Effects of 5-aminolevulinic acid supplementation on home-based walking training achievement in middle-aged depressive women: randomized, double-blind, crossover pilot study

Hiroshi Suzuki1,3,4, Shizue Masuki1,2, Akiyo Morikawa4, Yu Ogawa3, Yoshi-ichiro Kamijo1,2, Kiwamu Takahashi5, Motowo Nakajima5 & Hiroshi Nose1,2

Depressive patients often experience difficulty in performing exercise due to physical and psychological barriers. We examined the effects of 5-aminolevulinic acid (ALA) with sodium ferrous citrate (SFC) supplementation during home-based walking training in middle-aged depressive women. Nine outpatients [53 ± 8 (SD) yr] with major depressive disorder participated in the pilot study with randomized, placebo-controlled, double-blind crossover design. They underwent two trials for 7 days, each performing interval walking training (IWT) with ALA + SFC (ALA + SFC) or placebo supplement intake (PLC) intermittently with > a 10-day washout period. For the first 6 days of each trial, exercise intensity for IWT was measured by accelerometry. Before and after each trial, subjects underwent a graded cycling test, and lactate concentration in plasma ([Lac−])p, oxygen consumption rate (VO2), and carbon dioxide production rate (VCO2) were measured with depression severity by the Montgomery–Åsberg Depression Rating Scale (MADRS). We found that the increases in [Lac−]p, VO2 and VCO2 during the test were attenuated only in ALA + SFC ([before vs. after] × workload; all, P < 0.01), accompanied by increased training days, impulse, and time at fast walking during IWT (all, P < 0.05) with decreased MADRS-score (P = 0.001). Thus, ALA + SFC supplementation increased IWT achievement to improve depressive symptoms in middle-aged women.

Recently, the number of depressive patients has been increasing in Japan; the life-time prevalence is 6.7%, and women are affected twice as frequently as men1. Some depressed people feel unhappy and sad, while others have difficulty doing anything: going out, dressing, face-washing, and taking a bath2. One of their primary symptoms is fatigability, resulting in physical inactivity, reducing physical fitness, which would contribute to lifestyle-related diseases, including depression, falling into a viscous cycle3,4.

To prevent this, habitual exercise with moderate or higher intensity for ~30 min per day for 3–4 days per week has been recommended5–10. However, it may be difficult for depressive patients to perform exercise at the desired intensity, frequency and duration due to physical and psychological barriers11–14.

On the other hand, Masuki et al.15 recently examined the effects of oral supplementation of 5-aminolevulinic acid (ALA) + sodium ferrous citrate (SFC), a precursor of haeme that composes complex IV in the mitochondrial electron transport chain (ETC)16,17, on the voluntary achievement of high-intensity interval walking training (IWT) for 6 days in older women without depression and suggested that the achievement of the fast walking training time was enhanced by 69% in the experimental trial compared with the placebo trial. IWT is a
home-based walking training programme that repeats more than 5 sets of fast and slow walking for 3 min each per day, which is equivalent to more than 70% and 40% of peak aerobic capacity (V\textsuperscript{O2 peak}), respectively\textsuperscript{18–20}.

Regarding the mechanisms, since increases in oxygen consumption rate (\(V\textsuperscript{O2}\)), carbon dioxide production rate (\(V\textsuperscript{CO2}\)) and plasma lactate concentration ([Lac\textsuperscript{−}]) were reduced at a given exercise intensity during the graded cycling exercise in the experimental trial but not in the placebo trial after the intervention, these findings suggest that the higher achievement of IWT was attained by a reduced increase in [Lac\textsuperscript{−}] with improved oxygen utilization efficiency to produce adenosine-tri-phosphate (ATP) by activating ETC to improve subjective feeling against exercise stress.

Accordingly, we hypothesized that ALA\textsuperscript{+}SFC supplementation would increase the achievement of IWT by lowering barriers to higher intensity exercise in depressive patients and thereby improving their symptoms.

### Results

#### Subject characteristics.

Subject characteristics, clinical history of depression, and current medications are shown in Table 1. As shown in Table 2, body weight, blood pressure at rest, peak heart rate (HR\textsubscript{peak}), and VO\textsubscript{2peak} remained unchanged after the supplement intake in both trials (all, \(P > 0.08\)) (see Fig. 1 for experimental protocol). Resting heart rate (HR\textsubscript{rest}) decreased significantly after the ALA\textsuperscript{+}SFC supplement intake (ALA\textsuperscript{+}SFC trial) (\(P = 0.0014\)) but not after the placebo supplement intake (PLC trial) (\(P = 0.36\)). Haemoglobin concentration ([Hb]) decreased significantly in both trials after the supplement intake period (\(P < 0.05\)), with no significant interactive effects of (trials x before vs. after) (\(P > 0.7\)).

### Table 1. Physical characteristics, clinical history, and current psychological drug used for therapy in individual subjects. BMI, body mass index.

| Subject No. | Age, yr | Height, cm | BMI, kg/m\(^2\) | Clinical history of depression, mo | Psychotropic drugs                      |
|-------------|---------|------------|----------------|---------------------------------|---------------------------------------|
| 1           | 58      | 159.9      | 16.8           | 31                              | Ethyl loflazepate                      |
| 2           | 42      | 151.4      | 24.0           | 36                              | Paroxetine, Ethyl loflazepate, Rilmazafone |
| 3           | 47      | 155.1      | 30.5           | 55                              | Paroxetine, Trazodone, Bromazepam, Zopiclone |
| 4           | 61      | 159.7      | 24.0           | 4                               | Ethyl loflazepate                      |
| 5           | 65      | 154.1      | 16.8           | 24                              | Ethyl loflazepate                      |
| 6           | 52      | 154.8      | 29.4           | 48                              | Fluvoxamine, Trazodone, Bromazepam, Zolpidem |
| 7           | 42      | 166.5      | 37.2           | 36                              | Paroxetine, Mirtazapine, Trazodone, Sulpiride, Ethyl loflazepate, Zolpidem |
| 8           | 55      | 158.1      | 22.9           | 54                              | Sertraline, Zolpidem                   |
| 9           | 56      | 154.2      | 23.3           | 30                              | Duloxetine, Ethyl loflazepate, Lorazepam, Climazepam, Pregabaline, Brotizolam |
| Mean \(\pm\) SD | 53 \(\pm\) 8 | 157.1 \(\pm\) 4.5 | 25.0 \(\pm\) 6.5 | 35 \(\pm\) 16 | |

| Subject No. | Age, yr | Height, cm | BMI, kg/m\(^2\) | Clinical history of depression, mo | Psychotropic drugs                      |
|-------------|---------|------------|----------------|---------------------------------|---------------------------------------|

### Table 2. Physical characteristics of subjects. Values are the means \(\pm\) SE for 9 subjects in each trial. PLC, placebo intake condition; ALA\textsuperscript{+}SFC, 5-aminolevulinic acid + sodium ferrous citrate intake condition; SBP\textsubscript{rest} and DBP\textsubscript{rest}, resting systolic and diastolic blood pressure, respectively; HR\textsubscript{rest}, resting heart rate; HR\textsubscript{peak}, peak heart rate at VO\textsubscript{2peak}; VO\textsubscript{2peak}, peak oxygen consumption rate during the graded cycling test; [Hb], haemoglobin concentration in blood. * Compared with before supplement intake, \(P < 0.05\) and \(P < 0.01\), respectively.

|                | PLC Before | PLC After | ALA + SFC Before | ALA + SFC After |
|----------------|------------|-----------|------------------|----------------|
| Body weight, kg | 61.9 \(\pm\) 6.2 | 61.7 \(\pm\) 6.1 | 62.0 \(\pm\) 6.3 | 62.1 \(\pm\) 6.2 |
| SBP\textsubscript{rest}, mmHg | 116 \(\pm\) 3 | 123 \(\pm\) 4 | 120 \(\pm\) 2 | 116 \(\pm\) 4 |
| DBP\textsubscript{rest}, mmHg | 80 \(\pm\) 5 | 78 \(\pm\) 4 | 74 \(\pm\) 5 | 77 \(\pm\) 4 |
| HR\textsubscript{rest}, beats/min | 76 \(\pm\) 3 | 78 \(\pm\) 4 | 78 \(\pm\) 3 | 73 \(\pm\) 3* |
| HR\textsubscript{peak}, beats/min | 153 \(\pm\) 2 | 156 \(\pm\) 2 | 157 \(\pm\) 3 | 156 \(\pm\) 2 |
| VO\textsubscript{2peak}, l/min | 1.53 \(\pm\) 0.09 | 1.58 \(\pm\) 0.10 | 1.54 \(\pm\) 0.09 | 1.56 \(\pm\) 0.09* |
| [Hb], g/dl | 12.6 \(\pm\) 0.2 | 12.3 \(\pm\) 0.2* | 12.7 \(\pm\) 0.2 | 12.4 \(\pm\) 0.4* |

As shown in Fig. 2A, although VO\textsubscript{2}, VCO\textsubscript{2}, and [Lac\textsuperscript{−}] increased similarly as the intensity increased during the graded cycling test before the supplement
intake period in both trials, their increases were significantly attenuated in the ALA + SFC trial after the period with significant interactive effects of [(before vs. after) × time] (P = 0.0045, 0.0047, and 0.0068; 1-β = 0.978, 0.978, and 0.906, respectively) but not in the PLC trial (all, P > 0.2). Comparing the results before and after the period, [Lac]p decreased significantly at every intensity ≥45 W (P = 0.02, 1-β = 0.71) in the ALA + SFC trial but not in the PLC (P > 0.7). There was no significant change in ventilation volume (VE) response before and after the period (P > 0.4).

Figure 1. Experimental protocol. The study was conducted using a randomized, placebo-controlled, double-blind crossover design. Ex, graded cycling exercise test and depression severity assessment; IWT, high-intensity interval walking training. In each instance of supplement intake indicated by arrows, subjects ingested 250 mg of either 5-aminolevulinic acid (ALA) + sodium ferrous citrate (SFC) in the ALA + SFC trial or placebo supplement in the PLC trial at breakfast and at dinner (250 mg × 2 = 500 mg/day). See Supplemental Table S1 for details on the supplement compositions.

Figure 2. (A) Oxygen consumption rate (VO2), carbon dioxide production rate (VCO2), and plasma lactate concentration ([Lac]p) responses over intensities during the graded cycling exercise test in the PLC trial (left) and in the ALA + SFC trial (right). The average value per minute in VO2 and VCO2, and the values at every intensity of [Lac]p from rest to the highest workload of 75 W, at which all subjects could maintain the rhythm of pedalling at 60 cycles/min. Open symbols, before a supplement intake period; solid symbols, after a supplement intake period. Values are the means ± SE for 9 subjects. *P < 0.05 vs. before the supplement intake period. (B) The average changes in VO2, VCO2, and [Lac]p over intensities ≥45 W during graded cycling exercise after a supplement intake period. Values are the means ± SE for 9 subjects. *P < 0.05 and **P < 0.01 between the PLC and ALA + SFC trials.

Figure 2B shows the average changes in VO2, VCO2, and [Lac]p over intensities ≥45 W during graded cycling test after the supplement intake period; these values were determined by subtracting the pre-supplementation values from the post-supplementation values at every minute (VO2, VCO2) and every intensity ([Lac]p) ≥45 W in each subject, averaging them over the intensities, and presenting the mean and SE values.
for 9 subjects in each supplement trial. There were significant differences in the changes in \( \dot{V}O_2 \), \( \dot{V}CO_2 \), and \([\text{Lac}^-]\) between trials (all, \( P < 0.03, 1-\beta > 0.63 \)).

**Training achievement during the supplement intake period.** The training days (A), training impulse (B), and training time (C) during the supplement intake period (days 1–6) are shown in group mean (Fig. 3) and individual values (Fig. 4). As shown in Fig. 3, training days were 33% greater in the ALA + SFC trial than the PLC trial (\( P = 0.035, 1-\beta = 0.602 \)). The impulses for total and fast walking were both 46% higher in the ALA + SFC trial.
Table 3. Crossover analysis. \( \Delta V_O_2, \Delta V_C O_2, \Delta [Lac^-]_p, \) and \( \Delta M A D R S, \) changes in the average values of oxygen consumption rate \( (V_O_2)\), carbon dioxide production rate \( (V_C O_2)\), and lactate concentration in plasma \( ([Lac^-]_p)\) above 45 W during the graded cycling and in the Montgomery–Åsberg Depression Rating Scale \( (M A D R S)\) after supplement intake compared with before supplement intake. S1, PLC-(ALA + SFC) sequence; S2, (ALA + SFC)-PLC sequence. *S1 vs. S2 for each effect. Values are the means ± SE for 4 subjects for S1 and 5 subjects for S2. We tested the effects of 3 factors: carryover (physiological and other effects of the first supplement period are still present when the subject enters the second supplement period), period (the effect of stimulation order was present in PLC-(ALA + SFC) sequence group vs. (ALA + SFC)-PLC sequence group), and supplement effects on the average changes in \( V_O_2, V_C O_2, \) and \([Lac^-]_p\) ≥45 W during the graded cycling test (Fig. 2B). The analysis was performed by the method reported by Chow and Liu28. Similarly, the analysis was also performed to test the effects of 3 factors on training days, training impulse and time for fast walking, and the MADRS score with no carryover or period effects.

Figure 5. Montgomery–Åsberg Depression Rating Scale (MADRS) sum score in the PLC trial (left) and ALA + SFC trial (right), before and after a supplement intake period. Values are the means ± SE for 9 subjects. **P < 0.01 vs. before the supplement intake period.

than the PLC trial \( (P = 0.016 and 0.014, 1-\beta = 0.765 and 0.794, \) respectively). The training times for total and fast walking were 47% and 46% higher, respectively, in the ALA + SFC trial than the PLC trial \( (P = 0.022 and 0.009, 1-\beta = 0.703 and 0.859, \) respectively).

MADRS. As shown in Fig. 5, the Montgomery–Åsberg Depression Rating Scale (MADRS)21 score decreased significantly in the ALA + SFC trial \( (P = 0.0013, 1-\beta = 0.992)\) but not in the PLC trial \( (P > 0.1)\).

Crossover analysis. Table 3 shows the results of crossover analysis. We found significant supplement effects on the average changes in \( V_O_2, V_C O_2, \) and \([Lac^-]_p\) over the intensities ≥45 W, and training achievements, training days, impulse and time for fast walking, and the MADRS score with no carryover or period effects22.
Discussion

To our knowledge, this is the first study to investigate the effects of nutritional supplements on [Lac−]p during exercise and home-based walking training achievement in depressive patients. The major findings of this study are 1) the increases in [Lac−]p, VO2 and VCO2 during the graded cycling test were significantly suppressed in the ALA + SFC trial compared with in the PLC trial; 2) training days, training impulse, and time for fast walking significantly increased in the ALA + SFC trial compared with the PLC trial; and 3) the MADRS score was significantly decreased only in the ALA + SFC trial in middle-aged depressive women.

As shown in Fig. 2, increases in VO2, VCO2, and [Lac−]p during the graded cycling test were attenuated in the ALA + SFC trial but not in the PLC trial with no carryover or period effects (Table 3). Masuiki et al.15 reported similar results as in the present study in older women aged ~65 years old with no depression who had performed IWT for >12 months before participating in the study, assuming that their respiratory and [Lac−]p responses to graded cycling exercise had reached a steady state. They suggested that the increases in VO2, VCO2, and [Lac−]p during the graded cycling test were attenuated in the ALA + SFC trial. Regarding the mechanisms, they suggested that ALA + SFC supplementation improves mitochondrial functions to recover the age-associated decrease in transient O2 utilization rates for aerobic ATP production23,24, as well as exercise efficiency determined as work per the total metabolic cost of exercise, to reduce the O2 deficit at the onset of exercise. In the present study, although the subjects were ~10 years younger than those in the previous study, VO2 peak (Table 2) and probably mitochondrial function were reduced to a similar level as those in older subjects in the previous study15, which might have been caused by their lack of an exercise training habit and depression25,26. As a result, the same mechanisms likely worked in the ALA + SFC trial in the present study.

As shown in Fig. 3, training days, impulse, and time for 6 days in the ALA + SFC trial were significantly higher than those in the PLC trial with no carryover and period effects (Table 3), which are findings that are consistent with the previous study15. Regarding the mechanisms for the increased achievement for the ALA + SFC trial, since VO2 and VCO2 were saved above the intensities ≥45 W during graded cycling exercise after supplement intake and since the increase in [Lac−]p was significantly attenuated above the intensity, subjective feeling for fast walking might be improved due to reduced panting and muscle pain27,28, ultimately resulting in increases in impulse and time at fast walking in the trial.

As shown in Fig. 5, the MADRS score significantly decreased only in the ALA + SFC trial with no carryover and period effects (Table 3), in which subjects performed fast walking for IWT for ~50 min on average for 6 days, which agrees with previous studies7 suggesting a moderate or higher intensity of aerobic exercise (60–80% HR peak), 3 days/week, for 8 weeks improved depressive symptoms. Although the training period in the present study was shorter than in previous studies7, Dimeo et al.29 suggested that aerobic training at a moderate exercise intensity (Borg scale 13–14), which was 30 min/day for 10 days, significantly decreased the depression scores, the Hamilton Rating Scale by 33% and the self-assessed intensity score by 24%. Moreover, it was suggested that only one bout of high intensity exercise improved mood in depressive patients30 with an increase in the serum concentration of brain-derived neurotrophic factor (BDNF)31, which is suggested to decrease in depressive patients and increase when the symptoms are improved with anti-depressant drug administration32. In contrast, in the PLC trial, there was no significant change before and after the supplement intake period; rather, it tended to increase the MADRS score. This finding might be due to insufficient training achievement. Alternatively, the required exercise intensity might be too difficult for depressive patients. Indeed, Weinstein et al.33 suggested that when the demanded exercise intensity is perceived by depressive patients as being too hard, it evokes a negative mood. Therefore, ALA + SFC supplementation might lower the physical and psychological barriers to achieving moderate or higher intensity exercise training to decrease the MADRS score.

There are three experimental considerations that deserve additional discussion. First, we could not exclude any direct effects of the ALA + SFC supplementation on the central nervous system related to depression mechanisms. The association between depression and mitochondrial dysfunction in various brain regions has been suggested. For example, Omori et al.34 suggested that brain mitochondrial activity was enhanced by the 6-month administration of ALA in a mouse model of Alzheimer’s disease. Additionally, Perez et al.35 suggested that ALA + SFC supplementation of ~50 mg per day for 3 weeks improved the sleep quality score by ~30% (Pittsburgh Insomnia Rating Scale-20 question) in middle-aged and older people. Thus, ALA + SFC supplementation likely improves depression by activating mitochondrial function in the brain; however, since previous studies did not report physical activity during supplementation, it remains unclear how enhanced physical activity by ALA + SFC supplementation was involved in the results. In the present study, we found that the supplementation increased physical activity and improved depressive symptoms. Second, as shown in Table 2, the baseline HRrest significantly decreased in the ALA + SFC trial. Since sympathetic nervous system (SNS) activity is reportedly elevated in depressive patients36, the improved symptoms in the trial might decrease HRrest. Third, this study was conducted using only 9 subjects with a relatively short supplement intake period. Based on the present findings, a larger and longer trial to examine the effects of this treatment will be needed.

In summary, ALA + SFC supplementation improved respiratory and [Lac−]p responses during high-intensity exercise, increased fast-walking training achievement, and improved symptoms in middle-aged women with depression.

Methods

Subjects. This study protocol was approved by the Review Board on Human Experiments, Shinshu University School of Medicine, and conformed to the standards set by the Declaration of Helsinki. The trial was registered in UMIN (trial registration number: UMIN000013210) on February 21, 2014. We recruited female subjects aged 40–70 years who had no exercise habit from outpatients visiting our clinic in Tokyo for depression using a pamphlet at their scheduled examination. The recruitment was performed from 3.
February 22 to September 30, 2014. The inclusion criteria were that (1) they were diagnosed with major depressive disorder according to the Diagnostic and Statistical Manual of Mental Disorder (DSM ver.4)\(^\text{38}\) and had received psychotropic medication and psychotherapy, and the treatments were stabilized for 8 weeks; (2) they had recently experienced no drastic lifestyle changes; and (3) they were non-smokers and had no overt history of orthopaedic diseases to disturb IWT and iron deficiency anaemia to influence the results. We recruited female subjects to minimize any confounding effects of gender.

Eleven of 17 responders provided written informed consent and agreed to participate in this study. Since 2 subjects did not complete the protocol of the graded cycling test, we analysed the results in the remaining 9 subjects. However, no harmful events occurred during the intervention.

**Randomization.** Subjects were randomly assigned to PLC-(ALA + SFC) sequence or (ALA + SFC)-PLC sequence by an independent investigator (K.H.) using permuted-block randomization (block size: 4) with an allocation ratio of 1:1. The investigator was not involved in participant recruitment or any assessments. The random allocation sequence was generated using a computer.

**Protocol.** This study was carried out from February 22 to December 9, 2014 in a randomized, placebo-controlled, double-blind crossover design (Fig. 1) with an allocation ratio of 1:1. All subjects participated in two trials for 9 days each, followed by ≥10-day washout period; 7 days (days 1–7) for supplement intake and 2 days for graded cycling test before and after the trial (days 0 and 8). Subjects consumed either ALA + SFC (ALA + SFC trial) or the placebo supplement (PLC trial) for ≥1 hr before breakfast and dinner. During the graded cycling test, the cardiorespiratory responses and [Lac\(^-\)]\(_p\) were measured. During days 1–6, the training days, intensity, and time were recorded with a tri-axial accelerometer (JD Mate; Kissei Comtec, Matsumoto, Japan)\(^\text{39,40}\). There was no training on day 7 to avoid any acute influence of IWT on the graded cycling test. In subjects who still had a menstrual cycle, the experiments were scheduled during their follicular phases.

**Depression severity.** Before the graded cycling exercise test, a psychiatrist examined depression severity using the MADRS\(^\text{35}\); he was unaware of whether the subjects belonged to the ALA + SFC or PLC trials.

**Supplements.** The composition of supplements (SBI ALA Promo, Tokyo) is shown in Supplemental Table S1. The dose of ALA phosphate (100 mg/day) was the same as in the previous study using older women with no depression\(^\text{35}\). SFC, as a source of the iron ion in the supplements, was used to enhance the final step of haeme biosynthesis by the ABCB6 transporter and ferrochelatase in mitochondria\(^\text{31}\). The ALA + SFC and placebo supplements were similar in appearance, and all of the subjects and investigators who performed experiments and analyses were blinded to which trial the subjects actually underwent until all of the analyses were finished.

**Dietary intake.** Subjects in both trials were instructed to maintain their dietary habits including medications, except for the supplements, throughout the study while reporting food consumed during the period by answering a questionnaire prepared by a dietitian (FFQg version 4.0; Kenpakusya, Tokyo). We confirmed no significant differences in the values between the trials, including n-3 polyunsaturated fatty acid, which was reported to improve depressive symptoms\(^\text{41}\) (all, \(P > 0.4\)). Moreover, the amount of ALA contained in the diet\(^\text{43}\) was negligible compared to that in the supplement (Supplemental Tables S1, S2).

**Graded cycling test.** To minimize any inter-individual variation by different levels of food intake on the graded cycling test, we provided the standardized meals on the day before the test. At 0900 on the morning of the test, the subjects reported to the clinic that they were normally hydrated but had not eaten any food for more than 12 h before the experiment, except for a supplement 2 h before the visit. After measuring anthropological variables, the subjects entered a laboratory in the clinic controlled at ~25 °C and ~40% relative humidity. After a Teflon catheter was placed in the antecubital vein for blood sampling, the subjects rested quietly in an upright position on the saddle of the cycle ergometer for 15 min while all measurement devices were applied. After 10 min at rest, subjects started the cycling exercise at 60 revolutions/min at 0 W for 3 min, and then the intensity was increased to 15 W; then, the intensity was increased by 15 W every 2 min until the subjects were exhausted, during which time \(\text{VO}_2\), \(\text{VCO}_2\), and \(\dot{V}_{\text{E}}\) were measured with a respiratory gas analyser (Metamax3B; Cortex, Leipzig, Germany) (Fig. 2, Table 3) and heart rate (HR) was measured with a pulse rate monitor (Polar RS400; Vantage NV, Kempele, Finland) (Table 2). The criteria for judging whether exercise intensity reached \(\text{VO}_2\)\(_{\text{peak}}\) were that (1) subjects were not able to maintain the rhythm, (2) the respiratory quotient increased to over 1.1, and (3) HR reached the age-predicted maximal value. The \(\text{VO}_2\)\(_{\text{peak}}\) was determined by averaging the highest three consecutive values at the end of the exercise.

**Blood samples.** Blood samples were taken at rest and at the last minute of each intensity to determine [Lac\(^-\)]\(_p\) (YSI 2300 Stat Plus; Yellow Springs, OH) (Fig. 2, Table 3) and [Hb] at rest (Microsemi LC-660; HORIBA, Tokyo).

**Training achievement.** During the supplement intake period, except for the day before the second graded cycling test (days 1–6), subjects were instructed to perform IWT with the goal of repeating ≥5 sets of 3 min of slow walking at 40% \(\text{VO}_2\)\(_{\text{peak}}\), followed by 3 min of fast walking above 70% \(\text{VO}_2\)\(_{\text{peak}}\) per day, for ≥4 days/wk; during this period, the intensity and duration were recorded with a portable tri-axial accelerometer (JD Mate)\(^\text{56,57}\), and the measurements were transferred to the server computer at Shinshu University through the Internet after training\(^\text{56–20}\).
Training intensity was calculated from the product of body weight and average norm of three-dimensional accelerations and presented as the accumulated training impulse (N-min)\(^{39,40}\) for 6 days (Fig. 3).

**Analyses.** \(\text{VO}_2, \text{VCO}_2,\) and \([\text{Lac}]_p\) at the intensity \(\geq 45\) W. As shown in Fig. 2B, we analysed the average values of \(\text{VO}_2, \text{VCO}_2,\) and \([\text{Lac}]_p\) over the intensities \(\geq 45\) W in each subject for each trial since the increase in \([\text{Lac}]_p\) was significantly attenuated above the intensity in the ALA + SFC trial (Fig. 2A).

**Statistics.** Adequate sample size was determined based on the previous study assessing the effects of ALA + SFC supplementation on IWT achievements and \(\text{VO}_2, \text{VCO}_2,\) and \([\text{Lac}]_p\) during exercise in older women\(^{15}\) and the statistical power calculation.

One-way ANOVA for repeated measures was used to examine any significant differences in physical characteristics before vs. after the supplement intake period (Table 2) and examine any significant differences in dietary intake for the period between trials (Supplemental Table S2). This model was also used to examine any significant differences in training days, training impulse, and training time during the supplement intake period (Fig. 3); in the MADRS score before and after the period (Fig. 5); and in changes in \(\text{VO}_2, \text{VCO}_2,\) and \([\text{Lac}]_p\) \(\geq 45\) W (Fig. 2B) after the period between trials. Two-way ANOVA for repeated measures was used to examine any significant differences in the variables at every intensity during the graded cycling test before vs. after the supplement intake period in each trial, with a significant interactive effect of \([\text{before vs. after the supplement intake period}] \times \text{time}\) (Fig. 2A). As a subsequent post hoc test, the Tukey-Kramer test was used to perform any pairwise comparisons between trials.

In addition, because this study was conducted in a two-period crossover design, we performed a crossover analysis to examine the effects of 3 factors\(^{22}\) – carryover, period, and supplements – on the results, as described in the caption in Table 3.

The statistical power (1-\(\beta\)) is presented in the text as \(\alpha = 0.05\) when the key variables were significantly different between the PLC and ALA + SFC trials\(^{45}\). The null hypothesis was rejected when \(P < 0.05\). Values are expressed as the means ± SE, unless otherwise indicated.

**Data Availability.** The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

**References**

1. Kuehner, C. Why is depression more common among women than among men? *Lancet Psychiatry.* 4, 146–158 (2017).
2. Covinsky, K. E. *et al.* Depressive symptoms in middle age and the development of later-life functional limitations: the long-term effect of depressive symptoms. *J Am Geriatr Soc.* 58, 551–556 (2010).
3. Pinto Pereira, S. M., Geoffroy, M. C. & Power, C. Depressive symptoms and physical activity during 3 decades in adult life: bidirectional associations in a prospective cohort study. *JAMA Psychiatry.* 71, 1373–1380 (2014).
4. Azevedo Da Silva, M. *et al.* Bidirectional association between physical activity and symptoms of anxiety and depression: the Whitehall II study. *Eur J Epidemiol.* 27, 537–546 (2012).
5. Cooney, G. M. *et al.* Exercise for depression. *Cochrane Database Syst Rev.* CD004366 (2013).
6. Schuch, F. B. *et al.* Exercise as a treatment for depression: A meta-analysis adjusting for publication bias. *J Psychiatr Res.* 77, 42–51 (2016).
7. Perraton, L. G., Kumar, S. & Machotka, Z. Exercise parameters in the treatment of clinical depression: a systematic review of randomized controlled trials. *J Eval Clin Pract.* 16, 597–604 (2010).
8. Pedersen, B. K. & Saltin, B. Evidence for prescribing exercise as therapy in chronic disease. *Scand J Med Sci Sports.* 16(Suppl 1), 3–63 (2006).
9. Stanton, R. & Reaburn, P. Exercise and the treatment of depression: a review of the exercise program variables. *J Sci Med Sport.* 17, 177–182 (2014).
10. Rethorst, C. D., Wipfli, B. M. & Landers, D. M. The antidepressive effects of exercise: a meta-analysis of randomized trials. *Sports Med.* 39, 491–511 (2009).
11. Kramer, I. V., Helmes, A. W., Seelig, H., Fuchs, R. & Bengel, J. Correlates of reduced exercise behaviour in depression: the role of motivational and volitional deficits. *Psychol Health.* 29, 1206–1225 (2014).
12. Roshanaei-Moghadam, B., Katon, W. J. & Russo, I. The longitudinal effects of depression on physical activity. *Gen Hosp Psychiatry.* 31, 306–315 (2009).
13. Busch, A. M. *et al.* Preferences for Exercise as a Treatment for Depression. *Ment Health Phys Act.* 10, 68–72 (2016).
14. Patten, S. B., Williams, J. V., Lavorato, D. H. & Eliasziw, M. A longitudinal community study of major depression and physical activity. *Gen Hosp Psychiatry.* 31, 571–575 (2009).
15. Masuki, S. *et al.* Impact of 5-aminolevulinic acid with iron supplementation on exercise efficiency and home-based walking training achievement in older women. *J Appl Physiol* (1985). 120, 87–96 (2016).
16. Ogura, S. *et al.* The effect of 5-aminolevulinic acid on cytochrome c oxidase activity in mouse liver. *BMC research notes.* 4, 66 (2011).
17. Nishio, Y. *et al.* 5-aminolevulinic acid combined with ferrous iron enhances the expression of heme oxygenase-1. *Int Immunopharmacol.* 19, 300–307 (2014).
18. Nose, H. *et al.* Beyond epidemiology: field studies and the physiology laboratory as the whole world. *J Physiol.* 587, 5569–5575 (2009).
19. Masuki, S., Morikawa, M. & Nose, H. Interval Walking Training Can Increase Physical Fitness in Middle-Aged and Older People. *Exerc Sport Sci Rev.* 45, 154–162 (2017).
20. Morikawa, M. *et al.* Physical fitness and indices of lifestyle-related diseases before and after interval walking training in middle-aged and older males and females. *Br J Sports Physiol.* 45, 216–224 (2011).
21. Snaiti, R. P., Harrop, P. M., Newby, D. A. & Teale, C. Grade scores of the Montgomery-Asberg Depression and the Clinical Anxiety Scales. *The British journal of psychiatry: Br J Psychiatry.* 148, 599–601 (1986).
22. Chow S. C., Liu J. P. Statistical inferences for effects from a standard 2 × 2 crossover design. In: Design and Analysis of Bioavailability and Bioequivalence Studies. 48–69 (Dekker, 1992).
23. Troncze, I., Byrne, E. & Marzuki, S. Decline in skeletal muscle mitochondrial respiratory chain function: possible factor in ageing. *Lancet.* 1, 637–639 (1989).
24. Rooyackers, O. E., Adey, D. B., Ades, P. A. & Nair, K. S. Effect of age on in vivo rates of mitochondrial protein synthesis in human skeletal muscle. *Proc Natl Acad Sci USA* 93, 15364–15369 (1996).
25. Gram, M. et al. Skeletal muscle mitochondrial H2 O2 emission increases with immobilization and decreases after aerobic training in young and older men. J Physiol. 593, 4011–4027 (2015).
26. Russ, D. W. & Kent-Braun, J. A. Is skeletal muscle oxidative capacity decreased in old age? Sports Med. 34, 221–229 (2004).
27. Gregory, N. S., Whitley, P. E. & Sluka, K. A. Effect of Intramuscular Protons, Lactate, and ATP on Muscle Hyperalgesia in Rats. PLoS One. 10, e0138576 (2015).
28. Hoheisel, U., Reinhohl, I., Unger, T. & Mense, S. Acidic pH and capsaicin activate mechanosensitive group IV muscle receptors in the rat. Pain. 110, 149–157 (2004).
29. Dimeo, F., Bauer, M., Vaharam, I., Proest, G. & Halter, U. Benefits from aerobic exercise in patients with major depression: a pilot study. Br J Sports Med. 35, 114–117 (2001).
30. Bartholomew, J. B., Morrison, D. & Ciccolo, J. T. Effects of acute exercise on mood and well-being in patients with major depressive disorder. Med Sci Sports Exerc. 37, 2032–7 (2005).
31. Laske, C. et al. Exercise-induced normalization of decreased BDNF serum concentration in elderly women with remitted major depression. Int J Neuropsychopharmacol. 13, 595–602 (2010).
32. Sen, S., Duman, R. & Sanacora, G. Serum brain-derived neurotrophic factor, depression, and antidepressant medications: meta-analyses and implications. Biol Psychiatry. 64, 527–532 (2008).
33. Weinstein, A. A., Deuster, P. A., Francis, J. L., Beadling, C. & Kop, W. J. The role of depression in short-term mood and fatigue responses to acute exercise. Int J Behav Med. 17, 51–57 (2010).
34. Bansal, Y. & Kuhad, A. Mitochondrial Dysfunction in Depression. Curr Neuropharmacol. 14, 610–618 (2016).
35. Omori, C. et al. Facilitation of brain mitochondrial activity by 5-aminolevulinic acid in a mouse model of Alzheimer’s disease. Nutr Neurosci. 20, 538–546 (2017).
36. Perez, M. H. et al. The Role of 5-Aminolevulinic Acid (5-ALA) and Sleep. Int J Clin Med. 4, 1–7 (2013).
37. Veith, R. C. et al. Sympathetic nervous system activity in major depression. Basal and desipramine-induced alterations in plasma norepinephrine kinetics. Arch Gen Psychiatry. 51, 411–422 (1994).
38. American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders, 4th Edn. (American Psychiatric Association, 1994).
39. Yamazaki, T. et al. A new device to estimate VO2 during incline walking by accelerometer and barometry. Med Sci Sports Exerc. 41, 2213–2219 (2009).
40. Iwashita, S. et al. Triaxial accelerometer to evaluate walking efficiency in older subjects. Med Sci Sports Exerc. 35, 1766–1772 (2003).
41. Ota, U. et al. 5-Aminolevulinic acid (5-ALA): a precursor of heme fermentation, metabolism and usage. ALA-Porphyrin Science. 1, 3–17 (2013).
42. Giles, G. E., Mahoney, C. R. & Kanarek, R. B. Omega-3 fatty acids influence mood in healthy and depressed individuals. Nutr Rev. 71, 727–741 (2013).
43. Fueki, S., Ueda, Y., Dota, M. & Negishi, Y. Development of analytical method of 5-aminolevulinic acid in foods. Porphyrins. 19, 9–14 (2010).
44. Cohen J. Statistical power analysis for the behavioral sciences. 1–17 (Lawrence Erlbaum Associates, 1988).

Acknowledgements
This study was supported by grants from the Japan Society for the promotion of Science (24240089, 25670117 and 15H01830).

Author Contributions
H.S., K.T., M.N. and H.N. conception and design of research; H.S., A.M., Y.O., and Y.K. performed experiments; H.S., S.M., and A.M. analyzed data; H.S., S.M., and H.N. interpreted data; H.S., S.M., and H.N. prepared the manuscript; All authors approved the final version of the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-25452-2.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018