

# Cardiopulmonary exercise testing for patients with anorexia nervosa: a case-control study

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1 **Title**

2 Cardiopulmonary exercise testing for patients with anorexia nervosa: A case control study

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24 commented on the draft.

1 **Abstract**

2

3 **Purpose** Patients with anorexia nervosa (AN) require appropriate nutrient therapy and physical  
4 activity management. Eating disorder treatment guidelines do not include safe, evidence-based  
5 intensity criteria for exercise. This study used cardiopulmonary exercise testing (CPX) to evaluate the  
6 exercise tolerance of patients with AN.

7 **Methods** CPX was performed with 14 female patients with AN admitted to a specialized eating  
8 disorder unit between 2015 and 2019. Their anaerobic threshold (AT) was determined by assessing  
9 their exercise tolerance using CPX and compared with 14 healthy controls (HC). The metabolic  
10 equivalents (AT-METS) were compared when AT was reached. We examined factors related to AT  
11 (AN-AT) in the AN group, including age, body mass index (BMI), previous lowest weight, minimum  
12 BMI, past duration of BMI <15, exercise history, and  $\Delta$ HR (heart rate at the AT - resting heart rate).

13 **Results** The AT of the AN group (BMI: 15.7 [Mean]  $\pm$  1.8 standard deviation [SD]) was significantly  
14 lower than that of the HC group (BMI: 19.7  $\pm$  1.8) (AN: 10.0  $\pm$  1.8 vs. HC: 15.2  $\pm$  3.0 ml/kg/min,  
15 P<0.001). AT-METS was also significantly lower in the AN group than in the HC group (AN: 2.9  $\pm$   
16 0.52 vs. HC: 4.4  $\pm$  0.91, P<0.001). AN-AT was highly influenced by  $\Delta$ HR.

17 **Conclusion** This study showed that AT and AT-METS were lower in patients with AN than in HC.  
18 Patients with AN should be prescribed light-intensity aerobic exercise, and the current findings may  
19 help develop future physical management guidelines for patients with AN.

20

21 **Level of evidence:** III: Evidence obtained from case-control analytic studies

22

23 **Keywords**

24 Anorexia nervosa, Cardiopulmonary exercise testing, Anaerobic threshold, Metabolic equivalents,  
25 Physical activity

26

## 1 **Introduction**

2 Anorexia nervosa (AN) is a severe mental disorder characterized by malnutrition. It has a high  
3 prevalence of comorbid psychiatric disorders, marked resistance to treatment, and high risk of death  
4 from physical complications [1,2]. AN affects almost all body systems, including the brain, heart, liver,  
5 gastrointestinal tract, bones, and muscles [3,4]. For these reasons, early nutritional recovery is  
6 necessary in the treatment of patients with AN [5]. Hyperactivity is one of the symptoms of AN.  
7 Starvation and hyperactivity lead to weight loss, and they are associated with physiological changes  
8 that promote compulsive behavior [6]. Many patients are at increased risk for a variety of serious  
9 medical complications, such as fractures, electrolyte imbalances or sudden death, that can be caused  
10 or exacerbated by hyperactivity [7].

11 Both appropriate nutrition therapy and physical activity management are necessary. Hyperactivity  
12 during refeeding in patients with AN is associated with increased energy demands to achieve weight  
13 gain, poorer clinical outcomes, longer hospital stays, and increased psychiatric complications [8]. In  
14 this regard, physical activity management refers to the teaching of exercise limitations and appropriate  
15 physical activity levels necessary to prevent further weight loss in patients with AN. The importance  
16 of nutritional therapy for patients with AN is well established [5]. On the other hand, the treatment  
17 guidelines for eating disorders do not include criteria for exercise intensity based on the specific  
18 exercise physiology traits of patients with AN with regard to safe levels of physical activity [9-13].  
19 Systematic reviews and proposed recommendations in the guidelines for exercise in the treatment of  
20 AN are also not evidence-based with regard to exercise physiology [14].

21 Cardiopulmonary exercise testing (CPX) can provide information on biological responses based on  
22 exercise intensity. We applied CPX to patients with AN to evaluate their exercise tolerance. Among  
23 the parameters obtained from CPX, the anaerobic threshold (AT) was used as an index of exercise  
24 tolerance. The AT is the upper limit of aerobic exercise intensity, and values below the AT indicate that  
25 exercise can be performed without the accumulation of lactic acid and without acidosis [15]. Thus, AT  
26 is a useful indicator of exercise tolerance, and exercise intensity below AT, i.e., the state of aerobic  
27 metabolism, is considered to be safe. Exercise intensity below AT is associated with less increase in  
28 sympathetic activity, a smaller degree of blood pressure elevation, and fewer arrhythmias.

29 The components of AT comprise a complex system. AT is a comprehensive metabolic index of  
30 ventilation (external respiration), circulation, and metabolism (internal respiration), and is defined by  
31 gas exchange, oxygen transport to skeletal muscle, and oxygen availability in skeletal muscle [16].  
32 Exercise performance requires an appropriate heart rate (HR) response during exercise, based on  
33 normal autonomic nervous system function [17].

34 CPX can provide data on AT and metabolic equivalents (AT-METS) when AT is reached during an  
35 exercise load. MET is a unit that expresses the intensity of physical activity relative to that expended  
36 by the body at rest. METS are used clinically to prescribe physical activities that a patient can safely  
37 perform [18]. AT-METS is the MET when AT is reached, and any activity above this value indicates  
38 anaerobic metabolism.

1 The purpose of this study was to evaluate the exercise tolerance of patients with AN compared to  
2 HC using CPX.

### 4 **Methods**

5 The participants were 14 patients with AN and 14 healthy controls (HC). All participants were female.  
6 The patients were admitted to a specialized eating disorder unit at Kyushu University Hospital between  
7 March 2015 and January 2019. Patients admitted to the eating disorders unit were those with AN who  
8 had low body weight and were scheduled to undergo treatment by a specialist (eating disorder  
9 therapist) for the purpose of regaining weight and receiving psychotherapy, especially cognitive-  
10 behavioral therapy. Patients with bulimia nervosa admitted for eating behavior stabilization were  
11 excluded from this study. In the final study cohort, we included patients with AN who provided  
12 informed consent for participation in the study. All participants were admitted for medical instability  
13 due to eating disorders. Six had anorexia nervosa restricting type and eight had the binge  
14 eating/purging type. AN was diagnosed by a specialized eating disorder therapist according to  
15 Diagnostic and Statistical Manual of Mental Disorders (DSM)-5 criteria [19]. Healthy adult females  
16 with no underlying medical conditions were recruited as HC. All participants were confirmed by their  
17 physicians to have no contraindications to CPX [20]: none had acute myocardial infarction, unstable  
18 angina, uncontrolled arrhythmias, symptomatic severe aortic stenosis, uncontrolled symptomatic heart  
19 failure, acute pulmonary infarction, acute myocarditis, acute pericarditis, acute aortic dissection, or  
20 uncommunicative mental illness. All participants were informed by the examiner about the purpose of  
21 CPX and the possibility of complications.

22 This study was approved by the Ethics Committee of Kyushu University (Ethical Approval Number;  
23 26-191, October 21, 2014.). All participants provided written informed consent before participating in  
24 the study.

25 Age, duration of illness (years), body mass index (BMI) ( $\text{kg}/\text{m}^2$ ), previous minimum BMI ( $\text{kg}/\text{m}^2$ ),  
26 past duration (years) of BMI  $<15$  ( $\text{kg}/\text{m}^2$ ), and exercise history were extracted from medical records.  
27 We defined exercise history as habit of exercising via extracurricular activities, athletic clubs, etc., for  
28 at least two years in the past.

#### 30 *Cardiopulmonary exercise testing*

31 Nutritional therapy was administered to the patients with AN after hospitalization, and all of them  
32 regained weight and experienced improvement in their physical condition. The patients with AN in  
33 this study were hospitalized for an average of 103.5 days and gained an average of 5.3 kg. CPX was  
34 performed prior to discharge. During CPX, none of the participants were suffering from severe  
35 hypotension or bradycardia, severe anemia, respiratory disturbances, electrolyte abnormalities, severe  
36 liver damage, hypoglycemia, or gait disturbance. Moreover, a physician checked the physical condition  
37 of all participants when performing CPX. Emergency equipment was always available and ready for  
38 use when CPX was performed. CPX was performed by the first author, who received training from a

1 CPX specialist.

2 The laboratory was set up in an environment with good lighting and ventilation, temperature of 20–  
3 25 °C, and humidity of 40–60%. CPX was performed using a continuous expiratory gas analyzer  
4 (AE310-S Aero monitor; Minato Medical Science Co., Ltd, Osaka, Japan). This device comprises a  
5 respiratory flow meter, oxygen analyzer, and carbon dioxide analyzer, with breath gas measured by the  
6 breath-by-breath method. Before each CPX, the flowmeter and gas analyzer were calibrated. The  
7 participant wore a face mask for exhaled gas analysis. A cycle ergometer (AERO BIKE 75XLIII;  
8 Konami Sports Life Co., Ltd, Kanagawa, Japan) was used as a load device during CPX implementation.  
9 During CPX, all participants were instructed to keep the cycle ergometer revolution constant at 60  
10 revolutions per minute. The CPX was performed using a ramp load protocol [21]. The ramp load  
11 intensity was set to 5 Watt/min<sup>-1</sup> for the patients with AN, and 20 Watt/min<sup>-1</sup> for the HC.

12 The procedure for the exercise loading was as follows: first, rest for 4 minutes, then warm up for 4  
13 minutes, following which exercise loading was performed. After discontinuing the exercise loading, a  
14 cool-down was performed for 4–6 minutes. Once AT was reached, the exercise load was stopped and  
15 CPX was terminated. Before and after CPX, all participants underwent measurements with regard to  
16 HR, blood pressure, and arterial oxygen saturation, and a 12-lead electrocardiogram (ECG) was  
17 performed. During CPX, the HR, blood pressure, and arterial oxygen saturation were monitored every  
18 minute. Further, the examiner confirmed subjective symptoms such as shortness of breath, chest pain,  
19 malaise, and lower limb fatigue. The discontinuation criteria for CPX were based on subjective and  
20 objective symptoms [20]. All participants could stop exercising at any time. None of the participants  
21 experienced complications during or after CPX.

22

### 23 *CPX parameters*

24 In the breath gas analysis of CPX, oxygen uptake ( $\dot{V}O_2$ ) (ml/min), carbon dioxide output ( $\dot{V}CO_2$ )  
25 (ml/min), respiratory rate (RR) (f/min), and minute ventilation ( $\dot{V}E$ ) (ml/min) were measured. AT  
26 was determined using the trend method and confirmed by the V-Slope method. METS (AT-METS)  
27 was calculated at the time AT was achieved. Since 1 METS corresponds to an oxygen uptake of  
28 approximately 3.5 ml/min/kg, AT-METS was calculated by dividing the oxygen uptake at AT by 3.5.

29

### 30 *Statistical analysis*

31 A Shapiro–Wilk test was performed to assess normality. Continuous variables are represented by  
32 mean  $\pm$  standard deviation (SD) or median (range: minimum–maximum) according to the distribution  
33 of the variables.

34 First, multiple regression analysis by the stepwise method was performed with the AT of both groups  
35 as the dependent variable and age, BMI (kg/m<sup>2</sup>), group (AN or HC), and  $\Delta$ HR as the independent  
36 variables. For age, BMI, and  $\Delta$ HR, the normality of the variables was confirmed in advance by the  
37 Shapiro-Wilk test, and the shape of the distribution was confirmed by histogram analysis. Since none  
38 of these variables deviated significantly from the normal distribution or had a biased frequency, they

1 were not converted into dummy variables or changed. The presence or absence of exercise history was  
2 converted into a dummy variable, with exercise history = “1” and the absence of exercise history =  
3 “0”. Furthermore, we developed a correlation matrix table, but since there were no variables with  $|r| >$   
4 0.9, we targeted all variables.

5 The mean differences in AT (ml/kg/min), AT-METS, and  $\Delta$ HR (bpm) between the AN and HC groups  
6 were compared by an unpaired t-test. For the unpaired t-test, power was calculated for each variable.  
7 We defined  $\Delta$ HR (bpm) as (HR at the AT - resting HR) (bpm). The resting HR is the HR at the start of  
8 CPX.

9 Multiple regression analysis by the stepwise method was performed with the AT of the AN group  
10 (AN-AT) as the dependent variable and age, BMI (kg/m<sup>2</sup>), record low BMI (kg/m<sup>2</sup>), the period (years)  
11 with past BMI <15 (kg/m<sup>2</sup>), the presence or absence of exercise history, and  $\Delta$ HR as the independent  
12 variables. The total duration during which BMI was <15 in the past (period with past BMI <15) among  
13 the participants was  $1.5 \pm 1.3$  years. Patients with AN with BMI <15 (kg/m<sup>2</sup>) were classified as the  
14 most severe cases [19].

15 In all analyses, a difference was considered significant at  $P < 0.05$ . SPSS (IBM) ver. 28 software was  
16 used for all statistical analyses.

## 18 Results

19 The clinical background of the participants is shown in **Table 1**. No difference in age was found  
20 between the two groups. The weight (kg), BMI (kg/m<sup>2</sup>), and body surface area (BSA[m<sup>2</sup>]) were  
21 significantly lower in the AN group than in the HC group ( $P < 0.001$ ). Three participants in the AN  
22 group had a history of exercise. All participants were able to perform CPX with no adverse events.  
23 The data are expressed as means  $\pm$  standard deviation.

24 The parameters obtained in CPX are presented in Table 2. The AT of the AN group was significantly  
25 lower than that of the HC group (AN:  $10.0 \pm 1.8$  vs. HC:  $15.2 \pm 3.0$  ml/kg/min,  $P < .001$ ) (**Fig. 1**). The  
26 AT-METS of the AN group was also significantly lower than that of the HC group (AN:  $2.9 \pm 0.52$  vs.  
27 HC:  $4.4 \pm 0.91$ ,  $P < .001$ ) (**Fig. 2**). The powers in the unpaired t-test were 1.00 (AT), 0.99 (AT-METS),  
28 and 0.99 (HR at the AT), respectively.

29 No significant difference was found between the resting HR of the AN group and that of the HC  
30 group (AN:  $72.6 \pm 11.1$  vs. HC:  $75.1 \pm 8.7$  bpm,  $P = .524$ ). Contrastingly, the HR at the AT of the AN  
31 group was significantly lower than that of the HC group (AN:  $94.6 \pm 12.6$  vs. HC:  $125.1 \pm 17.0$   
32 ml/kg/min,  $P < .001$ ) (**Table 2**), and the  $\Delta$ HR of the AN group was significantly lower than that of the  
33 HC (AN:  $20.6 \pm 8.0$  vs. HC:  $50.1 \pm 18.8$  bpm,  $P < .001$ ).

34 The results of the multiple regression analysis of both groups are shown in Table 3. Age and BMI  
35 were excluded from the independent variables. The results of the analysis of variance (ANOVA) were  
36 significant, and the coefficient of determination ( $R^2$ ) was 0.650, indicating a highly adequate fit. The  
37 standard partial regression coefficient ( $\beta$ ) of  $\Delta$ HR and group were .449 and .420, respectively,  
38 indicating that  $\Delta$ HR and the difference of groups make a moderate contribution to AN-AT. The Durbin–

1 Watson ratio of 1.814 was acceptable, and there were no outliers where the predicted value exceeded  
2 3 SD of the measured value.

3 The results of multiple regression analysis of only the AN group are shown as follows. Age, BMI,  
4 previous minimum BMI, past duration of BMI <15, and exercise history were excluded from the  
5 independent variables. The result of ANOVA was significant, and the coefficient of determination ( $R^2$ )  
6 was 0.605, indicating a highly adequate fit. The standard partial regression coefficient ( $\beta$ ) of  $\Delta$ HR  
7 was .778, indicating that  $\Delta$ HR makes a significant contribution to AN-AT. The Durbin–Watson ratio  
8 of 2.503 was acceptable, and there were no outliers where the predicted value exceeded 3 SD of the  
9 measured value.

## 11 Discussion

12 In this study, the exercise tolerance of patients with AN was lower than that of HC, in line with the  
13 findings of a previous study by Biadi et al [22]. The exercise tolerance of patients with AN in this study  
14 was moderate-to-severe according to the Weber–Janicki classification [23], which cannot be explained  
15 by clinical factors such as BMI, duration of disease, or previous lowest weight. Age, previous  
16 minimum BMI, past duration of BMI <15, and history of exercise were not explanatory factors  
17 contributing to the lower AT of our patients with AN. It is particularly interesting that the decreased  
18 AT of our patients was not explained by the various clinical measures we estimated. It would be useful  
19 for clinicians to be able to estimate the exercise tolerance of patients with AN using more readily  
20 available clinical data without performing CPX. Further investigation of factors such as body  
21 composition and autonomic function test results is warranted.

23 For patients with AN, a low BMI has been shown to be associated with a higher risk of underweight-  
24 related death [24]. A BMI <13 kg/m<sup>2</sup> has also been proposed as a cutoff value for poor prognosis [24].  
25 Low BMI (BMI <13–14 kg/m<sup>2</sup>) is associated with a faster rate of decrease in BMI and an increase in  
26 the number of patients who require emergency hospitalization due to difficulty in walking or impaired  
27 consciousness [25]. A BMI of 13–14 kg/m<sup>2</sup> represents the boundary at which body composition  
28 changes significantly [25], and as the BMI falls below this value, the source of energy changes from  
29 fat to protein [25, 26]. During starvation, a BMI of 13 kg/m<sup>2</sup> may be the tipping point at which the  
30 homeostatic mechanism breaks down [26]. Thus, anaerobic metabolism is presumed to be very  
31 dangerous for patients with AN. Therefore, in this study, CPX was terminated when AT was reached.  
32 When comparing exercise intensity above and below AT, that below AT has the following  
33 characteristics: no sustained increase in lactate concentration, the ability to perform the exercise for a  
34 longer period of time without fatigue, less increase in sympathetic nerve activity, a smaller increase in  
35 blood pressure, and reduced incidence of arrhythmia. Among other factors, a metabolic state with  
36 reduced incidence of arrhythmia is physically safer for patients with AN, who often have coexisting  
37 electrolyte abnormalities. Elevated lactate concentration constitutes a risk among patients with AN,  
38 since it implies that the glycolytic system is facilitated. In other words, greater utilization of



1 carbohydrates is an additional risk for patients with AN who present with low body weight and low  
2 nutrition.

3  
4 The AT-METS of our patients with AN was  $2.9 \pm 0.52$ . Physical activity of 2.8 METS is the  
5 equivalent of playing with children or caring for animals. Further, 3.0 METS represents activities like  
6 normal walking or indoor cleaning [27,28]. The AT-METS of HC in this study was  $4.4 \pm 0.91$ . Physical  
7 activity of 4.0 METS is equivalent to activities like jogging or riding a bicycle. Finally, 4.5 METS  
8 represents activities like sapling planting and garden weeding [27,28].

9  
10 Patients with AN often have a blunted sympathetic response to maximal exercise, i.e., a variable  
11 response insufficiency [29]. Chronotropic incompetence (CI) is defined as an inability to raise the HR  
12 appropriately and to match cardiac output to metabolic demands during activity. In the present study,  
13 the increase in HR from the start of exercise to AT was suppressed in our patients with AN. This study  
14 suggests that the AN group was exercising below 50% of their age-adjusted maximum HR at the time  
15 that their AT was reached. Further, the  $\Delta$ HR of our patient group was shown to have a significant effect  
16 on AT. CI is associated with exercise intolerance [30]. The autonomic abnormalities of patients with  
17 AN may persist even after weight regain, and more careful management of physical activity is needed  
18 for patients with AN with CI [31].

19  
20 In this study, the respiratory rate at AT attainment was not significantly different between the two  
21 groups, but the minute ventilation rate was significantly lower in the AN group than in the HC group.  
22 This may be attributed to the AN group's body size, i.e., emaciation. It is suggested that the oxygen  
23 content of the blood may not increase sufficiently in patients with AN, with decreased minute  
24 ventilation comprising one factor. Reduced systolic ventricular function is not a major factor  
25 contributing to the reduced exercise capacity of patients with AN [32]. It has been reported that older  
26 patients have decreased oxygen availability in skeletal muscle [33] and decreased lean mass [34], both  
27 of which are determinants of decreased exercise tolerance. However, the association between AT,  
28 muscle mass, and skeletal muscle strength of patients with AN remains unclear. Another factor that  
29 may define AT is exercise habits [35], but this study did not show an association between exercise and  
30 AT in our patients with AN. It is particularly interesting that the decreased AT in AN was not explained  
31 by the various clinical measures we estimated.

32  
33 It is not advisable to manage the range of activity of patients with AN only with bed rest. In a study  
34 by Ibrahim et al, bed rest was not supported for the inpatient treatment of patients with AN [36]. Other  
35 studies have shown that physical activity has a positive impact on AN treatment. Exercise and physical  
36 therapy can help patients with AN recover from their physical and mental problems [37]. Maintaining  
37 safe physical activity during the refeeding period for patients with AN is beneficial for restoring body  
38 composition, maintaining bone mineral density, and mental status [8]. Furthermore, previous studies

1 have shown that exercise with nutritional support improves the quality of life and psychological well-  
2 being [38]. However, there is no consensus or recommendation regarding how physical activity should  
3 be managed in patients with AN, and implementation varies among specialized centers [8]. One  
4 systematic review of exercise in patients with eating disorders included 18 studies [14], only one of  
5 which was a meta-analysis focusing on AN treatment. The present study focuses on patients with AN  
6 who were underweight but not because of eating disorders alone. The findings of our study, which  
7 assessed the exercise physiology of patients with AN, provide an important basis for guiding the  
8 physical activity of patients with AN.

9  
10 A previous study showed that the BMI percentile was independently associated with the exercise  
11 endurance of adolescents with AN [32]. A BMI <14 (kg/m<sup>2</sup>) was noted as an indicator of high  
12 medical risk in patients with AN [39], and in some institutions, patients with a BMI <15 (kg/m<sup>2</sup>) are  
13 restricted from physical activity. In other centers, patients with a BMI of <18.5 (kg/m<sup>2</sup>) are  
14 prohibited from physical activity [40]. A BMI of ≤13–14 (kg/m<sup>2</sup>) has been reported to be a marker  
15 for consciousness impairment and gait disturbances [26]. However, the BMI of our patients with AN  
16 was not related to their AT level, indicating that the physical activity of patients with AN should not  
17 be defined and managed by BMI alone.

18  
19 In conclusion, the exercise tolerance of patients with AN was lower than that of HC. AN-AT was  
20 highly influenced by  $\Delta$ HR, but not influenced by age, BMI, previous minimum BMI, past duration of  
21 BMI <15, or exercise history. Patients with AN should be prescribed light-intensity aerobic exercise,  
22 and the current findings may help develop future physical management guidelines for patients with  
23 AN.

### 24 25 **Strength and limitations**

26 To our knowledge, this study is the first to demonstrate the AT-METS level of patients with AN  
27 based on CPX.

28 The study has several limitations. First, the sample size was small. AT and AT-METS each showed  
29 significant differences between the two groups, but both powers were too high. With a larger sample  
30 size, the AT and AT-METS of patients with AN might have been better defined. Second, the study was  
31 limited to Japanese female patients with AN and did not include older patients or HC. Third, our  
32 definition of exercise history is not supported by sufficient scientific evidence. Fourth, during CPX,  
33 the exercise load applied was limited to an AT level deemed to be safe for patients with AN. Therefore,  
34 peak  $\dot{V}O_2$  was not obtained, and the full exercise capacity of these patients was not assessed. Fifth, we  
35 did not measure cardiac function, autonomic nervous system activity, respiratory function, or skeletal  
36 muscle function during exercise; thus, factors that might contribute to poor exercise tolerance by  
37 patients with AN could not be identified. Finally, CPX is not readily available in all clinical settings.  
38 In the future, we will attempt to include more cases with more data on body composition and detailed

1 exercise history to help clinicians assess exercise tolerance without the use of CPX.

### 3 **What is already known on this subject?**

4 Exercise tolerance is decreased in patients with AN, and chronotropic incompetence is known to be  
5 a factor in the decreased exercise tolerance of patients with AN.

### 7 **What this study adds?**

8 The results of this study show that AT-METS in patients with AN is lower than that in HC.

### 10 **Declarations**

### 12 **Funding**

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15 Health and Medicine (20A 3001) and Grants-in-Aid for Scientific Research from the Japanese  
16 Ministry of Health, Labor, and Welfare (17K09340).

### 18 **Conflicts of interest**

19 The authors have no conflicts of interest or competing interests to declare. The authors have no  
20 relevant financial or non-financial interests to disclose.

### 22 **Availability of data and material**

23 The datasets generated and analyzed during the current study are available from the corresponding  
24 author on reasonable request.

### 26 **Code availability**

27 Not applicable.

### 29 **Author contributions**

30 Kawai Keisuke and Sudo Nobuyuki contributed to the study conception and design. Material  
31 preparation and data collection were performed by Yamashita Makoto, Toda Kenta, Aso Suzuyama  
32 Chie, Suematsu Takafumi, Yokoyama Hiroaki, Hata Tomokazu, and Takakura Shu. Analysis was  
33 performed by Yamashita Makoto and Kawai Keisuke. The first draft of the manuscript was written  
34 by Yamashita Makoto. All authors have read and approved the final manuscript.

### 36 **Ethics approval**

37 This study was approved by the Ethics Committee of Kyushu University (Ethical Approval Number  
38 26-191) and the National Center for Global Health and Medicine (Ethical Approval Number NGM-G-

1 003071-00).

2

3 **Consent to participate**

4 All participants provided written informed consent prior to participation in the study.

5

6 **Consent for publication**

7 All authors have given their consent for submission of the manuscript.

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1 **Tables**

2

3 Table 1. Clinical characteristics of the study population

	<b>Patients with AN (n = 14)</b>	<b>Healthy controls (n = 14)</b>	<b>P value</b>
Age (years)	25.7 ± 8.5	26.5 (24–47)	.077 <sup>b</sup>
Height (cm)	156.8 (152.5–172.5)	160.4 ± 4.8	.227 <sup>b</sup>
Body weight (kg)	39.4 ± 5.2	50.6 ± 4.1	<.001 <sup>a</sup>
BMI (kg/m <sup>2</sup> )	15.7 ± 1.8	19.7 ± 1.8	<.001 <sup>a</sup>
Body surface area (m <sup>2</sup> )	1.3 ± 0.10	1.5 ± 0.07	<.001 <sup>a</sup>

4 Data are expressed as mean ± SD or median (range: min.–max.) depending on data distribution.

5 P values were calculated with an unpaired t test (a) and Mann–Whitney U test (b).

6 Abbreviations: AN, anorexia nervosa; BMI, body mass index.

7

1 Table 2. Parameters at the time AT was achieved in CPX

	<b>Patients with AN (n = 14)</b>	<b>Healthy controls (n = 14)</b>	<b>P value</b>
Oxygen uptake (ml/min)	398.9 ± 90.7	788.8 ± 178.3	<.001 <sup>a</sup>
Carbon dioxide output (ml/min)	414.5 ± 98.2	750.2 ± 162.3	<.001 <sup>a</sup>
Gas exchange ratio	1.04 ± 0.06	0.95 ± 0.04	<.001 <sup>a</sup>
Respiratory rate (f/min)	23.1 (14.3–35.4)	24.9 ± 4.8	.23 <sup>b</sup>
Minute ventilation (l/min)	14.9 ± 3.0	23.4 ± 4.6	<.001 <sup>a</sup>
Heart rate (bpm)	94.6 ± 12.6	125.1 ± 17.0	<.001 <sup>a</sup>

2 Data are expressed as mean ± SD or median (range: min.–max.) depending on data distribution.

3 P values were calculated with an unpaired t test (a) and Mann–Whitney U test (b).

4 Abbreviations: AT, anaerobic threshold; CPX, cardiopulmonary exercise testing.

5

1 Table 3. Variables predictive of AT in all of the participants

	<b>B</b>	<b>β</b>	<b>P</b>	<b>95% CI</b>	
				Lower limit	Upper limit
Const.	7.962		<.001	6.076	9.848
ΔHR	.078	.449	.015	.017	.140
group	2.968	.420	.022	.464	5.473

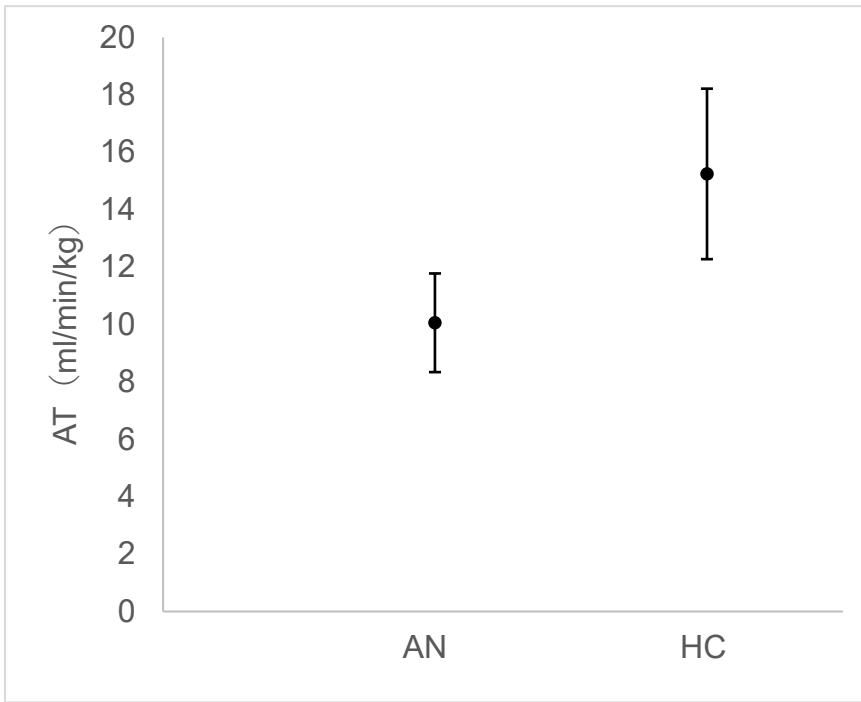
2  $R^2 = .650$ , ANOVA:  $P < .001$

3 Abbreviations: AT, anaerobic threshold; B, partial regression coefficient; β, standardized partial  
 4 regression coefficient; P, significance probability; CI, confidence interval; Const., constant; HR, heart  
 5 rate.

1 Figures

2

3 **Fig. 1**

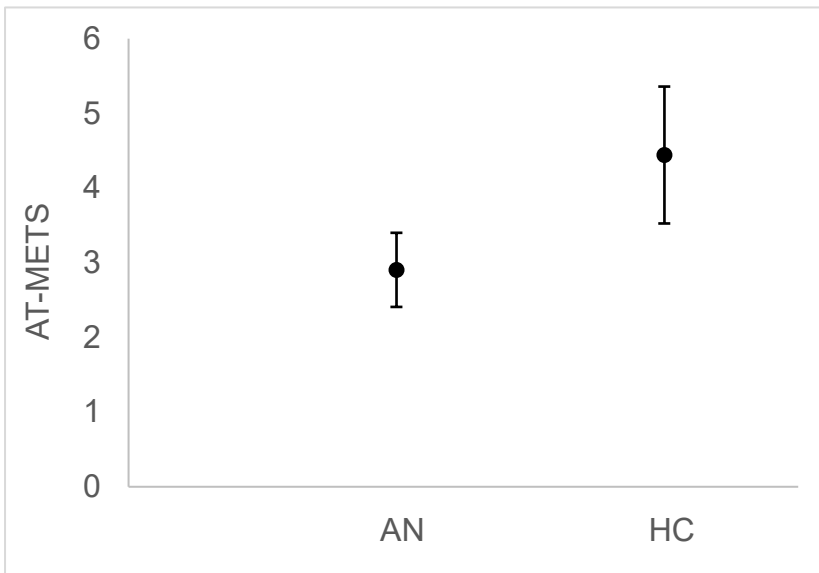


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**Fig. 2**



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1 **Figure captions**

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3 **Fig. 1** AT of the AN and HC groups

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5 Dots and error bars express means and 95% confidence intervals.

6 Abbreviations: AT, anaerobic threshold; AN, anorexia nervosa; HC, healthy control.

7

8 **Fig. 2** AT-METS of the AN and HC groups

9

10 Dots and error bars express means and 95% confidence intervals.

11 Abbreviations: AT, anaerobic threshold; METS, metabolic equivalents; AN, anorexia nervosa; HC,  
12 healthy control.