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Published in:
Journal of Pain Research

DOI (link to publication from Publisher):
[10.2147/JPR.S306504](https://doi.org/10.2147/JPR.S306504)

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Publication date:
2021

Document Version
Publisher's PDF, also known as Version of record

[Link to publication from Aalborg University](#)

Citation for published version (APA):

Vaegter, H. B., Christoffersen, L. O., Enggaard, T. P., Holdgaard, D. E. M., Lefevre, T. N., Eltved, R., Reisenhus, C. H., Licht, T. W., Laustsen, M. M., Hansson, S. H., Jensen, P. F., Larsen, T. R. F., Alpiger, S., Mogensen, B. G., & Høybye, M. T. (2021). Socio-Demographics, Pain Characteristics, Quality of Life and Treatment Values Before and After Specialized Interdisciplinary Pain Treatment: Results from the Danish Clinical Pain Registry (PainData). *Journal of Pain Research*, 14, 1215-1230. <https://doi.org/10.2147/JPR.S306504>

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Socio-Demographics, Pain Characteristics, Quality of Life and Treatment Values Before and After Specialized Interdisciplinary Pain Treatment: Results from the Danish Clinical Pain Registry (PainData)

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Background and Aims: PainData is an electronic internet-based clinical pain registry established to improve the understanding and treatment of high-impact chronic pain. The primary aim of this paper is to describe socio-demographics, pain characteristics, quality of life, and treatment values at baseline and follow-up in individuals referred to public and private interdisciplinary pain centers in Denmark between 2018 and 2020.

Methods: Self-reported patient-reported outcomes collected through PainData before (n=12,257) and after (n=4,111) treatment across 13 public and private interdisciplinary specialized pain centers in Denmark (87% of all pain centers in Denmark) are described.

Results: Mean duration of pain was 10 years, and one in three patients reported chronic widespread pain. More than 40% reported opioid use, and 50% had tried four or more different treatment modalities prior to referral. More than 60% reported poor sleep, severe fatigue, and memory and/or concentration deficits. Mean scores on pain catastrophizing, fear of movement, and pain-related disability were high, whereas scores on pain acceptance and self-efficacy were low. Physical and mental health were rated as poor and fair, respectively. One in four patients reported being very much improved or much improved after treatment. Items commonly reported after treatment were increased knowledge about pain, emotions and mood (66.5%), being better at accepting life with chronic pain (63.1%), and improved activity pacing (60.6%).

Conclusion: The PainData registry, containing data from a large cohort of individuals, can help to improve the understanding and treatment of high-impact chronic pain, and collaborations with other researchers are welcome.

Keywords: chronic pain, registry, PainData, questionnaires

Introduction

Chronic pain, as recognized by the International Association for the Study of Pain (IASP) as pain that persists past normal healing time,¹ is one of the leading causes of disability in the western world,² and it is associated with high rates of opioid consumption,³ healthcare utilization,⁴ depression and anxiety,⁵ as well as exclusion from the workforce.⁶ Similarly to many other countries, chronic pain in Denmark is a costly health problem, with an estimated annual expenditure for low back pain alone of over \$1 billion dollars in 2015.⁷

The majority of individuals with chronic pain are treated in the primary care sector, for example by their general practitioner, a physiotherapist, or a chiropractor,

with a smaller part attending hospital departments (eg, rheumatology or neurology) in secondary care. For individuals who do not benefit from primary or secondary care treatments, clinical guidelines recommend referral to interdisciplinary pain management programs in tertiary care settings.^{8,9} Based on recent data from the USA, approximately 8% of the population report high-impact chronic pain, defined as persistent pain with substantial restriction of life activities lasting 6 months or more,¹⁰ indicating that more than 350,000 Danes above 18 years of age may need treatment in a specialist interdisciplinary pain center.¹¹

Across a range of health conditions, including chronic pain,^{12,13} large-scale clinical registries are established to provide a data resource for quality assurance and research to inform clinical decision making and health policy.^{14,15} PainData is a clinical pain registry that aims to improve the understanding and treatment of individuals experiencing high-impact chronic pain. This involves the collection of patient-reported outcome (PRO) data from individuals referred to one of the public or private pain centers that offer specialized interdisciplinary pain treatment in Denmark. The collection of PRO data in the registry has three main aims: 1) to serve as a clinical tool to facilitate the dialogue between the individual and the interdisciplinary team during consultations, 2) to serve as a tool for national benchmarking and quality assurance, and 3) to facilitate national and international research in the field of high-impact chronic pain. PainData was developed and first implemented at the Pain Center at Odense University Hospital (Smertecenter Syd) in 2015. At that time, there was no equivalent national pain registry, and that is still the case. In 2018, the Danish Regions, which is the association and interest organization for the five regions in Denmark, launched a series of projects on value-based healthcare (Værdibaseret Sundhed [VBS]),¹⁶ including a project focusing on VBS within interdisciplinary pain treatment. A series of workshops were hosted to progress the outcome evaluation and benchmarking agenda according to what gives the greatest value and effect for the individual patient. In each workshop, there was representation from many different stakeholders, including the Danish Pain Society, the Danish Rheumatology Association, the Association of Chronic Pain Patients, the Fibromyalgia Association, the Association of General Practitioners, the Danish Psychology Association, leaders from public and private pain centers, as well as representatives from different interdisciplinary professions (social workers,

physiotherapists, pain physicians, nurses, and psychologists). The major outcomes of the workshops were 1) to explore patient values and goals in relation to interdisciplinary pain treatment, 2) to develop and test a battery of questions to assess the patient's values and goals, and 3) to agree on a minimum core data set, which would give insights into the patient's situation while reducing the burden of the patient, and allow for meaningful comparisons between centers. By June 1st 2019, questions related to patients' values and goals were developed and subsequently implemented in PainData in 10 out of 11 public interdisciplinary pain centers in Denmark, as well as in four out of five private interdisciplinary pain centers in Denmark, along with the Patient-Reported Outcomes Measurement Information System (PROMIS) Scale v1.2 – Global Health. The aims of this paper are to introduce the PainData registry and to summarize the self-reported characteristics of individuals with high-impact chronic pain who had an initial consultation, as well as the characteristics of individuals who completed treatment in a public or private interdisciplinary pain center in Denmark from 2018 to 2020.

Methods

Design

The included data are taken from an observational study of individuals who had their initial consultation or individuals who completed treatment in a public or private interdisciplinary pain center in Denmark between January 1st 2018 and December 31st 2020. Reporting of the study follows the STROBE guidelines (STrengthening the Reporting of OBservational studies in Epidemiology).¹⁷

Study Population

Participants are Danish individuals with chronic pain registered in the PainData registry at one of the following interdisciplinary pain centers: 1) Pain Center, University Hospital Odense; 2) Interdisciplinary Pain Center University Hospital Aalborg; 3) Pain Clinic, Regional Hospital Silkeborg; 4) Interdisciplinary Pain Center University Hospital Køge; 5) Interdisciplinary Pain Center Næstved Hospital; 6) Interdisciplinary Pain Center Holbæk Hospital; 7) Interdisciplinary Pain Center Gentofte Hospital; 8) Interdisciplinary Pain Center Rigshospitalet; 9) Pain Clinic, Friklinikken, Grindsted; 10) Interdisciplinary Pain Center, Lillebaelt Hospital; 11)

Interdisciplinary Pain Center Allévia; 12) CFR Capiro, Skørping; or 13) The Private Pain Clinic, Herlev. Pain centers/clinics 1–10 are public hospitals, and clinics 11–13 are private pain clinics with public funding. All interdisciplinary pain centers using PainData are outpatient hospital settings treating patients with chronic (>6 months) non-cancer pain conditions. Patients referred to the pain centers have previously tried a number of treatments in primary and secondary care settings with an unsatisfactory clinical response. Treatments across the different pain centers can be diverse, but commonly consist of one or more of the following elements: medical treatment with a specialist pain consultant (ie, individual adjustment of analgesics to improve effects and reduce side effects), individual consultations with a pain psychologist, physiotherapist, or social worker with cognitive-behavioral therapy training, participation in group sessions with relaxation therapy, acceptance and commitment therapy, or mindfulness-based stress reduction programs, which have shown moderate effects in this population.

Inclusion and Exclusion Criteria

Individuals with chronic pain,¹ who were scheduled for an initial consultation, completed the baseline questionnaire and gave data consent, or individuals who had completed treatment and completed the follow-up questionnaire and gave data consent in one of the involved public or private interdisciplinary pain centers in Denmark between January 1st 2018 and December 31st 2020, were included in the study.

The PainData Registry

The patient part of the PainData registry (see <https://pain-data-test.rsyd.dk>) is designed to capture PRO data at different time points related to the clinical contact in one of the interdisciplinary pain centers. Patients can access the registry 24 hours per day, 7 days per week, via different web browsers. After referral to a pain center but before the initial consultation, all referred patients are invited to answer questions about their clinical pain characteristics and adaptations to pain through a web-based questionnaire system sent via a personal link to the patients' official inbox, e-Boks (the channel that the Danish State and municipalities use to send official documents to citizens). Questionnaires are completed at home before the first consultation. None of the questions are mandatory. In addition to basic demographics, data include questionnaires for measures of pain intensity, disability, physical

and mental health, quality of life, several psychological constructs, and patient values in relation to treatment. During completion of the questionnaires, patients are invited to give consent so that the questionnaire data can be stored in the PainData research database and used for later research. The Danish Data Protection Agency approved the data collection (18/35221). The collected data are protected in accordance with the "Act on the Processing of Personal Data" (Act No. 429 of 31/05/2000) and the "Law on the Status of Patients" (Act No. 482 of 01/07/1998). Participants' personal data are protected in accordance with the Personal Data Processing Act and the Health Act. The duration of data retention is in accordance with notification to the Danish Data Inspectorate. The PainData registry runs on servers managed by the governmental administration of the Region of Southern Denmark. In addition to pain-related questionnaires at baseline and after treatment, PRO data are collected 12, 24, 36, 48, and 60 months after completed treatment, if patients consent to being contacted, to establish a unique prospective cohort over time. As several of the centers/clinics that use PainData have not been collecting data for more than 1–2 years, we only report data from baseline and immediately after treatment in this paper.

In the clinician part of the PainData registry, a summary report from each individual patient's self-reported questionnaire is generated, so that the interdisciplinary team can see the report before seeing the patient during the consultations. All information processing is in real time so that summary data are instantly available to the interdisciplinary team. The registry also has the possibility for clinicians to enter information (eg, pain diagnosis, comorbidities, or planned treatment); however, a consensus between the involved clinics on collection of clinician data has not yet been reached.

The PainData registry was developed by the Pain Center, University Hospital Odense, with the support of the Department of Anaesthesiology and Intensive Care Medicine, University Hospital Odense. The ongoing maintenance costs are divided evenly between the involved pain centers/clinics based on a cooperation agreement. The registry has been built to meet all current Danish health data security standards for data access, storage, back-up, and the tracking of who accesses, inputs, or modifies data. This is achieved via login requirements, passwords, two-factor authentication, firewalls, and logging of access.

Baseline and Follow-Up Questionnaires

Based on the biopsychosocial model of pain,¹⁸ PRO data are collected across several biopsychosocial domains, including pain, disability, work participation, psychological factors, quality of life, contextual factors, and patient values. Wherever possible, the choice of questions and questionnaires was based on recommendations and evidence of their role in chronic pain.^{19–21} The 35 questions in the minimum core data set include sex, age, height, weight, education, and work situation, as well as items related to manifestations of pain: pain chart (any locations of pain during the previous week [Figure 1A], which are used in other clinical registries in Denmark²²), onset date of pain, pain intensity (worst, average, and least in the last 24 hours, using numeric rating scales [NRSs] ranging from 0 [no pain] to 10 [worst imaginable pain],²³ and the average during the last 7 days item from the PROMIS-10 Global Health²⁴), use of analgesics (type, brand, dose, frequency), analgesic effect in the last 24 hours, screening for 1) sleep disturbance using the four items adapted from the Karolinska Sleep Questionnaire (KSQ) sleep quality subscale, assessing frequency of sleep disturbance, frequency of early awakenings, frequency of difficulties falling asleep and frequency of night-time awakenings with difficulty returning to sleep,²⁵ 2) stress, pain catastrophizing, and fear of movement using three-item screening questions,²⁶ and 3) general health, physical and mental health, social activities, and fatigue using the PROMIS-10 Global Health.²⁴ Items related to patient values in relation to treatment were based on workshops in the value-based healthcare project (see Table 3).

When the treatment course has been completed, patients are invited to complete a follow-up questionnaire, which contain approximately two-thirds of the questions in the baseline questionnaire. Further details about the questionnaires in the additional larger data set used by the 10 public pain centers/clinics are provided in [Supplementary Table S1](#).

Statistical Analysis

Descriptive statistics (mean and standard deviation [SD], median and percentiles) were used to describe socio-demographic status, pain characteristics, psychological distress, quality of life, and responses to value-based healthcare items at baseline and follow-up for women and for men. As the aim of this paper was not to compare PRO measures between men and women but to describe the population in the registry, no sex-related hypotheses were established a priori, and thus no formal statistical tests were performed. For the purpose of the description, the duration of pain was also grouped into less than 1 year, 1–3 years, 3–5 years, 5–10 years, and more than 10 years. The variable on education was grouped into the following categories: “Primary school education”, “Upper secondary education”, “Vocational education and training”, “Short cycle higher education”, “Vocational bachelor education”, and “Master’s program”. Baseline differences between responders and non-responders to the follow-up questionnaire were analyzed with independent *t*-tests or χ^2 tests, where appropriate. Differences in PRO measures at baseline and follow-up for patients who started and completed treatment within the period 2018–2020, who completed

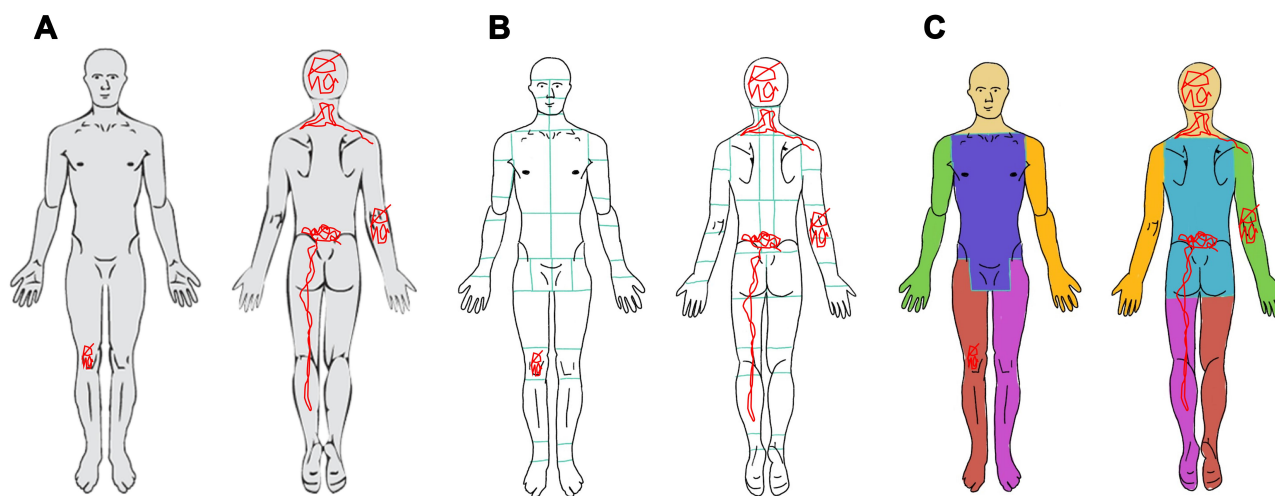


Figure 1 Spatial pain distribution. (A) Body chart from a patient questionnaire; (B) body chart divided into 71 body areas with the pain drawing distributed across 16 body areas; (C) body chart divided into seven body regions with the pain drawing distributed across four body regions.

both questionnaires and gave data consent, are presented with 95% CI, and effect sizes were calculated as Cohen's d , with $d=0.2$ reflecting a small, $d=0.5$ a medium, and $d=0.8$ a large effect.²⁷ All statistical tests were two tailed and the level of significance was set at $p<0.05$.

Results

Baseline Characteristics

In all, 17,351 patients were registered in PainData, with an initial consultation date in a public or private interdisciplinary pain center in Denmark between January 1st 2018 and December 31st 2020. How many of these patients showed up for their initial consultation is not registered in the database. The baseline questionnaire completion rate was 76.6% ($n=13,291$) of all registered patients. The consent rate for the use of personal questionnaire data for quality assurance and research use among completers of the baseline questionnaire was 92.3%. In total, 12,257 patients completed the baseline questionnaire and gave data consent. The amount of missing data on individual questions among these patients was relatively low (Table 1). Characteristics of the population at baseline in the 13 interdisciplinary pain centers/clinics are illustrated in Table 1 for women and for men. Percentiles for each variable are illustrated in Table 2 for women and for men. Responses to value-based healthcare questions are illustrated in Table 3.

Socio-Demographics

The mean age of all individuals was 50.0 ± 14.6 years (range 18–99 years) (Table 1). In total, 67.9% of patients in the registry were women. More than 30% of the individuals had vocational education and training as the highest level of education (higher proportion of men) and less than 5% had an educational level equivalent to a master's program. Almost 30% of individuals were on a pension, and 20% of patients not on a pension were on sick leave.

Pain Characteristics

The mean duration since the pain debut was 10.0 years (range 0.25–84 years). On average, the peak pain intensity level during the last 24 hours corresponded to moderately severe pain (~ 7.8 out of 10) (Table 1). Of all the individuals, back pain, followed by pain in the thorax/abdomen/genital area within the past week were reported most frequently (Table 1). Based on the body drawing illustrated in Figure 1B, the mean number of painful body areas was 21.6, and the median number of painful body

regions was 5 (Figure 1C). Based on the American College of Rheumatology's definition of chronic widespread pain (CWP) as pain localized in both the right and left sides of the body, both above and below the waist plus in the axial skeleton,²⁸ 30.9% of patients had CWP. Almost 90% reported using analgesics, with 43% reporting opioid use. Most patients reported that several different treatment modalities had previously been tried to relieve the pain, with more than 40% reporting having had surgery. The most common non-invasive treatments were exercise, massage, and acupuncture. More than 50% had tried four or more different treatment modalities prior to referral to an interdisciplinary pain center/clinic.

Psychological Distress and Quality of Life

In general, mean scores on pain catastrophizing, fear of movement, and pain-related disability were high, whereas scores on pain acceptance and self-efficacy were low (Table 1). However, mean scores on depression and anxiety were low to moderate. Physical and mental health assessed with the PROMIS Scale v1.2 – Global Health averaged 33.7 and 38.3 (T-scores), respectively, corresponding to a classification of poor and fair physical and mental health, respectively (Table 1). More than 40% reported emotional problems often or always during the last 7 days.

Other Symptoms

More than 80% of all individuals reported disturbed sleep several times per week or more. Sixty percent reported severe or very severe fatigue during the last 7 days, and moderate to severe memory and/or concentration deficits were reported by almost 70% of the individuals.

Value-Based Healthcare Items

Since the implementation of the value-based items in 2019, the items that most individuals reported being of value to them in relation to treatment (scores from “to a moderate degree” to “to a very great degree”) were “being able to participate in social activities” (93.9%), “improved activity pacing” (89.6%), and “help to learn pain coping strategies” (83.8%). The items that fewest individuals reported being of value to them (scores from “to a moderate degree” to “to a very great degree”) were “help with work and income” (58.3%), “improved communication about living with chronic pain” (69.3%), and “gain knowledge about pain, emotions and mood” (75.5%) (Table 3).

Table 1 Characteristics (Mean ± Standard Deviation and Range or Proportions) of Individuals Who Had an Initial Consultation (Baseline) and Individuals Who Completed Treatment (Follow-Up) in a Public or Private Interdisciplinary Pain Center in Denmark from 2018 to 2020.

Variables in the Minimum Core Data Set	Baseline (n=12,257)		Follow-Up (n=4111)	
	Women (n=8324; 67.9%)	Men (n=3933)	Women (n=2765; 67.3%)	Men (n=1346)
Age (years), M±SD (range), n=12,257 (missing=0.0%)	49.1±14.7 (18–99)	51.9±14.3 (18–93)	51.5±14.3 (18–97)	55.0±13.5 (20–89)
Height (cm), M±SD (range), n=11,551 (missing=5.8%)	166.9±6.6 (130–198)	179.7±7.2 (152–210)		
Weight (kg), M±SD (range), n=11,396 (missing=7.0%)	78.5±19.2 (35–185)	91.8±19.1 (42–200)		
BMI (kg/m ²), M±SD (range)	28.2±6.6 (12.9–70.8)	28.4±5.5 (13.6–50.1)		
Highest education, n=11,531 (missing=5.9%)				
Primary school education (%)	18.6	20.4		
Upper secondary education (%)	6.2	4.9		
Vocational education and training (%)	26.1	40.9		
Short cycle education (%)	13.2	7		
Vocational bachelor education (%)	25	14.6		
Master's program (%)	4.2	5		
Others (%)	6.7	7.2		
Work situation, n=11,436 (missing=6.7%)				
Normal working hours (%)	11.4	13.2		
Reduced working hours (%)	10.5	9.4		
Sick leave (part time or full time) (%)	18.7	20.9		
Pension (%)	27.8	31.3		
Studying (%)	3.7	1.9		
Unemployed (%)	4.9	4.4		
Public welfare support (other than sick leave and unemployed) (%)	23	18.9		
Pain duration (years), mean +/- SD (range), n=11,586 (missing=5.5%)	10.3±10.2 (0.25–84)	9.5±10.3 (0.25–68.4)		
<1 year (%)	5.7	7.9		
1–3 years (%)	21.9	24.6		
3–5 years (%)	14.2	14.6		
5–10 years (%)	20.2	19.1		
>10 years (%)	38	33.8		
Peak pain intensity last 24 hours (NRS: 0–10), M±SD (range), higher is worse, n=11,728 (missing=4.3%)	7.8±1.6 (0–10)	7.8±1.6 (0–10)	6.9±2.1 (0–10)	7.0±2.2 (0–10)
Average pain intensity last 24 hours (NRS: 0–10), M±SD (range), higher is worse, n=11,703 (missing=4.5%)	6.5±1.8 (0–10)	6.5±1.9 (0–10)	5.8±2.1 (0–10)	5.8±2.2 (0–10)
Least pain intensity last 24 hours (NRS: 0–10), M±SD (range), higher is worse, n=11,645 (missing=5.0%)	4.8±2.3 (0–10)	4.8±2.4 (0–10)	4.3±2.4 (0–10)	4.4±2.5 (0–10)
Average pain intensity last 7 days (NRS: 0–10), M±SD (range), higher is worse, n=9,291 (missing=6.6%)	5.1±2.9 (1–10)	5.1±2.9 (0–10)	4.9±2.4 (0–10)	4.8±2.5 (0–10)
Pain areas (0–71), M±SD (range), higher is worse, n=11,833 (missing=3.5%)	23.7±16.7 (0–71)	17.2±13.4 (0–71)		
Painful body areas (0–7), M±SD (range)	4.8±2.0 (0–7)	4.0±2.0 (0–7)		
Head and neck pain (%)	68.6	55.2		
Back pain (%)	79.8	77.8		
Thorax/abdominal/genital pain (%)	72.3	59.6		
Right leg pain (%)	67	56.5		
Left leg pain (%)	67.2	59.1		
Right arm pain (%)	64.5	47.4		
Left arm pain (%)	63.1	48		

(Continued)

Table 1 (Continued).

Variables in the Minimum Core Data Set	Baseline (n=12,257)		Follow-Up (n=4111)	
	Women (n=8324; 67.9%)	Men (n=3933)	Women (n=2765; 67.3%)	Men (n=1346)
Analgesic use (%), n=11,763 (missing=4.0%)	88.1	89.6	79.8	80.7
Opioids (%)	39.1	51	24	33.2
Tricyclic antidepressant (%)	19.9	18	22.8	19.7
Anticonvulsants (%)	23.3	27	24.6	27.1
NSAIDs (%)	28.8	25.7	15.2	14.1
Paracetamol (%)	71.5	64.5	48.3	42.4
Muscle relaxants (%)	15.8	14.3	21.9	17.5
Analgesic effect (effect in %), n=11,763 (missing=4.0%)	37.6±23.3 (0–100)	38.1±23.6 (0–100)	44.9±24.3 (0–100)	41.9±24.2 (0–100)
Stress (NRS: 0–10)*, M±SD (range), higher is worse, n=11,600 (missing=5.4%)	4.8±3.2 (0–10)	4.6±3.2 (0–10)	4.2±2.9 (0–10)	4.3±3.1 (0–10)
Pain catastrophizing (NRS: 0–10)*, M±SD (range), higher is worse, n=11,591 (missing=5.4%)	6.9±2.7 (0–10)	7.2±2.7 (0–10)	5.4±3.0 (0–10)	5.9±2.9 (0–10)
Fear of movement (NRS: 0–10)*, M±SD (range), higher is worse, n=11,368 (missing=7.3%)	5.1±3.4 (0–10)	5.8±3.4 (0–10)	3.9±3.4 (0–10)	5.0±3.6 (0–10)
PROMIS-10 Physical Health, M±SD (range), higher is better, n=9291 (missing=6.6%)	33.4±5.6 (16.2–61.9)	34.2±6.2 (16.2–54.10)	35.8±6.4 (16.2–67.7)	35.7±6.5 (16.2–57.7)
PROMIS-10 Mental Health, M±SD (range), higher is better, n=9291 (missing=6.6%)	38.5±7.5 (21.2–67.6)	37.8±7.8 (21.2–67.6)	41.3±7.9 (21.2–67.6)	39.3±8.1 (21.2–67.6)
General health (item 1 in PROMIS-10), n=9,291 (missing=6.6%)				
Excellent (%)	0.4	0.6	0.8	0.8
Very good (%)	3.7	3.7	5.6	3.9
Good (%)	19	19.2	30.1	23.9
Fair (%)	41.8	39.5	44.5	44.2
Poor (%)	35.1	36.9	19	27.3
Emotional problems in last 7 days (item 10 in PROMIS-10), n=9291 (missing=6.6%)				
Never (%)	5.3	6.2	7.4	7.5
Rarely (%)	15.1	15.2	21.6	19.4
Sometimes (%)	37.4	34.9	42.8	37.5
Often (%)	34.7	35	23.2	28.6
Always (%)	7.4	8.7	5	7.1
Sleep, n=11,709 (missing=4.5%)				
Difficulties falling asleep (% often or always)	67.6	65.4	56.9	57.5
Disturbed/poor sleep (% often or always)	80.5	80.2	67.7	70.9
Repeated awakenings (% often or always)	70.2	70.8	59.5	66.2
Early awakenings (% often or always)	62	67.1	55.9	65.8
Fatigue in last 7 days (item 8 in PROMIS-10), n=9291 (missing=6.6%)				
No fatigue (%)	0.8	1.1	1.4	2.3
Mild fatigue (%)	6.5	9.1	13.2	13.3
Moderate fatigue (%)	31.5	35	41.6	39.4
Severe fatigue (%)	43	41.4	33.8	34.8
Very severe fatigue (%)	18.2	13.4	10	10.2

(Continued)

Table 1 (Continued).

Variables in the Minimum Core Data Set	Baseline (n=12,257)		Follow-Up (n=4111)	
	Women (n=8324; 67.9%)	Men (n=3933)	Women (n=2765; 67.3%)	Men (n=1346)
Patient Global Impression of Change, n=3540 (missing=13.9%)				
Very much improved (%)			10.1	8.6
Much improved (%)			17.6	13.8
Minimally improved (%)			20.7	19.7
No change (%)			39.7	42
Minimally worsened (%)			4.5	7.9
Much worsened (%)			4.4	5.4
Very much worsened (%)			3	2.7
Variables in the Additional Data Set	Baseline (n=9303)		Follow-Up (n=3147)	
	Women (n=6432)	Men (n=2871)	Women (n=2156)	Men (n=991)
Fear of movement (TSK 17 items: 17–68), M±SD (range), higher is worse, n=8392 (missing=9.8%)	39.0±7.8 (17–67)	42.1±8.3 (17–68)		
Pain catastrophizing (PCS: 0–52), M±SD (range), higher is worse, n=8768 (missing=5.8%)	25.8±11.1 (0–52)	27.5±11.3 (0–52)	18.3±11.4 (0–52)	20.8±12.0 (0–52)
Depression (PHQ9: 0–27), M±SD (range), higher is worse, n=8628 (missing=7.3%)	10.5±5.5 (0–27)	10.4±5.9 (0–27)	8.8±5.5 (0–27)	9.5±5.9 (0–27)
Anxiety (GAD7: 0–21), M±SD (range), higher is worse, n=8594 (missing=7.2%)	6.0±4.9 (0–21)	6.3±5.3 (0–21)		
Stress (PSS 10 items: 0–40), M±SD (range), higher is worse, n=8710 (missing=6.4%)	19.1±6.9 (0–40)	19.0±7.3 (0–40)		
Disability (PDI 5 items: 0–50), M±SD (range), higher is worse, n=8868 (missing=4.7%)	34.9±9.5 (0–50)	35.4±10.1 (0–50)	30.5±10.8 (0–50)	31.7±11.1 (0–50)
Accept (CPAQ 8 items: 0–48), M±SD (range, higher is better, n=8600 (missing=7.6%)	21.8±8.5 (0–48)	19.5±9.1 (0–48)	23.1±8.1 (0–47)	20.3±9.0 (0–48)
Self-efficacy (PSEQ 2 items: 0–12), M±SD (range), higher is better, n=8509 (missing=8.6%)	5.9±3.2 (0–12)	5.3±3.3 (0–12)	6.4±3.3 (0–12)	5.6±3.3 (0–10)
Self-perceived general health (0–100)#, M±SD (range), higher is better n=8639 (missing=7.2%)	45.4±21.6 (0–100)	42.3±22.4 (0–100)	54.0±22.6 (0–100)	48.4±23.7 (0–100)
Memory and/or concentration deficit, n=8832 (missing=5.1%)				
None (%)	8	12.5		
Mild (%)	20.5	24.2		
Moderate (%)	41.9	39.2		
Severe (%)	29.6	24.1		
Treatments tried for current pain condition prior to referral, n=8327 (missing=10.5%)				
Acupuncture (%)	58	54.2		
Chiropractor/osteopath (%)	56.4	56.7		
Exercise (%)	90.3	88		
Massage (%)	75	68.8		
Mindfulness (%)	45.8	29.2		
Relaxation (%)	54.7	42.5		
Surgery (%)	43	48.3		

(Continued)

Table 1 (Continued).

Variables in the Minimum Core Data Set	Baseline (n=12,257)		Follow-Up (n=4111)	
	Women (n=8324; 67.9%)	Men (n=3933)	Women (n=2765; 67.3%)	Men (n=1346)
Reflexology (%)	31.6	22.6		
Tried 4 or more treatments (%)	58.4	51.5		
Exposed to traumatic event (%), n=8747 (missing=6.0%)	59.4	57.6		
Pain related to trauma (%), only asked to patients reporting exposure to traumatic event, n=5146	38.3	50.6		
Insurance claim (%), n=8742 (missing=6.1%)	16.7	24.9		

Notes: *Assessed using one-item psychosocial screening questions validated in this setting for each construct; #assessed using the EuroQoL health thermometer.

Abbreviations: BMI, body mass index; NRS, numerical rating scale; NSAID, non-steroidal anti-inflammatory drug; PROMIS-10, 10-item Patient-Reported Outcomes Measurement Information System; TSK, Tampa Scale of Kinesiophobia; PCS, Pain Catastrophizing Scale; PHQ9, 9-item Patient Health Questionnaire; GAD7, 7-item Generalized Anxiety Disorder; PSS, Cohen's Perceived Stress Scale; PDI, Pain Disability Index; CPAQ, Chronic Pain Acceptance Questionnaire; PSEQ, Pain Self-Efficacy Questionnaire.

Characteristics at Follow-Up

In total, 10,452 patients were registered in PainData as having completed treatment in a public or private interdisciplinary pain center in Denmark between January 1st 2018 and December 31st 2020. The follow-up questionnaire completion rate was 41.5% (n=4,341) of all completed patients. The consent rate for use of personal questionnaire data for quality assurance and research use among completers of the follow-up questionnaire was 94.4% (n=4,111). Characteristics of the population at follow-up in the interdisciplinary pain centers/clinics are shown in Table 1 for women and for men, and the responses to the value-based healthcare questions at follow-up are shown in Table 3.

Impression of Change

Individuals at follow-up had improved scores compared with baseline scores for all domains, and one out of four individuals reported being very much improved or much improved after treatment. The proportion of individuals using analgesics at follow-up was reduced, and, specifically, the proportion of opioid users was decreased at follow-up compared with baseline. For the patients who started and completed treatment within the period 2018–2020, who completed both questionnaires and gave data consent, the mean treatment duration was 217±170 days (range 0–1043 days). The within-group changes in outcomes were mostly of small to moderate effect sizes, with the largest effect observed for pain catastrophizing (Table 4).

Value-Based Healthcare Items

The achievements that most individuals reported (scores from “to a moderate degree” to “to a very great degree”)

were “gained knowledge about pain, emotions and mood” (66.5%), “had become better at accepting life with chronic pain” (63.1%), and “improved activity pacing” (60.6%).

The achievements that fewest individuals reported (scores from: “to a moderate degree” to “to a very great degree”) were “help with work and income” (24.4%), “helped to sleep better” (38.1%), and “received help to learn pain coping strategies” (49.2%) (Table 3).

Comparison of Follow-Up Questionnaire Completers and Non-Completers

Individuals who completed the follow-up questionnaire were not different regarding sex, pain duration, pain intensity, pain-related disability, opioid use, and fear of movement scores compared with non-responders (Table 5). However, individuals who did not complete the follow-up questionnaire were on average 3 years younger, with higher scores on pain catastrophizing, stress, anxiety and depression, and with lower scores on pain acceptance, self-efficacy, and physical and mental health at baseline.

Discussion

In this paper, we have described the aims and content of the clinical PainData registry as well as the participant recruitment/response rate across public and private interdisciplinary pain centers in Denmark. We have also reported socio-demographics, pain characteristics, quality of life, and treatment values before and after treatment in a large sample of individuals with high-impact chronic pain referred to public and private interdisciplinary pain centers in Denmark. A higher proportion of women

Table 2 Baseline Characteristics Presented with Cumulative Percentiles for Individuals Who Had an Initial Consultation in a Public or Private Interdisciplinary Pain Center in Denmark from 2018 to 2020.

Variables in the Minimum Core Data Set	Women (n=8324)							Men (n=3933)						
	5th	10th	25th	50th	75th	90th	95th	5th	10th	25th	50th	75th	90th	95th
Age	25	30	39	49	58	70	75	28	32	42	52	62	71	76
Weight	156	158	163	167	171	175	178	168	171	175	180	184	189	192
Height	53	57	65	76	89	104	115	65	70	79	90	102	115	125
BMI	19.4	20.7	23.3	27.3	31.9	36.7	40.1	20.9	22.4	24.7	27.7	31.4	35.3	38.0
Pain duration (years)	0.9	1.3	2.7	6.7	14.4	25.0	31.7	0.8	1.1	2.2	5.3	13.0	24.0	31.0
Peak pain intensity last 24 hours (NRS: 0–10)	5	6	7	8	9	10	10	5	6	7	8	9	10	10
Average pain intensity last 24 hours (NRS: 0–10)	3	4	5	7	8	9	10	3	4	5	7	8	9	9
Least pain intensity last 24 hours (NRS: 0–10)	1	2	3	5	6	8	9	1	2	3	5	6	8	9
Average pain intensity last 7 days (NRS: 0–10)	2	2	2	5	8	9	9	2	2	2	5	8	9	9
Pain areas (0–71)	3	5	10	20	34	49	58	2	4	7	14	24	36	45
Painful body regions (0–7)	1	2	3	5	7	7	7	1	1	2	4	6	7	7
Analgesic pain relief (%)	0	10	20	40	50	70	80	0	10	20	40	50	70	80
Stress (NRS: 0–10)	0	0	2	5	8	9	10	0	0	2	5	7	9	10
Pain catastrophizing (NRS: 0–10)	2	3	5	8	9	10	10	2	3	5	8	9	10	10
Fear of movement (NRS: 0–10)	0	0	2	5	8	10	10	0	0	3	6	9	10	10
PROMIS-10 Physical Health	23.5	26.7	29.6	32.4	37.4	39.8	42.3	23.5	26.7	29.6	34.9	37.4	42.3	44.9
PROMIS-10 Mental Health	25.1	28.4	33.8	38.8	43.5	48.3	50.8	25.1	28.4	33.8	36.3	43.5	48.3	50.8
Variables in the Additional Larger Data Set	Women (n=6432)							Men (n=2871)						
	5th	10th	25th	50th	75th	90th	95th	5th	10th	25th	50th	75th	90th	95th
Fear of movement (TSK 17 items: 17–68)	26	29	34	39	44	49	52	28	32	37	42	47	53	56
Pain catastrophizing (PCS: 0–52)	8	11	18	26	34	41	45	9	13	19	28	36	43	46
Depression (PHQ9: 0–27)	2	4	6	10	14	18	21	2	3	6	10	14	19	22
Anxiety (GAD7: 0–21)	0	0	2	5	9	14	16	0	0	2	5	9	14	17
Stress (PSS 10 items: 0–40)	8	10	14	19	24	28	31	7	9	14	19	24	28	31
Disability (PDI 5 items: 0–50)	17	22	29	36	42	46	48	15	21	30	37	43	47	49
Accept (CPAQ 8 items: 0–48)	7	11	16	22	27	33	36	4	8	13	20	25	31	35
Self-efficacy (PSEQ 2 items: 0–12)	0	2	4	6	8	10	11	0	1	3	5	8	10	11
Self-perceived general health (EQ5D-VAS: 0–100)	10	18	30	45	60	75	80	9	14	25	40	60	75	80

Abbreviations: BMI, body mass index; NRS, numerical rating scale; PROMIS-10, 10-item Patient-Reported Outcomes Measurement Information System; TSK, Tampa Scale of Kinesiophobia; PCS, Pain Catastrophizing Scale; PHQ9, 9-item Patient Health Questionnaire; GAD7, 7-item Generalized Anxiety Disorder; PSS, Cohen's Perceived Stress Scale; PDI, Pain Disability Index; CPAQ, Chronic Pain Acceptance Questionnaire; PSEQ, Pain Self-Efficacy Questionnaire; EQ5D, EuroQoL 5 Dimensions; VAS, visual analogue scale.

compared with men attended specialized interdisciplinary pain centers in Denmark. Most patients had had pain for more than 5 years and one in three patients reported chronic widespread pain. The use of opioids was common and most patients had tried a number of treatments in primary and secondary care settings with an unsatisfactory

clinical response. Poor sleep, severe fatigue, and memory and/or concentration deficits were reported by more than 60% of patients. The patient group had high scores on pain catastrophizing, fear of movement, and pain-related disability, whereas scores on pain acceptance and self-efficacy were low. Scores on depression and anxiety

Table 3 Responses to Value-Based Healthcare Questions from Individuals Who Had an Initial Consultation (Baseline) and Individuals Who Completed Treatment (Follow-Up) in a Public or Private Interdisciplinary Pain Center in Denmark from 2018 to 2020.

Baseline (n=9045)				Follow-Up (n= 2819)			
Items and Questions	Response Options	Women (n=6135; 67.8%)	Men (n=2910)	Items and Questions	Response Options	Women (n=1902; 67.5%)	Men (n=917)
Pain acceptance				Pain acceptance			
'Is it important for you to get better at accepting your life with chronic pain?'	Not at all (%)	6.9	9.6	'Have you become better at accepting your life with chronic pain?'	Not at all (%)	13.6	20.5
	To a slight degree (%)	10.6	12.1		To a slight degree (%)	17.2	24.3
	To a moderate degree (%)	21.3	25.2		To a moderate degree (%)	43.1	35.9
	To a great degree (%)	30.1	29.7		To a great degree (%)	22.4	15.5
	To a very great degree (%)	31.1	23.4		To a very great degree (%)	3.7	3.8
Social activities				Social activities			
'Is it important for you to be able to be with family and friends and participate in other social events?'	Not at all (%)	1.6	2.4	'Have you become better at being with family and friends and attending other social events?'	Not at all (%)	22	29.1
	To a slight degree (%)	3.7	5.4		To a slight degree (%)	22.9	27.5
	To a moderate degree (%)	11.3	13.7		To a moderate degree (%)	40.3	31.6
	To a great degree (%)	30.2	35.2		To a great degree (%)	12.4	9.1
	To a very great degree (%)	53.2	43.4		To a very great degree (%)	2.4	2.7
Sleep				Sleep			
'Is it important for you to get help to sleep better?'	Not at all (%)	10.7	11.6	'Have you been helped to sleep better?'	Not at all (%)	39.4	45
	To a slight degree (%)	10.1	9		To a slight degree (%)	19	22.5
	To a moderate degree (%)	16.7	16		To a moderate degree (%)	23.9	19.9
	To a great degree (%)	24.6	25.6		To a great degree (%)	13.8	10.5
	To a very great degree (%)	37.9	37.7		To a very great degree (%)	4	2.1
Analgesics				Analgesics			
'Is it important for you to try to relieve your pain by trying different painkillers?'	Not at all (%)	6.6	6.2	'Have you been helped to try to relieve your pain by trying different painkillers?'	Not at all (%)	25.7	24.5
	To a slight degree (%)	11.1	10.1		To a slight degree (%)	15.4	15.7
	To a moderate degree (%)	23.2	21.3		To a moderate degree (%)	24.4	27.5
	To a great degree (%)	27.1	28		To a great degree (%)	24.2	23.7
	To a very great degree (%)	32	34.3		To a very great degree (%)	10.3	8.5
Pain coping strategies				Pain coping strategies			
'Is it important for you to get help to cope with your pain with e.g. physical and mental exercises and techniques?'	Not at all (%)	6	7.2	'Have you received help to cope with your pain with e.g. physical and mental exercises and techniques?'	Not at all (%)	26.6	32.1
	To a slight degree (%)	9.2	10.9		To a slight degree (%)	19.4	22.2
	To a moderate degree (%)	19.4	21.8		To a moderate degree (%)	31.5	28.2
	To a great degree (%)	30.1	30		To a great degree (%)	19	14.4
	To a very great degree (%)	35.4	30.2		To a very great degree (%)	3.4	3.2
Mood and worries				Mood and worries			
'Is it important for you to gain knowledge about how your pain can affect your emotions and your mood?'	Not at all (%)	11.3	10.2	'Have you gained knowledge about how your pain can affect your emotions and your mood?'	Not at all (%)	16.5	21.9
	To a slight degree (%)	13.7	13		To a slight degree (%)	13.5	16.9
	To a moderate degree (%)	21.6	23		To a moderate degree (%)	27.5	32.2
	To a great degree (%)	25	27.2		To a great degree (%)	31.5	21.6
	To a very great degree (%)	28.4	26.6		To a very great degree (%)	11	7.5

(Continued)

Table 3 (Continued).

Baseline (n=9045)				Follow-Up (n= 2819)			
Items and Questions	Response Options	Women (n=6135; 67.8%)	Men (n=2910)	Items and Questions	Response Options	Women (n=1902; 67.5%)	Men (n=917)
Communication				Communication			
'Is it important for you to be able to tell your loved ones and others what it is like to live with chronic pain?'	Not at all (%)	11.8	15.3	'Have you become better at being able to tell your loved ones and others what it is like to live with chronic pain?'	Not at all (%)	20	24.9
	To a slight degree (%)	17.9	17.6		To a slight degree (%)	24.2	21.4
	To a moderate degree (%)	25.9	25.4		To a moderate degree (%)	32.4	32.1
	To a great degree (%)	22.4	23.9		To a great degree (%)	18.8	17.4
	To a very great degree (%)	22	17.8		To a very great degree (%)	4.5	4.2
Work and income				Work and income			
'Is it important for you to get help with what you can do in relation to your finances and your opportunities on the labor market?'	Not at all (%)	30.2	30.1	'Have you received help with what you can do in relation to your finances and your opportunities on the labor market?'	Not at all (%)	59.6	64.5
	To a slight degree (%)	11.3	12.2		To a slight degree (%)	13.1	12.6
	To a moderate degree (%)	14.2	16.5		To a moderate degree (%)	15.5	14
	To a great degree (%)	17.2	18.9		To a great degree (%)	8.3	6.4
	To a very great degree (%)	27.1	22.3		To a very great degree (%)	3.6	2.5
Activity pacing				Activity pacing			
'Is it important for you to achieve an appropriate balance between activity and rest?'	Not at all (%)	3.3	4.6	'Have you become better at finding an appropriate balance between activity and rest?'	Not at all (%)	13.8	18.4
	To a slight degree (%)	6	8.3		To a slight degree (%)	21.6	27
	To a moderate degree (%)	19.2	26.1		To a moderate degree (%)	38.9	35.3
	To a great degree (%)	33.1	34.3		To a great degree (%)	21.1	16
	To a very great degree (%)	38.4	26.7		To a very great degree (%)	4.7	3.3

were low to moderate, indicating that patients were more distressed than depressed. Physical and mental health were rated as poor and fair, respectively. One in four patients reported being very much improved or much improved after treatment. Items commonly reported after treatment were increased knowledge about pain, emotions and mood (66.5%), better at accepting life with chronic pain (63.1%), and improved activity pacing (60.6%).

The socio-demographics (age, sex, work status) and pain characteristics (duration, patient-reported cause, localization, intensity, use of analgesics, mood and cognitions) of the PainData sample in Denmark are in line with the characteristics of populations with high-impact chronic pain in, for example, Australia and New Zealand¹² and Sweden.¹³ In addition, the improvements reported after interdisciplinary pain rehabilitation in Denmark are of similar magnitude to those reported in the EPOCC registry (Australia and New Zealand)¹² and the SQRP registry (Sweden).¹³ Among the patients referred to specialized interdisciplinary pain

treatment in Denmark, the proportion of women was larger than the proportion of men. Some differences in PROs between women and men were noted. Women reported a higher number of painful body areas, whereas men reported higher scores on fear of movement and pain catastrophizing, and lower scores on acceptance and self-perceived general health. In addition, a higher proportion of men reported opioid use and pain related to trauma.

To date, several research projects^{26,29–33} have used data from the PainData registry, resulting in a number of publications, with more in press or under preparation. These projects have included collaborations between researchers from Denmark, Australia, Germany, and Ireland. The functionality of the registry allows easy implementation of additional questions and questionnaires for future research projects.

Strengths and Limitations

The PainData registry has a number of strengths. One strength is that the population is a consecutive cohort of

Table 4 Patient-Reported Outcome Measures at Baseline and Follow-Up for Patients Who Started and Completed Treatment Within the Period 2018–2020, Who Completed Both Questionnaires and Provided Data consent.

Baseline Characteristics	Baseline	Follow-Up	Difference	Cohen's d Effect Size
			Mean ± SD (95% CI)	
Peak pain intensity (NRS: 0–10), higher is worse, M±SD, n=2661	7.8±1.6	6.9±2.1	0.84±2.0 (0.77 to 0.92)	0.42
Average pain intensity (NRS: 0–10), higher is worse, M±SD, n=2658	6.4±1.8	5.8±2.1	0.68±2.0 (0.60 to 0.75)	0.34
Least pain intensity (NRS: 0–10), higher is worse, M±SD, n=2639	4.7±2.3	4.3±2.4	0.43±2.3 (0.34 to 0.51)	0.19
Analgesic use (%), n=2564	90.5	80.5	–	
Opioid use (%), n=2564	42.7	26.8	–	
Stress (NRS: 0–10)*, higher is worse, M±SD, n=2470	4.5±3.2	4.2±3.0	0.28±2.7 (0.17 to 0.38)	0.10
Pain catastrophizing (NRS: 0–10)*, higher is worse, M±SD, n=2460	6.9±2.8	5.5±3.0	1.33±2.8 (1.22 to 1.44)	0.48
Fear of movement (NRS: 0–10)*, higher is worse, M±SD, n=2366	5.3±3.4	4.3±3.5	1.0±3.4 (0.86 to 1.1)	0.29
PROMIS-10 Physical Health, higher is better, M±SD, n=1780	35.4±5.7	35.2±6.6	0.23±5.9 (–0.04 to 0.51)	0.04
PROMIS-10 Mental Health, higher is better, M±SD, n=1786	38.9±7.7	40.4±8.1	–1.50±6.9 (–1.83 to –1.18)	0.22
Pain catastrophizing (PCS: 0–52), higher is worse, M±SD, n=1660	25.9±11.1	19.4±11.9	6.48±9.3 (6.03 to 6.93)	0.70
Depression (PHQ9: 0–27), higher is worse, M±SD, n=1654	10.0±5.6	9.0±5.7	1.00±4.9 (0.76 to 1.24)	0.20
Disability (PDI 5 items: 0–50), higher is worse, M±SD, n=1736	34.9±9.7	30.8±11.0	4.09±9.3 (3.65 to 4.52)	0.44
Accept (CPAQ 8 items: 0–48), higher is better, M±SD, n=1681	21.4±8.6	22.3±8.5	–0.89±7.54 (–1.26 to –0.53)	0.12
Self-efficacy (PSEQ 2 items: 0–12), higher is better, M±SD, n=1635	6.0±3.2	6.2±3.4	–0.17±2.87 (–0.31 to –0.04)	0.06
Self-perceived general health (EQ5D-VAS: 0–100), higher is better#, M±SD, n=1681	46.8±22.0	50.7±23.2	–3.86±23.9 (–5.0 to –2.71)	0.16

Notes: *Assessed using one-item or two-item psychosocial screening questions and thresholds validated in this setting for each construct; #assessed using the EuroQoL health 0–100 VAS thermometer.

Abbreviations: NRS, numerical rating scale; PROMIS-10, 10-item Patient-Reported Outcomes Measurement Information System; PCS, Pain Catastrophizing Scale; PHQ9, 9 items Patient Health Questionnaire; PDI, Pain Disability Index; CPAQ, Chronic Pain Acceptance Questionnaire; PSEQ, Pain Self-Efficacy Questionnaire; EQ5D, EuroQoL 5 Dimensions; VAS, visual analogue scale.

all patients referred to the public and private interdisciplinary pain centers/clinics in Denmark, which improves its generalizability. Another strength is the recording of each patient's central person registration number, which facilitates data linkage with other Danish registries. In the context of the Danish healthcare system, population-based registers include those containing data on primary and secondary healthcare utilization, hospital diagnoses, prescription drugs, social conditions, work participation, and public welfare support.^{14,15}

The registry also has a number of weaknesses. First, less than half the patients complete follow-up questionnaires. This may reflect that the registry is not a discrete research project where only people who agree to full participation are included. It may also reflect that the main focus within the interdisciplinary pain centers/clinics so far has been on completion of the baseline questionnaire, as an integrated part of clinical practice and used during the initial consultation with the patient. Several measures to improve follow-up participation are currently being implemented, including providing automated reminders in the official e-Boks to those who had

not completed the questionnaire, and a reduction in the number of questions at follow-up to reduce the response burden. Despite this, the follow-up completion rate may also remain a weakness of the registry in the future. Routine data collection as part of clinical practice differs from collecting data to answer a specific research question, as it involves a trade-off between using detailed and validated questionnaires and being able to cover a number of potentially important clinical aspects of chronic pain. For example, in the PainData registry minimum core data set, one-item screening questions to assess fear of movement, pain catastrophizing, and stress are used to reduce the length of the survey and responder burden. Although these screening questions have known concurrent validity in this clinical setting,²⁶ relative to full reference validated standard questionnaires, their test–retest reliability, relative responsiveness, and prognostic value are still unknown. The inclusion of the full validated questionnaires in some of the pain centers will allow further investigation of this in the future. Other limitations may be the use of online questionnaires, as some patients may have limited access to a computer, and

Table 5 Baseline Comparison of Completers and Non-Completers of the Follow-Up Questionnaire.

Baseline Characteristics	Non-Completers	Completers	p-Value
Sex (women) (%)	68.3	66.8	0.10
Age (years), mean±SD	49.2±14.6	52.5±14.4	<0.001
BMI (kg/m ²)	28.2±6.2	28.4±6.3	0.05
Pain duration (years), mean±SD	9.9±10.2	10.3±10.4	0.10
Peak pain intensity (NRS: 0–10), mean±SD	7.8±1.6	7.8±1.6	0.09
Average pain intensity (NRS: 0–10), mean±SD	6.5±1.8	6.5±1.8	0.14
Least pain intensity (NRS: 0–10), mean±SD	4.8±2.3	4.8±2.3	0.36
Analgesic use (%)	87.9	90.4	<0.001
Opioid use (%)	42.8	43.1	0.77
Stress (NRS: 0–10)*	4.8±3.2	4.5±3.2	<0.001
Pain catastrophizing (NRS: 0–10)*	7.0±2.7	6.9±2.8	0.04
Fear of movement (NRS: 0–10)*	5.4±3.5	5.3±3.4	0.30
PROMIS-10 Physical Health	33.1±5.9	35.4±5.8	<0.001
PROMIS-10 Mental Health	38.1±7.6	39.0±7.8	<0.001
Fear of movement (TSK 17 items: 17–68)	40.0±8.1	39.8±8.0	0.52
Pain catastrophizing (PCS: 0–52)	26.5±11.2	25.9±11.1	0.04
Depression (PHQ9: 0–27)	10.7±5.6	9.9±5.6	<0.001
Anxiety (GAD7: 0–21)	6.3±5.1	5.5±4.9	<0.001
Stress (PSS 10 items: 0–40)	19.2±7.0	18.6±7.0	<0.001
Disability (PDI 5 items: 0–50)	35.2±9.7	34.8±9.8	0.14
Accept (CPAQ 8 items: 0–48), higher is better	21.1±8.8	21.2±8.7	0.50
Self-efficacy (PSEQ 2 items: 0–12), higher is better	5.7±3.2	5.9±3.2	0.01
Self-perceived general health (EQ5D-VAS: 0–100), higher is better#	43.8±21.7	46.4±22.3	<0.001

Notes: *Assessed using one-item or two-item psychosocial screening questions and thresholds validated in this setting for each construct; #assessed using the EuroQoL health 0–100 VAS thermometer.

Abbreviations: BMI, body mass index; NRS, numerical rating scale; PROMIS-10, 10-item Patient-Reported Outcomes Measurement Information System; TSK, Tampa Scale of Kinesiophobia; PCS, Pain Catastrophizing Scale; PHQ9, 9-item Patient Health Questionnaire; GAD7, 7-item Generalized Anxiety Disorder; PSS, Cohen's Perceived Stress Scale; PDI, Pain Disability Index; CPAQ, Chronic Pain Acceptance Questionnaire; PSEQ, Pain Self-Efficacy Questionnaire; EQ5D, EuroQo5 Dimensions; VAS, visual analogue scale.

some elderly patients may have difficulties in handling a computer. However, according to the Danish Agency of Digitization, almost 93% of the Danish population above 15 years of age uses digital mail, although the proportion of citizens not using digital mail is larger among individuals above 65 years of age,³⁴ and the proportion of citizens not using digital mail among this group of severely affected patients may be even larger.

Conclusion

The PainData registry contains PRO data from a large and comprehensive observational cohort of individuals attending public and private interdisciplinary pain centers in Denmark for the assessment and treatment of high-impact chronic pain. The registry contains detailed baseline and outcomes data on a broad range of biopsychosocial factors, and new data (eg, new questionnaires for clinical or research purposes) can be collected in the future. These data are in line with the characteristics of populations with high-impact chronic pain in other

countries and can be linked to the Danish population-based registries. Collaborations with other researchers are welcome.

Disclosure

Mr Torsten Wentzer Licht reports personal fees from Grünenthal, outside the submitted work. The authors report no conflicts of interest in this work.

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