Rethinking aerobic exercise intensity prescription in adults with spinal cord injury: time to end the use of “moderate to vigorous” intensity?

Michael J. Hutchinson and Victoria L. Goosey-Tolfrey

© The Author(s) 2021

INTRODUCTION

For adults with spinal cord injury (SCI), aerobic exercise is beneficial for improving indices of physical [1] and mental [2] health. On this theme, scientific guidelines published in 2018 describe the dose of aerobic exercise required to improve cardiorespiratory fitness and cardiometabolic health in adults with SCI [3]. Central to the guidelines is information on the exercise intensity. If aerobic exercise is performed at too low an intensity, without sufficient exercise volume, it will not lead to beneficial physiological adaptations [4]. Despite this, the guidelines provide no clear prescription of the exercise intensity, other than to say that aerobic exercise should be of a “moderate to vigorous” intensity [3]. The lack of clarity with the exercise intensity terminology is a hindrance to adults with SCI using the guidelines to inform their exercise habits; practitioners actively prescribing exercise training; and researchers investigating the effects of exercise training interventions on markers of health in adults with SCI. There is, therefore, an urgent need to better understand aerobic exercise intensity prescription in adults with SCI.

Guidelines for non-disabled adults define thresholds for five intensity zones (very light, light, moderate, vigorous, near-maximal/maximal) according to many physiological variables [5]. These variables include percentage maximum oxygen uptake (%VO₂peak) and heart rate (%HRpeak) with RPE differenced between PARA and TETRA. At LT1 and LT2, %VO₂peak and %HRpeak were significantly greater in TETRA compared to PARA and NON-SCI (P < 0.05). The variation in %VO₂peak and %HRpeak at lactate thresholds resulted in large variability in the domain distribution at fixed %VO₂peak and %HRpeak.

RESULTS: Associations between percentage of peak VO₂ (%VO₂peak) and HR (%HRpeak) with RPE differed between PARA and TETRA. At LT1 and LT2, %VO₂peak and %HRpeak were significantly greater in TETRA compared to PARA and NON-SCI (P < 0.05). The variation in %VO₂peak and %HRpeak at lactate thresholds resulted in large variability in the domain distribution at fixed %VO₂peak and %HRpeak.

CONCLUSIONS: Fixed %VO₂peak and %HRpeak should not be used for aerobic exercise intensity prescription in adults with SCI as the method does not lead to uniform exercise intensity domain distribution.

Spinal Cord (2022) 60:484–490; https://doi.org/10.1038/s41393-021-00733-2

STUDY DESIGN: Cohort study.

OBJECTIVES: To investigate and critique different methods for aerobic exercise intensity prescription in adults with spinal cord injury (SCI).

SETTING: University laboratory in Loughborough, UK.

METHODS: Trained athletes were split into those with paraplegia (PARA; n = 47), tetraplegia (TETRA; n = 20) or alternate health condition (NON-SCI; n = 67). Participants completed a submaximal step test with 3 min stages, followed by graded exercise test to exhaustion. Handcycling, arm crank ergometry or wheelchair propulsion were performed depending on the sport of the participant. Oxygen uptake (VO₂), heart rate (HR), blood lactate concentration ([BLa]) and ratings of perceived exertion (RPE) on Borg’s RPE scale were measured throughout. Lactate thresholds were identified according to log-VO₂ plotted against log-[BLa] (LT1) and 1.5 mmol L⁻¹ greater than LT1 (LT2). These were used to demarcate moderate (<LT1), heavy (LT1, <LT2) and severe (>LT2) exercise intensity domains.

RESULTS: Associations between percentage of peak VO₂ (%VO₂peak) and HR (%HRpeak) with RPE differed between PARA and TETRA. At LT1 and LT2, %VO₂peak and %HRpeak were significantly greater in TETRA compared to PARA and NON-SCI (P < 0.05). The variation in %VO₂peak and %HRpeak at lactate thresholds resulted in large variability in the domain distribution at fixed %VO₂peak and %HRpeak.

CONCLUSIONS: Fixed %VO₂peak and %HRpeak should not be used for aerobic exercise intensity prescription in adults with SCI as the method does not lead to uniform exercise intensity domain distribution.

Spinal Cord (2022) 60:484–490; https://doi.org/10.1038/s41393-021-00733-2
domain (between LT and critical power/speed (CP/CS)) there is a delayed steady state response due to the VO$_2$ “slow component”, whilst in the severe domain (above CP/CS) no steady state response is observed [8].

To satisfy the aim of producing a homogenous exercise intensity, the fixed percentage approach is only valid if it is demonstrated that equal relative intensities result in individuals exercising in the same intensity domain [7]. However, in a recent study of non-disabled participants, no fixed %VO$_2$max or %HR$_{max}$ typically used for exercise prescription, resulted in all participants being in the same intensity domain [9]. This has led to assertions that using fixed %VO$_2$max or %HR$_{max}$ for prescribing exercise intensity is inaccurate and will lead to significant inter-individual physiological responses, precluding homogenous exercise intensity prescription [7, 9]. Furthermore, with evidence that individual participant %VO$_2$R-%HR$_R$ relationships diverges from the assumed linear trajectory, there are also questions over how appropriate % VO$_2$R and %HR$_R$ are for prescribing exercise intensity at the individual level [10].

For adults with SCI there is currently nothing more to inform aerobic exercise intensity prescription than the arbitrary use of “moderate to vigorous” intensity [3]. Furthermore, evidence in non-disabled adults would suggest a need to rethink the traditional use of fixed percentages [7, 9, 10]. Therefore, this study aimed to investigate and critique potential methods for prescribing aerobic exercise intensity in adults with SCI.

METHODS

This study was performed via a retrospective analysis of athlete data collected in the author's laboratory. All procedures were approved by the Human participants ethical sub-committee at Loughborough University, and participants provided written, informed consent.

Participants

Data were available for 134 individuals (male: 98; female: 36). Participants were split into those with paraplegia (PARA), tetraplegia (TETRA), or alternate health condition (NON-SCI), see Table 1. Examples of health conditions for NON-SCI included spina bifida, limb deficiency, cerebral palsy, and arthrogryposis. Participants were competitive athletes, competing at a national or international level, from one of the following sports: handcycling, para-alpine ski, paratriathlon, wheelchair basketball, wheelchair rugby or wheelchair tennis.

Exercise testing

Participants completed a submaximal step test followed by graded exercise test (GXT) to exhaustion. Handcycle (HC) tests were performed in the participants own handcycle attached to a Cyclus 2 ergometer (Avantronic Richter, Leipzig, Germany). For some Paratriathlon, and all para-alpine ski athletes, arm crank ergometry (ACE) was used (Lode Angio, Lode B. V., Groningen, the Netherlands). The ergometer was positioned vertically so the crank axis centre was level with the shoulder, and horizontally to allow slight elbow flexion at the furthest point of the crank cycle. Wheelchair basketball, rugby and tennis players performed a wheelchair propulsion (WCP) test using a motorised treadmill (HP Cosmos, Traunstein, Germany) and their own custom sports wheelchair.

Submaximal tests were individualised based on the sport, sex, training status and level of impairment of the participant, with the goal of completing 6-8 stages (average: 6; range: 4–10). HC and ACE tests started at 15–60 W, with 10–20 W increments every 3 min. WCP tests started at 0.7–2.8 m s$^{-1}$ and were increased by 0.2–0.4 m s$^{-1}$ every 3 min. VO$_2$ (Metalyzer 3B, Cortex, Leipzig, Germany) and HR (RS400, Polar, Kempele, Finland) were continually monitored throughout. The Metalyzer was calibrated before each participant against ambient air and a mix of 15% O$_2$, 5% CO$_2$, with the volume calibrated using a 3 L syringe. RPE was verbally reported in the final minute of each stage using Borg's 6–20 RPE scale [11]. A capillary blood sample from the ear lobe was collected at the end of each stage for measurement of [BLA] (Biosen C-line, EKF Diagnostics, Barleben, Germany). HC and ACE tests were continuous, however, WCP tests were discontinuous as the treadmill needed to be slowed between stages to facilitate blood sampling. For discontinuous tests, the typical interval between stages was 45–60 s. Submaximal tests continued until [BLA] exceeded 4 mmol L$^{-1}$ or RPE was rated as 17. The RPE criteria was used in TETRA where there may have been blunted lactate responses [12].

Following the submaximal test, participants received 15 min of active recovery or rest before performing a GXT to exhaustion. The starting workloads were set to that from the prior submaximal (BLA) increased by 0.5 mmol L$^{-1}$ above rest. Participants performed 1 min at this load, before the exercise intensity were increased in a stepwise manner by 10–20 W min$^{-1}$ (HC/ACE) or 0.1 m s$^{-1}$ min$^{-1}$ (WCP) until participants reached volitional exhaustion. This was defined as an inability to maintain their preferred cadence at the required PO for HC/ACE, or the required speed on the treadmill, despite verbal encouragement. VO$_2$ and HR were again monitored throughout, as RPE and [BLA] measured at the end of the test.

Data processing

VO$_2$ and HR data were subjected to a 30 s rolling average, with the greatest of these from the GXT recorded as peak values (VO$_2$peak, HRpeak). VO$_2$ and HR in the final 30 s of each submaximal stage were extracted and calculated as percentages of peak (%VO$_2$peak, %HRpeak). Using the submaximal data, the lactate thresholds were identified as the intersection of the horizontal and ascending sections of the plot of log-[BLA] against log-VO$_2$ (LT$_1$) [13], and at [BLA] equal to LT$_1$ plus 1.5 mmol L$^{-1}$ (LT$_2$) [14]. The inverse of the log-VO$_2$ at these points were calculated to give the VO$_2$ at LT$_1$ and LT$_2$. HR at LT$_1$ and LT$_2$ was identified by interpolation of the linear VO$_2$-HR relationship for each participant. RPE was modelled against [BLA] using a quadratic function for each participant, with the resultant coefficients used to calculate the RPE at LT$_1$ and LT$_2$ [15]. Exercise intensity domains were defined as moderate (<LT$_1$), heavy (between LT$_1$ and LT$_2$) and severe (>LT$_2$).

Statistical analyses

Analyses were performed using IBM SPSS Statistics Version 23.0 (IBM Corp., Armonk, NY) and MLWIN Version 3.05 [16]. Data are presented as mean (standard deviation) with statistical significance accepted at P < 0.05. Data were checked for normal distribution using the Shapiro Wilk statistic.

All individual RPE data points were modelled against the corresponding %VO$_2$peak and %HR$_{peak}$ using dynamic multilevel models with lagged independent variable, whilst accounting for the initial condition. Separate models were created for %VO$_2$peak and %HR$_{peak}$ which served as the independent variable, with RPE as the dependent variable. Models were multilevel to adjust for the repeated stages performed by each participant and were used due to their ability to characterise group- and individual-level effects [17]. Stage was defined as the first, and participant as the second level. Models accounted for the initial condition (e.g., stage (i) = 1) as it was thought that RPE would depend on the %VO$_2$peak and %HR$_{peak}$ when i = 1. The need for them to be dynamic and incorporate a lagged independent variable was thought to be important as it was thought that RPE at subsequent measurement occasions (i.e., i = 1) would be dependent on %VO$_2$peak and %HR$_{peak}$ for that, as well as the previous, measurement occasion (i.e., i−1). Potential confounding variables were added to the models to assess whether they improved the model fit with fixed effects, or random effects for between- and within-individual variation. Confounding variables were sex (male/female), group (PARA/TETRA/NON-SCI) and exercise mode (ACE/HC/WCP). The resultant models were used to calculate the %VO$_2$peak and %HR$_{peak}$ Corresponding to each value on Borg's RPE scale.

Differences in VO$_2$ (L min$^{-1}$, ml kg$^{-1}$ min$^{-1}$, %VO$_2$peak), HR (beats min$^{-1}$, %HR$_{peak}$) and RPE between groups at LT$_{1}$ and LT$_{2}$ were assessed via one-way analysis of variance with Bonferroni post-hoc correction for multiple comparisons. Standardised effect sizes (ES) were calculated to describe the magnitude of differences and categorised as trivial (< 0.2), small (0.2–0.6), moderate (0.6–1.2), large (1.2–2.0) and very large (> 2.0) [18]. For each group, the percentage of participants in each intensity domain (moderate, heavy, severe) were calculated at 5% intervals from 35 to 95% VO$_2$peak and %HR$_{peak}$.

RESULTS

The associations between RPE and both %VO$_2$peak and %HR$_{peak}$ were not significantly affected by sex or exercise mode, so stratification based on these variables was not needed. The %
VO2peak and %HRpeak coinciding with each rating on Borg’s RPE scale for PARA and TETRA can be found in Table 2. The full RPE models against %VO2peak and %HRpeak can be found in the Supplementary Material.

**Table 1.** Participant characteristics by group.

|                         | PARA        | TETRA       | NON-SCI     |
|-------------------------|-------------|-------------|-------------|
| Sample size (n)         | 47          | 20          | 67          |
| Sex (M/F)               | 30/17       | 18/2        | 50/17       |
| Age (years)             | 33 ± 8 a    | 32 ± 7 a    | 27 ± 7      |
| Body mass (kg)          | 70.9 ± 14.1 a | 70.8 ± 12.9 | 64.5 ± 12.8 |
| Neurological level of injury | T4-L2     | C3-C7       | -           |
| Injury completeness     | Complete: 21 | Complete: 8 | -           |
|                        | Incomplete: 23 | Incomplete: 4 | -           |
| Time since injury (years) | 12 ± 9      | 12 ± 6      | -           |
| Peak oxygen uptake      | (L·min⁻¹)   | 2.5 ± 0.6 b | 1.7 ± 0.5   |
|                        | (ml·kg⁻¹·min⁻¹) | 35.1 ± 8.2 b | 23.4 ± 5.8 |
| Peak heart rate         | (beats·min⁻¹) | 188 ± 9 b   | 134 ± 20    |
| Sport (n)               | Handcycling | 8           | 0           |
|                        | Paratriathlon | 11         | 0           |
|                        | Para alpine ski | 2         | 1           |
|                        | Wheelchair basketball | 20     | 1           |
|                        | Wheelchair rugby | 0         | 15          |
|                        | Wheelchair tennis | 6         | 3           |
| Test mode (n)           | Arm crank ergometry | 11    | 1           |
|                        | Handcycling   | 10          | 0           |
|                        | Wheelchair propulsion | 26     | 19          |
| a: significantly greater than NON-SCI; b: significantly greater than TETRA, P < 0.05.

**Table 2.** Resultant calculations of percentage peak oxygen uptake and heart rate by group based on multilevel modelling.

| RPE | %VO2peak | %HRpeak |
|-----|----------|---------|
|     | PARA     | TETRA   | PARA | TETRA |
| 6   | 16       | 22      | 40   | 43    |
| 7   | 22       | 28      | 44   | 48    |
| 8   | 28       | 34      | 49   | 52    |
| 9   | 34       | 40      | 54   | 57    |
| 10  | 40       | 46      | 58   | 62    |
| 11  | 46       | 52      | 63   | 66    |
| 12  | 52       | 58      | 68   | 71    |
| 13  | 58       | 64      | 73   | 76    |
| 14  | 64       | 70      | 77   | 81    |
| 15  | 70       | 76      | 82   | 85    |
| 16  | 76       | 82      | 87   | 90    |
| 17  | 82       | 88      | 91   | 95    |
| 18  | 88       | 94      | 96   | 99    |
| 19  | 94       | 100     | 100  | -     |
| 20  | 100      | -       | -    | -     |

VO2peak and %HRpeak coinciding with each rating on Borg’s RPE scale for PARA and TETRA can be found in Table 2. The full RPE models against %VO2peak and %HRpeak can be found in the Supplementary Material.

**RPE and %VO2peak model**

RPE was significantly affected by the initial %VO2peak (i = 1) (P < 0.01), by %VO2peak at subsequent occasions when i > 1 (P < 0.01), as well as by the lagged %VO2peak (i.e., i-1) (P < 0.01). Each of these variables also showed significant between-individual variation, which was incorporated into the model. There was also an effect of Group at occasions when i > 1 (P = 0.01). TETRA showing significantly greater within-individual variation for the effect of %VO2peak on RPE compared to PARA and NON-SCI. As such, PARA and NON-SCI remained grouped, as there was no difference between these groups.

**RPE and %HRpeak model**

RPE was significantly affected by the initial %HRpeak (i = 1) (P < 0.01), by %HRpeak at subsequent occasions when i > 1 (P < 0.01), as well as by the lagged %HRpeak (i.e., i-1) (P < 0.01). These effects were fixed and showed no significant between- or within-individual variation. There was a fixed effect for Group, with the association between RPE and %HRpeak being significantly different for PARA (P = 0.03). There was no difference between TETRA and NON-SCI, so these remained grouped in this model.

**Responses at LT1 and LT2**

The VO2, HR and RPE at LT1 and LT2 are shown in Fig. 1. At LT1, there was a significant group effect for absolute (F2 = 7.11, P < 0.01; Fig. 1a) and relative VO2 (F2 = 17.65, P < 0.01; Fig. 1c), %VO2peak (F2 = 9.86, P < 0.01; Fig. 1e), HR (F2 = 42.79, P < 0.01; Fig. 1i) and %HRpeak (F2 = 7.94, P < 0.01; Fig. 1k). LT1 occurred at a significantly smaller absolute and relative VO2 in TETRA compared to PARA (ES = 0.86, 1.00) and NON-SCI (ES = 1.00, 1.62). However, %VO2peak at LT1 was significantly greater in TETRA compared to PARA (ES = 0.96) and NON-SCI (ES = 0.94). Similarly, HR at LT1 was smaller in TETRA compared to PARA (ES = 2.33) and NON-SCI (ES = 2.74), whereas %HRpeak was greater in TETRA compared to PARA (ES = 1.00) and NON-SCI (ES = 0.69). There was no significant difference between groups for RPE (F2 = 0.48, P = 0.62) at LT1 (Fig. 1g).

There was also a significant group effect at LT2 for absolute (F2 = 9.96, P < 0.01; Fig. 1b) and relative VO2 (F2 = 19.75, P < 0.01;
Fig. 1  Group physiological responses at lactate thresholds. a Absolute VO2 at LT1. b Absolute VO2 at LT2. c Relative VO2 at LT1. d Relative VO2 at LT2. e %VO2peak at LT1. f %VO2peak at LT2. g RPE at LT1. h RPE at LT2. i HR at LT1. j HR at LT2. k %HRpeak at LT1. l %HRpeak at LT2.

Data are presented at mean (SD) with individual points overlaid. Within each group, numbers refer to the same participant in each figure. Asterisk (*) indicates significantly greater than the identified group, P < 0.05.

Table 3. Classification of exercise intensity for individuals with paraplegia and tetraplegia, compared to non-disabled guidelines.

|                          | Very light (RPE ≤ 8)* | Light (RPE 9–11)* | Moderate (RPE 12–13)* | Vigorous (RPE 14–17)* | Near maximal-maximal (RPE ≥ 18)* |
|--------------------------|------------------------|-------------------|------------------------|------------------------|----------------------------------|
| %VO2peak                 | ≤36% VO2peak           | 37–45% VO2peak    | 46–63% VO2peak         | 64–90% VO2peak         | ≥91% VO2peak                     |
| Non-disabled*            |                        |                   |                        |                        |                                  |
| Paraplegia               | 31% VO2peak            | 32–49% VO2peak    | 50–61% VO2peak         | 62–85% VO2peak         | ≥86% VO2peak                     |
| Tetraplegia              | 37% VO2peak            | 38–55% VO2peak    | 56–67% VO2peak         | 68–91% VO2peak         | ≥92% VO2peak                     |
| %HRpeak                  | ≤56% HRpeak            | 57–63% HRpeak     | 64–76% HRpeak          | 77–95% HRpeak          | ≥96% HRpeak                      |
| Non-disabled*            |                        |                   |                        |                        |                                  |
| Paraplegia               | 51% HRpeak             | 52–65% HRpeak     | 66–75% HRpeak          | 76–93% HRpeak          | ≥94% HRpeak                      |
| Tetraplegia              | 54% HRpeak             | 55–68% HRpeak     | 69–78% HRpeak          | 79–97% HRpeak          | ≥98% HRpeak                      |

* data from Riebe et al. [5].

Fig. 1d), %VO2peak (F2 = 14.80, P < 0.01; Fig. 1f), HR (F2 = 58.99, P < 0.01; Fig. 1l) and %HRpeak (F2 = 6.10, P < 0.01; Fig. 1l). Absolute and relative VO2 at LT2 were significantly smaller in TETRA compared to PARA (ES = 0.83, 0.97) and NON-SCI (ES = 1.03, 1.72). However, %VO2peak at LT2 was significantly greater in TETRA compared to PARA (ES = 1.26) and NON-SCI (ES = 0.60). Furthermore, HR at LT2 was significantly smaller in TETRA than in PARA (ES = 3.00) and NON-SCI (ES = 3.55), while %HRpeak was significantly greater in TETRA compared to PARA (ES = 0.93) and NON-SCI (ES = 0.62). There was no significant difference between groups in RPE (F2 = 2.18, P = 0.19) at LT2 (Fig. 1h).

Intensity classification

Thresholds for %VO2peak and %HRpeak corresponding with intensity classifications used in non-disabled exercise guidelines are shown in Table 3. These data suggest there are differences between non-disabled individuals, PARA and TETRA in the thresholds for intensity classifications. Frequency distribution of individuals within moderate, heavy and severe intensity domains for discrete percentages of %VO2peak and %HRpeak are shown in Figs. 2 and 3, respectively. These show that no %VO2peak or %HRpeak typically used for exercise prescription purposes leads to all participants being in the same domain, with many %VO2peak including participants spread across all three domains.

DISCUSSION

This study aimed to investigate potential methods of aerobic exercise intensity prescription in adults with SCI. Findings demonstrate that there are differences between PARA and TETRA for the %VO2peak and %HRpeak corresponding with the
descriptions of “moderate” and “vigorous” exercise intensity, as used by the exercise guidelines for adults with SCI [1, 3]. However, the findings also show that using fixed %VO\textsubscript{2peak} or %HR\textsubscript{peak} cannot guarantee a homogenous domain-specific exercise intensity prescription for adults with SCI.

Fixed percentages and intensity domains for exercise intensity prescription

The finding of adults with SCI being in different intensity domains, as defined in this study according to LT\textsubscript{1} and LT\textsubscript{2}, despite being at the same %VO\textsubscript{2peak} or %HR\textsubscript{peak} supports similar evidence in non-disabled participants [9]. The domain-specific distribution is also arguably more variable in adults with SCI. For VO\textsubscript{2peak}, Iannetta et al. [9] report participants in moderate, heavy, and severe domains only at 70% VO\textsubscript{2peak}, whereas in this study this was shown at several fixed percentages, 55-70% VO\textsubscript{2peak} in PARA and 60-70% VO\textsubscript{2peak} in TETRA. It would, therefore, given recent calls to stop using fixed percentages for exercise intensity prescription in non-disabled participants [7, 9, 10], seem appropriate that this is expanded to apply to adults with SCI. For VO\textsubscript{2peak}, Iannetta et al. [9] report participants in moderate, heavy, and severe domains only at 70% VO\textsubscript{2peak}, whereas in this study this was shown at several fixed percentages, 55-70% VO\textsubscript{2peak} in PARA and 60-70% VO\textsubscript{2peak} in TETRA. It would, therefore, given recent calls to stop using fixed percentages for exercise intensity prescription in non-disabled participants [7, 9, 10], seem appropriate that this is expanded to apply to adults with SCI. Inter-individual variation will exist in sedentary or low-active participants as much, if not more, than in athletic groups, further limiting the use of fixed percentages for exercise intensity prescription.

Instead, more attention should be given to methods that can lead to participants exercising within the same exercise intensity domain, each of which are characterised by distinct VO\textsubscript{2} kinetic and blood lactate profiles [19, 20]. This study utilised LT\textsubscript{1} to identify the transition between moderate and heavy intensity domains, in accordance with the literature [19, 20]. However, a limitation within the current study was the use of LT\textsubscript{2} to identify the heavy-severe domain transition, due to a lack of evidence supporting this, as well as the number of different methods used to measure and identify LT\textsubscript{2} [7, 21]. As such, firm conclusions cannot be made regarding the heavy-severe domain transition from this study. It would have been more appropriate to measure CP/CS for this purpose [19–21], however, only data from a GXT were available in this study.

This highlights an important limitation to the widespread implementation of intensity domain-related exercise prescription. Specifically, the suitability of different testing protocols for identifying different threshold concepts, as well as data collection and threshold identification methods used [21]. This poses the challenge of how to simply prescribe domain-specific exercise intensity. Data from the present study may support the use of RPE for this purpose, as no difference in RPE was found between groups at LT\textsubscript{1} (Fig. 1g) and LT\textsubscript{2} (Fig. 1h), in support of previous findings [15]. Mean RPE at LT\textsubscript{1} and LT\textsubscript{2} in this study were found to be 11 and 15, respectively, suggesting that these values could be
used to guide exercise prescription using a simple and easy to implement method. However, the SD for these values ranged from 1 to 2 units dependent on group. This inter-individual variation serves, therefore, as a limitation to the use of fixed RPE values for exercise intensity prescription and highlights the importance of individualisation in this context. Individualisation, though, poses a further challenge, due to the trade-off between the precision required for research purposes versus the simple messaging for population-level exercise guidelines.

Implications for research
Our results highlight the need to prescribe exercise intensity in a way that ensures a homogenous intensity domain distribution between participants within both acute and longitudinal study designs. Previously, studies have conducted training interventions with an intensity of either a target range, or fixed value, using variables such as %VO₂peak, %HRpeak and RPE, e.g. [22–26], which will have led to significant domain heterogeneity. This means the dose of exercise stimulus would not have been controlled between participants. Moving forwards, researchers should identify the domain transitions and prescribe intensity in relation to these. That being said, the authors are not aware of any investigations into CP/CS using participants with SCI, so initial studies need to investigate the validity and reliability of identifying domain transitions in participants in SCI. Furthermore, VO₂ kinetic responses to exercise in each domain should be investigated in participants with SCI, due to potential differences in VO₂ kinetics between non-disabled participants and those with SCI [27]. Finally, these investigations must also account for any differences based on the mode of exercise [28].

Our findings also emphasise the need to individualise the exercise intensity prescription. In non-disabled participants, standardised intensity prescription (e.g., 55–75% VO₂peak) has been shown to lead to significant heterogeneity in responsiveness [29] leading to participants being described as either “responders” or “non-responders” to the intervention. However, an individualised exercise prescription (relative to ventilatory threshold) in non-disabled participants resulted in 100% responsiveness, compared to 60% in the standardised intervention [30]. Subsequently, future exercise research in participants with SCI should individualise the intensity prescription according to intensity domains, whilst also report individual responsiveness to an intervention. This will improve methodological control while also increasing confidence in conclusions made based on the data.

Implications for exercise guidelines using “moderate to vigorous” exercise intensity
Exercise guidelines must balance the precision required for a specific intensity prescription, against the need for a simple population-level recommendation. Furthermore, scientific guidelines must undergo a knowledge translation process to ensure that the scientific integrity of the guidelines are maintained, whilst also incorporating the varied needs of all potential end-users [31].

In adults with SCI, the scientific guidelines [1, 3] recommend performing aerobic exercise at a “moderate to vigorous” intensity, without providing any specific details on what this means. The same intensity prescription is also used in a community and clinical practice version of the guidelines [31]. Results from the current study would suggest the scientific integrity of such an intensity prescription is questionable. Table 3 shows equivalent thresholds for PARA and TETRA for “moderate” and “vigorous” intensity, based on the guidelines for non-disabled adults [5]. Combining these with the intensity domain distributions in Fig. 2, shows participants were spread across moderate, heavy, and severe domains at both “moderate” and “vigorous” intensities. It should also be noted that participants in the present study were competitive athletes, and that responses would likely be even more variable for sedentary or low-active populations. This shows how the use of “moderate” to “vigorous” intensity will not lead to anything close to resembling a uniform exercise intensity prescription between individuals. It also shows how someone expecting to perform “moderate” intensity exercise may actually be much closer to their maximum capacity. This will likely decrease the pleasure the person feels during the exercise, which could ultimately impact on whether they decide to continue doing it [32].

As it is operationally more difficult to define compared to frequency (e.g., 3 times a week) and duration (e.g., 30 min), it is possible that exercise intensity becomes an ignored piece of exercise guidelines. Perhaps research could seek to understand end-user perceptions of “exercise intensity”, or needs for interpreting and monitoring intensity, before using that information alongside physiological principles to underpin an evidence-informed intensity prescription. Alternatively, maybe the focus for exercise guidelines should shift from exercise intensity to also acknowledge factors that might help individuals become and stay active, such as their pleasure when performing exercise [32].

CONCLUSION
Prescribing a “moderate to vigorous” exercise intensity will not lead to a uniform intensity domain distribution in adults with SCI. Neither will the use of fixed percentages of VO₂peak or HRpeak or generic values of RPE, due to inter-individual variation. Such methods of exercise intensity prescription should not be used in this population. Future research should individualise the intensity prescription to ensure a homogenous inter-individual domain distribution. However, the accurate testing required to ensure an individualised intensity prescription poses a challenge to exercise guidelines aimed at informing behaviour at the population-level.

Data archiving
The datasets generated and analysed during the current study are available from the corresponding author on reasonable request.

REFERENCES
1. van der Scheer JW, Martin Ginis KA, Ditor DS, Goosey-Tolfrey VL, Hicks AL, West CR, et al. Effects of exercise on fitness and health of adults with spinal cord injury. Neurology 2017;89:736–45.
2. Kim DL, Lee J, Park H, Jeon YJ. The relationship between physical activity levels and mental health in individuals with spinal cord injury in South Korea. Int J Environ Res Public Health. 2020;17:1–9.
3. Martin Ginis KA, van der Scheer JW, Latimer-Cheung AE, Barrow A, Bourne C, Carruthers P, et al. Evidence-based scientific exercise guidelines for adults with spinal cord injury: an update and a new guideline. Spinal Cord. 2018;56:308–21.
4. van der Scheer JW, de Groot S, Tepper M, Faber W, ALLRISC Group, Veeger DH, et al. Low-intensity wheelchair training in inactive people with long-term spinal cord injury: a randomized controlled trial on fitness, wheelchair skill performance and physical activity levels. J Rehabil Med. 2016;48:33–42.
5. Riebe D, Ehrman JK, Liguori G, Magal M ACSM’s guidelines for exercise testing and prescription. Tenth edit. (Wolters Kluwer: Philadelphia, PA, 2018).
6. Theisen D. Cardiovascular determinants of exercise capacity in the Paralympic athlete with spinal cord injury. Exp Physiol. 2012;97:319–24.
7. Jammick NA, Pettitt RW, Granata C, Pyne DB, Bishop DJ. An examination and critique of current methods to determine exercise intensity. Sports Med. 2020;50:1729–56.
8. Rumley M, Jones AM. Power–duration relationship: physiology, fatigue, and the limits of human performance. Eur J Sport Sci. 2018;18:1–12.
9. Iannetta D, Inglis EC, Mattu AT, Fontana FY, Pogliaghi S, Keir DA, et al. A critical evaluation of current methods for exercise prescription in women and men. Med Sci Sport Exerc. 2020;52:466–73.
10. Ferri Marini C, Sisti D, Leon AS, Skinner JS, Sarzynski MA, Bouchard C, et al. HRR and VO2R fractions are not equivalent: is it time to rethink aerobic exercise prescription methods? Med Sci Sport Exerc. 2021;53:174–82.
11. Borgan GA Borg’s Perceived Exertion and Pain Scales. (Human Kinetics: Champaign, IL, 1998).
12. Hutchinson MJ, Kilgallon JW, Leicht CA, Goosey-Tolfrey VL. Perceived exertion responses to wheelchair propulsion differ between novice able-bodied and trained wheelchair sportspersons. J Sci Med Sport. 2019;23:403–7.
13. Beaver WL, Wasserman K, Whipp BJ. Improved detection of lactate threshold during exercise using a log-log transformation. J Appl Physiol. 1985;59:1936–40.
14. Dickhuth HH, Huonker M, Münzel T, Drexler H, Berg A. Keul J Individual anaerobic threshold for evaluation of competitive athletes and patients with left ventricular dysfunction. In: Bachl N, Graham TE, Lollgren H (eds). Advances in Ergometry. (Springer: Berlin Heidelberg, 1991), pp 173-9.
15. Hutchinson MJ, Kouwijzer I, de Groot S, Goosey-Tolfrey VL. Comparison of two Borg exertion scales for monitoring exercise intensity in able-bodied participants, and those with paraplegia and tetraplegia. Spinal Cord. 2021. https://doi.org/10.1038/s41393-021-00642-4.
16. Chartlon C, Rasbash J, Browne WJ, Healy M, Cameron B MLWIN Version 3.05. 2020.
17. Kristjansson SD, Kircher JC, Webb AK. Multilevel models for repeated measures research designs in psychophysiology: an introduction to growth curve modeling. Psychophysiology 2007;44:728–36.
18. Batterham AM, Hopkins WG. Making meaningful inferences about magnitudes. Int J Sports Physiol Perform. 2006;1:50–57.
19. Poole DC, Jones AM. Oxygen uptake kinetics. Compr Physiol. 2012;2:933–96.
20. Poole DC, Burnley M, Vanhatalo A, Rossiter HB, Jones AM. Critical power: an important fatigue threshold in exercise physiology. Med Sci Sport Exerc. 2016;48:320–34.
21. Poole DC, Rossiter HB, Brooks GA, Gladden LB. The anaerobic threshold: 50+ years of controversy. J Physiol. 2021;599:737–67.
22. Bakkum AJ, de Groot S, Stolwijk-Swuste JM, van Kuppevelt DJ, ALLRISC, van der Woude LH, et al. Effects of hybrid cycling versus handcycling on wheelchair-specific fitness and physical activity in people with long-term spinal cord injury: a 16-week randomized controlled trial. Spinal Cord. 2015;53:395–401.
23. Hicks AL, Martin KA, Ditor DS, Latimer AE, Craven C, Bugaresti J, et al. Long-term exercise training in persons with spinal cord injury: effects on strength, arm ergometry performance and psychological well-being. Spinal Cord. 2003;41:34–43.
24. Kim DI, Lee H, Lee BS, Kim J, Jeon JY. Effects of a 6-week indoor hand-bike exercise program on health and fitness levels in people with spinal cord injury: a randomized controlled trial study. Arch Phys Med Rehabil. 2015;96:2033–40.
25. Pelletier CA, Totosy de Zepetnek JO, MacDonald MJ, Hicks AL. A 16-week randomized controlled trial evaluating the physical activity guidelines for adults with spinal cord injury. Spinal Cord. 2015;53:363–7.
26. Valenti LJ, Dallmeijer AJ, Houdijk J, Slooman HJ, Janssen TW, Post MWM, et al. Effects of hand cycle training on physical capacity in individuals with tetraplegia: a clinical trial. Phys Ther. 2009;89:1051–60.
27. Gollie JM, Herrick JE, Keyser RE, Chien LMK, Collins JP, Shields RK, et al. Fatigability, oxygen uptake kinetics and muscle deoxygenation in incomplete spinal cord injury during treadmill walking. Eur J Appl Physiol. 2017;117:1989–2000.
28. Barstow TJ, Scremin AME, Mutton DL, Kunkel CF, Cagle TG, Whipp BJ. Peak and kinetic cardiorespiratory responses during arm and leg exercise in patients with spinal cord injury. Spinal Cord. 2000;38:430–5.
29. Bouchard C, An P, Rice T, Skinner JS, Gagnon J, et al. Familial aggregation of VO2(max) response to exercise training: Results from the HERITAGE family study. J Appl Physiol. 1999;87:1003–8.
30. Weathermax RM, Harris NK, Kilding AE, Dalleck LC. Incidence of VO2max responders to personalized versus standardized exercise prescription. Med Sci Sport Exerc. 2019;51:681–91.
31. Hoekstra F, McBride CB, Borisoff J, Fetterly MJ, Ginis S, Latimer-Cheung AE, et al. Translating the international scientific spinal cord injury exercise guidelines into community and clinical practice guidelines: a Canadian evidence-informed resource. Spinal Cord. 2020;58:647–57.
32. Ekkekakis P, Parfitt G, Petruzzello SJ. The pleasure and displeasure people feel when they exercise at different intensities: decennial update and progress towards a tripartite rationale for exercise intensity prescription. Sports Med. 2011;41:641–71.

ACKNOWLEDGEMENTS
We would like to thank the various members of the Peter Harrison Centre for Disability Sport who were involved in the collection of data used in this study. These include Dr John Lenton, Dr Christof Leicht, Dr Tom Paulson, Dr Terri Paulson, Dr Katy Griggs, Dr Ben Stephenson, Dr Ben Stone, Tom O’Brien, and Conor Murphy.

AUTHOR CONTRIBUTIONS
MJH and VLGT were both responsible for developing the concept of the study and were both involved in the data collection. MJH performed the data analysis, with MJH and VLGT involved in the interpretation of the data. MJH drafted the original version of the manuscript, with VLGT providing comments. MJH and VLGT were both responsible for reviewing and approving the final version of the manuscript.

FUNDING
This work was supported by funding from the Peter Harrison Foundation.

STATEMENT OF ETHICS
We certify that all applicable institutional regulations concerning the ethical use of human volunteers were followed during the course of this research.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41393-021-00733-2.

Correspondence and requests for materials should be addressed to Victoria L. Goosey-Tolfrey.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2021