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1 **Effect of number of sprints in a SIT session on change in VO₂max: a meta-analysis**

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16 **ABSTRACT**

17

18 **Purpose:** Recent meta-analyses indicate that sprint interval training (SIT) improves
19 cardiorespiratory fitness ($\dot{V}O_{2\max}$), but the effects of various training parameters on the
20 magnitude of the improvement remain unknown. The present meta-analysis examined the
21 modifying effect of the number of sprint repetitions in a SIT session on improvements in
22 $\dot{V}O_{2\max}$.

23 **Methods:** The databases PubMed and Web of Science were searched for original studies
24 that have examined pre- and post-training $\dot{V}O_{2\max}$ in adults following ≥ 2 weeks of training
25 consisting of repeated (≥ 2) Wingate-type cycle sprints, published up to 1 May 2016. Articles
26 were excluded if they were not in English, involved patients, athletes, or participants with a
27 mean baseline $\dot{V}O_{2\max}$ of $>55 \text{ mL}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$ or a mean age <18 years, and if a SIT trial was
28 combined with another intervention or used intervals shorter than 10 s. A total of 38 SIT trials
29 from 34 studies were included in the meta-analysis. Probabilistic magnitude-based
30 inferences were made to interpret the outcome of the analysis.

31 **Results:** The meta-analysis revealed a likely large effect of a typical SIT intervention on
32 $\dot{V}O_{2\max}$ (mean \pm 90 CL %: $7.8\% \pm 4.0\%$) with a possibly small modifying effect of the
33 maximum number of sprint repetitions in a training session ($-1.2 \pm 0.8\%$ decrease per 2
34 additional sprint repetitions). Apart from possibly small effects of baseline $\dot{V}O_{2\max}$ and age,
35 all other modifying effects were unclear or trivial.

36 **Conclusion:** We conclude that the improvement in $\dot{V}O_{2\max}$ with SIT is not attenuated with
37 fewer sprint repetitions, and possibly even enhanced. This means that SIT protocols can be
38 made more time-efficient, which may help SIT to be developed into a viable strategy to
39 impact public health.

40

41 **Key words:** systematic review; cardiorespiratory fitness; aerobic capacity; sprint interval
42 training

43 1 INTRODUCTION

44

45 The global increase in prevalence of noncommunicable diseases over the past decades (33)
46 can be attributed, at least in part, to the low levels of physical activity undertaken by the
47 majority of the general population (16). In light of this, a key aim of public health
48 organisations is to increase population physical activity levels (20). Of the health markers
49 that can be improved by physical activity, maximal aerobic capacity ($\dot{V}O_2\text{max}$) is consistently
50 shown to be the strongest prognostic marker for future cardiovascular health and premature
51 death in cross-sectional studies (36, 54). Furthermore, longitudinal studies demonstrate that
52 improvements in $\dot{V}O_2\text{max}$ are associated with substantial reductions in all-cause and
53 cardiovascular mortality during follow-up (9, 41).

54 Over the past two decades, relatively high volumes of moderate-intensity aerobic exercise
55 (total time commitment ≥ 150 min per week) have consistently been recommended for
56 improving health markers (20). However, uptake of and adherence to these
57 recommendations remains low in the general population (25), with lack of time identified as
58 one of the main perceived barriers to becoming and remaining physically active (37, 39, 68).
59 Therefore, the seminal finding by Burgomaster *et al.* (12) that a training protocol consisting
60 of repeated brief 'all-out' cycle sprints (i.e. Wingate sprints) is associated with aerobic
61 adaptations, has led to substantial interest in the use of (sub)maximal high-intensity interval
62 training (HIIT) and supramaximal sprint interval training (SIT) as time-efficient
63 alternative/adjunct exercise strategies for improving $\dot{V}O_2\text{max}$ (21). The most commonly
64 studied SIT protocol consists of 4-7 repeated 30-s Wingate sprints, thus resulting in less
65 than 4 minutes of high-intensity exercise per session (72). Over the past few years, several
66 meta-analyses have reported the efficacy of SIT in increasing $\dot{V}O_2\text{max}$ (24, 51, 62, 72).
67 These have concluded that in healthy individuals, SIT improves $\dot{V}O_2\text{max}$ to a similar (24) or
68 greater extent (51) than traditional aerobic training, with greater benefits for individuals with
69 lower pre-training $\dot{V}O_2\text{max}$ (51, 72).

70 Although these findings provide strong support for the effectiveness of SIT in improving
71 $\dot{V}O_2\text{max}$, surprisingly few efforts have been made to identify 'optimal' SIT protocols, e.g.
72 protocols which will either provide the greatest increase in $\dot{V}O_2\text{max}$, or a set increase with
73 the lowest total training volume or time commitment. Weston *et al.* (72) reported a likely
74 small effect of increasing the intervention duration and a possibly moderate effect of
75 increasing the work-to-rest ratio, but no studies have meta-analysed or directly investigated
76 the potential effects of the number of sprint repetitions in a SIT session. This parameter is
77 particularly important as it has a large influence on the total duration of a training session, as
78 well as the level of fatigue (42) and affective responses (19) experienced by the participant,
79 thus influencing the likelihood of individuals taking up and adhering to a specific SIT
80 intervention (26). As the main aim of investigating SIT protocols is generally to identify a
81 time-efficient alternative to aerobic exercise, there is a need to identify the effect of this
82 training parameter on the associated increase in $\dot{V}O_2\text{max}$. Recent evidence suggests that
83 the positive effects of SIT on $\dot{V}O_2\text{max}$ can be attained with fewer sprints (22, 23, 34, 48), and
84 therefore the aim of the present study was to perform a meta-analysis to provide estimates
85 of the modifying effect of the number of sprint repetitions in SIT protocols on the increase in
86 $\dot{V}O_2\text{max}$ in untrained adult participants following training.

87 2 METHODS

88

89 2.1 Literature Search Criteria and Study Selection

90 This study was undertaken in accordance with the Preferred Reporting Items for Systematic
91 Reviews and Meta-Analyses (PRISMA) statement guidelines (52). We aimed to identify all
92 studies that have examined pre- and post-training $\dot{V}O_2\text{max}$ following a period of at least 2
93 weeks of training consisting of repeated (≥ 2) 'all-out' Wingate cycle sprints or modifications
94 thereof (e.g. studies using 10-s, 15-s, or 20-s 'all-out' sprints instead of 30-s Wingate
95 sprints). For this purpose, the electronic databases PubMed and Web of Science were
96 searched for relevant available records up to 1 May 2016, using the 28 possible
97 combinations of the independent variable search terms 'Wingate', 'all-out', 'sprint', and
98 'interval training', and the dependent variable search terms 'fitness', 'aerobic capacity',
99 'aerobic power', ' $\dot{V}O_2\text{max}$ ', ' $\dot{V}O_2\text{peak}$ ', 'oxygen uptake', and 'oxygen consumption'. Relevant
100 studies cited in recent meta-analyses were also used (24, 51, 62, 72), as well as our own
101 recent work (50). The following articles were excluded: 1) review articles / commentaries, 2)
102 articles not written in English, 3) studies concerning patients, athletes, or participants with a
103 mean baseline $\dot{V}O_2\text{max}$ of $>55 \text{ mL}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$ or a mean age <18 years, 4) animal studies, 5)
104 study-trials in which SIT was combined with another intervention; and 6) SIT studies using
105 non-cycling exercise, intervals shorter than 10 s, or intervals that were not 'all-out'. Two
106 authors (NBJV and RSM) independently conducted the literature search and data extraction,
107 and any discrepancies were resolved by consensus. The reviewers were not blinded to
108 manuscript journals or authors. After removal of duplicate records, the titles and abstracts of
109 all identified articles were screened for records that were clearly not relevant. These articles
110 were omitted before assessing the full-text versions of the remaining articles for eligibility to
111 be included in the meta-analysis. If more than one article reported data for the same
112 experiment, duplicate data for these participants were only included once. The final dataset
113 included the results of 38 trials from 34 studies (**Figure 1**).

114

115 **2.2 Data Extraction**

116 Full papers were assessed for mean absolute pre- and post-training $\dot{V}O_2\text{max}$. Absolute
117 $\dot{V}O_2\text{max}$ ($\text{L}\cdot\text{min}^{-1}$) was used rather than relative $\dot{V}O_2\text{max}$ ($\text{mL}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$) as this provides an
118 estimate of true changes in the ability to take up and use oxygen, independent of possible
119 concomitant changes in body mass. Relative $\dot{V}O_2\text{max}$ was used for the five studies for which
120 absolute $\dot{V}O_2\text{max}$ data were not available (8, 40, 46, 55, 65). Any data for $\dot{V}O_2\text{max}$ obtained
121 at intermediate time-points during the intervention were excluded. The corresponding
122 authors of papers without the required data were contacted by email; authors from 23
123 studies were contacted (1, 2, 5, 6, 10-13, 22, 23, 27, 28, 30, 31, 34, 38, 47, 55, 59-61, 65,
124 67, 74) and we received raw data from 17 studies (5, 10-13, 22, 23, 27, 28, 30, 38, 47, 55,
125 59-61, 74). Graph digitizer software (Digitizelt, Braunschweig, Germany) was used to obtain
126 the data from one study for which absolute pre- and post-training $\dot{V}O_2\text{max}$ data were only
127 available in a figure (67). The effect of training was expressed as a percentage change-
128 score. Percentage effects of SIT on $\dot{V}O_2\text{max}$ were converted to factors ($= 1 + \text{effect} / 100$),
129 log transformed for the analysis, and then back transformed to percentages. Effects were
130 weighted using percentage standard errors derived from exact p-values, or from estimated
131 errors of measurement as recommended by Weston *et al.* (72). Under the assumption that
132 studies with similar test protocols and subject characteristics would have similar typical
133 errors of measurement, the typical errors from these studies were averaged (via the
134 weighted mean variance) and assigned to the studies that did not report an exact p value (1,
135 2, 6, 34, 44, 65, 67). The SE was then calculated via the relationship between typical error
136 and SE (69). Finally, data for the following potential moderators were extracted for each
137 study: participant characteristics (sex, age, body mass index (BMI), baseline $\dot{V}O_2\text{max}$),
138 training parameters (intervention duration, total number of training sessions, maximal
139 number of sprint repetitions per training session, sprint duration, sprint/recovery ratio, sprint
140 resistance), and study-type (controlled / uncontrolled; dummy variable). For trials with a no-

141 exercise control group, the effect entered into the meta-analysis was intervention minus
142 control. Data for aerobic exercise comparator groups were not included in the meta-analysis.

143

144 **2.3 Statistical Analysis**

145 To evaluate the extent of publication bias, a funnel plot of model residuals versus their
146 corresponding standard errors was inspected for evidence of asymmetrical scatter (72). This
147 approach takes into account any heterogeneity explained by the meta-regression, which is
148 not accounted for in standard funnel plots of observed effects vs. their standard errors. No
149 evidence of asymmetrical scatter was apparent (**Figure 2**).

150 A mixed effects meta-regression model was conducted using the 'metafor' package in R
151 (version 3.2.4, R Foundation for Statistical Computing, Vienna, Austria) (70). The overall
152 effect of SIT on $\dot{V}O_2\text{max}$ was evaluated using the mean values of the covariates. The
153 modifying effects of covariates were evaluated as the difference between levels (e.g.
154 male/female) for nominal variables. For numeric variables, effects were evaluated as the
155 change in $\dot{V}O_2\text{max}$ associated with a two standard deviation (SD) change in the predictor
156 (i.e. a typically low vs. a typically high value (32)), or a practically relevant value (e.g. three
157 additional SIT sessions would typically constitute an additional week of training). The
158 random effects in the model specified a between-study SD, representing the typical
159 difference in the true value of the effect in different study settings, plus a within-study
160 random effect to account for within-study repeated measurements (a control treatment
161 and/or more than one training treatment) (72). The SD was doubled before interpreting its
162 magnitude with the scale used to interpret fixed effects (63), for the same reason that the
163 magnitude of the effect of a linear covariate is evaluated with two SD of the covariate (32).
164 We performed a sensitivity analysis to determine whether the inference relating to the
165 modifying effect of maximum number of sprints was substantially altered when two
166 potentially influential studies (with 12 and 15 maximum sprints, respectively (31, 61)) were
167 removed from the analysis.

168 We used magnitude-based inferences to provide an interpretation of the real-world
169 relevance of the outcomes. Uncertainty in effect estimates was expressed as $\pm 90\%$
170 confidence limits, and as the likelihood that the true value was beneficial, trivial, or harmful in
171 relation to threshold values for benefit (improved fitness) and harm (reduced fitness) (32).
172 The overall effect of SIT on $\dot{V}O_2\text{max}$ was interpreted as a clinical outcome, whereby an
173 effect was deemed unclear if the chance that the true value was beneficial was $>25\%$, with
174 odds of benefit relative to odds of harm (odds ratio) of <66 . Modifying effects were evaluated
175 mechanistically and deemed unclear if the likelihood that the true value was beneficial *and*
176 harmful were both $>5\%$. Otherwise, the effect was deemed clear, and was qualified with a
177 probabilistic term using the following scale: $<0.5\%$, most unlikely; $0.5\text{-}5\%$, very unlikely; 5-
178 25% , unlikely; $25\text{-}75\%$, possible; $75\text{-}95\%$, likely; $95\text{-}99.5\%$, very likely; $>99.5\%$, most likely.
179 As robust anchors for the smallest worthwhile clinical and practical effects relating to
180 $\dot{V}O_2\text{max}$ were not available, standardised effect thresholds of 0.2, 0.6 and 1.2 SD were
181 adopted for small, moderate and large effects, respectively (72). Here, the SD related to the
182 average between-subject variances for baseline $\dot{V}O_2\text{max}$; these corresponded to magnitude
183 thresholds of 1.0%, 2.9% and 5.8%.

184 **3 RESULTS**

185

186 Data available for the 34 studies and 38 trials included in the meta-analysis are shown in
187 **Table 1** and **Figure 3**. The meta-analysis indicated an overall likely large effect of an
188 'average' SIT protocol on $\dot{V}O_{2\max}$ (mean \pm 90 CL % effect on the increase in $\dot{V}O_{2\max}$: $7.8 \pm$
189 4.0% ; **Table 2**). A possibly small effect was evident for the modifying effect of the maximum
190 number of sprint repetitions in a training session ($-1.2 \pm 0.8\%$ decrease per 2 additional
191 sprint repetitions; **Figure 4a**). The percentage chances that the modifying effect was
192 negative, trivial or positive were calculated to be 62.7%, 37.3% and 0.0% respectively. There
193 were possibly small effects of baseline $\dot{V}O_{2\max}$ ($-1.5 \pm 1.9\%$ decrease per $10 \text{ mL}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$
194 higher baseline $\dot{V}O_{2\max}$; **Figure 4b**) and age ($-1.1 \pm 1.2\%$ decrease per 7 y increase;
195 **Figure 4c**). All other modifying effects (intervention duration, number of sessions, sprint
196 duration, recovery time, sprint resistance, BMI, sex, and study type) were unclear or trivial
197 (**Table 2**). Unexplained variance between studies was $2.2 \pm 0.8\%$ (likely moderate). The
198 inference relating to the effect of maximum number of sprint repetitions was not altered when
199 the two studies with the highest number of sprint repetitions (31, 61) were removed from the
200 analysis ($-1.0 \pm 1.1\%$; possibly small decrease; chances that the modifying effect was
201 negative, trivial or positive of 51.6%, 48.2% and 0.0% respectively).

202 4 DISCUSSION

203

204 The main aim of the present meta-analysis was to examine the modifying effect of the
205 number of sprint repetitions in a SIT session on the increase in $\dot{V}O_2\text{max}$ following training.
206 Using data from 34 training studies and 418 participants we demonstrate that the
207 improvement in $\dot{V}O_2\text{max}$ with SIT is not attenuated with fewer sprint repetitions, and possibly
208 even enhanced. Considering the low physical activity levels in the general population (25),
209 and the fact that lack of time is consistently identified as one of the main perceived barriers
210 to becoming and remaining physically active (37, 39, 68), this finding has implications for the
211 design of practical SIT interventions for improving general health. SIT protocols have the
212 potential to be the most time-efficient interventions that are associated with improvements in
213 key health markers, but due to the need for recovery intervals following sprints, this potential
214 can only truly be achieved if the number of sprint repetitions is low. Therefore, our
215 observation that reducing the number of sprint repetitions will not attenuate the increase in
216 $\dot{V}O_2\text{max}$ associated with SIT, and in fact may possibly improve the effect, is an important
217 novel finding.

218 Based predominantly on the results of studies investigating the dose-response relationship
219 between regular *aerobic* exercise and improvements in health markers, it has generally been
220 accepted that at a given exercise intensity a greater volume of exercise training (in terms of
221 training duration and frequency) is associated with greater improvements in $\dot{V}O_2\text{max}$ (20).
222 For example, in a clinical trial comparing low or high-intensity aerobic training protocols with
223 matched energy expenditure (Studies of a Targeted Risk Reduction Intervention through
224 Defined Exercise (STRRIDE I)) the magnitude of change in $\dot{V}O_2\text{max}$ was greater in the
225 group exercising at a higher intensity (15). Although the volume of exercise used in HIIT and
226 SIT protocols is generally reduced compared to aerobic exercise programmes (11, 45, 60),
227 the principle of a dose-response relationship has not been challenged in these studies
228 directly; it is the interaction between training volume and intensity that is used to justify the

229 lower volume. Thus, HIIT and SIT studies investigating the effects of protocols with a lower
230 intensity or a shorter sprint duration tend to increase the number of sprint repetitions (43,
231 66). Apart from two studies that demonstrated that reducing sprint duration from 30 s to
232 either 15 s (74) or 10 s (30) does not attenuate the improvement in $\dot{V}O_2\text{max}$ with SIT, there
233 have been no HIIT or SIT studies that have specifically investigated the dose-response
234 relationship between the volume of high-intensity exercise and health outcomes. Our meta-
235 analysis provides the first evidence that at 'all-out' supramaximal exercise intensities the
236 generally accepted positive association between volume of training and magnitude of
237 training adaptations does not hold true. Thus, research into the health benefits of SIT should
238 increase the focus on protocols with fewer sprints.

239 Due to the relatively low number of studies examining the effects of SIT protocols with fewer
240 than six sprint repetitions, the present meta-analysis was not powerful enough to make
241 conclusions on the optimal number of all-out sprint repetitions. Only two studies have
242 investigated the effects of a SIT protocol incorporating just two sprints (48, 50). As one of
243 these used the largest sample size of all the studies included in the review ($n=34$ (50)), the
244 mean 10% increase in $\dot{V}O_2\text{max}$ observed with this protocol (termed reduced-exertion high-
245 intensity interval training, REHIT) appears to be robust. The greatest improvement in
246 absolute $\dot{V}O_2\text{max}$ (17%) was reported by Gibala's group (22), who modified the original
247 REHIT protocol to include a third sprint. However, the total duration of this intervention was
248 12 weeks, whereas at an intermediate measurement-point after 6 weeks the increase in
249 $\dot{V}O_2\text{max}$ was 12%, very similar to the 10% and 14% improvements observed with the
250 original REHIT protocol (48, 50). Although future studies should determine whether the
251 magnitude of the response for $\dot{V}O_2\text{max}$ is different between SIT protocols incorporating 2-4
252 sprints, the data presented in the present manuscript suggest that this difference will be
253 small. If this is indeed the case, then a number of considerations support the use of the
254 smallest number of sprints, i.e. the two sprints used in the REHIT protocol. Firstly, including
255 a warm-up, recovery, and cool-down, this protocol has the potential to be the most time-

256 efficient protocol. Furthermore, a drawback of the use of SIT as a public health intervention
257 is the potential for high associated perceived exertion and negative affective responses (8,
258 21). In this light it is important to point out that the number of sprint repetitions has been
259 shown to negatively affect both of these parameters (19, 42). Therefore, effective SIT
260 protocols with fewer sprint repetitions will likely offer the best chance of sedentary target
261 populations taking up and adhering to a SIT intervention for improving health (18). With this
262 in mind, the available evidence suggests that two sprints can be recommended as effective
263 at improving the important health marker of $\dot{V}O_{2max}$. It could be argued that considering the
264 apparent linear association between the number of sprint repetitions and improvement in
265 $\dot{V}O_{2max}$ (**Figure 4a**), a single sprint could be expected to produce similar improvements with
266 a lower time-commitment. However, we have recently performed the first study to investigate
267 the effects of a single supramaximal sprint on $\dot{V}O_{2max}$, and observed no significant increase
268 compared to a no-exercise control condition in response to 4 weeks of training with a sample
269 size of $n=15$ (64). Further studies are required to confirm whether supramaximal sprints only
270 improve $\dot{V}O_{2max}$ if they are repeated. Furthermore, in light of the fact that the majority of
271 studies that have studied the effects of SIT protocols incorporating 2 or 3 sprint repetitions
272 have used 20-s sprints rather than the more commonly used 30-s sprints (22, 23, 48, 50),
273 further studies are required to investigate the shortest sprint duration that can be used
274 without attenuating the adaptations to $\dot{V}O_{2max}$.

275 Our present analysis does not provide an explanation for the possibly negative effect of
276 reducing the maximal number of sprint repetitions on improvements in $\dot{V}O_{2max}$, but a
277 discussion of possible mechanisms is warranted. The main limiting factor of $\dot{V}O_{2max}$ is
278 generally assumed to be maximal cardiac output, possibly through increased blood volume
279 (7, 53). To date no studies have examined the effect of SIT on blood volume, but there is
280 evidence in favour (17, 71) and against (35) increases in blood volume in response to HIIT.
281 Similarly, there is evidence in favour (3) and against (45) increased maximal cardiac output
282 with SIT, with the latter finding suggesting that the adaptations to SIT for $\dot{V}O_{2max}$ may be

283 peripheral in origin. Indeed, several authors have proposed that improvements in $\dot{V}O_{2\max}$
284 with SIT are caused by improved skeletal muscle oxygen extraction due to increased
285 mitochondrial density (22, 35, 55, 62, 74). Although it remains unclear whether the
286 improvement in $\dot{V}O_{2\max}$ with SIT is due to central or peripheral adaptations, we propose
287 that both increased blood volume and increased mitochondrial density could plausibly be
288 explained by the rapid glycogen depletion associated with supramaximal exercise (49).
289 Firstly, maximal rates of glycogenolysis in the initial 15 seconds of a supramaximal sprint
290 (56) are associated with the accumulation of metabolic derivatives, resulting in a hypertonic
291 intramyocellular environment, influx of water, and a transient ~15-20% drop in plasma
292 volume within a timespan of just a few minutes (49). This severe disturbance of circulatory
293 homeostasis could be a stimulus for the body to increase blood volume in response to
294 repeated SIT sessions. Secondly, glycogenolysis is associated with the release and
295 activation of glycogen-bound 5' AMP-activated protein kinase (AMPK) (57), which through
296 downstream signalling pathways involving peroxisome proliferator-activated receptor gamma
297 coactivator 1-alpha (PGC1 α , a proposed master regulator of aerobic adaptations), could be
298 a mechanism leading to increased mitochondrial density (29). Glycogen breakdown during
299 repeated supramaximal sprints has been shown to be completely attenuated by the time of
300 the third sprint (56), and it is therefore plausible, for both of these speculated mechanisms,
301 that performing just two repeated supramaximal sprints is sufficient to 'saturate' (i.e.
302 maximally activate) the adaptive response. In other words, if either increased blood volume
303 or mitochondrial density underpins the changes in $\dot{V}O_{2\max}$ with SIT, and if rapid glycogen
304 breakdown regulates those adaptations, then no additional improvements would be
305 expected if more than 2-3 sprints are performed in a training session.

306 Apart from this hypothesis it is also possible that increasing the number of sprint repetitions
307 will result in 'pacing' strategies that affect the 'all-out' nature of the sprints (e.g. reduction of
308 peak and mean power output in initial sprints), or that accumulated fatigue may reduce the
309 effectiveness of later sprints. Furthermore, the fact that increasing the number of sprint

310 repetitions does not enhance the improvement in $\dot{V}O_2\text{max}$ with SIT provides strong evidence
311 against a role for the magnitude of the acute effects of supramaximal sprints on oxygen
312 transfer, energy turnover, or total energy use, as part of the stimulus for improving $\dot{V}O_2\text{max}$
313 with SIT, because for each of these factors the stimulus should be greater with more sprint
314 repetitions.

315 A number of limitations to the present meta-analysis should be noted. Firstly, in order to be
316 of use as a practical intervention for preventing and/or treating inactivity-related chronic
317 disease, SIT interventions should also be effective at improving for example insulin
318 sensitivity and glycaemic control, blood pressure, blood lipid profile, and body composition.
319 Therefore, one limitation is that only $\dot{V}O_2\text{max}$ was included as an outcome measure in the
320 present analysis. Whereas insufficient data for a meta-analysis is available for the effects of
321 SIT on blood pressure (14, 23, 73), blood lipid profile (4, 73), and body composition (66, 73),
322 the effect of SIT on insulin sensitivity and glycaemic control has received more attention (4,
323 22, 23, 48, 50, 58, 73). However, the methods used to assess the effects of SIT on these
324 parameters have varied, with different studies using oral glucose tolerance tests (4, 48, 50,
325 73), intravenous glucose tolerance tests (22), euglycemic hyperinsulinemic clamps (58), or
326 continuous glucose monitoring (23). This means that a meta-analysis of the effects of the
327 number of sprint repetitions in a SIT protocol on insulin sensitivity and glycaemic control is
328 also currently not feasible. Nonetheless, the improvements in insulin sensitivity and
329 glycaemic control observed to date with SIT protocols incorporating two (48) or three sprints
330 (22, 23) are encouraging.

331 Secondly, due to the number of available SIT studies the power of our meta-analysis is
332 insufficient to conclude with certainty that the modifying effect of the number of sprint
333 repetitions is negative; i.e. it remains possible that in reality performing more sprints will
334 result in the same improvements in $\dot{V}O_2\text{max}$ (a chance of approximately 1 in 3). However,
335 this is not of major importance to the significance of our findings: even 'no effect' of the
336 number of sprint repetitions would lead to the logical conclusion that performing SIT

337 protocols with more than 2 or 3 sprints is unnecessary for improving $\dot{V}O_{2\max}$ in sedentary
338 individuals. Based on the present analysis, the chance that in reality the effect of performing
339 more sprints is positive was calculated as 0.0%.

340 A final limitation of our meta-analysis is that only SIT interventions using all-out intensities
341 were included. Optimising time-efficient interventions aimed at improving general health
342 requires consideration of various parameters, and exercise intensity is undoubtedly one of
343 the key parameters affecting the effectiveness of HIIT and SIT protocols. However, due to
344 the large range of intensities used in SIT and HIIT protocols (~80%-350% of $\dot{V}O_{2\max}$) we felt
345 it was important to attempt to 'control' for this variable in the present analysis by including
346 only studies that used 'all-out' cycling exercise. Nonetheless, there is a clear need for
347 studies examining the effect of the number of sprint repetitions at lower exercise intensities,
348 e.g. in HIIT studies.

349 In conclusion, in the present meta-analysis we demonstrate that SIT is possibly more
350 effective at improving $\dot{V}O_{2\max}$ if fewer sprint repetitions are performed in a training session.
351 Considering the proclaimed aim of SIT to provide a time-efficient alternative / adjunct to high-
352 volume moderate-intensity aerobic exercise, this finding has important implications for the
353 design of practical SIT interventions. We put forward that SIT research should move away
354 from further characterising the commonly used 4-7 x 30-s Wingate protocol, and towards
355 establishing acceptable and effective protocols. This will require more studies to examine the
356 modifying effects of a range of training parameters (including number of sprint repetitions,
357 sprint duration, sprint intensity, and training frequency) on adaptations to key health markers,
358 as well as exercise enjoyment and acceptability, perceived exertion, and the potential to
359 cause negative affective responses.

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364 clearly, honestly, and without fabrication, falsification, or inappropriate data manipulation.
365 The results of the present study do not constitute endorsement by ACSM.

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Figure 1: Flow diagram of the study selection process

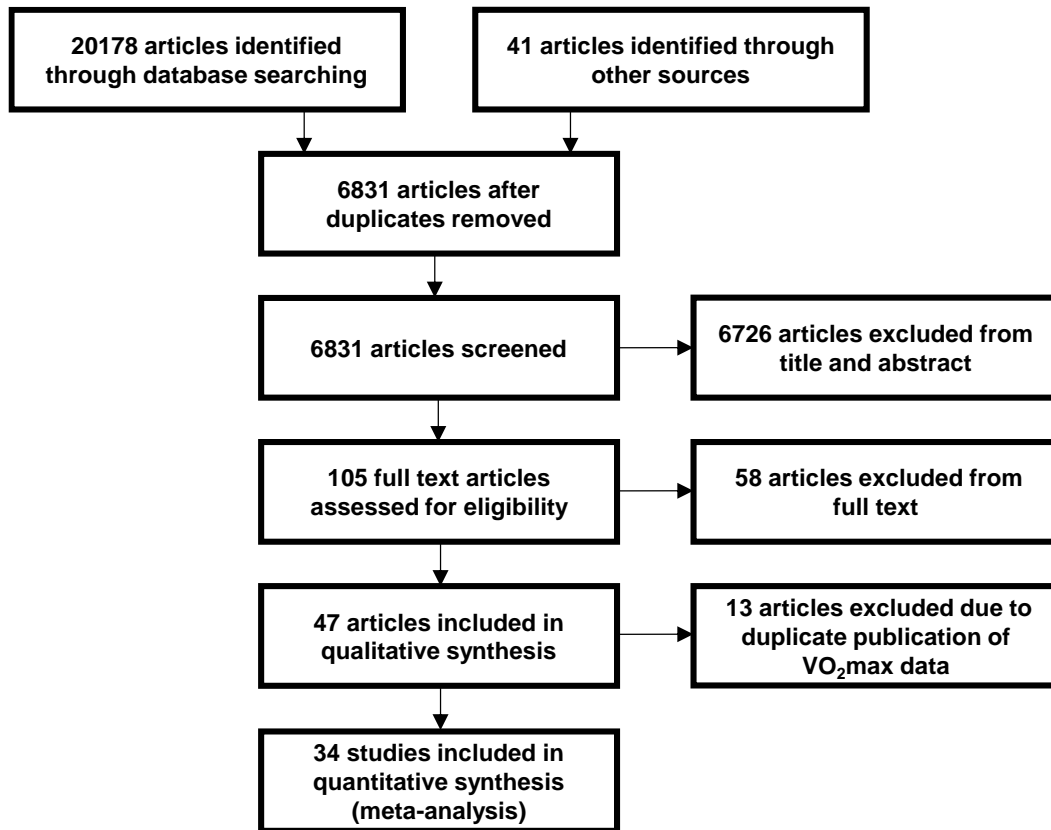


Figure 2: Funnel plot of model residuals versus their corresponding standard errors, with 90% confidence interval region

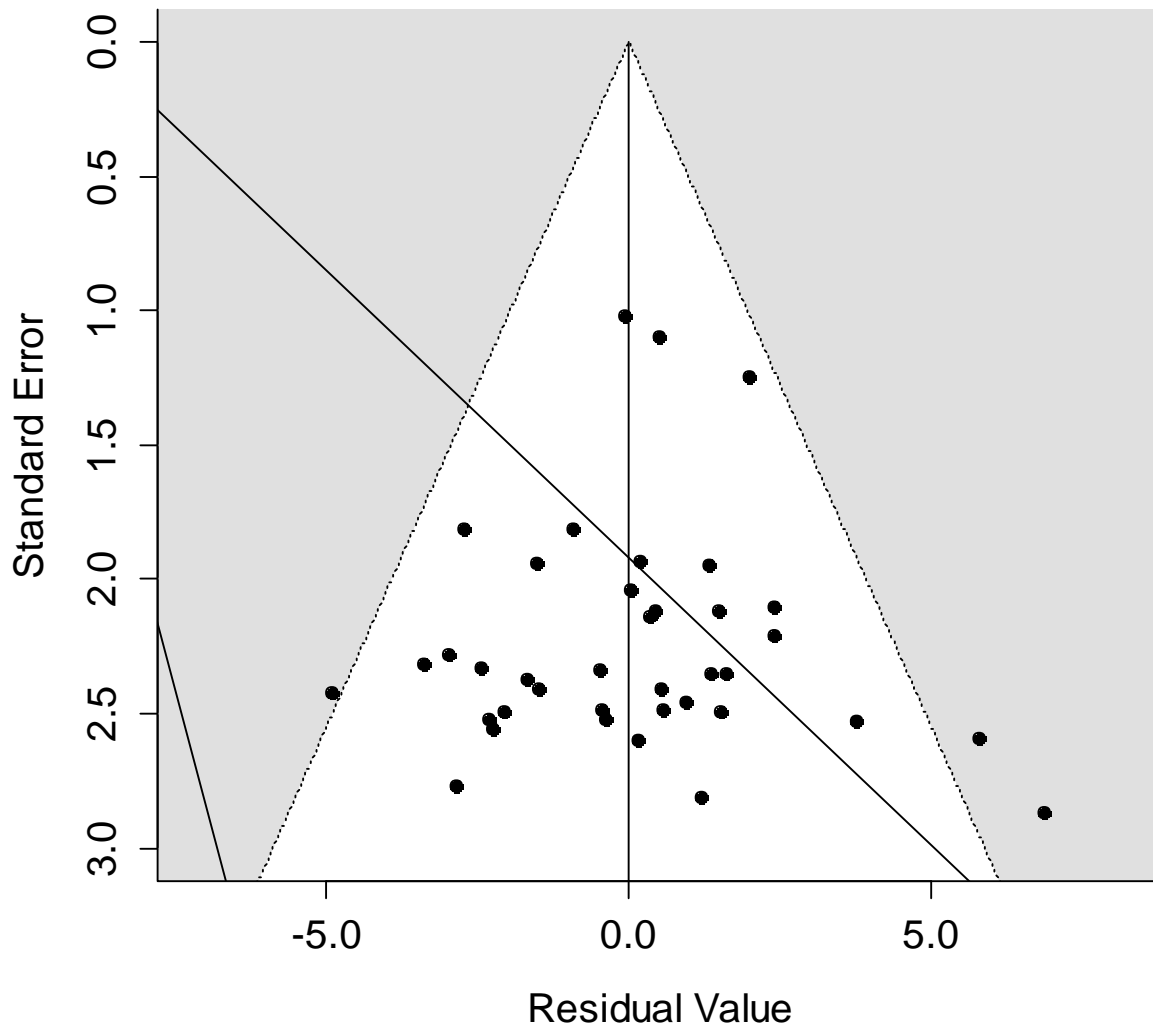


Figure 3: Main effects of SIT on $\dot{V}O_{2\max}$

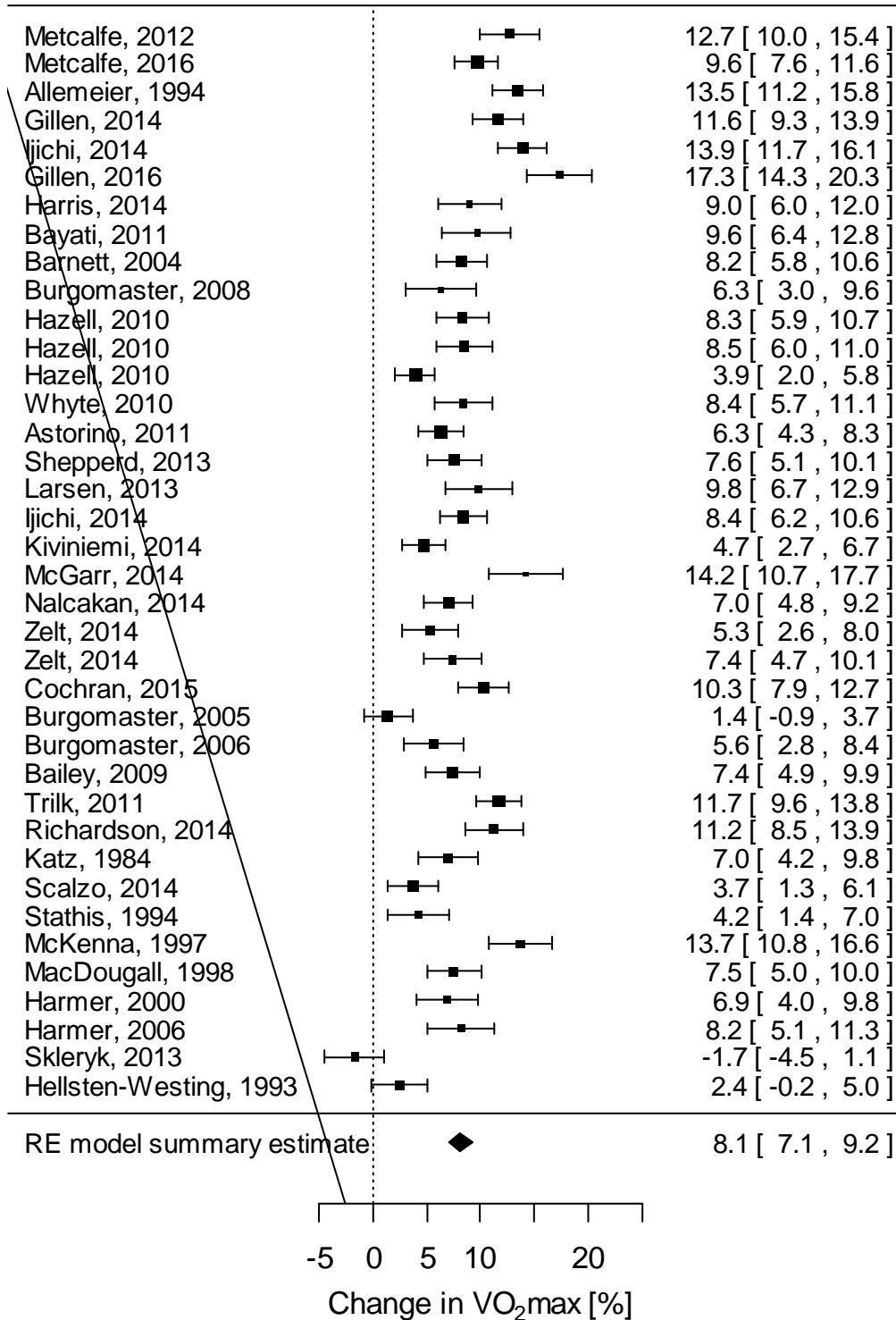


Figure 4: Modifying effects of number of sprint repetitions (A), baseline $\dot{V}O_2\text{max}$ (B), and age (C) on the effect of SIT on $\dot{V}O_2\text{max}$. Data-points represent individual trials included in the meta-analysis, and the size of the data-point is proportional to study weighting. Solid and dotted lines represent the effect of the moderator \pm 90% confidence limits respectively.

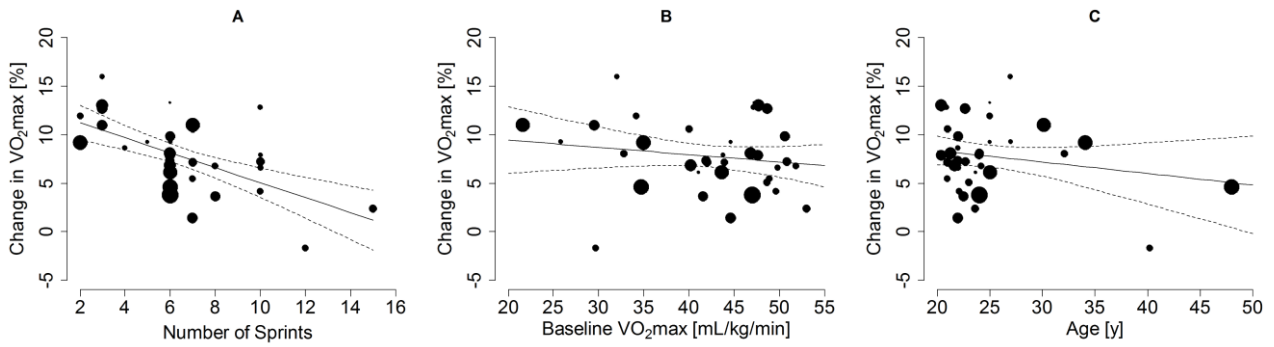


Table 1: Training effects, training protocol parameters, and participant characteristics for the studies included in t

Reference	Study design	SIT-group sample size (n)	Proportion of men	Mean baseline $\dot{V}O_2\text{max}$ ($\text{mL}\cdot\text{kg}^{-1}\cdot\text{min}^{-1}$)	Mean age (y)	Mean BMI ($\text{kg}\cdot\text{m}^{-2}$)	Training duration (weeks)	Total training sessions	Sprint duration (s)	Recovery duration (s)
Metcalfe (47)	C	11	0.45	34.2	25.0	23.5	6	18	20	200
Metcalfe (48)	NC	34	0.50	35.0	34.1	24.6	6	18	20	200
Allemeier (1)	C	11	1.00	48.7	22.7	24.8	6	15	30	120
Gillen (21)	NC	14	0.50	29.5	30.0	-	6	18	20	120
Ijichi (33)	C	10	1.00	47.7	20.4	21.0	4	20	30	600
Gillen (20)	C	9	1.00	32.0	27.0	27.0	12	36	20	120
Harris (27)	C	6	0.00	35.0	22.0	23.6	4	12	30	270
Bayati (7)	C	8	1.00	44.6	25.0	23.7	4	12	30	240
Barnett (5)	C	8	1.00	47.6	20.4	-	8	24	30	180
Burgomaster (10)	C	10	0.50	41.0	23.6	23.6	6	18	30	270
Hazell (28)	C	13	0.81	47.0	24.0	24.7	2	6	30	240
Hazell (28)	C	13	0.81	47.0	24.0	24.7	2	6	10	240
Hazell (28)	C	13	0.81	47.0	24.0	24.7	2	6	10	120
Whyte (68)	NC	10	1.00	32.8	32.1	30.3	2	6	30	270
Astorino (2)	C	20	0.55	43.6	25.0	24.1	2	6	30	300
Shepperd (56)	C	8	1.00	41.9	22.0	24.8	6	18	30	270
Larsen (39)	NC	8	1.00	25.8	27.0	26.8	2	6	30	240
Ijichi (33)	C	10	1.00	46.8	21.3	22.2	4	10	30	600
Kiviniemi (37)	C	13	1.00	34.7	48.0	25.6	2	6	30	240
McGarr (45)	C	8	0.75	47.2	25.0	-	2	8	30	240
Nalcakan (52)	C	8	1.00	40.2	21.7	25.5	7	21	30	270
Zelt (69)	C	11	1.00	48.6	23.0	25.0	4	12	30	270
Zelt (69)	C	12	1.00	43.9	22.0	26.0	4	12	15	285
Cochran (12)	C	12	1.00	50.6	22.0	25.7	6	18	30	240
Burgomaster (11)	C	8	0.75	44.6	22.0	25.6	2	6	30	240
Burgomaster (9)	C	8	1.00	48.9	21.0	23.8	2	6	30	240
Bailey (4)	C	8	0.63	42.0	21.0	23.7	2	6	30	240
Triik (63)	C	14	0.00	21.6	30.1	35.7	4	12	30	240
Richardson (54)	C	9	0.56	40.0	21.0	23.8	2	6	30	240
Katz (34)	NC	8	1.00	51.8	24.2	-	8	32	30	240
Scalzo (55)	NC	21	0.52	41.5	22.5	22.4	3	9	30	240
Stathis (61)	NC	8	0.75	49.6	22.1	-	7	21	30	180
McKenna (46)	NC	8	1.00	47.1	20.9	23.7	7	21	30	180
MacDougall (43)	NC	12	1.00	50.8	22.7	24.0	7	21	30	180
Harmer (25)	NC	7	1.00	49.8	22.0	23.5	7	21	30	180
Harmer (26)	C	7	0.71	43.7	24.0	23.8	7	21	30	180
Sklieryk (57)	C	8	1.00	29.7	40.2	32.2	2	6	10	80
Hellsten-Westing (29)	NC	11	1.00	53.0	23.6	-	6	18	10	50

Abbreviations: BM - body mass, BMI - body mass index, C - controlled, NC - not controlled, SE - standard error, SIT - sprint interval tra

Table 2 Main effect of SIT on $\dot{V}O_{2\max}$ and modifying effects

	Effect on $\dot{V}O_{2\max}$ (mean %, \pm 90% CL)	Inference
Main effect:	7.8 \pm 4.0	Likely large increase
Modifying effects:		
2 more sprint repetitions*	-1.2 \pm 0.8	Possibly small decrease
3 more training sessions*	0.7 \pm 0.4	Likely trivial change
10 s longer sprint duration*	0.6 \pm 1.3	Possibly trivial change
60 s longer recovery interval duration*	0.2 \pm 0.3	Most likely trivial change
3% of BM greater sprint resistance	1.0 \pm 2.3	Unclear
10 mL·kg ⁻¹ ·min ⁻¹ lower baseline $\dot{V}O_{2\max}$	1.5 \pm 1.9	Possibly small increase
7 years higher age	-1.1 \pm 1.2	Possibly small decrease
6.2 kg·m ⁻² higher BMI	0.8 \pm 2.7	Unclear
Female sex	-0.2 \pm 3.5	Unclear
Uncontrolled study	-0.9 \pm 2.1	Unclear

*The reference condition is an intervention using 14 SIT sessions and a maximum of 7 repeated 30-s sprints with 240 s recovery. Effects of SIT are presented as the % change compared to pre-training. *, indicates a practically relevant value was chosen to evaluate the effect magnitude; other numeric modifiers were evaluated as a 2 x SD change in the parameter. Abbreviations: BMI: body mass index, CL: confidence limits, SIT: sprint interval training, $\dot{V}O_{2\max}$: maximal aerobic capacity.*