Acute effects of whole body vibration exercise on post-load glucose metabolism in healthy men: a pilot randomized crossover trial

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Abstract

Purpose Exercise on a whole body vibration (WBV) platform, namely WBV exercise (WBVE), has long-term beneficial effects on glucose metabolism, similarly to conventional moderate-intensity exercise. Conventional moderate-intensity exercise reduces post-load plasma glucose levels at the acute phase. This study aimed to reveal acute effects of WBVE on post-load glucose metabolism.

Methods This randomized crossover trial enrolled 18 healthy men. They completed the following three interventions in a random order: (1) a 2-hour 75-g oral glucose tolerance test (OGTT) without WBVE (OGTT-alone), (2) 20-minute WBVE before an OGTT (WBVE→OGTT), and (3) 20-minute WBVE during an OGTT (OGTT→WBVE). Post-load glucose metabolism in the WBVE→OGTT and OGTT→WBVE interventions were compared with that in the OGTT-alone intervention.

Results Plasma glucose levels in the WBVE→OGTT and OGTT→WBVE interventions were not significantly different from those in the OGTT-alone intervention at any time point except 15 min, wherein the WBVE→OGTT intervention had higher glucose levels (111 [interquartile range, 102–122] mg/dL vs 122 [111–134] mg/dL, P = 0.026). Higher plasma glucagon levels were observed at 0 min in the WBVE→OGTT intervention and at 60 min in the OGTT→WBVE intervention (P = 0.010 and 0.015). Cortisol, Growth hormone, and adrenaline levels were significantly increased after WBVE, whereas noradrenaline levels were not. Serum insulin levels in the WBVE→OGTT intervention were significantly higher than those in the OGTT-alone intervention at 0 min (P = 0.008).

Conclusions WBVE did not decrease post-load plasma glucose levels at the acute phase. Acute effects of WBVE on post-load glucose metabolism would not be identical to those of conventional exercise.

The unique trial number and the name of the registry: UMIN000036520, www.umin.ac.jp, date of registration, June 12, 2019.

Keywords Whole body vibration exercise · Healthy men · Glucose metabolism · Acute phase

Introduction

Physical exercise is a key factor to reduce the risk of type 2 diabetes mellitus [1–3]. Exercise on a whole body vibration (WBV) platform, namely WBV exercise (WBVE), is a convenient, safe, and effective exercise modality, and has become popular in recent years. WBV enhances muscle activity, using mechanical loading of alternative gravitational acceleration with three-dimensional vibration [4, 5]. WBV platforms for home use are now commercially available, and attract increasing attention during this coronavirus pandemic, when citizens are asked to stay at home and refrain from going out. Clinical studies demonstrated that WBV had long-term beneficial effects, including
decrease of body fat mass [6], increase of muscle mass [7], and improvement of insulin sensitivity and glucose regulation [8], similarly to conventional moderate-intensity exercise [9]. However, it remained unrevealed how WBVE would affect glucose metabolism at the acute phase. The literature [10–14] shows that in comparison with rest, conventional moderate-intensity exercise, either following or followed by glucose ingestion, lowers post-load plasma glucose levels at the acute phase. By contrast, no data were so far available about the acute phase influence of WBVE on post-load glucose levels. The aim of the current pilot study was to investigate the acute effects of WBVE either following or followed by glucose ingestion on post-load glucose metabolism in healthy men, in comparison with rest as the control.

Materials and methods

Study design

This pilot study was an open-label, randomized, 6-sequence, 3-period, 3-intervention crossover trial with a 1:1:1:1:1:1 allocation, and was conducted at Osaka University, Suita City, Osaka, Japan, between July 2019 and July 2020. This study compared the acute effect on glucose metabolism between WBVE and rest in healthy men. Eighteen healthy male volunteers were enrolled. Each study participant received all of the following three interventions in a random order: (1) performing a 2-hour 75-g oral glucose tolerance test (OGTT) without any exercise including WBVE (OGTT-alone), (2) performing 20-minute WBVE before an OGTT (WBVE → OGTT), and (3) performing 20-minute WBVE during an OGTT (OGTT → WBVE) (Fig. 1). OGTTs were started at approximately 10:00 AM after an overnight fast. During an OGTT, blood samples were collected at 0 min and thereafter every 15 min for 2 h. In the OGTT → WBVE intervention, WBVE was started 30 min after glucose ingestion. The timing was determined in reference to the optimal timing of post-meal conventional exercise [15]. By contrast, we found no general consensus about the optimal timing of pre-load or pre-meal exercise [15, 16]. We therefore assigned different WBVE-to-OGTT intervals to the study participants in the WBVE → OGTT intervention, to supplementarily analyze the association between the interval and study outcomes. The interval was ranged from 10 to 60 min, which was randomly set in each participant. A blood sample was additionally collected 10 min after
WBVE in the WBVE → OGTT intervention. The interval between each intervention was at least two days.

The study was in accordance with the Declaration of Helsinki, and approved by the Institutional Review Board of Osaka University Hospital (date of approval, May 10, 2019; approval number, 18452-3). All of the participants provided written informed consent prior to participation. The trial was registered as UMIN000036520 at UMIN Clinical Trials Registry.

**Study participants and allocation**

Eighteen healthy male volunteers aged 20–59 years were enrolled to the study. The exclusion criteria included the history of diabetes or glucose intolerance, diseases potentially related to abnormal glucose homeostasis, symptoms potentially deteriorated by vibration (dizziness etc.), and a risk of fall. The current cross-over trial had 6 sequences, each of which consisted of 3 interventions (OGTT-alone, WBVE → OGTT, and OGTT → WBVE) in a different order. The study participants were allocated to one of the 6 sequences based on a 1:1:1:1:1:1-ratio randomization. The allocation concealment was ensured by using sequentially numbered opaque sealed envelopes (SNOSE). In brief, sealed opaque envelopes, each of which contained a card indicating one of the 6 sequences, were prepared and shuffled in advance of the clinical trial by one researcher (M.T.). Once a participant consented to participate in the clinical trial, a sealed opaque envelope was opened by another researcher (H.W.), and the participant was assigned to the sequence that was indicated by the card enclosed in the envelope. Sequences and interventions were opened (i.e., not blinded) after the assignment.

**WBVE**

WBVE was performed using a WBV platform (Personal Power Plate®, Performance Health Systems, LLC, Northbrook, IL), which mechanically generated three dimensional (i.e., vertical, horizontal, and sagittal) WBV load. The duration and the type of WBVE were determined by reference to previous clinical trials of conventional exercise, which demonstrated a significant acute phase effect of conventional exercise on post-load plasma glucose levels, with the sample size between 6 and 18 participants [10–14].

**Measurement**

All blood samplings were performed from the indwelling venous catheter. The samples were immediately centrifuged at 1000 rpm for 10 min at 4°C, and extracted plasma and serum were stored at −20°C. Plasma glucose levels were measured at 0 min and thereafter every 15 min, and serum insulin and plasma glucagon levels were measured at 0 min and thereafter every 30 min during a 2-hour OGTT. Plasma glucose was measured using a hexokinase activity assay kit (Kanto Chemical Co., Inc., Tokyo, Japan), and serum insulin was measured using a chemiluminescent enzyme immunoassay kit (Fujirebio Inc., Tokyo, Japan). Glucagon levels were measured using a sandwich enzyme-linked immunosorbent assay kit (Mercodia AB, Uppsala, Sweden) [17, 18]. Plasma cortisol, growth hormone (GH), noradrenaline and adrenaline levels were measured 10 min after WBVE in the WBVE → OGTT and the OGTT → WBVE interventions, and at the corresponding time points in the OGTT-alone intervention (Fig. 1). Cortisol and growth hormone levels were measured using an electro chemiluminescence immunoassay kit (Roche Diagnostics K.K, Tokyo, Japan), and noradrenaline and adrenaline levels were measured using a high performance liquid chromatography method (Tosoh Corporation, Tokyo, Japan). All of the measurements were performed by SRL Inc. (Tokyo, Japan)

**Study outcome**

The primary outcome measure was plasma glucose levels during an OGTT. Secondary outcome measures included serum insulin, plasma glucagon levels, and endocrine hormone levels (cortisol, GH, noradrenaline and adrenaline levels).

The sample size of the current pilot study was determined with reference to previous clinical trials of conventional exercise, which demonstrated a significant acute phase effect of conventional exercise on post-load plasma glucose levels, with the sample size between 6 and 18 participants [10–14].

**Statistical analysis**

We primarily treated data without parametric assumptions and therefore reported the data as medians interquartile ranges (IQRs). However, parametric statistics (means and standard deviations [SDs]) would be sometimes more informative and therefore the values were additionally reported. Glucose, insulin, and glucagon levels at each time point in the WBVE → OGTT and OGTT → WBVE interventions were compared to the corresponding levels in the OGTT-alone intervention, using the Wilcoxon signed rank
test. The effect size \( r \) for the Wilcoxon signed rank test (\( z \) score divided by the square root of \( n \), and Cohen’s \( d \) (the standardized difference calculated from means and SDs) were also demonstrated. Effect size \( r \) was categorized as small (0.1), medium (0.3), or large (0.5), and \( d \) was categorized as small (0.2), medium (0.5), or large (0.8) [19]. As the sensitivity analysis, the between-intervention differences were tested using the linear mixed model in which random effects for participants, fixed effects for the period, the allocated sequence, and the intervention were included. We also compared the change (delta) from baseline (0 min) between the interventions, using the Wilcoxon signed rank test.

Cortisol, GH, noradrenalin and adrenaline levels after WBVE were compared to those at the corresponding time point in the control intervention, using the Wilcoxon signed rank test. The association between the WBVE-to-OGTT interval and study outcomes in the WBVE \( \rightarrow \) OGTT intervention were investigated using the Spearman’s rank correlation method.

All statistical analyses were performed using R version 4.0.3 (R Development Core Team, Vienna, Austria).

**Results**

All of the 18 study participants completed the study, and were included in the analysis. The population had median and mean age of 33 (31–36) and 34 ± 5 years, and median and mean body mass index levels of 21 (19–23) and 22 ± 5 kg/m\(^2\). The median and mean WBVE-to-OGTT interval was 35 (20–50) and 35 ± 18 min in the WBVE \( \rightarrow \) OGTT intervention.

Figure 2A shows plasma glucose levels during an OGTT. Plasma glucose levels in the WBVE \( \rightarrow \) OGTT and OGTT \( \rightarrow \) WBVE interventions were not significantly different from those in the OGTT-alone intervention at any time point except 15 min, wherein plasma glucose levels were significantly higher in the WBVE \( \rightarrow \) OGTT intervention (111 [102–122], 111 ± 14 mg/dL vs 122 [111–134], 124 ± 19 mg/dL, \( P = 0.026 \) by the Wilcoxon signed rank test). Serum insulin levels in the WBVE \( \rightarrow \) OGTT intervention were significantly higher than those in the OGTT-alone intervention at 0 min (4.7 [3.7–5.7], 5.3 ± 3.3 μU/mL vs 3.7 [2.6–4.9], 4.1 ± 2.2 μU/mL, \( P = 0.008 \) by the Wilcoxon signed rank test), but not at any other time point (all \( P > 0.05 \) by the Wilcoxon signed rank test) (Fig. 2B). Higher plasma glucagon levels were observed at 0 min in the WBVE \( \rightarrow \) OGTT intervention and at 60 min in the OGTT \( \rightarrow \) WBVE intervention (14.7 [11.8–20.3], 16.2 ± 6.0 pg/mL vs 19.3 [15.3–27.0], 23.7 ± 16.3 pg/mL, \( P = 0.015 \) by the Wilcoxon signed rank test, respectively) (Fig. 2C). The sensitivity analysis using the linear mixed model showed similar results (Supplementary Table 1). The effect sizes in the sample are demonstrated in Supplementary Table 2. All measurements with statistically significant difference consistently had an effect size \( r \) of larger than 0.5, categorized as “large”. On the other hand, those without significant difference consistently had a smaller \( r \), categorized as “medium” or smaller; none of them were categorized as “large”. The effect size \( d \) was larger than 0.5, categorized as “medium” or larger, in measurements with significant difference, and smaller in those without it; the
exception was 60-minute glucagon levels in the OGTT → WBVE versus OGTT-alone intervention, which reached statistical significance (P < 0.05 by the Wilcoxon signed rank test) but had a “small” effect size d (equal to 0.384). The comparison of deltas is shown in Supplementary Table 3.

Cortisol, GH, and adrenaline levels after WBVE were significantly higher than those at the corresponding time points in the OGTT-alone interventions, whereas noradrenaline levels were not. (Table 1).

In the WBVE → OGTT intervention, the interval between WBVE and OGTT had no significant correlation with any study outcomes (Supplementary Table 4).

There were no adverse events during the trial.

**Discussion**

The current pilot study demonstrated that plasma glucose levels in the WBVE → OGTT and OGTT → WBVE interventions were not significantly different from those in the OGTT-alone intervention at any time point except 15 min, wherein plasma glucose levels were significantly higher in the WBVE → OGTT intervention. In the WBVE → OGTT intervention, serum insulin levels at 0 min were significantly higher, and plasma glucagon levels at 0 min were significantly higher compared to the OGTT-alone intervention. In the OGTT → WBVE intervention, post-load plasma glucagon levels were significantly higher at 60 min than in the OGTT-alone intervention, whereas serum insulin levels were not significantly different at any point compared to the OGTT-alone intervention. Plasma cortisol, GH and adrenaline levels at 10 min after WBVE were significantly higher than those at the corresponding time points in the OGTT-alone interventions; on the other hand, noradrenaline levels were not significantly changed with WBVE. The effect size in the sample r was categorized consistently as “large” in the measurements with significant difference and “medium” or smaller in those without significant difference. Furthermore, the effect size in the sample d was categorized as “medium” or larger in those with significant difference and as smaller in those without significant difference; the exception was 60-minute glucagon levels in the OGTT → WBVE vs OGTT-alone intervention, which reached statistical significance but had a “small” effect size d.

The additional analysis on the change (delta) from baseline showed that delta plasma glucose levels in the WBVE → OGTT and OGTT → WBVE interventions were not significantly different from those in the OGTT-alone intervention at any time point except 15 min, wherein plasma glucose levels were significantly higher in the WBVE → OGTT intervention. These results were similar to the primary analysis. Delta serum insulin levels in the WBVE → OGTT and OGTT → WBVE interventions were not significantly different from those in the OGTT-alone intervention at any time point. Higher delta plasma glucagon levels were observed at 60 min in the OGTT → WBVE intervention, which was similar to the primary analysis. In contrast, lower delta plasma glucagon levels were observed at 60, 90, and 120 min in the WBVE → OGTT intervention, which would be caused by the elevation of absolute plasma glucagon levels at 0 min in the WBVE → OGTT intervention.

Previous studies demonstrated that conventional moderate-intensity exercise both before and after glucose

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Table 1 Endocrine hormone levels 10 min after WBVE

|                     | OGTT-alone intervention | WBVE → OGTT intervention | OGTT → WBVE intervention |
|---------------------|-------------------------|---------------------------|---------------------------|
| Growth hormone (ng/mL) | 0.26 [0.14, 0.98], 1.05 ± 1.90 | 3.26 [2.72, 6.92], 5.25 ± 5.70, (P < 0.001) | N/A                      |
| Cortisol (μg/dL)     | 7.0 [5.5, 8.8], 6.98 ± 2.07 | 8.5 [7.2, 13.5], 10.6 ± 5.5, (P = 0.026) | N/A                      |
| Noradrenaline (pg/mL) | 269 [201, 369], 310 ± 147 | 336 [303, 406], 351 ± 75, (P = 0.167) | N/A                      |
| Adrenaline (pg/mL)   | 29 [24, 32], 32 ± 13 | 55 [42, 77], 59 ± 25, (P < 0.001) | N/A                      |

During OGTT: 10 min after WBVE in the OGTT → WBVE intervention and the corresponding time point in the control intervention

| Growth hormone (ng/mL) | 0.11 [0.04, 0.20], 0.15 ± 0.16 | N/A                      | 0.28 [0.11, 0.68], 0.53 ± 0.66, (P = 0.006) |
| Cortisol (μg/dL)       | 7.4 [4.6, 9.2], 6.9 ± 2.6 | N/A                      | 8.8 [6.0, 12.7], 10.1 ± 5.4, (P = 0.024) |
| Noradrenaline (pg/mL)  | 344 [248, 441], 363 ± 156 | N/A                      | 306 [245, 446], 341 ± 120, (P = 0.551) |
| Adrenaline (pg/mL)     | 23 [18, 33], 25 ± 11 | N/A                      | 40 [24, 50], 42 ± 22, (P < 0.001) |

Date are medians [interquartile ranges], means ± standard deviations, as well as P values versus the OGTT-alone intervention, tested by the Wilcoxon signed rank test.

N/A not assessed.
ingestion decreased post-load glucose levels at the acute phase [10–14]. By contrast, WBVE did not decrease post-load glucose levels either before or after glucose ingestion; rather, post-load plasma glucose levels were increased at 15 min in the WBVE → OGTT intervention. WBVE might have a different acute phase effect on post-load glucose metabolisms from conventional exercise.

The difference might be at least partially explained by altered endocrinological responses. After WBVE, GH levels were elevated compared with rest, and this response was apparently consistent with that observed after conventional exercise [20]. However, a recent study demonstrated that GH levels were higher after WBVE (i.e., exercise using WBV) than after exercise without WBV, suggesting that the stimulation of WBV per se would additionally increase GH levels [21]. Glucagon and cortisol also appeared to be secreted somewhat differently. Previous studies reported that conventional moderate-intensity exercise did not significantly affect glucagon or cortisol levels [22, 23]. By contrast, in the WBVE → OGTT intervention, glucagon levels at 0 min were significantly higher than those in the OGTT-alone intervention, and similar trend was seen at 30 min although the difference was not significant. Similarly, in the OGTT → WBVE intervention, glucagon levels at 60 min were significantly higher than the OGTT-alone intervention, although the parametric effect size in the sample was “small”, suggesting that from the parametric viewpoint, its difference in the sample might be smaller than that of the other measurements with statistical significance. Furthermore, cortisol levels after WBVE in the WBVE → OGTT and OGTT → WBVE interventions were significantly higher than those at the corresponding time points in the OGTT-alone intervention. The increase of these counterregulatory hormones after WBVE might contribute to a non-decrease or even increase of post-load glucose levels.

Cardinale M and colleagues demonstrated that cortisol levels were higher after WBVE (i.e., exercise using WBV) than after exercise without WBV, suggesting that the stimulation of WBV would increase cortisol levels [24]. The increase of cortisol levels after WBVE compared with rest in the current study would reflect the body response to the WBV stimulation. Several studies failed to detect a significant elevation of cortisol levels after WBVE [21, 25, 26], which might come from their small and insufficient sample size (n = 6–8). Other studies reported that decreased cortisol levels after WBVE [27, 28]. However, those studies compared cortisol levels after WBVE to those before the WBVE, and not to those in the control at the corresponding time point. The decrease of cortisol in their studies might reflect the circadian rhythm of cortisol, which has a peak around wake-up time and a subsequent drop [29]. The current study, comparing cortisol levels after WBVE with those at rest at the same time point, would successfully detect the increase of cortisol levels by the intervention.

Catecholamine levels are another important factor increasing plasma glucose levels. Previous studies demonstrated that noradrenaline rather than adrenaline levels were predominantly increased after conventional exercise [20]. Conversely, in the current study, adrenaline levels were significantly elevated 10 min after WBVE, whereas noradrenaline levels were not. Previous studies reported that adrenaline had a stronger effect on the increase of plasma glucose levels than noradrenaline [30]. The adrenaline-dominant elevation might be also associated with altered post-load glucose metabolisms after WBVE. Loreto CD et al. reported a significant elevation of noradrenaline levels 35 min after the end of WBVE [31]. Although we did not measure catecholamine levels at the same time point, such late alternations of catecholamine levels might also contribute to a non-decrease or even increase of post-load glucose levels. The current study observed a temporal increase of insulin levels after WBVE, which would reflect an increased insulin requirement against those elevated counter-regulatory hormones levels. This was in contrast to conventional exercise, where post-load insulin levels were decreased [10, 11, 13, 14], indicating decreased insulin requirements.

Why GH, glucagon, cortisol, and adrenaline levels were excessively increased after WBVE remained unknown. However, these findings suggest the body’s overreaction to the exercise, or a somewhat stronger response than expected from conventional exercise. During exercise, contracting skeletal muscle rapidly increases glucose uptake via glucose transporter type 4 (GLUT4) translocation to the plasma membrane [32, 33]. The body can appropriately balance glucose supply with the sudden, arbitrary demand, and successfully avoid hypoglycemia, wherein the liver plays a major role [34]. The physiological mechanisms of the delicate systemic balance, however, has not been completely understood. Humans might have acquired the delicate balance partially in an empirical manner during the process of evolution. WBVE is a novel exercise modality that humans have never experienced before. The body’s empirical endocrinological response might result in an overreaction relative to the muscle’s glucose consumption. These endocrinological overreactions might induce excess hepatic glucose release, leading to an insufficient suppression of post-load glucose levels.

Another factor potentially relevant to the altered glucose metabolisms is the glucose demand of the skeletal muscle. Previous studies reported that glucose uptake of the skeletal muscle during isometric contraction was different from that during isotonic contraction accompanied by a larger change of the muscle length [35]. During WBV static squatting, the skeletal muscle kept isometric from a macro point of view, while it was exposed to three-dimensional vibration load from a micro point of view. Muscle contraction during WBVE might be different from that during conventional
exercise, and the difference might affect the glucose demand of the skeletal muscle.

The current findings indicate that WBVE, expected to have long-term beneficial effects on glucose metabolism like conventional moderate-intensity exercise, would affect post-load glucose metabolisms in a different way from conventional moderate-intensity exercise at the acute phase. WBVE might not be an equivalent or alternative modality to conventional exercise; potential benefits and risks might be different from those expected from abundant evidence of conventional exercise. Many more physiological and clinical studies will be needed to accumulate the evidence of WBVE.

Our study has some limitations. First, the sample size was limited. The effect size in the sample size (n=18) could detect the population effect size (not the sample effect size) r>0.720 by the Wilcoxon signed rank test (and d>0.701 by the paired t test) with a power of 80% (calculated by G*Power version 3.1.9.2, Heinrich-Heine-Universität Düsseldorf, Düsseldorf, Germany), meaning that a difference with a smaller effect size in the population might be overlooked. Future studies with a larger sample size will be needed to detect such differences with a smaller effect size. Second, the current study did not include an intervention of conventional (i.e., WBV-free) exercise. We were therefore unable to compare WBVE with conventional exercise directly, and to distinguish the effect of the WBV stimulation from that of exercise per se. Third, the current study did not assess hepatic glucose production, muscle glucose uptake, and actual muscle contraction. Fourth, the subjects of the current study were healthy volunteers. It remained unknown whether the similar results would be obtained from patients with diabetes. Fifth, the effect of unmeasured endocrine hormones remained unclear. Some studies examined testosterone, insulin-like growth factor 1 levels after WBVE [24, 36, 37], which might contribute to post-load glucose metabolism after WBVE. Future studies will be needed to validate the current findings.

In conclusions, WBVE either following or followed by glucose ingestion did not decrease post-load plasma glucose levels like conventional moderate-intensity exercise.

Availability of data

The datasets generated and analysed during the current study are not publicly available due to ethical reason, but available from the corresponding author on reasonable request and with permission of the Institutional Review Board of Osaka University Hospital.

Author contributions H.W. designed the study, wrote the manuscript and researched date. M.T. designed the study and wrote the manuscript. N.K. contributed to discussion and reviewed/edited the manuscript. T.K., K.N. and I.S. reviewed/edited the manuscript. M.T. is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the date analysis.

Compliance with ethical standards

Conflict of interest The authors declare no competing interests.

Ethics approval The study was in accordance with the Declaration of Helsinki, and approved by the Institutional Review Board of Osaka University Hospital (date of approval, May 10, 2019; approval number, 18452-3).

Consent to participate Informed consent was obtained from all individual participants included in the study.

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