The responsiveness of quantitative sensory testing-derived sensory phenotype to disease-modifying intervention in patients with entrapment neuropathy: a longitudinal study

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Abstract

The German Research Network on Neuropathic Pain (DFNS) quantitative sensory testing (QST) method for sensory phenotyping is used to stratify patients by mechanism associated sensory phenotype, theorised to be predictive of intervention efficacy. We hypothesised that change in pain and sensory dysfunction would relate to change in sensory phenotype. We investigated the responsiveness of sensory phenotype to surgery in patients with an entrapment neuropathy.

With ethical approval and consent, this observational study recruited patients with neurophysiologically confirmed carpal tunnel syndrome. Symptom and pain severity parameters and DFNS QST were evaluated prior to and after carpal tunnel surgery. Surgical outcome was evaluated by patient-rated change. Symptom severity score of the Boston Carpal Tunnel Questionnaire and associated pain and paraesthesia subgroups were comparators for clinically relevant change.

QST results (n=76) were compared to healthy controls (n=54). At 6 months post-surgery 92% participants reported a good surgical outcome and large decrease in pain and symptom severity (p<.001). Change in QST parameters occurred for thermal detection, thermal pain and mechanical detection thresholds with a moderate to large effect size. Change in mechanical pain measures were not statistically significant. Change occurred in sensory

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phenotype post-surgery (p<.001); sensory phenotype was associated with symptom subgroup (p=.03) and patient-rated surgical outcome (p =.02).

QST derived sensory phenotype is sensitive to clinically important change. In an entrapment neuropathy model, sensory phenotype was associated with patient-reported symptoms and demonstrated statistically significant, clinically relevant change after disease modifying intervention. Sensory phenotype was independent of disease severity and may reflect underlying neuropathophysiology.

Keywords: Quantitative sensory testing (QST); sensory phenotype; neuropathic pain; responsiveness; stratification

Background
Quantitative sensory testing (QST) is a clinical examination method for evaluating nerve fibre function in response to graded multi-modal stimuli [3;28;36;37;42;67]. Historically, variability in QST tests have impeded extrapolating findings across studies [27]. To improve standardization and interpretability, the German Research Network on Neuropathic Pain (DFNS) published a comprehensive QST protocol (DFNS QST)[70] that is now widely employed.

DFNS QST has been implemented to define reference values in healthy participants [57;64;69]. This has enabled the interrogation of sensory dysfunction and description of heterogeneous sensory profiles in a range of neuropathic pain conditions [43;58;65;76;79].
Somatosensory dysfunction measured using DFNS QST can be clustered into three sensory phenotypes, composites of the 13 DFNS QST tests, primarily characterised by thermal or mechanical hyperalgesia or sensory loss [6]. To enhance the utility of DFNS QST as a stratification tool, an algorithm based on these three phenotypes was developed [81]; thereby affording greater sensitivity and precision in elucidating somatosensory dysfunction.

While somatosensory profile is hypothesised to reflect underlying pathophysiology of pain pathways and mechanisms in neuropathic pain conditions [2;4;18;31;59], the pathophysiology of neuropathic pain is variable, complex and not fully elucidated. The science of QST derived somatosensory profiling is, relatively speaking, in its infancy and it’s clinical importance remains under investigation. Ambiguities exist, for example there is an observed discordance for patient reported pain and pain sensitivity and the results of evoked pain measures employed in quantitative sensory testing [32;34]; these equivocal findings warrant further exploration.

Establishing that neuro-pathophysiology varies within a neuropathic pain condition is thought to be germane to improving treatment and outcomes. This has created the impetus for prescribing to target pathophysiological mechanisms [5;31]. It is unsurprising, therefore, that DFNS QST is now widely employed in clinical trials. In pain trials, it has been recommended that QST derived sensory phenotype be incorporated as a method for patient stratification [23;25;30;35;44;66]. In surgical prognosis design studies [68] somatosensory phenotype is incorporated as an exploratory risk factor for the development of neuropathic or persistent post-operative pain [1;13;83;84].
Despite this implementation of DFNS QST and QST derived sensory phenotype in research, there is no evidence supporting their responsiveness. Fundamentally, for a tool to be psychometrically robust, it is essential that it be both internally and externally responsive; detecting small, clinically important change [26;40;47]. Accepted methods for assessing internal responsiveness include assessing patients prior to and following a treatment known to be efficacious.

A carpal tunnel surgery cohort provides an elegant model for evaluating the responsiveness of QST and QST derived sensory phenotype in patients with compression neuropathy following disease modifying treatment. Carpal Tunnel Syndrome (CTS) is a compression neuropathy of the median nerve at the wrist leading to measurable sensory disturbance and pain[19;62]. DFNS QST studies of somatosensory function in CTS patients demonstrate that whereas sensory loss to thermal and mechanical stimuli are hallmarks of CTS[7;71]; hyperalgesia to thermal and mechanical stimuli are also observed[85]. Decompression surgery is efficacious[46;72;80] and essentially, there exists a well validated comparator measure of clinically relevant change; the Symptom Severity Scale of the Boston Carpal Tunnel Questionnaire, a disease specific, reliable and responsive patient-completed questionnaire[24;50].

**Study Aims:**

- Investigate change in DFNS QST and QST derived sensory phenotype before and after surgery in patients with carpal tunnel syndrome to determine internal responsiveness.
• Investigate association between DFNS QST and sensory phenotype and participant-reported measures of pain and sensory dysfunction before and after surgery to determine external responsiveness.

Methods

Ethical approval was granted by the Camberwell St Giles National Research Ethics Committee (14/LO/1436) on 29 August 2014 for a prospective, repeated measures observational study in a convenience sample of CTS participants (https://www.hra.nhs.uk/planning-and-improving-research/application-summaries/research-summaries/impact-of-pain-and-somatosensory-phenotype-on-carpal-tunnel-surgery/). Study findings are reported according to STROBE guidelines for observational studies [82]; this manuscript reports a secondary analysis of an existing dataset. Adult patients listed for open carpal tunnel decompression surgery at two London National Health Service (NHS) hospitals were recruited by poster, in person at their hospital clinic appointment and by post. Participants were not paid for study participation however travel was reimbursed.

Surgical outcome measurement

The primary measure used to classify surgical outcome as good or poor (binary outcome) was a patient-reported global rating of change (PGRC) at 6 months post-surgery. Using a 5 point ordinal scale, 1 = worse; 2 = unchanged; 3 = slightly better; 4 = much better and 5 = completely cured [10;11;51;60] a grade 3 or above was interpreted as treatment success. Where previous studies [10;11] have defined treatment success as 4 or above using the same ordinal scale, these investigators concede that their patients are selected for surgery based on good prognosis and as a consequence their findings are less generalizable to the wider
population of patients with CTS. However, comparable to the current study, Jerosch-Herold et al., (2014) investigated CTS surgical outcome in a pragmatic prospective cohort including subjects with multiple comorbidities and known risk factors for poor outcome and identified a grade of 3 (slightly better) to define treatment success. In the United Kingdom, patients with long-standing, severe median nerve compression are routinely advised they may not have complete resolution of symptoms following surgery [16]. This is in keeping with evidence of patient expectations of carpal tunnel surgery outcome; patients with long-standing and/or severe symptoms report limited expectations from surgery [49].

**Case definition of neuropathic pain**

To document the presence of a median nerve mononeuropathy, nerve conduction studies (NCS) were performed by the respective hospital neurophysiology departments and severity graded according to Bland [9] criteria. If this criterion was fulfilled, then pain was categorised as neuropathic where both of the following conditions were met [41;48;77]:

- a score of $\geq 4$ on the Douleur Neuropathique 4 questions (DN4) questionnaire [14].
  
The DN4 consists of seven symptom questions and three sensory examination measures and is a validated patient and examiner completed measure for the evaluation of neuropathic pain symptoms, signs and descriptors.

- pain present in a median nerve or extra-median nerve distribution. Pain distribution was classified from a patient-completed hand symptoms diagram as originally described by Katz et al. [52], with modification based on the work of Zanette et al. [85;86]. Pain within any aspect of the median nerve distribution of the hand (thumb, index, middle or ring fingers, including the dorsal digits), distal to the carpal tunnel was defined as median nerve distribution pain. Pain localised to the median nerve distribution distal to the carpal tunnel AND the dorsal radial hand (radial nerve
distribution) OR any portion of the small finger or ulnar nerve distribution of the hand was defined as extra-median nerve pain. Pain restricted to the ulnar nerve distribution of the hand, and/or pain occurring only proximal to the carpal tunnel was defined as non-median nerve distribution pain.

This two-stage triage for categorization of neuropathic pain was repeated at 3- and 6-month visits. However, in lieu of repeat electrophysiological testing, two or more abnormal quantitative sensory testing findings indicative of loss of sensory function (i.e. cold detection threshold; warm detection threshold; thermal sensory limen; mechanical detection threshold; vibration detection threshold [z x ±1.96]) was taken as a confirmatory diagnostic test [41].

**Participation Criteria**

Exclusion criteria were significant cognitive dysfunction or lack of English language, a history of potentially confounding conditions (rheumatoid arthritis, renal failure, peripheral neuropathy of any origin other than CTS), steroid injection of the study limb within the previous four weeks or previous carpal tunnel surgical release in the study hand, anatomic abnormalities of the wrist or hand, median nerve injury or compression secondary to traumatic injury and pregnancy.

**Schedule of study visits**

Baseline measures were completed within 6 weeks prior to surgery; 3- and 6-months postsurgery assessments were completed within ± 21 days.
Procedure

At baseline, demographic data and medical history was recorded. All tests and questionnaires (described below) were delivered in the same order across the participants, across visits. Pain and symptom severity parameters and median nerve somatosensory function were evaluated at three time points; prior to and at 3- and 6-months post-surgery. At three- and six-month follow-up visits, participants sealed their completed surgical outcome measures in an envelope coded with their participant identification number. Surgical outcome measures remained sealed and retained with participant case report forms until participants completed the study.

Pain and sensory dysfunction comparator measures

Symptom severity and frequency was assessed with the Symptom Severity Scale (SSS) of the Boston Carpal Tunnel Questionnaire (BCTQ) [54]. Eleven symptoms are rated on a 5-point scale with lower scores implying milder symptoms. It has been demonstrated in CTS that the SSS can be exploited to identify pain-dominant and paraesthesia-dominant subgroups with concomitant, distinct alterations in brain morphometry on structural MRI [56]. Therefore, in addition to total symptom severity score, the SSS was used to stratify participants to pain-dominant, paraesthesia-dominant and mixed-symptom subgroups for comparison with DFNS QST derived sensory phenotypes. BCTQ questions 1-5 were averaged to generate a pain score, questions 6-10 a paraesthesia score and participants were stratified based on the larger of the two. Where pain and paraesthesia scores were equivalent participants were stratified to the mixed symptom subgroup [56].
Pain dimensions were assessed with the Neuropathic Pain Symptom Inventory (NPSI) [15], a validated patient-completed measure. Total NPSI scores range from 0 to 100 with greater scores implying more severe symptom severity; item scores of 1–3 indicate mild pain severity, 4-6 moderate and 7-10 severe [33]. Pain severity was assessed with the validated [75] Brief Pain Inventory (BPI) whereby a Pain Severity Score is calculated as the mean of four questions quantifying present pain and the least, worst, and average pain over the last week. Pain is rated on an 11-point scale ranging from 0 (no pain) to 10 (pain as bad as you can imagine) [20].

**Median nerve somatosensory function**

QST was performed by a trained investigator (DK) according to the German Research Network on Neuropathic Pain (DFNS) protocol [70]. All equipment was new and calibrated prior to testing. All tests were performed in the same order, as described in Table 1. Participants were seated in a quiet, temperature-controlled room, with the test hand (surgical hand) supported on a table. Tests were first demonstrated on the dorsal contralateral forearm. Pressure pain threshold was tested at the thenar eminence, all other tests were performed at the volar distal phalanx of the middle finger. Median nerve function was tested at the middle finger because although it innervates the volar thumb, index, middle and radial half of the ring fingers, there is evidence that the middle finger is more symptomatic and more sensitive to tests of mechanical detection in patients with CTS [12;29;50].

Full details of the DFNS QST testing procedure are reported in **Supplementary Material 1** (https://doi.org/10.6084/m9.figshare.12860003.v2). In summary, a Somedic MSA thermal stimulator (Sweden) with an 18 mm² metal Somedic thermode was used for thermal detection.
and pain threshold tests. Thermal thresholds were tested using a baseline-temperature of 32°C and ramping at 1°C/second with limits of 5°C and 50°C. Mechanical detection threshold used glass monofilaments (Optihair2-Set, Marstock Nervtest, Germany) with nominal bending forces between 0.25 and 512 mN. Mechanical pain threshold, mechanical pain sensitivity and wind up ratio were tested using blunt probes with forces ranging from 8 to 512 mN (pinprick stimulator, MRC, Heidelberg, Germany). Dynamic mechanical allodynia (DML) was tested with a cotton wisp, a cotton bud (Q-Tip) and a standardised brush designed to produce minimum friction (Somedic, Sweden). Vibration detection threshold testing used a Rydel–Seiffer graded tuning fork (64 Hz, 8/8 scale). Pressure pain threshold was tested with a pressure algometer (FDN100, Wagner Instruments, Greenwich, CT, USA) with a surface area of 1 cm² and by applying pressure at a rate of 1 kg/cm² per second.

Insert table 1

Median Nerve QST Normative Data

Published normative data for DFNS QST in the hand pertain to the dorsal hand (radial nerve)[57]. Therefore, in parallel, we undertook a QST study in healthy volunteers to determine whether dorsal hand reference data is generalizable to the median nerve innervated volar hand and if not, to establish median nerve normative reference data. Imperial College Research Ethics Committee approval (IREC_13_1_10) was received on May 13th, 2014.

QST testing was performed by a single trained investigator (DK) and consistent with the DFNS QST procedure [70] employed in patients with CTS, as described in Supplementary Material 1 (https://doi.org/10.6084/m9.figshare.12860003.v2). Tests were performed at the
dorsal hand (radial nerve innervation) and volar distal middle finger (median nerve) in one session. All Healthy Volunteer QST study procedures and results are reported in Supplementary Material 2 (https://doi.org/10.6084/m9.figshare.12860066.v2).

Significant differences were found between the dorsal and volar hand test sites for most parameters, therefore DFNS normative data for the dorsal hand [57] are not generalisable to the median nerve innervated volar hand. To generate median nerve reference data for comparison with clinical cohorts, data that was not normally distributed was transformed based on distribution properties and transformed to create a normal distribution [38;45;73]. Data was reported (not back transformed) with the mean and standard deviation as consistent with DFNS QST data analysis [57].

**Statistical methods**

All continuous data were tested for normality of distribution. Patient characteristics and distribution of symptom severity measures were summarized using descriptive statistics. Change in continuous measures across three time points (baseline; 3 months; 6 months) was investigated with one-way repeated measures analysis of variance (ANOVA) with post hoc Tukey HSD test and pairwise comparisons or the non-parametric Friedman test with post-hoc Wilcoxon Signed Rank Tests with Bonferroni correction, as appropriate. Magnitude of change in repeated measures was investigated with a within-subjects effect size, calculated as $r=Z/\sqrt{N}$ [22;53] and interpreted as 0.1=small, .3=medium and 0.5=large [21;63].

The distribution of QST data was assessed with skewness and kurtosis distribution parameters, statistically with the Kolmogorov-Smirnov statistic and visually with inspection
of histograms and Normal QQ plots. Raw QST data were described with the median and interquartile range. At the group level, CTS and control QST data were compared with the Mann-Whitney U test at baseline, 3- and 6-months post-surgery. CTS QST data underwent the same data transformations as performed in healthy volunteer reference data and as reported in Supplementary Material 2 (https://doi.org/10.6084/m9.figshare.12860066.v2).

Standardised values (z-scores), were calculated for QST tests, whereby the mean control value was subtracted from the mean value of the CTS participant and divided by the standard deviation of the controls. The use of z scores enables interpretation of QST results on the individual level, values outside the range $z \times \pm 1.96$ were interpreted as abnormal [57]. Z scores with a positive value denote a gain in function (hyperalgesia) whereas a negative value a loss of function.

$$z = \frac{\text{value of participant} - \text{mean of controls}}{\text{standard deviation of controls}}$$

Using the German Research Network on Neuropathic Pain (DFNS) algorithm [81], participants were defined as “healthy” or stratified to one of three sensory phenotypes primarily characterised by; 1. thermal and mechanical sensory loss (Sensory Loss); 2. preserved sensory function, associated with mild heat or cold hyperalgesia (Thermal Hyperalgesia) and 3. loss of thermal sensation, combined with mechanical hyperalgesia or allodynia (Mechanical Hyperalgesia)[6]. For clarity, and in the context of sensory testing in patients with CTS, “healthy” suggests that sensory function is not characterised by small fibre dysfunction as would be consistent with neuropathy; there may however be loss of large sensory fibre function as evidenced by reduced mechanical or vibration detection.
Chi-square test for independence was used to investigate the relationship between categorical variables (QST derived sensory phenotype, symptom severity sub-group and surgical outcome) at baseline and 6 months post-surgery; significance is reported for the Pearson Chi-Square value or Fisher’s Exact Probability Test where cell counts were less than 5. McNemar’s test was used to investigate change in repeated binary measures. Spearman’s Rank Order Correlation Coefficients were performed to explore associations between symptom severity parameters and QST derived sensory phenotype. Where correlation coefficients were statistically significant (p≤.05), the strength of relationship was interpreted as small r=.10 to .29; medium r=.30 to .49; large r=.50 to 1.0 [21]. Differences in symptom severity parameters, between sensory phenotypic groups, were investigated with the one-way between groups analysis of variance (ANOVA) with post hoc Tukey HSD test.

Results
Seventy-six CTS participants were enrolled between October 2014 and December 2016 and completed baseline study measures; however, 4 participants did not undergo surgery (one patient declined surgery; three patients had surgery cancelled due to ongoing medical investigations) (Figure 1). Baseline assessments were completed at [median (IQR)] 1(7) days prior to surgery, 3-month assessments were aimed at 90 days post-surgery and were completed at [median (IQR)] 91(31) days; 6- month assessments were aimed at 180 days post-surgery and were completed at [median (IQR)] 193(29) days. Of participants who dropped out of the study (n=3), one moved away and two reported they were too busy with work to attend.
Demographic data and key health parameters are reported in Table 2. In the majority of participants (76%) severity of nerve compression was graded from moderately to extremely severe [9]. For the generation of median nerve QST reference data, a convenience sample of fifty-four participants was selected from the Healthy volunteer QST study (Supplementary Material 2 https://doi.org/10.6084/m9.figshare.12860066.v2). Group differences for age and sex were not statistically significant (p>.05).

Surgical Outcome

At 6 months post-surgery, 59 (92%) participants reported a good surgical outcome (completely cured 28%; much better 52%; slightly better 13%) while 5 (8%) reported a poor outcome (unchanged 5%; worse 3%), demonstrating that for the vast majority of participants median nerve decompression surgery was indeed an effective intervention.

Incidence of Neuropathic Pain

Prior to surgery, 58 (76%) of participants met the case definition for neuropathic pain based on nerve conduction studies, symptom distribution and DN4 score (Table 3). At three months post-surgery, 11 (17%) participants had two or more abnormal QST scores indicating loss of sensory function consistent with a classification of neuropathic pain; at 6 months post-surgery...
11 (18%) had two or more abnormal QST scores. At all time points, localisation of pain within the median nerve distribution was most common. Participant reported pain distribution was not associated with the severity of nerve compression on nerve conduction studies (p=.44) or duration of symptoms prior to surgery (p=.89). Improvement (decrease) in DN4 scores was statistically significant at three months post-surgery (p<.001) and from baseline to 6 months post-surgery (p<.001) with a large effect size. There was a comparable reduction in the number of patients meeting the case definition for neuropathic pain; likewise, change was statistically significant from baseline to 3 months and baseline to six months (p<.001).

Insert Table 3

Pain parameters
All pain and symptom severity parameters were normally distributed. There was a significant effect for time (p < .001) and the magnitude of the effect size was large for all pain parameters (Table 4). Pairwise comparisons demonstrate a statistically significant decrease in pain and symptom severity for all pain parameters from baseline to 3 months and baseline to 6 months. Symptom Severity Scale score was not associated with severity of nerve compression on electrophysiological testing (p=.88).
Somatosensory function and sensory phenotype

Descriptive data (raw, not transformed) and differences in QST results for CTS participants and controls at baseline, 3- and 6-month assessments are reported in Table 5. Statistically significant differences for CTS compared to controls persist from baseline to 6 months postsurgery for cold detection threshold, thermal sensory limen, heat pain threshold, pressure pain thresholds, mechanical detection threshold and vibration detection threshold, demonstrating persistent small (Aδ, C) and large sensory fibre (Aβ) dysfunction. Dynamic mechanical allodynia is a pathological sensory response and was not exhibited by controls or CTS participants at any time point. Paradoxical heat sensations, also a pathological response, were not observed in controls and only infrequently observed in CTS participants (pre-surgery 14% participants; 3 months post-surgery 17%; 6 months post-surgery 10%).

Internal Responsiveness

CTS QST data was transformed, and Z scores calculated using transformed control group mean and standard deviations. Change in Z scores for repeated-measures QST thermal modalities from baseline to 6 months are illustrated in Figure 2; change in mechanical measures in Figure 3. Change in QST parameters across evaluations was statistically significant for thermal detection thresholds, thermal pain thresholds and mechanical detection thresholds from baseline to 3 months and baseline to 6 months. Changes in mechanical pain measures (mechanical pain threshold, mechanical pain sensitivity and pressure pain threshold) were not statistically significant at any time point. The magnitude of change (effect
size) was large for warm detection threshold, thermal sensory limen and mechanical detection threshold. A moderate effect size was identified for cold detection threshold, cold pain threshold, heat pain threshold and vibration detection threshold; change in pressure pain threshold was statistically significant only from baseline to 3 months post-surgery and magnitude of change was small (Table 6).

QST Derived Sensory Phenotype

Prior to surgery, 21% of participants were classified as having a “healthy” QST derived sensory phenotype [81]. At all evaluations, healthy profile, thermal and mechanical hyperalgesia phenotypes were more common than sensory loss phenotypes (Fig 4).

Statistically significant change in sensory phenotype was detected from baseline to 3 months post-surgery (p<.001) and 3 months to 6-months post-surgery (p<.001). The association of disease severity measures with sensory phenotype at baseline and 6 months was investigated.
Neither the duration of symptoms prior to surgery, localisation of symptoms (median, extra-median or non-median distribution) or the severity of nerve compression based on nerve conduction studies was associated with sensory phenotype at either time point (p> .05). At 6 months post-surgery, recovering sensory function is demonstrated by normalising QST values (Figs 2;3) and improvement in participant reported pain and symptom severity parameters (Tables 4;7). Healthy sensory profile (23%) and thermal hyperalgesia phenotype (46%) predominate, however, mechanical hyperalgesia (18%) and sensory loss (13%) phenotypes persist in one third of the sample.

**External Responsiveness**

To investigate the external responsiveness of QST derived sensory phenotype we explored the association of sensory phenotype with corresponding reference measures of pain and symptom severity at baseline, 3- and 6-months post-surgery. Baseline pain and symptom severity parameters were not found to be associated with sensory phenotype, however, there was a significant correlation between pain parameters and sensory phenotype at 3- and 6-months post-surgery (Table 7). Dimensions of neuropathic pain, by QST sensory phenotype, were further explored with the NPSI subscales (burning pain; pressing pan; paroxysmal pain; evoked pain; paraesthesia) at baseline and 6 months. Pain burden, again, is lower across all pain dimensions for participants with a healthy QST profile, however differences in pain dimensions between phenotypes were not statistically significant. Paraesthesia was the most common and most severe pain dimension, reported at baseline by 96% of participants and
rated as severe by 65% of those. At 6 months, the frequency of reporting of each dimension was reduced, with 58% of participants reporting no symptoms.

**Insert Table 7**

Baseline Symptom Severity Scale scores (SSS) were used to stratify participants into pain dominant (n=27; 36%), paraesthesia dominant (n=37; 49%) or mixed symptom (n=12; 16%) subgroups. In participants with a baseline healthy sensory profile (21% of cohort), thermal hyperalgesia (29%) and sensory loss phenotype (18%); symptoms of paraesthesia predominate (in 69%, 50% and 71% of phenotypic group participants, respectively). In contrast, in participants with a mechanical hyperalgesia phenotype (32% of cohort), symptoms of pain predominate (in 54% of phenotype). A statistically significant association was found for QST derived sensory phenotype and SSS symptom subgroups (Fisher’s Exact Test, 2-sided, p=.027) (*Fig 5*), demonstrating that QST derived sensory phenotype is consistent with or reflects patient reported pain and sensory dysfunction.

**Insert Figure 5**

Differences in pain and symptom severity parameters between sensory phenotypic groups were explored with the DN4, BPI Pain Severity Score, NPSI total score and Symptom Severity Scale score at base line and 6 months post-surgery (*Table 8*). At baseline, pain and symptom severity score ratings are lowest in participants with a healthy profile phenotype and highest in those with a sensory loss phenotype, however group differences were not
statistically significant. Group differences persist at 6 months post-surgery, with statistically significant differences detected between sensory phenotypic groups for the DN4, NPSI total score and symptom severity score. Differences in pain and symptom severity parameters, between phenotypic groups at baseline, 3- and 6-months post-surgery are illustrated in Figure 6. Statistically significant differences were found in DN4 scores between healthy profile and sensory loss phenotypes at 6 months post-surgery. Pain severity scores are greatest in those with mechanical hyperalgesia and sensory loss phenotypes as compared to those with a healthy profile at 3- and 6-months post-surgery, however phenotypic differences do not reach statistical significance (healthy profile compared to sensory loss; p=.06). For the NPSI total score and Symptom Severity Scale score, statistically significant differences were detected at both 3- and 6-months post-surgery (healthy profile compared to sensory loss phenotype; thermal hyperalgesia compared to sensory loss phenotype).

The association of sensory phenotype and patient-reported surgical outcome was explored with the Chi-square test for independence. Considering baseline, pre-operative phenotype, differences in outcome between phenotypic groups were observed; a good surgical outcome was reported by 100% of those with a healthy sensory profile, 95% with thermal
hyperalgesia, 91% sensory loss and 85% with mechanical hyperalgesia, nonetheless group differences were not statistically significant (Fisher’s Exact test p=.51). Sensory phenotype at 6 months was found, however, to be associated with patient reported surgical outcome (Fig 7). Of participants with a healthy sensory profile at 6 months, 93% reported a good surgical outcome, as did 100% of those with thermal hyperalgesia and 88% of those with sensory loss, while only 73% of participants with a mechanical hyperalgesia phenotype reported a good surgical outcome. Fisher’s Exact Probability Test indicated a significant association between sensory phenotype and good versus poor outcome at 6 months, $\chi^2 (3, n = 61) = 7.46, p = .02$. The effect size for this finding, Cramers’s V, was moderate, .36 [21].

Discussion

This novel study is the first to evaluate the responsiveness of DFNS QST and QST derived sensory phenotype in a longitudinal study of patients with compressive neuropathy following disease modifying intervention. A carpal tunnel surgery model was chosen as patients with CTS present with mixed pain and paraesthesia symptomology; there is a psychometrically robust comparator assessment of pain and paraesthesia in this clinical cohort and there is a disease modifying treatment of established efficacy for CTS. Comprehensive evidence for the responsiveness of DFNS QST and sensory phenotype were identified.
Internal responsiveness refers to the ability of an instrument to capture change over time. In the present study, statistically significant change was detected for the majority of DFNS QST modalities and in sensory phenotype at 3- and 6-months post-surgery. External responsiveness is the degree to which change in an instrument is associated with change in a corresponding reference measure. Clinically relevant reference measures, the Symptom Severity Scale, DN4, NPSI and BPI pain severity score, demonstrated significant improvement post-surgery. Pain and symptom severity reference measures were associated with sensory phenotype at 3- and 6-months post-surgery. At 3- and 6-months post-surgery, differences in symptom severity, between sensory phenotypic groups, were statistically significant. Additionally, pain and paraesthesia subgroups, derived from the patient-reported baseline SSS, were associated with sensory phenotype. Therefore, in patients with carpal tunnel syndrome, change in sensory phenotype reflects clinically relevant change.

Establishing the responsiveness of QST derived sensory phenotype has important implications for clinical practice and research. Instrument responsiveness suggests sensory phenotype might be employed to determine if treatment is resulting in clinically important change, to evaluate the effectiveness of programs of care and might underpin the identification of subgroups of patients who most benefit from care. For research purposes, the responsiveness of sensory phenotype suggests that statistically significant, clinically important change can be identified longitudinally. Furthermore, the magnitude of change, or effect size, is relevant for determining requisite study sample size.

Previous investigations in patients with CTS report an association between the severity of nerve compression on nerve conduction studies and the distribution of paraesthesias [17;85]. This association was not identified here, however, we explored pain distribution,
rather than paraesthesia, as our aim was to establish if pain distribution supported the
categorisation of pain as neuropathic. In patients with CTS, there is good evidence that the
severity of physiologic nerve compression is not associated with patient reported symptom
severity [39;55;78]; this dissociation was observed in the present cohort. Sensory phenotype,
similarly, while not associated with the severity of physiologic nerve compression, was
associated with patient reported symptoms. In patients with CTS, symptom severity and
sensory phenotype are independent of electrophysiologic measures of disease severity and
may reflect heterogeneity in underlying neuropathophysiology.

In this sample, a good surgical outcome was reported by 92% of participants. However, in
this pragmatic cohort, a lenient cut-point of 3 (slightly better) was chosen a priori to
determine surgical success. If a more stringent cut-point of grade of 4 (much better) was
taken, then a good outcome would have been reported by 80% of participants and is more in
keeping with the literature [10].

In patients with peripheral neuropathic pain, the frequency of DFNS QST derived sensory
phenotypes, thought to reflect different neurobiological mechanisms, differ between
aetiologies [6]. Tampin et al. [74] described the distribution of QST derived sensory
phenotypes in a cohort of 103 patients with CTS (healthy profile 23%; thermal hyperalgesia
20%; mechanical hyperalgesia 32%; sensory loss 18%). Except for a higher frequency of
thermal hyperalgesia (29%) in our cohort at baseline, phenotype distribution was similar and
comparable to that reported in patients with peripheral nerve injury [6]. Variability in the
distribution of phenotypes between CTS cohorts may result from differences in median nerve test site or in control data.

Intriguingly, we observed a decrease in the frequency of participants with a healthy sensory profile and increase in thermal hyperalgesia between 3 and 6-months post-surgery. Heat pain threshold demonstrated significant change up to 6 months post-surgery, and differences between CTS patients and controls become significant at 6 months. Change in sensory phenotype has not previously been reported in a carpal tunnel surgery cohort, however Baskozos et al. [7] similarly reported an increase in thermal sensitivity in CTS patients at 6 months post-surgery. Baskozos et al. demonstrate, histologically and electrodiagnostically, that while recovery of large and small fibre function is observed following median nerve decompression, recovery remains incomplete at 6 months. It is unclear, beyond 6 months, if small fibre function recovers further and thermal hyperalgesia resolves.

In the present cohort, prior to median nerve decompression surgery, loss of thermal detection (small fibre) and/or mechanical detection (large fibre) was more commonly observed than gain to thermal or mechanical stimuli. After surgery, while measures of large fibre function demonstrate significant improvement, at 6 months they remain reduced relative to reference data suggesting recovering but persistent dysfunction. In contrast, a mixed pattern of small fibre function recovery is observed. Cold detection threshold and thermal sensory limen improve significantly but remain reduced at 6 months, whereas warm detection threshold and cold pain threshold normalise at 3 months post-surgery.
While less common, gain of function was observed at baseline in mechanical pain measures (wind up ratio, mechanical pain threshold, mechanical pain sensitivity and pressure pain threshold) (Fig 3). Wind-up ratio, a measure of pathological response of small Aδ fibre function, normalises at 3 months post-surgery while pressure pain threshold decreases across assessments and remains significantly reduced at 6 months post-surgery. However, as pressure pain threshold was tested at the thenar eminence, near to the surgical incision, increase in pressure pain sensitivity may be consistent with post-surgical nociceptive pain.

As noted, Baskozos et al. [7] previously reported longitudinal change in DFNS QST in patients following carpal tunnel surgery. Consistent with our findings, their sample presented with reduced thermal and mechanical detection thresholds pre-surgery. While improvement in small and large fibre encoded modalities was observed post-surgery, sensory function does not normalise relative to control data. Their work provides further evidence for the responsiveness of DFNS QST. While 83% of their sample report a good surgical outcome, there is persistent somatosensory dysfunction on QST that is consistent with findings on post-surgical electrophysiological studies. Furthermore, somatosensory dysfunction is consistent with participant reported symptom severity; symptoms improve however approximately 50% of patients continue to report pain and 38% paraesthesia or numbness.

Equivocal findings for individual QST modalities between the Baskozos et al. sample and our results may arise from differences in administration of the DFNS QST protocol or in control data. Whereas Baskozos et al. performed sensory testing at the volar proximal phalanx of the index finger, the present sample was tested at the volar distal phalanx of the middle finger, purported to be more sensitive to sensory change in patients with CTS [29]. We demonstrated, in our healthy volunteer QST study, that published DFNS QST reference data for the dorsum of the hand [57] are not generalisable to the median nerve innervated volar
hand. In the absence of published median nerve reference data, investigators are reliant upon generating comparator QST control data, possibly resulting in discrepancies between studies. CTS is the most common of the entrapment neuropathies [61] and median nerve decompression surgery is one of the most commonly performed surgical procedures in the hand [8]. Future studies of median nerve somatosensory function can be anticipated and will benefit from the availability of published reference data for the DFNS QST protocol [70] for the median nerve distribution of the hand.

Study limitations

Addressing risk of bias
This study was reported according to STROBE guidelines [82] to reduce reporting bias. All patients scheduled for carpal tunnel surgery were invited to participate, however it is impossible to control for participant-selection bias. In clinical studies of this nature, the clinician-patient interaction cannot be ruled out and may influence or bias the patients’ perception or judgment of outcome. To reduce the likelihood of assessor bias the investigator (D.K.) was blinded to patient reported results of surgical outcome until patients completed the trial, however, no post hoc analysis was conducted to determine if or how often blinding was broken. The completed measures were placed in pre-labeled, coded envelopes by study participants and secured in the written case report form. As the baseline and post-surgical measures were completed by one investigator in the present study, there is risk of investigator bias.
Conclusions

In a carpal tunnel surgery model, DFNS QST and QST derived sensory phenotype identify statistically significant and clinically relevant change in somatosensory function after disease modifying intervention. Sensory phenotype at 6 months post-surgery was associated with and identified statistically significant differences in patient-reported surgical outcome. Following an efficacious intervention, significant differences were detected in persistent pain and symptom severity between phenotypic groups. QST derived sensory phenotype was not associated with disease severity measures including the duration or localisation of symptoms or severity of nerve compression and may reflect underlying neuropathophysiology. Future investigations of variables associated with sensory phenotype heterogeneity may yield evidence informing the elucidation of variability in response to intervention and underpin advances in personalised medicine. Our findings demonstrate QST derived sensory phenotype is responsive, enabling the identification of clinically important change.

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This report is independent research and the views expressed in this publication are those of the authors and not necessarily those of the NHS, the National Institute for Health Research or the Department of Health.

Contributions: DLK and ASCR designed the study. DR, JV CMA contributed to study concept and interpretation. DLK performed all participant testing. JV, DR and DLK performed statistical analysis. All authors contributed to drafting and critically revising the manuscript.
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Conflicts of interest:
DLK; DR; CMA Nothing to declare
JV: Received personal fees from Casquar, outside of the submitted work.
ASCR:
- IASP Council Member and Chair 18th World Congress on Pain Scientific Programme Committee
- ASCR undertakes consultancy and advisory board work for Imperial College Consultants- in the last 24 months this has included remunerated work for: Abide, Pharmanovo, Lateral, Novartis, Pharmaleads, Mundipharma, Orion, Asahi Kasei, Toray & Theranexis
- ASCR was the owner of share options in Spinifex Pharmaceuticals from which personal benefit accrued upon the acquisition of Spinifex by Novartis in July 2015 and from which payments continued until 2019.
- ASCR is named as an inventor on patents:
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Figure legends

Figure 1. Participant recruitment. Flow diagram of the recruitment of study participants

Figure 2. Change in QST thermal measures

Boxes represent the standard deviation, the centre line the mean. The black upper dotted line represents + 1.96z, the bottom dotted line - 1.96z. Scores between the two are interpreted as normal, those above as gain of function and below as loss of function. Significance is denoted as * at the 0.05 probability level; ** at 0.01; *** at 0.001. CDT, cold detection threshold; CPT, cold pain threshold; HPT, heat pain threshold; TSL, thermal sensory limen; WDT, warm detection threshold.

Figure 3. Change in QST mechanical measures

Boxes represent the standard deviation, the centre line the mean. The black upper dotted line represents + 1.96z, the bottom dotted line - 1.96z. Scores between the two are interpreted as
normal, those above as gain of function and below as loss of function. Significance is denoted as * at the 0.05 probability level; ** at 0.01; *** at 0.001. NS; not statistically significant. MDT, mechanical detection threshold; MPS, mechanical pain sensitivity; MPT, mechanical pain threshold; PPT, pressure pain threshold; VDT, vibration detection threshold; WUR, wind-up ratio.

Figure 4. Sensory phenotypes at baseline and 3- and 6-months post-surgery.

Figure 5. Percentage of participants with pain or paraesthesia dominant symptoms or mixed symptoms (pain and paraesthesia in equal measures) by sensory phenotype, at baseline.

Figure 6. Pain and symptom severity parameters explored by sensory phenotype at baseline, 3- and 6-months post-surgery. A. DN4; B. BPI Pain Severity Score; C. Neuropathic Pain Symptom Inventory (NPSI); D. Symptom Severity Scale score. Significance for Multiple Comparisons with post hoc Tukey test is denoted as * at the 0.05 probability level; ** at 0.01. NS, not statistically significant.

Figure 7. The association of sensory phenotype at 6 months and surgical outcome.
Table 1. DFNS Quantitative Sensory Testing Battery (8)

<table>
<thead>
<tr>
<th>Stimulus</th>
<th>Fibre</th>
<th>Measure</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Thermal Measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cold Detection Threshold (CDT)</td>
<td>Aδ</td>
<td>°C</td>
</tr>
<tr>
<td>Warm Detection Threshold (WDT)</td>
<td>C</td>
<td>°C</td>
</tr>
<tr>
<td>Thermal Sensory Limen (TSL)</td>
<td>Aδ &amp; C</td>
<td>°C</td>
</tr>
<tr>
<td>Paradoxical Heat Sensations (PHS)</td>
<td>Aδ &amp; C (PR)</td>
<td>x3</td>
</tr>
<tr>
<td>Cold Pain Threshold (CPT)</td>
<td>Aδ</td>
<td>°C</td>
</tr>
<tr>
<td>Heat Pain Threshold (HPT)</td>
<td>C</td>
<td>°C</td>
</tr>
<tr>
<td><strong>Mechanical Measures</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mechanical Detection Threshold (MDT)</td>
<td>Aβ</td>
<td>mN</td>
</tr>
<tr>
<td>Dynamic Mechanical Allodynia (DMA)</td>
<td>Aβ (SRF)</td>
<td>0-100</td>
</tr>
<tr>
<td>Mechanical Pain Threshold (MPT)</td>
<td>Aδ</td>
<td>mN</td>
</tr>
<tr>
<td>Mechanical Pain Sensitivity (MPS)</td>
<td>Aδ (SRF)</td>
<td>0-100</td>
</tr>
<tr>
<td>Windup Ratio (WUR)</td>
<td>Aδ (PR)</td>
<td>ratio</td>
</tr>
<tr>
<td>Vibration Detection Threshold (VDT)</td>
<td>Aβ</td>
<td>x8</td>
</tr>
<tr>
<td>Pressure Pain Threshold (PPT)</td>
<td>C</td>
<td>kg/cm²</td>
</tr>
</tbody>
</table>

*PR, pathological response; SRF, stimulus response function*
Table 2.
Baseline demographic parameters

<table>
<thead>
<tr>
<th></th>
<th>CTS (n=76)</th>
<th>Controls (n=54)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age mean years (SD)</td>
<td>58.5 (13.5)</td>
<td>54.9 (11.3)</td>
</tr>
<tr>
<td>Female sex n (%)</td>
<td>65 (86)</td>
<td>38 (70)</td>
</tr>
<tr>
<td>BMI mean (SD)</td>
<td>28.8 (6.8)</td>
<td></td>
</tr>
<tr>
<td>Duration of CTS symptoms</td>
<td>36 (42)</td>
<td></td>
</tr>
<tr>
<td>median months (IQR)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nerve conduction study severity</td>
<td>normal (3 (4))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>very mild (3 (4))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mild (16 (21))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>moderately severe (22 (29))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>severe (14 (18))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>very severe (16 (21))</td>
<td></td>
</tr>
<tr>
<td></td>
<td>extremely severe (2 (3))</td>
<td></td>
</tr>
</tbody>
</table>
Table 3

Incidence of neuropathic pain across assessments

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline (n=76)</th>
<th>3 months</th>
<th>6 months</th>
<th>ANOVA sig</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>DN4 mean (sd)</td>
<td>5.39 (2.05)</td>
<td>1.81 (1.77)</td>
<td>1.29 (1.91)</td>
<td>&lt;.001</td>
<td>.76</td>
</tr>
<tr>
<td>Pain distribution n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain free</td>
<td>7 (9)</td>
<td>17 (25)</td>
<td>25 (40)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median nerve</td>
<td>44 (60)</td>
<td>39 (57)</td>
<td>28 (45)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Extra-median</td>
<td>24 (32)</td>
<td>9 (12)</td>
<td>8 (13)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-median</td>
<td>1 (1)</td>
<td>4 (5)</td>
<td>1 (2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neuropathic pain n (%)</td>
<td>58 (76%)</td>
<td>7 (11%)</td>
<td>7 (12%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Change in DN4, pain distribution and percentage of participants categorised as having neuropathic pain at baseline; 3- and 6-months post-surgery. Change in DN4 between time points was investigated with the Paired-Samples T-test. Change from baseline to 3 months and baseline to 6 months was statistically significant (p <.001); change from 3 months to 6 months was not (p=.09). 3-month sample size: DN4 and neuropathic pain n=65; pain distribution n=69. 6-month sample size: DN4 and neuropathic pain n=61, pain distribution n=62. Sd, standard deviation; sig, statistical significance.
Table 4

Change in pain parameters across assessments

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>3 months</th>
<th>6 months</th>
<th>ANOVA sig</th>
<th>Effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline- 3 months</td>
<td>3 - 6 months</td>
<td>Baseline- 6 months</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>BPI pain severity (1-10)</strong></td>
<td>4.17 (2.73)</td>
<td>1.89 (2.48)</td>
<td>1.45 (2.15)</td>
<td>&lt;.001</td>
<td>.49</td>
</tr>
<tr>
<td>Paired t-test sig</td>
<td>&lt;.001</td>
<td>.35</td>
<td>&lt;.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>NPSI (0-100)</strong></td>
<td>35.3 (24.2)</td>
<td>14.52 (18.97)</td>
<td>9.55 (15.0)</td>
<td>&lt;.001</td>
<td>.57</td>
</tr>
<tr>
<td>Paired t-tests sig</td>
<td>&lt;.001</td>
<td>.006</td>
<td>&lt;.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Symptom Severity Score (1-5)</strong></td>
<td>3.18 (.81)</td>
<td>1.78 (.73)</td>
<td>1.64 (.74)</td>
<td>&lt;.001</td>
<td>.78</td>
</tr>
<tr>
<td>Paired t-test sig</td>
<td>&lt;.001</td>
<td>.052</td>
<td>&lt;.001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Baseline, 3 month and 6-month data reported with mean and standard deviation. Sig, statistical significance.
<table>
<thead>
<tr>
<th>Test</th>
<th>Controls n=54</th>
<th>CTS Baseline n=76</th>
<th>CTS 3 months n=65</th>
<th>CTS 6 months n=61</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>sig</td>
<td>sig</td>
<td>sig</td>
<td></td>
</tr>
<tr>
<td>CDT, °C</td>
<td>-3.09 (3.02)</td>
<td>-5.65 (5.39)</td>
<td>&lt;.001</td>
<td></td>
</tr>
<tr>
<td>WDT, °C</td>
<td>5.75 (4.99)</td>
<td>7.19 (5.0)</td>
<td>.004</td>
<td></td>
</tr>
<tr>
<td>TSL, °C</td>
<td>9.85 (7.06)</td>
<td>15.01 (10.6)</td>
<td>&lt;.001</td>
<td></td>
</tr>
<tr>
<td>CPT, °C</td>
<td>10.53 (10.96)</td>
<td>5.0 (8.72)</td>
<td>.003</td>
<td></td>
</tr>
<tr>
<td>HPT, °C</td>
<td>47.22 (5.31)</td>
<td>46.79 (5.87)</td>
<td>.67</td>
<td></td>
</tr>
<tr>
<td>PPT, kg/cm²</td>
<td>401 (208.75)</td>
<td>376 (173.5)</td>
<td>.02</td>
<td></td>
</tr>
<tr>
<td>MPT, mN</td>
<td>152.43 (171.6)</td>
<td>100.49 (195.25)</td>
<td>.06</td>
<td></td>
</tr>
<tr>
<td>MPS, 0-100</td>
<td>.29 (.37)</td>
<td>.36 (.92)</td>
<td>.15</td>
<td>.48</td>
</tr>
<tr>
<td>WUR, ratio</td>
<td>.195 (1.53)</td>
<td>1.56 (.95)</td>
<td>.04</td>
<td>1.60 (1.81)</td>
</tr>
<tr>
<td>MDT, mN</td>
<td>.25 (.25)</td>
<td>1.14 (3.44)</td>
<td>&lt;.001</td>
<td>.71 (.78)</td>
</tr>
<tr>
<td>VDT x/8</td>
<td>8 (1)</td>
<td>6.67 (3)</td>
<td>&lt;.001</td>
<td>7.33 (2.17)</td>
</tr>
</tbody>
</table>

Data is reported with median (interquartile range). Statistical significance is reported for results of the Mann-Whitney U test comparing healthy volunteer control data to CTS patients at 3 time points; baseline, 3- and 6-months post-surgery. CDT, cold detection threshold; CPT, cold pain threshold; HPT, heat pain threshold; IQR, interquartile range; MDT, mechanical detection threshold; MPS, mechanical pain sensitivity; MPT, mechanical pain threshold; PPT, pressure pain threshold; sig, statistical significance; TSL, thermal sensory limen; VDT, vibration detection threshold; WDT, warm detection threshold; WUR, wind-up ratio. Bold represents statistical significance.
Table 6

Responsiveness of QST

<table>
<thead>
<tr>
<th></th>
<th>Baseline to 3 months p=</th>
<th>z=</th>
<th>effect size</th>
<th>Baseline to 6 months p=</th>
<th>z=</th>
<th>effect size</th>
</tr>
</thead>
<tbody>
<tr>
<td>CDT</td>
<td>&lt;.001</td>
<td>3.51</td>
<td>0.44</td>
<td>0.005</td>
<td>2.79</td>
<td>0.36</td>
</tr>
<tr>
<td>WDT</td>
<td>&lt;.001</td>
<td>4.03</td>
<td>0.50</td>
<td>0.007</td>
<td>2.69</td>
<td>0.34</td>
</tr>
<tr>
<td>TSL</td>
<td>&lt;.001</td>
<td>4.25</td>
<td>0.53</td>
<td>0.005</td>
<td>2.83</td>
<td>0.36</td>
</tr>
<tr>
<td>CPT</td>
<td>0.003</td>
<td>2.30</td>
<td>0.37</td>
<td>0.007</td>
<td>2.70</td>
<td>0.35</td>
</tr>
<tr>
<td>HPT</td>
<td>0.007</td>
<td>2.69</td>
<td>0.33</td>
<td>0.004</td>
<td>2.91</td>
<td>0.37</td>
</tr>
<tr>
<td>MDT</td>
<td>&lt;.001</td>
<td>4.73</td>
<td>0.59</td>
<td>&lt;.001</td>
<td>5.28</td>
<td>0.68</td>
</tr>
<tr>
<td>VDT</td>
<td>&lt;.001</td>
<td>3.52</td>
<td>0.44</td>
<td>&lt;.001</td>
<td>5.28</td>
<td>0.68</td>
</tr>
<tr>
<td>PPT</td>
<td>0.03</td>
<td>2.17</td>
<td>0.27</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

N=76 at baseline, N=61 at 6 months. CDT, cold detection threshold; CPT, cold pain threshold; HPT, heat pain threshold, MDT, mechanical detection threshold; PPT, pressure pain threshold; TSL, thermal sensory limen; VDT, vibration detection threshold; WDT, warm detection threshold.
<table>
<thead>
<tr>
<th>Pain Parameter</th>
<th>BPI pain severity score</th>
<th>NPSI total score</th>
<th>Symptom Severity Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensory Phenotype</td>
<td>r (p)</td>
<td>r (p)</td>
<td>r (p)</td>
</tr>
<tr>
<td>Baseline</td>
<td>0.142 (0.22)</td>
<td>0.132 (0.26)</td>
<td>0.188 (0.10)</td>
</tr>
<tr>
<td>3 months</td>
<td>0.230 (0.06)</td>
<td>0.313 (0.01)</td>
<td>0.327 (0.008)</td>
</tr>
<tr>
<td>6 months</td>
<td>0.266 (0.03)</td>
<td>0.439 (&lt;0.001)</td>
<td>0.346 (0.006)</td>
</tr>
</tbody>
</table>

Table 7. Association of sensory phenotype with pain and symptom severity parameters at baseline, 3- and 6-months post-surgery. $r =$ Spearman’s rank order correlation coefficient, $p =$ statistical significance. Significant associations in **bold**.
<table>
<thead>
<tr>
<th></th>
<th>Healthy Profile</th>
<th>Thermal Hyperalgesia</th>
<th>Mechanical Hyperalgesia</th>
<th>Sensory loss</th>
<th>p=</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DN4</strong></td>
<td>Baseline</td>
<td>5.2 (2.43)</td>
<td>5.3 (1.96)</td>
<td>5.8 (1.91)</td>
<td>5.8 (1.76)</td>
</tr>
<tr>
<td></td>
<td>6 months</td>
<td>.43 (.51)</td>
<td>1.25 (1.71)</td>
<td>1.82 (2.56)</td>
<td>3.0 (2.73)</td>
</tr>
<tr>
<td><strong>BPI Pain Severity</strong></td>
<td>Baseline</td>
<td>3.1 (2.05)</td>
<td>4.6 (2.97)</td>
<td>4.5 (2.83)</td>
<td>4.8 (3.13)</td>
</tr>
<tr>
<td></td>
<td>6 months</td>
<td>1.04 (1.60)</td>
<td>.67 (1.37)</td>
<td>2.41 (2.81)</td>
<td>3.38 (3.33)</td>
</tr>
<tr>
<td><strong>NPSI</strong></td>
<td>Baseline</td>
<td>28.2 (20.6)</td>
<td>41.6 (28.13)</td>
<td>38.7 (25.12)</td>
<td>40.6 (23.81)</td>
</tr>
<tr>
<td></td>
<td>6 months</td>
<td>3.21 (7.32)</td>
<td>6.07 (9.94)</td>
<td>14.73 (20.48)</td>
<td>25.5 (23.07)</td>
</tr>
<tr>
<td><strong>Symptom Severity Score</strong></td>
<td>Baseline</td>
<td>3.0 (0.59)</td>
<td>3.2 (0.88)</td>
<td>3.4 (0.81)</td>
<td>3.4 (1.01)</td>
</tr>
<tr>
<td></td>
<td>6 months</td>
<td>1.34 (.30)</td>
<td>1.44 (.52)</td>
<td>2.01 (.99)</td>
<td>2.44 (1.10)</td>
</tr>
</tbody>
</table>

Differences in pain and symptom severity parameters between phenotypic groups at baseline and 6 months post-surgery reported with mean (standard deviation). Significance is reported for one way between groups Analysis of Variance (ANOVA) with post hoc Tukey HSD test; at 6 months Welch test is reported due to homogeneity in variance. Distribution of the sample across sensory phenotypes; number and percentage of patients with neuropathic pain reported in brackets [n; %]: baseline N=76 [58; 76%]; healthy profile n=16; 21% [10; 17%]; thermal hyperalgesia n=24; 32% [15; 26%]; mechanical hyperalgesia n= 24; 32% [22; 38%]; sensory loss n=14; 18% [11; 19%]. 6 months N= 61 [7; 12%]; healthy profile n=14; 23% [0]; thermal hyperalgesia n=28; 46% [1; 14%]; mechanical hyperalgesia n=11; 18% [2; 27%]; sensory loss n=8; 13% [4; 57%].