The role of the cervical spine in post-concussion syndrome

Cameron M. Marshall, Howard Vernon, John J. Leddy and Bradley A. Baldwin

Introduction

Concussion injuries, or mild traumatic brain injury (mTBI), have an estimated prevalence of 3.8 million per year in the United States [1] and are considered one of the least understood injuries facing the sports medicine and neuroscience communities today [2]. In the majority of cases, concussion symptoms resolve within 7–10 days; however, ~10–15% of these patients develop persistent symptomatology lasting weeks, months or even years after injury [3-5]. Depending on the classification system used, post-concussion syndrome (PCS) is defined as the persistence of three or more symptoms for 4 weeks (International Classification of Diseases, Tenth Revision), or 3 months (Diagnostic and Statistical Manual of Mental Disorders), following a minor head injury [6-9].

The pathophysiology of mTBI during the acute phase of injury is becoming increasingly well-known suggesting ion imbalance, metabolic disruptions, blood flow abnormalities and autonomic nervous system (ANS) dysfunction as the main culprits [6,10-14]. When attempting to search for potential mechanisms surrounding PCS; however, findings have been inconsistent and there remains no clear physiologic explanation for this disorder [9,15]. In fact, most studies examining the physiologic dysfunctions found during the acute phase of concussion have demonstrated a return to baseline or control levels within 2–4 weeks following injury [6,12,13,16,17]. This has led to great debate within the medical community as to whether PCS is mainly a physiological or psychological manifestation following concussion injuries.

One proposed mechanism for persisting symptomatology that has not been examined in great detail is concomitant low-grade sprain–strain injury of the cervical spine occurring concurrently with significant head trauma. Based on the obvious anatomical association between head and neck, it is reasonable to expect that any significant blunt impact and/or acceleration/ deceleration of the head will also result in some degree of inertial loading of the neck potentially resulting in strain injuries to the soft tissues and joints of the cervical spine. Studies in high school and college football players using the Head Impact Telemetry system have demonstrated that the range of linear impact accelerations causing concussion injury is between 60 and 160 g (where ‘g’ represents gravity), with the highest predictive occurrence occurring at 96.1 g [18]. Studies examining whiplash associated disorders (WAD) have demonstrated two very important features for our discussion: 1) biomechanical studies have demonstrated that as little as 4.5 g of neck acceleration can cause mild strain injury to the tissues of the cervical spine [19] and 2) the signs and symptoms reported by these patients, with the exception of a few key differences (i.e. radicular symptoms), appear strikingly similar to those experienced in mTBI (Table 1) [20-28].

If an individual is to sustain an injury where the head is accelerated between 60 and 160 g, it is highly likely that the tissues of the cervical spine have also reached their injury threshold of 4.5 g. A 2006 study from the University of Guelph examined junior ‘A’ hockey players immediately following injuries that either appeared as whiplash mechanisms (i.e. a blow to the body with a whipping motion of the neck...
and head), or injuries that appeared as concussion mechanisms (i.e. a direct blow to the head). It was discovered that 100% of the injured athletes had signs and symptoms of both WAD and concussion indicating that these injuries are happening concurrently [29].

Injury or dysfunction of the cervical spine has been shown to cause headaches [30,31], dizziness and loss of balance [32-34], nausea, visual and auditory disturbances [35], reduced cognitive function [19,26], and many other signs and symptoms considered synonymous with concussion. Interestingly, the limited studies involving functional imaging of the brain in these patients do not demonstrate any significant findings, which could indicate either psychological overlay or at risk cervicogenic etiology [36,37].

This paper presents a review of the literature surrounding PCS, as well as cervical spine dysfunction to propose a theoretical hypothesis for mild cervical strain injury, suffered concurrently at the time of the concussion, acting as a major symptomatic culprit in many PCS patients. This paper also discusses the management of five PCS patients using both active rehabilitation and passive manual therapy of the cervical spine resulting in a significant reduction in symptoms as well as a significant improvement in function.

### Pathophysiology of concussion and post-concussion syndrome

During concussive injury to the brain, previous literature demonstrates that neurons undergo stretch and shear forces that cause brief mechanical disruption of the cell structure, which leads to an indiscriminate exchange of ions, potassium ($K^+$) efflux, glutamate release and subsequent mitochondrial calcium ($Ca^{2+}$) overload resulting in impaired oxidative metabolism and reduced ATP production. In an effort to restore proper ion balance within the cell, the ATP pump is activated to a greater capacity requiring large amounts of ATP. The imbalance between increased ATP requirements, decreased ATP production and reduced cerebral blood flow quickly leads to an energy crisis within the brain known as spreading depression (Figure 1) [10,14].

It is believed that the symptoms of concussion are due to reversible neuronal dysfunction and energy mismatch and not due to structural damage of the involved neurons, which is why conventional structural imaging techniques such as CT and MRI are typically unremarkable [3].

The vast majority of mTBI patients experience a resolution of symptoms within 7–10 days following injury [3]. There are a number of proposed reasons as to why roughly 10–15% of patients go on to experience longer lasting symptomatology, including both physiological and psychological etiology. Continued reductions of ATP, continued axonal dysfunction,
altered blood flow, continued ANS dysfunction have all been suggested as possible physiological alterations accompanying PCS; however, evidence surrounding each of these mechanisms is limited.

Theory 1: Continued metabolic dysfunction

Animal studies have demonstrated reductions in ATP as early as 1-min following impacts equivalent to concussion [38,39]. Unfortunately, ATP levels cannot be measured directly in vivo and therefore culling and processing must take place prior to direct ATP measurement. Proton magnetic resonance spectroscopy (MRS-H), however, is able to measure the concentration of metabolite levels, in particular N-acetyl aspartate (NAA), a high correlate of ATP levels \( R^2 = 0.84 \) [17,39], within the brain of human subjects.

Human spectroscopy studies have demonstrated significant decreases in NAA: creatine and NAA: choline ratios in the acute phase of concussion; however, most return to non-injured control levels by 30 days post-injury (Figure 2) [13,40-43]. All subjects in these studies were asymptomatic within 8 days following injury; however, metabolite levels did not normalize until day 30. This indicates the likelihood of a weak correlation between symptoms and metabolic dysfunction.

One conflicting study came from Henry et al., in 2011 [44] where NAA: creatine ratios were still significantly reduced at 6 months post-injury in a group of 10 concussed football players as compared to an uninjured control group. Again, all concussed subjects were asymptomatic by the 6-month follow-up and had returned to full play. The authors posited that demands placed on the university athlete (i.e., course load coupled with intense practice and game schedules) may have been a hindrance to complete metabolic recovery [44].

The evidence of continued metabolic dysfunction in subjects diagnosed with PCS is limited [45] as most MRS-H studies have focused on the acute stages of injury or those with chronic impairment following more severe brain injuries. There have been two studies that have demonstrated metabolic reductions in symptomatic PCS patients; one in adult PCS patients [46] as well as pediatric PCS patients [47].

As previously mentioned, most studies demonstrate a return of concussion-related metabolites to control levels by 30 days following injury; however, suffering a second concussion during this metabolically depressed state has been shown to increase the time to NAA normalization in both animal and human subjects. In two human studies, symptomatic recovery took between 3 and 8 days following a single concussion. Following a second injury, suffered prior to NAA normalization, subjects reported significantly increased symptom durations, lasting between 24 and 59 days. A substantial delay in NAA normalization was also found in these doubly concussed athletes; ranging from 60 to 120 days post-injury [41,48]. These two studies not only highlight the importance of achieving full recovery before returning to sporting activity but also provide a potential physiologic basis for the increased propensity of PCS in patients with a previous concussion history.

The literature presented above demonstrates limited evidence for a metabolic justification for persistent concussion symptoms following a single concussive episode. Experiencing a second concussion prior to full metabolic recovery after the first has demonstrated persistent symptomatology and metabolic dysfunction in two separate case-series. Interestingly, in each of these cases (nine total), symptomatic recovery long preceded metabolic recovery, which leads one to question whether symptom recovery and metabolic recovery are closely related.

Figure 2. Cerebral metabolite ratios in controls versus sports concussion patients. Bar graph showing the metabolite ratios of N-acetyl aspartate/choline-containing (NAA/Cho) compounds, NAA/creatine-containing (NAA/Cr) compounds and choline-containing compounds/creatine-containing compounds (Cho/Cr) in controls and concussed patients. Each histogram is the mean value determined in 30 healthy controls and 40 concussed athletes in the study by Vagnozzi et al., 2010. At 3-days post-injury NAA/Cr is reduced by 17.6%, NAA/Cho is reduced by 21.4% with both gradually recovering to complete normalization at 30 days. The Cho/Cr ratio did not show any significant variation.

\( ^a p < 0.01 \) with respect to controls.

\( ^b p < 0.01 \) with respect to values determined at the previous time points.

Figure reprinted from [13] with permission from Oxford University Press.
Theory 2: Continued axonal dysfunction

Diffusion tensor imaging (DTI), an advanced imaging technique that assesses the integrity of white matter tracts by measuring the rate of water diffusion along, and perpendicular to, axonal fibers [49-51], has increased the understanding of concussion injuries substantially in recent years. The neuronal shearing that occurs with impact as well as the subsequent ion imbalance and Ca⁺ overload, can alter the cellular environment and brain function due to suggested myelin loss, microscopic lesions and axonal degeneration [49,52,53]. The three main measures of interest in DTI are fractional anisotropy (FA), mean diffusivity (MD) and axial diffusivity. Studies examining acutely concussed patients have found significant decreases in FA and significant increases in MD compared to controls [51,52], particularly in the posterior corpus callosum according to a recent meta-analysis [49]. On the contrary, other studies have shown increases in FA and decreases in radial diffusivity and MD [51,54] in almost equal numbers [55]. This inverse relationship between FA and MD is thought to represent damage to the microfilaments and microtubules of the axon [56]. However, like much of the research surrounding concussion injuries, as the scanning takes place further away from the initial injury most of the studies have shown variable results possibly attributed to inconsistent measurement methodology [49,53] or potential flaws within the measurement itself [55].

There have been a number of studies examining DTI in PCS patients in recent years [50,56-59], with mixed results when compared to uninjured controls. Comparing patients with ongoing symptomatology with uninjured, matched control groups is problematic for a number of reasons, especially when trying to elucidate the source of post-concussion symptoms. Lange et al., in one of the largest and most recent DTI PCS studies, examined 60 patients with recent mTBI (both asymptomatic and symptomatic patients) and compared them to 34 patients with orthopedic/soft tissue injuries (excluding cervical spine) between 6 and 8 months post-injury. Mild TBI subjects also completed a post-concussion symptom checklist and, based-upon the presence of three or more symptoms between 6 and 8 months post-injury, were classified as either being post-concussional disorder (PCD) present (n = 21), or PCD absent (n = 39). FA and MD values for various regions of the corpus callosum showed differences in all recent mTBI patients versus the orthopedic injury group; however, there were no significant differences found between those subjects classified as PCD present and those classified as PCD absent. The authors concluded that their data does not support an association between white matter integrity in the corpus callosum and post-concussion symptoms 6–8 months post-injury [60]. Recent research has also suggested that being involved in contact sports creates significant changes in various DTI parameters over the course of a season in athletes who have not suffered diagnosable injury [54,61,62].

In summary, despite early indications that microstructural brain injury can be evaluated using DTI, this imaging modality cannot currently predict PCS symptoms, clinical outcome and/or clinical recovery in individual subjects [51]. Furthermore, as the imaging takes place further from the time of the injury, results become less consistent and may not be associated with symptoms of PCS, but rather with a history of recent mTBI and/or involvement in contact sports.

Theory 3: Psychological factors

The diagnosis of PCS is complicated due to multiple symptoms that represent a myriad potential differential diagnoses with a large degree of psychological overlay. Among others, PCS has been shown to have similar presentations as depression, somatization and chronic pain [6]. It has also been suggested by numerous authors that preexisting psychological disturbances such as depression and anxiety increase the risk of developing PCS [3,63]. Although there is extensive research in this area, it is beyond the scope of this paper to provide a detailed review of this literature.

Theory 4: Altered cerebral blood flow

A common symptom observed in concussed athletes is the onset of post-concussion-like symptoms, particularly headache, when there is an increase in cardiovascular output during exercise [15]. One of the major hypotheses suggests that this is due to continued reductions in cerebral blood flow and oxygenation [6]. Reductions in cerebral blood flow have been found immediately following impact [10]. Other studies have found both increases as well as no change when examined within a day or two following injury [64,65]. Studies examining mTBI patients in the acute phase have also found ANS dysfunction such as altered cerebrovascular reactivity [11,66,67], cerebral autoregulation, cerebral oxygenation and heart rate variability [68] when compared to uninjured controls; however, these findings seem to normalize within 4–14 days post-injury [12,67,68]. Research surrounding these phenomena in PCS patients is, however, somewhat limited. To the authors’ knowledge there have been no studies to date that have demonstrated similar autonomic dysfunctions in PCS subjects.

Single positron emission computed tomography (SPECT) is a resting state functional imaging technique used to examine cerebral blood flow using radioactive tracers, most commonly technetium-99 m-hexamethylpropyleneamine oxime (⁹⁹Tc-HMPAO) [45]. Studies using ⁹⁹Tc-HMPAO SPECT in heterogeneous populations of chronic mTBI patients have demonstrated that between 40 and 60% demonstrate significant hypoperfusion in various regions of the brain. The most common areas cited for reduced blood flow in these studies are the frontal and parietal lobes [69-72]. At first glance, hypoperfusion of various cerebral levels may seem like a plausible explanation for post-concussion symptomatology and dysfunction; however, studies examining other entities such as chronic neck and upper thoracic pain [73], late-whiplash [23,74], acute and chronic low back pain [75], obsessive-compulsive disorder [76], depression [77], chronic fatigue syndrome, AIDS dementia complex [78] and many others have all demonstrated similar findings on SPECT. Furthermore, many chronic mTBI SPECT studies have not been able to...
demonstrate consistent correlation with neuropsychological test results or post-concussion symptoms [45]. At this point, it seems that although cerebral hypoperfusion may be present in roughly half of PCS patients, this does not represent a specific etiology of chronic mTBI but rather a complex, non-specific finding that seems just as likely to appear in various chronic pain, anxiety and depression patients as it does in those diagnosed with chronic mTBI/PCS.

Case for a likely cervicogenic component to concussion injuries

Aside from the direct biomechanical associations and injury mechanics mentioned in the Introduction, the symptoms of headache and dizziness, so prevalent in concussion-type injuries, may actually be the result of cervicogenic mechanisms due to a concomitant whiplash injury suffered at the same time [22,29,79,80]. Two primary categories of mechanisms can be proposed: pain-related and proprioceptive-related. Pain-related mechanisms predominate in the development of headaches, while proprioceptive mechanisms predominate in cervicogenic dizziness and oculomotor control deficits.

Studies of the primary afferent connections of neurons in the C2 dorsal root ganglion [81] indicate that the following structures receive mono-synaptic inputs from these afferents:

- Lateral cervical nucleus
- Central cervical nucleus
- Caudal projections to C5 level
- Cuneate nucleus, lateral cuneate nucleus
- Nucleus tractus solitarius
- Intercalatus nucleus
- Nucleus X of the vestibular system

‘Upper cervical afferents’ are clearly capable of evoking activity in the vestibular system.

Richmond [81] reported that afferents from deep spinal structures such as those likely to be injured in cervical acceleration injuries terminate in deeper laminae in the dorsal horn than do cutaneous afferents. It is well known that afferents from the C2 and C3 dorsal root ganglia terminate on the same second-order dorsal horn neurons as do many nociceptive afferents arising from the trigeminal sensory nucleus, especially those in the subnucleus caudalis of the descending tract of the trigeminal nerve [82-85]. In fact, Bogduk has termed this continuous column of cells the ‘trigemino-cervical nucleus,’ while Gobel has called it the ‘medullary dorsal horn’ [86]. This critical convergence phenomenon provides a mechanism for referred pain from upper cervical structures (from C0 to at least C3) to the upper cranium and forehead, mimicking headache.

Human studies of pain provocation [87-89] and pain abolition [90,91] have replicated these patterns of cranial pain referral, which can so easily be described by patients as headache. Animal studies of pain provocation in the upper cervical spine [92-95] as well as lower cervical muscles [96,97] report activations of EMG in the cranio-cervical muscles as well as findings indicative of central sensitization, all of which could underlie the development of headache arising from injury to the deep tissues of the upper cervical spine.

The concept of ‘cervicogenic vertigo’ is less established in the mainstream neurologic literature. Mechanisms underlying this phenomenon rely on both monosynaptic and polysynaptic reflex pathways from the upper cervical spine afferents (associated with a rich innervation from joint and muscle proprioceptors in the cervical spine) to the brainstem structures associated with balance.

Both the cervico-collic and cervico-ocular reflexes work in conjunction with the vestibulo-ocular reflex to stabilize the head and trunk during head/neck motions (CCR) and to stabilize gaze during head rotations (COR). Disturbances in these reflexes can, therefore, initiate balance disturbances and symptoms associated with this problem, particularly those associated with disturbances of eye gaze control [98-100].

Investigating the connection between cervical spine injury and balance disturbances has included studies of increased postural sway and impaired head repositioning capability in whiplash patients [24,99-102] as well as numerous clinical reports of relief of vertigo-like symptoms following treatments to the neck in post-concussion or post-traumatic headache (PCH) patients [103-105]. However, direct evidence of such a link is somewhat scanty. Aside from the work of De Jong et al., mentioned above [106], Vuillerme and Pinsault [107] induced balance disturbances in 16 healthy males after experimental upper cervical pain. Additionally, studies employing vibration of the cervical muscles, presumably to activate muscle spindle output, and, especially in a unilateral fashion, have shown induced postural imbalance and dizziness in normal subjects [108,109].

Remarkably, only two studies have been found that directly examine the presence of cervical dysfunction findings in PCS patients who were involved in relatively low-energy injuries. Treleaven et al. [79] studied 12 post-concussion subjects with PCH and found that the PCH group was distinguished from the normal control group by the presence of painful upper cervical segmental joint dysfunction, less endurance in the neck flexor muscles and a higher incidence of moderately tight neck musculature. They concluded that upper cervical joint dysfunction, which is precisely the disturbance referred to above, should be assessed for in post-concussion headache. Second, a treatment study conducted by Jensen et al. [110], examined 23 patients with persistent PCH an average of 359 days post-concussion (range 302–423 days). The authors found that 18 of the 19 patients available for follow-up had painful upper cervical joint restrictions when compared to an uninjured control group (n = 19). The researchers then randomly assigned the PCH group to one of two treatment groups: the manual therapy group, which received mobilizations of the cervical spine as well as a form of assisted muscle stretching, or the cold-pack group, in which subjects were put on an ice pack under their neck and shoulders for 15–20 min. After only two visits, spaced 1 week apart, the manual therapy group had a 57% reduction of their pre-injury pain scores whereas the cold-pack group showed no reduction in headache frequency or intensity. Interestingly, the manual therapy group also had a reduction in analgesic use as well as a 52% reduction in dizziness and visual
disturbance ratings. A more recent randomized controlled trial from the University of Calgary highlighted very similar findings in which weekly treatments focusing on cervical spine and vestibular rehab in subacute and chronic concussion patients resulted in 73% of athletes becoming asymptomatic and cleared to return to sport versus only 7% in a group that did not receive this therapy [111].

Cases

The following is a case-series of five patients who presented to one of our Canadian multidisciplinary concussion clinics with persistent symptoms following a diagnosed concussion sustained anywhere from 5 weeks to 31 months prior to presentation. All subjects underwent concussion evaluation, full neurological examination, including cranial nerve screen, cerebellar testing, as well as upper and lower limb neurological examination. All neurological examinations were unremarkable, aside from one patient (patient 3) who complained of persistent tinnitus and showed conductive hearing loss on the right side. Subsequent referral to an otolaryngologist revealed no abnormalities in this patient. A summary of these patient cases can be found below (Table 2).

Case 1

A 25-year-old male professional mixed martial arts athlete presented 4 months after injury, which was sustained while sparring during a training session. The athlete was thrown to the ground striking the posterior aspect of his head on the mat. Symptoms of headache, dizziness and nausea persisted for roughly 1 week following injury and he attempted to return to training; however, was unable to do so due to symptom exacerbation and was subsequently referred to a neurologist. The patient saw two separate neurologists and was put on absolute physical and cognitive rest following each of his three follow-up visits spaced 1 month apart. MRI examinations were unremarkable. This patient had a history of four previous concussions; however, all had resolved completely within 1 week. Upon presentation to our clinic the patient complained of numerous post-concussion symptoms the most notable of which were daily headaches, dizziness, noise sensitivity, and an alternating cold and hot sensation in the posterior aspect of the head. The headaches were localized to the frontal area of the forehead as well as bilaterally in the temporal regions.

The patient was put through the Balke physical exertion protocol to assess for blood flow abnormalities according to Leddy et al., 2010 and 2011, and Baker et al., 2012 [112-114]. The patient was able to fully complete the test with no exacerbation of symptoms and was subsequently permitted to return to non-contact exercise and training. Examination of the cervical spine revealed a painful and restricted C2/3 joint on the left side with no referral to the cranium. Palpation of the longus colli, suboccipital triangle, sternocleidomastoid (SCM) and splenius capitis, however, all created referral into the head in the same pattern as the daily headaches experienced by the patient.

A course of treatment was initiated consisting of passive soft tissue therapy using Active Release Therapy (ART®) and localized vibration therapy over the affected muscles, as well as active deep-neck flexor endurance training utilizing a

Table 2. Summary chart of patient cases presenting with persistent post-concussion symptoms.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age, Sex</th>
<th>MOI</th>
<th>Time (weeks)</th>
<th>Symptoms</th>
<th>Balke</th>
<th>Treatment</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>25, m</td>
<td>Sparring – contact with posterior head on ground</td>
<td>16</td>
<td>Headache, dizziness, noise sensitivity</td>
<td>Pass</td>
<td>ART® of affected muscles, SMT of cervical spine, LV, DNFEE</td>
<td>- 3 tx significant reduction in symptoms</td>
</tr>
<tr>
<td>2</td>
<td>59, f</td>
<td>Slip and fall – contact with posterior head on ground</td>
<td>124</td>
<td>Headache, dizziness, anxiety</td>
<td>Pass</td>
<td>ART®, Mobilizations of cervical spine, DNFEE, Vestibular rehab with head and eye movement included</td>
<td>- 8 tx full resolution of all symptoms</td>
</tr>
<tr>
<td>3</td>
<td>19, m</td>
<td>Hockey – check to head</td>
<td>96</td>
<td>Headache, dizziness, visual problems, fatigue, sensitivity to light, mental fogging, concentration difficulties, irritability, memory difficulties</td>
<td>Pass</td>
<td>ART®, SMT of cervical spine, eventual referral to performance optometrist for co-management</td>
<td>- 3 months of treatment 2x/week showed significant subjective improvement</td>
</tr>
<tr>
<td>4</td>
<td>19, m</td>
<td>Car accident</td>
<td>14</td>
<td>Headache, irritability, sleep disturbances, visual difficulties, concentration problems, neck pain</td>
<td>Pass</td>
<td>ART®, SMT cervical spine, DNFEE, Visual tracking exercises, eventual referral to performance optometrist</td>
<td>- patient still seeking ongoing care on a monthly basis due to fear of aggravation</td>
</tr>
<tr>
<td>5</td>
<td>51, f</td>
<td>Hockey – blow to left side of head (temporal region)</td>
<td>5</td>
<td>Daily headache, neck pain</td>
<td>Not done</td>
<td>ART®, SMT cervical spine, home-based ROM exercises</td>
<td>- Case ongoing</td>
</tr>
</tbody>
</table>

Abbreviations: ART® = Active release therapy; DNFEE = Deep neck flexor endurance exercises; LV = Localized vibration therapy; m = Male; MOI = Mechanism of injury; ROM = Range of motion; SMT = Spinal manipulative therapy (High-velocity, low-amplitude); tx = Treatments.
Cervical spine and post-concussion syndrome

A 59-year-old female university counselor presented with persistent symptoms of 31 months duration following a slip-and-fall injury where she hit the posterior aspect of her head on the frozen ground. Her initial symptoms at the time of injury were being dazed and having a headache. Over the next 3 weeks her symptoms progressed and she was referred to a neurologist as well as a neurosurgeon. CT and MRI evaluations were unremarkable aside from a congenital fusion visualized at the C2–C3 level of the cervical spine. She was advised by the neurosurgeon to go on complete mental and physical rest. At each monthly follow-up she was given the same recommendations, which amounted to 9 months of lost work with brief periods of failed attempts at gradual workplace re-integration.

When she presented to our clinic she was finally back to work but was still finding it very difficult with her ongoing headaches and dizziness. Her main complaint of headache was described as ‘someone driving a screwdriver right through the top of my head,’ which increased as the day progressed. The patient was also very anxious and concerned for her condition believing that there was no hope for recovery. All neurological examinations were unremarkable and the patient was put through the Balke physical exertion test. Even though the patient had not exercised since her injury on recommendations from her previous physicians, she was able to pass the Balke treadmill reaching a peak heart rate of 180 beats per minute with no exacerbation of her symptoms.

Examination of her cervical spine revealed several painful zygapophyseal joint restrictions from C0/C1 down to the C4 level bilaterally. Palpation of the soft tissues revealed several painful trigger points, which caused referred pain into the head the worst of which was splenius capitis on the left side. Splenius capitis palpation recreated the screwdriver sensation at the top of her head. Vestibular testing revealed a negative Dix–Hallpike test for benign paroxysmal positional vertigo and a positive rotatory chair test for cervicogenic vertigo. Further vestibular testing revealed balance impairments when eye movements and neck and head rotations were incorporated. Passive and active treatment of the cervical spine were initiated and consisted of a similar treatment methodology and schedule as seen in Case 1 with the exception of high-velocity low-amplitude cervical spine manipulation as the patient did not wish to have this type of treatment. Low-velocity mobilizations were therefore used. The patient reported excellent symptomatic relief for 2–3 days following each treatment. Additional vestibular rehabilitation incorporating visual motion and head and neck movements during various balance progressions was also utilized.

Treatment visits began on a weekly basis, however, the patient elected to increase frequency to twice per week as they were very effective in significantly reducing her symptoms. At 3 months following the initiation from treatment the patient had continued working, was driving with no issue, and exercising with a personal trainer. Although the patient is currently asymptomatic and has returned to all activities of daily living, she elected to continue with therapy once per month due to personal fear of symptomatic return.
6/6 to 1/6, irritability reduced from 5/6 to a 0/6, and light sensitivity reduced from 3/6 to a 0/6. At this point, he still complained of some mild visual difficulties particularly with reading and watching television. He was subsequently referred to a performance vision optometrist for co-management of his visual symptoms. At the time of the writing of this paper, his care was still ongoing having only had four visits. This case highlights that some patients with persistent symptoms can have significant improvements in their PCS through various active and passive treatments of the cervical spine in a minimal amount of time. This particular patient had been highly symptomatic for a 2-year period and within 21 days after presentation he had an 80% reduction in his post-concussion symptom severity score.

Case 4
This 19-year-old male junior hockey player presented 14 weeks after sustaining a concussion in a car accident in which he rolled a small convertible several times. CT imaging at the hospital following the injury was unremarkable. Main symptom complaints on presentation were frontal headaches, visual problems, concentration difficulties, neck pain, irritability, emotional lability and sleep difficulties including taking up to 4 h to fall asleep at night. His PCSS at presentation demonstrated 9/22 total symptoms with an overall severity score of 36/132. At the time of presentation, the patient was out of school and sports for two full months on the advice of his family physician. Neurological examination was unremarkable; however, he had some difficulties with visual smooth pursuit, demonstrating saccadic eye movement. On treadmill testing, the patient was able to complete the test with a heart rate of 167 beats per minute with no exacerbation of symptoms.

Examination of the cervical spine revealed several trigger points in splenius capitis, longus colli, suboccipital triangle and the SCM all of which created referral into the head particularly the suboccipitals and SCM bilaterally. He also had painful joint restrictions in the C0/C1, C2/3 and C3/4 facet joints bilaterally.

Treatment of the cervical spine was initiated consisting of ART®, SMT, as well as deep neck flexor endurance and visual tracking exercises done at home. After the first treatment, the patient reported significant relief of his symptoms. At the beginning of the second visit, the patient’s headache score had dropped from 3/6 to 1/6. His total PCSS showed a total of 6 out of a possible 22 symptoms with a total severity score of 19, representing a 53% reduction in symptom severity following only one treatment. Two more treatments were provided the following week and on the fourth visit the patient reported a PCSS of 3/22 with a total severity score of 6/132 and a headache score of 0/6. The patient was also participating in non-contact sport-specific activity (skating) 3–4 h per day, had enrolled in an online adult education course, and reported that he was sleeping better at night. On the seventh visit, the patient reported a PCSS of 2 out of 22 with a total symptom severity score of 2, including a neck pain score of 1/6 as well as a 1/6 rating for blurred vision. At this point, the patient was referred to a performance vision optometrist for co-management. By the eighth visit, the patient reported that he was doing his visual exercises and that he was already feeling much better, citing a full resolution of all symptoms. He was subsequently put through the Chicago Blackhawks physical exertion testing program and passed with no exacerbation of any of his symptoms and thereby permitted to return to contact sport.

Case 5
This 51-year-old female human resources manager presented 5-weeks after sustaining a concussion while playing in a recreational hockey game in which she collided with an opposing player striking the left side of her head. The patient had no previous history of concussion and her ongoing symptoms were constant daily headaches on the left temporal region and left-sided neck pain on left rotation. All neurological testing was unremarkable. Due to the presence of neck pain, this patient underwent cervical spine examination without first undergoing the Balke treadmill test, which revealed a very painful facet joint restriction at C3/4 on the left. Soft tissue palpation revealed several trigger points in the left suboccipital triangle, longus colli and the SCM; all of which, referred pain into the left temporal region, recreating her headache. Treatment of the cervical spine was initiated including ART® to the affected soft tissue trigger points, spinal manipulation to the painful and restricted joints, as well as deep neck flexor endurance exercises in the form of seated chin-tucks performed at home. Thirty-minute treatments were initiated on a weekly basis and after the third treatment the patient was no longer experiencing any headaches but still had some mild neck stiffness from time to time. The patient was given further range of motion exercises and was scheduled for a follow-up 3 weeks later at which she reported that her symptoms of both headaches and neck pain had completely resolved. The patient was subsequently discharged and permitted to resume all activity.

Conclusion
The acute symptoms of concussion are widely considered to be the result of functional neuronal disturbance and an altered cerebral environment however; the etiology of chronic post-concussion symptoms remains unclear. The pathophysiological processes involved in the acute neurometabolic cascade of concussion have been shown by the majority of studies to resolve within a 30-day period. As previously discussed, diffusion tensor examinations of PCS patients also provides inconsistent evidence for neuronal dysfunction as a significant underlying cause of chronic symptoms. Studies utilizing SPECT have found hypoperfusion in various brain regions for up to 5 years following concussion, however, these do not seem specific to concussion injuries, as similar findings are present in myriad physical and psychological conditions. The above findings demonstrate that although the underlying pathophysiology of acute concussion is becoming increasingly well understood, the same cannot be said for the symptoms experienced in PCS.
Based on previously established tissue injury thresholds, acceleration/deceleration of the head–neck complex of sufficient magnitude to cause mTBI is also likely to cause concurrent injury to the joints and soft tissues of the cervical spine. It has also been well established that injury and/or dysfunction of the cervical spine can result in numerous signs and symptoms synonymous with concussion, including headaches, dizziness, as well as cognitive and visual dysfunction; making diagnosis difficult. Due the strong personal, social and economic burdens posed by PCS, we strongly suggest that future research be conducted to further examine the relationship of cervical spine injury contributing to symptomatology in these patients. It is important not only to improve diagnostic procedures but also study the outcome of treatment aimed at ameliorating cervical spine dysfunction in a randomized and controlled fashion.

In conclusion, management of persistent PCS symptoms through ongoing brain rest is outdated and demonstrates limited evidence of effectiveness in these patients. Instead, the cases presented above as well as previous literature in this area produce initial evidence that skilled, manual therapy-related assessment and rehabilitation of cervical spine dysfunction should be considered for chronic symptoms following concussion injuries.

Declaration of interest
The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.

References


Cervical spine and post-concussion syndrome


