Prolonged cycling reduces power output at the moderate-to-heavy intensity transition

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Abstract
Purpose To determine the effect of prolonged exercise on moderate-to-heavy intensity transition power output and heart rate.
Methods Fourteen endurance-trained cyclists and triathletes took part in the present investigation (13 males, 1 female, V·O₂peak 59.9 ± 6.8 mL.kg⁻¹.min⁻¹). Following a characterisation trial, participants undertook a five-stage incremental step test to determine the power output and heart rate at the moderate-to-heavy intensity transition before and after two hours of cycling at 90% of the estimated power output at first ventilatory threshold (VT₁).
Results Power output at the moderate-to-heavy intensity transition significantly decreased following acute prolonged exercise when determined using expired gases (VT₁, 217 ± 42 W vs. 196 ± 42 W, P < 0.0001) and blood lactate concentrations (LoglogLT, 212 ± 47 W vs. 190 ± 47 W, P = 0.004). This was attributable to loss of efficiency (VT₁, -8 ± 10 W; LoglogLT, -7 ± 9 W) and rates of metabolic energy expenditure at the transition (VT₁, −14 ± 11 W; LoglogLT, −15 ± 22 W). The heart rate associated with the moderate-to-heavy intensity transition increased following acute prolonged exercise (VT₁, 142 ± 9 beats.min⁻¹ vs. 151 ± 12 beats.min⁻¹, P < 0.001; LoglogLT, 140 ± 13 beats.min⁻¹ vs. 150 ± 15 beats.min⁻¹, P = 0.006).
Conclusion These results demonstrate the external work output at the moderate-to-heavy intensity transition decreases during prolonged exercise due to decreased efficiency and rates of metabolic energy expenditure, but the associated heart rate increases. Therefore, individual assessments of athlete ‘durability’ are warranted.

Keywords Cycling · Thresholds · Durability · Fatigue resistance · Exercise · Duration

Introduction
Endurance athletes commonly perform physiological assessments for estimating the external work outputs that demarcate the boundaries between exercise intensity domains. These intensity domain transitions are used to assess performance capability, regulate training and competition intensities, monitor training load, and quantify adaptations to training (Maunder et al. 2021). The moderate and heavy exercise intensity domains are defined by physiological responses, including distinct blood lactate and whole-body oxygen uptake (V·O₂) kinetics profiles (Burnley and Jones 2018; Jones et al. 2019).

An under-studied effect of prolonged exercise is the likely reduction in external work rates observed at the intensity transitions over time; the resilience to which we termed ‘durability’ (Maunder et al. 2021). A series of recent studies reported a reduction in the so-called critical power following prolonged heavy-intensity cycling, and indicated that the magnitude of the reduction was sensitive to carbohydrate availability (Clark et al. 2018, 2019a, b). Less is understood
regarding the effects of acute prolonged exercise on the moderate-to-heavy intensity transition, which may plausibly also decline with prolonged exercise in a manner related to carbohydrate availability. Specifically, depletion of the intramyofibrillar glycogen store during prolonged exercise appears to impair muscle contractile function (Ortenblad et al. 2013). Therefore, following prolonged exercise, intramyofibrillar glycogen depletion-induced impairment of specific muscle fibres may increase the mechanical and metabolic burden of a given power output on a smaller pool of fully functional fibres. Speculatively, this could increase the fibre-specific work rate at a given power output, reducing the power output achieved at the moderate-to-heavy intensity transition following prolonged exercise.

A reduction in gross cycling efficiency with prolonged moderate-intensity exercise has been observed (Passfield and Doust 2000; Hopker et al. 2017). This would theoretically contribute to reduced power output at intensity transitions following prolonged exercise, even if the rate of metabolic energy expenditure associated with the transitions is maintained over time. The contributions made by changes in gross cycling efficiency and rates of metabolic energy expenditure to prolonged exercise-induced changes in the power output at the moderate-to-heavy intensity transition could be quantified by measurement of the energy expenditure vs. power output relationship, and identification of the rate of energy expenditure associated with the moderate-to-heavy intensity transition, before and after prolonged exercise.

Given the possibility that power outputs at intensity domain transitions may decrease during prolonged exercise, the use of data generated in well-rested physiological profiling assessments to regulate training intensity could result in inadvertent drift into higher intensity domains over time (Maunder et al. 2021). A common strategy employed by endurance athletes to combat this is the use of heart rates in inadvertent drift into higher intensity domains over time.

Methods

Participants

Fourteen endurance-trained cyclists and triathletes took part in the present investigation (13 males, 1 female, age 34 ± 10, height 178.1 ± 5.6 cm, mass 71.2 ± 6 kg, \( V\cdot O_{2\text{peak}} \) 59.9 ± 6.8 mL kg\(^{-1}\) min\(^{-1}\), training volume 9 ± 3 h week\(^{-1}\)). A priori sample size estimation indicated that a total sample size of 15 was required to detect a large magnitude (ES = 0.7) reduction in power output at the moderate-to-heavy intensity transition with 80% statistical power using the G*Power software package. A large magnitude effect size was used for this calculation based on previous studies showing the effect of prolonged exercise on the heavy-to-severe intensity transition (Clark et al. 2018, 2019a, b). A one-tailed test was used as it seemed implausible that the moderate-to-heavy intensity power output would increase following acute prolonged exercise. Data collection was interrupted by a COVID-19 lockdown and one participant dropped out. All participants were free of recent (<3 months) musculoskeletal injury and chronic disease and habitually training > 5 h week\(^{-1}\) in cycling-based endurance sports. This study was performed in accordance with the standards of the Declaration of Helsinki, 2013, and the Auckland University of Technology Ethics Committee approved all procedures (21/253).
Study design

Participants visited the laboratory on two occasions, ~7 d apart, for: (i) a characterisation trial involving a maximal, incremental cycling test after an overnight fast and (ii) a prolonged trial, which involved a prolonged cycling assessment with estimation of the first ventilatory threshold (VT1) before and after two hours of moderate-intensity cycling, after an overnight fast (Fig. 1). The order of visits was not randomised as the incremental test data were used to define the parameters of the prolonged trial.

Characterisation trial

Participants initially reported to the laboratory for an incremental cycling test. Participants arrived after a 10-h overnight fast having refrained from vigorous exercise for 24 h and having ingested ~ 1 L of plain water ~ 2 h beforehand. Height and body mass was first measured. Cycling subsequently commenced on an electromagnetically braked ergometer at 95 W, and the power output initially increased by 35 W every 3 min (Excalibur Sport, Lode BV, Groningen, NET). Expired gases were collected continuously using indirect calorimetry (TrueOne 2400, ParvoMedics, UT, USA). Heart rate was measured continuously using a chest-strap heart rate monitor (Tickr, Wahoo Fitness, Atlanta, USA).

Once the respiratory exchange ratio exceeded 1.0 and clear signs of increased VE V·O2−1 emerged, power output was increased by 35 W every minute until task failure. The V·O2peak was identified as the highest 15-s average V·O2, and VT1 was identified as the V·O2 at which a systematic rise in VE V·O2−1 occurred. This V·O2 was converted to a power output by linear fit of the power output vs V·O2 relationship, using the last minute of V·O2 data from each 3-min stage.

The last minute of expired gas data in each 3-min stage was used to determine whole-body fat oxidation rates through standard calculations (Jeukendrup and Wallis 2005) (Eq. 1). The highest observed rate of whole-body fat oxidation was accepted as the peak fat oxidation rate (PFO), as per our recent work (Maunder et al. 2022).

\[
\text{Whole body fat oxidation rate (g \cdot min}^{-1}) = 1.695 \times \dot{V}O_2 - 1.701 \times \dot{V}CO_2
\]

where V·O2 and VCO2 are in L min−1.

Prolonged trial

Participants arrived for the prolonged trial after a 10-h overnight fast, having refrained from vigorous exercise for 24 h, and having ingested ~ 1 L of plain water ~ 2 h beforehand. Following measurement of body mass, the experimental trial commenced on the same electromagnetically braked ergometer as the initial assessment with a 5-min warm-up at 100 W, followed by a five-stage incremental assessment to determine the power output and heart rate at the moderate-to-heavy intensity transition. The incremental test began with 4-min at 50 W below the previously estimated VT1 power output, and power output increased by 25 W per increment.

Expired gases and heart rate were measured continuously (TrueOne 2400, ParvoMedics, UT, USA; Tickr, Wahoo Fitness, Atlanta, USA), and a fingerprick capillary blood sample was obtained in the last 30-s of each increment for measurement of blood lactate concentration using an automated analyser (Lactate Pro 2, Arkray). These data were used to quantify the power output and heart rate at the moderate-to-heavy intensity transition prior to prolonged exercise (PRE, see Data analysis section below). Participants then cycled for 5 min at 100 W, and then at 90% of the previously estimated power output at VT1 for 2 h. Heart rate was recorded throughout, and participants consumed plain water ad libitum. Expired gases were collected for 4 min, every 15 min.

Following the two-hour constant work-rate phase, participants again cycled for 5 min at 100 W before repeating the five-step incremental exercise assessment. These data were used to quantify the power output and heart rate at the moderate-heavy intensity transition following prolonged exercise (POST). Sweat loss was also assessed by measurement of pre- and post-exercise body mass, and water consumption. Total water consumption was recorded by measuring the mass of the bottle before and after use and refilling, and was then added to changes in body mass.

Fig. 1 Schematic overview of the two laboratory visits
to calculate total sweat loss. If participants used the toilet during the trial, body mass was recorded before and after and accounted for in sweat loss calculations.

**Estimation of the moderate-to-heavy transition**

The PRE and POST moderate-to-heavy intensity transitions were estimated using expired gas and blood lactate data. Specifically, using expired gas data, the moderate-to-heavy intensity transitions in the PRE and POST assessments of the experimental trial were estimated using the VT1 method in accordance with the procedures described above for the initial assessment. The V̇O₂ at VT1 was converted to a power output by linear fit of the power output vs. V̇O₂ relationship, using the last minute of V̇O₂ data from each of the five 4-min stages. Power output was then matched with a heart rate value by linear fit of the power output vs. heart rate relationship, using the average heart rate during the last minute of each 4-min stage. The VT1 power output was then converted to a rate of whole-body energy expenditure by linear fit of the whole-body energy expenditure vs. power output relationship. The whole-body rate of energy expenditure was calculated for each power output in the incremental assessment using the average V̇O₂ and VCO₂ in the last minute of each of the five 4-min stages with a stoichiometric equation (Jeukendrup and Wallis 2005) (Eq. 2)

\[
\text{Whole body rate of energy expenditure (kcal x min}^{-1} = 0.550 \times \dot{VCO}_2 + 4.471 \times \dot{VO}_2
\]

Where V̇O₂ and V̇ĊO₂ are in L min⁻¹.

Using blood lactate data, the PRE and POST moderate-to-heavy intensity transitions were estimated using the LoglogLT method. The LoglogLT method models a blood lactate concentration vs. power output curve using two segments, and the intersection point of the two lines with the lowest residuals sum of squares is taken as the moderate-to-heavy intensity transition (Jamnick et al. 2018). The LoglogLT power output in the PRE and POST assessments were converted to heart rate, V̇O₂, and whole-body rate of energy expenditure values by linear fit of the relationships between these values and power output, as per above. As LoglogLT data produced essentially the same results as VT1, and the same inferences, only VT1 data are reported. The LoglogLT data can be found in Supplementary Figs. 1–2.

To quantify the proportion of prolonged exercise-induced changes in moderate-to-heavy intensity transition power output associated with changes in gross cycling efficiency and changes in rates of metabolic energy expenditure achieved at the moderate-to-heavy transition, rates of energy expenditure observed at VT1 and LoglogLT in the POST assessment were converted to a power output value using linear regression of the power output vs. energy expenditure relationship for each participant in the PRE assessment (denoted as POSTEEPREEff). The POSTEEPREEff therefore indicates the power output that the rate of metabolic energy expenditure observed at the moderate-to-heavy transition in the POST assessment would have achieved with the level of gross cycling efficiency in the PRE assessment. Accordingly, the proportion of prolonged exercise-induced changes in VT1 and LoglogLT power output associated with changes in energetic efficiency and rates of metabolic energy expenditure achieved at the transition was calculated using the below equation (Eq. 3).

\[
\text{Contribution of change in energetic efficiency to change in power output at the moderate to heavy intensity transition = POST - POSTEEPREEff}
\]

**Statistical analysis**

Data are presented as mean ± standard deviation (SD), unless otherwise stated. Normality of data distributions were assessed using the Shapiro–Wilks test. The effect of prolonged exercise on moderate-to-heavy intensity transition power output, heart rate, V̇O₂, and rate of energy expenditure was assessed using paired t-tests (or the non-parametric equivalent Wilcoxon test). Relationships between PRE to POST changes in moderate-to-heavy transition power output and PFO, V̇O₂peak, the moderate-to-heavy intensity transition in the PRE assessment, sweat loss, and dehydration were assessed using Pearson’s (r) or Spearman’s rank-order (r_s) correlation coefficients, depending on normality, and expressed with 95% confidence intervals. Changes in heart rate, whole-body energy expenditure, V̇O₂, and respiratory exchange ratio (RER) over-time during the two-hour constant work-rate phase were analysed using repeated measures one-way analyses of variance. Whole-body fat
oxidation rates during the first three stages of the PRE and POST incremental tests were compared using a mixed model analysis of variance due to missing data-points. Variance was located post-hoc using Holm-Bonferroni stepwise correction. Analyses were performed in GraphPad Prism Version 9.3.1. Significance was inferred when $P \leq 0.05$.

**Results**

**Constant work-rate phase**

The estimated power output at VT$_1$ in the initial assessment was 216 ± 45 W. Consequently, the two-hour constant work-rate phase in the experimental trial was completed at 194 ± 41 W. From 15 to 120 min of the two-hour constant work-rate phase, heart rate significantly increased ($P < 0.0001, 8.2 \pm 2.7\%$, Fig. 2a). Rates of whole-body energy expenditure ($P = 0.07, 3.4 \pm 4.3\%$, Fig. 2b) and VO$_2$ ($P = 0.08, 3.4 \pm 4.9\%$, Fig. 2c) did not increase during the two-hour constant work-rate phase, although both effects approached significance. The RER did not change during the two-hour constant work-rate phase ($P = 0.61$, Fig. 2d).

**Moderate-to-heavy intensity transition**

The power output at VT$_1$ significantly decreased from PRE to POST (217 ± 42 W vs. 196 ± 42 W, $\Delta = -21 \pm 12$ W, $\Delta = -10.0 \pm 5.8\%, P < 0.0001$, Fig. 3). The magnitude of PRE to POST change in VT$_1$ power output was not significantly associated with PFO, VO$_2$peak, prolonged exercise-induced sweat loss, or prolonged exercise-induced dehydration (Table 1). However, the magnitude of the reduction in VT$_1$ power output from PRE to POST was be related to the VT$_1$ expressed as %VO$_2$peak in the PRE assessment.
Table 1: Bivariate associations between durability of the moderate-to-heavy transition (Δ VT1) expressed in absolute units, W, and as a percentage of PRE values and outcome measures in this study. Data are reported as Spearman’s rank-order (rs) correlation coefficients (95% confidence intervals), with accompanying P values.

|          | Δ VT1 (W) | Δ VT1 (%) |
|----------|-----------|-----------|
| PFO (g min⁻¹) | r_s = −0.13 | r_s = 0.09 |
| (g min⁻¹) | (−0.63, 0.44) | (−0.47, 0.61) |
| Δ FO step 1 | P = 0.66 | P = 0.75 |
| (g min⁻¹) | r_s = −0.14 | r_s = 0.17 |
| (−0.63, 0.43) | (−0.43, 0.67) |
| Δ FO step 2 | P = 0.63 | P = 0.58 |
| (g min⁻¹) | r_s = −0.24 | r_s = 0.22 |
| (−0.69, 0.45) | (−0.39, 0.70) |
| Δ FO step 3 | P = 0.40 | P = 0.46 |
| (g min⁻¹) | r_s = −0.18 | r_s = 0.20 |
| (−0.67, 0.43) | (−0.44, 0.71) |
| V̇O₂peak (mL.kg⁻¹.min⁻¹) | P = 0.57 | P = 0.42 |
| (−0.69, 0.36) | (−0.41, 0.66) |
| V̇O₂peak (L.min⁻¹) | P = 0.43 | P = 0.54 |
| (−0.75, 0.24) | (−0.52, 0.56) |
| PRE VT1 (%V̇O₂peak) | P = 0.22 | P = 0.92 |
| (−0.00, 0.84) | (−0.03, 0.83) |
| 0.54 | 0.52 |
| Sweat loss | P = 0.05 | P = 0.06 |
| (L) | r_s = −0.17 | r_s = 0.23 |
| (−0.71, 0.49) | (−0.45, 0.74) |
| Dehydration | r_s = 0.43 | r_s = 0.50 |
| (% of BM) | (−0.25, 0.82) | (−0.16, 0.85) |
| P = 0.19 | P = 0.19 |

Δ FO step 1–3 = change in whole-body fat oxidation rate from PRE to POST in step 1–3. BM body mass, PFO peak fat oxidation rate observed in the initial assessment, V̇O₂peak = peak oxygen uptake, and Δ VT1 = prolonged exercise-induced change in the first ventilatory threshold power output. Significant relationships (P ≤ 0.05) are highlighted in bold.

There was an effect of prolonged exercise (P = 0.04), and prolonged exercise by intensity interaction (P = 0.002), on whole-body fat oxidation rates during the PRE and POST assessments. Specifically, whole-body fat oxidation rates were greater in the POST vs. PRE assessment during the first and second steps (Fig. 5). The loss of power output at the moderate-to-heavy intensity transition associated with decreased energetic efficiency was significantly associated with changes in whole-body fat oxidation rates from PRE to POST, but these relationships were not present for the loss of power output associated with rates of metabolic energy expenditure at the transition (Table 2, Supplementary Fig. 3).

Discussion

The aim of this study was to determine the effects of prolonged moderate-intensity cycling on the moderate-to-heavy intensity transition power output and heart rate. Our primary observations were that: (i) the power output at the moderate-to-heavy intensity transition significantly decreased following prolonged cycling, (ii) this loss of power output was attributable to reduced gross cycling energetic efficiency and reduced rates of metabolic energy expenditure at the transition, and (iii) the heart rate associated with the moderate-to-heavy intensity transition increased following prolonged cycling. These data have implications for athlete profiling, training load monitoring, and training programming, and indicate that the ‘durability’ of the moderate-to-heavy intensity transition warrants attention at the individual level.

The observed reduction in moderate-to-heavy intensity transition power output following prolonged exercise was partially accounted for by reduced energetic efficiency, or the translation of metabolic energy expenditure to mechanical power output (Fig. 4d). Loss of efficiency was also demonstrated during the two-hour constant work-rate phase, as whole-body rates of energy expenditure (P = 0.07, Fig. 2b) and V̇O₂ (P = 0.08, Fig. 2b) at the fixed power output increased over-time, albeit not significantly. These observations are in line with previous work demonstrating decreased gross cycling efficiency following prolonged exercise (Passfield and Doust 2000; Hopker et al. 2017). Mechanistically, the rising energy and V̇O₂ cost of given power outputs following prolonged exercise may reflect progressive loss of skeletal muscle contractile efficiency due to increasing recruitment of less efficient type II muscle fibres (Jones et al. 2011). The increased V̇O₂ cost of producing a fixed power output would have been exacerbated beyond the loss of energetic efficiency by the increased fat oxidation (Fig. 5), as fat oxidation requires more V̇O₂ per unit of energy produced than carbohydrate oxidation (Frayn 1983).
The remainder of the reduction in moderate-to-heavy intensity transition power output was accounted for by decreased rates of metabolic energy expenditure at the transition (Fig. 4d). This is demonstrated by the reduction in rates of energy expenditure as well as \( V\dot{O}_2 \) at the moderate-to-heavy intensity transition from PRE to POST (Fig. 4ab). Plausibly, the observed reduction in energy expenditure at the moderate-to-heavy intensity transition with prolonged exercise may also be at least partially attributable to decreased endogenous carbohydrate availability. Specifically, localised glycogen depletion in the intramyofibrillar compartment has been linked to impaired excitation–contraction coupling, manifesting as reduced \( Ca^{2+} \) release from the sarcoplasmic reticulum under neural innervation (Ørtenblad et al. 2013). Therefore, intramyofibrillar glycogen depletion during the prolonged exercise of the current study may plausibly have diminished the function of individual working muscle fibres. Evidence for depletion of endogenous carbohydrate availability is provided by the observed increase in whole-body fat oxidation rates from PRE to POST (Fig. 5), given the autoregulatory nature of muscle glycogen metabolism (Hargreaves et al. 1995). In turn, impaired contractile activity of specific muscle fibres due to intramyofibrillar glycogen depletion may have increased the metabolic burden that a given power output placed on the smaller number of active, fully functional fibres. More specifically, greater burden may have been placed on less oxidative type IIAB and IIB later in the prolonged exercise bout, as evidenced by prior work on fibre type-specific glycogen depletion patterns during prolonged exercise (Vøllestad et al. 1984). These proposed effects of glycogen depletion on the moderate-to-heavy transition may
have been exacerbated by the exercise of the present study being conducted after an overnight fast and without carbohydrate intake during exercise, and may plausibly therefore be lessened in training and competition scenarios in which exercise is performed postprandially and with carbohydrate feeding. However, as muscle glycogen depletion, and more specifically compartmental muscle glycogen depletion, was not measured in this study, this mechanism remains speculative and could be interrogated in future work. Additionally, the importance of glycogen availability for durability of the moderate-to-heavy transition could be further explored through repetition of the present protocol with experimental manipulation of pre-exercise glycogen availability through exercise and/or nutrition interventions. Interestingly, the contribution made by decreased efficiency, but not decreased rates of metabolic energy expenditure at the transition, to prolonged exercise-induced changes in the moderate-to-heavy intensity transition power output was related to the magnitude of PRE to POST changes in whole-body fat oxidation rates; that is, those for whom decreases in energetic efficiency were large exhibited the largest PRE to POST increases in whole-body fat oxidation rates (Table 2). Larger increases in fat oxidation following prolonged exercise may reflect greater muscle glycogen depletion (Hargreaves et al. 1995), and glycogen depletion may as discussed negatively impact contractile function and in turn efficiency at the level of the muscle fibre (Ørtenblad et al. 2013). These data may therefore indirectly support that the loss of energetic efficiency with prolonged exercise was at least partially attributable to the degree of glycogen depletion. However, these data are associational and muscle glycogen was not measured in the present study, and so this mechanism should be interrogated directly in future work.

In contrast, the reduction in moderate-to-heavy intensity transition power output was not significantly associated with PFO (Table 1). The PFO is a marker of an individual’s capacity for fat oxidation during exercise (Maunder et al. 2018), meaning that having a greater capacity to oxidise fatty acids in a fresh state during exercise was not related to ‘durability’ of the moderate-to-heavy intensity transition. If muscle glycogen depletion was the primary mechanism behind the observed prolonged exercise-induced reduction in moderate-to-heavy intensity transition power output, one might have predicted that possessing a greater capacity to oxidise fatty acids during exercise would have mitigated this decline. The absence of a relationship between PFO and the durability of the moderate-to-heavy intensity transition therefore appears to counter this proposed mechanism. However, PFO may not completely reflect the degree of muscle glycogen depletion induced by the prolonged exercise, and therefore to test this hypothesis future work may consider replicating the design of the present study, but with measurements of muscle glycogen content.

As the power output at the moderate-to-heavy intensity transition decreased following acute prolonged exercise, these data suggest that using a well-rested assessment of power output at the moderate-to-heavy intensity transition for programming prolonged exercise risks inadvertent drift from the moderate into the heavy intensity domain. This may have implications for training prescription; specifically, drift into the heavy domain may extend the recovery required after sessions intended to be of moderate intensity and therefore low physiological stress (Seiler et al. 2007; Stanley et al. 2013). Similarly, training load models may need to consider accounting for the durability of intensity domain transitions to better quantify training load.

During the two-hour constant work-rate phase, heart rate increased by $8.2 \pm 2.7\%$ from 15 to 120 min (Fig. 2a). This may have been related to increases in core temperature and therefore cutaneous blood flow, progressive dehydration and therefore reduced stroke volume (Coyle and Gonzalez-Alonso 2001), as well as the increased metabolic demand.

### Table 2 Bivariate associations between PRE to POST changes in whole-body fat oxidation rates in steps 1, 2, and 3 of the incremental exercise tests with the contributions to prolonged exercise-induced changes in moderate-to-heavy intensity transition power output made

|                      | Δ FO step 1 (g min⁻¹) | Δ FO step 2 (g min⁻¹) | Δ FO step 3 (g min⁻¹) |
|----------------------|------------------------|------------------------|------------------------|
| Δ Efficiency (VT₁, W) | $r = -0.72$ (−0.90, −0.30) $P = 0.004$ | $r = -0.74$ (−0.91, −0.34) $P = 0.003$ | $r = -0.57$ (−0.85, −0.02) $P = 0.04$ |
| Δ Metabolic EE (VT₁, W) | $r = 0.36$ (0.21, 0.75) $P = 0.20$ | $r = 0.31$ (−0.26, 0.72) $P = 0.28$ | $r = 0.15$ (−0.44, 0.65) $P = 0.63$ |

Δ Efficiency = contribution made by changes in efficiency to prolonged-exercise-induced changes in the moderate-to-heavy intensity transition, Δ FO step 1–3 = change in whole-body fat oxidation rate from PRE to POST in step 1–3, Δ Metabolic EE = contribution made by changes in metabolic energy expenditure at the transition to prolonged-exercise-induced changes in the moderate-to-heavy intensity transition. Significant relationships ($P \leq 0.05$) are highlighted in bold.
of the fixed work rate (Fig. 2b). More importantly, and in contrast to our hypothesis, it was observed that the heart rate associated with VT1 significantly increased from PRE to POST (6.3 ± 5.8%, Fig. 4c). These data demonstrate that the cardiovascular drift that occurred with acute prolonged exercise was proportionally larger than the downward drift in the power output associated with the moderate-to-heavy intensity transition, and therefore that the heart rate associated with the moderate-to-heavy intensity transition increases over time during acute prolonged exercise. These results therefore suggest use of well-rested assessments of the heart rate at the moderate-to-heavy intensity transition to prescribe prolonged exercise may risk ‘undertraining’, or downward drift within the moderate-intensity domain over time.

In the present study there was inter-individual variation in the degree of reduction in moderate-to-heavy intensity transition power output following prolonged exercise, with the reduction in VT1 power output ranging from ~9–44 W (Fig. 3). This suggests that the durability of the moderate-to-heavy intensity transition is not a uniform characteristic between-athletes, and thus that profiling the effects of prolonged exercise at the individual level may be useful for capturing an endurance athlete’s physiological profile. Moderate strength relationships were observed between the initial VT1 power output, expressed as %V·O2peak, and the durability of the moderate-to-heavy intensity transition, although this relationship was not present when the moderate-to-heavy intensity transition was estimated using blood lactate data (Table 1). Future work may consider exploring the implications of this characteristic for endurance performance, and also the training-related and physiological characteristics that differentiate athletes with high vs. low durability.

In conclusion, the present investigation demonstrated prolonged moderate-intensity cycling significantly reduced the power output observed at the moderate-to-heavy intensity transition. This reduction was associated with decreased gross cycling efficiency and rates of metabolic energy expenditure at the transition. The heart rate associated with this transition increased following prolonged exercise. Therefore, it may be important for endurance athletes to understand how intensity transitions are affected by prolonged exercise at an individual level to refine physiological profiling, training prescription, and load monitoring.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s00421-022-05036-9.

Authors’ contributions JDS, EM, AEK, and DJP: conceived and designed the research. JDS and EM: conducted experiments and collected the data. JDS and EM: analysed the data. JDS and EM: drafted the manuscript. All authors read, revised, and approved the manuscript.

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Data availability Data are available from the corresponding author upon reasonable request.

Code availability None used.

Declarations

Conflict of interest The authors declare no competing interests associated with this manuscript.

Ethics approval The Auckland University of Technology Ethics Committee approved all procedures (21/253).

Consent to participate All participants provided written informed consent.

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