1	Title
2	Low-volume HIIT and MICT speed $\dot{V}O_2$ kinetics during high-intensity "work-to-work" cycling with a
3	similar time-course in type 2 diabetes.
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18	Running head:
19	HIIT vs MICT on $\dot{V}O_2$ kinetics during w-to-w exercise in T2D
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24

25 Abstract

26 We assessed the rates of adjustment in oxygen uptake (VO₂) and muscle deoxygenation (i.e., 27 deoxygenated haemoglobin and myoglobin, [HHb+Mb]) during the on-transition to high-intensity cycling 28 initiated from an elevated baseline (work-to-work) before training and at weeks 3, 6, 9 and 12 of low-29 volume high-intensity interval training (HIIT) and moderate-intensity continuous training (MICT) in type 30 2 diabetes (T2D). Participants were randomly assigned to MICT (n=11, 50 min of moderate-intensity 31 cycling), HIIT (n = 8, 10x1 min of high-intensity cycling separated by 1-min of light cycling) or non-32 exercising control (n=9) groups. Exercising groups trained 3 times per week. Participants completed two 33 work-to-work transitions at each time point consisting of sequential step increments to moderate- and 34 high-intensity work-rates. [HHb+Mb] kinetics were measured by near-infrared spectroscopy at the vastus 35 lateralis muscle. The pretraining time constant of the primary phase of $\dot{V}O_2$ ($\dot{V}O_2\tau_n$) and the amplitude of 36 the $\dot{V}O_2$ slow component ($\dot{V}O_2A_s$) of the high-intensity w-to-w bout decreased (P<0.05) by a similar 37 magnitude at wk 3 of training in both MICT (from, 56±9 to 43±6s, and from 0.17±0.07 to 0.09±0.05 38 L.min⁻¹, respectively) and HIIT (from, 56 ± 8 to $42\pm6s$, and from 0.18 ± 0.05 to 0.09 ± 0.08 L.min⁻¹, 39 respectively) with no further changes thereafter. No changes were reported in controls. The parameter 40 estimates of Δ [HHb+Mb] remained unchanged in all groups. MICT and HIIT elicited comparable 41 improvements in VO₂ kinetics without changes in muscle deoxygenation kinetics during high-intensity 42 exercise initiated from an elevated baseline in T2D despite training volume and time commitment being 43 ~50% lower in the HIIT group.

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45 New & Noteworthy

Three weeks of high-intensity interval training and moderate-intensity continuous training decreased the time constant of the primary phase of oxygen uptake ($\dot{V}O_2$) and amplitude of the $\dot{V}O_2$ slow component during a high-intensity exercise initiated from an elevated baseline, a protocol that mimics the abrupt metabolic transitions akin to those in daily life, in type 2 diabetes. These $\dot{V}O_2$ kinetics improvements 50 were maintained until the end of the 12-week intervention without changes in muscle deoxygenation

51 kinetics.

- 52
- 53 Keywords: exercise transitions, near-infrared spectroscopy, oxygen extraction, exercise tolerance,
- 54 oxygen uptake slow component

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58 Introduction

In healthy people, the initiation of a transition to high-intensity, constant work-rate upright cycling from moderate-intensity baseline cycling, referred to as work-to-work (w-to-w), elicits a significantly longer time constant of the primary phase of the oxygen uptake ($\dot{V}O_2$) kinetics response ($\dot{V}O_2 \tau_p$) than initiating the same transition from rest or 'unloaded' cycling (1-4). This prolonged $\dot{V}O_2 \tau_p$ translates to a compromised rate of oxidative energy transfer upon transition to the higher-intensity step of this protocol and has been attributed to constrained cellular respiration in the already active muscle fibers (5) and/or a larger recruitment of fast twitch (type II) muscle fibers to meet the augmented metabolic demand (6).

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67 Recently, Gildea et al. (7) observed that this slowing of $\dot{VO}_2 \tau_p$ during high-intensity w-to-w transitions is 68 significantly greater in middle-aged individuals with type 2 diabetes (T2D) compared with their healthy 69 counterparts, and that this effect is, at least in part, due to diabetes-induced limitations in peripheral 70 oxygen (O₂) delivery to the working muscle. This is in agreement with consistent observations of blunted or slowed $\dot{V}O_2$ τ_p responses during on-transitions to moderate-intensity exercise from an unloaded 71 72 baseline in young and middle-aged individuals with T2D (8-12), that also appear to be influenced by 73 impairments in O_2 delivery to active muscles (7, 12-16). W-to-w transitions replicate metabolic 74 transitions from moderate to higher metabolic rates akin to those in daily life (such as abrupt velocity 75 changes in walking/running/stair climbing, or changes in speed and/or gradient during cycling), and thus, 76 interventions that may enhance \dot{VO}_2 kinetics during w-to-w transitions in T2D are of great relevance and 77 warrant investigation. In this regard, short-term (\sim 12-weeks), traditional endurance training interventions, 78 involving ~150 min of continuous exercise per week [intensities ranging from ~60 to 80% maximum 79 heart rate (HR_{max})], have been shown to be effective at improving $\dot{V}O_2 \tau_p$ during moderate-intensity 80 transitions initiated from an unloaded baseline in T2D (17-19). However, to our knowledge the effect of 81 exercise training on $\dot{V}O_2 \tau_p$ during high-intensity w-to-w transitions in T2D has not been explored.

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83 Accordingly, the purpose of the current study was to investigate the effects of 12 weeks of two commonly 84 employed exercise training interventions, on VO₂ kinetics during high-intensity w-to-w cycling 85 transitions in individuals with uncomplicated T2D. Specifically, we compared the effects of moderate-86 intensity (< ventilatory threshold, VT) continuous training (MICT) with low-volume, high-intensity 87 interval training (HIIT), which typically involves ~75 min per week of intermittent vigorous exercise, 88 including less than 15 min of high-intensity efforts per session (20). Low volume HIIT was chosen for its 89 time efficient nature (~50% lower time commitment) given "lack of time" is frequently cited as a key 90 barrier for the well reported poor exercise adherence to current time-oriented physical activity guidelines 91 in T2D (21). While we have recently reported that low-volume HIIT and MICT elicit similar benefits in $\dot{V}O_2 \tau_p$ during moderate-intensity exercise transitions (22), interval training promotes greater oxidative 92 93 enzyme adaptations in type II fibers (23), which are predominantly recruited during high-intensity efforts and might be expected to result in faster $\dot{V}O_2 \tau_p$ responses during high-intensity exercise transitions. 94 95 Thus, we hypothesized that HIIT would be more effective at speeding VO₂ kinetics during the high-96 intensity bouts of the w-to-w transitions. In an attempt to explore the mechanistic basis of any exercise-97 induced effect on VO₂ kinetics in T2D, the rate of muscle deoxygenation (i.e., deoxygenated 98 haemoglobin and myoglobin, HHb+Mb) was measured to assess the alterations on muscle fractional O_2 99 extraction. In addition, to assess the time course effects of these adaptations, physiological measurements 100 were taken every 3 weeks throughout the intervention (i.e., before training and at weeks 3, 6, 9 and 12).

101

102 Methods

103 Participants

Participants were recruited from the Diabetes Outpatient Clinics of St. Columcille's and St. Vincent's University Hospitals (Dublin). Participant's eligibility was initially checked following chart review. Specifically, participants were included if they had a clinical history of diabetes < 11 yr, were sedentary [≤ 1.5 h/week of moderate-intensity exercise (<VT) and ≤ 1 structured exercise/week in the preceding 6 months, see *testing*] (24) and had HbA_{1c} levels of <10%. Participants were excluded if they were treated by exogenous insulin, were smokers, had a disease contraindicating physical training, or demonstrated evidence of renal, liver or cardiovascular disease. All individuals completed a 12-lead electrocardiogram treadmill stress test (Bruce protocol) at St. Columcille's Hospital prior to attending the laboratory tests.

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113 Thirty-four participants completed the baseline laboratory assessments (see testing) and were given 114 opaque sealed envelopes randomly allocating them to one of the 3 intervention groups (MICT, initially n115 = 13; HIIT, initially n = 9; or Control, initially n = 12). Eight participants dropped out of the study for personal reasons unrelated to the experiment (MICT, n = 2; HIIT, n = 3; Control, n = 3). Participants in 116 117 the Control group were offered re-randomization to one of the exercise training groups after the 118 intervention period, of which 2 accepted (HIIT, n = 2) and subsequently completed the training 119 intervention. The final study population consisted of 26 participants undergoing the intervention, of 120 whom 2 underwent both Control and HIIT. Thus, 28 completed responses from the study intervention 121 were included for statistical analysis (MICT, n = 11; HIIT, n = 8; Control, n = 9). All participants 122 provided written informed consent prior to participation. The study was approved by the Faculty of 123 Health Sciences' Research Ethics Committee, Trinity College Dublin, and St Vincent's Healthcare Ethics 124 and Medical Research Committee, and conducted in accordance with the principles outlined by the 125 Declaration of Helsinki.

126

127 Supervised exercise interventions

Overview. Participants in the HIIT and MICT groups carried out a 12-week supervised intervention, training 3 times per week on non-consecutive days at a local health and fitness center in Co. Dublin.
Participants in the Control group received no intervention and continued with their normal daily routine.
All exercise training sessions were supervised by a study investigator. Training intensity was adjusted at
3-week intervals (i.e., every 9 sessions) to reflect changes in fitness levels. Both exercise groups
completed a 5 min warm up and 5 min cool down before and after each session on an aerobic machine of their choice (elliptical, treadmill, rower or cycle ergometer). The main component of each training session
was completed on a cycle ergometer as follows:

136 Low-volume high-intensity interval training: The HIIT group completed 10 x 1-min bouts of high-137 intensity cycling interspersed with 1-min of light cycling. The high-intensity bout was completed at a 138 power output equivalent to 70% of the difference between participant's peak power output (PO_{peak}) and 139 the power output at ventilatory threshold (VT) (70% Δ) achieved during the ramp exercise test (see 140 *testing*), whereby participants were expected to exercise in the severe-intensity domain.

Moderate-intensity continuous training: Each MICT session comprised of 50 minutes of cycling at a power output equivalent to ~80% VT as calculated from the ramp test (see *testing*). The energy expenditure from the supervised exercise sessions was estimated based on the American College of Sports Medicine's equation (25).

- 145
- 146 Testing

147 Prior to the commencement of, and every 3 weeks throughout the intervention, participants were required 148 to attend the exercise testing laboratory on two separate occasions to complete a ramp incremental 149 cycling test to exhaustion, 3 high-intensity calf plantar-flexion transitions, 2-4 moderate- and high-150 intensity cycling exercise transitions, and 2 w-to-w step transitions to high-intensity cycling exercise 151 commencing from a baseline of moderate-intensity exercise. Data presented in the current manuscript are 152 based on the cycling high-intensity w-to-w step transitions. Data on peak exercise responses obtained 153 from the cycling ramp test (26) and moderate-intensity transitions (22) have been reported previously, 154 while data on calf plantar-flexion transitions are not presented herein. For each participant, all tests were 155 performed at the same time of day. All exercise tests were carried out in an upright position on an 156 electrically braked cycle ergometer (Excalibur Sport; Lode B.V., Groningen, Netherlands). Participants 157 were asked to refrain from consuming alcohol, caffeine and non-prescribed nutritional supplements as 158 well as avoiding any strenuous exercise in the 24 hours prior to testing. Prior to the intervention activity 159 levels were assessed by the use of 5-day RT3 triaxial accelerometry (Stayhealthy Inc, CA) (Table 1). The

160 threshold for sedentary or inactive behavior (<1.5 metabolic equivalents or METs) was set as < 100 161 counts/min, counts/min between 101 and 1317 were considered light activity (1.5-3 METs); and 162 counts/min >1317 corresponded to moderate-to-vigorous physical activity (>3 MET) (27). At baseline 163 (pretraining) and at the end of the intervention period (posttraining) fasting venous blood samples were 164 collected to assess glycosylated haemoglobin (HbA_{1c}). Participants were familiarized with the ramp 165 incremental test and constant work-rate tests prior to commencing the intervention.

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167 Ramp incremental cycling tests: The test started with an initial work-rate of 10 W for 2 min (i.e., 168 'unloaded' cycling). This was followed by a progressive increase in power output at 10-25 W/min based 169 on participants' activity levels. Pedalling rate was held constant at an individually selected cadence 170 between 60-75 revolutions per minute (rpm) and was maintained throughout all further testing. 171 Failure/exhaustion in a test was determined as a drop in cadence exceeding 10 rpm for >5 s. Peak work-172 rate was the power output achieved at the point of failure. \dot{VO}_{2peak} was the highest \dot{VO}_2 value (15-s 173 average) attained during the test. The first ventilatory threshold (VT) was determined using the V-slope 174 method (28); whereas the respiratory compensation point (RCP) was determined by identifying the second non-linear increase of \dot{V}_E and $\dot{V}CO_2$, whereby an increase in $\dot{V}_E/\dot{V}O_2$ is accompanied by an 175 176 increase of $\dot{V}_E/\dot{V}CO_2$ (29).

177

178 High-intensity work-to-work cycling exercise transitions. All participants performed two separate w-to-w 179 transitions to constant work-rate high-intensity cycling at 50% delta (Δ 50%; the sum of the power output 180 at VT and 50% of the difference between the power output at VT and $\dot{V}O_{2peak}$ obtained during the ramp 181 incremental test at the pretraining time point) each commencing from an elevated baseline of 80% VT 182 (80% of each participant's VT). Therefore, for each participant the same absolute power output was used 183 at all 5 time points during the intervention. The order of these bouts was fixed for all participants. Each 184 transition consisted of 3 min of "unloaded" cycling at 10W, immediately followed by 6 min of moderate-185 intensity (80% VT) cycling which in turn was immediately followed by 6 min of high-intensity ($\Delta 50\%$) 186 cycling. Exercise was performed continuously with changes in power output initiated as a step function 187 without giving prior warning to the individual. There was a 45-60 min rest period between each of the 188 cycling bouts. This resting period was sufficient for physiological parameters to return to baseline levels 189 and subsequently not to influence \dot{VO}_2 kinetics parameters (measured in a subgroup of 12 participants 190 with T2D, albeit employing a single high-intensity w-to-w transition), and this is consistent with previous 191 reports in healthy active individuals (30). Given that in the present study the mean response times of VO_2 192 during the ramp cycle exercise (31) were not accounted for when calculating these target power outputs, 193 power outputs at VT were overestimated. Five participants (MICT, n = 1; HIIT, n = 2; Control, n = 2) 194 failed to complete 6 min of exercise at $\Delta 50\%$ during the w-to-w bouts at baseline, so only physiological 195 responses collected over the same period (i.e., ≤ 6 min, range 3-5 min) during the subsequent time points 196 were analyzed. Heart rate (HR), gas exchange/ventilatory variables and muscle oxygenation & 197 deoxygenation were continuously measured during each cycling bout.

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199 Measurements

200 During exercise, participants wore a facemask to continuously collect expired air using an online 201 metabolic system (Innocor, Innovision A/S, Odense, Denmark) that measured airflow using a 202 pneumotachometer. Carbon dioxide analysis was performed by using a photoacoustic gas analyzer and 203 oxygen was analyzed using an oxygen sensor (Oxigraf Inc., USA) based on the principle of laser diode 204 absorption spectroscopy. The system was calibrated prior to each test as per manufacturer's 205 recommendations. Both the oxygen sensor and photoacoustic gas analyzer require multi-point calibration 206 that is routinely performed by the manufacturer every 6-12 months. Analysis of expired air allowed 207 determination of the rate of pulmonary O₂ uptake (\dot{VO}_2), CO₂ output (\dot{VCO}_2), minute ventilation (\dot{V}_F) and 208 the respiratory exchange ratio (RER) breath-by-breath. Heart rate (HR) was recorded every 5 s (Polar 209 S610i, Polar Ltd, Finland), with peak HR defined as the highest HR attained within the last 15 s of 210 termination of the test.

211 A continuous wave NIRS system (Hamamatsu Niro 200Nx; Hamamatsu Photonics, Hamamatsu, Japan), 212 was used to determine muscle oxygenation status non-invasively through the spatially resolved 213 spectroscopy technique and modified Beer-Lambert principle with three wavelengths of emitting light (λ 214 = 735, 810, and 850 nm). The theoretical basis of NIRS and its use in exercise measurements have been 215 described in detail elsewhere (32) but briefly, this technique estimates the optical density changes of 216 oxygenated (O₂Hb+Mb) and deoxygenated haemoglobin and myoglobin (HHb+Mb) based on the oxygen 217 dependency of absorption changes for near-infrared light in these proteins. As the vastus lateralis (VL) 218 muscle is a dominant locomotor muscle during cycling, the present study examined the concentration of 219 HHb+Mb (Δ[HHb+Mb]), and tissue oxygenation index (TOI) of the right vastus lateralis (VL) muscle. 220 After shaving, cleaning and drying the skin, the probes were placed on the belly of the muscle, 10-16 cm 221 above the lateral femoral condyle, parallel to the major axis of the thigh with a 3 cm spacing between the 222 emitter and receiver. The probes were housed in a black rubber holder and secured on the skin surface 223 with bi-adhesive tape and then covered with a dark elastic bandage, which minimized extraneous 224 movement and the intrusion of stray light throughout the exercise protocol. Since the depth of the 225 measured area was estimated to be approximately one-half the distance between the emitter and the 226 receiver (~1.5 cm), the present study determined the thickness of the skin and adipose tissue at the site of 227 the probe placement via 2D ultrasound operating in B-mode (Zonare Ultra Smart Cart, Software version 228 4.7, USA), to ensure that data largely represented absorption of near-infrared light in muscle tissue and 229 not in subcutaneous fat. All individuals presented with adiposity <1.5 cm over the site of interrogation on 230 the vastus lateralis.

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233 Data Analysis

 \dot{VO}_2 *Kinetics:* The breath-by-breath \dot{VO}_2 data for each transition were linearly interpolated to provide second-by-second values and time aligned such that time 0 represented the onset of exercise. Data from each transition were ensemble-averaged to yield a single, average response for each individual and further 237 time-averaged into 5 s bins (33). Data were then fitted to a monoexponential function (Eq. 1) or 238 biexponential function (Eq. 2). During the high-intensity exercise bouts responses were fitted to Eq. 2. 239 During the moderate-intensity bouts, the majority of the 140 responses (90%) consisted of a single 240 (primary) phase (visual inspection) and were fitted to Eq. 1. The remaining responses (10%) displayed a 241 second phase ("slow component") and were fitted to Eq. 2. This second phase was observed in 14 242 responses (from 9 participants, Control, n = 3; HIIT, n = 3; MICT, n = 3), had a mean amplitude of 76 243 mL/min (SD = 21 mL/min), was only observed among control participants beyond week 3 of the 244 intervention, and was likely due to the fact that in the present study the mean response times of $\dot{V}O_2$ 245 during the ramp cycle exercise were not accounted for when calculating the target power outputs (31). 246 The equations are as follows:

247

248 Equation 1
$$\dot{VO}_2(t) = \dot{VO}_2$$
 baseline $+ A_p[1-e^{(t-TDp)/\tau p}]F1$
249 Equation 2 $\dot{VO}_2(t) = \dot{VO}_2$ baseline $+ A_p[1-e^{(t-TDp)/\tau p}]F1 + A_s[1-e^{(t-TDs)/\tau s}]F2$

250

251 where $\dot{VO}_2(t)$ represents the absolute \dot{VO}_2 at a given time t; \dot{VO}_2 baseline (for moderate-intensity, in Eq's 252 1 & 2) is the mean \dot{VO}_2 in the final 30 s of unloaded cycling, whereas \dot{VO}_2 baseline (for high-intensity, in 253 Eq. 2) is the mean \dot{VO}_2 in the final 60 s of the moderate-intensity cycling exercise preceding the step 254 transition to high-intensity cycling exercise. Ap and As, are the amplitudes of the increase in VO2 for the 255 primary and slow component phases; TD_p and TD_s are the time delays of these phases, and τ_p and τ_s are 256 the time constants of the phases, defined as the duration of time for which VO_2 increases to a value 257 equivalent to 63% of the amplitude. The conditional expressions F1 and F2 limit the fitting of the phase to 258 the period at and beyond the time delay associated with that phase. The first 20 s of data after the onset of 259 exercise (i.e., the phase I VO₂ response) were deleted, while still allowing TD_p to vary freely (to optimize 260 accuracy of parameter estimates (34)). However, TD_s was constrained to avoid the possibility of 261 including the slow component in the modelled fit for the fundamental phase of VO2. VO2 data were 262 modelled from 20 s to 360 s of each step transition. The MRT was calculated through the fitting of a

monoexponential curve from exercise onset to provide information on the "overall" VO2 kinetics during 263 264 the high-intensity exercise bout, with no distinction made for the various phases of the response. The VO_2 265 data were fit using a weighted least-squares non-linear regression procedure (TableCurve 2D, Systat, 266 USA). Data points lying outside the 95% prediction interval during the initial fit of a model were 267 excluded. For moderate-intensity exercise, only estimates representing the primary phase are presented. 268 Whilst the presence of a slow component was detected in 14 responses during moderate-intensity exercise 269 transitions, the presence of this phase does not appear to significantly affect the parameter estimates of 270 the earlier phases (35). The end-exercise $\dot{V}O_2$ response, referred to as End A, was calculated as the 271 averaged $\dot{V}O_2$ over the last 30 s. Because the asymptomatic value (A_s) of the exponential term describing 272 the VO₂ slow component may represent a higher value than is actually reached at the end of the exercise, 273 the actual amplitude of the slow component was calculated as the absolute difference between the End A 274 and VO₂ baseline + A_p. The amplitude of the slow component was also described relative to the entire $\dot{V}O_2$ response [i.e., $A_s / (A_p + A_s)$]. The functional "gain" of the primary $\dot{V}O_2$ response (G_p) was 275 276 calculated as the difference between $\dot{V}O_2$ A_p and $\dot{V}O_2$ baseline normalized to the difference in power 277 outputs between the moderate-intensity exercise and unloaded cycling; and the functional gain of the 278 entire response at the end of the high-intensity exercise bout (i.e., end-exercise gain) was calculated in a 279 similar manner.

280

281 [HHb+Mb] kinetics and TOI. To provide information on muscle deoxygenation throughout the protocol, 282 we modelled the [HHb+Mb] response for moderate- and high-intensity exercise.-As per the $\dot{V}O_2$ data, the 283 NIRS-derived Δ [HHb+Mb] data for each transition were linearly interpolated to provide second-by-284 second values and time aligned. Data from each transition were ensemble-averaged to yield a single 285 average response for each individual, and further time-averaged into 5 s bins. A time delay (TD) at the 286 onset of exercise occurs in the [HHb+Mb] profile before it increases with an exponential like time course 287 (36). This was determined in the present study via visual inspection as a systematic increase above the 288 pre-transition level. For the moderate-intensity transitions, [HHb+Mb] data were fitted from the end of 289 this TD to 180 s using a monoexponential (Eq. 1) function as per VO_2 . The shorter fitting window of 180 290 s was selected to counteract the previously reported variations in the NIRS signal between 180-360 s from 291 exercise onset (also observed herein), from impacting the fitting of the on-transient response whilst 292 permitting the reaching of a steady-state (37, 38). For the high-intensity transitions, [HHb+Mb] data were 293 fitted from the end of the TD to the end of the exercise bout using a biexponential (Eq. 2) function as per 294 VO₂. For the moderate- and high-intensity exercise, the time course for the primary phase of the 295 Δ [HHb+Mb] response, referred to as the effective response time ($\tau'\Delta$ [HHb+Mb]), was determined from 296 the sum of the TD and τ from the onset of exercise. The amplitude change in TOI (TOI A) was calculated 297 as the difference between baseline (30 s prior to each transition) and end-exercise (final 30 s) values.

298

299 Statistical Analysis

300Physical characteristics and activity levels at baseline among groups were compared using a one-way301ANOVA. Peak physiological responses, training intensity, TOI values and kinetics parameter estimates302for $\dot{V}O_2$ and [HHb+Mb] throughout the intervention were compared using a two-factor [time (pretraining,303week 3, week 6, week 9, posttraining) vs. group (HIIT, MICT, CON)] mixed ANOVA. Body mass and304HbA_{1c} results were also compared using a two-factor [time (pretraining, posttraining) vs. group (HIIT,305MICT, CON)] mixed ANOVA. Differences were detected using a Student-Newman-Keuls *post hoc* test.306Significance was set at P < 0.05. All values are expressed as mean \pm standard deviation (SD).

307

308 Results

309 *Physical characteristics, pretraining peak exercise values and activity levels.*

310 Participants' physical characteristics, peak exercise values and activity levels at baseline are presented in

311 Table 1. HbA_{1c} (%) (time x group interaction, P < 0.012) was reduced in the MICT (pre = $6.9 \pm 0.5\%$,

312 post = $6.6 \pm 0.5\%$) and HIIT groups (pre = $7.3 \pm 0.5\%$, post = $7.0 \pm 0.6\%$) but not in the control (pre = 6.8

313 $\pm 1.0\%$, post = 7.0 $\pm 1.0\%$) group.

314

315 *Exercise adherence and caloric expenditure*

316 The mean exercise adherence was $94 \pm 6\%$ (range 31-36 sessions) and $97 \pm 4\%$ (range 32-36 sessions) in 317 the HIIT and MICT groups respectively. The average training intensity (power output) increased 318 significantly (P < 0.05) every 3 weeks (i.e. after each laboratory testing session) in the MICT group 319 (weeks 1–3, 84 ± 33 W; weeks 4–6, 102 ± 39 W; weeks 7–9, 113 ± 43 W; weeks 10-12, 122 ± 44 W) 320 while it also significantly increased every 3 weeks until week 9, but not between week 9 and 12 in the 321 HIIT group (weeks 1–3, 176 ± 35 W; weeks 4–6, 192 ± 37 W; weeks 7–9, 203 ± 38 W; weeks 10-12, 206 322 \pm 40 W). The average energy expenditure and total work done per training session (including the warm 323 up) was ~228 kcal and ~165 kJ for the HIIT group, and ~478 kcal and ~326 kJ for the MICT group. No 324 adverse training effects to training were observed throughout the intervention period in either exercising 325 group.

326

327 \dot{VO}_{2peak} from ramp incremental cycling

There was a significant time x group interaction (P < 0.001) for absolute $\dot{V}O_{2peak}$, so that $\dot{V}O_{2peak}$ did not increase in the control group ($\dot{V}O_{2peak}$ at pretraining = 1.86 ± 0.52 L/min), but it significantly increased after 3 weeks of MICT (from 2.08 ± 0.68 to 2.39 ± 0.68 L/min) and HIIT (from 2.42 ± 0.44 to 2.61 ± 0.47 L/min), with no further significant changes thereafter ($\dot{V}O_{2peak}$ at posttraining = 2.55 ± 0.73 L/min and 2.71 ± 0.54 L/min, respectively). Additional peak physiological responses have been reported in a companion paper (26).

334

335 *VO*₂ kinetics and NIRS-derived responses during high-intensity exercise of the w-to-w transition

The parameter estimates of the $\dot{V}O_2$ kinetics response for the high-intensity exercise bouts throughout the intervention period are shown in Table 2, and responses for representative individuals are shown in Fig 1. Individual $\dot{V}O_2$ τ_p and $\dot{V}O_2$ A_s responses throughout the intervention period are shown in Fig 2. Pretraining $\dot{V}O_2$ τ_p and MRT values were similar among the 3 groups. After 3 weeks of training, $\dot{V}O_2$ τ_p and MRT were significantly reduced in both the HIIT and MICT groups with no further significant 341 changes thereafter. In contrast, $\dot{V}O_2 \tau_p$ and MRT were not changed throughout the 12-week period in the 342 control group (time x group interaction, P < 0.01). Similarly, $\dot{V}O_2 A_s$ was significantly reduced after 3 343 weeks of MICT and HIIT with no further changes thereafter, but it did not change (time x group 344 interaction, P < 0.01) in the control group. The $\dot{V}O_2 A_p$ or the functional $\dot{V}O_2$ gain were not different 345 among groups and did not change throughout the intervention.

346

The kinetics parameters for Δ [HHb+Mb] as well as TOI values are displayed in Table 3 and Δ [HHb+Mb] responses for representative individuals are shown in Fig 3. The effective response times of muscle deoxygenation (Δ [HHb + Mb] τ_{p} , Δ [HHb +Mb] A_{p} , Δ [HHb +Mb] A_{s} and the ratio of the modelled amplitudes of the primary phase Δ [HHb + Mb]/ Δ VO₂ were not different among groups and did not change throughout the intervention (Table 3). The magnitude of the change in TOI during the highintensity exercise transitions was not affected by the intervention in either group.

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354 *VO*₂ kinetics and NIRS-derived responses during moderate-intensity exercise of the w-to-w transition

355 The parameter estimates of the $\dot{V}O_2$ kinetics response for the moderate-intensity exercise bouts 356 throughout the intervention period are shown in Table 2. For $\dot{V}O_2 \tau_p$, there was a significant time x group 357 interaction (P < 0.001), so that $\dot{VO}_2 \tau_p$ did not change in the control group, but it was reduced after 3 358 weeks of MICT and HIIT with no further changes thereafter. There was a main effect of group (P < 0.05) 359 for $\dot{V}O_2$ A_p so that it was larger in the HIIT group compared with the other 2 groups. Kinetics parameters 360 for Δ [HHb+Mb] as well as TOI values are displayed in Table 3. Exercise training did not affect the 361 effective response time of the Δ [HHb+Mb] response or the ratio of the modelled amplitudes of the 362 Δ [HHb + Mb]/ Δ VO₂ in either group. There was a main effect of group (P < 0.05) for Δ [HHb + Mb] A_p so 363 that it was larger in the HIIT compared with the control groups. The magnitude of the change in TOI 364 during the moderate-intensity exercise transitions were not affected by the intervention in either group, 365 and they were larger in the HIIT compared with the other 2 groups (main effect, group, P = 0.025).

366

367 **Discussion**

368 To our knowledge this is the first study to investigate the time-course effects of low-volume HIIT and 369 MICT on \dot{VO}_2 kinetics during high-intensity exercise initiated from an elevated baseline in individuals 370 with uncomplicated T2D. The principal findings were that both HIIT and MICT significantly reduced $\dot{V}O_2 \tau_p$ as well as the amplitude of $\dot{V}O_2 A_s$ during the transition to high-intensity cycling by week 3 of 371 372 training and that these effects occurred in the absence of changes in the dynamic response of Δ [HHb+Mb] 373 suggesting an improved microvascular blood flow delivery. In contrast with our hypothesis, these 374 adaptations were of a magnitude that was not different between exercising groups and were maintained 375 without further improvements until the end of the 12-week intervention period.

376

377 *Time-course effects of exercise training on* $\dot{V}O_2 \tau_p$ *during high-intensity exercise of the w-to-w transition* 378 In the present study, despite training volume and time commitment being ~50% lower in the HIIT 379 compared with the MICT group, both interventions significantly reduced $\dot{V}O_2 \tau_p$ after the 12-week 380 intervention period (31% MICT; 35% HIIT), with the reductions already apparent at the 3-week time 381 point (24% MICT; 26% HIIT). While in a companion paper of the current investigation we have recently 382 shown that the performance of both HIIT and MICT interventions elicit rapid (i.e., within 3 weeks) 383 adaptations in $\dot{V}O_2 \tau_p$ during transitions to moderate-intensity efforts from an unloaded/resting baseline in 384 T2D (22), herein we report for the first time the effects of these interventions on \dot{VO}_2 kinetics upon step 385 transitions to high-intensity exercise initiated from elevated metabolic rates in T2D. Among healthy participants, a number of studies have shown that HIIT and MICT interventions speed $\dot{V}O_2 \tau_p$ during 386 387 transitions to moderate- and high-intensity efforts from an unloaded baseline (39-43); but to our 388 knowledge, only one previous study has assessed VO2 kinetics responses during severe-intensity 389 transitions initiated from a moderate-intensity baseline following HIIT and/or MICT. Specifically, 390 consistent with our findings, Da Boit et al. (44) reported significant reductions in $\dot{VO}_2 \tau_p$ (26% and 22%) 391 subsequent to 2 weeks of either repeated sprint training (RST) (4-7, 30 s 'all-out' sprints interspersed by 4 392 mins of recovery) or MICT (60-110 mins cycling at 90% VT). Additionally, in agreement with Da Boit

et al. (44) albeit during w-to-w exercise in the moderate-intensity domain (i.e. transitions from 45% VT to 90% VT), Williams et al. (45) reported a 40% reduction in $\dot{V}O_2 \tau_p$ (45s to 25s) in healthy untrained young males subsequent to 4 weeks of HIIT (8-12 1 min cycling intervals at 110% WR_{max} interspersed by 1 min of unloaded cycling).

397

398 In the present study the observed speeding of VO₂ kinetics occurred without changes in the adjustment of 399 muscle deoxygenation suggesting that these training-induced reductions in $\dot{V}O_2 \tau_p$ could partly be due to 400 an improvement in microvascular O_2 delivery and/or enhanced intracellular O_2 utilization. Similarly, we 401 have recently reported that the accelerated $\dot{V}O_2 \tau_p$ responses during transitions to moderate-intensity 402 exercise following both HIIT and MICT in T2D were accompanied by no changes in [HHb + Mb] 403 kinetics and with a simultaneous reduction in the normalized Δ [HHb + Mb]/ Δ VO₂ ratio, indicative of an 404 increase in O₂ delivery relative to utilization within the microvasculature (22). These findings are also in 405 agreement with Williams et al. (45) who showed in healthy individuals that the enhanced $\dot{V}O_2 \tau_p$ upon 406 transition to w-to-w exercise in the moderate-intensity domain following HIIT was induced without 407 changes in the adjustment of local muscle deoxygenation. It is possible that training enhanced blood flow 408 kinetics and local blood flow distribution contributed to the faster VO2 kinetics. In this regard, substantial 409 evidence exists to suggest that T2D is associated with impairments in the dynamic response of 410 vasodilation (13, 16) and matching of capillary blood flow to metabolism (46) in contracting myocytes, 411 while a short term continuous endurance training intervention enhances leg vascular conductance kinetics 412 at low contractile intensities, at least in females with T2D (47). This is consistent with previous reports of 413 healthy populations showing faster conduit artery blood flow kinetics subsequent to a continuous aerobic 414 training intervention (48).

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416 Effect of exercise training on $\dot{V}O_2A_s$ during high-intensity exercise of the w-to-w transition

417 Alongside reductions in $\dot{V}O_2 \tau_{p}$, both training interventions also significantly reduced the amplitude of the

418 $\dot{V}O_2$ slow component ($\dot{V}O_2 A_s$) during the high-intensity bout of the w-to-w transition at the same 3-week

419 time point and remained that way until the end of the interventions. This is in contrast to findings in 420 healthy individuals, whereby 2 weeks (i.e. 6 exercise sessions) of either RST or MICT did not elicit any 421 changes in the VO₂ A_s during severe-intensity work-to-work transitions despite eliciting significant reductions in $\dot{V}O_2 \tau_p$ (44). However, during transitions from unloaded to severe-intensity exercise, Bailey 422 423 et al. (40) reported that only 2 weeks of RST (4-7, 30 s 'all-out' sprints interspersed by 4 min rest), but 424 not MICT (cycling at 90% VT for a duration that resulted in an equal work volume to RST), were 425 sufficient to reduce $\dot{V}O_2 A_s$ in healthy individuals. The fact that $\dot{V}O_2 A_s$ is larger in severe-intensity 426 transitions initiated from a resting baseline compared with a moderate-intensity baseline might suggest 427 the potential to reduce $\dot{V}O_2 A_s$ in the former, is likely larger. Therefore, authors suggested that a longer 428 duration training programme may be needed to allow for training induced adaptations in the VO₂ A_s 429 during the moderate to severe-intensity w-to-w transitions (44). On the other hand, in agreement with our 430 findings, 6 weeks of both HIIT (20 x 1-min at 90% VO_{2peak} interspersed by 60s rest) or continuous 431 aerobic training (30 mins at 60% VO_{2peak}) significantly decreased the amplitude of the VO₂ A_s (from 0.41 to 0.30 L.min⁻¹; and from 0.38 to 0.29 L.min⁻¹, respectively) during severe-intensity exercise initiated 432 433 from an unloaded cycling baseline in healthy populations (43).

434

The exercise-induced changes in the amplitude of the $\dot{V}O_2 A_s$ herein can be, at least partially, attributable to adaptations in the skeletal muscle properties given the working skeletal muscle accounts for ~80% of the $\dot{V}O_2 A_s$ (49). In this regard, in individuals with T2D, short-term continuous aerobic training has been shown to significantly increase oxidative enzyme activity and mitochondrial size and protein content (50). On the other hand, 2 weeks of low-volume HIIT also increased mitochondrial oxidative activity (51) and, stimulated activity of peroxisome-proliferator activated Υ coactivator (PGC-1 α), shown to regulate mitochondrial content and respiration in diabetic skeletal muscle.

442

We had hypothesized that HIIT would speed $\dot{V}O_2 \tau_p$ to a greater extent than MICT given that during HIIT a greater proportion of type II muscle fibers are recruited during the repeated intervals above the VT. This 445 would induce greater oxidative enzyme adaptations (23) compared with MICT, that predominantly 446 involves the recruitment of Type I oxidative muscle fibers. However, this was not the case herein, as both 447 interventions speeded $\dot{V}O_2 \tau_{p_s}$ (as well as reduced the $\dot{V}O_2 A_s$) by a magnitude not different among them. 448 Importantly, participants herein were exercising at a lower relative exercise intensity at each testing 449 timepoint throughout the interventions compared with pretraining, which likely reduced the proportion of 450 type II fibers recruited. Similarly, in healthy populations, both continuous endurance training and HIIT 451 interventions that provide sufficient stimulus for adaptation have also been shown to be equally effective 452 at speeding VO_2 kinetics during high-intensity transitions initiated from a moderate- intensity (44) or 453 resting (43) baseline, as well as during moderate-intensity transitions initiated from a resting baseline 454 (41). As herein, in these studies participants used the same absolute power output during exercise 455 transitions at all testing time points. Thus, it is plausible that in the present study both training 456 interventions provoked rapid increases in the oxidative capacity of Type I and II fibers and/or stimulated 457 phenotypical shifts in type II muscle fibers, or indeed mechanism intrinsic to individual muscle fibers, 458 and as such improved mitochondrial function or respiratory capacity. Such improvements would 459 plausibly serve to improve metabolic stability, and subsequently negate the need to recruit higher level 460 glycolytic fibers and thus, reducing the amplitude of the $\dot{V}O_2 A_s$.

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462 Effect of exercise training on $\dot{VO}_2 \tau_p$ during moderate-intensity exercise of the w-to-w transition

463 Consistent with findings from our recent companion paper (22), HIIT and MICT accelerated the $\dot{V}O_2$ 464 kinetics during the moderate-intensity exercise transition after 3 weeks of training with no additional 465 changes thereafter, while there were no changes in any $\dot{V}O_2$ parameters in the control group. In addition, 466 muscle deoxygenation kinetics responses were not altered throughout the intervention, suggestive of an 467 improvement in the balance of O_2 delivery and utilization being a likely underlying mechanism of the 468 accelerated $\dot{V}O_2$ kinetics. Indeed, it is likely that the training-enhanced $\dot{V}O_2$ kinetics and possibly the 469 metabolic/fatiguability of muscle during the moderate-intensity baseline, contributed to speeding the $\dot{V}O_2$ 470 kinetics of the high-intensity transition and reducing the fatigue-related and time-dependent increase in 471 motor unit recruitment which underpins the slow component of $\dot{V}O_2$ during high-intensity exercise.

472

473 Limitations

474 A number of limitations of the present study must be acknowledged. First, the NIRS-derived findings 475 herein relate to a single muscle, the VL, and therefore, interpretation of these data is limited to the site of 476 interrogation (i.e. superficial sample of the VL). The established heterogeneity extant within a single 477 muscle in terms of vascularity and fiber type, fiber recruitment, vascular control, and blood flow (52), 478 likely extends to the VL, as well as the temporal and spatial heterogeneity in NIRS-derived responses 479 extant both among and within muscles (53). Second, five participants did not complete the required 6 min 480 of high-intensity cycling exercise during the w-to-w transitions at the pre-training time point. However, 481 we believe this had little influence on the interpretation of our findings given that all participants showed 482 a clear $\dot{V}O_2$ slow component phase, they were similarly distributed among groups (2-3 in each group) and 483 only physiological responses collected over the same period during the subsequent time points were 484 analyzed. In this regard, future studies should attempt to identify each individual's critical power to 485 confirm that high-intensity exercise transitions were carried out within the same intensity domain for all 486 participants (i.e. heavy or severe domain). Third, while in 14 responses (from 9 participants) a small VO₂ 487 slow component phase was observed during the moderate-intensity transitions, these participants were 488 also similarly distributed among groups (3 in each group), thus, the influence on the interpretation of the 489 current findings is likely minor. Fourth, herein the VO₂ slow component was estimated using a second 490 exponential response (Eq 2), but it is also a common practice to identify the onset of the slow component 491 by fitting a monoexponential equation (Eq1) up to the point where residuals deviate from Gaussian 492 distribution. We therefore carried out additional analyses to identify the onset of the slow component in 493 line with the latter method, and these estimates coincided very closely (not shown) with subsequent 494 statistical outcomes unaffected. Finally, given that the current study is the first to report training-induced 495 changes in VO₂ kinetics during high-intensity w-to-w transitions in T2D, the overall trial was powered to

496 detect changes in $\dot{V}O_{2peak}$ (26), so, we cannot exclude the possibility that the limited number of 497 participants that completed the study precluded the observation of additional benefits in $\dot{V}O_2$ kinetics 498 beyond the 3rd week of training.

499

500 Conclusions

501 The present study primarily demonstrated that both HIIT and MICT are safe and effective interventions 502 that accelerate the VO₂ kinetics response during high-intensity exercise initiated from an elevated baseline 503 in individuals with uncomplicated T2D. Both forms of training induced a reduction in the amplitude of 504 the $\dot{V}O_2$ A_s and an acceleration of $\dot{V}O_2$ τ_p without changes in [HHb + Mb] kinetics responses. 505 Improvements in O_2 delivery during exercise are likely to have contributed to the observed reduction in 506 $\dot{V}O_2 \tau_p$ with training, while the reduction in the amplitude of the $\dot{V}O_2 A_s$ may have been caused by 507 exercise-induced changes in skeletal muscle properties and motor unit recruitment patterns. From a 508 practical perspective, investigating the training effects on the w-to-w protocol is of great relevance as it 509 mimics the abrupt metabolic transitions akin to those in daily life such as abrupt walking/running/stair 510 climbing velocity changes when for instance, people need to arrive on time to a place or an appointment. 511 Moreover, individuals with T2D are being encouraged to actively commute to work by healthcare 512 practitioners given the effectiveness of active commuting to improve body composition and 513 cardiovascular health. In this regard, when people cycle to work, sudden changes in gradient and/or speed 514 also mimic the w-to-w protocol investigated in the present study. Furthermore, given individuals with 515 T2D perceive even light to moderate exercise as being more difficult than healthy counterparts (42), the 516 perception of these w-to-w transitions is also likely harder which can ultimately result in a more sedentary 517 lifestyle. Therefore, the present study yields promising results supporting the efficacy of time-saving low-518 volume HIIT in eliciting increases in exercise tolerance given a faster provision of aerobic metabolism 519 will serve to reduce muscle fatigue during abrupt w-to-w transitions.

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524

525 Disclosures

- 526 No conflicts of interest, financial or otherwise, are declared by the authors.
- 527

528 Author Contributions

- 529 N.G., J.R., D.O., S.G., and M.E. conceived and designed research; N.G., A.M., D.C., A.N., and J.R.
- 530 performed experiments; N.G., A.M., and M.E. analyzed data; N.G., A.M., S.G., and M.E. interpreted
- 531 results of experiments; N.G. and M.E. prepared figures; N.G. and M.E. drafted manuscript; N.G., A.M.,
- 532 D.C., A.N., J.R., D.O., S.G., and M.E. edited and revised manuscript; N.G., A.M., D.C., A.N., J.R., D.O.,
- 533 S.G., and M.E. approved final version of manuscript.
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- 702

703 Figure captions

704 **Figure 1.** Representative time course of changes for the adjustment in normalized oxygen uptake (\dot{VO}_2 ; 705 open circles) during the work-to-work cycling transitions for individuals in the moderate-intensity 706 continuous training (MICT), high-intensity interval training (HIIT) and non-exercising control groups. 707 The vertical line illustrates the abrupt transition to the higher work-rate. The continuous black lines of 708 best fit illustrate the primary phase of the VO_2 response. Note the relatively faster time constant of the 709 primary phase of the $\dot{V}O_2$ response ($\dot{V}O_2 \tau_p$) and a reduced $\dot{V}O_2$ slow component beyond week 3 of 710 training in the participants from the HIIT and MICT groups, while $\dot{V}O_2 \tau_p$ and $\dot{V}O_2$ slow component are 711 not affected by training in the participant from the control group.

712

Figure 2. Individual time course of changes in the time constant of the primary phase of the oxygen uptake ($\dot{V}O_2 \tau_p$) and amplitude of the $\dot{V}O_2$ slow component ($\dot{V}O_2 A_s$) in the moderate-intensity continuous training (MICT, panels A and D; n = 11), high-intensity interval training (HIIT, panels B and E; n = 8) and non-exercising control groups (panels C and F; n = 9). Thin lines represent individual participants and thick lines, the mean change in each group. A two-factor (time vs group) mixed ANOVA was used for the analysis.

719 * Significantly different from pretraining (P < 0.05).

720

Figure 3. Representative time course of changes for the adjustment in normalized deoxygenated hemoglobin and myoglobin concentration (Δ [HHb+Mb]; open circles) during the work-to-work cycling transitions for representative individuals in the moderate-intensity continuous training (MICT), highintensity interval training (HIIT) and non-exercising control groups. The vertical line illustrates the abrupt transition to the higher work-rate. The continuous grey lines of best fit illustrate the primary phase of the Δ [HHb+Mb] response. Note the time constant of the primary phase of the Δ [HHb+Mb] response (Δ [HHb+Mb] τ_p) is not affected by training in any of the three group.















	MICT	HIIT	Control
n	11	8	9
Sex (male, female), <i>n</i>	7, 4	6, 2	4, 5
Age, yr	54 ± 10	51 ± 10	54 ± 9
BMI, kg/m ²	31.0 ± 5.7	28.8 ± 3.2	30.5 ± 3.6
Time since diabetes diagnosis, yr	6.6 ± 3.7	6.8 ± 3.7	6.6 ± 3.3
HbA _{1c} , %	6.9 ± 0.5	7.3 ± 0.5	6.8 ± 1.0
Fat layer of VL, mm	7.9 ± 4.2	6.5 ± 2.7	8.6 ± 3.2
Diabetes medication			
Diet only, <i>n</i>		1	1
Metformin, n	9	7	6
Sulfonylurea, <i>n</i>	2	3	2
DPP-4 inhibitor, <i>n</i>			2
GLP-1 analogues, <i>n</i>	1		1
Anti-hypertensive medication			
Angiotensin converting enzyme		1	
inhibitor, <i>n</i>			
Angiotensin II receptor blocker,	1		1
n			
Statins, <i>n</i>	5	3	3
Aspirin, <i>n</i>	3	1	2
PO _{peak} , W	160 ± 54	198 ± 41	148 ± 49
PO@ Δ50%, W	126 ± 43	161 ± 31	115 ± 36
PO@ 80% VT, W	74 ± 27	$99 \pm 17^{*\dagger}$	66 ± 20
Habitual physical activity			
Inactive, h/day	17.4 ± 2.0	17.4 ± 2.9	17.9 ± 1.9
Light, h/day	5.8 ± 1.7	5.8 ± 2.6	5.4 ± 1.2
MVPA, h/day	0.8 ± 0.7	0.8 ± 0.3	0.7 ± 0.9

Table 1. Physical characteristics, pretraining peak exercise values, and activity levels.

Data are mean \pm SD. n = no. of participants; MICT, moderate-intensity continuous training; HIIT, high-intensity interval training; BMI, body mass index; HbA_{1c}, glycosylated haemoglobin; VL, vastus lateralis; DPP-4, Dipeptidyl-peptidase 4; GLP-1, Glucagon-like peptide 1. PO, power output; VT, ventilatory threshold; MVPA, moderate-to-vigorous physical activity. A one-way ANOVA was used for the analysis.

* Significantly different than Control (P < 0.05).

† Significantly different than MICT (P < 0.05).

	Ductus	W1-2	W/1- (W/1- 0	De etters in in e		
	Pretraining	week 3	week b	week 9	Posttraining		
Moderate intensity	Moderate intensity						
Baseline VO_2 , L/min MICT ^a 0.04 + 0.21 0.04 + 0.17 0.04 + 0.24 0.06 + 0.21 0.02 + 0.17							
	0.94 ± 0.21	0.94 ± 0.17	0.94 ± 0.24	0.96 ± 0.21	0.92 ± 0.17		
HIII	0.80 ± 0.23	0.84 ± 0.16	0.83 ± 0.09	0.84 ± 0.14	0.83 ± 0.12		
$\begin{array}{ c c c c c c c c c c c c c c c c c c c$							
$VO_2 A_p, L/min$	0 (4 + 0 22	0.00 + 0.20	0 (1 + 0 04	0 (1 + 0 07	0 (2 + 0 20		
MIC1	0.64 ± 0.32	0.60 ± 0.30	0.61 ± 0.24	0.61 ± 0.27	0.62 ± 0.29		
HIIT	0.92 ± 0.29	0.88 ± 0.27	0.90 ± 0.18	0.87 ± 0.24	0.89 ± 0.18		
Control	0.52 ± 0.22	0.55 ± 0.24	0.52 ± 0.25	0.51 ± 0.24	0.53 ± 0.22		
$VO_2 G_p$ gain, mL.mir	n ⁻¹ .W ⁻¹						
MICT	9.8 ± 1.9	9.3 ± 1.7	9.8 ± 1.7	9.4 ± 1.1	9.6 ± 1.1		
HIIT	10.2 ± 2.2	9.7 ± 1.8	10.1 ± 1.0	9.7 ± 1.5	10.0 ± 0.8		
Control	9.3 ± 1.6	9.7 ± 1.2	9.2 ± 1.3	9.0 ± 0.9	9.5 ± 0.8		
$VO_2 \tau_p, s$							
MICT	46 ± 12	$33 \pm 5^{*\dagger}$	$29 \pm 11^{*\dagger}$	$27 \pm 6^{*\dagger}$	$28 \pm 6^{*\dagger}$		
HIIT	41 ± 7	$32\pm4^{*\dagger}$	$27\pm4^{*\dagger}$	$26\pm4^{*\dagger}$	$27 \pm 4^{*\dagger}$		
Control	43 ± 7	41 ± 6	40 ± 7	41 ± 8	46 ± 7		
$CI_{95}\dot{V}O_{2}\tau_{p},s$							
MICT	4.4 ± 1.2	4.0 ± 1.5	4.1 ± 1.0	3.4 ± 1.1	3.3 ± 1.0		
HIIT	4.4 ± 0.4	4.2 ± 1.1	4.0 ± 0.9	3.3 ± 0.9	3.8 ± 0.7		
Control	4.0 ± 1.1	3.7 ± 0.5	3.8 ± 0.7	4.0 ± 1.2	4.8 ± 1.4		
$\dot{V}O_2 \tau_p, s$							
MICT	46 ± 12	$33\pm5^{*\dagger}$	$29 \pm 11^{*\dagger}$	$27\pm6^{*\dagger}$	$28\pm6^{*\dagger}$		
HIIT	41 ± 7	$32\pm4^{*\dagger}$	$27\pm4^{*\dagger}$	$26\pm4^{*\dagger}$	$27\pm4^{*\dagger}$		
Control	43 ± 7	41 ± 6	40 ± 7	41 ± 8	46 ± 7		
High Intensity							
Baseline VO ₂ , L/min							
MICT	1.58 ± 0.39	1.55 ± 0.35	1.54 ± 0.38	1.57 ± 0.38	1.54 ± 0.36		
HIIT ^a	1.75 ± 0.31	1.73 ± 0.22	1.74 ± 0.19	1.71 ± 0.21	1.72 ± 0.20		
Control	1.31 ± 0.32	1.30 ± 0.28	1.30 ± 0.31	1.31 ± 0.32	1.30 ± 0.30		
$\dot{V}O_2 A_p$, L/min							
MICT	0.38 ± 0.15	0.44 ± 0.14	0.48 ± 0.16	0.47 ± 0.13	0.49 ± 0.16		
HIIT	0.51 ± 0.17	0.54 ± 0.18	0.55 ± 0.16	0.56 ± 0.15	0.57 ± 0.13		
Control	0.42 ± 0.15	0.41 ± 0.18	0.42 ± 0.15	0.40 ± 0.18	0.41 ± 0.21		
$\dot{V}_{02} \tau_{\rm m} s$							
MICT	56 ± 9	$43 \pm 6^{*\dagger}$	$41 \pm 5^{*\dagger}$	$41 \pm 7^{*\dagger}$	$39 \pm 7^{*\dagger}$		
HIIT	$\frac{56 \pm 9}{56 \pm 8}$	$42 + 6^{*\dagger}$	$40 + 5^{*\dagger}$	$38 + 4^{*\dagger}$	$\frac{37}{37+4^{*\dagger}}$		
Control	50 ± 0 54 + 6	53+4	52 + 6	52 - 7	52 + 6		
Clos VO2 Tr. S	01-0			<u> </u>	<i>52</i> ± 0		
MICT	86+25	89 + 24	90 + 25	83+24	88+17		
	0.0 - 2.0		, <u>-</u> <u>-</u>	0.0 - 2.1	0.0 - 1.7		

Table 2. Dynamic response characteristics of $\dot{V}O_2$ during moderate-intensity and highintensity cycling exercise of the work-to-work transitions for the MICT, HIIT and Control groups.

Control	8.8 ± 2.3	8.8 ± 2.3	9.0 ± 2.8	9.0 ± 2.5	8.8 ± 2.2		
$\dot{V}O_2 A_s, L/min$							
MICT	0.17 ± 0.07	$0.09\pm0.05^{*\dagger}$	$0.08\pm0.06^{*\dagger}$	$0.08\pm0.06^{*\dagger}$	$0.07\pm0.05^{*\dagger}$		
HIIT	0.18 ± 0.05	$0.09\pm0.08^{*\dagger}$	$0.11\pm0.06^{*\dagger}$	$0.10\pm0.07^{*\dagger}$	$0.10\pm0.08^{*\dagger}$		
Control	0.17 ± 0.05	0.17 ± 0.04	0.17 ± 0.06	0.17 ± 0.08	0.17 ± 0.05		
$\dot{V}O_2 A_s, \%$							
MICT	32 ± 11	$16 \pm 5^{*\dagger}$	$16 \pm 9^{*\dagger}$	$14\pm9^{*\dagger}$	$14\pm9^{*\dagger}$		
HIIT	26 ± 5	$14 \pm 9^{*\dagger}$	$16\pm7^{*\dagger}$	$15\pm8^{*\dagger}$	$14\pm9^{*\dagger}$		
Control	29 ± 7	30 ± 8	29 ± 9	31 ± 9	32 ± 10		
VO ₂ MRT, s							
MICT	115 ± 8	$73\pm10^{*\dagger}$	$73 \pm 11^{*\dagger}$	$70\pm14^{*\dagger}$	$73\pm15^{*\dagger}$		
HIIT	112 ± 8	$76 \pm 10^{*\dagger}$	$78\pm13^{*\dagger}$	$80\pm14^{*\dagger}$	$78\pm12^{*\dagger}$		
Control	114 ± 13	118 ± 12	120 ± 11	119 ± 9	121 ± 11		
End-exercise $\dot{V}O_2$ gain, mL.min ⁻¹ .W ⁻¹							
MICT	10.2 ± 0.9	9.6 ± 1.2	10.1 ± 1.7	9.9 ± 0.8	10.1 ± 1.0		
HIIT	10.8 ± 2.1	10.0 ± 1.6	10.4 ± 0.9	10.1 ± 1.1	10.3 ± 0.7		
Control	10.7 ± 1.4	10.8 ± 1.3	10.6 ± 1.3	10.4 ± 0.6	10.8 ± 1.5		

Data are mean (SD). VO2, oxygen consumption; MICT, moderate-intensity continuous

training (n = 11 participants); HIIT, high-intensity interval training (n = 8 participants); Control (n = 9 participants); A, amplitude; τ , time constant , $\dot{V}O_2$, oxygen consumption; p, primary response; CI₉₅, 95% confidence interval; s, slow component phase; MRT, mean response time. A two-factor (time vs group) mixed ANOVA was used for the analysis.

* Significantly different from pretraining (P < 0.05); † significantly different from Control (P < 0.05); ^a significantly different than Control (P < 0.05); ^b significantly different than MICT (P < 0.05).

Table 3. Dynamic response characteristics of [HHb + Mb] and TOI during moderate-

intensity and high-intensity cycling exercise of the work-to-work transitions for the MICT,

	Pretraining	Week 3	Week 6	Week 9	Posttraining		
Moderate intensity							
Baseline Δ [HHb + Mb] μ Mol.cm							
MICT	-67 ± 42	-72 ± 68	-84 ± 60	-60 ± 54	-62 ± 32		
HIIT	-59 ± 44	-69 ± 35	-66 ± 41	-66 ± 38	-59 ± 35		
Control	-55 ± 37	-52 ± 30	-53 ± 30	-60 ± 29	-54 ± 31		
Δ [HHb + Mb] A _p , µN	Mol.cm						
MICT	93 ± 36	104 ± 57	95 ± 40	91 ± 54	89 ± 32		
HIIT ^a	183 ± 109	170 ± 105	169 ± 108	181 ± 108	179 ± 111		
Control	73 ± 59	71 ± 52	66± 54	68 ± 55	68 ± 44		
Δ [HHb + Mb] τ , s	•						
MICT	29 ± 7	28 ± 5	27 ± 10	27 ± 12	27 ± 4		
HIIT	23 ± 3	27 ± 5	23 ± 2	24 ± 9	23 ± 3		
Control	26 ± 4	24 ± 6	26 ± 7	27 ± 5	26 ± 7		
Primary phase Δ [HH	$(b + Mb)/\Delta \dot{V}O_2 \mu Mc$	ol.cm.(L/min)					
MICT	149 ± 84	158 ± 76	153 ± 71	141 ± 72	140 ± 66		
HIIT	174 ± 129	193 ± 110	170 ± 119	186 ± 133	186 ± 141		
Control	122 ± 86	110 ± 67	113 ± 85	114 ± 77	120 ± 74		
Baseline TOI, %	•						
MICT	71 ± 3	72 ± 5	72 ± 7	70 ± 5	73 ± 6		
HIIT	71 ± 7	71 ± 8	71 ± 6	71 ± 9	72 ± 6		
Control	71 ± 6	72 ± 7	71 ± 6	71 ± 7	74 ± 7		
TOI A, %	·						
MICT	4.1 ± 3.5	4.9 ± 3.9	3.2 ± 2.3	2.8 ± 3.8	3.2 ± 3.8		
HIIT ^{ab}	7.1 ± 6.5	9.1 ± 5.9	7.1 ± 5.2	7.4 ± 6.0	8.6 ± 6.0		
Control	2.9 ± 3.6	3.4 ± 3.4	2.9 ± 3.3	2.7 ± 2.4	3.2 ± 4.8		
High Intensity	•						
Baseline Δ [HHb + M	lb]μMol.cm						
MICT	32 ± 42	31 ± 60	12 ± 47	36 ± 60	31 ± 46		
HIIT ^{ab}	122 ± 130	123 ± 100	115 ± 130	125 ± 129	122 ± 137		
Control	9 ± 68	23 ± 67	12 ± 84	7 ± 71	15 ± 50		
Δ [HHb + Mb] A _p , μ M	Δ [HHb + Mb] A _n , µMol.cm						
MICT	60 ± 38	68 ± 31	69 ± 32	75 ± 48	65 ± 26		
HIIT	78 ± 42	77 ± 51	86 ± 44	91 ± 97	80 ± 32		
Control	42 ± 49	33 ± 36	39 ± 51	39 ± 48	35 ± 42		
Δ [HHb + Mb] τ , s							
MICT	30 ± 10	32 ± 5	33 ± 6	34 ± 7	33 ± 7		
HIIT	32 ± 11	32 ± 11	32 ± 11	32 ± 9	32 ± 9		
Control	28 ± 6	26 ± 11	29 ± 9	29 ± 13	28 ± 13		
Primary phase Δ [HH	$(b + Mb)/\Delta \dot{V}O_2 \mu Mc$	ol.cm.(L/min)	·	·	·		
MICT	163 ± 111	148 ± 58	141 ± 66	149 ± 84	132 ± 67		

 110 ± 68

 134 ± 100

 136 ± 97

 122 ± 64

HIIT and Control groups.

HIIT

 135 ± 93

Control	83 ± 70	65 ± 53	82 ± 97	74 ± 60	73 ± 64		
Δ [HHb + Mb] A _s , μ Mol.cm							
MICT	10 ± 6	15 ± 10	15 ± 15	14 ± 11	13 ± 10		
HIIT	12 ± 7	12 ± 8	19 ± 12	17 ± 12	12 ± 8		
Control	11 ± 7	12 ± 6	12 ± 8	13 ± 10	12 ± 10		
Baseline TOI, %							
MICT	67 ± 5	67 ± 8	69 ± 8	68 ± 7	70 ± 9		
HIIT	64 ± 12	62 ± 12	64 ± 11	63 ± 12	63 ± 11		
Control	68 ± 9	68 ± 10	68 ± 8	68 ± 8	70 ± 10		
TOI A, %							
MICT	3.1 ± 2.3	3.0 ± 2.6	4.6 ± 1.9	4.6 ± 2.2	3.6 ± 2.7		
HIIT	3.7 ± 2.3	4.5 ± 3.4	6.3 ± 4.2	5.7 ± 4.9	4.3 ± 3.3		
Control	3.0 ± 2.4	2.3 ± 2.4	2.0 ± 3.6	2.2 ± 2.9	2.2 ± 3.3		

Data are mean (SD). MICT, moderate-intensity continuous training (n = 11 participants);

HIIT, high-intensity interval training (n = 8 participants); Control (n = 9 participants); A, amplitude; τ , time constant, $\dot{V}O_2$, oxygen consumption; p, primary response; s, slow component phase. [HHb + Mb], deoxygenated haemoglobin and myoglobin concentration; τ ' [HHb + Mb], effective time constant (τ + TD); TOI, tissue oxygenation index. A two-factor (time vs group) mixed ANOVA was used for the analysis.

^a significantly different than Control (P < 0.05); ^b significantly different than MICT (P < 0.05).

Low-volume HIIT and MICT speed $\dot{V}O_2$ kinetics during high-intensity "work-to-work" cycling with a similar time-course in type 2 diabetes



Both forms of training induced a rapid (by wk 3) acceleration of primary \dot{VO}_2 time constant and a reduction in the amplitude of the \dot{VO}_2 slow component despite training volume and time commitment being 50% lower in the HIIT group.