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3 High intensity interval training (HIIT) increases insulin-
4 like growth factor-I (IGF-I) in sedentary aging men but
5 not masters' athletes: An observational study.

6

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8

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22

23 **Abstract**

24 *Introduction:* The aim of this investigation was to examine the impact high intensity interval
25 training (HIIT) on serum insulin-like growth factor-I (IGF-I) in active compared with sedentary
26 aging men.

27 *Methods:* 22 lifetime sedentary (SED; 62 ± 2 years) and 17 masters' athletes (LEX; 60 ± 5
28 years) were recruited to the study. As HIIT requires preconditioning exercise in sedentary
29 cohorts, the study required three assessment phases; enrolment (phase A), following
30 preconditioning exercise (phase B), and post-HIIT (phase C). Serum IGF-I was determined by
31 electrochemiluminescent immunoassay.

32 *Results:* IGF-I was higher in LEX compared to SED at baseline ($P=0.007$, Cohen's $d=0.91$),
33 and phase B ($P=0.083$, Cohen's $d=0.59$), with only a small difference at C ($P=0.291$, Cohen's
34 $d=0.35$). SED experienced a small increase in IGF-I following preconditioning from 13.1 ± 4.7
35 to 14.2 ± 6.0 $\text{ug}\cdot\text{dl}^{-1}$ ($P=0.376$, Cohen's $d=0.22$), followed by a larger increase post-HIIT (16.9
36 ± 4.4 $\text{ug}\cdot\text{dl}^{-1}$), which was significantly elevated compared with baseline ($P=0.002$, Cohen's
37 $d=0.85$), and post-preconditioning ($P=0.005$, Cohen's $d=0.51$). LEX experienced a trivial
38 change in IGF-I from A to B (18.2 ± 6.4 to 17.2 ± 3.7 $\text{ug}\cdot\text{dl}^{-1}$ [$P=0.538$, Cohen's $d=0.19$]), and
39 a small change post-HIIT (18.4 ± 4.1 $\text{ug}\cdot\text{dl}^{-1}$ [$P=0.283$, Cohen's $d=0.31$]). Small increases were
40 achieved in fat free mass in both groups following HIIT ($P<0.05$, Cohen's $d=0.32-0.45$).

41 *Conclusions:* In conclusion, HIIT with preconditioning exercise abrogates the age associated
42 difference in IGF-I in SED and induces small improvements in both SED and LEX.

43

44 **Key words:** Exercise, growth hormone, HIIT, IGF-I, training

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47

48 INTRODUCTION

49 Insulin-like growth factor-I (IGF-I) is an endocrine and autocrine/paracrine growth
50 factor expressed by multiple cell types. In humans, serum IGF-I peaks during adolescence and
51 displays a gradual decline during middle-age. The reduction in circulating anabolic hormones,
52 namely growth hormone (GH) and IGF-I, has been termed ‘somatopause’, and suggested as a
53 potential mechanism for the atrophic sequelae of aging [1]. IGF-I is considered to play a central
54 role in the age-associated compromise of both skeletal [2], and muscle [3] integrity.

55 Exercise plays an integral role in maintaining muscle mass and function during
56 advanced age [4]. Furthermore, exercise can improve cardiovascular function, metabolic
57 health, muscular function, body composition, and quality of life, in aging cohorts [5, 6, 7]. For
58 example, Fiatarone and colleagues [8] reported significant increases in maximal strength and
59 gait speed following eight weeks of high-intensity resistance training in nonagenarians.
60 Favorable adaptations to endurance training in elderly populations center on cardioprotective
61 benefits [9], whilst high intensity interval training (HIIT) is a form of exercise largely untested
62 in aging populations despite resurgent interest younger cohorts [10, 11]. HIIT involves repeated
63 bouts of high-intensity exercise, interspersed with recover periods, proclaimed as a time-
64 efficient healthogenic strategy [10, 11] despite falling short of the recommended exercise
65 volume to improve and maintain cardiovascular health [12]. However, prior to undertaking
66 HIIT in sedentary aging cohorts, it is prudent to undertake a programme of preconditioning
67 exercise [13].

68 Whilst acute exercise-induced elevations in IGF-I are consistently reported [14], the
69 effect of exercise training on basal IGF-I is poorly understood, particularly in aging cohorts.
70 For example, eight weeks of endurance training resulted in a 19% increase in systemic IGF-I
71 in males aged 66 ± 2 years [15], yet Vitiello et al. [16] observed no change in IGF-I following
72 six months of endurance training in males aged 67 ± 1 years. Furthermore, a recent

73 investigation reported decreased systemic IGF-1 following 12 weeks resistance exercise in
74 older adults (74 ± 6 years), yet an increase in lean mass [17]. As such, the role of IGF-I in the
75 adaptive process to exercise during middle and older age remains unclear.

76 One way to identify whether a relationship exists between basal IGF-I and exercise
77 during advancing age is to compare masters' athletes with age matched sedentary counterparts.
78 Similarly, subjecting a sedentary cohort to structured exercise training may establish whether
79 basal IGF-I is influenced by an exercise intervention. A single study to date has directly
80 compared serum IGF-I in masters' athletes with controls [18], where the authors outline a lack
81 of difference between endurance runners, speed-power athletes compared with moderately
82 active controls. However, comparisons between masters' athletes and sedentary aging men
83 have not been established. Similarly, no study has examined the influence of HIIT, either with
84 or without conditioning exercise on serum IGF-I in sedentary aging men compared with
85 masters' athletes.

86 With these aspects in mind, the aims of this study were to; 1) establish whether masters'
87 athletes and sedentary controls have different serum IGF-I concentrations, and 2) determine
88 whether HIIT preceded by preconditioning exercise would impact basal IGF-I concentrations
89 in aging men. We hypothesized that: 1) IGF-I would be greater in masters' athletes compared
90 to sedentary older males, and 2) six weeks of HIIT, preceded by 6 weeks of preconditioning
91 would increase IGF-I in sedentary aging men compared with masters' athletes.

92

93 **METHODS AND MATERIALS**

94 **Participants**

95 Following approval to exercise by their general practitioner, participants provided
96 informed written consent prior to the study which was approved by the institutional ethics
97 committee. Twenty two males (62 ± 2 years, with a stature of 175 ± 6 cm, body mass of $91 \pm$

98 16 kg, and peak oxygen uptake of 28 ± 6 ml·kg·min⁻¹) comprised the lifelong sedentary group
99 (SED). Seventeen males (60 ± 5 years, with a stature of 173 ± 6 cm, body mass of 78 ± 12 kg,
100 and peak oxygen uptake of 39 ± 6 ml·kg·min⁻¹) were enrolled as lifelong exercisers (LEX) and
101 acted as a positive control group. Participants recruited for the SED group did not participate
102 in organized exercise programmes and had not done so for >30 years. The LEX group were
103 highly active exercisers and had been so for the previous >30 years. They consisted of current
104 masters' athletes in sports including water-polo, triathlon, track cycling, road cycling, and
105 distance running.

106

107 **Exercise Training**

108 To account for the contribution of aerobic conditioning exercise, participants were
109 tested at three time points (phase A, B, and C), interspersed with two six weeks training blocks
110 (12 weeks training in total [Figure 1]). As preconditioning, (training block 1), a six week
111 exercise programme that reflected the ACSM guidelines of 150 mins·wk⁻¹ of moderate to
112 vigorous exercise was prescribed to SED. SED were advised to achieve a minimum of two
113 sessions per week in accordance with the ACSM guidelines for older persons [19]. Participants
114 were given verbal instructions on the use of a Polar FT1 heart rate monitor (Polar, Kempele,
115 Finland) and exercise intensities were self-monitored, enabling recording of exercise time, and
116 average and peak heart rate. The aim was to achieve an average heart rate reserve (HRR) of
117 approximately 55% for the first two weeks of the intervention. This was increased to 60% of
118 HRR for the subsequent weeks including 5-10 s of increased intensity every 10 min. The final
119 two weeks required vigorous periods of exercise every 5 min achieving a HRR of 60-65%. The
120 mode of training was optional, and included walking, walk/jogging, jogging, and cycling. Over
121 the six week intervention, 160 ± 15 min·wk⁻¹ exercise were achieved. Whilst SED underwent
122 preconditioning, LEX maintained their habitual training. LEX recorded their weekly exercise,

123 which included type, frequency, intensity (recorded by heart rate telemetry), and duration of
124 training. Time spent in low to medium intensity (<65% heart rate reserve [HRR]), and high-
125 intensity (>65% HRR) training totalled $214 \pm 131 \text{ min}\cdot\text{wk}^{-1}$ and $67 \pm 52 \text{ min}\cdot\text{wk}^{-1}$ respectively.

126 Both groups undertook supervised HIIT programmes from phase B to C. HIIT sessions
127 were performed every five days, for six weeks (nine sessions in total). Rationale for this
128 programme is provided by our previous work which identified that five days recovery was
129 required for recovery of peak power output (PPO) post-HIIT amongst older males [20].
130 Sessions consisted of 6 x 30 s sprints at 40% PPO (determined during familiarization)
131 interspersed with 3 min active recovery on a cycle ergometer (Wattbike Ltd., Nottingham, UK).
132 Sessions were conducted in groups of 4-6 and were the sole exercise performed by both groups
133 during this time. To allow for comparison with other literature, training intensities were
134 compared with power achieved at $\text{VO}_{2\text{peak}}$. In the majority of cases, 40% of PPO exceeded
135 power at $\text{VO}_{2\text{peak}}$; in 4 cases (1 SED; 3 LEX), it exceeded 90% of power at $\text{VO}_{2\text{peak}}$ (92; 92; 96;
136 98% respectively). In SED, mean training intensity equated to $141 \pm 27\%$ of power at $\text{VO}_{2\text{peak}}$,
137 whilst in LEX, mean training intensity equated to $126 \pm 22\%$ of power output at $\text{VO}_{2\text{peak}}$.

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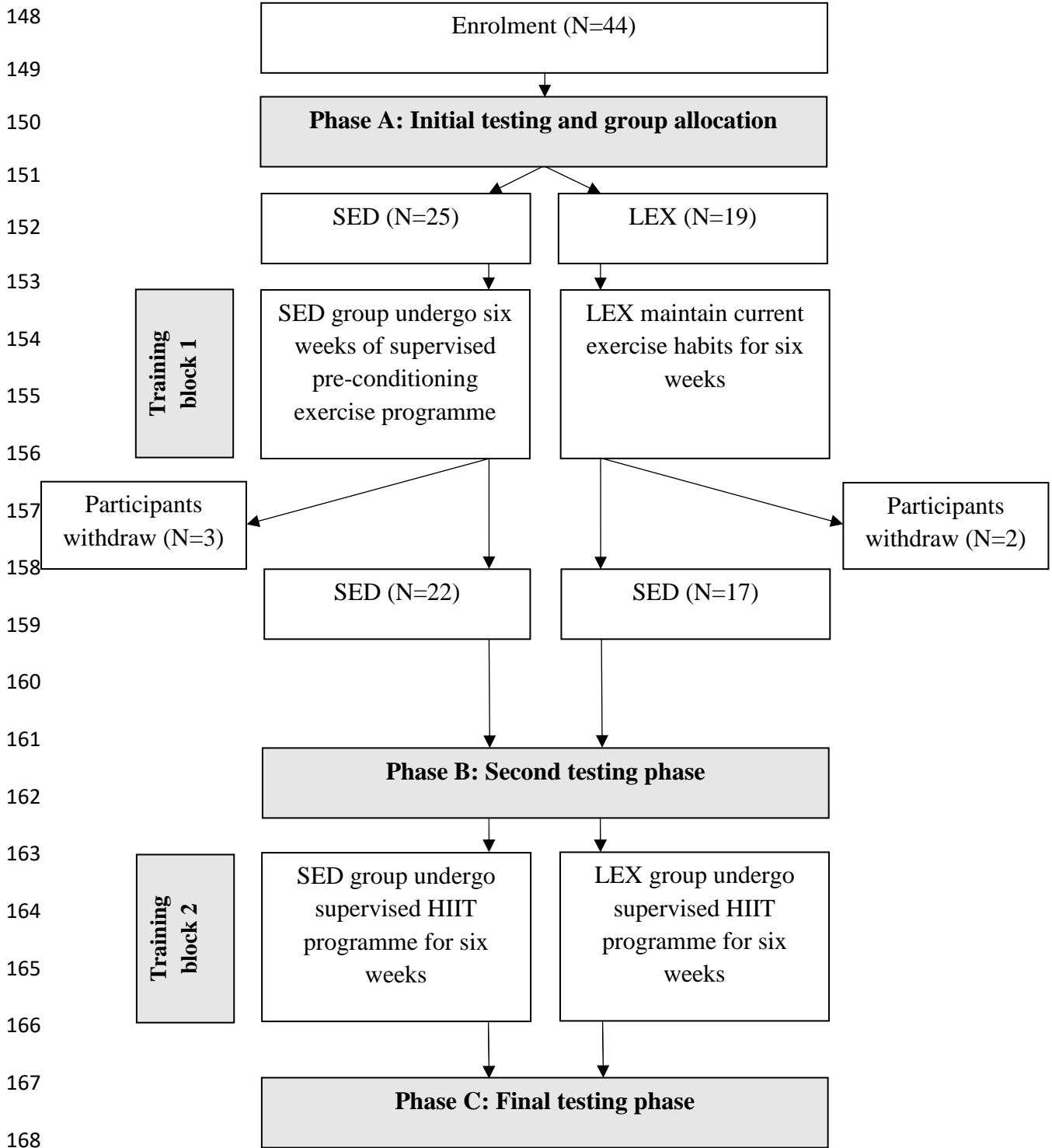
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169 **Figure 1:** The CONSORT (Consolidated Standards of Reporting Trials) flow chart depicting
 170 transit of lifelong sedentary (SED) and lifelong exercising (LEX) participants through the study.
 171 HIIT = high intensity interval training.

172

173 **Blood draws and analysis**

174 Blood samples from each participant were collected at each phase between 07:00-09:00
175 h, 48-72 hours following the last exercise session as previously described [21, 22]. Samples
176 were obtained using a 20-gauge disposable needle equipped with Vacutainer tube holder
177 (Becton Dickinson, Oxford, UK) following an overnight fast and 20 min supine rest. Blood
178 draws were conducted from the antecubital vein, by the same phlebotomist to control for
179 biological variation, and inter- and intra-subject variation. Approximately 14 mL of blood was
180 drawn into two 10 mL serum separator tubes and allowed to clot at room temperature prior to
181 being centrifuged at 6,000 rpm at 15°C for 15 min. Resultant serum was divided into
182 appropriate aliquots and stored at -80°C until analysis. Serum concentrations of IGF-I were
183 measured by electrochemiluminescent immunoassay on the E601 module of the Roche Cobas
184 6000 (Burgess Hill, West Sussex, U.K.). Inter-assay coefficients of variation (CV) over a 6
185 month period were <5%. Analysis was conducted in the Clinical Biochemistry Laboratory at
186 Royal Glamorgan Hospital (Wales, UK).

187

188 **Body composition**

189 Stature was measured to the nearest 0.1 cm using a stadiometer (Seca, Birmingham,
190 UK), and body mass and body composition was determined by a multi frequency bioelectrical
191 impedance analyzer (BIA [Tanita MC-180MA Body Composition Analyzer, Tanita UK Ltd.]).
192 GMON software (v1.7.0, Tanita UK Ltd.) was used to determine absolute and relative body
193 fat. Fat free mass (FFM) was calculated by subtracting fat mass from total body mass.

194

195 **Peak oxygen uptake and peak power output**

196 Peak oxygen uptake (VO_{2peak}) was determined using a Cortex II Metalyser 3B-R2
197 (Cortex, Biophysik, Leipzig, Germany) utilizing methods previously described [23] and a

198 modified Storer Test [24]. PPO was established using the 6 s Herbert test [25] on an air-braked
199 cycle ergometer (Wattbike Ltd., Nottingham, UK). Order of measurement was; blood
200 sampling, body composition, PPO determination, and VO_{2peak} assessment.

201

202 **Statistical Analysis**

203 Following a Shapiro-Wilk test of normality and Levene's test for homogeneity of
204 variance, a 2 x 3 (group [SED, LEX] x time [phase A, B, C]) repeated measures analysis of
205 variance (ANOVA) with *post hoc* Tukey's LSD tested for differences between groups and
206 between time points. To determine relationships between variables, a Pearson's correlation
207 coefficient was conducted. Alpha level was set *a priori* at $P \leq 0.05$, and effect size is displayed,
208 and classified as <0.2 =trivial, $0.2-0.49$ =small, $0.5-0.79$ =moderate, and >0.8 =large. Data are
209 presented as mean \pm standard deviation (SD). As LEX maintained their current exercise habits
210 between phase A and B, we used these two samples to determine the absolute minimum
211 threshold for a meaningful change in IGF-I (expressed as a percentage).

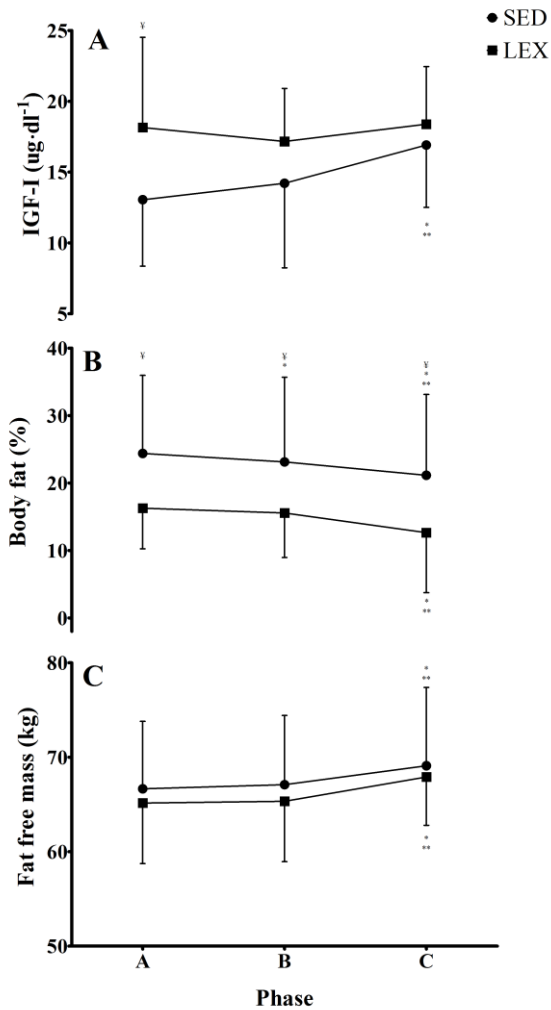
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213 **RESULTS**

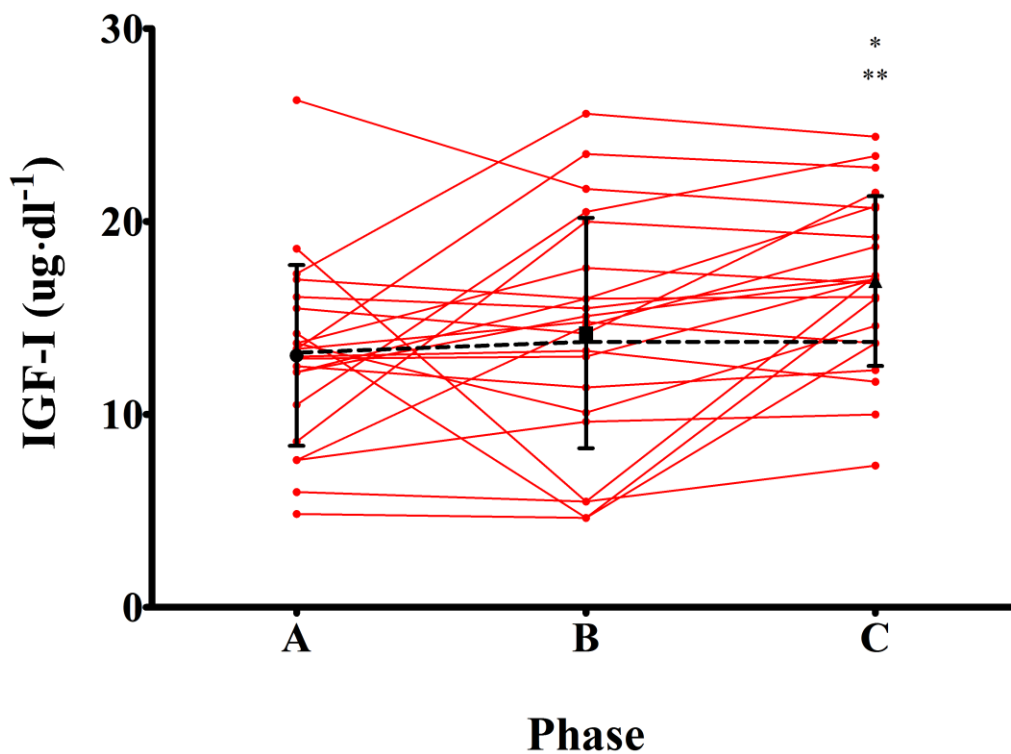
214 Basal IGF-I concentrations at phase A, B, and C for both groups are displayed in Figure
215 2A, and in SED in Figure 3. IGF-I was largely higher in LEX compared to SED at baseline
216 ($P=0.007$, Cohen's $d=0.91$), moderately higher after preconditioning ($P=0.083$, Cohen's
217 $d=0.59$), whilst a small difference existed post-HIIT ($P=0.291$, Cohen's $d=0.35$).

218 SED IGF-I increased post-HIIT compared to baseline (16.9 ± 4.4 and 13.1 ± 4.7 $ug \cdot dl^{-1}$
219 ¹ respectively [$\sim 29\%$ increase; $P=0.002$, Cohen's $d=0.85$]) and compared to preconditioning
220 ($\sim 21\%$ increase; 14.2 ± 6.0 $ug \cdot dl^{-1}$ [$P=0.005$, Cohen's $d=0.51$]). Preconditioning accounted for
221 $\sim 8\%$ of the increase from baseline ($P=0.376$, Cohen's $d=0.22$).

222 LEX experienced a trivial ~1% difference in IGF-I post-HIIT compared to baseline
 223 (18.4 ± 4.1 and 18.2 ± 6.4 $\mu\text{g}\cdot\text{dl}^{-1}$ respectively [$P=0.901$, Cohen's $d=0.04$]), and a ~7% increase
 224 post-HIIT compared to phase B (17.2 ± 3.7 $\mu\text{g}\cdot\text{dl}^{-1}$ [$P=0.283$, Cohen's $d=0.31$]). A trivial change
 225 in IGF-I was observed in LEX from phase A to B, equal to 5% ($P=0.538$, Cohen's $d=0.19$).
 226



227
 228 **Figure 2:** (A) Insulin-like growth factor (IGF-I), (B), body fat percentage, and (C) fat free
 229 mass in a group of sedentary (SED) and lifelong exercising (LEX) older males. Data are
 230 presented as mean \pm SD. *Denotes significant differences from phase A ($P<0.05$). **Denotes
 231 significant difference between phase B and C ($P<0.05$). †Denotes significant difference
 232 between groups at this experimental phase.



233

234 **Figure 3:** Insulin-like growth factor (IGF-I) in a group of sedentary older males. Data are
 235 presented as individual data points in addition to mean \pm SD. The minimum threshold for a
 236 meaningful change compared to baseline has been added as a dashed line. *Denotes significant
 237 differences from phase A ($P < 0.05$). **Denotes significant difference between phase B and C
 238 ($P < 0.05$).

239

240 Body composition is displayed in Figure 2B and 2C. Body fat percentage was greater
 241 in SED than LEX at baseline ($P = 0.013$, Cohen's $d = 0.862$), after preconditioning ($P = 0.031$,
 242 Cohen's $d = 0.74$), and post-HIIT ($P = 0.020$, Cohen's $d = 0.80$).

243 HIIT decreased SED body fat percentage by $\sim 3.3\%$ compared to baseline ($21.1 \pm 12.0\%$
 244 and $24.4 \pm 11.6\%$ respectively [$P < 0.001$, Cohen's $d = 0.28$]) and $\sim 2.2\%$ compared to
 245 preconditioning ($23.1 \pm 12.6\%$ [$P = 0.008$, Cohen's $d = 0.16$]). A $\sim 1.3\%$ decrease occurred as a
 246 result of preconditioning alone ($P = 0.006$, Cohen's $d = 0.10$). LEX body fat percentage decreased

247 from $16.3 \pm 6.0\%$ at baseline to $12.6 \pm 8.9\%$ post-HIIT ($P=0.006$, Cohen's $d=0.48$), which was
248 also lower than at phase B ($15.6 \pm 6.6\%$ [$P=0.020$, Cohen's $d=0.37$]). LEX body fat percentage
249 was trivially decreased from phase A to B ($P=0.079$, Cohen's $d=0.11$).

250 FFM was not significantly different between SED and LEX at A, B, or C ($P=0.439$ -
251 0.61). SED FFM was similar at baseline and following preconditioning (66.7 ± 7.1 kg and 67.1
252 ± 7.3 kg respectively [$P=0.336$, Cohen's $d=0.06$]). This was followed by a $\sim 3.0\%$ increase post-
253 HIIT (69.1 ± 8.3 kg [$P=0.005$, Cohen's $d=0.26$]), which was $\sim 3.6\%$ greater than at baseline
254 ($P=0.001$, Cohen's $d=0.32$). LEX FFM was unchanged from phase A to B (65.2 ± 6.4 kg and
255 65.3 ± 6.4 kg respectively [$P=0.590$, Cohen's $d=0.03$]), followed by a $\sim 4.0\%$ increase post-
256 HIIT (67.9 ± 5.1 kg [$P=0.008$, Cohen's $d=0.45$]), which was $\sim 4.1\%$ greater than at baseline
257 ($P=0.006$, Cohen's $d=0.48$).

258 At baseline, a weak negative correlation was present between IGF-I and BMI ($P=0.016$,
259 $r=-0.385$), and IGF-I and body fat percentage ($P=0.030$, $r=-0.345$), whereas a moderate
260 relationship existed between IGF-I and FFM ($P=0.087$, $r=0.600$). The change in IGF-I from
261 pre- to post-HIIT was not significantly associated with change in FFM ($P=0.860$, $r=0.029$) or
262 body fat percentage ($P=0.860$, $r=-0.029$). There was a strong significant correlation between
263 change in FFM and body fat percentage from baseline to post-HIIT ($P<0.001$, $r=-0.904$).

264

265 **DISCUSSION**

266 The main findings from the present study were that 1) masters' athletes (LEX) have
267 higher basal IGF-I concentrations than age-matched sedentary (SED) counterparts and that 2)
268 a programme of HIIT training that includes preconditioning exercise increases IGF-I
269 concentrations in SED compared with LEX. These data provided preliminary evidence for the
270 positive influence of HIIT on tissue growth factors in lifelong sedentary aging men.

271 Our data are in agreement with some [26], but not all [17, 27] previous investigations
272 in reporting increased IGF-I following exercise interventions. However, it is evident that a)
273 previous exercise training, and b) exercise intensity, mediate the IGF-I response to training.
274 LEX exhibited greater IGF-I compared to SED at baseline, supporting our hypothesis that
275 lifelong exercise would be associated with higher basal IGF-I concentrations. Preconditioning
276 accounted for ~8% of the IGF-I increase in SED, whereas HIIT accounted for a further ~21%,
277 despite a reduction in training volume from ~160 min·wk⁻¹ to ~3-6 min·wk⁻¹. As such, it
278 appears that HIIT likely induces greater increases in basal IGF-I compared with a higher
279 volume of lower intensity exercise in SED. However, this remains preliminary until confirmed
280 by a randomized controlled trial.

281 Arnarson et al. [17] observed that following 12 weeks resistance training, lean body
282 mass increases were negatively associated with IGF-I changes, leading these authors to
283 hypothesize that during periods of anabolism, IGF-I was redistributed from circulation into
284 tissue. However, our data do not support such a redistribution, as SED FFM was increased
285 post-HIIT, concomitantly with an increase in systemic IGF-I. Moreover, changes in FFM were
286 not associated with changes in IGF-I in either group. Hofmann and colleagues [27] add further
287 ambiguity to the relationship between lean mass and IGF-I, reporting increased muscle quality
288 and chair stand performance after six months of resistance training, without serum IGF-I
289 perturbations at three, or six months. As such, despite the *in vitro* evidence demonstrating IGF-
290 I to be critical for muscle hypertrophy [28], further research is required to untangle the known
291 associations between exercise, IGF-I, and lean body mass, in older adults.

292 HIIT appears to induce a number of favorable adaptations in older adults, including
293 vascular function [23], quality of life [7], muscle power [29], and mitochondrial function [30].
294 Despite the evident benefits of HIIT to older adults, we suggest caution when inducting older
295 adults onto a HIIT programme, as we have previously demonstrated delayed recovery in older,

296 compared to young, males [20]. Moreover, Donath et al. [31] reported increased postural sway
297 following a single HIIT session in seniors (70 ± 3 years), suggesting an increased likelihood of
298 falls acutely post-HIIT. As such, pragmatic periodization is necessary to allow adequate
299 recovery and mitigate risk arising from HIIT [13].

300 One limitation of the present investigation is that groups were not BMI-matched (SED
301 were heavier than LEX). As such, differences in IGF-I between groups could have been caused
302 by differences in body composition, rather than differences in habitual exercise, which is
303 supported by the moderate correlations between IGF-I and body composition parameters.
304 However, given the interrelationship between exercise and body composition, delineating these
305 effects was outside the scope of the present study and requires further investigation.

306

307 **Conclusion**

308 In conclusion, lifelong sedentariness was associated with lower systemic IGF-I
309 compared to masters' athletes. However, a programme of HIIT training that includes
310 preconditioning exercise increases IGF-I concentrations in SED compared with LEX and
311 provides preliminary evidence for the positive influence of HIIT on tissue growth factors in
312 lifelong sedentary aging men.

313

314 Authors declare they have no conflict of interest.

315

316

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