Primed exercise accelerates oxygen uptake kinetics during high-intensity cycle exercise in middle-aged individuals with type 2 diabetes

Joel Rocha\textsuperscript{1}, Norita Gildea\textsuperscript{2*}, Donal O’Shea\textsuperscript{3}, Simon Green\textsuperscript{4} and Mikel Egaña\textsuperscript{2}

\textsuperscript{1}Division of Sport and Exercise Sciences, Abertay University, Dundee, United Kingdom, \textsuperscript{2}Department of Physiology, School of Medicine, Trinity College Dublin, The University of Dublin, Dublin, Ireland, \textsuperscript{3}Endocrinology, St Columcille’s and St Vincent’s Hospitals, Dublin, Ireland, \textsuperscript{4}School of Science and Health, Western Sydney University, Sydney, AU-NSW, Australia

Background: The primary phase time constant of pulmonary oxygen uptake kinetics ($V\cdot O_2 \tau_p$) during submaximal efforts is longer in middle-aged people with type 2 diabetes (T2D), partly due to limitations in oxygen supply to active muscles. This study examined if a high-intensity “priming” exercise (PE) would speed $V\cdot O_2 \tau_p$ during a subsequent high-intensity cycling exercise in T2D due to enhanced oxygen delivery.

Methods: Eleven (4 women) middle-aged individuals with type 2 diabetes and 11 (4 women) non-diabetic controls completed four separate cycling bouts each starting at an ’unloaded’ baseline of 10 W and transitioning to a high-intensity constant-load. Two of the four cycling bouts were preceded by priming exercise. The dynamics of pulmonary $V\cdot O_2$ and muscle deoxygenation (i.e. deoxygenated haemoglobin and myoglobin concentration [HHb + Mb]), were calculated from breath-by-breath and near-infrared spectroscopy data at the vastus lateralis, respectively.

Results: At baseline $V\cdot O_2 \tau_p$ was slower ($p < 0.001$) in the type 2 diabetes group (48 ± 6 s) compared to the control group (34 ± 2 s) but priming exercise significantly reduced $V\cdot O_2 \tau_p$ ($p < 0.001$) in type 2 diabetes (32 ± 6 s) so that post priming exercise it was not different compared with controls (34 ± 3 s). Priming exercise reduced the amplitude of the $V\cdot O_2$ slow component (As) in both groups (type 2 diabetes: 0.26 ± 0.11 to 0.16 ± 0.07 L/min; control: 0.33 ± 0.13 to 0.25 ± 0.14 L/min, $p < 0.001$), while [HHb + Mb] kinetics remained unchanged.

Conclusion: These results suggest that in middle-aged men and women with T2D, PE speeds $V\cdot O_2 \tau_p$ likely by a better matching of $O_2$ delivery to utilisation and reduces the $V\cdot O_2$ As during a subsequent high-intensity exercise.

Keywords: near-infrared spectroscopy, oxygen extraction, cycling, exercise tolerance, oxygen uptake slow component
1 Introduction

Type 2 diabetes (T2D) is a major chronic condition with a concerning rapidly increasing global prevalence. Importantly, men and women with T2D demonstrate a consistent impairment in cardiorespiratory capacity reflected by a decreased peak oxygen uptake (VO\textsubscript{2peak}) (Green et al., 2015), that is an independent predictor of all-cause mortality (Wei et al., 2000). In addition, at the onset of moderate-intensity exercise a slowed primary phase time constant of pulmonary oxygen uptake (VO\textsubscript{2}) kinetics (VO\textsubscript{2} \( \tau \text{p} \)) is observed in young and middle-aged people with T2D (Bauer et al., 2007; Mac Ananey et al., 2011; O’Connor et al., 2012; Kiely et al., 2015; O’Connor et al., 2015). Similarly, recent evidence suggests that compared with controls, in middle-aged individuals with T2D VO\textsubscript{2} \( \tau \text{p} \) is also slowed during exercise transitions from a moderate-intensity baseline to high-intensity (i.e work-to-work transitions) (Gildea et al., 2021b). While VO\textsubscript{2} \( \tau \text{p} \) is a well-established key determinant of exercise tolerance (Jones and Poole, 2005; Goulding et al., 2021), the mechanisms for the constrained VO\textsubscript{2} \( \tau \text{p} \) in T2D remain to be elucidated, but accumulating evidence suggests that impairments in oxygen supply to the active musculature (Padilla et al., 2006; MacAnaney et al., 2011; Kiely et al., 2014) and a subsequent mismatch of local O\textsubscript{2} delivery to muscle VO\textsubscript{2} (Bauer et al., 2007; Gildea et al., 2019; Rocha et al., 2019; Gildea et al., 2021b) play an important role.

The impediments in VO\textsubscript{2} kinetics in T2D are also apparent during high-intensity exercise transitions initiated from rest or ‘unloaded’ baseline, with Brandenburg et al. (1999) showing a significantly slower mean response time (MRT) of the VO\textsubscript{2} kinetics in females with T2D compared with BMI-matched controls. Nevertheless, Mac Ananey et al. (2011) showed a non-significant tendency for a slower VO\textsubscript{2} MRT and VO\textsubscript{2} \( \tau \text{p} \) (\textasciitilde 13\% and \textasciitilde 5\% respectively) during high-intensity cycling transitions initiated from static rest in females with T2D of similar characteristics, compared with BMI-matched controls (Mac Ananey et al., 2011). Noteworthy, when transitions are initiated from static (instead of ‘unloaded’) rest, VO\textsubscript{2} \( \tau \text{p} \) has been shown to be \textasciitilde 15\% longer when the time delay is not constrained (Whipp et al., 1982) as was the case therein (Mac Ananey et al., 2011), potentially influencing their findings.

Importantly, abrupt or sudden transitions to high-intensity activity (i.e. running, cycling or stair climbing) from rest or very light activity are akin to those in daily life (such as commuting to work), so, there is a need to examine interventions that may enhance the VO\textsubscript{2} dynamic response during these exercise transitions in T2D. In this regard, studies in healthy active individuals presenting with an initial fast VO\textsubscript{2} \( \tau \text{p} \) show that an acute prior bout of heavy-intensity “priming” exercise (PE) does not alter VO\textsubscript{2} \( \tau \text{p} \) during subsequent high-intensity upright cycling exercise initiated from rest (Burnley et al., 2000; Burnley et al., 2001; Burnley et al., 2002a; Koppo and Bouckaert, 2002; Sahlin et al., 2005). This is likely because PE appears to facilitate muscle oxygen delivery rather than specific metabolic pathways and in these healthy active individuals VO\textsubscript{2} \( \tau \text{p} \) seems limited by the later (i.e. intracellular energetics) (Gerbino et al., 1996; Sahlin et al., 2005; Jones et al., 2006). However, PE accelerates the MRT of the overall VO\textsubscript{2} dynamic response in these healthy participants typically through an increase in the primary phase amplitude of the VO\textsubscript{2} (VO\textsubscript{2} \( A_p \)) and/or reducing the slow component amplitude of the VO\textsubscript{2}, the latter being potentially related to the reduced requirement for type II muscle fiber activation after PE (DiMenna et al., 2008). On the contrary, when VO\textsubscript{2} kinetics are further slowed as a direct consequence of impaired O\textsubscript{2} delivery and reduced perfusion pressure to active muscles, as is observed during supine or prone high-intensity exercise (Rossiter et al., 2001; Jones et al., 2006; Goulding et al., 2017), PE accelerates VO\textsubscript{2} \( \tau \text{p} \) in the respective subsequent bouts of high-intensity exercise, possibly due to improved blood flow distribution, and/or reduced muscle fatigue to active muscles (DiMenna et al., 2010).

Thus, considering that O\textsubscript{2} supply to the muscle seems to be constrained in individuals with T2D, and high-intensity priming exercise has been proposed as an intervention that can augment the delivery of O\textsubscript{2} to the muscle, we tested the hypothesis that PE would reduce VO\textsubscript{2} \( \tau \text{p} \) in a subsequent bout of high-intensity exercise initiated from unloaded exercise in this population. Given that alterations exist in muscle fiber type in the T2D skeletal muscle, with individuals with T2D possessing larger proportions of type II and lower proportions of type I fibers than controls (Marin et al., 1994), we also hypothesized that in individuals with T2D PE would reduce the VO\textsubscript{2} \( A_p \). To shed light on contributions of muscle fractional O\textsubscript{2} extraction to any PE-induced changes in VO\textsubscript{2} kinetics, this study measured rates of muscle deoxygenation (i.e., deoxygenated haemoglobin and myoglobin, [HHb + Mb]) using near-infrared spectroscopy (NIRS). In addition, the age of participants was limited to less than 60 years to control for the potential confounding effects of age on the T2D-induced effects on exercise tolerance (Wilkerson et al., 2011; O’Connor et al., 2015).

2 Methods

2.1 Participants and recruitment

A total of 22 individuals, 11 with T2D (7 men/4 women) and 11 healthy controls (7 men/4 women) volunteered and provided written informed consent to take part in this study (Table 1). Recruitment for the control group was undertaken from the general population, whilst individuals with T2D were recruited from outpatient diabetes clinics of two hospitals in Dublin (i.e. St. Vincent’s University Hospital and St. Columcille’s Hospital). Four of the women participating in this study were premenopausal (2 T2D and 2 Control) and four were...
TABLE 1 Physical characteristics, peak exercise values, and activity levels.

<table>
<thead>
<tr>
<th>Physical characteristics</th>
<th>Controls</th>
<th>T2D</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>11</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Sex (male, female), n</td>
<td>7, 4</td>
<td>7, 4</td>
<td></td>
</tr>
<tr>
<td>Age, yr</td>
<td>40 (18)</td>
<td>43 (14)</td>
<td>0.21</td>
</tr>
<tr>
<td>Stature, m</td>
<td>1.70 ± 0.08</td>
<td>1.71 ± 0.09</td>
<td>0.77</td>
</tr>
<tr>
<td>BMI, kg/m²</td>
<td>30.9 ± 3.9</td>
<td>30.8 ± 4.6</td>
<td>0.97</td>
</tr>
<tr>
<td>Body Mass, kg</td>
<td>88 (17)</td>
<td>92 (33)</td>
<td>0.97</td>
</tr>
<tr>
<td>Fat layer VL, mm</td>
<td>7.9 (7.4)</td>
<td>6.0 (2.5)</td>
<td>0.25</td>
</tr>
<tr>
<td>HbA1c, %</td>
<td>5.1 ± 0.1*</td>
<td>7.4 ± 1.7</td>
<td>0.02</td>
</tr>
<tr>
<td>FPG, mmol/L</td>
<td>4.3 ± 0.7*</td>
<td>8.6 ± 3.5</td>
<td>0.01</td>
</tr>
<tr>
<td>Time since diagnosis, yr</td>
<td>4.5 ± 3.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VO₂, L/min</td>
<td>2.50 ± 0.52*</td>
<td>1.94 ± 0.49</td>
<td>0.02</td>
</tr>
<tr>
<td>VO₂, mL.kg. ·min⁻¹</td>
<td>28.4 ± 6.6*</td>
<td>21.3 ± 3.59</td>
<td>0.01</td>
</tr>
<tr>
<td>PO₃peak, W</td>
<td>199 ± 53*</td>
<td>149 ± 43</td>
<td>0.03</td>
</tr>
<tr>
<td>Habitual physical activity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inactive, h/day</td>
<td>19.4 ± 1.6</td>
<td>18.4 ± 0.3</td>
<td>0.19</td>
</tr>
<tr>
<td>Light, h/day</td>
<td>3.7 ± 1.3</td>
<td>5.0 ± 0.3</td>
<td>0.06</td>
</tr>
<tr>
<td>MVPA, h/day</td>
<td>0.86 ± 0.60</td>
<td>0.69 ± 0.29</td>
<td>0.56</td>
</tr>
</tbody>
</table>

Data are means ± SD, for variables that were normally distributed and median with interquartile range in parentheses for variables that showed skewness and were not normally distributed in one or both groups. BMI, body mass index; VL, vastus lateralis; HbA1c, glycosylated haemoglobin; FPG, fasting plasma glucose; MVPA, moderate-to-vigorous physical activity; VO₂, oxygen uptake; PO₃, power output. *Significantly different from T2D (p < 0.05).

2.2 Experimental procedures

2.2.1 Overview

Upon successful completion of a treadmill stress test (Bruce protocol) at St Columcille’s Hospital, participants attended the testing laboratories on two separate occasions. The participants in the control group completed all tests in the University’s human performance laboratory whilst participants with T2D did so in the exercise testing facility at St Columcille’s Hospital. Visit one consisted of participants completing a maximal cycling test to exhaustion to measure peak oxygen uptake (VO₂peak). In visit 2, participants completed four exercise transitions from a baseline of 10 W (i.e., unloaded) to high-intensity, of which two were preceded by PE. Cycling tests were completed in the upright position on an Excalibur Sport cycle ergometer (Lode BV, Groningen, Netherlands). Participants were asked to avoid the consumption of caffeine, alcohol and non-prescribed nutritional supplements together with any arduous physical activities during the 24 h preceding each visit. Menstrual cycle was controlled for when scheduling the visits of the premenopausal participants in this study, with testing taking place during the mid-follicular phase of their menstrual cycle (days 5–12, self-determined).

2.2.2 Visit 1: Maximal cycling test to exhaustion

All participants completed a ramp incremental cycling test to volitional exhaustion with an initial work rate of 10 W for 2 min (i.e., ‘unloaded’ cycling) followed by a progressive increase in power output of 10–25 W/min (based on each individual’s physical activity level). Participants were required to maintain a constant cadence throughout the test, self-selecting a pedalling rate between 60 and 75 rpm. Test termination occurred when participants had a cadence reduction of 10 rpm for more than 5s. At the end of the test peak work rate was determined as the highest power output achieved, whilst VO₂ was determined as the highest VO₂ value (15-s average) attained. The V-slope method was used to determine VT (Beaver et al., 1986).

2.2.3 Visit 2: Four cycling exercise transitions

All participants completed four identical 9-min cycling exercise bouts transitioning from an ‘unloaded’ power output of 10 W (3 min) to a constant-load of 50% delta (50%Δ) (6 min). The latter intensity was determined from the results of the maximal cycling test by adding the 50% difference between the power outputs at VO₂ and VT to the power output at VT. All participants completed the four bouts of exercise in the same order. Two of the bouts (bouts 1 and 3) were carried out without PE (50%Δ unprimed) and two bouts (bouts 2 and 4) were completed preceded by PE (50%Δ primed) (Figure 1). The unprimed 50%Δ bouts were used as PE. A resting period of

postmenopausal (2 T2D and 2 Control) not undergoing hormone replacement therapy. All participants were non-smokers (not smoking during the previous 12 months) and physically inactive [≤150 min week⁻¹ of moderate-intensity (<ventilatory threshold, VT) exercise in the preceding 6 months] (McKay et al., 2022). The latter status was confirmed by participants wearing RT3 triaxial accelerometers (Stayhealthy Inc., CA) over a 5-day period (Rowlands et al., 2004). Participants’ time since diagnosis of T2D was between 2 and 10 years (mean ± SD = 4.5 ± 3.2 yrs) and had HbA1c levels below 10%. In addition, exclusion criteria included the use of β-blockers or insulin, clinical evidence of liver or renal disease, if they suffered from persistent proteinuria (urine protein >200 mg/dl) or had high creatinine levels (suggestive of renal disease, which can alter exercise performance); autonomic insufficiency/dysfunction, symmetrical neuropathy, abnormal cardiac function, or evidence of ischaemic heart disease. Regarding medication, two individuals in the control group were on statins whilst participants with T2D were treated with either oral (n = 9) and/or subcutaneous (n = 2) hypoglycaemic agents (only metformin, n = 5; metformin and sulphonylurea, n = 1; SGLT-2 inhibitors, n = 1; GLP-1, n = 1), as well as antihypertensive medication (ACE inhibitor, n = 3; ARBs, n = 2; CCBs, n = 3) and statins (n = 6). The study received ethical approval from both Trinity College Dublin (TCD), and St Vincent’s Healthcare research ethics committees.
12 min was used between the first and second bouts, and the third and fourth bouts; whilst a 45–60 min seated rest period was used between the second and third bouts. The longer resting period was important to ensure that participant’s physiological parameters returned to a baseline state so as not to affect subsequent VO$_2$ kinetics parameters. This was determined by measuring these parameters in a subgroup of 11 participants with T2D, albeit employing a single high-intensity transition which is consistent with previous reports in healthy active individuals (Burnley et al., 2006). Participants’ gas exchange/ventilatory variables, muscle oxygenation/deoxygenation and heart rate (HR) data were measured continuously throughout each exercise bout.

2.3 Measures

2.3.1 Pulmonary gas exchange and heart rate

Breath-by-breath data was continuously obtained during exercise by participants wearing a facemask that was connected to a metabolic gas analysis system (Inn scor, Inn nov ish A/S, Odense, Denmark). Parameters analysed were oxygen consumption, carbon dioxide production, respiratory exchange ratio and minute ventilation. Calibration of equipment was undertaken before each use according to the recommendations of the manufacturer. In addition, calibration of the system’s oxygen and photoacoustic sensors is undertaken periodically (every 6–12 months) by the manufacturer. A heart rate monitor was used to measure HR at 5 s intervals (Polar S610i, Polar Ltd., Finland).

2.3.2 Muscle deoxygenation and tissue oxygenation index

Muscle oxygenation (O$_2$Hb + Mb), deoxygenation (HHb + Mb) and tissue oxygenation index (TOI) data were acquired using a continuous wave NIRS system (Niro 200NX; Hamamatsu, Japan). This device uses the spatially resolved spectroscopy (SRS) technique and modified Beer-Lambert (MBL) principle. Detailed information about this technique and its application during exercise is available elsewhere (Ferrari et al., 2011). In the present study this measurement was undertaken in the vastus lateralis (VL) muscle of the participant’s right quadriceps given the VL is the primary locomotor muscle during leg cycling (Laplaud et al., 2006; Okushima et al., 2016). In order to ensure good quality signals, necessary skin preparation was undertaken involving shaving any hair present and cleaning the area with alcohol. After the skin was dried, the probes in their rubber holder were securely positioned on the muscle, between 10 and 16 cm above the femoral condyle using transparent adhesive tape. A dark elastic bandage was also used to further protect the probes from external light and movement. The depth of the area being measured is approximately one-half the distance between the emitter and the receiver probes (~1.5 cm). Therefore, ultrasound measurements of the skin and adipose tissue at the probe location were taken in all participants using the B-mode of a 2D ultrasound (Zonare Ultra Smart Cart, Software version 4.7, United States) to ensure that data collected was representative of the muscle tissue. This was confirmed with all participants having less than 1.5 cm of adipose tissue thickness at the probe location.

2.4 Data analysis

2.4.1 Oxygen uptake kinetics

The linear interpolation method was used to estimate second by second values from the breath-by-breath VO$_2$ data for each transition. Data was then aligned to ensure that the start of the exercise bout was time 0. To achieve a single smoothed averaged response for each participant, data were ensemble- and time-averaged into 5s bins. A biexponential (Eq. 1) function was then used to fit these responses:

$$VO_2(t) = VO_2\text{baseline} + A_1\left[1 - e^{-(t-TDp)/\tau_p}\right]F1 + A_2\left[1 - e^{-(t-TDm)/\tau_m}\right]F2$$

In the above function $VO_2(t)$ is VO$_2$ (absolute) at a given time (t); VO$_2$ baseline represents the mean VO$_2$ in the last 30 s of the “unloaded” cycling; Amplitudes (A), time delays (TD) and time constants (τ) for VO$_2$ primary (p) and slow component (s) phases are represented as $A_p$, $A_s$, $TD_p$, $TD_s$, $\tau_p$, and $\tau_s$ respectively. Time constant is the time that VO$_2$ takes to reach 63% of the amplitude of the corresponding phase. F1 and F2 are two conditional expressions that ensure the fitting of the phase is restricted to the period at and beyond the time delay associated with that phase. The initial 20 s of the VO$_2$ response data from the start of the cycling bout (i.e. 
cardiodynamic phase) were omitted but the TD, was allowed to vary freely so that the fit could be optimised (Murias et al., 2011). A monoexponential curve was fitted to calculate mean response time (MRT) and ascertain the overall VO₂ kinetics during high-intensity cycling irrespective of the different VO₂ phases. A weighted nonlinear least-squares regression (TableCurve 2D, Systat, United States) was used to fit all VO₂ response data. During the initial fit of the model, only data points within the 95% prediction interval were included. The average of the final 30 s of the VO₂ was calculated to represent the end of exercise VO₂ response (i.e. VO₂ End A). The latter was then used to calculate the absolute As, that is (VO₂ baseline + Ap) subtracted from VO₂ End A, while TD was constrained. The As was also calculated relative to the entire response (As/(Ap + As)). The end of exercise VO₂ gain, representing the functional gain of the overall response, was also calculated from the pre-transition level since the TD could be optimised (Murias et al., 2011). A two-way mixed model ANOVA [condition (unprimed, primed) x diabetes status (T2D, Control)] and the post hoc Tukey test were used to analyse all the dynamic response data for oxygen uptake, heart rate and muscle deoxygenation as well as TOI responses, $p < 0.05$ was used to determine statistical significance. Results from parametric analyses are presented as mean ± SD whereas non-parametric results are presented using median and interquartile ranges.

### 3 Results

#### 3.1 Participant characteristics

Unsurprisingly, individuals with T2D had significantly higher HbA₁c and fasting plasma glucose levels than healthy individuals (Table 1). Importantly, the T2D and control groups were matched according to sex distribution, age, BMI, body mass and activity levels.

#### 3.2 Peak exercise responses

Individuals with T2D had significantly lower absolute VO₂peak, VO₂peak normalised to body mass, and peak power output than healthy individuals (Table 1).

#### 3.3 Effect of priming exercise on oxygen uptake kinetics

The primed and unprimed dynamic response characteristics of VO₂ at high-intensity cycling exercise transitions for each group are presented in Table 2. The VO₂ responses for a representative individual with T2D and a healthy control are presented in Figure 2. Individual VO₂ τp and VO₂ As responses are provided in Figure 3. The unprimed VO₂ τp and MRT were significantly longer in individuals with T2D compared with healthy controls ($p < 0.001$ for both parameters). PE significantly reduced VO₂ MRT ($p < 0.001$) in the T2D and control groups; however, no group difference was present ($p = 0.053$) during the subsequent exercise transition (diabetes status × condition interaction, $p < 0.001$). PE also elicited a reduction in VO₂ τp in individuals with T2D ($p < 0.001$) but not in the healthy controls ($p = 0.98$) so that VO₂ τp was not different between groups after PE (diabetes status × condition interaction, $p < 0.001$). In addition, subsequent to PE, VO₂ As was reduced while baseline VO₂ was increased in both the T2D and healthy control groups (condition effect, $p < 0.001$ for both parameters).
TABLE 2 Dynamic response characteristics of oxygen uptake (VO2) at high-intensity cycling exercise transitions.

<table>
<thead>
<tr>
<th></th>
<th>Unprimed Controls</th>
<th>Type 2 diabetes</th>
<th>Primed Controls</th>
<th>Type 2 diabetes</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>11</td>
<td>11</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>Baseline VO2 L/min</td>
<td>1.02 ± 0.23</td>
<td>0.90 ± 0.16</td>
<td>1.18 ± 0.23*</td>
<td>0.96 ± 0.20*</td>
</tr>
<tr>
<td>VO2 A0, L/min</td>
<td>1.04 ± 0.41</td>
<td>0.79 ± 0.28</td>
<td>1.09 ± 0.38</td>
<td>0.80 ± 0.29</td>
</tr>
<tr>
<td>VO2 τp, s</td>
<td>34 ± 2</td>
<td>48 ± 6</td>
<td>34 ± 3</td>
<td>32 ± 6*</td>
</tr>
<tr>
<td>VO2 A∞, L/min</td>
<td>0.33 ± 0.13</td>
<td>0.26 ± 0.11</td>
<td>0.25 ± 0.14*</td>
<td>0.16 ± 0.07*</td>
</tr>
<tr>
<td>VO2 A∞%</td>
<td>24.5 ± 7.0</td>
<td>25.6 ± 10.5</td>
<td>18.5 ± 7.0*</td>
<td>17.6 ± 6.6*</td>
</tr>
<tr>
<td>VO2 TD*, s</td>
<td>121 ± 24</td>
<td>138 ± 35</td>
<td>130 ± 26</td>
<td>139 ± 40</td>
</tr>
<tr>
<td>VO2 end A, L/min</td>
<td>2.39 ± 0.61†</td>
<td>1.95 ± 0.40</td>
<td>2.52 ± 0.59†</td>
<td>1.93 ± 0.45</td>
</tr>
<tr>
<td>VO2 MRT, s</td>
<td>67 ± 5†</td>
<td>81 ± 13</td>
<td>57 ± 4*</td>
<td>64 ± 9*</td>
</tr>
<tr>
<td>End-exercise VO2 gain, mL.min⁻¹.W⁻¹</td>
<td>9.7 ± 1.6</td>
<td>10.1 ± 1.7</td>
<td>9.5 ± 1.2</td>
<td>9.2 ± 1.8</td>
</tr>
</tbody>
</table>

Data are means ± SD; n = no. of participants. A, amplitude; τ, time constant; end A, steady-state oxygen uptake (VO2) response; TD, time delay; p, primary phase; s, slow component phase.

*p < 0.05 vs. unprimed within the same diabetes status group (i.e. within controls or within Type 2 diabetes); †p < 0.05 vs. participants with type 2 diabetes within the same condition (i.e. within unprimed or primed).

3.4 Effect of priming exercise on Δ [HHb + Mb] kinetics and tissue oxygenation index responses

Table 3 presents the parameter estimates for Δ[HHb + Mb] kinetics and TOI responses whereas Figure 4 shows the Δ[HHb + Mb] responses for representative individuals. No statistical group difference was observed in the unprimed parameter estimates. PE resulted in significantly elevated levels of baseline Δ [HHb + Mb] in T2D (p < 0.001) and a tendency for higher levels in controls (p = 0.08) (group × condition interaction, p = 0.04). Participants with T2D showed a larger ratio of the modelled amplitudes of Δ [HHb + Mb]/ΔVO2 than controls (main effect of group, p = 0.01). None of the remaining [HHb + Mb] kinetics parameters were affected by PE. Estimates of baseline TOI as well as ΔTOI were higher during the primed bout in individuals with T2D (p < 0.01 for both) but not controls (p = 0.7 and 0.9 respectively) (group × condition interaction, p = 0.02 and 0.01 respectively).

4 Discussion

In agreement with our primary hypothesis, this study presents for the first-time evidence that in middle-aged individuals with T2D PE reduces VO2 τp during high-intensity exercise initiated from unloaded exercise without changes in the dynamic response of Δ [HHb + Mb]. In addition, consistent with our second hypothesis, PE significantly reduced the VO2 A∞ during the high-intensity exercise bout. Together, these priming exercise-induced effects rendered a reduction in the VO2 MRT response.

3.5 Effect of priming exercise on heart rate kinetics

The HR τ was significantly longer in individuals with T2D compared with healthy controls (main effect of group, p < 0.01), but PE did not affect HR τ in any of the groups (T2D unprimed: 56 ± 3 s, T2D primed: 55 ± 2 s; controls unprimed: 51 ± 3 s, controls primed: 50 ± 5 s). Baseline HR was higher in T2D (main effect of group, p = 0.04) and subsequent to PE it increased (main effect of condition, p < 0.001) in both groups (T2D unprimed: 109 ± 14 beats. min⁻¹, T2D primed: 118 ± 13 beats. min⁻¹; controls unprimed: 98 ± 9 beats. min⁻¹, controls primed: 106 ± 10 beats. min⁻¹). End-exercise HR was not different among groups, but it increased following PE (main effect of condition, p = 0.017) in both groups (T2D unprimed: 157 ± 12 beats. min⁻¹, T2D primed: 159 ± 13 beats. min⁻¹; controls unprimed: 156 ± 16 beats. min⁻¹, controls primed: 162 ± 15 beats. min⁻¹).

4.1 Effect of priming exercise on oxygen uptake τp

In the present study, VO2 τp responses observed during high-intensity upright cycling transitions were significantly amplified in individuals with T2D (48 s) than controls (32 s) leading to a longer MRT in T2D compared with controls. This is consistent with previous results from Brandenburg et al. (1999) who showed a significantly longer MRT during high-intensity cycling transitions in females with T2D compared with BMI-matched
controls (Brandenburg et al., 1999), although Mac Ananey et al. (2011) only observed a tendency for longer VO$_2$ kinetics in females with T2D of similar characteristics (Mac Ananey et al., 2011). Importantly, we have recently reported in a subgroup of participants who took part in the present study, that T2D slows VO$_2$ $\tau_p$ during transitions to both, moderate-intensity (Rocha et al., 2019; Rocha et al., 2020) as well as high-intensity work-to-work (Gildea et al., 2021b) transitions. Hence, the present study extends the findings of a blunted VO$_2$ $\tau_p$ in T2D when compared with carefully matched healthy controls.

The performance of a PE bout herein resulted in a subsequent significant reduction in this VO$_2$ $\tau_p$ among individuals with T2D but not in those without, bringing the primed VO$_2$ $\tau_p$ in T2D in line with the control group. These findings suggest that when the dynamic response of VO$_2$ is impaired by limitations in O$_2$ delivery, as is the case in T2D (Bauer et al., 2007; MacAnaney et al., 2011; Kiely et al., 2014), PE speeds VO$_2$ $\tau_p$ in the subsequent exercise bout. This notion is supported by studies that have explored these responses when exercising in the prone and supine positions (Rossiter et al., 2001; Jones et al., 2006; Goulding et al., 2017), thus, compromising exercising muscle perfusion pressure and O$_2$ delivery (Egaña et al., 2010; Egaña et al., 2013). For instance, an investigation where healthy participants performed high-intensity cycling bouts
with and without PE in the supine posture, Jones et al. (2006) observed that PE induced a 37% reduction ($p < 0.05$) in $\tau V \cdot O_2 p$ (38 s ± 18 s vs. 24 ± 9, s) in the subsequent bout, that was in line with that reported in the unprimed upright posture (Jones et al., 2006). Thus, findings from the current study expand the recently reported findings by our group of a significant speeding in $V \cdot O_2 \tau p$ following priming exercise during moderate-intensity (Rocha et al., 2019; Rocha et al., 2020) as well as high-intensity work-to-work exercise (Gildea et al., 2021b) to that of high-intensity exercise initiated from light exercise in individuals with T2D who are younger than 60 years of age.

The notion that priming exercise enhanced $O_2$ supply in the subsequent exercise transition in T2D is evidenced by an increased TOI which suggests an increase in $O_2$ availability, likely mediated by a PE-induced increased vasodilation and muscle blood flow at the beginning of the subsequent exercise (Gerbino et al., 1996). However, given that the NIRS-derived overall muscle deoxygenation kinetics and/or amplitude were not affected by PE herein, there is the possibility that the priming-induced reduction in $V \cdot O_2 \tau p$ in T2D was also partly mediated by an improved intracellular $O_2$ utilization, likely mediated by the upregulation of rate-limiting mitochondrial oxidative enzymes (Gurd et al., 2006; Gurd et al., 2009) and elevated mitochondrial calcium concentrations (Wüst and Stienen, 2018). On the other hand, the fact that HR kinetics were not altered following PE suggests that central mechanisms (i.e. quicker delivery) did not influence the priming response.

### 4.2 Effect of priming exercise on oxygen uptake slow component

In the present study, PE significantly reduced both $V \cdot O_2 \tau p$, and $A_s$ during the high-intensity transition in participants with

![FIGURE 3](https://via.placeholder.com/150)

**FIGURE 3**

Individual and mean ± SD (bar graph) changes in time constant of the primary phase of oxygen uptake ($V \cdot O_2 \tau p$) (A) and amplitude of the $V \cdot O_2$ slow component ($V \cdot O_2 A_s$) (B) in participants with type 2 diabetes ($n = 11$) and healthy controls ($n = 11$) during high-intensity cycling transitions without priming exercise (unprimed) and with priming exercise (primed). $p < 0.05$ vs. unprimed within the same diabetes status group (i.e., within controls or within Type 2 diabetes). $p < 0.05$ vs. healthy controls within the same condition (i.e., within unprimed or primed).

<table>
<thead>
<tr>
<th>TABLE 3 Dynamic response characteristics of Δ [HHb + Mb] and TOI during high-intensity cycling exercise transitions.</th>
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<tbody>
<tr>
<td><strong>Unprimed</strong></td>
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<tr>
<td><strong>Controls</strong></td>
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<tr>
<td>$n$</td>
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<tr>
<td>Baseline $\Delta$ [HHb + Mb], μM/cm</td>
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<tr>
<td>$\Delta$ [HHb + Mb] $A_p$, μM/cm</td>
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<tr>
<td>$\Delta$ [HHb + Mb] $TD_p$, s</td>
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<tr>
<td>$\Delta$ [HHb + Mb] $\tau_p$, s</td>
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<tr>
<td>$\Delta$ [HHb + Mb] $\tau_s$, s</td>
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<tr>
<td>Primary phase $\Delta$ [HHb + Mb]/$\Delta V \cdot O_2 A_s$, μM/cm (L/min)</td>
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<tr>
<td>$\Delta$ [HHb + Mb] $A_o$, μM/cm</td>
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<tr>
<td>Baseline TOI, %</td>
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<td>$\Delta$ TOI %</td>
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</tbody>
</table>

Values are means ± SD; $n$ = no. of participants, $A_p$ amplitude, $\tau$, time constant; $TD_p$ time delay; $p$, primary phase; $s$ slow component phase; $\tau$, time constant; $\tau'$, effective response time ($\tau + TD_p$), deoxygenated haemoglobin and myoglobin concentration; TOI, tissue oxygenation index; $V \cdot O_2$, oxygen uptake. $p < 0.05$ vs. unprimed within the same diabetes status group (i.e. within controls or within Type 2 diabetes). $p < 0.05$ vs. participants with type 2 diabetes within the same condition (i.e. within unprimed or primed).
T2D. However, in the control group, although PE reduced the V\textsubscript{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{o}}}}}}}}}}}}}}\text{O}_2$\text{~}τ\text{p}$ remained unaffected. These findings in control individuals are consistent with the current evidence on the influence of PE on subsequent transitions to heavy/severe-intensity upright cycling exercise initiated from an unloaded baseline in healthy participants (Burnley et al., 2000; Burnley et al., 2001; Scheuermann et al., 2001; Burnley et al., 2002a; Burnley et al., 2002b; Fukuba et al., 2002; Burnley et al., 2006; Jones et al., 2006; Goulding et al., 2017).

The priming-induced reduction in the slow component of the present study can likely be attributed to alterations in motor unit recruitment patterns. For instance, our group has recently shown (Gildea et al., 2021b) a priming-induced reduction in iEMG between the end of exercise and the time of the onset of VO\textsubscript{2} A\text{±}, (ΔiEMG\textsubscript{end-TD}) upon transition to a subsequent high-intensity cycling bout, albeit from an elevated baseline (work-to-work), concomitant with a significant reduction in the VO\textsubscript{2} A\text{±}. Although, herein, iEMG was not measured, it is possible that PE induced a decreased requirement for additional type II muscle fiber activation during the subsequent high-intensity cycling bout, thereby reducing the associated VO\textsubscript{2} cost of muscle fiber activation (DiMenna et al., 2008). By reducing dependency on these less efficient muscle fibers, the increase in sustained metabolic acidosis, a likely mediator of [PCr] and VO\textsubscript{2} slow components, can be slowed (Poole et al., 1988; Poole et al., 1991; Barstow et al., 1996; Rossiter et al., 2002; Krustrup et al., 2004). Alternatively, PE could facilitate an increased and more homogenous muscle perfusion within the active musculature, which is supported by the observed elevated baseline TOI during the primed bout in T2D herein. Consequently, the reliance on [PCr] degradation and glycogenolysis would be reduced, attenuating the rate of fatigue development and thus, delaying motor unit recruitment (DiMenna et al., 2010). In addition, this altered muscle activation response to priming exercise concomitant with the elevated TOI is consistent with the ‘oxygen-conforming’ effect, which has been demonstrated under involuntary and voluntary small muscle activation (Fitzpatrick et al., 1996; Drouin et al., 2022), although the mechanisms governing the oxygen-conforming response remain to be elucidated.

The observed PE-induced reduction in the VO\textsubscript{2} A\text{±} during high-intensity cycling in T2D, is all the more pertinent given individuals with T2D possess a 2-fold increase in type IIb fibers (Mogensen et al., 2007), demonstrate attenuated motor unit firing patterns in the VL compared with healthy controls (Watanabe et al., 2012; Watanabe et al., 2013) and tend to have lower dissociating capacity of myoglobin at intensities above VT (Miyamoto et al., 2020). Nevertheless, it is important to note that not all studies support the association between neuromuscular activation and the VO\textsubscript{2} slow component (Scheuermann et al., 2001; Garland et al., 2006; Cannon et al., 2007), and this is possibly due to the variability associated with measurements and normalization of iEMG.

With this new physiological insight of impaired VO\textsubscript{2} kinetics during high-intensity exercise transitions in T2D that are affected by limitations in O\textsubscript{2} delivery, future studies should investigate if exercise training mitigates these impairments. This will be practically relevant as high-intensity exercise transitions replicate metabolic transitions akin to those in daily life such as initiating sudden transitions to rapid walking, running, or stair climbing. While recent studies have demonstrated that time-efficient high-intensity interval training as well as longer-duration moderate-intensity continuous exercise training

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**FIGURE 4**

Changes in deoxygenated hemoglobin and myoglobin concentration [$\Delta (\text{HHb} + \text{Mb})$] for a representative healthy control (A) and an individual with type 2 diabetes (B) during high-intensity cycling transitions without priming exercise (open circles) and with priming exercise (solid circles). The vertical line illustrates the abrupt transition to the higher work rate. The continuous black lines of best fit illustrate the primary phase of the $\Delta (\text{HHb} + \text{Mb})$ response. Note the time constant of the primary phase of the $\Delta [\text{HHb} + \text{Mb}]$ response is not affected by prior priming exercise in any of the 2 groups.
interventions seem effective in enhancing VO₂ $\tau_p$ during moderate-intensity transitions (Green et al., 2020; Gildea et al., 2021a) as well as high-intensity work-to-work transitions (Gildea et al., 2022) in T2D, new studies should explore if these exercise training interventions of different doses can influence the VO₂ kinetics response during high-intensity transitions.

5 Limitations

Our results are limited to middle-aged mixed groups of men and women, hence, further studies should explore sex- and/or age-related differences in these outcomes. Even if our protocol did not allow block randomization, the sequence of the unprimed and primed exercise transitions was the same for all participants; hence, this likely has a minor impact on the interpretation of the current findings.

6 Conclusion

The present study showed that a preceding high-intensity exercise (i.e. warm-up) or priming exercise accelerated the overall MRT of the VO₂ dynamic response during high-intensity transitions in middle-aged individuals with T2D. This finding was attributed to a speeding of the primary phase time constant of VO₂ and a reduction in the amplitude of the VO₂ slow component while PE did not affect the dynamic response of muscle deoxygenation. Thus, in the presence of the likely diminished vasomotor responses in T2D, it is likely that undertaking a prior high-intensity exercise bout resulted in a more appropriate distribution of blood flow within the working muscle microvasculature, serving to alleviate the metabolic debacle to maintain VO₂.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

References


Ethics statement

The studies involving human participants were reviewed and approved by Faculty of Health Science Research Ethics Committee, Trinity College Dublin and St Vincent’s Healthcare Ethics and Medical Research Committee. The patients/participants provided their written informed consent to participate in this study.

Author contributions

JR, NG, DO’S, SG and ME conceived and designed the research and analyzed data. JR and NG conducted experiments and collected all participant data. NG and ME drafted the paper. All authors interpreted data and contributed to the writing of the final paper. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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prior sprint exercise and passive warming on oxygen uptake kinetics during heavy type 2 diabetes.

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exercise subsequent to HIIT versus moderate-intensity continuous training in Physiol. "work-to-work" cycling with a similar time-course in type 2 diabetes.

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