











Sleep Disturbance in Musculoskeletal Conditions: Impact of a Digital Care Program

Justin K Scheer ¹, Fabíola Costa ², Dora Janela ², Maria Molinos ², Anabela C Areias ², Robert G Moulder ³, Jorge Lains ^{4,5}, Virgílio Bento ², Vijay Yanamadala ^{2,6,7}, Fernando Dias Correia ^{2,8}

¹Department of Neurological Surgery, University of California, San Francisco, CA, USA; ²Sword Health, Inc, Draper, UT, USA; ³Institute for Cognitive Science, University of Colorado Boulder, Boulder, CO, USA; ⁴Rovisco Pais Medical and Rehabilitation Centre, Tocha, Portugal; ⁵Faculty of Medicine, Coimbra University, Coimbra, Portugal; ⁶Department of Surgery, Quinnipiac University Frank H. Netter School of Medicine, Hamden, CT, USA; ⁷Department of Neurosurgery, Hartford Healthcare Medical Group, Westport, CT, USA; ⁸Neurology Department, Centro Hospitalar e Universitário do Porto, Porto, Portugal

Correspondence: Fernando Dias Correia, Sword Health Inc, 13937 Sprague Lane Ste 100, Draper, UT, 84020, USA, Tel +1 385-308-8034, Fax +1 801-206-3433, Email fcorreia@swordhealth.com

Background: Musculoskeletal (MSK) pain is highly prevalent worldwide, resulting in significant disability, and comorbid sleep disturbances. Digital therapy for MSK pain can provide significant improvements in care access, alongside pain and disability reductions. However, studies on the effect of such programs on sleep are lacking.

Purpose: To evaluate the impact on pain-related sleep impairment after a 12-week remote multimodal digital care program (DCP) for MSK conditions.

Patients and Methods: This is an ad-hoc analysis of a decentralized single-arm study into engagement and clinical outcomes after a DCP for MSK rehabilitation. Patients were stratified by baseline sleep disturbance, based on sleep questions in the questionnaires: Oswestry Disability Index, Neck Disability Index, and the Quick Disabilities of the Arm, Shoulder and Hand questionnaire. Additional outcomes were pain, Generalized Anxiety Disorder 7-item scale, Patient Health 9-item questionnaire, Work Productivity, and Activity Impairment, and program engagement.

Results: At baseline, 5749 patients reported sleep disturbance (78.0% of eligible patients). These reported significantly worse clinical outcomes at baseline than patients without sleep disturbance (all $p < 0.001$). Patients with comorbid sleep disturbance showed improvements in sleep, with a significant proportion reporting full recovery at program completion: 56% of patients with upper limb conditions (including 10% of patients with severe sleep disturbance at baseline), and 24% with spine conditions. These patients also reported significant improvements in all clinical outcomes at program completion. Engagement and satisfaction were high, and also higher than in patients without sleep impairment.

Conclusion: This is the first study of its kind investigating the effect of a completely remote DCP for MSK pain on sleep. Patients reporting comorbid sleep disturbance had significant improvement in sleep, alongside pain, mental health and work productivity at program completion. The results suggest that a DCP for MSK pain can improve sleep disturbances in patients with upper limb and spine conditions.

Keywords: pain, physical therapy, eHealth, telerehabilitation, remote care

Introduction

High quality restful sleep is essential to one's health and well-being. Sleep deprivation may arise from different etiologies and it is associated with many medical conditions, such as obesity,^{1,2} cancer,^{3,4} type 2 diabetes,⁵ and cardiovascular disease.^{6,7} Poor sleep and pain interaction share a complex and reciprocal relationship: poor sleep can exacerbate pain, and pain can disrupt restorative nights of sleep.⁸⁻¹⁰ Moreover, poor sleep is associated with the development of chronic pain, especially musculoskeletal (MSK) conditions.¹¹⁻¹³ On the other hand, MSK conditions result in significant disability and suffering, with a very high worldwide prevalence of 1.71 billion.^{14,15} Sleep disorders are highly comorbid with MSK conditions with a reported prevalence of 72.1% for chronic back pain, 65.4% for rheumatoid arthritis, 70.3%

for osteoarthritis, and 95.5% for fibromyalgia.¹² Patients with both chronic pain and sleep disturbances have a longer duration of pain and greater pain severity¹⁶ and in addition, these patients have more mental health comorbidities, such as pain catastrophizing, anxiety, and depression.¹⁶

Currently, exercise-based physical therapy is the mainstay initial treatment for MSK conditions.^{14,17–19} Not only is it generally effective in addressing MSK pain, it can also improve sleep quality.^{20–24} A recent meta-analysis by Kelley et al demonstrated that exercise interventions resulted in significant improvements in overall sleep quality, subjective sleep, and sleep latency.²² Moreover, a recent randomized controlled trial by Tseng et al found that moderate-intensity exercise training had a significant beneficial effect on sleep quality and cardio-autonomic function.²¹

Within physical therapy interventions, there is a recent trend towards telerehabilitation and digital physical therapy.^{25–27} These programs are effective alternatives to traditional physical therapy^{28–34} and can increase access to treatment, as well as affordability by reducing travel limitations, time constraints, and geographic restrictions. Digital therapy can also increase patient compliance, empowerment, and self-management.^{27,35} Overall, telerehabilitation has been well-accepted by patients with MSK conditions, as suggested by the reported high satisfaction rates, similar to those reported after in-person rehabilitation.^{36,37} To our knowledge, however, no study has been conducted to evaluate the impact of a telerehabilitation program for MSK conditions on pain-related sleep disturbances. In prior studies, we have reported that a multimodal digital care program (DCP) combining exercise-based physical therapy with a psychoeducational component provides a comprehensive approach to pain management with improvements in pain, mental health, and work impairment.^{31–33,38,39}

The purpose of the present study was to evaluate the impact on sleep impairment related to MSK pain after a 12-week remote, multimodal digital care program (DCP) for MSK conditions, with the hypothesis that patients would experience an improvement in sleep disturbance following completion of the program.

Material and Methods

Study Design

The current study is a post-hoc analysis of a decentralized, single-arm investigation to address the impact of a home-based, multimodal DCP on sleep in patients reporting MSK pain in the upper limb or spine. This study is part of a trial that was prospectively approved by the New England Institutional Review Board (number 120190313) and registered on ClinicalTrials.gov (NCT04092946) on September 17th 2019. The study was conducted in accordance with the Declaration of Helsinki. The DCP was administered at the patients' home and was delivered between June 29th 2020 and November 4th 2021.

Population

The study population included adults (>18 years of age) participating in employer health plans from a total of 50 states and the District of Columbia in the US. Patients that presented MSK pain were invited to apply for the Sword Health's DCP (Draper, Utah, USA) through a dedicated enrollment website (pre-selecting those with ability to interact with technologies). All participants completed a baseline form providing demographic data and details regarding the clinical condition, alongside specific questions to screen for potential clinical red flags, which was posteriorly assessed by an assigned physical therapist (PT) through an onboarding video call. Any patient that selected any level of MSK-related sleep problem at baseline on the validated metrics were included in the present study. Exclusion criteria included the following: (1) a health condition (eg cardiac, respiratory) or any contra-indication incompatible with performing at least 20 minutes of light to moderate exercise; (2) serious injury or clinical red flags not cleared by a physician; (3) receiving treatment for active cancer; and (4) reporting rapidly progressive loss of strength and/or numbness in the arms/legs or unexplained change in bowel or urinary function in the previous 2 weeks; (5) inability to understand simple and complex motor commands; and (6) not reporting sleep problems. Informed consent was obtained from all patients. Since the recommended validated scales to assess hip or knee conditions by International Consortium for Health Outcomes Measurement (ICHOM) – Hip or Knee and Osteoarthritis Outcome Score – do not account specifically for the sleep domain, we decided to not include patients with lower limb MSK conditions in the present

study and include only those with spine or upper limb pain. Any patient that selected any level of MSK-related sleep problem at baseline on the validated metrics were included in the present study. Patients not reporting sleep problems were excluded.

Intervention

The DCP has been previously described in a number of studies.^{33,39} In brief, the program consists of a 4-, 8-, or 12-week telerehabilitation intervention that includes exercise, education and cognitive behavioral therapy (CBT). This program digitally interfaced between the patient and an assigned physical therapist (PT) who monitors the patients for the study duration. An FDA-listed class II medical device that consists of inertial motion trackers, a dedicated tablet with a mobile app, and a cloud-based portal was made available to all patients. The exercise sessions were designed following current guidelines and were prescribed and tailored by the PT according to the patient's condition and progression. Exercise consisted of gradual progressive movement exposure, including exercises focused on mobility/stretching, strength, and balance. Progress would include an increase of: (i) number of repetitions and sets, (ii) external load, (iii) range of motion, and (iv) inclusion of multi-articular and multidirectional exercises. Patients performed personalized exercise sessions independently at their convenience through the tablet display, with 3 sessions per week being recommended as a base scenario. The tablet displayed an audio-video explanation prior to each exercise set. During the exercise, an execution interface was shown, with real-time audio-video biofeedback based on the motion trackers data. The devices connected to a cloud-based portal, enabling asynchronous and remote monitoring by the assigned PT, who performed data-driven adjustments to the exercise program as needed. The education and CBT components of the program were developed according to current clinical guidelines and prior research.^{40–43} Education was also personalized according to each MSK condition addressing the following topics: anatomy, physiology, symptoms, evidence-based treatments, fear-avoidance, and active coping skills, including dealing with feelings of anxiety, depression, and sleep hygiene. The CBT program was based on third generation CBT techniques – mindfulness, acceptance and commitment therapy, and empathy-focused therapy. The education and CBT materials were delivered to patients through written articles, audio content and interactive modules. Bi-directional communication with the assigned PT was ensured through a built-in secure chat within a smartphone application and video calls. Patients who did not engage in any exercise session for 28 consecutive days were considered dropouts.

Demographic Data

Demographic data collected included age, body mass index (BMI), patient gender, and employment status. The gender category included man, woman, and non-binary. The employment status categories were defined as employed (full time or part time) or unemployed.

Clinical Outcomes

Outcomes were collected at baseline, 4, 8 and 12 weeks.

1. Primary outcome was sleep metrics from the following validated questionnaires:
 - (a) Quick Disabilities of the Arm, Shoulder, and Hand questionnaire (QuickDASH), an 11-item questionnaire with a Likert scale addressing disability and symptom severity. Scores range from 0% to 100% (higher scores indicating poorer functioning).⁴⁴
 - i. Sleep question: “During the past week, how much difficulty have you had sleeping because of the pain in (your condition)? 1 – No difficulty; 2 – Mild difficulty; 3 – Moderate difficulty; 4 – Severe difficulty; 5 – So much difficulty that I can't sleep.”
 - (b) Oswestry disability index (ODI), a 10-item questionnaire with a Likert scale addressing the impact of back pain on multiple dimensions of the patients' life. Scores range from 0% to 100% (higher scores indicate more disability).⁴⁵

- i. Sleep question: “Sleeping: 1) My Sleep is never disturbed by pain; 2) My sleep is occasionally disturbed by pain; 3) Because of pain I have less than 6 hours sleep; 4) Because of pain I have less than 4 hours sleep; 5) Because of pain I have less than 2 hours sleep; 6) Pain prevents me from sleeping at all.”
 - (c) Neck Disability Index (NDI), a 10-item questionnaire with a Likert scale addressing the impact of neck pain on multiple dimensions of the patients’ life. Scores range from 0 to 50 (higher scores indicate more disability).⁴⁶
 - i. Sleep question: “Sleeping: 1) I have no trouble sleeping; 2) My sleep is slightly disturbed (less than 1 hour sleepless); 3) My sleep is mildly disturbed (1–2 hours sleepless); 4) My sleep is moderately disturbed (2–3 hours sleepless); 5) My sleep is greatly disturbed (3–5 hours sleepless); 6) My sleep is completely disturbed (5–7 hours sleepless).”
2. Secondary outcomes:
- (a) Self-reported pain, using the Numerical Pain Rating Scale (NPRS), through the question: “Please rate your average pain over the last 7 days” from 0 (no pain at all) to 10 (worst pain imaginable”).
 - (b) Generalized Anxiety Disorder 7-item scale (GAD-7) (range 0–21)⁴⁷ to assess anxiety, and Patient Health 9-item questionnaire (PHQ-9) (range 0–27) to assess depression (higher scores indicate worse mental health).⁴⁸
 - (c) Work Productivity and Activity Impairment (WPAI) for general health questionnaire, evaluated employed patients to assess overall work impairment (WPAI overall: total presenteeism and absenteeism from work), presenteeism (WPAI work), absenteeism (WPAI time) and non-work-related activities impairment (WPAI activity). Scores range from 0% to 100% (higher scores indicate declined productivity).⁴⁹
 - (d) Engagement: measured through the following: A) completion of the program (considered as the retention rate); B) total number of completed exercise sessions; C) number of sessions per week, D) total time spent performing exercise sessions; E) total articles read, F) total messages exchanged between the physical therapist and the patient, and G) overall satisfaction through the question: “On a scale from 0 to 10, how likely is it that you would recommend this intervention to a friend or neighbor?”.

Statistical Analysis

Descriptive statistics were applied to characterize the study population and included continuous variables being described with the mean and standard deviation. Categories were described with frequencies (percentage). Comparison of baseline means between the groups initially included an analysis of variance (ANOVA). Frequency analyses for categorical variables were conducted via Pearson’s χ^2 analysis. Patients that completed the program were defined as “completers” and those who did not were defined as “non-completers”. Comparisons were made between completers and non-completers.

For the longitudinal data, latent growth curve analysis (LGCA) was used to estimate clinical outcome trajectories across the digital program based on the individual patient trajectories and considering time as a continuous variable.⁵⁰ This methodology is in the same family of linear mixed-effects modeling however, it is estimated as a structural equation model.⁵¹ This type of model considers repeated measures on the same individual to be correlated. LGCA includes measures of model fit and full information maximum likelihood (FIML) to deal with any missing data.⁵² FIML considers all available data at each time point from all patients. Prior studies have demonstrated the superiority of FIML compared to other imputation methods.⁵² For the sleep disturbance analysis, ordinal regression was conducted with the R package “MASS”, taking into account the different item-ranged questions. Ordinal regression is a method for modeling dependent variables that are ordinal in nature by assuming an underlying latent continuous distribution behind the observed ordinal data. This latent distribution is assumed to be Gaussian and has a mean of 0 and a standard deviation of 1. Ordinal regression models determine threshold values on this latent distribution that best estimate the observed proportions of each ordinal category. Independent variables were tested for statistical significance by allowing them to influence the location of the mean of this distribution, thus changing the observed proportions of each ordinal category based upon the estimated thresholds. Significant independent variables indicate a reliable shift in the underlying latent distribution

greater than expected by chance. In the case of the current data, we evaluated if the latent distribution of sleep quality ordinal categories changed significantly from pre to post.

To facilitate interpretation of the results, 3 subgroups were created post-analysis. For QuickDASH: No disturbance = 1; Mild/Moderate disturbance = 2 and 3, and Severe disturbance = 4 and 5. For ODI and NDI: No disturbance = 1; Mild/Moderate disturbance = 2 and 3, and Severe disturbance = 4, 5 and 6.

All statistical analyses were conducted using commercially available software (SPSS v22, IBM, Armonk, NY) or coded using R (version 1.4.1717, R Foundation for Statistical Computing). The level of significance was set at $p < 0.05$ for all tests.

Results

A total of 5749 patients reported some level of sleep disturbance at baseline (78.0%) and were therefore selected for the study. From these, 1250 (21.7%) were dropouts and 448 (7.8%) were excluded, with 4051 (70.5%) of patients completing the program. The study flow diagram is presented in Figure 1. No serious adverse events^{53,54} were reported during the study.

Baseline Characteristics

Baseline demographics of both the entire cohort and stratified by upper limb or spine pain are presented in Table 1. Overall, patients were middle aged (50.8 ± 12.7 years), overweight, with mean BMI 29.1 ± 6.5 , majorly composed of women (56.7%) and employed individuals (81.9%, Table 1). In addition, a majority of patients reported chronic MSK pain (79.1%).

Patients with upper limb conditions (wrist/hand, elbow and shoulder), were significantly younger (49.6 ± 10.6 vs 51.4 ± 13.4 years, $p < 0.001$), had a lower mean BMI (28.5 ± 6.4 vs 29.3 ± 6.6 , $p < 0.001$), and a higher proportion of acute

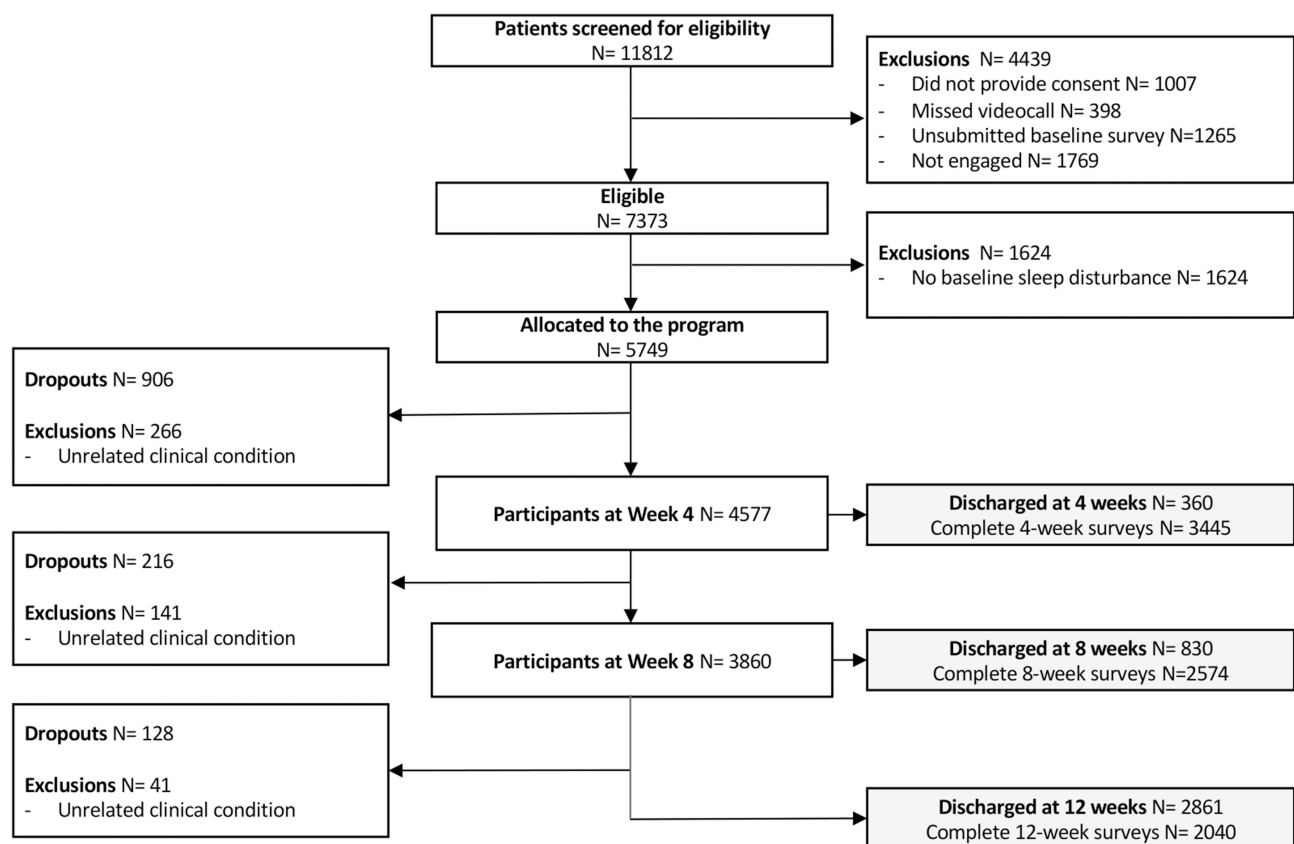


Figure 1 Flowchart of the study.

Table I Baseline Characteristics of Study Patients

Characteristic	Upper Limb (N=1724)	Spine (N=4025)	P value	Entire Cohort (N=5749)
Age (years), mean (SD)	49.6 (10.6)	51.4 (13.4)	<0.001	50.8 (12.7)
Age categories, N (%):			<0.001	
<25	12 (0.7)	33 (0.8)		45 (0.7)
25–40	357 (20.7)	936 (23.2)		1293 (22.5)
40–60	1071 (62.1)	2000 (49.6)		3071 (53.4)
> 60	284 (16.5)	1056 (26.2)		1340 (23.3)
Gender, N (%) ^a :			0.305	
Female	943 (54.7)	2320 (57.6)		3263 (56.7)
Male	776 (45.0)	1693 (42.1)		2469 (42.9)
Non-binary	3 (0.2)	9 (0.2)		12 (0.2)
BMI, mean (SD) ^b	28.5 (6.4)	29.3 (6.6)	<0.001	29.1 (6.5)
BMI categories, N (%) ^b :			<0.001	
Underweight (<18.5)	17 (0.9)	27 (0.7)		44 (0.7)
Normal (18.5–25)	512 (29.7)	1084 (27)		1596 (27.8)
Overweight (25–30)	660 (38.3)	1346 (33.5)		2006 (34.9)
Obese (30–40)	427 (24.8)	1271 (31.7)		1698 (30)
Obese grade III (>40)	106 (6.2)	287 (7.1)		393 (6.8)
Employment status, N (%) ^c			<0.001	
Employed (part-time or full-time)	1566 (90.9)	3143 (78.1)		4709 (81.9)
Unemployed	114 (6.6)	777 (19.3)		891 (15.4)
Acuity, N (%) ^d			<0.001	
Acute (< 12 weeks pain)	546 (32.1)	646 (16.2)		1192 (20.9)
Chronic (> 12 weeks pain)	1154 (67.8)	3344 (83.8)		4498 (79.1)
Outcome measures, mean (SD)				
Sleep problems, N(%):			<0.001	
Moderate	1527 (89)	3924 (97.5)		5451 (95)
Severe	197 (11)	101 (2.5)		298 (5)
Pain Level ^e	5.00 (1.8)	5.2 (1.9)	<0.001	5.1 (1.9)
GAD-7 ≥5 ^f	8.5 (3.8)	9.1 (4.1)	0.010	8.9 (4.1)
GAD-7	2.9 (4.1)	3.7 (4.6)	<0.001	3.5 (4.5)
PHQ-9 ≥5 ^g	8.5 (3.5)	9.6 (4.5)	<0.001	9.3 (4.4)
PHQ-9	2.2 (3.6)	3.2 (4.7)	<0.001	2.9 (4.5)
WPAI Overall >0 ^h	28.7 (18.9)	30.9 (20.8)	0.005	30.3 (20.3)
WPAI Overall ⁱ	17 (20.3)	20.8 (22.4)	<0.001	19.5 (21.8)
WPAI Work >0 ^j	27.3 (17.8)	29.2 (19.1)	0.014	28.6 (18.7)
WPAI Work ⁱ	15.9 (19.1)	19.3 (20.8)	<0.001	18.16 (20.3)
WPAI Time >0 ^k	18.6 (16.0)	18.4 (18.0)	0.922	18.5 (17.6)
WPAI Time ⁱ	1.7 (7.2)	2.9 (9.8)	<0.001	2.5 (9.0)
WPAI Activity >0 ^l	35.4 (21.4)	38.7 (22.9)	<0.001	37.7 (22.5)
WPAI Activity	29.3 (23.7)	31.7 (25.5)	0.008	31 (25.0)

Notes: ^a5 missing values; ^b10 missing values; ^c149 missing values; ^d59 missing values; ^e51 missing values; ^fN= 1807 (entire cohort), N=458 (upper limb), N=1349 (spine); ^gN=1451 (entire cohort), N= 337 (upper limb) N= 1114 (spine); ^hN=2831 (entire cohort), N=862 (upper limb), N=1969 (spine); ⁱN=4390; ^jN=2787 (entire cohort), N=850 (upper limb), N=1937 (spine); ^kN=595 (entire cohort), N=134 (upper limb), N=462 (spine); ^lN=4719 (entire cohort), N=1426 (upper limb), N=3293 (spine).

Abbreviations: BMI, body mass index; GAD-7, Generalized Anxiety Disorder 7-item scale; PHQ-9, Patient Health 9-item questionnaire; WPAI, Work Productivity and Activity Impairment Questionnaire.

conditions than patients with spine pain (32.1% vs 16.2%, $p<0.001$). By contrast, there was a significantly larger proportion of unemployed individuals within those with spine (neck and back) conditions (19.3% vs 6.6%, respectively, $p<0.001$). This group also reported worse mean mental health scores at baseline, particularly when filtering for scores ≥ 5 (GAD 9.1 ± 4.1 vs 8.5 ± 3.8 and PHQ 9.6 ± 4.5 vs 8.5 ± 3.5 , $p=0.01$ and $p<0.001$, respectively) and work impairment (WPAI overall >0 30.9 ± 20.8 vs 28.7 ± 18.9 , $p=0.005$ [Table 1](#)).

Comparing completers with non-completers, the latter were significantly younger (46.2 ± 11.0 vs 51.8 ± 12.5 years, $p<0.001$) and reported higher levels of BMI (30.1 ± 7.2 vs 28.6 ± 6.2 , $p<0.001$, [Supplementary Table S1](#)), with no additional demographic differences between groups ($p>0.05$ for all).

When comparing patients with and without sleep disturbance at baseline, the former were significantly older, had higher mean BMI, and presented higher proportions of women, unemployed patients, chronic pain sufferers and conditions affecting the spine ($p<0.001$ for all, [Supplementary Table S2](#)). They also reported worse baseline mean scores for pain, anxiety, depression, and WPAI ($p<0.001$ for all, [Supplementary Table S2](#)).

Improvement in Sleep Disturbance

As depicted in [Figure 2](#) and [Table 2](#), we observed an improvement in patients reporting baseline sleep disturbance, with a shift in distribution onto lower sleep impairment categories. The highest improvements were observed in those with upper limb conditions, with 56% of patients reporting total recovery on sleep impairment, with 46% corresponding to patients with moderate disturbance at baseline and 10% to patients reporting severe disturbance at baseline. Among patients with spine conditions, 24% reported complete recovery, all from the group reporting moderate baseline disturbance.

Clinical Outcomes

Overall, patients reported improvements in all clinical outcomes at program end. Pain decreased on average 2.42 points (95% CI 2.33; 2.50), while anxiety and depression (among those with scores ≥ 5) decreased by 4.15 (95% CI 3.83; 4.49)

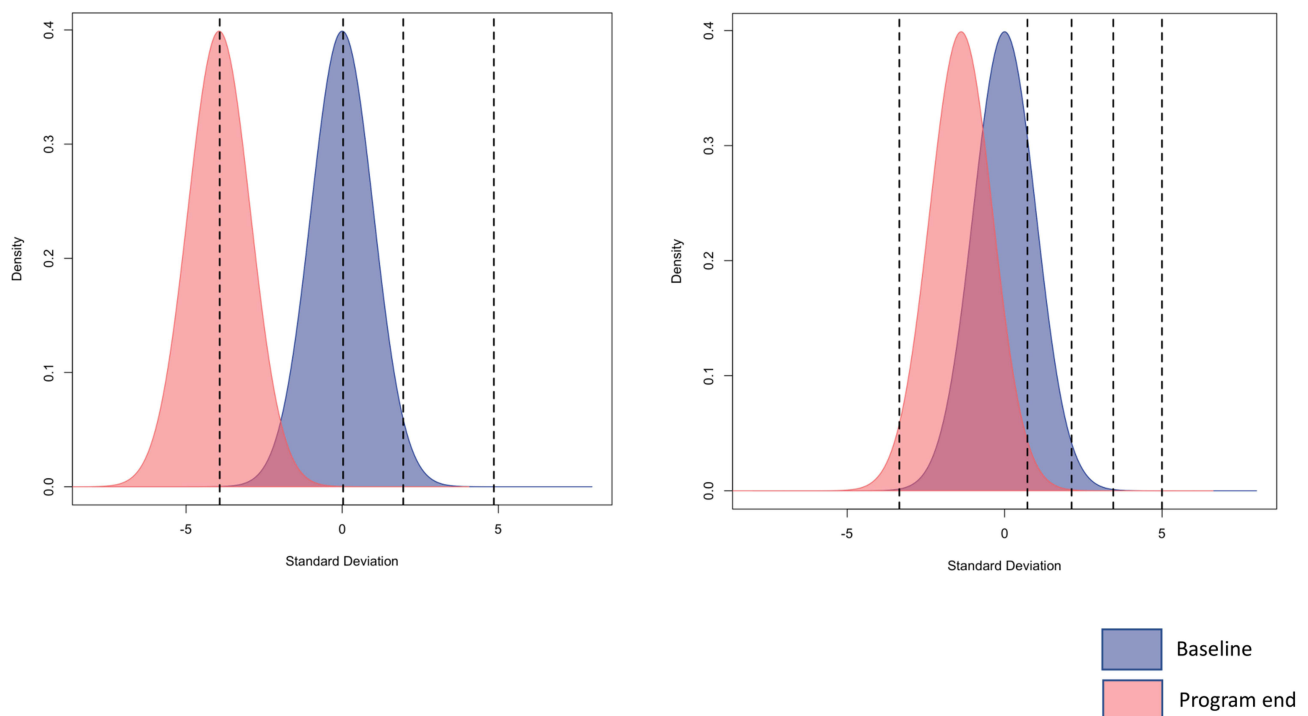


Figure 2 Results from the ordinal regression showing the distribution of the population at baseline and program end. The right plot refers to the results for those with upper limb conditions and the left for spine conditions. The dashed vertical lines represent the thresholds (cut-offs derived from ordinal regression analysis) which establish the boundaries between the response levels (the 5 and 6 items-questionnaires for upper limb and spine, respectively).

Table 2 Ordinal Regression Analysis of the Pre-Post Proportions of the Sleep Categories Distribution

Pre/post	Estimate	95% CI		t	P value
Upper limb	-3.94	-4.29	-3.61	-22.62	< 0.001
Spine	-1.38	-1.53	-1.23	-18.18	< 0.001

Notes: Estimate refers to the change of the different categories' mean distribution between baseline and treatment end, ie to what extent the mean distribution will shift towards a lower category (as seen here) or to a higher category (in that case the estimate would have a positive value).

and 4.42 (95% CI 3.98; 4.86) points, respectively. Regarding work productivity, among those with WPAI scores > 0 at baseline, we observed an improvement of 15.32 (95% CI 14.03; 16.61) points, as well as an improvement of 19.48 (95% CI 18.47; 20.49) points in non-work-related activity impairment ([Supplementary Table S3](#)).

Importantly, greater mean changes were observed for pain and impairment in both work- and non-work-related activities in patients with sleep disturbance at baseline, in comparison with those without sleep impairment ([Supplementary Table S4](#)).

Impact of Covariates on the Sleep Categories Distribution

The impact of baseline characteristics on changes in sleep disturbance category are presented in [Table 3](#). Patients with higher baseline pain scores presented lower chances of improvement on sleep disturbance ([Table 3](#)). In addition, within patients with spine conditions, those with chronic pain, higher depression (PHQ-9) scores at baseline and women were less likely to improve ([Table 3](#)). None of the other demographic characteristics had a significant impact on sleep disturbance changes.

Table 3 Impact of Covariates (Demographic and Clinical) on Sleep Disturbance Changes

Therapy Area	Variable Type	Variable	Estimate	95% CI		t	P value
Upper limb	Demographic	Post	-6.32	-7.30	-5.48	-13.77	
		Age	0.01	-0.02	0.03	0.59	0.552
		Gender=woman	0.42	-0.10	0.94	1.58	0.115
		BMI	-0.01	-0.05	0.03	-0.56	0.574
	Clinical	Post	-5.62	-6.74	-4.62	-10.47	
		Pain	0.38	0.16	0.62	3.28	0.001
		GAD-7	-0.03	-0.16	0.10	-0.43	0.670
		PHQ-9	0.08	-0.08	0.24	0.97	0.331
Acuity=Chronic	0.57	-0.04	1.19	1.83	0.068		
Spine	Demographic	Post	-3.50	-3.94	-3.06	-15.56	
		Age	0.01	-0.01	0.02	0.75	0.451
		Gender=Woman	0.45	0.12	0.78	2.65	0.008
		BMI	0.03	0.00	0.05	1.87	0.062
	Clinical	Post	-3.49	-4.03	-2.97	-12.91	
		Pain	0.37	0.27	0.46	7.79	< 0.001
		GAD-7	0.04	-0.03	0.10	1.04	0.298
		PHQ-9	0.10	0.03	0.16	2.86	0.004
Acuity=Chronic	0.58	0.13	1.02	2.56	0.010		

Notes: Post refers to the average total change estimate from baseline (pre) until treatment end. The covariates influence on this mean change can be calculated by adding the covariate estimate to the post estimate. Eg. when accounting for having a chronic condition in upper limb, the final estimate would be -5.76, meaning that the shift would be less pronounced/probable and therefore a smaller number of patients would shift category (recover from sleep).

Abbreviations: BMI, body mass index; GAD-7, Generalized Anxiety Disorder 7-item scale; PHQ-9, Patient Health 9-item questionnaire.

Table 4 System Usability-Related Outcomes

Usability Outcomes	Upper Limb	Spine	P value	Entire Cohort
Number of sessions per week	2.8 (1.2)	2.8 (1.1)	0.651	2.8 (1.2)
Number of sessions	29.3 (21.3)	29.7 (21.2)	0.521	29.53 (21.2)
Total exercising time (min)	359.1 (263.9)	366.6 (263.5)	0.337	364.3 (263.6)
Number of messages exchanged	17.5 (23.9)	19.1 (25.8)	0.032	18.6 (25.3)
Total articles read	3.7 (6.8)	3.5 (6.3)	0.447	3.6 (6.7)
Satisfaction (0–10)	8.6 (1.7)	8.7 (1.7)	0.126	8.7 (1.7)

Engagement Outcomes

The engagement outcomes for the entire cohort and stratified by upper limb and spine conditions are listed in Table 4. Similar engagement was observed across the different body areas, except for the communication with the PT, where those with spine conditions presented a higher frequency of contacts. Similar satisfaction values were observed between therapy areas ($p=0.126$).

Notably, compared to those with no comorbid sleep disturbance, patients with baseline sleep impairment had a significantly higher mean number of sessions per week, total number of sessions, number of messages with the PT and total articles read ($p<0.05$ for all, [Supplementary Table S5](#)). They also had a significantly higher mean satisfaction score than those without baseline sleep disturbance (8.6 ± 1.7 vs 8.5 ± 1.8 , respectively, $p<0.001$).

Discussion

Main Findings

This is the first study of its kind investigating the effect of a completely remote digital care plan for MSK pain on sleep improvement. There was a prevalence of baseline sleep disturbance of 78%. Overall, patients with comorbid sleep disturbance showed a recovery pattern, with a significant proportion of these patients reporting complete sleep impairment recovery at program completion. Patients with upper limb conditions had the highest improvements in sleep, with 56% of patients reporting total recovery of sleep impairment (including 10% of patients with severe sleep disturbance at baseline), while 24% of patients with spine conditions recovered completely from their baseline sleep disturbance, despite the higher prevalence of poor prognostic factors in this cohort (female gender, having chronic pain and higher depression scores at baseline). Chances of improvement in sleep disturbance were negatively impacted by baseline pain severity.

Overall, patients with comorbid sleep disturbance at baseline reported significant improvement in mean pain, anxiety, depression and work productivity scores at program completion. Engagement and satisfaction were high in these patients (higher than observed among patients without sleep impairment at baseline).

Comparison with Prior Work

Sleep disturbances are highly prevalent and comorbid with a number of different medical conditions.^{1,5,11,12}

Despite more wide investigation on the association between sleep and MSK pain in chronic widespread pain and chronic low back pain, evidence indicates that this association also exists for a broader range of conditions,^{55–57} including those affecting the upper limb.^{55,58,59} In fact, sleep disturbance is one of the main complaints among patients with shoulder disorders.^{60,61} Within MSK pain sufferers, sleep disorders have a reported prevalence between 65.4% and 95.5%.¹² The present study reported an overall prevalence of 78%, which is in line with the current literature.

MSK conditions with related comorbid sleep disturbance have been associated with poor clinical outcomes including longer pain duration and greater pain severity, as well as higher disability.¹⁶ This was also found in the present study, as patients with comorbid sleep disturbance had significantly worse mean pain, anxiety/depression, and work impairment scores, as well as a higher proportion of chronic pain sufferers, than those without sleep disturbance at baseline.

It has been well established that traditional exercise programs significantly improve MSK pain and sleep.^{20,22} A meta-analysis with a total of 950 patients found a statistically significant improvement in several sleep outcomes with exercise.²² Another meta-

analysis conducted by Kovacevic et al gathering 13 different studies found significant improvements in subjective sleep duration and sleep quality.²⁰ In the present study, patients reported improved outcomes in pain, with a mean reduction greater than the 2-points recommended by the IMMPACT guidelines as the minimal clinically important difference.⁶² Additionally, a high proportion of patients reported improvements in pain-related sleep disturbances, in line with the current literature on exercise and sleep improvement. A greater recovery of sleep impairment was observed in patients with upper limb conditions. Patients with spine conditions presented several poor prognostic factors, such as chronic conditions,¹⁶ higher baseline pain severity,⁶³ poorer mental state⁶⁴ and a larger proportion of women.⁶⁵ Women have been reported to have higher prevalence of MSK conditions,⁶⁶ associated with higher pain, mental distress and sleep disorders.^{67–69} These results emphasize the opportunity of taking advantage of the adaptability of digital multimodal programs, to better address the specific needs of high-risk patients.

Patients also reported improvement in both anxiety and depression, which are domains highly comorbid with both chronic MSK pain and sleep disorders,^{11,12,70,71} and whose association tends to result in lower improvement and worse outcomes.^{11,72} This is in agreement with the reported in the present study, as patients with spine pain and more severe mental health issues were more likely to end up in a more severe sleep category at the program completion. It is important to note that all of these comorbidities are likely interrelated, with varying synergistic effects on one another. This highlights the importance of adopting a biopsychosocial view in the management of MSK conditions, considering other aspects involved which influence health and recovery besides pain. Improving both pain and mental status on patients, through a DCP including not only exercise, but also education and CBT, might have contributed for synergistic improvements in all clinical outcomes measured.

Overall engagement in the program was high, with a high satisfaction rate similarly reported by patients with spine or upper limb pain. Patients with comorbid sleep disturbances had a significantly higher engagement in the program than those without baseline sleep disturbances at baseline, specifically with regard to more sessions, exercising time, interactions with the PT, and articles read. We speculate that higher baseline pain levels, together with existing sleep disturbances may have positively influenced the motivation of these patients to engage in the program.

Strengths and Limitations

There are many strengths to this study, one of which is the large sample size from a real-world context. Patients were enrolled through their employers' health plans from 50 states and the District of Columbia in the U.S. allowing for a diverse population and better generalizability. Another strength is the DCP itself, which is a multimodal approach that includes exercise, utilizing real-time biofeedback, regular communication with the same PT, as well as education and CBT, in a digital format. All these factors favor accessibility, engagement and can therefore maximize outcomes, as well as address both physical and mental health aspects jointly. An additional strength is the use of self-reported sleep outcomes, which are integrated in widely used and validated outcome scales, allowing for translational application and generalizability to other studies. Lastly, the novel idea of investigating sleep disturbance within a digital therapy program allows for a foundation for future work in this field, and helps identify areas of improvement for future telerehabilitation programs.

The major limitation of the study is the lack of a control group, which does not allow for any causal effect to be determined on sleep improvement. However, given the real-world context in which this was performed, the most obvious comparator would be a "waiting list" control, which was not ethical considering the high accessibility this technology affords. In addition, because the program includes patients that are beneficiary to their employers' health benefits, the current population may not be an exact representation of the general population in the U.S.. Thus any applicability of the results to clinical settings in which there are more uninsured, elderly, or work-disabled patients may be limited. In addition, patients with lower limb MSK conditions were excluded since ICHOM recommended validated metrics do not account specifically for the sleep domain. It is therefore unknown if these results apply to that population. However, given the improvement in pain and sleep in the other conditions, it is likely they will benefit too. Further studies on this population is warranted to confirm the present findings in that specific population. The sleep metrics used for analysis were from self-reported validated surveys completed by the patient and thus are subjective in nature by definition. However, no specific and dedicated scale was used that focused on sleep disturbance alone. Therefore, more research is needed to better characterize the types of sleep disturbance and improvements in patients with MSK, particularly under a telerehabilitation context. Finally, long-term follow-up was also not available to ascertain the benefits of the program at

later time points and to see if any of the sleep improvements remained. Further prospective controlled studies are warranted to better characterize the effect of digital therapy on sleep disturbance outcomes.

Conclusion

This is the first study of its kind investigating the effect of a completely remote digital care plan for MSK pain on sleep. Patients reporting comorbid sleep disturbance had significant improvement in MSK pain-related sleep disturbance, as well as pain, mental and work productivity. The results suggest that a digital care program for MSK pain can improve sleep disturbances in patients with upper limb and spine conditions.

Abbreviations

MSK, Musculoskeletal; DCP, Digital care program; US, United States; ICHOM, International Consortium for Health Outcomes Measurement; CBT, Cognitive behavioral therapy; PT, Physical Therapist; FDA, United States Food and Drug Administration; BMI, Body Mass Index; QuickDASH, Quick Disabilities of the Arm, Shoulder and Hand questionnaire; NPRS, Numerical Pain Rating Scale; GAD-7, Generalized Anxiety Disorder 7-item scale; PHQ-9, Patient Health 9-item questionnaire; WPAI, Work Productivity and Activity Impairment; ANOVA, Analysis of variance; LGCA, Latent growth curve analysis; FIML, Full information maximum likelihood; NDI, Neck Disability Index; ODI, Oswestry Disability Index; IMMPACT, Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials.

Data Sharing Statement

All data relevant to the study are included in the article or are available as Digital Content at [Supplementary Materials](#). Only de-identified individual participant data is provided. Further information, including the study protocol, can be found at ClinicalTrials.gov (NCT04092946).

Ethics Approval and Informed Consent

The study was approved by the New England IRB (protocol number 120190313) and prospectively registered in ClinicalTrials.gov, NCT04092946, 17/09/2019. This study was conducted in accordance with the approved guidelines. All patients were informed about the purpose and procedures of the study and provided informed consent.

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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

Fabiola Costa, Dora Janela, Maria Molinos, Anabela C. Areias, Virgílio Bento, Vijay Yanamadala and Fernando Dias Correia are employees at Sword Health, the study sponsor. Robert G. Moulder is an independent scientific consultant responsible for statistical analysis, while Jorge Lains and Justin K. Scheer are independent scientific/clinical consultants who were funded by Sword Health in connection with the development and execution of this article. Fernando Dias Correia, Vijay Yanamadala and Virgílio Bento also hold equity from Sword Health. The authors report no other conflicts of interest in this work.

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