Although concussion is recognised as one of the most frequently occurring injuries in contact sport, it is difficult for health practitioners to determine when an athlete can safely return to play. This difficulty primarily arises from the lack of clear, sensitive, and objective markers. More often than not, health practitioners must rely on the self-reported symptoms from the athlete and good clinical judgement as their primary tool for determining when an athlete should return to play. Unfortunately, relying on these symptoms is problematic, particularly in sport, and returning an athlete too soon may place them at risk for a further neurological injury, which could result in long term neurological sequelae.

A concussed athlete’s response to physiological stress may provide valuable insight into determining when they have recovered from concussion. However, few studies have investigated this relationship between mild head injury and physiological stress. In one study, concussed athletes demonstrated an increase in heart rate when exposed to mental stress, potentially indicating an autonomic dysfunction. In addition, neuroautonomic cardiovascular dysfunction has been well documented in severe head trauma. Thus, the purpose of this study was to assess the ability of athletes to perform exercise and their physiological response to that exercise, following an period of being asymptomatic at rest after a sport induced concussion.

**METHODS**

**Subjects**

A total 147 junior hockey players were given a written and oral briefing of the study and the risks involved prior to signing the informed consent. The study was approved by the university research ethics review committee. Baseline data collected included a medical questionnaire, concussion history, and a signs and symptoms questionnaire similar to that of Lovell and Collins.

Fourteen of the 147 participants had sustained a concussion. Their corresponding matched controls (n = 14) were team mates chosen to match, as closely as possible, the concussed athletes (table 1). Team trainers in collaboration with game observers used the concussion symptom guidelines of the Canadian Hockey Association to determine whether or not an athlete had sustained a concussion.

The concussed athletes performed the exercise protocol within 72 hours of being asymptomatic at rest. The test was repeated 5 days after the initial exercise assessment.

**Symptoms**

The same signs and symptoms questionnaire administered at baseline was given to the concussed athletes at 1.8 (0.2) days after injury. If the athlete was still symptomatic in relation to their baseline questionnaire, follow ups were conducted at 3 day intervals until the athlete was asymptomatic at rest or had returned to play. Once cleared to participate in exercise testing, the questionnaire was filled out prior to and 5 minutes following exercise at each testing session.

The questionnaire consisted of 20 signs and symptoms associated with concussion that are graded on a scale of 0–6 by the athlete. For the purpose of analysis between baseline and the symptoms reported at 2 days post-injury, a composite score was constructed using five symptoms that had been previously validated as indicators of concussion: headache, difficulty concentrating, difficulty remembering, nausea, and dizziness. Because the signs and symptoms questionnaire was administered 5 minutes following exercise, a physical composite score comprised of only nausea, dizziness, and headache was used in the analysis comparing before and after exercise.

**Exercise protocol**

All 28 athletes were required to fill out a modified PAR-Q before each testing session to ensure their health status had not changed from the start of the season. The exercise protocol was performed on a constant load cycle ergometer (Jaeger ER800, Germany). The participants began the protocol with a 2 minute warm up, pedalling at a frequency of 50–60 rpm with a load of 40 W. Following the warm up,
the athletes engaged in a 10 minute, low—moderate intensity, steady state exercise bout (80–90 rpm against a constant load of 1.5 W/kg of body weight). This was immediately followed by a high intensity interval session consisting of repeated 40 second high intensity bouts at 4.7 W/kg with a pedalling frequency of 90–100 rpm, followed by a 20 second free pedal (30 W) and a subsequent 20 second rest period. The high intensity exercise bouts were repeated until the athlete could no longer maintain the workload. Recovery consisted of a 1 minute free pedal with a load of 30 W, followed by a resting state for an additional 4 minutes. The number of high intensity exercise bouts completed was recorded. This exercise protocol was designed to reflect the high intensity interval activity observed in the sport of ice hockey.

Heart rate and blood lactate measures
A three lead CM5 configuration was used to record ECG data. The ECG signal was sampled at a frequency of 976 Hz through a Burdick EK-10 Electrocardiograph and simultaneously stored on a laptop using the WINDAQ® data acquisition system. ECG data was collected throughout the entire exercise protocol in addition to a 5 minute recovery sample immediately following the completion of the high intensity exercise protocol. To ensure that heart rate (HR) was at steady state, only minutes 2–9 from the low—moderate intensity steady state exercise bout were used for analysis. Five minute resting samples were also taken prior to exercise and at 30 min post-exercise. Blood was extracted from the fingertip prior to exercise and at 1 and 3 minutes post-exercise and analysed for lactate concentration (Accutrend™ Lactate analysis system).

Data analysis
The concussed athletes and their corresponding matched controls were divided into two cohorts as outlined in table 1; those who missed playing time (n = 9) for concussed and matched controls; no missed time cohort: n = 5 for concussed and matched controls. *The number of prior concussions does not include the concussion sustained during the course of this study.

<table>
<thead>
<tr>
<th>Missed time</th>
<th>Age (years)</th>
<th>Height (cm)</th>
<th>Weight (kg)</th>
<th>No. of prior concussions*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concussed</td>
<td>17.8 (0.5)</td>
<td>179.4 (1.4)</td>
<td>80.2 (4.1)</td>
<td>2.0 (0.5)</td>
</tr>
<tr>
<td>Matched controls</td>
<td>18.7 (0.4)</td>
<td>181.5 (1.6)</td>
<td>76.5 (1.8)</td>
<td>1.3 (0.5)</td>
</tr>
<tr>
<td>No missed time</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concussed</td>
<td>18.8 (0.8)</td>
<td>181.3 (2.7)</td>
<td>79.6 (2.2)</td>
<td>1.8 (0.6)</td>
</tr>
<tr>
<td>Matched controls</td>
<td>19.0 (0.8)</td>
<td>179.7 (2.5)</td>
<td>81.0 (3.2)</td>
<td>0.8 (0.6)</td>
</tr>
</tbody>
</table>

Values are means (SE). Missed time cohort: n = 9 for concussed and matched controls; no missed time cohort: n = 5 for concussed and matched controls.

RESULTS

Subjects
All of the 14 concussed athletes were deemed to have sustained a concussion based on the reports filed by the team trainers and game observers. Six of the concussed athletes from the MT cohort were unable to participate in the exercise protocol at 1.7 (0.2) days post-injury because their symptoms had not yet returned to baseline. Of the nine concussed athletes in this group, four experienced loss of consciousness, while four others exhibited post-traumatic amnesia immediately following their injury. The remaining athlete from this group reported no loss of consciousness, but had missed playing time as a direct result of his injury. In contrast, the symptoms from all concussed athletes of the NMT cohort had returned to baseline levels 2.0 (0.5) days after the injury. In addition, only one of the concussed athletes in the NMT group reported post-traumatic amnesia and none experienced loss of consciousness. An average of 6.7 (1.8) days for the MT cohort and 2.0 (0.5) days for the NMT group had elapsed from the time of injury to the first exercise test.

Exercise bouts and blood lactate levels
The number of high intensity exercise bouts completed by the concussed athletes was not significantly different from their matched controls in either cohort (table 2); however, a significant test by concussion interaction was detected. Specifically, the concussed group completed 8.0 (1.1) bouts at the first exercise test, and 6.8 (1.1) bouts at test 2. In contrast, their matched controls demonstrated the opposite response, completing 7.6 (0.7) bouts at test 1, and 8.2 (0.9) bouts at test 2. The concussed participants and the matched controls of the NMT group were able to perform an average of 9.4 (1.1) exercise bouts across both exercise tests, before reaching volitional fatigue. This was significantly higher than the 6.6 (0.7) bouts completed by the MT cohort.

In response to the high intensity exercise protocol, blood lactate levels rose significantly from an average resting value of 1.7 (0.1) mmol/l to 11.4 (0.4) mmol/l immediately after exercise. No significant difference in blood lactate levels at either rest or post-exercise was observed between the concussed group and their matched controls (table 2).

Symptoms
At baseline, both the MT and NMT concussed athletes had similar composite scores of 3.3 (1.4) and 3.6 (2.7) respectively. Following their reported concussion, the MT athletes had a score of 6.8 (2.2), whereas the NMT athletes had a composite score of 3.6 (2.0). Neither the main effect (3.6 (1.9) NMT; 5.0 (1.5) MT) or interactions were statistically significant.

At the first exercise test, the MT concussed athletes and their matched controls respectively reported a physical symptoms composite score of 1.9 (0.7) and 1.0 (0.5) at rest (fig 1). Following exercise at the same testing session, the
concussed athletes and their controls demonstrated a non-significant increase in symptom severity. The MT concussed athletes reported a score of 4.3 (1.8), whereas the MT controls reported less severe symptoms with a score of 2.3 (0.9). Similar results were also seen 5 days later (MT concussed 3.0 (1.1); MT control 2.5 (1.0)). No difference was found between the MT and NMT cohorts or between the concussed and control athletes of the NMT group (fig 2).

Heart rate
The maximum heart rate (MHR) achieved by the MT concussed athletes was higher, but not statistically different, at both testing sessions relative to their matched controls (table 3). Across both tests a significant rise in heart rate was observed during the course of the steady state exercise session (figs 3, 4). The rise in HR in the MT group over both exercise tests was found to be significantly greater than their matched controls (fig 3). The MT concussed athletes also had a significantly higher average heart rate than their matched controls over the 8 minute time period across both tests (126 (3.4) beats/min MT concussed; 116.0 (1.9) beats/min for MT control). This significant difference was not apparent between the concussed and control groups of the NMT cohort (fig 4).

HR dropped significantly during the 5 minutes immediately following exercise cessation (figs 5, 6). However, the average HR of the concussed athletes during this 5 minute time frame was not significantly different from their matched controls in either the MT or NMT cohort. Similarly no concussion by time interaction effect was detected, indicating that the decrease in HR over the 5 minute recovery period was similar in both groups (figs 5, 6).

Table 2  The number of exercise bouts completed and blood lactate levels (mmol/l) at rest and post-exercise during both testing sessions

<table>
<thead>
<tr>
<th>Test</th>
<th>Exercise bouts</th>
<th>Resting lactate</th>
<th>Post-exercise lactate</th>
<th>Exercise bouts†</th>
<th>Resting lactate</th>
<th>Post-exercise lactate*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Missed time</td>
<td>Concussed [n = 8]</td>
<td>6.6 (1.1)</td>
<td>2.2 (0.4)</td>
<td>10.9 (0.8)</td>
<td>5.3 (0.9)</td>
<td>1.7 (0.2)</td>
</tr>
<tr>
<td>Matched controls [n = 4]</td>
<td>7.4 (0.9)</td>
<td>1.6 (0.3)</td>
<td>11.1 (0.7)</td>
<td>7.3 (1.2)</td>
<td>1.8 (0.2)</td>
<td>11.1 (0.9)</td>
</tr>
<tr>
<td>No missed time</td>
<td>Concussed [n = 5]</td>
<td>10.2 (2.2)</td>
<td>1.5 (0.1)</td>
<td>12.2 (1.2)</td>
<td>9.2 (2.0)</td>
<td>1.6 (0.3)</td>
</tr>
<tr>
<td>Matched controls [n = 4]</td>
<td>8.0 (1.1)</td>
<td>1.4 (0.2)</td>
<td>10.0 (2.2)</td>
<td>10.0 (0.9)</td>
<td>1.5 (0.2)</td>
<td>12.9 (1.1)</td>
</tr>
</tbody>
</table>

Values are means (SE). *p<0.01 significant exercise effect across both tests; †p<0.05 missed time main effect across both tests.

DISCUSSION
The number of exercise bouts completed by the concussed athletes was not significantly different from their matched controls. However, concussed athletes who missed playing time had a significantly higher heart rate during the low–moderate, steady state exercise period, and they also exhibited a greater rise in heart rate over time. Despite the difference in cardiovascular response to exercise, concussed athletes and their matched controls reported similar symptoms before and after exercise. Contrary to our hypothesis, the capacity to perform high intensity interval activity was unaltered by concussion. The effort put forth by the concussed and control athletes at both testing sessions was...
The current study shows an 8% increase in submaximal HR in concussed athletes of the MT cohort conducted their first exercise test at slightly less than 1 week post-injury and within 72 hrs of being asymptomatic at rest. This may have been too late in the recovery phase for the effects of concussion to influence exercise capacity. However, it may be argued that a detraining effect, rather than concussed, n = 5; matched control, n = 3. *Significant time main effect (p<0.05).

Table 3 Heart rate (beats/min) at rest and post-exercise, and maximum heart rate achieved during high intensity exercise, at both testing sessions

<table>
<thead>
<tr>
<th>Missed time</th>
<th>Test 1</th>
<th>Test 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concussed (n=9)</td>
<td>68.2 (3.7)</td>
<td>183.4 (1.8)</td>
</tr>
<tr>
<td>Matched control (n=7)</td>
<td>63.2 (2.4)</td>
<td>173.6 (2.4)</td>
</tr>
<tr>
<td>No missed time</td>
<td>62.0 (3.1)</td>
<td>180.0 (9.8)</td>
</tr>
<tr>
<td>Concussed (n=4)</td>
<td>67.9 (2.1)</td>
<td>182.6 (6.7)</td>
</tr>
<tr>
<td>Matched control (n=4)</td>
<td>67.9 (2.1)</td>
<td>182.6 (6.7)</td>
</tr>
</tbody>
</table>

RHR, resting heart rate; PEHR, post-exercise heart rate; MHR, maximum heart rate. Values are reported as means (SE). *p<0.01 significantly different from both RHR and PEHR across both tests; †p<0.01 significantly different from RHR across both tests.

It is possible that the exercise capacity test used in this study may have been too gross a measure to detect a difference between the concussed athletes and their matched controls after a period of being asymptomatic at rest. An additional factor influencing the exercise capacity of the concussed athletes might be the elapsed time between the time of the injury and the first exercise test. Specifically, the concussed athletes of the MT cohort conducted their first exercise test at slightly less than 1 week post-injury and within 72 hrs of being asymptomatic at rest. This may have been too late in the recovery phase for the effects of concussion to influence exercise capacity.

The significant difference in HR during steady state exercise is of considerable interest. The use of symptoms to progress through a stepwise exercise programme is strongly recommended by the Concussion in Sports Group when managing a concussed athlete’s return to play.4 However, no detectable difference was found in the reported symptoms of our concussed athletes, despite these individuals demonstrating an adverse cardiovascular response to exercise. Consequently, while it appears that the concussed athlete has not yet fully recovered from their injury, this is not reliably detected by the reported symptoms.

It may be argued that a detraining effect, rather than concussion, is responsible for the elevated HR. However, previous studies suggest that an average elapsed time (from injury to the first exercise test) of less than 1 week is insufficient to induce a change in HR of this magnitude.19–21 The current study shows an 8% increase in submaximal HR in less than 1 week. This increase in HR is consistent with previous literature demonstrating abnormality in cardiovascular functioning following severe head trauma.11–13 It has been stated that athletes recovering from a sport induced concussion can probably return to play 5–7 days post-injury provided their symptoms have resolved.22 However, the speed of symptom resolution is dependent upon the individual.2 2 In the present study, HR was still elevated at the second exercise test 12 days post-injury in the concussed athletes who missed playing time. If indeed this elevated heart rate reflects cardiovascular autonomic dysfunction, a longer time frame is required to monitor these athletes to full recovery.

Of particular interest is a study by Goldstein et al,17 the results of which suggest that the degree of uncoupling...
WHAT IS ALREADY KNOWN ON THIS TOPIC

- Concussed athletes demonstrated an increase in heart rate when exposed to mental stress, potentially indicating a neuroautonomic dysfunction, which is well documented in severe head trauma.
- Study of concussion symptoms and responses to physical exercise is recommended in determining whether an athlete should return to play following a concussion.

WHAT THIS STUDY ADDS

- This is the first study to identify potential cardiovascular neuroautonomic control abnormalities during exercise in concussed athletes following period of being asymptomatic at rest.
- It also demonstrates that these athletes’ ability to perform maximum exercise is unaffected following period of being asymptomatic at rest after concussion.

between the autonomic and cardiovascular systems is proportional to the severity of the neurological injury. In contrast to the concussed athletes of the MT cohort, those from the NMT group did not demonstrate a difference in steady state heart rate in relation to their controls. Based on Goldstein’s findings, it could be suggested that the concussions sustained in the NMT group were not as severe. This assumption is further supported by the fact that the symptoms of this group had returned to baseline values at approximately 2 days post-injury. It is possible that, unlike traumatic brain injury, the severity of the neurological damage associated with concussion may be insufficient to induce cardiovascular changes at rest. However, under conditions of physiological stress, such as exercise, cardiovascular abnormalities may become more explicit.

In contrast to the anecdotal evidence, the concussed athletes from the present study demonstrated no difference in symptoms before and after exercise. It is unknown as to whether or not the lack of symptom aggravation is due to misrepresentation of symptoms from the athlete as a result of “contra-malingering”, or if indeed they were not perturbed as a function of exercise. Thus the use of symptoms, particularly following exercise, requires further research.

In summary, exercise capacity is unaffected in concussed athletes following a period of being asymptomatic at rest. However, the abnormal cardiovascular response to submaximal exercise appears to indicate impairment in neuroautonomic cardiovascular regulation. Furthermore, the degree of cardiovascular abnormality appears to be representative of concussion severity. Consequently, it is anticipated that this research will stimulate additional exploration into the relationship between neuroautonomic cardiovascular functioning following a concussion.

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Conflict of interest: none declared

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