Heart rate variability responses to a combined exercise training program: correlation with adiposity and cardiorespiratory fitness changes in obese young men

Jatuporn Phoemsathawee¹, Piyapong Prasertsri², Naruemon Leelayuwat²,³,⁴*¹

¹Department of Sports Science and Health, Faculty of Sports Science, Kasetsart University, Nakhon Pathom, Thailand ²Faculty of Allied Health Sciences, Burapha University, Chonburi, Thailand ³Department of Physiology, Faculty of Medicine, Khon Kaen University, Khon Kaen, Thailand ⁴Exercise and Sport Sciences Development and Research Group, Khon Kaen University, Khon Kaen, Thailand

Although the influence of adiposity indices and cardiorespiratory fitness (CRF) on heart rate variability (HRV) has been demonstrated extensively, the causal link between the changes in adiposity as well as in CRF and the alterations in cardiac autonomic function is unclear. Thus, this study aimed to assess the correlation between the changes in adiposity and CRF and the alterations in HRV after 12-week exercise training. Twenty obese sedentary men aged 20.5 ± 1.2 years were randomly assigned into 2 groups (n = 10 each): the control (CG) and the exercise group (EG). The EG trained 60 min of combined aerobic, anaerobic and strengthening exercise, 4 sessions/wk for 12 weeks, whilst the CG remained relatively inactive. Measurements of resting HRV, body composition, and peak oxygen consumption (VO₂peak) were obtained at baseline and after the 12-week training program. Compared with CG, the exercise training significantly reduced adiposity indices and improved vagal-related HRV variables and VO₂peak. Significant correlations were observed between changes in HRV variables and adiposity indices and VO₂peak changes. Stepwise regression analysis revealed that changes in a Poincaré plot index (SD1/SD2 ratio) predicted 32.4% of the variance in the relative VO₂peak changes. These findings suggest that obese sedentary young men achieved significant improvements in vagal activity, adiposity indices and aerobic fitness after the exercise training. The higher reduction in fat mass, especially central obesity, the greater alteration of vagal modulation. Moreover, the alteration in resting HRV is a possible predictor for adaptations to exercise training in obese sedentary young men.

Keywords: Exercise training, Central obesity, Heart rate, Exercise test

INTRODUCTION

The prevalence of obesity continues to rise and presents as a major public health concern in Thailand. Essentially, the prevalence of obesity in the 18–24 age group has been doubled for over 10 years (Aekplakorn and Mo-Suwan, 2009). Obesity rises the risks of cardiovascular diseases (CVD) and all-cause of mortality (Jensen et al., 2014). Obesity is also associated with dysregulation of cardiac autonomic function (Lindmark et al., 2005). It has been reported that autonomic dysregulation is an important mediator in the development of CVD risks in obese individuals (Lindmark et al., 2005; Sheema and Malipatil, 2015). Assessment of heart rate variability (HRV) has been recognized as a noninvasive method for assessing cardiac autonomic modulation (Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology, 1996). Reduced HRV has been associated with higher weight and adiposity (Gutin et al., 2005; Tian et al., 2015). On the other hand, weight reduction via exercise training and/or calorie restriction in obese subjects has been suggested to improve vagal activity (Facchini et al., 2003; Ito et al., 2001; Tian et al., 2015). The increase in vagal activity in response to weight loss program may consistently contribute to reduce the risk of cardiovascular morbidity and of sudden cardiac death (Facchini et al., 2003; Gutin et al., 2005; Ito et al., 2001).
However, little is known regarding how exercise-induced body composition changes HRV.

Many have reported that exercise training can alter parasympathetic activity (Facchini et al., 2003; Gutin et al., 1997; Gutin et al., 2005; Ito et al., 2001; Kiilavuori et al., 1995; Levy et al., 1998; Nagai and Moritani, 2004; Stein et al., 1999; Tan et al., 2015). In contrast, some researches have not shown such effect (Bourcher and Stein, 1995; Davy et al., 1997). The influence of exercise-induced HRV changes is controversial probably due to different exercise training programs and measures of HRV. In addition, most studies are cross sectional and have been conducted on children and adolescents (Gutin et al., 2000; Gutin et al., 2005; Lucini et al., 2013; Nagai and Moritani, 2004) or obese individuals with chronic diseases (Pagani and Lucini, 2001; Sjoberg et al., 2011). There has been very little research reported on the exercise-induced body composition and aerobic fitness changes HRV (Ito et al., 2001; Tan et al., 2015), particularly in obese sedentary young adults. There is also a lack of data in males (Ito et al., 2001). Therefore, this study attempted to clarify the casual link between the changes in body composition as well as in aerobic fitness and the alterations in cardiac autonomic modulation after a 12-week exercise training. Understanding how exercise-induced body composition and aerobic fitness alters HRV may lead to insights into the risks of CVD and a mean to predict the exercise adaptation in obese sedentary young adults.

**MATERIALS AND METHODS**

**Subjects and study design**

The randomized controlled trial conducted, for which 42 obese men aged 19 to 22 years with body mass index (BMI) ≥25 kg/m² and percentage of body fat (%BF) >24% (American College of Sports Medicine, 2010), were recruited. Subjects were no regular physical exercise in the last 6 months. Subjects were excluded if they had any clinical sign of CVD, neurological, musculoskeletal limitations to exercise or other overt chronic diseases, or current medication use. Subjects completed a Physical Activity Readiness Questionnaire (PAR-Q) before participating in the study. Subjects were informed of potential risks and the procedure of the study before they signed a written informed consent form. The study protocol was approved by the Kasetsart University Research Ethics Committee (COA61/033) and was conducted according to Declaration of Helsinki.

After the initial evaluation, the 20 eligible obese subjects (age, 20.5 ± 1.2 years; BMI, 31.4 ± 5.2 kg/m²; %BF, 30.0% ± 6.5%) were randomly assigned (1:1) to the exercise (EG, n = 10) or control (CG, n = 10) group, which were group-matched by age, BMI, and aerobic fitness. The EG group underwent a 12-week combined exercise training program with 4 × 60-min supervised sessions per week and was asked to maintain habitual diet throughout the study. The CG group did not participate in any weight control programs and was asked to maintain habitual diet and physical activity levels during the study. Measurements were performed in each subject at baseline and 1 week after the last exercise session. Anthropometrics and body composition, peak oxygen consumption (VO₂peak), and resting HRV were measured at the same time of the day for each subject. Prior to test, subjects were asked to abstain from consumption of alcohol or caffeine-containing beverages for 24 hr. Smoking was also not allowed for at least 4 hr. To ensure consistent baseline activity levels, they were also instructed to avoid intense and/or prolonged exercise for 2 days.

**Exercise intervention**

The EG subjects participated in a 4 × 60-min sessions/wk of combined exercise program for 12 weeks and was assigned to increase energy expenditure by 452 ± 48.1 kcal/session or 1,825 ± 112.4 kcal/week (Rippe and Hess, 1998). Training sessions consisted of 2 × 30-min sessions/wk with continuous moderate-intensity aerobic exercise (fast walking, jogging, ball games, or swimming) at an intensity between 50% to 70% of individual maximum heart rate (HRmax) and 10 × 1-min high-intensity interval training for 2 sessions/wk at an intensity between 80% to 90% of HRmax and interspersed with 2 min of recovery, followed by 20 min of strengthening exercises and 10 min of stretching and cooldown. The strengthening exercises were consisted of 2 to 3 series of 10 to 15 repetitions of the arms, legs, and trunk. The strengthening period was determined by the subject’s body weight. HR was monitored continuously during each training session using a HR monitor (Forerunner 220, Garmin Ltd., Schaffhausen, Switzerland) and work rate was adjusted to maintain target HR. The training sessions were carefully supervised by experienced trainers. In addition, besides the supervised training sessions, the EG subjects received a recommendation to undergo unsupervised physical activity (by walking 30 min every day). The frequency and duration of unsupervised activities were assessed by a weekly recall questionnaire. The CG subjects did not participate in any intervention and was asked to maintain their habitual diet and physical activity levels throughout the study.
Anthropometrics and body composition measurements

Body mass and body composition, including %BF, fat mass (FM), and fat free mass (FFM), were measured using a bioimpedance analysis device (Inbody 720, Biospace Inc., Seoul, Korea) with light clothing and without shoes. Height was measured without shoes using a standard stadiometer (Health o Meter Professional, Sunbeam Products Inc., Boca Raton, FL, USA). The BMI was calculated as body mass divided by height squared (kg/m²). Waist circumference (W) was measured form midway between the lower rib margin and the iliac crest at the end of inspiration, using a flexible and inextensible measuring tape (Hochetsmass Balzer GmbH, Sulzbach, Hessen, Germany). Hip circumference (H) was also measured at the level of trochanter major and the waist-to-hip circumferences (W/H) ratio was calculated.

VO2peak measurement

The VO2peak was measured with an incremental exercise test on an electromagnetically braked cycle ergometer (VIAsprint 150 P, Ergoline GmbH, Bitz, Germany). Briefly, the test was started at a 5-min seated rest on the cycle ergometer, followed by 3 min of baseline unloaded cycling. The incremental protocol was initiated with a work rate of 50 W, and the work rate was increased by 25 W every 2 min. The subjects were encouraged to maintain the fixed pedaling frequency at 50 to 60 rev/min until volitional exhaustion, after which the subjects completed an unloaded recovery period of 5 min.Expired gas samples were collected on a breath-by-breath basis using a portable metabolic device (JAEGER Oxycon Mobile, CareFusion, Hoechberg, Germany), which was calibrated before each test using a 3-L syringe and known concentrations of oxygen (15% O2) and carbon dioxide (5% CO2). Oxygen consumption (VO2), carbon dioxide production (VCO2), and HR were continuously recorded and analyzed throughout the test. Arterial O2 saturation was continuously monitored using a fingertip pulse oximeter (OLV-3100K, Nihon Kohden, Tokyo, Japan). Rating of perceived exertion was obtained using the Borg scale (6–20) at the end of each work rate. Blood pressure (BP) was recorded using an automatic sphygmomanometer (Tango M2, SunTech Medical Inc., Morrisville, NC, USA) in the last 30 sec of each workload. The VO2peak was defined as the highest 30-sec average value of VO2. The VO2peak was achieved when at least three of the following four criteria were met: (a) a plateau in VO2 despite an increasing work load, (b) a respiratory exchange ratio > 1.1, (c) a HR within 10/min of HRmax, and (d) volitional exhaustion (Howley et al., 1995).

HRV recordings and analysis

The RR intervals were recorded using a 5-min electrocardiogram (ECG) recording (eMotion Faros device, Mega Electronics, Kuopio, Finland) in lying position. The ECG recording was collected online at a sampling rate of 1,000 Hz, in real time, and stored on a computer. The ECG recording took place at a fixed time in the morning (between 8:00 a.m. and 10:00 a.m.) to avoid possible circadian influences on autonomic function. Prior to the recording, subjects rested comfortably supine for at least 20 min in a quiet air-conditioned room with a room humidity and temperature of 67%±1.8% and 25°C±1.1°C, respectively. The respiratory rate, determined by visual inspection of chest movement, was within the normal range (12–20/min).

The RR intervals were analyzed via a software program using the recommendations of the Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology (Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology, 1996). Time- and frequency-domain parameters together with the Poincare’ plot or nonlinear components of HRV were calculated. The time-domain parameters consisted of mean RR intervals, standard deviation of all normal RR intervals (SDNN) and squared differences between adjacent normal NN intervals (RMSSD). The mean RR intervals and the SDNN provided information on sympathetic and parasympathetic cardiac modulation, and the RMSSD subsequently provided information on parasympathetic cardiac modulation. The frequency-domain parameters consisted of low frequency power (LF power: 0.04–0.15 Hz) and high-frequency power (HF power: 0.15–0.4 Hz) and the low-frequency/high-frequency (LF/HF) ratio. The LF power indicated sympathetic and vagal modulations simultaneously, the HF power indicated cardiac vagal modulation, and the LF/HF ratio indicated cardiac sympathetic–vagal balance (Ca*9/Poincaré plot is a method to illustrate non-linear HRV components. Poincare’ parameters, SD1 (Poincaré plot standard deviation perpendicular the line of identity) and SD2 (Poincaré plot standard deviation along the line of identity), were analyzed quantitatively by calculating the standard deviations from the RR interval data. The SD1 related to the fast beat-to-beat variability, while the SD2 described the longer-term variability. The SD1/SD2 ratio reflected non-linear components of HRV (Rajendra Acharya et al., 2006; Tulppo et al., 1996).

Statistical analysis

All data are presented as mean±standard deviation. Normality of data distribution was assessed using the Shapiro–Wilk normal-
Heart rate variability responses to combined exercise training

Phoemsapthawee J, et al.

Heart rate variability variables measured at baseline and after 12-week exercise training

| Variable          | CG (n = 10) | EG (n = 10) |
|-------------------|------------|------------|
|                   | Baseline   | 12 Weeks   | Baseline   | 12 Weeks   |
| Resting HR (msec) | 67.8 ± 6.1 | 69.2 ± 7.1 | 72.1 ± 7.6 | 66.3 ± 3.7†† |
| Mean RR intervals (msec) | 6.8 ± 0.1 | 6.7 ± 0.1 | 6.7 ± 0.1 | 6.8 ± 0.1†† |
| SDNN (msec)       | 4.2 ± 0.4 | 4.1 ± 0.5 | 4.2 ± 0.3 | 4.2 ± 0.3 |
| RMSSD (msec)      | 3.9 ± 0.9 | 3.7 ± 0.6 | 3.9 ± 0.4 | 3.9 ± 0.3 |
| pNN50 (%)         | 25.7 ± 18.3 | 27.2 ± 24.0 | 30.2 ± 16.6 | 33.7 ± 15.4 |
| TP (msec²)        | 8.1 ± 0.9 | 7.8 ± 1.1 | 8.3 ± 0.5 | 8.1 ± 7.2 |
| HF (msec²)        | 6.9 ± 1.1 | 6.3 ± 1.3 | 6.7 ± 0.9 | 6.9 ± 0.1 |
| HF (nu)           | 30.6 ± 12.3 | 24.8 ± 11.9 | 24.6 ± 14.0 | 33.0 ± 12.6†† |
| LF (msec²)        | 7.2 ± 1.0 | 6.9 ± 1.2 | 7.3 ± 0.8 | 6.8 ± 1.4† |
| LF (nu)           | 40.8 ± 16.2 | 41.2 ± 17.7 | 39.3 ± 18.2 | 38.2 ± 14.2 |
| LF/HF ratio       | 1.9 ± 2.2 | 2.6 ± 3.1 | 2.4 ± 2.1 | 1.7 ± 1.5†† |
| SD1               | 47.0 ± 30.6 | 35.6 ± 25.7 | 38.2 ± 12.0 | 38.3 ± 13.0 |
| SD2               | 94.0 ± 36.5 | 85.7 ± 43.3 | 92.8 ± 21.4 | 88.1 ± 24.6** |
| SD1/SD2 ratio     | 0.4 ± 0.1 | 0.4 ± 0.1 | 0.4 ± 0.1 | 0.4 ± 0.1 |

Values are presented as mean ± standard deviation. CG, control group; EG, exercise group; HR, heart rate; SDNN, standard deviation of normal RR intervals; RMSSD, square differences between adjacent normal NN intervals; pNN50, percentage difference between adjacent normal RR intervals > 50 msec; TP, total power; HF, high-frequency power; LF, low-frequency power; LF/HF ratio, low-frequency to high-frequency ratio; SD1, Poincaré plot standard deviation perpendicular the line of identity; SD2, Poincaré plot standard deviation along the line of identity.

*P < 0.05, **P < 0.01, significant difference from baseline within group. †P < 0.05, significant difference in mean change between groups.

Adiposity and cardiorespiratory fitness measured at baseline and after 12-week exercise training

| Variable          | CG (n = 10) | EG (n = 10) |
|-------------------|------------|------------|
|                   | Baseline   | 12 Weeks   | Baseline   | 12 Weeks   |
| Age (yr)          | 20.7 ± 1.2 | -           | 20.3 ± 1.3 | -           |
| Height (cm)       | 174.6 ± 5.7 | -           | 174.6 ± 6.6 | -           |
| Body mass (kg)    | 97.8 ± 19.6 | 97.8 ± 19.6 | 97.8 ± 19.6 | 97.8 ± 19.6†† |
| BMI (kg/m²)       | 31.4 ± 5.3 | 31.4 ± 5.3 | 31.4 ± 5.3 | 31.4 ± 5.3** |
| Body fat (%)      | 29.6 ± 13.4 | 29.9 ± 13.3 | 30.5 ± 13.5 | 28.6 ± 13.5 |
| Fat mass (kg)     | 68.7 ± 8.2 | 69.0 ± 9.0 | 65.0 ± 8.9 | 65.0 ± 8.0** |
| FFM (kg)          | 39.3 ± 5.0 | 39.5 ± 5.6 | 37.1 ± 5.1 | 37.3 ± 4.5 |
| W (cm)            | 100.9 ± 15.6 | 102.3 ± 15.8 | 100.2 ± 15.2 | 97.7 ± 14.7†† |
| H (cm)            | 112.7 ± 8.5 | 112.6 ± 9.5 | 111.2 ± 10.5 | 108.1 ± 11.0†† |
| W/H ratio         | 0.91 ± 0.04 | 0.91 ± 0.04 | 0.91 ± 0.04 | 0.90 ± 0.04** |
| VO_peak (L/min)   | 2.6 ± 0.4 | 2.6 ± 0.4 | 2.5 ± 0.3 | 2.8 ± 0.3** |
| VO_peak (ml/kg/min) | 27.1 ± 5.3 | 27.0 ± 5.4 | 28.9 ± 5.4 | 30.5 ± 4.8** |

Values are presented as mean ± standard deviation. CG, control group; EG, exercise group; BMI, body mass index; FFM, fat free mass; SWM, skeletal muscle mass; W, waist circumference; H, hip circumference; W/H ratio, waist to hip circumference; VO_peak, peak oxygen consumption.

**P < 0.01, significant difference from baseline within group. †P < 0.05, ††P < 0.01, significant difference in mean change between groups.

Effects of combined exercise training

Body mass (P < 0.05), BMI (P < 0.01), FM (P < 0.01), W (P < 0.01), H (P < 0.01), and W/H ratio (P < 0.01) were significantly reduced in the EG group following the 12-week combined exercise training as compared to baseline levels. Meanwhile, there were no significant differences in any anthropometry or body composition variables in the CG group as compared to baseline levels. In the EG group, the mean reduction in body mass (P < 0.01), BMI (P < 0.01), FM (P < 0.05), W (P < 0.01), H (P < 0.05), and W/H ratio (P < 0.05) following the 12-week combined exercise training were significantly different from the CG group. The absolute (P < 0.01) and relative VO_peak (P < 0.01) following the 12-week combined exercise training increased significantly in the EG group compared to baseline values. Meanwhile, there was no significant difference in VO_peak in the CG group compared to baseline values (Table 1). In the EG group, the mean improvements in absolute (P < 0.01) and relative VO_peak (P < 0.01) following the 12-week exercise training were compared to baseline values.
Heart rate variability responses to combined exercise training

Phoemsapthawee J, et al. • Heart rate variability responses to combined exercise training

combined exercise training were significantly different from the CG group (Table 2).

The HRV variables following exercise training are presented in Table 1. Mean RR intervals ($P < 0.05$) and normalized values of HF ($P < 0.05$) following the 12-week combined exercise training increased significantly, while resting HR ($P < 0.05$), LF (ln msec$^2$) ($P < 0.01$), LF/HF ratio ($P < 0.05$), and SD2 ($P < 0.01$) following the 12-week combined exercise training was significantly reduced

**Table 3. Correlation coefficients ($r$) of changes between adiposity indices and aerobic fitness and heart rate variability variables**

| Variable          | $\Delta$ RR intervals (msec) | $\Delta$SDNN (msec) | $\Delta$RMSSD (msec) | $\Delta$pNN50 (%) | $\Delta$TP (msec$^2$) | $\Delta$HF (msec$^2$) | $\Delta$LF (msec$^2$) | $\Delta$LF/HF ratio | $\Delta$SD1/SD2 ratio |
|-------------------|-------------------------------|--------------------|---------------------|------------------|-----------------------|-----------------------|----------------------|---------------------|----------------------|
| Body mass (kg)    | -0.411                        | -0.112             | 0.093               | 0.033            | -0.129                | -0.234                | -0.081               | -0.286              | 0.008                |
| BMI (kg/m$^2$)    | -0.406                        | -0.092             | 0.082               | 0.077            | -0.092                | -0.259                | -0.081               | -0.286              | 0.008                |
| Fat mass (kg)     | -0.502*                       | 0.008              | 0.073               | -0.007           | -0.134                | -0.211                | -0.138               | -0.258              | 0.026                |
| FFM (kg)          | 0.085                         | -0.109             | 0.105               | 0.080            | 0.024                 | 0.056                 | 0.050                | -0.024              | 0.064                |
| Fat mass (%)      | -0.113                        | -0.013             | 0.059               | 0.495*           | 0.016                 | 0.022                 | 0.378                | -0.185              | 0.020                |
| BMI (kg/m$^2$)    | -0.450*                       | 0.028              | 0.045               | -0.029           | -0.109                | -0.101                | -0.165               | -0.284              | -0.066               |
| Waist circumference (cm) | 0.153                      | -0.001             | 0.169               | 0.360            | -0.033                | 0.240                 | 0.459*               | -0.373              | -0.233               |
| VO$_{2peak}$ (L/min) | 0.225                     | 0.100              | 0.270               | 0.471*           | 0.044                 | 0.310                 | 0.425*               | -0.349              | -0.212               |

Absolute change ($\Delta$: 12-week value – baseline value).

BMI, body mass index; FFM, fat free mass; W, waist circumference; W/H ratio, waist-to-hip circumference; VO$_{2peak}$, peak oxygen consumption; SDNN, standard deviation of normal RR intervals; RMSSD, square root of the mean squared difference between adjacent normal RR intervals; pNN50, percentage difference between adjacent normal RR intervals >50 msec; TP, total power; HF, high-frequency power; LF, low-frequency power; LF/HF, low-frequency to high-frequency ratio; SD1, Poincaré plot standard deviation perpendicular the line of identity; SD2, Poincaré plot standard deviation along the line of identity.

n=20. *$P<0.05$, **$P<0.01$.

**Fig. 1.** Correlation between (A) the change in waist circumference and the change in pNN50 ($r=0.495$, $P<0.05$), (B) the change in waist-to-hip ratio and the change in mean RR intervals ($r=-0.045$, $P<0.05$), (C) the change in fat mass and the change in mean RR intervals ($r=-0.502$, $P<0.05$), (D) the change in relative VO$_{2peak}$ and the change in pNN50 ($r=0.471$, $P<0.05$), (E) the change in relative VO$_{2peak}$ and the change in the SD1/SD2 Poincaré plot index ($r=0.57$, $P<0.01$), (F) the change in relative VO$_{2peak}$ and the change in the LF/HF ratio ($r=-0.444$, $P<0.05$). ● EG group; ○ CG group; pNN50, percentage of successive RR intervals that differ by more than 50 msec; SD1, Poincaré plot standard deviation perpendicular the line of identity; SD2, Poincaré plot standard deviation along the line of identity; LF, low-frequency; HF, high-frequency; VO$_{2peak}$, peak oxygen consumption.
Correlation between the changes in HRV and the changes in adiposity indices and aerobic fitness over the training period

The correlation coefficients (r) for changes between HRV variables and adiposity indices and aerobic fitness are presented in Table 3. Pearson correlation between the changes in adiposity and HRV changes showed significant positive correlation of W changes with Δ pNN50 (r = 0.495, P < 0.05) (Fig. 1A), and negative correlation of W/H ratio (r = -0.450, P < 0.05) (Fig. 1B) and FM changes (r = -0.502, P < 0.05) with Δ mean RR intervals (ln msec) (Fig.1C). Significant positive correlations were observed between changes of VO2peak and Δ pNN50 (r = 0.471, P < 0.05) (Fig. 1D), and Δ HFnu (r = 0.425, P < 0.05), and Δ SD1/SD2 ratio (r = 0.570, P < 0.01) (Fig. 1E). Meanwhile, the changes of VO2peak and Δ LF/HF ratio (r = -0.470, P < 0.05) (Fig. 1F) showed significant negative correlation (Table 3).

Subjects with greater individual changes in vagal-related HRV variables had more reduction in central obesity, W and W/H ratio, and a greater increase in VO2peak over the training period (Fig. 1). The results of the multiple stepwise regression analysis revealed that only the changes in SD1/SD2 ratio explained a significant proportion of the variance in the relative VO2peak changes (R2 = 0.324, P < 0.01) (Table 4).

### DISCUSSION

Our results indicate that vagal activity improves with fat loss and increased aerobic fitness after the exercise training in obese sedentary young men. Changes of the vagal-related HRV variables were significantly related to fat loss. Moreover, the subjects with a greater sympathovagal balance had greater improvement in the aerobic fitness (relative VO2peak). These finding suggest that the alteration in resting HR, especially the SD1/SD2 ratio is a possible predictor for adaptations to exercise training in obese sedentary young men.

Obesity is associated with dysregulation of cardiac autonomic function and rises the risks of CVD. The cardiac autonomic dysfunction has been attributed to excess fat and weight gain (Sheema and Malipatil, 2015). Our findings indicate that the higher reduction in FM and central obesity (measured by W and W/H ratio), the greater alteration of vagal-related HRV variables (mean RR intervals and pNN50). Our study confirms the results of previous studies in which cardiac parasympathetic activity increased with fat loss after exercise training (Tian et al., 2015). It was difficult to compare our results with the previous study because our subjects were all men, whereas the other analyzed men and women together (Tian et al., 2015). It has been suggested that gender influences HRV; healthy men at age < 30 years have been reported to have greater HRV variables than women, and age also affects cardiac autonomic control of HR (Umetani et al., 1998). A previous study (Rissans et al., 2001) reported that the vagal activity related to a large weight loss by an average of 9.5% (8.7 kg). However, in our present study, the vagal-related HRV variables correlated to the reduction in FM and central obesity instead of the weight loss (2.3% or 2.2 kg). The interindividual differences in mean RR intervals with fat loss (6.6% or 2 kg) were observed in this study. Indeed, the changes in central obesity (measured by W and W/H ratio) correlated with changes in vagal-related HRV variables (mean RR intervals and pNN50). The more central fat loss, the more increased vagal activity. Obese individuals with higher central fat were sympathovagal imbalance compared to those with lower central fat (Soares-Miranda et al., 2011). Another study confirmed the improvements in vagal-related HRV variables elicited by exercise training correlated with altered trunk fat percentage and/or

---

**Table 4. Stepwise liner regression analysis for factors associated with the relative VO2peak.**

|                          | Changes in VO2peak (mL/kg/min) |
|--------------------------|--------------------------------|
|                          | Unstandardized coefficients | Standardized coefficients | t  | P-value | R   | R2  |
| Constant                 | 2.039                        | 0.516                       | -  | 3.953 | 0.001 | -   | -  |
| ΔSD1/SD2 ratio           | 8.141                        | 2.769                       | 0.570 | 2.940 | 0.009 | 0.570** | 0.324** |

VO2peak, peak oxygen consumption; SD1, Poincaré plot standard deviation perpendicular the line of identity; SD2, Poincaré plot standard deviation along the line of identity; SEM, standard error of the mean. **P<0.01.
W in overweight and obese adults (Tian et al., 2015). Thus, the regional fat distribution might contribute to the individual differences in HRV response (Chen et al., 2008). These findings demonstrate that central obesity influences vagal activity after exercise training. Although weight reduction in obese subjects has been suggested to improve parasympathetic activity (Karason et al., 1999; Rissanen et al., 2001), such effect was not observed in (Mi-nami et al., 1999). Thus, the casual link between weight loss and beneficial alterations in cardiac autonomic function is still unclear. In addition, influence of continuous endurance training with moderate intensity on resting HRV was not observed in obese women without fat loss (Figueroa et al., 2007). Nevertheless, such influence of a progressive-intensity training was observed with fat loss (Tian et al., 2015). In our study, an improved vagal-related HRV variables was observed after the exercise training in obese seden-
tary young men with fat loss. This might demonstrate a correla-
tion between the fat loss due to exercise training and the improved HRV. The vagal-related HRV variables appear to be influenced in individuals with fat loss, especially central fat in our study. How-
ever, low R-squared values ($R^2 = 0.252, P<0.05$) revealed that re-
duced FM does not fully explain the improvement in HRV. Alter-
atations of potential contributing factors might be involved, such as insulin resistance (Lindmark et al., 2005), leptin (Quilliot et al.,
2008), and inflammation (Akinci et al., 2008).

Low aerobic fitness and abnormalities of cardiac autonomic modulation are significant contributors to increased CVD risk in obese individuals (Grassi et al., 2004; Oktay et al., 2017; Piccirillo et al., 1998; Wei et al., 1999). Our findings indicate that the 12-week combined exercise training program is able to improve aerobic fitness and to favorably modify sympathovagal balance, with increases in mean RR intervals and normalized HF and decreases in resting HR and the LF/HF ratio. These findings are consistent with previous studies supporting the idea that training improves autonomic function (Buchheit and Gindre, 2006; Facchini et al., 2003). Interestingly, the reduction in resting HR may explain why we observed significant increases in normalized HF values and a reduction in the LF/HF ratio, as the variables are primarily mediated by RR intervals. It has been suggested that lower resting HR can result from an increase in vagal and a reduction in sympathetic outflow. The adaptations to exercise training appear to affect both branches of the autonomic nervous system (ANS) (Carter et al., 2003). Our findings are consistent with the improvement in sympathovagal balance after the exercise training already suggested by the HRV variables (mean RR intervals, HFnu, and LF/HF ratio). The reduction in resting HR possibly contributes to the improvement observed in aerobic fitness (Buchheit and Gindre, 2006; Facchini et al., 2003). Our results confirm that moderately strong correlations were found between the changes in pNN50, HFnu, LF/HF ratio, and SD1/SD2 ratio versus the increased in aerobic fitness. These associations concur with previous studies which revealed that higher aerobic fitness is associated with a higher vagal cardiac control (Carter et al., 2003; Grant et al., 2013; Nagai and Moritani, 2004). Increased parasympathetic and decreased sympathetic outflow to the heart elicited by exercise training is typically considered a factor in cardioprotection (Billman and Kukiellka, 2007). Our findings are supported by the stepwise regression analysis revealing that changes in the SD1/SD2 ratio can predict 32.4% of the variance in the relative VO2peak changes. The SD1/SD2 ratio is the most important predictor of changes in aerobic fitness elicited by exercise training. The SD1 indicates the parasympathetic activity, the SD2 indicates the sympathetic modulation, and the SD1/SD2 ratio indicates the sympathovagal balance (Shaffer and Ginsberg, 2017). Moreover, the high SD1/SD2 ratio can be used as an indicator of healthy cardiac dynamics (Shaffer and Ginsberg, 2017). As the SD1/SD2 ratio is proposed to be a good measure of sympathovagal balance, it can be assumed that an increased parasympathetic and a decreased sympathetic outflow to the heart may be related to improved performance.

To our knowledge, this is the first study detailing the effect of exercise training in the ANS response to fat loss and aerobic capacity in obese sedentary young men. Sympathovagal balance elicited by the exercise training correlated with fat loss and aerobic fitness. These findings appeared to support a concept of cardiovascular protection that regular physical activity could reduce mortality and morbidity of CVD. Moreover, our results suggested that resting HRV, especially the SD1/SD2 ratio, might be a predictor of CRF in overweight and obese young men. However, the study had limitations. The number of subjects was small. Furthermore, there were no data in females; neither were sex-related dif-
finitions in HRV studied.

These findings suggested that obese sedentary young men achieved significant improvements in vagal activity, adiposity indices and aerobic fitness after 12 weeks of the combined exercise training. The improvements in vagal activity elicited by the exercise training program correlated with fat loss in obese sedentary young men. Moreover, the subjects with a greater sympathovagal balance had greater improvement in the aerobic fitness. These findings suggest that the alteration in resting HRV is a possible predictor for adaptations to exercise training in obese sedentary
young men.

**CONFLICT OF INTEREST**

No potential conflict of interest relevant to this article was reported.

**ACKNOWLEDGMENTS**

The authors would like to thank the Exercise and Sport Sciences Development and Research Group, Khon Kaen University, Khon Kaen, Thailand. The authors would also like to thank the participants for their enthusiastic participation in this study.

**REFERENCES**

Aekplakorn W, Mo-Suwan L. Prevalence of obesity in Thailand. Obes Rev 2009;10:589-592.

Akinci G, Akinci B, Coskun S, Bayindir P, Hekimsoy Z, Ozmen B. Evaluation of markers of inflammation, insulin resistance and endothelial dysfunction in children at risk for overweight. Hormones (Athens) 2008;7:156-162.

American College of Sports Medicine. ACSM’s guidelines for graded exercise testing and prescription. 8th ed. Philadelphia (PA): Lippincott Williams & Wilkins; 2010.

Billman GE, Kukielka M. Effect of endurance exercise training on heart rate onset and heart rate recovery responses to submaximal exercise in animals susceptible to ventricular fibrillation. J Appl Physiol (1985) 2007;102:231-240.

Boutcher SH, Stein P. Association between heart rate variability and training response in sedentary middle-aged men. Eur J Appl Physiol Occup Physiol 1995;70:75-80.

Buchheit M, Gindre C. Cardiac parasympathetic regulation: respective associations with cardiorespiratory fitness and training load. Am J Physiol Heart Circ Physiol 2006;291:H451-458.

Carter JB, Banister EW, Blaber AP. Effect of endurance exercise on autonomic control of heart rate. Sports Med 2003;33:33-46.

Chen GY, Hsiao TJ, Lo HM, Kuo CD. Abdominal obesity is associated with autonomic nervous derangement in healthy Asian obese subjects. Clin Nutr 2008;27:212-217.

Davy KP, Willis WL, Seals DR. Influence of exercise training on heart rate variability in post-menopausal women with elevated arterial blood pressure. Clin Physiol 1997;17:31-40.

Facchini M, Malfatto G, Sala L, Silvestri G, Fontana P, Lafortuna C, Sartorio A. Changes of autonomic cardiac profile after a 3-week integrated body weight reduction program in severely obese patients. J Endocrinol Invest 2003;26:138-142.

Figueroa A, Baynard T, Fernhall B, Carhart R, Kanaley JA. Endurance training improves post-exercise cardiac autonomic modulation in obese women with and without type 2 diabetes. Eur J Appl Physiol 2007;100:437-444.

Grant CC, Murray C, Janse van Rensburg DC, Fletcher L. A comparison between heart rate and heart rate variability as indicators of cardiac health and fitness. Front Physiol 2013;4:337.

Grassi G, Dell’Oro R, Facchini A, Quarti Trevano F, Bolla GR, Mancia G. Effect of central and peripheral body fat distribution on sympathetic and baroreflex function in obese normotensives. J Hypertens 2004;22:2363-2369.

Gutin B, Barbeau P, Litaker MS, Ferguson M, Owens S. Heart rate variability in obese children: relations to total body and visceral adiposity, and changes with physical training and detraining. Obes Res 2000;8:12-19.

Gutin B, Howe C, Johnson MH, Humphries MC, Sniæder H, Barbeau P. Heart rate variability in adolescents: relations to physical activity, fitness, and adiposity. Med Sci Sports Exerc 2005;37:1856-1863.

Gutin B, Owens S, Slavens R, Sjøgaard G, Treiber F. Effect of physical training on heart-period variability in obese children. J Pediatr 1997;130:938-943.

Howley ET, Bassett DR Jr, Welch HG. Criteria for maximal oxygen uptake: review and commentary. Med Sci Sports Exerc 1995;27:1292-1301.

Ito H, Ohshima A, Tsuzuki M, Ohto N, Yanagawa M, Maruyama T, Kaji Y, Kanaya S, Nishioka K. Effects of increased physical activity and mild calorie restriction on heart rate variability in obese women. Jpn Heart J 2001;42:459-469.

Jensen MD, Ryan DH, Apovian CM, Ard JD, Cornazzie AG, Donovan KA, Hu FB, Hubbard VS, Jakicic JM, Kushner RF, Loria CM, Millen BE, Monas CA, Pi-Sunyer FX, Stevens J, Stevens VL, Wadden TA, Wolfe BM, Yanovski SZ. American College of Cardiology/American Heart Association Task Force on Practice Guidelines; Obesity Society. 2013 AHA/ACC/TOS guideline for the management of overweight and obesity in adults: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines; Obesity Society. 2013 AHA/ACC/TOS guideline for the management of overweight and obesity in adults: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines; Obesity Society. J Am Coll Cardiol 2014;63(25 Pt B):2985-3023.

Karason K, Molgaard H, Wikstrand J, Sjöström L. Heart rate variability in obesity and the effect of weight loss. Am J Cardiol 1999;83:1242-1247.

Kilavuori K, Toivonen I, Näsöri H, Leinonen H. Reversal of autonomic derangements by physical training in chronic heart failure assessed by heart rate variability. Eur Heart J 1995;16:490-495.

Levy WC, Cerqueira MD, Harp GD, Johannesen KA, Abrass IB, Schwartz RS, Stratton JR. Effect of endurance exercise training on heart rate vari-
ability at rest in healthy young and older men. Am J Cardiol 1998;82:1236-1241.
Lindmark S, Lönn L, Wiklund U, Tufvesson M, Olsson T, Eriksson JW. Dysregulation of the autonomic nervous system can be a link between visceral adiposity and insulin resistance. Obes Res 2005;13:717-728.
Lucini D, de Giacomi G, Tosi F, Malacarne M, Respizzi S, Pagani M. Altered cardiovascular autonomic regulation in overweight children engaged in regular physical activity. Heart 2013;99:376-381.
Minami J, Kawano Y, Ishimitsu T, Matsuoka H, Takishita S. Acute and chronic effects of a hypocaloric diet on 24-hour blood pressure, heart rate and heart-rate variability in mildly-to-moderately obese patients with essential hypertension. Clin Exp Hypertens 1999;21:1413-1427.
Nagai N, Moritani T. Effect of physical activity on autonomic nervous system function in lean and obese children. Int J Obes Relat Metab Disord 2004;28:27-33.
Oktay AA, Lavie CJ, Kokkinos PF, Parto P, Pandey A, Ventura HO. The interaction of cardiorespiratory fitness with obesity and the obesity paradox in cardiovascular disease. Prog Cardiovasc Dis 2017;60:30-44.
Pagani M, Lucini D. Autonomic dysregulation in essential hypertension: insight from heart rate and arterial pressure variability. Auton Neurosci 2001;90:76-82.
Piccirillo G, Vetta F, Viola E, Santagada E, Ronzoni S, Cacciafesta M, Marigliano V. Heart rate and blood pressure variability in obese normotensive subjects. Int J Obes Relat Metab Disord 1998;22:741-750.
Quilliott D, Böhme P, Zannad F, Ziegler O. Sympathetic-leptin relationship in obesity paradox: effect of weight loss. Metabolism 2008;57:555-562.
Rajendra Acharya U, Paul Joseph K, Kannathal N, Lim CM, Suri JS. Heart rate variability: a review. Med Biol Eng Comput 2006;44:1031-1051.
Rippe JM, Hess S. The role of physical activity in the prevention and management of obesity. J Am Diet Assoc 1998;98(10 Suppl 2):S31-38.
Rissanen P, Franssila-Kallunki A, Rissanen A. Cardiac parasympathetic activity is increased by weight loss in healthy obese women. Obes Res 2001;9:637-643.
Shaffer F, Ginsberg JP. An overview of heart rate variability metrics and norms. Front Public Health 2017;5:258.
Sheema UK, Malipatil BS. A cross-sectional study on effect of body mass index on the spectral analysis of heart rate variability. Natl J Physiol Pharmaco15;2015;5:250-252.
Sjoberg N, Brinkworth GD, Wycherley TP, Noakes M, Saint DA. Moderate weight loss improves heart rate variability in overweight and obese adults with type 2 diabetes. J Appl Physiol (1985) 2011;110:1060-1064.
Soares-Miranda L, Alves AJ, Vale S, Aires L, Santos R, Oliveira J, Mota J. Central fat influences cardiac autonomic function in obese and overweight girls. Pediatr Cardiol 2011;32:924-928.
Stein PK, Ehsani AA, Domitrovich PP, Kleiger RE, Rottman JN. Effect of exercise training on heart rate variability in healthy older adults. Am Heart J 1999;138(3 Pt 1):567-576.
Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology. Heart rate variability: standards of measurement, physiological interpretation and clinical use. Circulation 1996;93:1043-1065.
Tian Y, Huang C, He Z, Hong P, Zhao J. Autonomic function responses to training: correlation with body composition changes. Physiol Behav 2015;151:308-313.
Tulppo MP, Mäikallio TH, Takala TE, Seppänen T, Huikuri HV. Quantitative beat-to-beat analysis of heart rate dynamics during exercise. Am J Physiol 1996;271(1 Pt 2):H244-252.
Umetani K, Singer DH, McCratty R, Atkinson M. Twenty-four hour time domain heart rate variability and heart rate: relations to age and gender over nine decades. J Am Coll Cardiol 1998;31:593-601.
Vesterinen V, Häkkinen K, Hynynen E, Mikkola J, Hokka L, Nummela A. Heart rate variability in prediction of individual adaptation to endurance training in recreational endurance runners. Scand J Med Sci Sports 2013;23:171-180.
Wei M, Kampert JB, Barlow CE, Nichaman MZ, Gibbons LW, Paffenbarger RS Jr, Blair SN. Relationship between low cardiorespiratory fitness and mortality in normal-weight, overweight, and obese men. JAMA 1999;282:1547-1553.