFA, confirmatory factor analysis; KMO, Kaiser-Meyer-Olkin; RMSEA, root-mean-square error approximation.

Author Contributions

All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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Disclosure

The authors declare that they have no conflicts of interest.

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