



Cronfa - Swansea University Open Access Repository

This is an author produced version of a paper published in: *Journal of Sports Sciences*

Cronfa URL for this paper: http://cronfa.swan.ac.uk/Record/cronfa48871

Paper:

McNarry, M., Lewis, M., Wade, N., Davies, G., Winn, C., Eddolls, W., Stratton, G. & Mackintosh, K. (2019). Effect of asthma and six-months high-intensity interval training on heart rate variability during exercise in adolescents. *Journal of Sports Sciences*, 1-8.

http://dx.doi.org/10.1080/02640414.2019.1626115

This item is brought to you by Swansea University. Any person downloading material is agreeing to abide by the terms of the repository licence. Copies of full text items may be used or reproduced in any format or medium, without prior permission for personal research or study, educational or non-commercial purposes only. The copyright for any work remains with the original author unless otherwise specified. The full-text must not be sold in any format or medium without the formal permission of the copyright holder.

Permission for multiple reproductions should be obtained from the original author.

Authors are personally responsible for adhering to copyright and publisher restrictions when uploading content to the repository.

http://www.swansea.ac.uk/library/researchsupport/ris-support/

1 Effect of Asthma and Six-Months High-Intensity Interval Training on Hea	rt Rate
---	---------

2 Variability during Exercise in Adolescents

- 3
- ⁴ ¹McNarry MA, ¹Lewis, M.J., ¹Wade N, ²Davies GA, ^{1,2}Winn CON, ¹Eddolls WTB,
- ⁵ ¹Stratton GS, ¹Mackintosh KA
- 6
- 7 ¹Applied Sports Technology Exercise and Medicine Research Centre, Swansea University,
- 8 ²Swansea University Medical School
- 9
- 10 Dr. M.A. McNarry
- 11 School of Sport and Exercise Sciences
- 12 Swansea University
- 13 Bay Campus
- 14 Swansea, SA1 8EN
- 15 Tel 01792 513069
- 16 Email: <u>m.mcnarry@swansea.ac.uk</u>
- 17
- 18
- 19 Running Title: HIIT and HRV in Children
- 20
- 21
- 22 Word Count: 4,471
- 23
- 24

- 25 Abstract
- Little is known regarding the influence of asthma and exercise, and their interaction, on
 heart rate variability in adolescents.
- 28Thirty-one adolescents with asthma $(13.7\pm0.9 \text{ years}; 21.9\pm3.9 \text{ kg} \cdot \text{m}^{-2}; 19 \text{ boys}, 12 \text{ girls})$ 29and thirty-three healthy adolescents $(13.8\pm0.9 \text{ years}; 20.3\pm3.2 \text{ kg} \cdot \text{m}^{-2}; 16 \text{ boys}, 17 \text{ girls})$ 30completed an incremental ramp test and three heavy-intensity constant-work-rate cycle31tests. Thirteen adolescents (7 boys, 6 girls; 6 asthma, 7 control) completed six-months32high-intensity interval training (HIIT) and were compared to age- and sex-matched33controls. Standard time-domain, frequency-domain and non-linear indices of heart rate34variability (HRV) were derived at baseline, three- and six-months.
- Asthma did not influence HRV at baseline or following HIIT. Total power, low frequency and normalised low frequency power, and sympathovagal balance increased at three-months in HIIT, subsequently declining towards baseline at six-months. Normalised high frequency power was reduced at three-months in both groups, which was sustained at six-months. No effects of HIIT were observed in the time-domain nor in the non-linear indices.
- HRV was not influenced by asthma, potentially because such derangements are a
 function of disease progression, severity or duration. HIIT may be associated with a
 short-term shift towards greater sympathetic predominance during exercise, perhaps
 caused by physiological overload and fatigue.
- 45 Keywords: Heart rate variability; Youth; Exercise; Non-linear; Children
- 46
- 47

48 Introduction

49	Heart rate variability (HRV) provides a non-invasive insight into the function of the
50	autonomic nervous system (TaskForce, 1996) by measuring changes in beat-to-beat
51	variability of the cardiac (RR) interval. Indeed, an altered HRV has been associated with a
52	multitude of pathological conditions, including respiratory conditions such as Chronic
53	Obstructive Pulmonary Disease and Cystic Fibrosis (e.g. Chang, Silberstein, Rambod,
54	Porszasz, & Casaburi, 2011; McNarry & Mackintosh, 2016).
55	Asthma, a chronic respiratory condition characterised by swelling of the bronchioles
56	that leads to breathlessness and wheezing (Carson et al., 2013), is one of the most common
57	respiratory diseases in the world affecting approximately 1 in 11 children (Wanrooij,
58	Willeboordse, Dompeling, & van de Kant, 2014). In adults, it is generally accepted that
59	asthma is associated with cardiac parasympathetic predominance (Lutfi, 2012, 2015), the
60	degree of which is associated with asthma severity (Lutfi, 2012). However, it is unclear
61	whether the same is true in youth with asthma, since some studies have reported a similar
62	parasympathetic predominance (Emin et al., 2012; Fujii et al., 2000; Gomes et al., 2013;
63	Ostrowska-Nawarycz, Wroński, Błaszczyk, Buczyłko, & Nawarycz, 2006), yet others have
64	found no influence of asthma on parasympathetic modulation (Rezvan, Dabidi Roshan, &
65	Mahmudi, 2015). These discrepancies might be related to methodological limitations which
66	preclude inter-study comparisons, such as the inclusion of obese participants (Rezvan, et al.,
67	2015), a wide range of participant ages with no consideration of maturational status (Emin,
68	et al., 2012; Fujii, et al., 2000; Ostrowska-Nawarycz, et al., 2006) or the use of absolute
69	exercise intensities (Astrup et al., 2007; Galinier et al., 2000; Ostrowska-Nawarycz, et al.,
70	2006). Indeed, a reliance on absolute work rates fails to account for inter-participant

71 differences in relative exercise intensities and the metabolic cost they engender. Since 72 characteristic changes in HRV with increasing exercise effort are well documented (Lewis, 73 Kingsley, Short, & Simpson, 2007), a reliance on absolute work rates might confound any 74 interpretation of differences in HRV during or following exercise. It should also be noted that Rezvan et al. (2015) quantified HRV during exercise, whilst most others examined it 75 76 pre to post. Indeed, given that autonomic dysfunction has been suggested to be associated 77 with the pathologic response to exercise in those with asthma (Lewis, Short, & Lewis, 78 2006), further research comparing HRV during exercise in youth with asthma and their 79 healthy counterparts is warranted.

Exercise is associated with improved lung function and mental health in those with asthma (Avallone & McLeish, 2013), as well as helping to prevent, or at least reduce the symptoms of asthma (Andrade, Britto, Lucena-Silva, Gomes, & Figueroa, 2014; Westergren et al., 2016). Furthermore, adults with asthma who engage in higher levels of moderate-tovigorous physical activity demonstrate a similar HRV to healthy adults (Yueh- Shia, Fu-Chih, Su- Ru, & Chii, 2011). However, no studies have investigated whether HRV dysfunction is influenced by exercise in adolescents with asthma.

A commonly cited barrier for those with asthma is a fear of exercise-induced bronchoconstriction (Carson, et al., 2013), which is more likely to occur during continuous aerobic exercise (Sidiropoulou, Fotiadou, Tsimaras, Zakas, & Angelopoulou, 2007). In contrast, intermittent exercise places a lower burden on the respiratory system (Beauchamp et al., 2010), suggesting that high-intensity interval training (HIIT) has potential as an exercise modality to help manage asthma. Indeed, in adults, HIIT has been more effective than aerobic endurance training at eliciting improvements in HRV (Heydari, Boutcher, &

94 Boutcher, 2013; Kiviniemi et al., 2014) although similar studies have produced no effects in 95 children, despite significant increases in aerobic fitness (Mandigout et al. (2002) and 96 Gamelin et al. (2009)). The reasons for these discrepancies are unclear and further research 97 is required to elucidate whether they reflect physiological or methodological differences. 98 Therefore, the purpose of this study was to investigate the influences of asthma and 99 HIIT, and their interaction, on the HRV of adolescents during exercise of a relative exercise 100 intensity. It was hypothesised that individuals with asthma would have a parasympathetic 101 predominance and reduced total HRV during light- and heavy-intensity exercise, and that 102 these differences would be ameliorated by the intervention.

103 Methods

104 Participants

105 Sixty-four adolescents (35 boys, 29 girls; 13.7 ± 0.9 years) were selected using stratified 106 randomisation from 618 participants who were involved in a larger randomised control trial 107 (The X4A trial: eXercise for Asthma with Commando Joe's). This sample included thirty-108 one adolescents with asthma (13.7±0.9 years; 19 boys, 12 girls) and thirty-three healthy 109 adolescents (13.8±0.9 years; 16 boys, 17 girls). Asthma severity was assessed using the 110 Global Initiative for Asthma guidelines (Global Initiative for Asthma, 2017) according to 111 the medication step required to achieve asthma control and classified the current participants 112 as having mild (n = 27) or severe (n = 4) as thma. Participants were excluded if they did not 113 have stable asthma. Ethical approval was granted by the institutional research ethics 114 committee (ref: 140515 and PG/2014/29). Parent/guardian consent and child assent were 115 obtained prior to participation.

116 Intervention

117 The intervention design was based on formative work (Winn et al., 2017). Participants 118 within the intervention group were required to attend a six-month HIIT intervention, three 119 days per week, in accord with recommendations from recent systematic reviews (Eddolls, 120 McNarry, Stratton, Winn, & Mackintosh, 2017). The 30-minute sessions consisted of 121 circuits and games-based activities designed to elicit a heart rate of >90% Heart Rate 122 maximum (HR_{max}) derived during the incremental ramp exercise test, with a 1:1 exercise to 123 rest ratio. The duration of the bouts was progressively increased from 10 s bouts initially to 124 30 s bouts. Throughout each session, participants' HRs were continuously monitored 125 (Activio AB, Stockholm, SWE). Maximal HR was predicted using the formula of Tanaka et 126 al. (2001), validated for use in adolescents (Mahon, Marjerrison, Lee, Woodruff, & Hanna, 2010). The intervention was delivered by a trained professional from Commando Joe's® 127 128 (Manchester, UK). Participants in the control group engaged in their usual day-to-day 129 activities.

130 **Procedures**

Participants were assessed at three time-points: baseline, mid-intervention (3-months), and post-intervention. Each participant was asked to attend the laboratory at the same time during the school day (\pm 2 hrs), in a hydrated state and at least two hours postprandial. All the exercise tests were performed on an electromagnetically braked cycle ergometer (Ergoselect 200, Ergoline GmbH, Lindenstrasse, Germany) and participants were asked to maintain a cadence of 75 \pm 5 revolutions per minute.

137 Anthropometrics

- 138 Stature and sitting stature were measured to the nearest 0.1 cm (Seca213, Hamburg,
- 139 Germany) and body mass to the nearest 0.1 kg (Seca876, Hamburg, Germany). Lower limb
- 140 length was calculated as the difference between stature and sitting stature and subsequently
- 141 used to estimate maturity offset using the equations of Mirwald et al. (2002).

142 Spirometry

- 143 Forced Expiratory Volume in 1 second (FEV1) was measured using a portable dry spirometer
- 144 (Vitalograph, Buckingham, UK). The best of three measurements was taken according to the
- 145 American Thoracic Society Guidelines (2005) and expressed as a percentage of the age-sex-
- 146 stature predicted value (Rosenthal et al., 1993).

147 Physical Activity

- 148 Participants wore the ActiGraph GT3X+ accelerometer (ActiGraph, Pensacola, Florida,
- 149 USA), set at 100 Hz, on their right mid-axillary line at the level of the iliac crest for seven
- 150 consecutive full days, only removing it if they undertook contact or water-based activities.
- 151 Wear-time diaries were used to log the reasons and duration of accelerometer removal.

152 Incremental Test

- 153 On the first visit, participants performed an incremental ramp test to volitional exhaustion to
- 154 determine peak oxygen uptake (VO₂) and the Gas Exchange Threshold (GET). The ramp
- 155 protocol consisted of 3 minutes of unloaded pedalling (0 W) followed immediately by an
- 156 increase in work rate at 12-24 W \cdot min⁻¹. Peak $\dot{V}O_2$ was taken as the highest 10-second

157 average. The GET was determined using the V-slope method (Beaver, Wasserman, &

- 158 Whipp, 1986) and the work rate that would elicit 40% of the difference between GET and
- 159 peak $\dot{V}O_2$ ($\Delta 40\%$) subsequently determined, accounting for the mean response time for $\dot{V}O_2$
- 160 during ramp incremental exercise (Whipp, Davis, Torres, & Wasserman, 1981).

161 Square Wave Exercise Tests

Participants repeated three square-wave exercise transitions on separate days, at least 24
hours apart, which comprised of six minutes baseline pedalling against no external work
rate (0W, "light intensity exercise") followed by an abrupt transition to the target work rate
(Δ40%) which was maintained for six minutes.

166 Measurements

Pulmonary ventilation (VE) and gas exchange ($\dot{V}O_2$ and $\dot{V}CO_2$) were measured on a breath-167 168 by-breath basis (Jaeger Oxycon Mobile, Jaeger, Germany) using a facemask with low dead-169 space connected via an impeller turbine assembly (Jaeger Triple V, Germany). Gas 170 analysers were calibrated prior to each test with gases of known concentrations and the 171 turbine volume transducer was calibrated using a built-in function calibrated using a 31 172 syringe (Hans Rudolph, Kansas City, MO). The volume and concentration signals were 173 time-aligned by accounting for the delay in capillary gas transit and analyser rise time 174 relative to the volume signal. 175 Beat-to-beat RR intervals were recorded continuously by a six-lead 176 electrocardiogram (Physio Flow PF-05 Lab1, Manatec Biomedical, France) at a sampling 177 frequency of 250 Hz. The electrodes were positioned on the forehead, neck, xiphoid process

- and on the left-hand side of the lower ribs, avoiding the abdominal muscles (Welsman,
- 179 Bywater, Farr, Welford, & Armstrong, 2005).

180 Data analysis

181 Using a MATLAB-based package developed by Physioflow, the ECG recording from each 182 constant-work-rate test was extracted and reformatted before using an independent software 183 package (Kubios HRV 2.1, Biomedical Signal Analysis Group, Finland) to detect R-wave 184 peaks from the ECG signal. The new signal was manually inspected for signal degradation 185 and physiological artefacts. This was then verified by automatic processes in the software, 186 ensuring removal of irregularly occurring large artefacts from the RR data prior to further 187 analysis without significantly affecting the spectral components of interest (<1% of RR 188 intervals were recorded as artefacts via both inspection techniques).

189 HRV variables from the final three minutes of each stage of exercise (unloaded and 190 $\Delta 40\%$) were quantified in the time domain (RMSSDRR: square root of the mean of the sum 191 of the squares of differences between adjacent RR intervals; SDNN: standard deviation of 192 all 'normal' RR intervals) according to the Task Force guidelines (1996). Frequency domain 193 and non-linear measures of HRV were also derived. Specifically, prior to spectral analysis, 194 RR interval data were re-sampled at 3.0 Hz to account for the mean HR_{max} during exercise 195 (168 beats \cdot min⁻¹ being equivalent to 2.80 Hz) and to remove non-uniformly spaced RR 196 intervals. Using Welch's periodogram method, re-sampled data were then linearly de-197 trended and segmented into consecutive 90 s Hamming windows, with a 50% overlap. This 198 was designed to reduce spectral leakage before power spectral density was estimated using a 199 fast Fourier transform algorithm. Data are presented as low-frequency (LF) power (0.04-

200 0.15 Hz), high-frequency (HF) power (0.15-0.40 Hz) and total power (TP; 0.01-0.40 Hz). In 201 addition, extended frequency bandwidths are presented for HF and TP (HF_{Bf} and TP_{Bf}, 202 respectively) to account for the high breathing frequencies (Bf) and RSA influences during 203 exercise (Lewis *et al.*, 2007), with the upper limit relative to individual tests (Bf Hz = peak 204 breathing frequency/60). LF, HF, and HF_{Bf} were also presented in normalised units (nu) and 205 as ratios (LF/HF; LF/HF $_{Bf}$), acting as indicators of sympathovagal balance. Non-linear 206 measures included the standard deviations of the Poincaré plot (SD1 and SD2). 207 Additionally, sample entropy (SampEn), the quantified rate of entropy within the RR data 208 sample, was derived, providing measures of signal complexity. The embedding dimension 209 for this was set at m = 2 and tolerance at r = 0.25 * SDNN. Finally, de-trended fluctuation 210 analysis was used to estimate the self-similarity within short- (DFA α 1 = 4-16 beats) and 211 long-term (DFA α 2 = 16-64 beats) HRV indices. 212 The physical activity data was analysed using KineSoft (version 3.3.67; KineSoft, 213 Saskatchewan, Canada) employing 1 second epochs with sustained periods of at least 20

214 minutes at zero counts considered non-wear-time (Catellier et al., 2005). A minimum daily

215 wear-time of 10 hours for 3 days, including 1 weekend day, was selected in order to provide

a more accurate overview of participants' habitual physical activity levels (Rich et al.,

217 2013). Evenson, Catellier, Gill, Ondrak and McMurray (2008). Cut-points, shown to be

218 valid and reliable determinants of physical activity intensity in children and adolescents

219 (Trost, Loprinzi, Moore, & Pfeiffer, 2011), were used to calculate time spent in the different

220 intensities.

221 Statistical Analysis

222 Individuals with moderate, mild and severe asthma were grouped since no differences 223 between study variables were observed according to asthma severity. On completion of the 224 intervention, 13 participants (7 healthy, 6 asthma (4 mild, 2 severe); 7 boys, 6 girls) met the 225 criteria of providing a complete dataset and having attended at least two sessions per week 226 for at least 70% of the intervention. This sub-sample of participants were therefore used for 227 the analysis of the effect of the HIIT intervention, along with 13 age- and sex-matched 228 controls (Table 2). This sub-sample did not differ anthropometrically from the wider study 229 population. 230 Normal distribution was confirmed using the Shapiro-Wilks test. Subsequently, a 231 mixed repeated measures ANOVA was conducted to investigate the influence of the 232 intervention and its interaction with asthma status, with post-hoc analyses using a 233 Bonferroni correction to identify the specific location of significant differences when a main 234 effect was observed. All statistical analyses were computed using SPSS Statistics 22 (IBM 235 Corp, Armonk, NY, 2013). Data are expressed as mean \pm SD and statistical significance 236 was accepted at $P \leq 0.05$.

237 **Results**

238 Baseline comparisons showed no significant anthropometric differences according to

asthma status (Table 1) or between intervention and control groups (Table 2). Over the

- training period, both intervention and control groups demonstrated a (non-significant)
- 241 upward trend for \dot{VO}_{2peak} from baseline to three-months, with the intervention group
- 242 demonstrating a further significant increase at six-months $(2.02 \pm 0.11 \text{ to } 2.36 \pm 0.14 \text{ l} \cdot \text{min}^{-1}$

¹, P < 0.001). Participants with asthma did not differ from their healthy peers in terms of the HRV parameters at baseline or in response to the intervention. Subsequently, the asthma and non-asthma groups were combined for further analyses to increase the statistical power for assessing the effect of intervention.

247 Influence of HIIT on HRV during light-intensity exercise

248 During light-intensity unloaded exercise, SDNN differed across time-points, with a

significant time by group interaction. Specifically, as presented in Table 3, in contrast to a

- significant reduction in SDNN from baseline to three-months in the control group, SDNN
- increased in the intervention group over the same period. A similar significant time by

252 group interaction was found for SD2, which increased at all time-points relative to baseline

in the intervention group (Table 3), but decreased from baseline to three-months in the

control group before returning to baseline at six-months.

255 Influence of HIIT on HRV during heavy-intensity exercise

256 As shown in Table 4, the six-month HIIT intervention was associated with significant 257 alterations in frequency-domain HRV during the constant work rate, heavy-intensity exercise, although no significant differences were observed in the time-domain or non-linear 258 259 measures. Specifically, in the intervention group, total power, low-frequency, normalised 260 LF and indices of sympathovagal balance demonstrated a significant time by group 261 interaction, increasing at three-months in the intervention group before declining towards 262 baseline values at six-months (Table 3). In the control group, indices of sympathovagal 263 balance were significantly reduced at three- and six-months compared to baseline. In

264 contrast, HFnu was significantly reduced at three- and six-months in the intervention group265 compared to an increased HFnu at six-months in the control group.

266

267 **Discussion**

268 The present study is the first to investigate the influence of asthma on ANS function in 269 adolescents during relative intensity exercise and whether a six-month HIIT programme can 270 ameliorate the any deleterious influences of asthma on HRV. In contrast to our hypotheses, 271 HRV was not influenced by asthma and, despite eliciting a significant increase in aerobic 272 fitness, the intervention did not influence HRV. Specifically, whilst no significant changes 273 were observed in time-domain or non-linear measures, the spectral parameters, which are 274 related, at least in part, to sympathetic activity, demonstrated significant training effects. 275 In adults with asthma, airway inflammation due to hyper-reactivity in response to 276 certain stimuli, such as exercise, is associated, amongst other factors, with an abnormal 277 ANS control, manifest in the form of a parasympathetic predominance associated with a 278 reduced global HRV and tonic bronchoconstriction (Lewis, et al., 2006). However, while 279 some authors report similar derangements in children with asthma at rest (Emin, et al., 280 2012; Ostrowska-Nawarycz, et al., 2006), differences during exercise are presently 281 equivocal, with a lack of research in this area despite the potential importance and 282 prognostic value of identifying altered ANS control in children and adolescents with 283 asthma. In contrast to the present study, two studies have reported similar findings to those 284 in adults (Fujii, et al., 2000; Gomes, et al., 2013), suggesting an increased vagal tone 285 following exercise. However, in agreement with the current results, the most recent study by 286 Rezvan and colleagues (2015) found no difference in parasympathetic tone between those 287 with and without asthma during and following incremental exercise, although they did 288 report a greater reduction in HRV immediately post-exercise in those with asthma. The 289 interpretation of these findings is limited by a number of methodological factors, such as the 290 added confounder of using obese children in the study by Rezvan et al. (2015), which could 291 be associated with independent influences on the ANS (Thayer, Yamamoto, & Brosschot, 292 2010), as well as a failure to use relative exercise intensities (Fujii, et al., 2000; Gomes, et 293 al., 2013; Rezvan, et al., 2015). The use of relative exercise intensities is important because 294 increasing intensity is associated with a global reduction in HRV (Lewis, et al., 2007; Perini 295 & Veicsteinas, 2003). Therefore, if participants are not exercising at the same relative 296 intensity, erroneous conclusions could be drawn regarding HRV which are really a 297 reflection of different exercise intensities, rather than disease related differences per se 298 between groups. Finally, all of the studies to date have involved pre-pubertal children and 299 therefore do not account for the potential influence of maturity on the ANS (Lenard, 300 Studinger, Mersich, Kocsis, & Kollai, 2004), or for the interaction between these maturity-301 related adaptations and the influence of asthma. These methodological differences might 302 explain the discrepant findings of the present study, which suggest that asthma does not 303 influence ANS control during exercise in pubertal adolescents, although it is pertinent that 304 previous studies have largely focussed on HRV following exercise (Emin, et al., 2012; Fujii, 305 et al., 2000; Gomes, et al., 2013; Ostrowska-Nawarycz, et al., 2006), thereby limiting 306 comparisons. It is also important to consider the potential interaction between disease 307 severity and ANS control as the relatively mild, self-reported, severity of asthma in the 308 present participants may have been insufficient to elicit significant derangements in ANS

309 control. Alternatively, the discrepancy between the present results and those in adults may 310 reflect disease progression or longevity such that alterations in ANS control are not manifest 311 until adulthood. Indeed, the degree of autonomic derangement is related to asthma duration 312 (Lutfi, 2012), as well as medication use. Specifically, participants with asthma severe 313 enough to potentially induce significant changes in sympathovagal tone were likely to be 314 medicated with either, or both, short- and long-acting beta-agonists. Lewis et al. (2006) 315 found that these medications increased cardiac sympathetic excitement and could explain 316 the appearance of similar sympathovagal balances between those with and without asthma. 317 Whilst others have reported significant differences in HRV between those with controlled 318 and uncontrolled asthma (Lutfi, 2015) further work is warranted to elucidate the relationship 319 between disease severity, duration, maturation and asthma with regards to the ANS response 320 during exercise.

321 The present findings are partly in accord with the increased LF and TP reported 322 following HIIT in adults (Heydari, et al., 2013; Kiviniemi, et al., 2014; Rakobowchuk, 323 Harris, Taylor, Cubbon, & Birch, 2013), although the unchanged HF and decreased HFnu 324 contradicts these studies. Furthermore, the current results disagree with the results from 325 other HIIT studies in children (Gamelin, et al., 2009; Mandigout, et al., 2002). Specifically, 326 Mandigout, et al. (2002) and Gamelin et al. (2009) reported no alterations in autonomic 327 balance, despite significant increases in aerobic fitness. The discrepancies between these 328 findings may be attributable to methodological differences, such as the duration and 329 conditions under which the HRV measures were derived, the types of exercises and the 330 duration of the intervention. Indeed, considerable differences in the duration of recordings 331 used to obtain HRV, which ranged from three-minutes in the present study to 24-hours in

332 Mandigout et al. (2002), limit inter-study comparisons, particularly given that it is widely 333 accepted to be inappropriate to compare parameters derived from different recording lengths 334 (TaskForce, 1996). It is also important to highlight that whilst Mandigout et al. (2002) used 335 high-intensity, intermittent exercises, they did not specifically implement a HIIT 336 intervention. The longer duration of the intervention in the present study may also explain 337 the significant influences on autonomic balance that were not reported in earlier studies in 338 children (Gamelin, et al., 2009; Mandigout, et al., 2002). However, significant HRV 339 adaptations can occur after two to three weeks in adults (Kiviniemi, et al., 2014; Seals & 340 Chase, 1989). Whilst the applicability of these findings to younger populations remains to 341 be elucidated, these findings suggest that training adaptations could have occurred in shorter 342 interventions. Interestingly, the present results indicate that the influence of HIIT on ANS is 343 age- or maturity-, dependant, with pre-pubertal children less sensitive and less able to 344 induce training-related adaptations (Gilliam & Freedson, 1980; Katch, 1983). This potential 345 influence of maturity may be particularly important when trying to understand the complex 346 interactions underpinning HRV responses, which are simultaneously influenced by both 347 neural and humoral effects (Binah, Weissman, Itskovitz-Eldor, & Rosen, 2013). 348 Interestingly, the present study found an increase in LF power with HIIT which 349 persisted even when changes in TP were accounted for, reflecting an increased 350 sympathovagal balance. Although the physiological underpinnings of the LF band still 351 remain to be conclusively determined (TaskForce, 1996), these changes suggest an 352 increased sympathetic tone following three-months of training which decreased towards 353 baseline values at six-months. These deleterious initial changes in autonomic balance may 354 be attributable to training load, which plays a key factor in eliciting autonomic adaptations

355	(Gamelin, et al., 2009); a high training load, to which participants are not accustomed, can
356	lead to overload and an accumulation of fatigue. Such fatigue is associated with a shift in
357	autonomic balance from parasympathetic- to sympathetic-predominance (Mourot et al.,
358	2004; Schmitt, Regnard, & Millet, 2015). Therefore, the change in LF and associated
359	variables indicates an initial fatigue-related autonomic shift towards sympathetic
360	predominance which, following adaptation to the exercise load, returned to baseline values
361	at six-months. Indeed, it could be postulated that a longer study duration may have
362	demonstrated a continued decline in LF, resulting in a parasympathetic predominance as
363	typically reported following HIIT in adults (Heydari, et al., 2013; Kiviniemi, et al., 2014)
364	and aerobic training in adolescents (Hedelin, Wiklund, Bjerle, & Henriksson- Larsén,
365	2000).
366	The degree of training-induced adaptation may be related to an individual's genetics,
367	and the pre-training level of HRV (Buchheit & Gindre, 2006). Specifically, pre-training
368	SDNN is strongly correlated to the magnitude of improvements observed post-training and
369	it has been recommended that training is conducted more frequently, or at a higher intensity,
370	
	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present
371	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present results may reflect a high baseline HRV, thereby limiting the potential for improvements to
371 372	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present results may reflect a high baseline HRV, thereby limiting the potential for improvements to be elicited through HIIT. The lack of comparable data regarding HRV during exercise limits
371372373	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present results may reflect a high baseline HRV, thereby limiting the potential for improvements to be elicited through HIIT. The lack of comparable data regarding HRV during exercise limits further conclusions, but the relatively poor aerobic fitness at baseline in the current
371372373374	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present results may reflect a high baseline HRV, thereby limiting the potential for improvements to be elicited through HIIT. The lack of comparable data regarding HRV during exercise limits further conclusions, but the relatively poor aerobic fitness at baseline in the current participants, the significant influence of the intervention on their aerobic fitness, and the
 371 372 373 374 375 	in those with poorer pre-training HRV (Buchheit & Gindre, 2006). Therefore, the present results may reflect a high baseline HRV, thereby limiting the potential for improvements to be elicited through HIIT. The lack of comparable data regarding HRV during exercise limits further conclusions, but the relatively poor aerobic fitness at baseline in the current participants, the significant influence of the intervention on their aerobic fitness, and the strong correlation reported between HRV and $\dot{V}O_{2peak}$ are relevant to note (Buchheit &

376 Gindre, 2006).

17

377 The decreased normalised HF (HFnu) observed in the present study at three- and 378 six-months may be a reflection of an increased LF component, decreasing the relative 379 contribution of HF when corrected for changes in TP, or a decrease in parasympathetic 380 activity. Although elevated parasympathetic activity is more commonly reported following 381 training (Buchheit & Gindre, 2006; Carter, Banister, & Blaber, 2003), studies employing 382 pharmacological autonomic blockades to assess the heart rate and HRV response have 383 reported decreased vagal activity in trained athletes (Furlan et al., 1993; Katona, McLean, 384 Dighton, & Guz, 1982). Nevertheless, the application of such studies to paediatric 385 populations is highly speculative and caution is required when drawing any further 386 conclusions.

387 The present study utilised multiple repeat constant-work-rate transitions at each 388 time-point, thus improving reliability of measures and counteracting the day-to-day 389 variation in HRV (Schroeder et al., 2004). Nonetheless, there are a number of limitations 390 that should be noted. Firstly, the length of the ECG recordings was shorter than 391 recommended (TaskForce, 1996) and the sample size may be deemed low, although it is 392 still comparable to previous studies which found significant effects of HIIT on ANS 393 function (Rakobowchuk, et al., 2013). It would also have aided in the interpretation of the 394 present findings to have resting values for HRV. Furthermore, asthma was self-reported and 395 relied on the participants' understanding of disease severity, which was potentially mild and 396 may not have been severe enough to elicit autonomic alterations (Emin, et al., 2012). The 397 study may also have been subject to selection bias as those with more severe asthma may 398 have chosen not to participate due to fear of exacerbation or exercise induced asthma.

18

399	In conclusion, the present findings suggest that HRV is not deleteriously influenced
400	by asthma in adolescents during relative intensity exercise and only spectral power was
401	influenced by the six-month HIIT intervention, despite eliciting significant increases in
402	aerobic fitness. Whilst highlighting the dissociation between aerobic fitness and HRV, these
403	results indicate that HIIT may be associated with short-term, deleterious shifts in autonomic
404	balance towards greater sympathetic predominance during exercise due to physiological
405	overload and fatigue, which are ameliorated within six-months.
406	

407 Acknowledgements

408 This work was funded by the Asthma UK Centre for Applied Research [AUK-AC-2012-01] 409 and Swansea University Medical School. Commando Joe's® supported the work and assisted 410 in funding co-author WTBE.

411

References

413	Andrade, L. B., Britto, M. C. A., Lucena-Silva, N., Gomes, R. G., & Figueroa, J. N. (2014). The efficacy
414	of aerobic training in improving the inflammatory component of asthmatic children
415	Randomized trial. Respiratory Medicine, 108(10), pp. 1438-1445
416	doi: <u>https://doi.org/10.1016/j.rmed.2014.07.009</u> Retrieved from
417	http://www.sciencedirect.com/science/article/pii/S0954611114002479
418	Astrup, A. S., Nielsen, F. S., Rossing, P., Ali, S., Kastrup, J., Smidt, U. M., & Parving, HH. (2007)
419	Predictors of mortality in patients with type 2 diabetes with or without diabetic
420	nephropathy: a follow-up study. Journal of Hypertension, 25(12), pp. 2479-2485
421	doi:10.1097/HJH.0b013e3282f06428 Retrieved from
422	https://journals.lww.com/jhypertension/Fulltext/2007/12000/Predictors of mortality in
423	patients_with_type_2.17.aspx
424	Avallone, K. M., & McLeish, A. C. (2013). Asthma and Aerobic Exercise: A Review of the Empirica
425	Literature. Journal of Asthma, 50(2), pp. 109-116. doi:10.3109/02770903.2012.759963
426	Retrieved from <u>https://doi.org/10.3109/02770903.2012.759963</u>
427	Beauchamp, M. K., Nonoyama, M., Goldstein, R. S., Hill, K., Dolmage, T. E., Mathur, S., & Brooks, D
428	(2010). Interval versus continuous training in individuals with chronic obstructive pulmonary
429	diseasea systematic review. Thorax, 65(2), pp. 157-164. doi:10.1136/thx.2009.123000
430	Retrieved from http://dx.doi.org/10.1136/thx.2009.123000
431	Beaver, W. L., Wasserman, K., & Whipp, B. J. (1986). A new method for detecting anaerobic threshold
432	by gas exchange. Journal of Applied Physiology, 60(6), pp. 2020-2027. Retrieved from
433	http://dx.doi.org/
434	Binah, O., Weissman, A., Itskovitz-Eldor, J., & Rosen, M. R. (2013). Integrating beat rate variability
435	From single cells to hearts. <i>Heart Rhythm, 10</i> (6), pp. 928-932
436	doi:10.1016/j.hrthm.2013.02.013 Retrieved from
437	http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3923529/
438	Buchheit, M., & Gindre, C. (2006). Cardiac parasympathetic regulation: respective associations with
439	cardiorespiratory fitness and training load. American Journal of Physiology - Heart and
440	Circulatory Physiology, 291(1), pp. H451-H458. doi:10.1152/ajpheart.00008.2006 Retrieved
441	from https://www.physiology.org/doi/abs/10.1152/ajpheart.00008.2006
442	Carson, K. V., Chandratilleke, M. G., Picot, J., Brinn, M. P., Esterman, A. J., & Smith, B. J. (2013)
443	Physical training for asthma. Cochrane Database Sys
444	<i>Rev</i> (9)doi:10.1002/14651858.CD001116.pub4 Retrieved from
445	http://dx.doi.org/10.1002/14651858.CD001116.pub4
446	Carter, J., Banister, E., & Blaber, A. (2003). Effect of Endurance Exercise on Autonomic Control o
447	Heart Rate. Sports Medicine, 33(1), pp. 33-46. Retrieved from
448	http://ejournals.ebsco.com/direct.asp?ArticleID=4C028F658A70E3A4D802
449	Chang, ET., Silberstein, D., Rambod, M., Porszasz, J., & Casaburi, R. (2011). Heart rate variability
450	during constant work rate exercise at and above the critical power in patients with severe
451	chronic obstructive pulmonary disease. Tzu Chi Medical Journal, 23(2), pp. 42-45
452	doi: <u>https://doi.org/10.1016/j.tcmj.2011.05.001</u> Retrieved from
453	http://www.sciencedirect.com/science/article/pii/S1016319011000450
454	Eddolls, W. T. B., McNarry, M. A., Stratton, G., Winn, C. O. N., & Mackintosh, K. A. (2017). High
455	Intensity Interval Training Interventions in Children and Adolescents: A Systematic Review
456	Sports Medicine, 47(11), pp. 2363-2374. doi:10.1007/s40279-017-0753-8 Retrieved from
457	http://dx.doi.org/10.1007/s40279-017-0753-8

458 Emin, O., Esra, G., Aysegül, D., Ufuk, E., Ayhan, S., & Rusen, D. M. (2012). Autonomic nervous system 459 dysfunction and their relationship with disease severity in children with atopic asthma. 460 Respiratory Physiology & Neurobiology, 183doi:10.1016/j.resp.2012.07.002 Retrieved from 461 https://doi.org/10.1016/j.resp.2012.07.002 462 Fujii, H., Fukutomi, O., Inoue, R., Shinoda, S., Okammoto, H., Teramoto, T., . . . Seishima, M. (2000). 463 Autonomic regulation after exercise evidenced by spectral analysis of heart rate variability 464 in asthmatic children. Annals of Allergy, Asthma, and Immunology, 85(3), pp. 233-237. 465 Retrieved from <Go to ISI>://WOS:000089561300013 466 Furlan, R., Piazza, S., Dell'Orto, S., Gentile, E., Cerutti, S., Pagani, M., & Malliani, A. (1993). Early and 467 late effects of exercise and athletic training on neural mechanisms controlling heart rate. 468 Cardiovascular Research, 27(3), pp. 482 - 488. 469 Galinier, M., Pathak, A., Fourcade, J., Androdias, C., Curnier, D., Varnous, S., . . . Bounhoure, J. P. 470 (2000). Depressed low frequency power of heart rate variability as an independent predictor 471 of sudden death in chronic heart failure. European Heart Journal, 21(6), pp. 475-482. 472 doi:10.1053/euhj.1999.1875 Retrieved from http://dx.doi.org/10.1053/euhj.1999.1875 473 Gamelin, F., Baquet, G., Berthoin, S., Thevenet, D., Nourry, C., Nottin, S., & Bosquet, L. (2009). Effect 474 of high intensity intermittent training on heart rate variability in prepubescent children. 475 [journal article]. European Journal of Applied Physiology, 105(5), pp. 731-738. 476 doi:10.1007/s00421-008-0955-8 Retrieved from https://doi.org/10.1007/s00421-008-0955-477 8 478 Gilliam, T. B., & Freedson, P. S. (1980). Effects of a 12-week school physical fitness programme on 479 peak VO₂, body composition and blood lipids in 7 to 9 year old children. International Journal 480 of Sports Medicine, 1, pp. 73-78. 481 Global Initiative for Asthma. (2017). Global strategy for asthma management and prevention. 482 Available from: www.ginasthma.org 483 Gomes, E. L., Sampaio, L. M., Costa, I. P., Dias, F. D., Ferneda, V. S., Silva, G. A., & Costa, D. (2013). 484 Analysis of autonomic modulation during maximal and submaximal work rate and functional 485 capacity in asthmatic children. Journal of Asthma, 50doi:10.3109/02770903.2013.793707 486 Retrieved from https://doi.org/10.3109/02770903.2013.793707 487 Hedelin, R., Wiklund, U., Bjerle, P., & Henriksson-Larsén, K. (2000). Pre- and post-season heart rate 488 variability in adolescent cross-country skiers. Scandinavian Journal of Medicine & Science in 489 Sports, 10(5), pp. 298-303. doi:doi:10.1034/j.1600-0838.2000.010005298.x Retrieved from 490 https://onlinelibrary.wiley.com/doi/abs/10.1034/j.1600-0838.2000.010005298.x 491 Heydari, M., Boutcher, Y. N., & Boutcher, S. H. (2013). High-intensity intermittent exercise and 492 cardiovascular and autonomic function. Clinical Autonomic Research, 23(1), pp. 57-65. 493 doi:10.1007/s10286-012-0179-1 Retrieved from https://doi.org/10.1007/s10286-012-0179-494 1 495 Katch, V. L. (1983). Physical conditioning of children. [Article]. J Adols Health, 3(4), pp. 241-246. 496 Retrieved from <Go to ISI>://A1983QE26900004 497 Katona, P. G., McLean, M., Dighton, D. H., & Guz, A. (1982). Sympathetic and parasympathetic cardiac 498 control in athletes and nonathletes at rest. Journal of Applied Physiology, 52(6), pp. 1652-499 1657. doi:10.1152/jappl.1982.52.6.1652 Retrieved from 500 https://www.physiology.org/doi/abs/10.1152/jappl.1982.52.6.1652 501 Kiviniemi, A. M., Tulppo, M. P., Eskelinen, J. J., Savolainen, A. M., Kapanen, J., Heinonen, I. H. A., . . . 502 Kalliokoski, K. K. (2014). Cardiac Autonomic Function and High-Intensity Interval Training in 503 Middle-Age Men. Medicine and Science in Sport and Exercise, 46(10), pp. 1960-1967.

504	doi:10.1249/mss.000000000000000000000000 Retrieved from <u>https://journals.lww.com/acsm-</u>
505	msse/Fulltext/2014/10000/Cardiac Autonomic Function and High Intensity.11.aspx
506	Lenard, Z., Studinger, P., Mersich, B., Kocsis, L., & Kollai, M. (2004). Maturation of Cardiovagal
507	Autonomic Function From Childhood to Young Adult Age. <i>Circulation, 110</i> (16), pp. 2307-
508	2312. doi:10.1161/01.cir.0000145157.07881.a3 Retrieved from
509	http://circ.ahajournals.org/content/circulationaha/110/16/2307.full.pdf
510	Lewis, M. J., Kingsley, M., Short, A. L., & Simpson, K. (2007). Rate of reduction of heart rate variability
511	during exercise as an index of physical work capacity. Scandinavian Journal of Medicine &
512	Science in Sports, 17(6), pp. 696-702. doi:doi:10.1111/j.1600-0838.2006.00616.x Retrieved
513	from https://onlinelibrary.wiley.com/doi/abs/10.1111/j.1600-0838.2006.00616.x
514	Lewis, M. J., Short, A. L., & Lewis, K. E. (2006). Autonomic nervous system control of the
515	cardiovascular and respiratory systems in asthma. Respiratory Medicine, 100(10), pp. 1688-
516	1705. doi:10.1016/j.med.2006.01.019 Retrieved from <go isi="" to="">://WOS:000241040000003</go>
517	Lutfi, M. F. (2012). Autonomic modulations in patients with bronchial asthma based on short-term
518	heart rate variability. Lung India, 29doi:10.4103/0970-2113.99111 Retrieved from
519	https://doi.org/10.4103/0970-2113.99111
520	Lutfi, M. F. (2015). Patterns of heart rate variability and cardiac autonomic modulations in controlled
521	and uncontrolled asthmatic patients. [journal article]. BMC Pulmonary Medicine, 15(1), p
522	119. doi:10.1186/s12890-015-0118-8 Retrieved from https://doi.org/10.1186/s12890-015-
523	0118-8
524	Mahon, A. D., Marierrison, A. D., Lee, J. D., Woodruff, M. E., & Hanna, L. E. (2010), Evaluating the
525	prediction of maximal heart rate in children and adolescents. Research Quarterly for Exercise
526	and Sport, 81(4), pp. 466-471, doi:10.1080/02701367.2010.10599707 Retrieved from
527	http://dx.doi.org/10.1080/02701367.2010.10599707
528	Mandigout, S., Melin, A., Fauchier, L., N'Guven, L. D., Courteix, D., & Obert, P. (2002). Physical training
529	increases heart rate variability in healthy prepubertal children. European Journal of Clinical
530	Investigation, 32(7), pp. 479-487, doi:doi:10.1046/i.1365-2362.2002.01017, x Retrieved from
531	https://onlinelibrary.wiley.com/doi/abs/10.1046/j.1365-2362.2002.01017.x
532	McNarry, M. A., & Mackintosh, K. A. (2016). Reproducibility of Heart Rate Variability Indices in
533	Children with Cystic Fibrosis. <i>PloS One</i> , 11(3), p e0151464.
534	doi:10.1371/journal.pone.0151464 Retrieved from
535	https://doi.org/10.1371/journal.pone.0151464
536	Miller, M. R., Hankinson, J., Brusasco, V., Burgos, F., Casaburi, R., Coates, A., Wanger, J. (2005).
537	Standardisation of spirometry. European Respiratory Journal, 26(2), pp. 319-338.
538	doi:10.1183/09031936.05.00034805 Retrieved from
539	http://erj.ersjournals.com/content/26/2/319.abstract
540	Mirwald, R. L., Baxter-Jones, A. D., Bailey, D. A., & Beunen, G. P. (2002). An assessment of maturity
541	from anthropometric measurements. <i>Medicine and science in sports and exercise, 34</i> (4), pp.
542	689-694. Retrieved from http://www.ncbi.nlm.nih.gov/pubmed/11932580
543	Mourot, L., Bouhaddi, M., Perrey, S., Cappelle, S., Henriet, M., Wolf, J., Regnard, J. (2004).
544	Decrease in heart rate variability with overtraining: assessment by the Poincaré plot analysis.
545	Clinical Physiology and Functional Imaging, 24(1), pp. 10-18. doi:10.1046/j.1475-
546	0961.2003.00523.x Retrieved from http://dx.doi.org/10.1046/i.1475-0961.2003.00523.x
547	Ostrowska-Nawarycz, L., Wroński, W., Błaszczyk, J., Buczyłko, K., & Nawarycz, T. (2006). The heart
548	rate variability analysis in youth and children with bronchial asthma. Polski Merkuriusz

550 Perini, R., & Veicsteinas, A. (2003). Heart rate variability and autonomic activity at rest and during 551 exercise in various physiological conditions. [journal article]. European Journal of Applied 552 *Physiology*, *90*(3), pp. 317-325. doi:10.1007/s00421-003-0953-9 Retrieved from 553 https://doi.org/10.1007/s00421-003-0953-9 554 Rakobowchuk, M., Harris, E., Taylor, A., Cubbon, R. M., & Birch, K. M. (2013). Moderate and heavy 555 metabolic stress interval training improve arterial stiffness and heart rate dynamics in 556 humans. [journal article]. European Journal of Applied Physiology, 113(4), pp. 839-849. 557 doi:10.1007/s00421-012-2486-6 Retrieved from https://doi.org/10.1007/s00421-012-2486-558 6 559 Rezvan, K., Dabidi Roshan, V., & Mahmudi, S. A. (2015). Short-term heart rate variability in asthmatic 560 obese children: effect of exhaustive exercise and different humidity conditions. Journal of Sports Medicine and Physical Fitness, 55(11), pp. 1390 - 1396. 561 562 Rosenthal, M., Bain, S. H., Cramer, D., Helms, P., Denison, D., Bush, A., & Warner, J. O. (1993). Lung 563 function in white children aged 4 to 19 years: I--Spirometry. Thorax, 48(8), pp. 794-802. 564 Schmitt, L., Regnard, J., & Millet, G. P. (2015). Monitoring Fatigue Status with HRV Measures in Elite 565 Athletes: An Avenue Beyond RMSSD? Frontiers in Physiology, 6, р 343. 566 doi:10.3389/fphys.2015.00343 from Retrieved 567 http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4652221/ 568 Schroeder, E. B., Whitsel, E. A., Evans, G. W., Prineas, R. J., Chambless, L. E., & Heiss, G. (2004). 569 Repeatability of heart rate variability measures. Journal of Electrocardiology, 37(3), pp. 163-570 172. doi:10.1016/j.jelectrocard.2004.04.004 Retrieved from 571 http://www.sciencedirect.com/science/article/pii/S0022073604000421 572 Seals, D. R., & Chase, P. B. (1989). Influence of physical training on heart rate variability and 573 baroreflex circulatory control. [Journal Article]. Journal of Applied Physiology, 66(4), pp. 574 1886-1895. Retrieved from http://jap.physiology.org/jap/66/4/1886.full.pdf 575 Sidiropoulou, M. P., Fotiadou, E. G., Tsimaras, V. K., Zakas, A. P., & Angelopoulou, N. A. (2007). The 576 effect of interval training in children with exercise-induced asthma competing in soccer. 577 Journal of Strength and Conditioning Research, 21(2), pp. 446-450. Retrieved from 578 http://onlinelibrary.wiley.com/o/cochrane/clcentral/articles/666/CN-579 00613666/frame.html 580 Tanaka, H., Monahan, K. D., & Seals, D. R. (2001). Age-predicted maximal heart rate revisited. Journal 581 of the American College of Cardiology, 37(1), pp. 153-156. Retrieved from http://dx.doi.org/ 582 TaskForce. (1996). Heart rate variability: standards of measurement, physiological interpretation, 583 and clinical Circulation, 1043-1065. Retrieved from use. 93(5), pp. 584 http://circ.ahajournals.org/content/93/5/1043.short 585 Thayer, J. F., Yamamoto, S. S., & Brosschot, J. F. (2010). The relationship of autonomic imbalance, 586 heart rate variability and cardiovascular disease risk factors. International Journal of 587 Cardiology, 141(2), pp. 122-131. doi:<u>http://dx.doi.org/10.1016/j.ijcard.2009.09.543</u> 588 Retrieved from https://www.clinicalkey.com.au/#!/content/1-s2.0-S0167527309014879 589 Wanrooij, V. H., Willeboordse, M., Dompeling, E., & van de Kant, K. D. (2014). Exercise training in 590 children with asthma: a systematic review. British Journal of Sports Medicine, 48(13), pp. 591 from 1024-1031. doi:10.1136/bjsports-2012-091347 Retrieved 592 http://bjsm.bmj.com/content/bjsports/48/13/1024.full.pdf 593 Welsman, J., Bywater, K., Farr, C., Welford, D., & Armstrong, N. (2005). Reliability of peak VO₂ and 594 maximal cardiac output assessed using thoracic bioimpedance in children. [Article]. 595 European Journal of Applied Physiology, 94(3), pp. 228-234. doi:10.1007/s00421-004-1300-596 5 Retrieved from <Go to ISI>://000229506500002

597	Westergren, T., Fegran, L., Nilsen, T., Haraldstad, K., Kittang, O. B., & Berntsen, S. (2016). Active pla
598	exercise intervention in children with asthma: a PILOT STUDY. British Medical Journal Oper
599	6(1)doi:10.1136/bmjopen-2015-009721 Retrieved from
600	http://bmjopen.bmj.com/content/bmjopen/6/1/e009721.full.pdf
601	Whipp, B. J., Davis, J. A., Torres, F., & Wasserman, K. (1981). A test to determine parameters of
602	aerobic function during exercise. Journal of Applied Physiology, 50, pp. 217-221.
603	Winn, C. O. N., Mackintosh, K. A., Eddolls, W. T. B., Stratton, G., Wilson, A. M., Rance, J. Y., Davies
604	G. A. (2017). Perceptions of asthma and exercise in adolescents with and without asthma
605	Journal of Asthma, 55(8), pp. 868-876. doi:10.1080/02770903.2017.1369992 Retrieved from
606	http://dx.doi.org/10.1080/02770903.2017.1369992
607	Yueh-Shia, T., Fu-Chih, L., Su-Ru, C., & Chii, J. (2011). The influence of physical activity level on hear
608	rate variability among asthmatic adults. Journal of Clinical Nursing, 20(1-2), pp. 111-118
609	doi:doi:10.1111/j.1365-2702.2010.03397.x Retrieved from
610	https://onlinelibrary.wiley.com/doi/abs/10.1111/j.1365-2702.2010.03397.x
611	

Table 1 Baseline participant characteristics

	Asthma	Healthy
n	31	32
Age (years)	13.5 ± 0.9	13.4 ± 0.9
Height (cm)	161.0 ± 10.0	157.9 ± 8.2
Mass (kg)	57.6 ± 14.6	50.7 ± 11.6
BMI (kg·m ⁻²)	22.0 ± 3.9	20.0 ± 3.1
$\dot{V}O_{2peak}$ (l·min ⁻¹)	2.10 ± 0.49	2.04 ± 0.50
MVPA (mins)	60.3 ± 33.5	57.0 ± 18.2

Mean \pm SD. *n*, sample size; BMI, body mass index; $\dot{V}O_{2peak}$, peak oxygen uptake; MVPA, moderate-to-vigorous physical activity

	Intervention	Control
n	13	13
Age (years)	13.3 ± 0.9	13.6 ± 1.0
Height (cm)	160.7 ± 8.8	159.5 ± 8.8
Mass (kg)	52.3 ± 11.2	53.7 ± 12.8
BMI (kg·m ⁻²)	20.1 ± 3.2	20.8 ± 3.7
$\dot{V}O_{2peak}$ (l·min ⁻¹)	2.00 ± 0.42	2.09 ± 0.47
$\dot{V}O_{2peak}$ (ml·kg ^{-0.45} ·min ⁻¹)	332 ± 51	362 ± 52
MVPA (mins)	47.9 ± 14.3	58.5 ± 24.8
$FEV_1(l \cdot min^{-1})$	2.96 ± 0.79	2.86 ± 0.82
FEV ₁ (%)	98 ± 15	91 ± 15

Table 2 Sub-sample characteristics of those who met the minimum adherence criteria

620 Mean \pm SD. *n*, sample size; BMI, body mass index; $\dot{V}O_{2peak}$, peak oxygen uptake; MVPA, 621 moderate-to-vigorous physical activity; FEV₁, forced expiratory volume in one second

Table 3 HRV	⁷ during	light-in	tensity	exercise	over six	-months

	Baseline	Three-months	Six-months	Baseline	Т
SDNN (ms)	37.3 ± 6.0	$42.0 \pm 3.2^{*}$	44.3 ± 4.7	45.6 ± 6.3	
RMSSD (ms)	32.7 ± 7.8	35.5 ± 5.4	37.7 ± 6.5	40.2 ± 8.1	
$LF (ms^2)$	564.5 ± 181.5	735.1 ± 126.7	661.3 ± 104.7	725.8 ± 188.3	6
LFnu	64.1 ± 3.1	62.5 ± 3.3	61.2 ± 3.7	60.5 ± 3.2	
HF (ms ²)	459.7 ± 201.2	457.4 ± 105.8	608.6 ± 149.3	505.6 ± 208.8	4
HFnu	35.3 ± 3.2	37.2 ± 3.3	38.5 ± 3.7	39.3 ± 3.3	
$\mathrm{HF}_{Bf}~(\mathrm{ms}^2)$	706 ± 305	707 ± 173	864 ± 234	751 ± 316	
HF _{Bf} nu	47 ± 4	47 ± 4	49 ± 4	51 ± 4	
$TP (ms^2)$	1648 ± 519	1761 ± 313	1889 ± 276	1767 ± 538	
TP_{Bf} (ms ²)	$1887{\pm}590$	2012 ± 344	2149 ± 342	2070 ± 612	
LF:HF	2.22 ± 0.35	2.23 ± 0.32	2.12 ± 0.31	2.08 ± 0.36	
LF:HF _{Bf}	1.30 ± 0.18	1.38 ± 0.22	1.32 ± 0.19	1.23 ± 0.19	
SD1 (ms)	23.4 ± 5.5	25.2 ± 3.8	27.2 ± 4.6	28.5 ± 5.7	
SD2 (ms)	46.9 ± 6.8	$53.4\pm3.9^*$	$55.7\pm4.2^*$	55.23 ± 7.1	
SampEn	1.68 ± 0.07	1.63 ± 0.07	1.68 ± 0.08	1.61 ± 0.07	
DFAa1	1.07 ± 0.06	1.11 ± 0.07	1.04 ± 0.06	1.02 ± 0.07	
DFAa2	0.84 ± 0.05	0.86 ± 0.03	0.83 ± 0.04	0.79 ± 0.05	

Mean \pm SD. SDNN, standard deviation of all the RR intervals; RMSSD, Root mean square of successive RR in low frequency power (0.04-0.15 Hz); HF, high frequency power (0.15-0.4 Hz); HFnu, high frequency power in , expanded high frequency power (0.15-*Bf* Hz); TP, total power; TP_{*Bf*}, total expanded power; LF:HF, low and hig a ratio; LF:HF_{*Bf*}, low and expanded high frequency powers as a ratio; nu, normalized power; SD1 and SD2, Star Poincaré Plot; SampEn; Sample entropy within the RR data; DFA, detrended fluctuation analysis of short-term (HRV indices. * Significant difference to baseline within group (p < 0.05)

		Intervention			
	Baseline	Three-months	Six-months	Baseline	T
SDNN (ms)	9.8 ± 0.9	9.3 ± 1.3	10.3 ± 4.3	10.4 ± 0.9	
RMSSD (ms)	10.6 ± 0.9	7.7 ± 1.3	9.7 ± 1.7	9.8 ± 1.5	
LF (ms ²)	6.0 ± 3.9	$10.3\pm1.9^*$	4.0 ± 0.6	10.9 ± 4.1	
LFnu	61.1 ± 5.2	$72.0\pm5.3^*$	68.6 ± 5.9	62.8 ± 5.4	
$HF (ms^2)$	3.8 ± 1.4	3.9 ± 1.2	2.3 ± 0.7	3.9 ± 1.4	
HFnu	37.7 ± 5.1	$29.0\pm5.1^{*}$	$31.0\pm0.7^*$	36.7 ± 5.3	
$\mathrm{HF}_{Bf}~(\mathrm{ms}^2)$	23 ± 6	21 ± 5	20 ± 7	20 ± 6	
HF _{Bf} nu	75 ± 5	79 ± 11	72 ± 4	71 ± 5	
TP (ms ²)	26 ± 16	$49\pm9^*$	17 ± 3	43 ± 16	
TP_{Bf} (ms ²)	46 ± 17	65 ± 11	35 ± 7	60 ± 18	
LF:HF	2.60 ± 0.64	$5.19\pm0.83^*$	3.80 ± 0.59	3.98 ± 0.66	/
LF:HF _{Bf}	0.45 ± 0.15	$0.75\pm0.14^{*}$	0.46 ± 0.09	0.59 ± 0.15	(
SD1 (ms)	7.5 ± 1.0	5.4 ± 1.0	6.8 ± 1.2	7.1 ± 1.1	
SD2 (ms)	11.1 ± 1.1	10.2 ± 1.0	12.0 ± 1.0	12.4 ± 1.2	
SampEn	1.50 ± 0.09	1.59 ± 0.10	1.35 ± 0.11	1.50 ± 0.09	
DFAa1	0.43 ± 0.06	0.56 ± 0.05	0.42 ± 0.04	0.45 ± 0.06	
DFAa2	1.03 ± 0.07	1.07 ± 0.08	1.04 ± 0.09	1.03 ± 0.07	

Table 4 HRV during heavy-intensity exercise over six-months

Mean \pm SD. SDNN, standard deviation of all the RR intervals; RMSSD, Root mean square of successive RR in low frequency power (0.04-0.15 Hz); HF, high frequency power (0.15-0.4 Hz); HFnu, high frequency power in , expanded high frequency power (0.15-*Bf* Hz); TP, total power; TP_{*Bf*}, total expanded power; LF:HF, low and hig a ratio; LF:HF_{*Bf*}, low and expanded high frequency powers as a ratio; nu, normalized power; SD1 and SD2, Star Poincaré Plot; SampEn; Sample entropy within the RR data; DFA, detrended fluctuation analysis of short-term (HRV indices. * Significantly different to baseline within group (p < 0.05)