Heterogeneity and incidence of non-response for changes in cardiorespiratory fitness following time-efficient sprint interval exercise training.

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Abstract

Interindividual variability for training-induced changes in maximal oxygen uptake ($\text{VO}_2\text{max}$) has been well described in response to continuous aerobic and high-intensity interval exercise. However, whether similar variability is observed following time-efficient sprint interval training (SIT) protocols with a minimal total training volume (i.e. reduced-exertion high-intensity interval training; REHIT) is not known. We conducted a pooled analysis of $n=117$ (68 men) training participants (mean±SD: age: 30±10 y; $\text{VO}_2\text{max}$: 34.8±7.5 ml·kg$^{-1}$·min$^{-1}$), who completed a $\text{VO}_2\text{max}$ assessment before and 3 days after 6 weeks of REHIT comprising of two 10-20 second ‘all-out’ cycling sprints per session, and $n=40$ no-intervention control participants (age: 30±13 y; $\text{VO}_2\text{max}$: 31.5±6.5 ml·kg$^{-1}$·min$^{-1}$) who completed repeated $\text{VO}_2\text{max}$ tests over a comparable time frame. Individual responses estimated using 50% confidence intervals derived from the technical error were interpreted against a smallest worthwhile change (SWC) of 1.75 ml·kg$^{-1}$·min$^{-1}$. The standard deviation of individual responses ($\text{SD}_{\text{IR}}$) was 2.39 ml·kg$^{-1}$·min$^{-1}$ (Cohens $d=0.32$, i.e. ‘moderate’) demonstrating clinically meaningful interindividual differences in training-induced changes in $\text{VO}_2\text{max}$ following REHIT that exceed the technical, biological and random within-subjects variability of $\text{VO}_2\text{max}$ assessment. The likely (75% probability) non-response rate was 18% (21/117), and 49% (57/117) of individuals demonstrated an increase in $\text{VO}_2\text{max}$ likely higher than the SWC. We conclude that the well-described increase in $\text{VO}_2\text{max}$ observed following REHIT at the group level is subject to substantial variability in magnitude at an individual level. This has important implications for exercise prescription and can be harnessed by future studies aiming to elucidate mechanisms of adaptation.
Keywords: Aerobic Capacity; High-Intensity Interval Training; Sprint Interval Training; Individual Responses; Individual Variability; Cardiorespiratory Fitness

1. Introduction

The maximal attainable rate of oxygen uptake (VO₂max) is amongst the most important physiological traits that determine long term health and longevity. Indeed, a low VO₂max predicts cardiovascular and all-cause mortality to a similar or greater extent compared with other established risk factors, including body mass index, smoking, hypertension and type 2 diabetes (Ross et al., 2016). Although VO₂max is a partly heritable trait (Bouchard et al., 1998), it can also be improved (on average) through regular exercise training (Bouchard et al., 1999; Sisson et al., 2009), and those who are able to improve their VO₂max over several years lower their risk of cardiovascular and all-cause mortality in a dose-dependent manner (Lee et al., 2011).

Whilst VO₂max improves on average in response to both continuous endurance and high-intensity interval exercise training (Bacon et al., 2013; Weston et al., 2014), it has been recognised for over 3 decades that individual measured changes following standardised exercise training can be highly variable and a proportion of people will demonstrate no measurable change (so called, ‘non-responders’) (Bouchard et al., 1999; Lortie et al., 1984). Even following several months of high-volume aerobic exercise training, measured individual changes in VO₂max can range from decreases of 100 ml·min⁻¹ to gains of more than 1100 ml·min⁻¹ (Bouchard et al., 1999). This interindividual variability in response is thought to be explained by a range of factors, including random or technical error, the method of standardising relative exercise intensity, and genetic and epigenetic variance (Sarzynski et al., 2017). Some have also argued that ‘non-responders’ to exercise training in general do not exist and instead are an artefact of an insufficient training stimulus for those individuals (Bacon et al., 2013; Montero and Lundby, 2017). Nonetheless, some individuals may be non-responders.
to set training interventions (characterised by training intensity, duration, frequency, and mode) that are efficacious at inducing training effects in others.

Over the last 15 years, sprint interval training (SIT) has emerged as an efficacious exercise training stimulus for improving VO$_2$max in previously inactive individuals (Gillen and Gibala, 2014; Sultana et al., 2019; Vollaard and Metcalfe, 2017; Vollaard et al., 2017). A particularly interesting finding to emerge from SIT research is that the training-induced change in VO$_2$max does not appear to increase (and possibly decreases) with an increasing number of sprint repetitions (Vollaard et al., 2017). Indeed, at a group level, improvements in VO$_2$max have been observed with as few as two or three, 20-second, all-out cycling sprints performed regularly over a 6-12-week training intervention (termed ‘reduced-exertion high-intensity interval training’ or REHIT) (Gillen et al., 2016; Metcalfe et al., 2016, 2012). These findings have particular relevance in the search for effective, ‘real-world’, time-efficient exercise strategies to overcome lack of time as a barrier to exercise initiation and adherence in low active individuals (Vollaard and Metcalfe, 2017).

The variability in training-induced changes in VO$_2$max has been well described in response to continuous endurance training (Bouchard et al., 1999; Sisson et al., 2009) and more recently in response to high-intensity (HIIT) and sprint interval training (SIT) (Astorino and Schubert, 2014; Bonafiglia et al., 2016; Gurd et al., 2016; Islam et al., 2020; Phillips et al., 2017; Williams et al., 2019). However, studies examining interindividual variability in response to SIT to date have involved arduous SIT protocols requiring a relatively high number of sprint repetitions (Astorino and Schubert, 2014; Bonafiglia et al., 2016; Gurd et al., 2016; Islam et al., 2020; Williams et al., 2019), have combined HIIT and SIT protocols together (Williams et al., 2019) or, in some cases, the analysis has been limited by not utilising relevant information from a time-matched control condition (Astorino and Schubert, 2014). No study has characterised the
heterogeneity in response or the incidence of non-responders to a genuinely time-efficient SIT exercise protocol such as REHIT. Given the low overall dose of exercise involved with REHIT (a total of <10 minutes of sprint exercise within 3 hours of training time over a 6-week training period, e.g. (Metcalf et al., 2016, 2012)), alongside suggestions that ‘non-responders’ may be an artefact of an insufficient training stimulus (Bacon et al., 2013; Montero and Lundby, 2017), it is reasonable to question whether the incidence of non-response would be high following this training intervention, and what proportion of individuals (if any) are likely to show a change that would be considered clinically meaningful. Individual variability in the training-induced change in VO$_2$max in response to REHIT has been alluded to (Metcalf et al., 2016), but not definitively demonstrated using an adequate sample size, or appropriate experimental and statistical methods. The inclusion of data from no-exercise control group is particularly important when assessing individual responses to exercise training in order to account for the variance caused by technical error, day-to-day biological and random within subjects variability (Atkinson and Batterham, 2015; Bonafiglia et al., 2019). Thus, the aim of this study was firstly to establish whether true individual variability in changes in VO$_2$max in response to REHIT exists and, if so, to characterise the heterogeneity of response and incidence of non-responders to this extremely low-volume and time-efficient exercise intervention.
2. Methods

2.1. Participants / Sample

We conducted a secondary analysis of six published studies conducted in our laboratories between 2012 and 2020 (Metcalfe et al., 2020, 2016, 2012; Nalçakan et al., 2018; Songsorn et al., 2016; Thomas et al., 2020). This included a sample of 117 pooled training participants (68 male, 49 female) who underwent an almost identical SIT intervention (described in full below), and 40 pooled no-intervention control participants (16 male, 24 female) from three of these studies who underwent two assessments of maximal oxygen uptake either ~4 (n=14; (Songsorn et al., 2016)) or ~6 (n=26; (Metcalfe et al., 2020, 2012)) weeks apart. Training participants from two of the included studies who underwent substantially different SIT interventions, either involving single 20-s sprints (Songsorn et al., 2016) or reduced sprint duration (Nalçakan et al., 2018), were excluded because these studies demonstrated that these interventions either do not alter VO$_2$max (single 20-s sprints; (Songsorn et al., 2016)) or results in a significantly lower mean increase (reduced sprint duration; (Nalçakan et al., 2018)). The inclusion and exclusion criteria were similar across all six included studies. All participants were classified as either sedentary or moderately physically active on enrolment onto the study according to the criteria of the International Physical Activity Questionnaire. Participants with any contraindication to exercise based on a self-report health history questionnaire and an assessment of high resting blood pressure (>140/90 mmHg) or high resting heart rate (>100 bpm) were excluded. The pooled participant characteristics are shown in Table 1. Ethical approval was obtained for all included experiments (details and approval references are available in the original articles (Metcalfe et al., 2020, 2016, 2012; Nalçakan et al., 2018;
Songsorn et al., 2016; Thomas et al., 2020) and all participants provided their written consent to take part after they received information about the study both verbally and in writing. All experiments were conducted in accordance with the Declaration of Helsinki.

<table>
<thead>
<tr>
<th>Table 1 Baseline Participant Characteristics</th>
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<tr>
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<tr>
<td><strong>Male / Female (n)</strong></td>
</tr>
<tr>
<td>Age (y)</td>
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<td>Height (m)</td>
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<tr>
<td>Body Mass (kg)</td>
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<tr>
<td>BMI (kg·m⁻²)</td>
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<tr>
<td>VO₂max (ml·kg⁻¹·min⁻¹)</td>
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</table>

Data is presented as mean ± SD. p-values derived from independent t-test.

2.2. Assessment of Maximal Oxygen Uptake

All participants underwent an incremental cycling test to their limit of tolerance to determine maximal oxygen uptake. In participants in the SIT groups, these were conducted in the week prior to training and then ~3 days following the final training session. The specific protocols have been described previously (Metcalfe et al., 2020, 2016, 2012; Nalçakan et al., 2018; Songsorn et al., 2016; Thomas et al., 2020). Although there were slight variations in the protocol and equipment used across studies, the mean and standard deviation of the change in VO₂max are strikingly similar (Table 3) suggesting these are ‘typical’ for studies of this nature and hence appropriate for a pooled analysis. In the majority (n=103) of training participants and all (n=40) of the control participants, breath by breath measurements of pulmonary gas exchange were captured continuously during the test using a metabolic cart. Breath by breath measurements of oxygen uptake (VO₂) were converted into 15-breath rolling averages and VO₂max was taken as the highest value for a 15-breath rolling average achieved during the test. This method was used in all but one of the included experiments which was reanalysed for the present analysis (Metcalfe et al., 2012). For a small sample of training participants
(n=14) from one study (Metcalf et al., 2016), pulmonary gas exchange was measured using
the Douglas Bag technique. It was considered appropriate to retain these participants in this
analysis because previous independent studies have reported similar test-retest reliability for
VO₂max measured using Douglas Bags or breath by breath methods (Katch et al., 1982;
Phillips et al., 2017). The following secondary criteria were used to verify a maximal effort: 1)
volutional exhaustion, 2) a plateau in VO₂ despite increasing workload, 3) RER >1.15 and 4) a
maximal heart rate within 10 beats of age predicted maximum (220-age). All participants
achieved two or more of these criteria.

2.3. Sprint Interval Training and No-Intervention Control

All training participants underwent a 6-week cycling based REHIT intervention with only
small differences across studies (Table 2). The majority of participants (n=104) completed this
supervised in an exercise physiology laboratory on a mechanically braked cycle ergometer
(Monark, Vansbro, Sweden) (Metcalf et al., 2016, 2012; Nalçakan et al., 2018; Thomas et al.,
2020). Participants (n=13) in one study completed the intervention unsupervised on a
commercially available electronically braked cycle ergometer (CAROL™, Integrated Health
Partners Ltd, London, UK) (Metcalf et al., 2020). Each SIT session lasted ~10 min and
consisted of low intensity cycling interspersed with two ‘all-out’ sprints against a fixed
resistance (between 10 and 20 s). In the 2-3 seconds prior to each ‘all-out’ sprint, participants
increased their pedal cadence to their maximal speed, the braking resistance was then applied
to the bike, and participants cycled as fast as they could for the duration of the sprint. The
majority of participants completed 3 sessions/week (n=76) but a subset completed either 2
(n=29) or 4 (n=12) sessions/week. We recently demonstrated that the mean change in VO₂max
with SIT was not different across training frequencies of 2, 3 or 4 sessions/week (Thomas et
al., 2020), so it was deemed appropriate to pool them together in this analysis. All no-
intervention control participants from the 3 separate studies were given the same instructions to maintain their current physical activity levels and dietary patterns for the duration of the study.

Table 2 Training interventions applied in included studies

<table>
<thead>
<tr>
<th>Intervention Duration (weeks)</th>
<th>Metcalfe 2012</th>
<th>Metcalfe 2016</th>
<th>Nalcakan 2018</th>
<th>Thomas 2020</th>
<th>Metcalfe 2020</th>
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</thead>
<tbody>
<tr>
<td>Frequency (sessions / week)</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>2, 3 or 4</td>
<td>2</td>
</tr>
<tr>
<td>Total Session Duration (mins)</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>8:20-8:40</td>
</tr>
<tr>
<td>Sprints Per Session (n)</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Braking Mass (% BM)</td>
<td>7.5</td>
<td>5</td>
<td>7.5</td>
<td>7.5</td>
<td>5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sprint Duration(s)</th>
<th>Week 1</th>
<th>Week 2</th>
<th>Week 3</th>
<th>Week 4</th>
<th>Week 5</th>
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<tr>
<td></td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
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</tbody>
</table>

| Warm-Up Duration (s)          | 180          | 180          | 120          | 100-110      | 120          |
| Recovery Duration (s)         | 200-220      | 200-220      | 200-220      | 200-220      | 180          |
| Cool Down Duration (s)        | 180          | 180          | 240          | 240          | 180          |

| Intensity of Warm-Up, Recovery and Cool Down | ~60 W | Unloaded | Unloaded | Unloaded | ~25 W |

Note: in the Metcalfe et al (2012 and 2016) studies, the first session only contained 1 x 10 s sprint. Where a range of warm-up and recovery durations are presented, this is due to the increase in sprint duration during initial training weeks.

2.4. Statistical Analysis

2.4.1. Group Responses

The effect of SIT on VO_2max was analysed using a two-way repeated measures analysis of variance (group x time), performed in Graphpad Prism 8 for macOS (Version 8.4.2, San Diego, CA, USA). Alpha was set at p<0.05 and effect size was calculated using Cohens d (Hopkins et al., 2009).

2.4.2. Individual Variability
To characterise whether true individual responses to SIT were present, the standard deviation (SD) of individual responses (SDIR) was calculated from the square root of the difference between the square of the SD of the change (post minus pre) in the exercise training group (SDex) and the control group (SDcon) (Atkinson and Batterham, 2015):

$$SD_{IR} = \sqrt{(SD_{EX})^2 - (SD_{CON})^2}$$

The SDIR was subsequently interpreted against thresholds for standardised mean change of 0.1 (small effect), 0.3 (moderate effect) and 0.6 (large effect) (Hopkins, 2015) and against the smallest clinically worthwhile change (see below).

### 2.4.3. Classification of Non-, Uncertain and Positive Responders to SIT

The statistical procedures recommended by Swinton et al (Swinton et al., 2018) were followed to classify individual changes in VO2max following SIT using 50% confidence intervals (CIs) that were calculated using the typical error for repeated measurements of VO2max from the control participants. The typical error (TE) was calculated using the formula:

$$TE = \frac{SD_{CON}}{\sqrt{2}}$$

Typical error for relative VO2max was 1.30 ml·kg⁻¹·min⁻¹. For completeness, we classified individual responses against a zero and clinically relevant response thresholds as this is more likely to be informative of individual responsiveness in short term (e.g. 6 weeks) exercise training studies where adaptations may still be accumulating (Islam et al., 2020). However, when interpreting and discussing the data we take the more conservative approach of defining responses against the smallest worthwhile change, alluding to responses against the zero threshold where relevant. Responses were considered against two clinically relevant thresholds of change to provide additional information on the proportion of individual responses at different magnitudes: thresholds of 1.75 ml·kg⁻¹·min⁻¹ (equivalent to ~0.5 METs) and 3.5
ml·kg⁻¹·min⁻¹ (equivalent to ~1 MET) were applied based on the data of Lee et al who

demonstrated an ~8% and ~16% decrease in relative risk of CVD, respectively, over ~11 years

of follow up (Lee et al., 2011). Both of these thresholds have been applied in previous studies

on this topic (Bonafiglia et al., 2018). A responder was classified if the entire 50% CI lay above

the specified response threshold. In these instances, the use of a 50% CI means that there is a

75% probability of a response for this individual, i.e. ‘likely’ (Hopkins, 2015). If the 50% CI

crossed the response threshold then this was classified as an ‘uncertain response’, whilst a ‘non-

responder’ was defined if the entire 50% CI lay below the response threshold (Bonafiglia et

al., 2018). All analysis of individual responses was performed in Microsoft Excel and

Graphpad Prism 8 for macOS (Version 8.4.2, San Diego, CA, USA).
3. Results

3.1. Group Effects and SDIR

For relative VO$_{2\text{max}}$, there were main effects of group ($F=13.2$, $p<0.001$) and time ($F=35.7$, $p<0.001$) as well as a group x time interaction ($F=39.8$, $p<0.001$): mean relative VO$_{2\text{max}}$ increased in the SIT group compared to the control group (Table 3 and Figure 1, $d=0.43$). The SDIR for the change in relative VO$_{2\text{max}}$ following SIT compared to the control group was 2.39 ml·kg$^{-1}$·min$^{-1}$ with an effect size of 0.32 (‘moderate’). The SDIR was not substantially altered (2.30 ml·kg$^{-1}$·min$^{-1}$) when one potentially influential SIT participant with a -5.99 ml·kg$^{-1}$·min$^{-1}$ decrease in VO$_{2\text{max}}$ was removed from the analysis ($d=0.30$, ‘moderate’). Thus, there is evidence of individual differences in VO$_{2\text{max}}$ trainability in response to REHIT that exceed the smallest clinically worthwhile effect of 1.75 ml·kg$^{-1}$·min$^{-1}$.

Table 3 Changes in absolute and relative VO$_{2\text{max}}$ following SIT

<table>
<thead>
<tr>
<th>Study</th>
<th>Pre</th>
<th>Post</th>
<th>Delta</th>
<th>Pre</th>
<th>Post</th>
<th>Delta</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Training</strong></td>
<td></td>
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<tr>
<td>Metcalfe 2012</td>
<td>2.19±0.57</td>
<td>2.48±0.60</td>
<td>0.29±0.13</td>
<td>33.5±4.9</td>
<td>38.3±4.8</td>
<td>4.7±2.7</td>
</tr>
<tr>
<td>Metcalfe 2016</td>
<td>2.54±0.65</td>
<td>2.78±0.67</td>
<td>0.24±0.23</td>
<td>35.0±7.8</td>
<td>38.1±7.9</td>
<td>3.0±3.3</td>
</tr>
<tr>
<td>Nalcakan 2018</td>
<td>2.77±0.75</td>
<td>3.04±0.75</td>
<td>0.27±0.28</td>
<td>39.0±6.9</td>
<td>42.3±6.5</td>
<td>3.3±3.3</td>
</tr>
<tr>
<td>Thomas 2020</td>
<td>2.77±0.77</td>
<td>3.01±0.82</td>
<td>0.24±0.24</td>
<td>35.4±7.2</td>
<td>38.5±7.5</td>
<td>3.1±2.8</td>
</tr>
<tr>
<td>Metcalfe 2020</td>
<td>2.25±0.75</td>
<td>2.42±0.82</td>
<td>0.17±0.21</td>
<td>28.0±6.7</td>
<td>29.8±7.6</td>
<td>1.8±2.5</td>
</tr>
<tr>
<td><strong>Control</strong></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Metcalfe 2012</td>
<td>2.32±0.64</td>
<td>2.37±0.77</td>
<td>0.05±0.18</td>
<td>33.8±5.5</td>
<td>34.2±6.0</td>
<td>0.3±2.3</td>
</tr>
<tr>
<td>Songsorn 2015</td>
<td>2.07±0.69</td>
<td>2.08±0.68</td>
<td>0.01±0.10</td>
<td>32.0±5.8</td>
<td>32.2±6.0</td>
<td>0.2±1.6</td>
</tr>
<tr>
<td>Metcalfe 2020</td>
<td>2.26±0.70</td>
<td>2.20±0.71</td>
<td>-0.06±0.09</td>
<td>28.1±7.3</td>
<td>27.4±7.2</td>
<td>-0.7±1.4</td>
</tr>
<tr>
<td><strong>Pooled</strong></td>
<td>2.59±0.73</td>
<td>2.83±0.77</td>
<td>0.24±0.23</td>
<td>34.8±7.5</td>
<td>37.9±7.9</td>
<td>3.1±3.0</td>
</tr>
</tbody>
</table>
Data is presented as mean ± SD

### 3.2. Classification of Non-, Uncertain and Positive Responders to SIT

When considered against a threshold of zero, 5/117 (4%) of participants were classified as likely non-responders, 29/117 (25%) were classified as uncertain responders, and 83/117 (71%) of the participants showed an increase in VO$_2$max that was likely higher than zero (Figure 2A). When considered against the minimal clinically relevant threshold, 18% (21/117) of participants were likely non-responders, 39/117 (33%) were classified as uncertain responders, and 57/117 (49%) showed an increase likely higher than the minimal clinically relevant threshold of 1.75 ml·kg$^{-1}$·min$^{-1}$ (Figure 2B). Furthermore, 33 out of those 57 (i.e. 33/117; 28%) of those participants showed an increase likely higher than 3.5 ml·kg$^{-1}$·min$^{-1}$ (Figure 2B).
4. Discussion

Individual variability in the change in VO$_2$max following exercise training has important implications for optimising and personalising exercise prescriptions to improve health. This is the first study to report the heterogeneity of response and the incidence of non-response to a genuinely time-efficient SIT protocol that has been shown to be efficacious at a group level in both supervised lab (Metcalfe et al., 2016, 2012; Naçakan et al., 2018; Thomas et al., 2020) and unsupervised real-world (Metcalfe et al., 2020) settings. Importantly, for the first time, we demonstrate statistically that inter-individual differences in training-induced changes in VO$_2$max following REHIT exceed the technical, day-to-day biological, and random within-subjects variability of VO$_2$max assessment over a similar time frame. Of particular note, we report a non-response rate of 18% and that 49% of individuals demonstrate an increase in VO$_2$max that is likely higher than the smallest clinically worthwhile difference.

Several previous studies have quantified non-response rates to a high dose of aerobic exercise training and to HIIT and SIT protocols involving a greater number/duration of high-intensity or all-out sprint efforts (Astorino and Schubert, 2014; Bonafiglia et al., 2016; Gurd et al., 2016; Islam et al., 2020; Phillips et al., 2017; Sisson et al., 2009; Williams et al., 2019). Direct comparisons with these HIIT and SIT studies are somewhat challenging because different thresholds have been applied to quantify non-response, including a change <1xTE (i.e. <0.86 ml·kg$^{-1}$·min$^{-1}$; (Phillips et al., 2017)), <2xTE (i.e. <1.74 ml·kg$^{-1}$·min$^{-1}$; (Gurd et al., 2016)), <1 coefficient of variation (CV) of repeated measurement (i.e. <1.3 ml·kg$^{-1}$·min$^{-1}$; (Astorino and Schubert, 2014)), or <SWC minus 1 x TE (i.e. <1.80 ml·kg$^{-1}$·min$^{-1}$; (Williams et al., 2019)). In the largest pooled analysis of HIIT (n=299) and SIT (n=116) to date, Williams reported a non-response rate of 35% for HIIT and 52% for SIT, using a threshold of <1.8 ml/kg/min (Williams
et al., 2019). Applying the same criteria to the current study would yield a non-response rate of 38%. Thus, it is possible to conclude that the proportion of likely non-responders observed with REHIT is similar to or less than with other SIT and HIIT protocols involving a greater number/duration of high-intensity or all-out sprint efforts (Williams et al., 2019). The fact that similar rates of non-response are observed when applying HIIT and SIT protocols with varying numbers and durations of sprints strongly suggests that non-response for VO\textsubscript{2}max is not an artefact caused by an insufficient dose of exercise training. If this were the case, then the non-response rate would be expected to decrease with an increased number/duration of sprints and hence a greater training ‘stimulus’. The application of sprints of an ‘all-out’ intensity adds further weight to this argument because this likely overcomes any issues regarding the standardisation of exercise intensity (or more specifically the homeostatic disruption) across individuals during each acute training session. Whilst individual differences in the homeostatic disturbance can be expected with MICT standardised using a % of HRmax or VO\textsubscript{2}max, and this may have subsequent implications for the adaptive response (Jamnick et al., 2020; Montero and Lundby, 2017; Preobrazenski et al., 2018), the disruption in homeostasis with SIT / REHIT is always likely to be severe and hence sufficient to switch on signalling pathways that underpin adaptation (if possible for that individual). Indeed, the fact that group level increases in VO\textsubscript{2}max are similar between SIT protocols applying 2-3 sprints compared with protocols applying 6-8 sprints (Vollaard et al., 2017) implies that the adaptive signalling pathways responsible for increasing VO\textsubscript{2}max in response to SIT become ‘saturated’ with only a small number of acute ‘all-out’ sprint efforts (Vollaard and Metcalfe, 2017). Taken together, we contend that non responders for VO\textsubscript{2}max in response to exercise training are a real physiological phenomenon and our data show they are observed in response to REHIT/SIT. That said, it remains unknown whether non-responders to REHIT/SIT for changes in VO\textsubscript{2}max are also non-responders for other health markers (i.e. universal non responders), but previous
work looking at interindividual differences in response to aerobic exercise training suggests that this may not be the case (Vollaard et al., 2009). Future studies need to clearly establish whether non-responders to REHIT/SIT would demonstrate adaptations to different types of exercise, such as MICT. Although previous studies have attempted to examine this (Bonafiglia et al., 2016), the small sample size limits the conclusions that can be made at this stage.

Our data show that ~half of individuals are likely to show an increase in VO\(_2\)max that is greater than the smallest worthwhile change after only 6 weeks of training. This is a striking observation given that REHIT involves less than 10 min of sprint exercise and a total exercise time of 3 hours over that 6-week period. Nevertheless, it is worth noting that this represents a relatively short-term training intervention. At a group level, the magnitude of increase in VO\(_2\)max following SIT that occurs from 6 to 12 weeks is similar to that observed from 0 to 6 weeks of the intervention (Gillen et al., 2016) and this raises important questions about individual differences in the rate at which adaptations are accrued. Indeed, in our analysis, a proportion of individuals (26/117; 22%) demonstrated a change that was likely >0 ml·kg\(^{-1}\)·min\(^{-1}\) but not of a magnitude which exceeded the minimal clinically important difference threshold of 1.75 ml·kg\(^{-1}\)·min\(^{-1}\). Whilst it would be expected that non-responders would continue to show limited adaptation to a longer intervention, it will be important to determine whether individuals showing a slower rate of adaptation at 6 weeks would continue to accrue adaptations (at this lower rate) and would demonstrate a clinically meaningful magnitude of response at a later time point.

The mechanisms that explain the individual variation in response to REHIT are unclear and cannot be determined from the present study. As the training interventions applied were unable (due to logistical reasons) to control the majority of environmental factors outside of the training intervention (e.g. each participants pattern of nutrition, sleep, stress etc.), it can be
expected that some of the variability in adaptation between individuals is explained by (unquantified) environment-training interactions. As an example, the dose, type and timing of nutrition can have a powerful impact on the skeletal muscle signalling response to acute exercise (Cluberton et al., 2005; Guerra et al., 2010; Stocks et al., 2018) and, hence, can modify adaptation to exercise training. If such variables are not controlled across individuals, then it can be expected that this will introduce a level of individual variability in adaptation. This may also explain, at least in part, the poor reproducibility of individual responses to repeated (identical) exercise training interventions in the same sample of participants (Del Giudice et al., 2020).

On the other hand, it is also clear that there is a heritable component to exercise trainability (Bouchard, 2019; Bouchard et al., 1999; Sarzynski et al., 2017; Timmons et al., 2010). The majority of this evidence comes from studies of the training response to aerobic and resistance exercise and the relevance of this information to SIT remains unknown. Indeed, we still do not know whether REHIT/SIT enhances VO$_2$max through mechanisms that are similar to or distinct from MICT (Gibala and Little, 2019; Vollaard and Metcalfe, 2017). However, the inter-individual variability observed in the present study can be used by future investigations interested in identifying (molecular) predictors of response. If responders vs. non-responders to a specific intervention (e.g. SIT/REHIT) can be identified, then contrasting traits / molecular signatures in groups of responders vs. non-responders provides a strong approach to identify potential physiological / genetic / epigenetic factors that determine the interindividual variability in training response (Keller et al., 2007). Such studies are needed to enable personalised medicine, for example, to enable personalised advice on effective interventions (Keller et al., 2007; Timmons et al., 2010) and can be a powerful way to elucidate molecular mechanisms of training adaptations (Keller et al., 2011, 2007). An improved understanding of the molecular mechanisms of adaptation to SIT would be invaluable in the effort to optimise
SIT protocols to enable the greatest adaptations with minimal required effort and time-commitment.

There are a number of limitations to the current analysis that should be considered. Firstly, and most importantly, whilst the large sample size is a strength of this study, this is a pooled dataset from five independent studies and there were minor differences in training protocols, testing procedures, and the duration of the control intervention (4-weeks for n=14 and 6-weeks for n=26) between some of the studies (described in full in the methods). It is possible that these differences may affect the validity of the SDir estimate, which assumes that all sources of variability are equal between the exercise and control groups except that the exercise group underwent exercise training (Atkinson and Batterham, 2015; Bonafiglia et al., 2019). However, two pieces of information can help to mitigate these concerns. Firstly, the SD of the training-induced change in VO$_2$max was comparable between studies (Table 3). Secondly, we performed a sensitivity analysis to examine how excluding the 4-week control participants impacts the SDir calculation. We found a similar SDir estimate of 2.30 ml·kg$^{-1}$·min$^{-1}$ compared to 2.39 ml·kg$^{-1}$·min$^{-1}$ when all control participants were included. Similarly, a comparable SDir of 2.36 ml·kg$^{-1}$·min$^{-1}$ was found when training participants undergoing VO$_2$max assessment with Douglas Bags (n=14) were excluded from the analysis. Thus, these differences in methodology across the pooled independent studies to do not appear to greatly impact the validity of our findings.

Another limitation is that most of the studies involved supervised, lab-based exercise, so it remains unclear whether the results would be the same in real-world settings and this will be important to address in future studies. Furthermore, this analysis is also largely limited to young sedentary but healthy men and women and it is not possible to determine whether different populations (e.g. lean vs overweight, young vs old, men vs women) may show different levels
of response / non-response. Finally, it should also be noted that other important health markers were not considered in this analysis and so at present it remains unknown whether non-responders for VO$_2$max in response to SIT would be able to improve markers of cardiometabolic health.

In conclusion, we demonstrate for the first time that the well described increase in VO$_2$max observed following REHIT at the group level, is subject to substantial variability in magnitude at an individual level. This is an important observation with potential future implications for prescribing SIT/REHIT as an intervention for improving health and can be harnessed by future studies aiming to elucidate the mechanisms by which REHIT improves VO$_2$max.
References


**Figure Legends:**

**Figure 1** Changes in VO$_2$max following SIT (panel A) and following a no-intervention control (panel B). Data is presented as mean and SD (bars) on the primary y-axis, or as individual change scores (black dots) on the secondary y-axis.

**Figure 2** Individual changes in VO$_2$max following SIT classified against either a zero (panel A) or clinically relevant thresholds of 1.75 ml·kg$^{-1}$·min$^{-1}$ (light blue dashed and dotted line) and 3.5 ml·kg$^{-1}$·min$^{-1}$ (dark blue dashed line) (panel B). Dots are individual changes and error bars are 50% confidence intervals. Red square = likely non-responder, orange diamond = uncertain responder, light blue circle = likely responder (>0 ml·kg$^{-1}$·min$^{-1}$ in panel A and >1.75 ml·kg$^{-1}$·min$^{-1}$ in panel B). In panel B, dark blue circle = likely responder >3.5 ml·kg$^{-1}$·min$^{-1}$.

Pie charts show the absolute proportion (n) of participants in each category.

**Conflict of Interest Statement:**

All authors confirm they have no conflict of interest to declare.