Effects of sprint interval and continuous endurance training on serum levels of inflammatory biomarkers

Fariborz Hovanloo¹, Tahereh Arefirad¹,²* and Sajad Ahmadizad¹

Abstract

Chronic and inflammatory diseases are major causes of mortality. Although the anti-inflammatory effects of exercise have been confirmed, but the effect of different types of exercise on inflammatory markers is different. The aim of this study is comparing the effects of two types of sprint interval (SIT) and continuous endurance (CET) training on inflammatory markers. Sixteen students who had recreational activities participated in this study and were randomly assigned to one of the two protocols. The SIT protocol consisted of four to six 30-s “all-out” Wingate tests separated by 4 minutes of recovery and The CET protocol included 90–120 minutes of cycling at 65% Vo2max. The two protocols were performed 3 days per week and for two weeks. In each group, two blood samples were collected before and 2 days (24 and 48 hrs) after the training. Results showed that there was no significant difference between the two training protocols on all measured parameters (p>0.05). The results of present study showed that the SIT and CET have identical effects on inflammatory markers.

Keywords: Inflammatory biomarkers, Sprint interval training, Continuous endurance training

Introduction

Chronic and inflammatory diseases are major causes of mortality [1]. On the other hand, anti-inflammatory effects of exercise have been confirmed [2-4] and its effect has been shown on pro and anti-inflammatory cytokines [5,6], and thus, the protective effect of exercise against mortality is quite plausible. One study has demonstrated a strong and consistent inverse relationship between physical fitness and leukocyte count and markers of inflammation such as serum IL-6 and CRP, whereas serum level IL-10 is positively correlated with fitness [2]. Fischer et al. (2004) have shown that mRNA expression content of exercise-induced IL-6 is markedly reduced in human skeletal muscle after 10 wk endurance training [7], but Robson-Ansley et al. (2007) have shown that plasma level IL-6 content is significantly elevated following intensified training [8]. Although current recommendations for physical fitness involve performing aerobic and resistance exercise with moderate to vigorous intensity for several hours per week, people generally fail to follow such regimes due to lack of time [9]. Nowadays, experts have focused on interval training program with high intensity and low volume [10]. Previous studies have shown that this training induces fat oxidation during training in women [11], improves insulin action in young sedentary men [9] and improves muscle oxidative capacity [10,12,13] and it is superior to moderate intensity exercise in rats [14]. Rakobowchuk et al. (2008) have also confirmed that 6-week sprint interval training with low volume is a time-efficient strategy to elicit improvements in peripheral vascular structure and function that are comparable to endurance training [15]. Some researchers have compared the effects of sprint interval training (SIT) with continuous endurance training (CET) on inflammatory markers, but the results have been controversial. The aim of the current study is to examine whether there is any significant difference between the effect of sprint interval and continuous endurance training on inflammatory markers.
Methods

Subjects
In this randomized trial, 16 students (8 men, 8 women) who had regular recreational activities participated. All subjects took some form of recreational exercise two to three times per week (jogging, cycling, etc.). Subjects' completed two questionnaires before starting the training program: one about baseline information and the other about the type and intensity of their physical activity. Subjects were randomly assigned to either the SIT or the CET training program. Following routine medical screening, the purpose of the study and associated risks were explained for all subjects and written informed consent was obtained from them. The study was approved by the Research Ethics Committee of Shahid Beheshti University.

Pre-experimental procedures
Prior to baseline measurements, subjects were made several familiarization visits to the laboratory in order to be informed of the testing procedures and training devices. During one of these visits, subjects performed an incremental test to exhaustion on an electronically braked cycle ergometer using an online gas collection system to determine \( \text{VO}_2\text{max} \). Following a 5-minute warm-up, the test began at 50W with the workload increasing by 25W every 2 minutes until volitional exhaustion. The value recorded for \( \text{VO}_2\text{max} \) corresponded to the highest value achieved over a 30-second collection period [13]. All exercise tests were performed at least 1 week prior to baseline testing. During the test, heart rate was measured continuously using a heart rate monitor. \( \text{VO}_2\text{max} \) was confirmed using established physiological criteria defined by the British Association of Sport and Exercise Science (BASES), and included a respiratory exchange ratio (RER) above 1.15, oxygen uptake reaching a plateau with increasing work rate, and a heart rate close to age-predicted maximal values [16].

Wingate test
Subjects in the SIT group completed a 30-second maximal effort on an electronically braked cycle ergometer at a resistance equivalent to 7.5% of their body mass.

Training protocol
CET consisted of continuous cycling on an ergometer, 3 days per week for 2 weeks, at a power output corresponding to 65% \( \text{VO}_2\text{max} \). Subjects did exercise training in the first two sessions for 90 minutes, in the second and third sessions for 105 minutes, and in the last two sessions for 120 minutes. The numbers of Wingate tests performed during each training session were increased from four in sessions 1 and 2 to five in sessions 3 and 4, and finally to six in sessions 5 and 6. For all training sessions, the recovery interval between Wingate tests was fixed at 4 minutes during which, subjects rested [10] (Table 1).

Blood sampling and analyses
Venous blood samples were obtained at 8–10 am in the Obesity Research Center, Research Institute for Endocrine Sciences of Shahid Beheshti University of Medical Sciences. Subjects had slept and had fasted for 8–9 hours preceding blood sampling, eat food before 2 blood sampling as same as possible, and were asked not to do heavy exercise, take medications, or get exposed to sensitizers within 24 hours preceding blood sampling. To be included in the study, subjects ought to have had no infection within one week before sampling. Blood samples were taken after 15 minutes of rest in sitting position. The first samples were taken 24 h before the first session and the second samples were taken 48 h after the last session (after 2 weeks). Blood samples (10 ml) were obtained from antecubital vein. Two ml of the blood sample was poured into EDTA tubes for WBC measurement and the rest was centrifuged at 4°C for 5 minutes at 3000 g. After centrifugation, serum was stored at −70°C for subsequent analyses. CRP, IL-10, IL-6 were measured by ELISA kit and insulin resistance was calculated as HOMA-IR.

Statistical analyses
The distribution of variables was assessed by Kolmogorov-Smirnov test. To compare groups, we calculated the difference (after-before) for each group and tested it by T test if the distribution was normal and by Mann-Whitney U test if the distribution was not normal. The before and after mean in each group was tested by Paired T test if the distribution was normal and by Wilcoxon test if the distribution was not normal. P-values less than 0.05 were considered statistically significant. Analysis was done with SPSS, PC program, version 16. (SPSS Inc., Illinois Chicago, USA).

Results
All variables with the exception of CRP had normal distribution. The baseline characteristics of subjects were not statistically different between two groups (Table 2).
Results showed that the type and duration of training had no significant effect on IL-6, IL-10, CRP, WBC and insulin resistance index (P>0.05). The level of IL-6 was different between the two training groups. Although mean level of IL-6 increased after CET and decreased after SIT, it was not statistically significant (P>0.05). Results showed that serum levels of IL-10 and CRP decreased after both training programs but the decrease was not statistically significant (P>0.05). WBC also decreased after both training programs, but the decrease was significant only after CET (P:0.03). In addition, the decrease in insulin resistance index was close to significance after SIT (P:0.053) but it showed no change in CET (P: 0.94) (Tables 3, 4).

Discussion

The major finding of the present study was that neither CET nor SIT has any significant effect on serum levels of IL-6, IL-10, CRP, WBC, and insulin resistance index. To the best of our knowledge, this is the first study on the effects of two types of interval and continuous training with identical format (cycling), frequency (3 days in the week), and duration (2 weeks) but different volume. Previous studies have investigated effects of either interval or continuous training on these markers [7,17-22] but none has studied the effects of both trainings simultaneously.

Our results are consistent with results reported by Rakowchuk et al. (2008) and Gibala et al. (2006) that showed no difference between these two training types [10,15]. However, Rakowchuk had studied untrained subjects with 4.5 min recovery periods. Tejona et al. (2008) had reported that interval training is more efficient than continuous training in improving metabolic parameters in patients with metabolic syndrome [23]. However, the results may have been biased by differences between subjects, protocol and time (16 weeks). Most studies have showed improvements in these cytokines by 6 to 24 weeks of training [7,19-21,24]; therefore, the time required for these changes should be more than 2 weeks. Muscles are sources of IL-6 during exercise [25]. In the present study; we have not taken biopsies from muscles. Previous results have shown that IL-6 response is different between the 2 types of training that may be due to gender as a determinant [26,27]. Another justification is that myofibrils have differences in expression of cytokines [24] and differences between 2 types of training may be due to difference in involvement of slow and fast fibrils. Similarly, Gibala et al. (2006) and Burgomaster et al (2005) reported that SIT can increase glycogen content and muscle oxidative capacity [10,13], for this reason muscles didn’t rely on to IL-6 and its effects about provided substrate from liver and fat tissue [28] and decreased levels IL-6 in this research but CET increased IL-6 that its reason can decrease glycogen content. Low glycogen content is most important factor for this increase probably [29-31] that this situation mostly body rely on fatty acid

### Table 2 Baseline characteristics of subjects in two group

| Variable          | CET       | SIT       | P-value |
|-------------------|-----------|-----------|---------|
| Age(year)         | 25(1.69)  | 22(2.16)  | 0.07    |
| Weight(kg)        | 65.38(1.14)| 65.91(7.88)| 0.96    |
| Height(Cm)        | 1.70(0.09)| 1.71(0.09)| 0.70    |
| BMI(kg/m2)        | 22.44(2.02) | 22.37(2.50) | 0.95    |
| VO2max(ml/kg/min)| 34.17(5.52) | 33.75(6.08) | 0.88    |

The data are expressed as mean (SD).

### Table 3 Inflammatory biomarkers mean in before and after of two group

| Type of training | CET (n=8) | SIT (n=8) | P-value |
|------------------|-----------|-----------|---------|
| Before           | After     | Before    | After   |   |
| IL-6(pg/ml)     | 2.88(0.21) | 2.91(0.45) | 0.87 | 3.00(0.41) | 2.78(0.26) | 0.33 |
| IL-10(pg/ml)    | 7.95(0.96) | 7.85(1.15) | 0.87 | 7.60(1.64) | 7.21(1.46) | 0.64 |
| CRP(ng/ml)      | 461.40(294) | 333(361.34) | 0.20 | 373.5(144.66) | 341.66(325.06) | 0.09 |
| WBC(×10^3/μl)   | 5.97(0.99) | 5.35(0.89) | 0.03 | 6.08(1.12) | 6.02(1.55) | 0.90 |
| HOMA-IR         | 1.17(0.57) | 1.17(0.70) | 0.94 | 1.78(0.58) | 1.24(0.32) | 0.053 |

The data are expressed as mean (SD).

### Table 4 Inflammatory biomarkers mean in before and after of two groups

| Type of training | CET (n=8) | SIT (n=8) | P-value |
|------------------|-----------|-----------|---------|
| Difference (After – Before) |           |           |         |
| IL-6(pg/ml)     | 0.03(0.46) | -0.21(0.54) | 0.38 |
| IL-10(pg/ml)    | 0.10(1.80) | -0.38(2.11) | 0.78 |
| CRP(ng/ml)      | -160(220) | -31.83(347.70) | 0.75 |
| WBC(×10^3/μl)   | -0.62(0.69) | -0.06(1.50) | 0.35 |
| HOMA-IR         | 0.01(0.22) | -0.53(0.65) | 0.06 |

The data are expressed as mean (SD).
and IL-6 that increased lipolysis. In the other hand, long activity induced injured mussels and there is an inflammatory response and produced this cytokine [8]. CRP decreased in both types of training, because IL-6 can induce CRP [32], therefore the decrease in CRP is expected in SIT but not in CET. As Oberbach et al. (2006) reported, we can conclude that CRP is not induces only by IL-6 and other factors can also induce production of CRP [33]. In the present study, IL-10 decreased after both types of training, this cytokine controls inflammation by suppressing the production of pro-inflammatory cytokines [34] but other changes related to regular training such as decreased weight or percent body fat can justify the anti-inflammatory effects of training that has not been investigated in this study [33]. Moreover, IL-10 is produced by T cells, B cells, monocytes and macrophages as well [35], therefore decreased WBC after both types of training can induce decrease in IL-10. The decrease in WBC in CET was more prominent than SIT which may be due to Fitness [36] and increased plasma volumes after endurance training [37] can explain decreased WBC. Finally, insulin resistance index decreased in SIT but did not change in CET. Increased oxidative capacity of muscles, less dependence of body on IL-6, and increased sensitivity to IL-6, lead to decreased requirement of the body to glucose in blood. However, increased IL-6 after CET showed that adoptions after training have not been sufficient for the required activity within muscles and substrate should have been provided to cells from outside, which explains why no decrease in insulin resistance index was observed. Effects of CET remained after the last session for a short time period because half-life of 4 GLUT is short [38]. Therefore, slightly blood sampling has been too late for this measurement. The main limitation of this study is low sample size which decrease power of current study. The decrease in IL-6 and CRP serum levels after SIT requires studies with longer follow-up (more than 6 sessions) in order to improve health status of people by means of shorter but more efficient training.

**Conclusion**

Our results shows that SIT with high intensity and low volume, and CET with moderate intensity and high volume have identical effects on the body. There were no significant improvements in inflammatory biomarkers, but the decrease in insulin resistance index was marginally significant.

**Endnote**

aAny substance to which subjects were sensitive and had mentioned it in the questionnaire.

**Competing interests**

The authors declared that they have no competing interests.

**Authors’ contribution**

FH participated in study designed and drafted the manuscript. TA participated in study designed performing laboratory tests, data acquisition and drafted the manuscript. SA participated in study designed and statistical analysis. All authors read and approved the final manuscript.

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