CASE REPORT

Pain relief as a primary treatment goal: At what point does functioning and well-being become more important? A case study of an adolescent with debilitating chronic pain

Andrew S Tseng BA/BS1, Karen Weiss PhD2, Tracy Harrison MD2, Dan Hansen RN2, Barbara Bruce PhD2

BACKGROUND: Pediatric chronic pain is a common problem with significant economic implications and devastating consequences on quality of life. The present report describes a case involving a 15-year-old girl with severe and debilitating chronic pain.

RESULTS: Before her referral to a pain rehabilitation program, the patient saw numerous specialists who treated her with an aggressive medical regimen and two spinal cord stimulators. She was then referred for intensive interdisciplinary treatment and, after three weeks of rehabilitation, she reported clinically significant changes in anxiety, pain catastrophizing and functional disability. The patient was successfully titrated off all of her opioid medications and, eventually, both neurostimulator implants were removed.

DISCUSSION: Interdisciplinary pain rehabilitation is a useful treatment for patients with chronic pain. With its primary emphasis on functional restoration as opposed to strictly pain reduction, patients can regain a higher quality of life with reduced pain and fewer medications, surgeries and hospitalizations.

Key Words: Behavioural therapy; Chronic pain; Pain rehabilitation

Despite an abundance of research investigating treatment for chronic pain conditions, chronic pain continues to be a common problem in adult and pediatric populations. Patients with chronic pain often do not achieve substantial relief from a number of medical and nonmedical interventions, and their struggles with pain continue to have a significant effect on quality of life (1). Patients and providers go to great lengths to achieve pain reduction through the use of multiple medications, physical therapy, integrative medicine and medical procedures including surgeries. As a result of extended medical evaluation and treatment, delayed referral to interdisciplinary treatment, employment difficulties, disability and economic burden occur (2,3). Some patients will spend many years searching for additional explanations for the etiology of their pain, and treatments to decrease and/or resolve their pain (4,5).

It is well established that chronic pain is a biopsychosocial phenomenon (6). Nonetheless, many patients continue to receive only medical treatments for their chronic pain, without consideration for the psychosocial factors that are likely contributing to pain and disability. This sort of singular approach is not only minimally effective, but may also lead to increased disability and decreased psychological well-being.

Chronic pain may be particularly devastating for children and adolescents. Many pediatric patients with chronic pain experience reduced functioning and are believed to have a quality of life lower than that of cancer patients (4,7). For patients with chronic pain having no apparent physical etiology, the response to medication and therapy is often poor (8). Debilitating chronic pain in adolescents has been associated with family dysfunction and emotional struggles. Pediatric patients often withdraw from school for prolonged periods of time, during which they not only fall behind in academics but also miss important social interactions (1,7). This can lead to disruptions in development, particularly in the areas of social and emotional functioning.

The purpose of the present case study is to describe an adolescent girl who struggled with significant difficulties with chronic pain to highlight the negative consequences aggressive pain management treatment can have on a patient’s functioning and well-being. We also discuss the utility of nonpharmacological pain rehabilitation management and the importance of considering all aspects of a patient’s functioning, rather than pain relief being the only treatment outcome goal for pediatric pain patients.

CASE PRESENTATION

Presenting problem
Andrea (not patient’s real name) was a 15-year-old girl who had developed severe bilateral pain of her upper and lower extremities, as well as facial pain with transient erythema over 22 months, all of which had begun as pain and redness in her left second toe.
Developmental and medical history
Andrea grew up in a household with two parents and two younger siblings (a brother and a sister). Her father worked for the government and her mother worked from home. She described her family situation as being comfortable and supportive, with no history of abuse. Before the onset of her pain, Andrea had started public high school after being home-schooled for four years. She was an honors student who earned mostly As. She was involved in many activities, including piano, flute, guitar and soccer. She believed that she made friends easily and had a few close confidants who were very supportive of her.

Andrea was reportedly healthy until she began noticing redness and pain in her left second toe while playing soccer at the age of 14 years. Within days, she noticed further discolouration and swelling that worsened over the next three months. At that time, Andrea refused to wear shoes, even in the winter snow, because of this pain.

As part of standard clinical practice, Andrea completed several measures before and on discharge from the pain program, in addition to three and six months following the program. These measures included functional disability because this is the primary treatment goal of the pain program; pain catastrophizing, depression and anxiety because all of these variables can influence pain and disability; and pain intensity.

Andrea completed the Functional Disability Inventory (FDI) (9), a well-established, 15-item, self-report measure that assesses difficulty in physical and psychosocial functioning due to health status. On this form, she rated the difficulties in engaging in activities of daily living due to her pain. Subsequently, her mother became more involved in caring for these basic needs, including bathing her. This regression in autonomy caused some strain on their relationship, which was also distressing to Andrea. Despite the development of significant psychosocial concerns, Andrea was never referred to a psychologist or cognitive behavioural therapist.

Initial presentation and behavioural conceptualization
Andrea came to the Mayo Clinic (Rochester, Minnesota, USA) for an additional opinion for her multiple pain issues as well as urinary and defecation complaints, multiple food sensitivities and tachycardia. She saw providers in neurology, sleep medicine, dermatology, pediatric pain medicine, adolescent psychology and psychiatry, and underwent electromyography, polysomnography and autonomic reflex screening while at the institution. Numerous physicians became highly concerned with her extensive medication list, particularly given that she was frequently drowsy during visits with physicians and often fell asleep during interviews.

Andrea’s treatment team determined she was struggling with a chronic pain condition and given that multiple treatments had not provided long-term relief and her functioning was severely impaired, she was referred to the Pediatric Pain Rehabilitation Program at the Mayo Clinic.

Measurement of treatment outcomes
As part of standard clinical practice, Andrea completed several measures before and on discharge from the pain program, in addition to three and six months following the program. These measures included functional disability because this is the primary treatment goal of the pain program; pain catastrophizing, depression and anxiety because all of these variables can influence pain and disability; and pain intensity.

Andrea completed the Functional Disability Inventory (FDI) (9), a well-established, 15-item, self-report measure that assesses difficulty in physical and psychosocial functioning due to health status. On this form, she rated the difficulties in engaging in activities of daily living with regard to home, school and social tasks on a five-point scale ranging from 0 (no trouble) to 4 (impossible). FDI scores range between
Pain relief versus functionality

Andrea completed a numerical rating scale to indicate current pain intensity on a scale from 0 (indicating ‘no pain’) to 10 (indicating ‘worst pain imaginable’). Such rating scales are commonly used to assess pain intensity and have been shown to have good reliability and validity in pediatric populations (8,19).

RESULTS

Intensive outpatient pain rehabilitation

Andrea and her family participated in a three-week, hospital-based outpatient interdisciplinary pediatric pain rehabilitation program aimed at helping adolescents with chronic pain learn coping skills to manage chronic symptoms to improve functioning. The program included physical therapy, occupational therapy, recreational therapy, biofeedback, cognitive behavioural therapy and parent education. The program occurs in a group setting with approximately 12 adolescents and young adults between 13 and 21 years of age. In this program, there is a significant amount of time dedicated to support and education of parents with 10 h to 15 h of parent programming each week.

By the time she had entered the program, Andrea had titrated off some medications as recommended by her physicians. She was using a cane to walk and wore sandals that would not exacerbate her foot and toe pain. During intake assessments, although oriented to time and place, Andrea was fatigued and fell asleep multiple times during consultation. Her mood was dysphoric and affect was flat. Her thought processes were tangential and content was appropriate to session. Her speech was slow and slurred. Eye contact was lacking at times because she had difficulty keeping her eyes open. Andrea and her mother reported being very motivated to decrease medications and learn coping skills to manage pain and return to regular functioning.

Andrea and her treatment team identified major lifestyle functional goals she would work on during treatment: independent performance of daily cares, walking independently, relaxing independently, eating regular meals, sleeping restfully/waking independently, regular school attendance, socialization with peers and medication tapering.

Throughout the course of her rehabilitation, Andrea was self-directed and participated fully in all activities. However, before the start of medication tapering, Andrea expressed extreme anxiety. She was reluctant to stop many of her opioid medications for fear of pain exacerbation. She cried throughout consultations and was in clear emotional distress. However, she was gradually able to taper off all opioid medications including tapentadol, fentanyl patch and oxycodone, as well as sleep medications.

Her mental status gradually improved and she was more fully able to participate in daily programming and interactions with her peers. She worked at decreasing pain behaviours and cooperated with the treatment team. Within one week, she was able to walk without her cane. Initially, she had extreme difficulty with compliance with wearing socks and shoes due to allodynia. However, by the second week of therapy, she was able to wear shoes full time, in part as a result of the intensive physical therapy and desensitization that occurred on a daily basis as part of programming.

Andrea was asked to turn off her spinal cord stimulators for short periods of time and to use the coping strategies she was learning to manage any symptoms. Andrea complied, but experienced increased anxiety while doing so and was not able to discontinue use of the neurostimulators while in the three-week program. Nonetheless, she made drastic improvements in all five major lifestyle goals, and once again was able to function independently. She was encouraged to consider weaning off the neurostimulators at some point in the future when she believed her coping skills were more solidified.

In the parent program, Andrea’s mother learned strategies to be supportive of her daughter while encouraging developmentally appropriate autonomy. She was asked to ignore pain behaviours and discontinue talking about symptoms with her daughter. She also developed a behaviour plan to provide incentives for progress and decreased privileges when Andrea demonstrated inappropriate behaviours such as being disrespectful toward parents, or missing school or other activities. Andrea’s mother reported this aspect of the program was very helpful and she felt confident that she could return home and implement these strategies.

At the end of the program, Andrea reported clinically significant decreases in pain catastrophizing, anxiety and functional disability (Figure 1). At the six-month follow-up, she reported an absence of pain, pain catastrophizing and functional disability. Depression and anxiety symptoms were in the nonclinical range (Figure 1). She was also successfully titrated off all of her narcotic medications and most of her sleep medications by discharge and by three- and six-month follow-up (Table 1).

One-year follow-up

One year later, Andrea’s mother telephoned the program staff to report that Andrea had continued to do well once returning home from the Pediatric Pain Rehabilitation Program. Andrea was attending school full time and was involved in a variety of extracurricular activities.
including volleyball. She had returned to her normal weight. Eventually, Andrea decided the neurostimulators bothered her and decided to have them both removed, despite some hesitation from her medical providers at home due to concerns about relapse. Her mother reported that Andrea returned to her baseline level of functioning before pain onset and was now able to perform all of her activities of daily living.

**DISCUSSION**

The present case illustrates the devastating effects of chronic pain and how aggressive medical treatments may not provide adequate pain relief and may actually contribute to increased disability. The nature of Andrea's symptoms and her parents' desires to help their child led to intensive evaluation by multiple physicians to rule out the possibility of life-threatening disorders. When a specific diagnosis was eventually suggested, she attempted numerous medications and implantation of two spinal cord stimulators. Regardless of etiology, Andrea's pain developed into a chronic pain syndrome that involved severe disability. An interdisciplinary approach that included decreasing medications, physical reconditioning, and cognitive and behavioural therapy proved to be highly effective in decreasing anxiety, depression and disability for this patient.

**The search for a definitive diagnosis in the setting of chronic pain**

The present case highlights the inherent difficulties of treating patients with chronic pain. Patients with chronic pain may feel frustrated and hopeless without a definitive diagnosis, and this may fuel further preoccupation with their pain. Patients often feel dissatisfied and/or invalided when informed that multiple tests have not revealed a cause for their pain. They may then feel misunderstood and continue to search relentlessly for an answer and satisfactory treatment (20). They often believe providers are suggesting that their pain is 'not real' or is psychological in nature. In addition, because of the physical disability and depression that are often comorbid with chronic pain, patients and caretakers may often be more desperate for an answer to resolve these issues. Thus, one of the greatest dangers occurs when desperate patients are driven to pursue drastic therapies in attempts to eliminate their pain.

For Andrea's parents, their well-intentioned concerns and their desire to find a cure for their daughter's pain led them on a time-consuming journey through doctors' offices and operating rooms. In their situation, it may have been reasonable to seek a second opinion, especially when the diagnosis was inconclusive and their daughter continued to experience significant pain. Their persistence may appear to have paid off when well-meaning physicians prescribed a new intervention that may have worked temporarily, only to relapse back into the cycle of continued pain and poor functioning. When one therapy did not work, they sought yet another treatment from another provider. In the end, these aggressive and experimental treatments accumulated and proved to be deleterious.

**Pain reduction versus functioning and well-being**

Many health care providers may define success in pain management as a reduction in pain score. Although this is certainly an appropriate target in the treatment of acute pain, targets for the treatment of chronic pain should also include improvement in functioning and overall well-being because complete pain relief may not be an attainable goal. More importantly, consuming preoccupation with pain relief may lead to years of searching for the next best treatment and avoidance of regular activities. Unfortunately, patients who decide to be home-schooled and stop activities that exacerbate their pain often experience increased pain and worsened functioning (7).

In the case of Andrea, it was likely the combination of her distress about her pain, her parents' concern about her well-being and her health care providers' desire to help her feel better that led to multiple medications that impacted her ability to function from a cognitive standpoint. The continued progression of her disability then led to unhealthy changes in the family dynamic, difficulty attending school, decreased participation in preferred activities and eventual isolation from same-age peers. All of this then contributed to Andrea struggling with self-esteem issues (self-efficacy about being able to perform most normal daily tasks), depression and suicidal ideation. By focusing primarily on the treatment of her pain, her health care providers may not have realized the negative effects this treatment was having on her overall functioning. Her treatment course appeared to have contributed to the transformation of a healthy and active 15-year-old girl into a minimally responsive, depressed and dysfunctional patient.

**Cognitive behavioural therapy, interdisciplinary approaches, and functional restoration**

Cognitive behavioural therapy and interdisciplinary approaches are the most widely accepted biopsychosocial treatments for chronic pain (21,22). These treatments often do not focus on elimination of pain but on functional restoration (23). When compared with opioid and other medical treatments, cognitive behavioural therapy and interdisciplinary approaches not only yield comparable pain reduction rates, but are also more cost-effective and lead to significant increases in physical functionality, and reductions in future hospitalizations and surgeries (22,24–27). In the pediatric population, it is particularly important to minimize this disruption in development by allowing children to return to school and to hopefully begin the process of normalizing their lives (28).

Given the utility of the biopsychosocial conceptualization and approach to chronic pain, we recommend health care providers consider interdisciplinary screening at initial assessment for all providers with chronic pain. In addition, at any point in the treatment of chronic pain where functioning starts to be adversely affected, an interdisciplinary treatment would likely be helpful. Moreover, patients tend to be more accepting of such an approach when providers emphasize that reported symptoms are real, but chronic pain has a complex etiology that requires treatment of the problem(s) from multiple angles.

Andrea's story highlights the importance of consideration of all aspects of a patient's well-being, including involvement in typical activities (eg, school), social interaction, family relationships and psychological well-being. Attempts at pain reduction should not be made without considering the possible effects on these other areas of functioning. Notably, in Andrea's case, we observed an improvement in functioning before an improvement in pain intensity. It is also important to note that an improvement in functioning without any improvement in pain intensity would have still been considered a treatment success.

**CONCLUSION**

Although Andrea's case is an extreme example, her experiences of spending years seeking a satisfactory medical diagnosis and treatment to eliminate pain, and progressive disability in multiple areas of functioning mirror that of many adolescents who experience chronic pain. Fortunately, Andrea was ultimately referred to an interdisciplinary treatment program that encouraged her to discontinue medications causing side effects and helped her and her family learn and implement strategies of increased coping in the context of pain and reconceptualizing the treatment goal to be fully functioning. For the millions of individuals experiencing chronic pain, it is imperative to have discussions about comprehensive approaches to pain management and the importance of including quality of life and overall functioning as treatment goals, which are just as important, if not more so, as alleviating the pain itself.

**DISCLOSURES:** None.
REFERENCES


