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Patient and Disease Characteristics Associate With Sensory Testing Results in Chronic Pancreatitis

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Background: Abdominal pain is the most common symptom in chronic pancreatitis (CP) and has an extensive impact on patients’ lives. Quantitative sensory testing (QST) provides information on sensitivity to pain and mechanisms that can help quantify pain and guide treatment. The aims of this study were (1) to explore sensitivity to pain in patients with CP using QST and (2) to associate patient and disease characteristics with QST results.

Methods: Ninety-one patients with painful CP and 28 healthy control participants completed a QST paradigm using static tests (muscle pressure stimulation and electrical skin stimulations) to unravel segmental and widespread hyperalgesia as a consequence of visceral pain. A dynamic conditioned pain modulation (CPM) paradigm was used as a proxy of pain modulation from the brainstem to inhibit incoming nociceptive barrage, and questionnaires were used to gather information on pain experience and quality of life.

Results: Patients had impaired CPM compared with controls (18.0 ± 29.3% vs. 30.9 ± 29.3%, P = 0.04) and were hypersensitive to pressure stimulation, specifically in the pancreatic (Th10) dermatome (P < 0.001). The capacity of CPM was associated with clinical pain intensity (P = 0.01) and (in the univariate analysis only) the use of opioids was associated with hyperalgesia to pressure stimulation (P = 0.05).

Conclusions: Sensitivity to pain in CP patients can be characterized by a simple bedside QST. Severe clinical pain in CP was associated with reduced CPM function and should be targeted in management.

Key Words: chronic pancreatitis, pain, pain measurement, hyperalgesia

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Abdominal pain is the most common symptom in chronic pancreatitis (CP) and is present during the clinical course of the disease in up to 90% of patients.1,2 The pain is often intense and long-lasting, and it is frequently associated with malnutrition, opioid addiction, physical and emotional disability, and major socioeconomic problems.3 Pain is a symptom that has an extensive influence on patients’ well-being and functional ability. Treatment of pancreatic pain can, however, be difficult with unpredictable outcome.4

Studies on pancreatic pain are difficult to interpret, as visceral pain is a highly individual perception, which is difficult to quantify.5 In the clinic, pain is affected by several confounders such as comorbidities, anxiety, and side effects to medications, which complicates the assessment of pain. To enable health care providers to better characterize sensitivity to pain, experimental models based on quantitative sensory testing (QST) can be used. These techniques are based on the rationale that different neural pathways and networks can be explored using standardized stimulation with simultaneous recording of the evoked pain response by psychophysical and/or objective methods.6–9

Because of convergence between visceral and somatic afferent nerves at the spinal level, somatic QST can be used indirectly to obtain information on pain sensitivity including segmental (spinal) and central sensitization in the context of visceral pain. In addition, QST may also provide knowledge on the dynamic function of the pain system using, for example, the CPM paradigm. This is an experimental paradigm designed to activate endogenous pain inhibitory systems, whereby centers in the brainstem gate incoming stimuli at the spinal cord. Both static and dynamic QST paradigms have previously been used in patients with CP and provided evidence for a malfunctioning pain system with signs of central sensitization and deficient inhibitory pain modulation.10–12 However, the association between QST assessment parameters and patients’ and disease characteristics remains unexplored.

In recent studies, QST has been used to characterize pain sensitivity in different diseases including neuropathy, chronic pelvic pain, irritable bowel syndrome, and functional dyspepsia.10–13 In these studies, “transetiological” patterns of sensory symptoms and deficits have been observed, thus emphasizing that abnormal pain sensitivity is universally observed across chronic pain conditions.14 Moreover, past studies have shown correlations between

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QST parameters and clinical pain characteristics, and QST has been used in predictive contexts, wherein pretreatment QST profiles were able to predict the outcome of surgery, and treatment with analgesics.

We hypothesized that somatic QST can be used to characterize pain sensitivity in CP and that putative changes would associate with clinical pain characteristics. The aims of the study were as follows: (1) to show that a simple somatic QST paradigm reveals significant and plausible differences in pain sensitivity between patients with painful CP and controls and (2) to investigate associations between clinical characteristics of pain and pain sensitivity in CP patients.

**MATERIALS AND METHODS**

**Study Oversight**

This was a cross-sectional study including patients with painful CP and healthy, pain-free volunteers as controls. All participants provided written informed consent before the examinations, which were conducted at the Centre for Pancreatic Diseases at Aalborg University Hospital, Denmark, and the Department of Surgery at Radboud University Nijmegen Medical Centre, Netherlands. The protocol was approved by the local ethical committees and medical agencies.

**Patients**

**CP Patients**

A total of 91 patients with CP from chronic abdominal pain were included in the study. Inclusion criteria included a minimum age of 18 years, CP diagnosis based on the Lünenburg criteria with a score of ≥4, and chronic abdominal pain typical for CP (ie, dull epigastric pain at least 3 days a week). Exclusion criteria were generalized chronic painful conditions other than CP and cognitive impairment hindering their ability to follow instructions. Patients were instructed to continue their usual analgesic treatment on the day of the examination.

A flowchart of the patient inclusion process is provided in Figure 1.

**Healthy Controls**

Twenty-eight healthy, pain-free adults were included as control group. Inclusion criteria included a minimum age of 18 years. Exclusion criteria included chronic or acutely painful conditions, regular use of any kind of analgesics, and pregnancy.

**Procedures**

**Screening**

Initially, all patients were screened according to the inclusion and exclusion criteria; this included a detailed patient history in the CP group to determine pain localization and characterization, comorbidities, alcohol and tobacco use, and medications. Opioid doses were converted to morphine equivalents in milligrams. All patients completed a pain questionnaire (Brief Pain Inventory—short form) and a quality of life questionnaire (EORTC QLQ-C30).

The patients were instructed to keep a pain diary record on an 11-point Numeric Rating Scale every day for 1 week before examination. The patients reported the highest pain score and the average pain score experienced during the last 24 hours.

**QST**

QST was performed on all study participants using a standardized test sequence. Three investigators were trained in QST and performed the examinations.

**Static QST Assessments.** The static QST paradigm included pressure stimulation and electrical stimulation. Pressure stimulation thresholds were tested 1 time using an algometer with a 1.0 cm² probe (Somedic AB, Stockholm, Sweden). The pressure thresholds were examined on the participant’s right side at 5 different sites: below the midline of the clavicle (C5 dermatome), pancreatic abdominal area, above the umbilicus (abdominal Th10 dermatome), pancreatic site (just lateral of the spine in the dorsal Th10 dermatome), hip region on the anterior superior iliac spine (L1 dermatome), and on the quadriceps muscle 5 cm proximal to the patella (L4 dermatome). Two thresholds were measured: pressure pain detection threshold and pressure pain tolerance threshold (pPTT).

Thresholds to electric constant current skin stimulation (Digistim; Biometer A/S, Copenhagen, Denmark) with tetanic stimulation at 100 Hz were measured in the same dermatomes, using 2 electrodes placed 3 cm apart. The equivalents of the 2 thresholds were determined (electrical pain detection threshold and electrical pain tolerance threshold). These methods have previously been described in detail.

**Dynamic QST Assessment.** CPM is a clinically measurable proxy of endogenous pain modulation. It can be experimentally induced using a conditioning stimulus (eg, the cold pressor test) and quantified by applying a test stimulation before and after the conditioning stimulus. In this study, the conditioning stimulus consisted of immersion of the dominant hand in cold water (2.0 ± 0.3°C, continuously stirred) for 2 minutes. If the pain became intolerable before this point, the participants were allowed to remove their hand from the water. The duration of cold pressor stimulation was noted. The test stimulus was pressure stimulation (pPTT) measured on the L4 dermatome on the nondominant side, before the cold pressor test and immediately after its completion. The CPM capacity was quantified as the absolute and relative changes (%) in pPTT before and after the conditioning stimulation.

Comprehensive QST batteries have previously been recommended when examining pain sensitivity in chronic pain patients. These batteries consisted of up to 13 examination modalities and were very time consuming. In this study, we have focused on developing a QST paradigm that was possible to perform at the bedside in a limited time.

**FIGURE 1.** Flowchart of the inclusion process.
consisting only of 4 elements designed to evaluate the most important pain mechanisms. The need for specialized equipment in this paradigm has also been kept at a minimum.

Patient and Disease Characteristics

To examine whether clinical characteristics were associated with QST results, the patient group was subdivided according to pain intensity (mild to moderate: mean Visual Analog Scale score ≤ 5/severe: > 5 in pain diary), presence of diabetes mellitus, opioid consumption (yes/no), and pain pattern (continuous/intermittent). Pain patterns were based on the pain diaries; constant pain was defined as persistent (daily) pain and intermittent pain as short periods of pain separated by pain-free days.

Statistical Analysis

All data are presented as mean ± SD or number (%) unless otherwise indicated. Demographics, clinical data, and CPM parameters of CP patients and controls were compared by the Fisher exact test, Student t test, and 1-way analysis of variance, as appropriate. The 1-way analysis of variance tests were Bonferroni-corrected post hoc. Electrical and pressure stimulation data were log transformed to obtain a secondary normal distribution, and a mixed effects model was used with CP/healthy participants as fixed effects and stimulation site as a random effect, allowing us to obtain an analysis of the overall difference on pressure/electrical stimulation independent of the stimulation site. A mixed effects model was also used for subanalysis of electrical and pressure stimulation data, with clinical subgroups as fixed effects and stimulation site as a random effect. Univariate and multivariate regression analyses were performed to investigate the association of clinical subgroups with QST parameters. Correlations between electrical and pressure stimulation data were analyzed using the Pearson correlation coefficients.

A P-value ≤ 0.05 was considered statistically significant. The software package STATA, version 15.1 (StataCorp LP, College Station, TX) was used for statistical calculations.

RESULTS

A total of 91 CP patients and 28 controls completed the study. Clinical and demographic characteristics of the 2 groups are presented in Table 1. The mean age of patients was 52.6 ± 11.5 years compared with 44.1 ± 9.0 years in controls (P = 0.01). The distribution of sex was proportionate between patients and controls (male patients: 58% vs. 50%; P = 0.44).

Sensitivity to Pain in Patients With CP and in Healthy Controls

Pressure Stimulation

Overall, patients were hypersensitive to pressure stimulation compared with healthy controls (P < 0.001). When comparing the examined dermatomes separately, patients had lower pressure pain threshold at all examined sites, albeit significant differences were only seen for the dorsal pancreatic dermatome (P = 0.01), abdominal pancreatic dermatome (P < 0.001), and L4-control dermatome (P = 0.045) (Fig. 2).

TABLE 1. Clinical and Demographic Characteristics

<table>
<thead>
<tr>
<th></th>
<th>CP</th>
<th>Healthy</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>91</td>
<td>28</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>53 (58)</td>
<td>14 (50)</td>
</tr>
<tr>
<td>Age, mean (±SD)</td>
<td>52.6 (±12)</td>
<td>44.1 (±9)</td>
</tr>
<tr>
<td>BMI, mean (±SD)</td>
<td>22.8 (±5)</td>
<td>24.4 (±4)</td>
</tr>
<tr>
<td>Duration of CP, mean (±SD)</td>
<td>14 (±7)</td>
<td></td>
</tr>
<tr>
<td>Etiology (TIGAR-O), n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toxic-metabolic</td>
<td>43 (47)</td>
<td></td>
</tr>
<tr>
<td>Idiopathic</td>
<td>26 (29)</td>
<td></td>
</tr>
<tr>
<td>Genetic</td>
<td>12 (13)</td>
<td></td>
</tr>
<tr>
<td>Autoimmune</td>
<td>2 (2)</td>
<td></td>
</tr>
<tr>
<td>Recurrent and severe acute pancreatitis</td>
<td>3 (3)</td>
<td></td>
</tr>
<tr>
<td>Obstructive</td>
<td>5 (6)</td>
<td></td>
</tr>
<tr>
<td>Alcohol consumption</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ongoing abuse, n (%)</td>
<td>11 (12)</td>
<td></td>
</tr>
<tr>
<td>Amount, mean units per week (±SD)</td>
<td>6 (±13)</td>
<td></td>
</tr>
<tr>
<td>Smoking</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current smoker, n (%)</td>
<td>65 (71)</td>
<td></td>
</tr>
<tr>
<td>Weekly amount, mean (±SD)</td>
<td>14 (±12)</td>
<td></td>
</tr>
<tr>
<td>Analgetics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients treated with opioids, n (%)</td>
<td>69 (76)</td>
<td></td>
</tr>
<tr>
<td>Daily amount, mean (±SD) (mEq)</td>
<td>94 (±124)</td>
<td></td>
</tr>
<tr>
<td>Range</td>
<td>0-750</td>
<td></td>
</tr>
<tr>
<td>Diabetes mellitus, n (%)</td>
<td>24 (26)</td>
<td></td>
</tr>
<tr>
<td>Patients treated with enzymes for pancreatic exocrine insufficiency, n (%)</td>
<td>48 (53)</td>
<td></td>
</tr>
</tbody>
</table>

*Alcohol-abusing patients were defined as female patients drinking > 7 U of alcohol per week or male patients drinking > 14 U of alcohol per week. BMI indicates body mass index; CP, chronic pancreatitis.

Electrical Stimulation

There were no significant overall differences in electrical pain thresholds between controls and patients (P = 0.13), but an interaction between group and stimulation site was seen (P = 0.002). Hence, differences in electrical pain thresholds were confined to specific dermatomes, and, when comparing the examined dermatomes separately, patients had lower electrical pain threshold in the dorsal pancreatic dermatome (P = 0.01) and L1-control dermatome (P = 0.045) (Fig. 3).
Sensitivity to Pain in Patients With CP by Patient and Disease Characteristics

The clinical and demographic characteristics of the different subgroups are presented in Table 2.

Sensitivity to Pain by Pain Pattern

Patients with severe clinical pain had impaired mean CPM capacity compared with patients with mild to moderate pain and with controls (9.7 ± 23.2% vs. 22.7 ± 28.1% vs. 30.9 ± 29.3%, P = 0.01) (Fig. 4). There was no difference in the pain experienced during the cold pressor test in the subgroups (P = 1.00). No other differences in QST parameters were observed between mild to moderate clinical pain and severe clinical pain; pressure stimulation (P = 0.96) and electrical stimulation (P = 0.97).

Sensitivity to Pain by Opioid Treatment

Opioid-treated patients were hypersensitive to pressure stimulation compared with their opioid-naive counterparts (P < 0.05). When comparing the examined dermatomes, opioid-treated patients had lower pressure pain threshold at the abdominal pancreatic dermatome (208 ± 102 vs. 181 ± 145; P = 0.01), while borderline significant differences were seen for the dorsal pancreatic dermatome (432 ± 225 vs. 354 ± 207; P = 0.07) and L4-control dermatome (467 ± 162 vs. 400 ± 248;
P = 0.07) (Fig. 5). No other differences in QST parameters were observed between opioid-treated and opioid-naive patients; electrical stimulation (P = 0.20) and CPM capacity (17.5 ± 25.4 vs. 18.2 ± 30.6; P = 0.15).

There were no differences in QST parameters between diabetic and nondiabetic patients; pressure stimulation (P = 0.97), electrical stimulation (P = 0.14), and CPM (19.5 ± 25.1 vs. 18.2 ± 30.4; P = 0.15).

Table 3 summarizes the result of the multivariate analysis for CPM. Clinical pain intensity (coefficient = −3.7 ± 1.8%; P = 0.047) was independently associated with less CPM capacity after adjusting for age, sex, diabetes, and opioid consumption (Fig. 6). In contrast, the significant difference in pressure stimulation thresholds when comparing opioid-naive and opioid-treated patients, was lost in the multivariate analysis (P = 0.30).

**Sensitivity to Pain by Diabetes**

There were no differences in QST parameters between diabetic and nondiabetic patients; pressure stimulation (P = 0.97), electrical stimulation (P = 0.14), and CPM (19.5 ± 25.1 vs. 18.2 ± 30.4; P = 0.15).

**Multivariate Analysis**

Table 3 summarizes the result of the multivariate analysis for CPM. Clinical pain intensity (coefficient = −3.7 ± 1.8%; P = 0.047) was independently associated with less CPM capacity after adjusting for age, sex, diabetes, and opioid consumption (Fig. 6). In contrast, the significant difference in pressure stimulation thresholds when comparing opioid-naive and opioid-treated patients, was lost in the multivariate analysis (P = 0.30).

**Correlations Between QST Parameters and Questionnaires**

Average pressure pain thresholds (mean of all stimulation sites) were significantly correlated to the average electrical pain thresholds (r = 0.38, P < 0.001) and the pain interference score on the Brief Pain Inventory (r = −0.24, P = 0.03). No other correlations were seen between the different QST parameters and questionnaire scores, as can be seen in Table 4.
DISCUSSION

We investigated central sensitivity to pain in patients with 
CP and in pain-free controls using a static and dynamic 
QST paradigm and examined its association with patient and 
disease characteristics. Patients were characterized by pressure 
hyperalgesia, with the most pronounced changes observed for 
the pancreatic dermatome and in patients on opioid-based 
pain medication. Furthermore, a decreased CPM capacity 
was seen in CP patients, which was independently associated 
with clinical pain intensity. Sensitivity to pain was unrelated 
to diabetes. These findings attest to the growing body of evidence 
suggestive of abnormal sensitivity to pain in CP patients and 
also support the use of somatic QST for pain character-
ization in this context.

Sensitivity to Pain in CP and Healthy Controls

We found that CPM capacity was impaired in the CP 
group compared with controls. In addition, CP patients were 
hypersensitive to pressure and electrical stimulations in the 
examined locations. The reduced CPM capacity in CP 
patients corresponds to results from previous studies.9,32,33 
Systemic hyperalgesia in CP patients could indicate a change 
in sensitivity to pain. It has previously been suggested that 
patients with chronic pain have systemic sensitization second-
ary to central reorganization and structural brain 
changes.34–37 The patients in our study had been diagnosed 
with CP at a mean of 14 years before testing. Their long 
duration of disease may have resulted in a longer period of 
modeling, and it may be that patients with shorter duration 
of disease have shown lesser evidence of these changes.33 

We have compared the results of this study with knowledge 
gained in the clinic from assessing the results of more extensive 
QST paradigms. We have found that these results are quite 
similar to what has previously been shown in QST examinations; 
however, no statistical validation has been performed.

Sensitivity to Pain and Clinical Parameters

Pain intensity was significantly associated with CPM 
capacity in CP patients, as patients with severe pain had 
more reduced CPM capacity compared with patients with 
mild pain. Similar results have been seen in other diseases 
such as irritable bowel syndrome, osteoarthritis, spinal cord 
innjury, and functional dyspepsia.13,15,38,39 Prolonged duration 
of the painful condition has resulted in lower CPM 
capacity in both patients with CP and patients with 
osteoarthritis.9,15 Taken together with our findings of associ-
ation between CPM capacity and pain intensity, the data 
indicate that impaired pain modulation is an important 
mechanism in chronic pancreatic pain. However, the ques-
tion of whether reduced CPM capacity is caused by 
persistent severe pain, or is actually the cause for developing 
severe pain, remains unsolved. The predictive value of CPM 
function in the context of treatment has previously been 
studied. Bouwense et al40 found that CPM function pre-
dicted the efficacy of treatment with pregabalin in CP.

Diabetes can cause hyposensitivity due to neuropathy,41,42 but 
we found no association between the presence of diabetes 
and pressure stimulation, electrical stimulation, or CPM 
capacity, and all patients with symptoms of other painful 
conditions were excluded. These data suggest that, if any of the 
patients had an undiagnosed neuropathy, it did not seem to 
have an effect on the results, as the QST results of the diabetic 
group and the nondiabetic group were quite similar.

Of note, patients on opioid pain management regimens 
were found to be hypersensitive to pressure stimulation in 
univariate analysis compared with their opioid-naïve counter-
parts. One explanation for this could be that these patients 
experience more severe pain (the reason for the opioid therapy) 
or, in some cases, opioid-induced hyperalgesia. In our study, 
the association was lost in the multivariate analysis, potentially 
due to type II error in the setting of a small sample size.

When looking at pain patterns over time, a previous 
study has found that patients with intermittent pain were 
more likely to respond to treatment than those with a more 
stable pain profile.43 It has been suggested that patients with 
continuous pain have a more stabilized and irreversible central 
sensitization, which is a sign of end-stage pain chronification 
with severe central neuroplastic changes that are less prone to 
respond to treatment.44 These findings were not reproducible 
in this study, as pain pattern was not significantly associated 
with QST parameters. This could be explained by the rela-
tively short baseline pain registration, which might not be 
sufficient to illustrate the pain pattern thoroughly.

In the current study, there was no screening for comorbidities such as anxiety and depression, which may 
influence the results. However, the lack of correlation 
between life quality and QST results suggest that such 
symptoms are likely not of major importance in this study.

Pain sensitivity has been quantified using somatic QST, 
although the pain in CP is visceral. This could, in theory, be 
a source of error, as it is an indirect measurement. However, 
variations with QST similar to those found in the clinical 
subgroups of the CP patients have previously been seen in 
various somatic pain conditions, implying that this variation is 
not caused by the indirect measurements relating to the pan-
creas, but rather to central changes in the pain system.10,15,45,46

Strengths and Limitations

QST has been widely used in pain research over the 
past 40 years, and, especially, the static tests have shown a

TABLE 4. Intercorrelations Between QST Parameters and Correlations With BPI and QOL Scores

<table>
<thead>
<tr>
<th>CPM</th>
<th>ePDT</th>
<th>pPDT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Correlation Coefficient</strong></td>
<td><strong>Correlation Coefficient</strong></td>
<td><strong>Correlation Coefficient</strong></td>
</tr>
<tr>
<td><strong>P</strong></td>
<td><strong>P</strong></td>
<td><strong>P</strong></td>
</tr>
<tr>
<td>BPI pain severity score</td>
<td>−0.03</td>
<td>0.78</td>
</tr>
<tr>
<td>BPI pain interference score</td>
<td>0.08</td>
<td>0.45</td>
</tr>
<tr>
<td>EORTC QLQ score</td>
<td>0.03</td>
<td>0.77</td>
</tr>
<tr>
<td>ePDT</td>
<td>0.07</td>
<td>0.46</td>
</tr>
<tr>
<td>pPDT</td>
<td>−0.01</td>
<td>0.93</td>
</tr>
</tbody>
</table>

Bold values indicates significant values.

BPI indicates Brief Pain Inventory; CPM, Conditioned Pain Modulation; EORTC QLQ, the European Organization for Research and Treatment Quality of Life Questionnaire; ePDT, electrical pain detection threshold; pPDT, pressure pain detection threshold; QOL, quality of life; QST, Quantitative Sensory Testing.
high degree of standardization and reproducibility. Dynamic QST tests in chronic pain patients have suffered from large variability between patients and poor day-to-day reproducibility in patients with CP. Although dynamic tests such as CPM are encumbered by this variability, an argument to be made is that substantial variability over time in the CPM paradigm must be expected, given the necessarily reactive and dynamic nature of the modulation of nociceptive inputs. The fluctuating nature of chronic pancreatic pain may also contribute to the variability. We have considered this in our analysis of dynamic QST results and performed our testing in a standardized manner to limit unnecessary variability.

In addition, an age difference was seen between patients and controls, with the patients being of more advanced age. Previous studies have shown that advancing age can affect the QST results, but when examining whether the age differences influenced the results of the tests, no connection was found. We, therefore, conclude that the age difference did not affect the results of this study.

There was also a difference in the duration of the cold pressor test between patients and controls. However, no correlation between the duration of the cold pressor test and the magnitude of CPM response in patients was found, and this could suggest that the patients’ CPM response is already chronically activated to its maximum, thereby not allowing for further increase, and this could even result in facilitation of the pain.

As a comparator, one could reflect on whether it is optimal to use healthy volunteers or whether a control group consisting of patients with similar comorbidities and other characteristics (ie, alcohol consumption, smoking habits, etc.), for example, a group consisting of patients with pain-free CP, would be better suited. We will focus on elucidating this aspect in future studies.

The data concerning the patients compared with healthy volunteers are, however, not the main finding of this study. It is a result in line with previous studies in CP and QST and is, therefore, mainly interesting in the fact that it confirms that the results of this study probably can be transferred to the general population of patients with painful CP.

In this study, we compared CP patients on opioid management regimes with their opioid-naive counterparts, but we did not include a control group on opioid management regimes due to nonabdominal pain. It is, therefore, difficult to elucidate the full effects of opioids on QST results, and we plan to include this in future studies.

In this study, we did not examine psychiatric comorbidities such as anxiety and depression, although this could influence the patient pain report, as psychiatric comorbidities can change the patients’ experience of pain. All results should, therefore, be evaluated with this in mind. As a consequence, we have included a screening of psychiatric comorbidities in the examination at our clinic to ensure that the examination is thorough and exhaustive.

Clinical Perspective: Mechanism-based Treatment

The choice of analgesic used to treat pancreatic pain, is typically based on local traditions and the treating physician’s personal experience and preferences, and, in severe pain, opioids are often needed. For some practitioners and patients, opioids play an integral part in the treatment of persistent severe pain. Although opioids can provide effective analgesia, they are also associated with considerable side effects, and a high risk of addiction. Several studies point to the fact that patients can benefit from a more individualized treatment approach with a focus on pain mechanisms. QST can be used to characterize the patient’s sensitivity to pain and thereby improve the possibility of individualizing treatment according to affected pain mechanisms. For example, it has been shown that pregabalin has moderate inhibitory effects on central sensitization, and this effect can be predicted by static QST. Likewise, a reduced CPM capacity can be potentiated with either tapentadol or duloxetine, and the clinical effect of such treatment is predictable by dynamic QST. CP function has also shown to be predictive of pregabalin’s treatment effect in patients with CP.

Our study provides a simple QST protocol that can be applied in the clinic, with results that are comparable to prior literature comparing CP patients with controls. Its simplicity makes it amenable to bedside examination in the clinical setting. Incorporating this protocol into clinical evaluation of pancreatic pain represents a feasible and important improvement in the evaluation of pancreatic pain, as recommended in international treatment guidelines (Drewes and colleagues). This study does not provide sufficient control material to estimate reliable QST-based cutoff values, but this will be the next step in developing a validated QST-based treatment guideline.

CONCLUSIONS

The clinical characteristics of CP patients associate with objective quantitative testing of the pain system. The main finding was that severe clinical pain was associated with decreased CPM capacity. This discovery, via the use of quantitative sensory testing, may represent a paradigm shift in clinical evaluation and treatment of CP. These findings further emphasize that changes in sensitivity to pain are important. Future studies should address how quantitative sensory testing can be used to predict disease course and treatment response.

REFERENCES


