KEY WORDS: Straphizing; Neuropathic pain; Topical analgesics; Treatment efficacy

Pain catastrophizing predicts poor response to topical analgesics in patients with neuropathic pain. T Mankovsky BA1, Mary E Lynch MD2, AJ Clark MD3, J Sawynok PhD4, Michael JL Sullivan PhD1

BACKGROUND: Previous research suggests that high levels of pain catastrophizing might predict poorer response to pharmacological interventions for neuropathic pain.

OBJECTIVE: The present study sought to examine the clinical relevance of the relation between catastrophizing and analgesic response in individuals with neuropathic pain. Clinically meaningful reductions were defined in terms of the magnitude of reductions in pain through the course of treatment, and in terms of the number of patients whose end-of-treatment pain ratings were below 4/10.

METHODS: Patients (n=82) with neuropathic pain conditions completed a measure of pain catastrophizing at the beginning of a three-week trial examining the efficacy of topical analgesics for neuropathic pain.

RESULTS: Consistent with previous research, high scores on the measure of pain catastrophizing prospectively predicted poorer pain response to treatment. Fewer catastrophizers than noncatastrophizers showed moderate (≥2 points) or substantial reductions in pain ratings through the course of treatment. Fewer catastrophizers than noncatastrophizers achieved end-of-treatment pain ratings below 4/10.

CONCLUSIONS: The results of the present study suggest that the development of brief interventions specifically targeting catastrophic thinking might be useful for enhancing the effects of pharmacological interventions for neuropathic pain. Furthermore, failure to account for the level of catastrophizing might contribute to null findings in clinical trials of analgesic medication.

Key Words: Catastrophizing; Neuropathic pain; Topical analgesics; Treatment efficacy

It has been estimated that there are more than 300,000 patients suffering from neuropathic pain in Canada and more than 3.8 million in the United States (1,2). Neuropathic pain has been defined as “pain arising as a direct consequence of a lesion or disease affecting the somatosensory system” (3). As a result of the aging population and the increase in survival rates following interventions that give rise to neuropathic pain conditions (eg, cancer), the prevalence of neuropathic pain conditions is expected to increase significantly over the next two decades (4).

Pharmacological interventions are the dominant treatment approach for neuropathic pain (5,6). Unfortunately, many clinical trials of available treatments for neuropathic pain have yielded disappointing results, and many patients with neuropathic pain conditions continue to experience ongoing distressing and disabling symptoms of pain (7,8). It has been suggested that as many as one-half of the patients with neuropathic pain conditions are incompletely or totally refractory to available treatments (9).

Recent studies have examined the relationship between pain catastrophizing and pain experience in individuals suffering from neuropathic pain conditions (10). Pain catastrophizing has been defined as a negative cognitive set brought to bear during actual or anticipated pain experience, comprising elements of rumination, magnification and helplessness (11,12). Pain catastrophizing has been associated with heightened pain experience in patients with postherpetic neuralgia (13), phantom limb pain (14), post-traumatic neuralgia (15) and pain associated with spinal cord injury (10).

There are also indications that pain catastrophizing might represent a risk factor for poor outcomes in clinical interventions for neuropathic pain. Haythornthwaite et al (13) reported the findings of a study in which patients with postherpetic neuralgia were enrolled in a crossover randomized clinical trial comparing an opioid, an anti-depressant and a placebo. Analyses revealed that initial pain catastrophizing scores predicted higher post-treatment pain ratings, even when controlling for baseline pain. It has been suggested that catastrophizing might be associated with dysregulation or dysfunction in endogenous opioid pain-control systems that might compromise the effectiveness of pharmacological interventions for pain (16,17).

Although the relationship between catastrophizing and poor treatment outcomes has been observed in several studies, the clinical relevance of this relationship remains unclear. Before advocating for treatment, a more complete understanding of the relationship between pain catastrophizing and pain is required. This study was designed to examine the clinical relevance of the relation between pain catastrophizing and analgesic response in individuals with neuropathic pain.

An additional objective of this study was to determine whether the development of brief interventions specifically targeting catastrophic thinking might contribute to null findings in clinical trials of analgesic medication.

OBJECTIVE: The present study sought to examine the clinical relevance of the relation between catastrophizing and analgesic response in individuals with neuropathic pain. Clinically meaningful reductions were defined in terms of the magnitude of reductions in pain through the course of treatment, and in terms of the number of patients whose end-of-treatment pain ratings were below 4/10.

METHODS: Patients (n=82) with neuropathic pain conditions completed a measure of pain catastrophizing at the beginning of a three-week trial examining the efficacy of topical analgesics for neuropathic pain.

RESULTS: Consistent with previous research, high scores on the measure of pain catastrophizing prospectively predicted poorer pain response to treatment. Fewer catastrophizers than noncatastrophizers showed moderate (≥2 points) or substantial reductions in pain ratings through the course of treatment. Fewer catastrophizers than noncatastrophizers achieved end-of-treatment pain ratings below 4/10.

CONCLUSIONS: The results of the present study suggest that the development of brief interventions specifically targeting catastrophic thinking might be useful for enhancing the effects of pharmacological interventions for neuropathic pain. Furthermore, failure to account for the level of catastrophizing might contribute to null findings in clinical trials of analgesic medication.

Key Words: Catastrophizing; Neuropathic pain; Topical analgesics; Treatment efficacy

HISTORIQUE: Selon des recherches antérieures, de forts taux de catastrophisation de la douleur pourraient prédire une moins bonne réponse aux interventions pharmacologiques pour soulager la douleur neuropathique.

OBJECTIF: La présente étude visait à examiner la pertinence clinique du lien entre la catastrophisation et la réponse analgésique des personnes ayant une douleur neuropathique. Des réductions cliniquement significatives étaient définies d’après la magnitude des réductions de la douleur pendant la durée du traitement et le nombre de patients dont la douleur en fin de traitement se situait sous les 4/10.

MÉTHODOLOGIE: Les patients (n=82) ayant des maladies provoquant une douleur neuropathique ont rempli une mesure de catastrophisation de la douleur au début d’un essai de trois semaines portant sur l’efficacité des analgésiques topiques pour soulager la douleur neuropathique.

RÉSULTATS: Conformément aux recherches antérieures, les indices élevés sur la mesure de catastrophisation de la douleur ont prédit prospectivement une moins bonne réponse au traitement. Moins de personnes faisant de la catastrophisation que de personnes n’en faisant pas ont montré une réduction modérée (≥2 points) ou substantielle d’évaluations de la douleur pendant le traitement. Moins de personnes faisant de la catastrophisation que de personnes n’en faisant pas sont parvenues à une évaluation de la douleur de moins de 4/10 en fin de traitement.

CONCLUSIONS: D’après les résultats de la présente étude, l’élaboration de brèves interventions ciblant expressément la pensée catastrophique pourrait être utile pour améliorer les effets des interventions pharmacologiques pour soulager la douleur neuropathique. De plus, le fait de ne pas tenir compte du niveau de catastrophisation peut contribuer à des observations nulles dans des essais cliniques sur les analgésiques.

1Department of Psychology, McGill University, Montreal, Quebec; 2Pain Management Unit, Queen Elizabeth II Health Sciences Centre, Dalhousie University, Halifax, Nova Scotia; 3Chronic Pain Centre, Calgary Health Region, Calgary, Alberta; 4Department of Pharmacology, Dalhousie University, Halifax, Nova Scotia

Correspondence: Dr Michael Sullivan, Department of Psychology, McGill University, 1205 Docteur Penfield, Montréal, Québec H3A 1B1.

Telephone 514-398-4455, fax 514-398-4896, e-mail michael.sullivan@mcgill.ca

10 ©2012 Pulsus Group Inc. All rights reserved
catastrophizing-reduction interventions to improve pain outcomes, it is necessary to determine whether the relationship between catastrophizing and treatment response is sufficiently robust to be considered clinically meaningful.

One approach to defining the clinical meaningfulness of pain reduction has been to assess the magnitude of pain reduction achieved by an analgesic agent. It has been suggested that a 2-point reduction on an 11-point numerical rating scale (NRS) represents moderate improvement in pain, and that a 4-point reduction represents a substantial improvement in pain (18,19). Another approach to defining clinical meaningfulness of pain reduction has been to examine the proportion of patients whose pain levels falls below 4 on an 11-point NRS (20). A value of 4/10 corresponds to the boundary between mild and moderate pain.

The primary objective of the present study was to examine the clinical meaningfulness of the relationship between catastrophizing and analgesic response in a sample of patients with neuropathic pain. Patients with varied neuropathic pain conditions completed a measure of catastrophizing before initiating a three-week course of a topical analgesic. Analyses addressed whether high catastrophizers were less likely than low catastrophizers to show a 2-point or greater reduction in pain ratings in response to treatment, and whether fewer catastrophizers than noncatastrophizers achieved post-treatment pain ratings less than 4/10.

**METHODS**

**Participants**

The study sample comprised 82 patients (41 men, 41 women) who were enrolled in one of two clinical trials examining the efficacy of topical analgesics for neuropathic pain. Patients had diagnoses of diabetic neuropathy (n=16), postherpetic neuralgia (n=10) or postsurgical or post-traumatic neuropathic pain (n=56). The mean age of participants was 51.1 years with a range of 24 to 84 years. The mean (± SD) duration of pain was 70.9±66.8 months.

**Procedure**

The study sample included participants who were recruited from three hospital outpatient pain management units in eastern Canada for enrollment in two clinical trials examining the efficacy of topical analgesics for neuropathic pain. Patients with varied neuropathic pain conditions completed a measure of catastrophizing before initiating a three-week course of a topical analgesic. Analyses addressed whether high catastrophizers were less likely than low catastrophizers to show a 2-point or greater reduction in pain ratings in response to treatment, and whether fewer catastrophizers than noncatastrophizers achieved post-treatment pain ratings less than 4/10.

**Measures**

**Pain catastrophizing:** The Pain Catastrophizing Scale (PCS) was used to measure catastrophic thinking related to pain. The PCS comprises 13 items describing different thoughts and feelings associated with pain. The PCS has been shown to have high internal consistency (coefficient alpha = 0.87) and to be associated with heightened pain, pain behaviour and pain-related disability (11,15,23).

**Pain severity:** Patients completed a daily pain diary in which they were asked to rate the severity of their pain on an 11-point NRS with the end points of no pain (score = 0) and severe pain (score = 10). Pain severity ratings were averaged to yield four pain scores: baseline, week 1 of treatment, week 2 of treatment and week 3 of treatment. For the purposes of the present study, moderate pain reduction was defined as a reduction in pain ratings from baseline to week 3 of ≥2 points, and substantial pain reduction was defined as a reduction in pain ratings of ≥4 points.

**RESULTS**

**Correlates of initial pain severity and treatment response**

Correlates of initial pain severity and treatment response are presented in Table 2. Age and pain duration were not significantly correlated with initial pain severity or change in pain. Consistent with previous research, the PCS was positively correlated with initial pain severity (12). The PCS was negatively correlated with pain reduction.

**Pain severity:**

Pain severity was analyzed using logistic regression analyses. Table 3 summarizes the mean pre- and post-treatment pain ratings as a function of level of catastrophizing. Figure 1 shows the number of catastrophizers (56%) to show a 2-point or greater reduction in pain ratings. Again, catastrophizers (7%) were less likely than noncatastrophizers to show a ≥2-point reduction in pain ratings. Overall, 42% of patients showed a reduction of ≥2 points in pain ratings. Treatment condition assignment (χ²=1.14; P=NS) did not vary significantly as a function of sex.

There was a tendency for men to be more likely to have a diagnosis of diabetic neuropathy, but the difference did not attain statistical significance (χ²=5.1, P=0.07).

**Correlates of initial pain severity and treatment response**

Correlates of initial pain severity and treatment response are presented in Table 2. Age and pain duration were not significantly correlated with initial pain severity or change in pain. Consistent with previous research, the PCS was positively correlated with initial pain severity (12). The PCS was negatively correlated with pain reduction.

**Catastrophizing and treatment response**

Table 3 summarizes the mean pre- and post-treatment pain ratings as a function of level of catastrophizing. Figure 1 shows the number of catastrophizers and noncatastrophizers who showed a 2-point or greater reduction in pain ratings through the course of treatment. Overall, 42% of patients showed a reduction of ≥2 points in pain ratings. Catastrophizers (30%) were significantly less likely than noncatastrophizers (56%) to show a ≥2-point reduction in pain ratings (χ²=6.0; P<0.01).

Overall, 16% of patients showed a reduction of ≥4 points in pain ratings. Again, catastrophizers (7%) were less likely than noncatastrophizers (24%) to show a ≥4-point reduction in pain ratings (χ²=4.5; P<0.05) (Figure 1).

Figure 1 shows the number of catastrophizers and noncatastrophizers whose pain ratings fell below 4 at the end of treatment. Overall, 24% of patients’ pain ratings were below 4/10 at the end of treatment. Low scores on catastrophizing were associated with a threefold increase (36%) in the probability of achieving end-of-treatment pain ratings below 4/10 compared with high scores on catastrophizing (12%) (χ²=6.2; P<0.01).

Results from the logistic regression analyses revealed that the relationship between catastrophizing and poor treatment response remained significant even when controlling for initial pain ratings (OR 0.16 [95% CI 0.03 to 0.77]; P<0.05). Similarly, the relationship between
Consequently, as a function of regression to the mean, both catastrophizers typically be higher than noncatastrophizers’ pretreatment pain scores. At the end of treatment, catastrophizers’ pretreatment pain scores will continue to be higher than noncatastrophizers’ pretreatment pain scores. This finding extends previous research by showing that the relationship between catastrophizing and poor treatment response can also be considered clinically meaningful.

The findings of the present study join growing literature highlighting the importance of psychological factors in the experience of neuropathic pain (15,24-26). The findings are consistent with previous research showing that high levels of catastrophizing predict poorer response to pharmacological interventions for neuropathic pain (12), and extend previous research by showing that the relationship between catastrophizing and poor treatment response can also be considered clinically meaningful.

Overall, 42% of patients showed moderate improvement in response to treatment and 16% showed substantial improvement. The magnitude of treatment response is similar to that reported in other studies examining the efficacy of analgesic agents in individuals with neuropathic pain (27-29). Studies examining the efficacy of analgesic agents in individuals with neuropathic pain have shown that high levels of persistent pain in spite of the treatment they receive may contribute to inaccurate conclusions regarding their efficacy.

The mechanisms by which catastrophizing interferes with the response to analgesia remain unclear. However, accumulating evidence suggests that catastrophic thinking may be associated with dysregulation of endogenous opioid pain-control systems (31). In a study investigating postsurgical pain following breast cancer surgery, Jacobsen et al (30) found that individuals with high levels of catastrophizing required higher doses of postoperative opioid analgesics to control their postsurgical pain. A more recent study by Goodin et al (31) found that pain catastrophizing was negatively associated with diffuse nociceptive inhibitory controls, a psychophysical measure of endogenous pain inhibition, suggesting that high levels of pain catastrophizing may be related to a disruption in the endogenous modulation of pain.

Several psychological and rehabilitative interventions have been shown to yield reductions in catastrophic thinking (32-34). The results of the present research suggest that interventions specifically designed to reduce catastrophizing might similarly yield better treatment response for patients with neuropathic pain conditions. It is possible that emotional disclosure techniques might be useful in reducing pain patients’ tendency to focus excessively on pain symptoms. Previous research has shown that emotional disclosure manipulations can lead to decreases in emotional distress and reduced preoccupation with emotionally laden information (35). Similarly, the absence of emotional disclosure has been linked to increased thought intrusions and a variety of negative health outcomes (36). Sullivan and Neish (37) showed that high catastrophizers who were provided with an opportunity to disclose their emotional concerns before undergoing dental treatment, reported less pain than high catastrophizers who did not disclose. Emotional disclosure techniques can be effectively incorporated into clinical interviews, and have been shown to have beneficial impact even after one session (37,38).

### Table 1
Sample characteristics (n=82)

<table>
<thead>
<tr>
<th></th>
<th>Women</th>
<th>Men</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>41</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>Age, years</td>
<td>49.0±3.4</td>
<td>53.3±4.7</td>
<td>NS</td>
</tr>
<tr>
<td>Diagnosis (phn/pit/pdn), n</td>
<td>5/4/2</td>
<td>5/4/2</td>
<td>NS</td>
</tr>
<tr>
<td>Treatment condition (ket/ami/ami-ket), n</td>
<td>11/10/20</td>
<td>7/11/23</td>
<td>NS</td>
</tr>
<tr>
<td>Pain duration, months</td>
<td>65.7±8.4</td>
<td>76.9±6.4</td>
<td>NS</td>
</tr>
<tr>
<td>Baseline pain severity</td>
<td>7.0±1.0</td>
<td>6.9±1.2</td>
<td>NS</td>
</tr>
<tr>
<td>PCS</td>
<td>22.8±11.2</td>
<td>26.2±10.7</td>
<td>NS</td>
</tr>
</tbody>
</table>

Data presented as mean ± SD unless otherwise indicated. amti Amtriptyline; ami-ket Amitriptyline + ketamine; dn Diabetic neuropathy; ket Ketamine; NS Not significant; PCS Pain Catastrophizing Scale; phn Postherpetic neuralgia; pstn Postsurgical/traumatic neuropathy

### Table 2
Correlates of initial pain severity and treatment response (n=82)

<table>
<thead>
<tr>
<th></th>
<th>Baseline pain severity</th>
<th>Change in pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.05</td>
<td>0.04</td>
</tr>
<tr>
<td>Pain duration</td>
<td>0.05</td>
<td>0.01</td>
</tr>
<tr>
<td>Pain catastrophizing scale</td>
<td>0.22*</td>
<td>0.42**</td>
</tr>
</tbody>
</table>

*p<0.05; **P<0.01

### Table 3
Mean pain intensity ratings at baseline and at week 3

<table>
<thead>
<tr>
<th>PCS</th>
<th>n Mean±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain severity at baseline (0 to 10) Noncatastrophizers</td>
<td>41 6.69±0.99</td>
</tr>
<tr>
<td>Catalystrophizers</td>
<td>41 7.27±1.32</td>
</tr>
<tr>
<td>Pain severity week 3 (0 to 10) Noncatastrophizers</td>
<td>41 4.34±2.09</td>
</tr>
<tr>
<td>Catalystrophizers</td>
<td>41 6.11±1.97</td>
</tr>
</tbody>
</table>

PCS Pain Catastrophizing Scale

Figure 1: Percentage of catastrophizers and noncatastrophizers with 2-point and 4-point or greater reductions in pain ratings.

Their pain. However, when assessed in terms of a threshold for treatment success (ie, achieving scores in the mild range of pain), the higher initial pain scores of catastrophizers reduces the likelihood that the treatment success threshold will be attained. As such, catastrophizers are more likely than noncatastrophizers to continue suffering from high levels of persistent pain in spite of the treatment they receive.

The ineffectiveness of pharmacological interventions for neuropathic pain may be due to inaccurate conclusions regarding their efficacy. The mechanisms by which catastrophizing interferes with the response to analgesics remain unclear. However, accumulating evidence suggests that catastrophic thinking may be associated with dysregulation of endogenous opioid pain-control systems. In a study investigating postsurgical pain following breast cancer surgery, Jacobsen et al (30) found that individuals with high levels of catastrophizing required higher doses of postoperative opioid analgesics to control their postsurgical pain.

The mechanisms by which catastrophizing interferes with the response to analgesics remain unclear. However, accumulating evidence suggests that catastrophic thinking may be associated with dysregulation of endogenous opioid pain-control systems. In a study investigating postsurgical pain following breast cancer surgery, Jacobsen et al (30) found that individuals with high levels of catastrophizing required higher doses of postoperative opioid analgesics to control their postsurgical pain. A more recent study by Goodin et al (31) found that pain catastrophizing was negatively associated with diffuse nociceptive inhibitory controls, a psychophysical measure of endogenous pain inhibition, suggesting that high levels of pain catastrophizing may be related to a disruption in the endogenous modulation of pain.

Several psychological and rehabilitative interventions have been shown to yield reductions in catastrophic thinking. The results of the present research suggest that interventions specifically designed to reduce catastrophizing might similarly yield a better treatment response for patients with neuropathic pain conditions.

It is possible that emotional disclosure techniques might be useful in reducing pain patients’ tendency to focus excessively on pain symptoms. Previous research has shown that emotional disclosure manipulations can lead to decreases in emotional distress and reduced preoccupation with emotionally laden information. Similarly, the absence of emotional disclosure has been linked to increased thought intrusions and a variety of negative health outcomes. Sullivan and Neish (37) showed that high catastrophizers who were provided with an opportunity to disclose their emotional concerns before undergoing dental treatment, reported less pain than high catastrophizers who did not disclose. Emotional disclosure techniques can be effectively incorporated into clinical interviews, and have been shown to have beneficial impact even after one session (37,38).
Fear reduction techniques might also be an important component of interventions designed to target catastrophizing because mechanisms related to fear might contribute to alarmist reactions to pain stimuli (39,40). Because pain catastrophizing and pain-related fear are partially overlapping constructs, the fear component of catastrophizing might lead individuals to process information about pain-related stimuli preferentially (41,42). Exposure interventions have been shown to be effective in yielding reductions in fear and catastrophic thinking (43). Cognitive techniques designed to reduce the threat value of pain symptoms might also be useful in reducing the pain focus of high catastrophizers (44).

Finally, it is possible that activity-mobilization interventions might provide a useful alternative to cognitive distraction strategies for assisting high catastrophizers in reducing their attentional focus on pain (33). Because activity involvement demands attention, focus on activity might limit the cognitive resources that can be used to attend to pain-related stimuli. Although clinical trials evaluating the most effective approaches to reducing catastrophic thinking have yet to be conducted, there are indications that activity-based interventions can yield significant reductions in catastrophic thinking (45,46).

The bulk of studies demonstrating the efficacy of catastrophizing-reduction techniques involved treatment programs that typically extended over a period of 10 to 12 weeks and used functional improvement as the main objective of treatment. Considered either in terms of duration or treatment focus, these interventions might not be entirely applicable in the context of improving response to analgesic medication. The development and evaluation of new approaches to reducing catastrophic thinking in individuals with neuropathic pain may be a fruitful line of inquiry.

Caution must be exercised in the interpretation of the present findings. First, the data were drawn from patients who participated in clinical trials for the treatment of neuropathic pain. Volunteers for clinical trials likely differ in important ways from patients who typically seek treatment for neuropathic pain. In addition, factors associated with treatment response to topical analgesics might also differ in important ways from factors associated with treatment response to other modes of administration. Similarly, the mechanisms of action of topical agents might differ from mechanisms of action of other routes of medication administration. Finally, it is difficult to address whether the exclusion of high catastrophizers from the data would lead to a significant effect of the drug treatment within the current study sample. Some of the cases were drawn from an open-label trial in which there was no comparison group. However, in the secondary data analysis of a subset to the data presented in the present article, the removal of high catastrophizing scores led to a significant effective treatment for individuals who received a combination of amitriptyline and ketamine. These analyses are reported in a separate publication (26).

In spite of these limitations, the findings of the present study join growing literature showing that catastrophizing interferes with the effectiveness of analgesics. This is likely to have significant implications for the detection of treatment effects for the relief of pain. The present study supports the importance of including psychological measures in clinical trials, not only as important outcome measures, but also as determinants of treatment response (47). In the present study, the adverse influence of catastrophizing on treatment response was demonstrated using three different indexes of clinical significance. The effect sizes observed may be sufficient to obscure actual treatment effects, resulting in null findings in trials that do not control for the level of catastrophizing. Future research will need to examine the techniques that are most effective in reducing the pain focus of high catastrophizers. Future research will also need to examine whether interventions designed to specifically target catastrophic thinking influence treatment response to analgesic medication of neuropathic pain.

ACKNOWLEDGEMENTS: The authors thank Paulette Nauss for her assistance in data collection. The authors also thank Nathalie Gauthier and Isabelle Tremblay who completed data entry for this project. The work reported in this article is drawn from two clinical trials funded by EpiCep Corporation. This work was partially supported by a grant from the Canadian Institutes for Health Research.

REFERENCES


