THESIS

1 MONTH EFFECT OF BREAKING UP SEDENTARY ACTIVITY ON INSULIN SENSITIVITY AND GLUCOSE HOMESOTASIS IN FREE-LIVING OVERWEIGHT/OBESE ADULTS

Submitted by
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ABSTRACT

1 MONTH EFFECT OF BREAKING UP SEDENTARY ACTIVITY ON INSULIN SENSITIVITY AND GLUCOSE HOMESTASIS IN FREE-LIVING OVERWEIGHT/OBESE ADULTS

Sedentary behavior (SB) triggers an inability to adjust substrate use to substrate availability (low metabolic flexibility, MF), which may precede glucose intolerance in the pathogenesis of insulin resistance. We and others have shown that frequent interruptions in SB leads to improved glycemic control, however the underlying role of MF in this process is unknown. This study examined the effects of breaking up SB on MF and glucose metabolism in free-living overweight and obese adults. To distinguish effects of breaking up SB from being physically active, we also studied a group where participants performed a single energy matched continuous bout of exercise. Physically inactive, adults (12F/9M, mean±SD, age: 33±8 yr, BMI: 29.5±3.3 kg/m$^2$) were randomly assigned to a 4 week intervention consisting of brisk walking for 5 min each hour for 10h, 5 d/wk (MICRO, n=10), or 4 weeks of an intervention consisting of one continuous 45 min bout of exercise per day, 5d/wk (ONE, n=9). Outcomes assessed at baseline and after each intervention included: MF (waking respiratory quotient, RQ, minus sleeping RQ as measured in a whole room calorimeter), insulin sensitivity (SI, IVGTT), 24h glycemia (continuous glucose monitor), 24h glucose oxidation ($^{13}$C glucose tracer), SB, time spent standing, time spent stepping (ActivPAL) and TEE (double labeled water). Groups were similar on all outcome variables at baseline. Linear mixed models evaluated intervention and intervention-by-group effects. MICRO and ONE decreased time sitting and increased time stepping with no significant changes in TEE. Compared to ONE, MICRO decreased 24h
glycemic variability (p=0.06), improved the acute whole body insulin sensitivity (p=0.08) and acute insulin response to glucose (AIRg) (p=0.02), maintained exogenous glucose oxidation (p<0.03) and improved MF (p=0.02). Independent of time sitting and stepping, breaking up SB improves glucose homeostasis and MF. The effects of such an intervention in persons with type 2 diabetes warrants further study.
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1. INTRODUCTION

Overweight and obese individuals are major health concerns in most Westernized countries\(^1\). Experts claim a pandemic of obesity with tremendous burdens on societies and health care systems\(^1\). In 2016 nearly 72% of all adults were overweight or obese\(^2\). Excess weight and obesity are risk factors for developing type 2 diabetes (T2D).\(^3\) 9.4% of the US population has been diagnosed with T2D, and an additional 84.1 million have prediabetes\(^3\). T2D cost US taxpayers $245 billion in 2012, a 41% increase from 2007\(^1\). Because the epidemic has mainly occurred over the past 50 years, blaming genetics alone is inappropriate. With the agricultural, industrial and technological revolutions, both the built and social environmental factors have however drastically been modified, which have influenced diet and physical activity. Poor diet (high fat, fructose, high saturated fat, high energy density), passive transportation and general adoption of sedentary behaviors have become the new standards of the modern life.

Physical inactivity (i.e. not meeting the physical activity guidelines of 150 minutes of moderate to vigorous physical activity) and sedentary behaviors (i.e. having a MET value between 1 and 1.5; equivalent to sitting or lying down)\(^4\) are recognized risk factors for developing metabolic diseases including obesity and T2D\(^5\). Our research group showed that physical inactivity induced by bed rest studies in healthy normal-weight adults lead to multiple metabolic alterations including whole-body and muscle insulin resistance, even when energy balance is maintained\(^6\). This is associated with the development of metabolic inflexibility, defined as the inability of the body to adjust substrate use to changes in fuel availability and energy demand. Metabolic inflexibility is a core component of metabolic diseases and a predictor of long term weight gain\(^7\). Importantly, we showed that metabolic inflexibility induced by
physical inactivity precedes the development of glucose intolerance\textsuperscript{6}. Glucose tolerance is essential to metabolic health. Chronic hyperglycemia and also large fluctuations in blood glucose are both risk factors for developing T2D and the associated cardiovascular complications for people with diabetes \textsuperscript{8}.

Based on a large body of epidemiological and experimental evidence, the key to healthy body composition, physiology, and T2D management is largely dependent on physical activity. However, while most US adults know exercise is beneficial and 62\% self-reported meeting the ACSM physical activity guidelines of 150 minutes of moderate to vigorous physical activity (MVPA), only 9.6\% of those surveyed actually met the guidelines based on accelerometry\textsuperscript{9}. Some studies have even found that independent of meeting the American College of Sports Medicine physical activity guidelines, sedentary time showed associations with waist circumference, insulin, and HOMA-IR \textsuperscript{10}. Furthermore, population studies showed that adults who frequently interrupt their sedentary (sitting) time short bouts of activity have healthier metabolic profiles than adults who engage in long periods of uninterrupted sitting\textsuperscript{10}. An increasing number of intervention studies showed that acute exposure to breaking up sedentary behaviors decreases post-prandial glycemia and insulinemia in adults who are normal-weight, overweight, obese and attained of T2D\textsuperscript{11}. We and others showed that these metabolic effects were more pronounced than when physical activity was performed as a single continuous match for total active time and energy expenditure\textsuperscript{12,13}. We further reported that interrupting sedentary time with short bouts of walking primarily increases 24-h carbohydrate oxidation, where an energy matched single continuous bout increased 24-h fat oxidation\textsuperscript{11}. Finally, performing microbouts (\(<\text{or equal}5\text{-min})\) of activity also lowers glycemic variability the following night\textsuperscript{5}. Based on these epidemiological and experimental evidence, the US Office of Disease Prevention
and Health Promotion updated in 2018 the US physical activity guidelines. They state that “adults should move more and sit less throughout the day. Some physical activity is better than none. Adults who sit less and do any amount of moderate-to-vigorous physical activity gain some health benefits.”

However, no study to date has proven with a long term study that “sitting less and moving more” has positive effects on health via a study in a free living condition or the exact degree of effectiveness that this has. The purpose of this study was to examine the sustained effect of breaking up sedentary behavior with short bouts of brisk walking over four weeks on glucose metabolism, insulin sensitivity, and metabolic flexibility in free-living overweight and obese male and female adults. To distinguish the metabolic effects of breaking up sedentary behavior from time spent active, this intervention was compared to a second 4-weeks one consisting in the performance of brisk walking as a single continuous bout matched for total active time.
2. METHODS

2.1 Participants

Eligible participants were males and females (premenopausal follicle-stimulating hormone $\leq 30$ mIU/mL), between ages 18-45 years old, with a body mass index (BMI) of 25-35 kg/m$^2$, weight stable for the 6 months leading up to study (+/- 3 kg), physically inactive (self-reported < 150-min/week of moderate to vigorous physical activity), who walked less than 6,500 steps per day (measured over 7 days via pedometer in free-living conditions) and who were sedentary (self-reported > 9 h sitting per day). Exclusion criteria includes history of renal, cardiovascular, or hepatic diseases, type 1 or type 2 Diabetes, cancer, smoking, consumption of drugs (>40 g/d), HIV positive, psychiatric disorders, or any medications that interfere with energy metabolism. Participants were recruited via newspapers and online advertisements, flyers, and emails. All subjects signed a written informed consent. This study was performed in accordance with the Declaration of Helsinki and approved by the Colorado Multiple Institutional Review Board.

2.2 Study Design and Procedures

2.2.1 Overall study design
Following a screening visit and baseline measures, participants were randomly assigned to one of two 4-week intervention groups. Participants consumed their normal diets throughout the intervention with the exception of three days prior to both pre and post chamber visits, during which time meals were provided. In pre and post intervention, insulin sensitivity was measured by an intra-venous glucose tolerance test (IVGTT), and nutrient metabolism and metabolic flexibility were measured during a 24h stay in a whole room calorimeter. During two weeks prior to the intervention and during the final two weeks of the intervention, daily glycemic variability was assessed by continuous glucose monitoring, total energy expenditure (TEE) and activity energy expenditure (AEE) were measured by the combination of doubly labelled water (DLW) and indirect calorimetry, time spent in light, moderate and vigorous physical activity and time spent sedentary were measured by 3D-accelerometry. All study visits were done at the University of Colorado Hospital Clinical Trial Research Center (CTRC). Overview of the study design is summarized in Figure 1.

**Figure 1.** Overall study design
2.2.2 Screening visit

During the screening visit, participants were screened for exclusion criteria. Medical history was obtained, and a physical examination were conducted. Fasting blood sample was drawn. The International Physical Activity Questionnaire (IPAQ) was completed to inform on participant’s habitual physical activity. Total body mass was measured with a calibrated scale and body composition was determined using Dual energy X-ray absorptiometry (DXA, Hologic Delphi-W, Bedford, MA). Pedometers were given to the participants to record their daily steps for 7 days to determine their average daily steps.

2.2.3 Physical Activity Interventions

Participants were assigned to one of the two 4 week-interventions, which were as follows:

- **One bout of traditional exercise (ONE):** Participants were asked to perform moderate-intensity activity (brisk walking) for 45min/day, 5 days/week. Exercise days and time of day were determined by the participant.

- **Microbouts of exercise (MICRO):** Participants were asked to break up prolonged sitting time by completing 5 minutes of moderate-intensity activity (brisk walking) each hour for 10 consecutive hours, 5 days/week. Exercise days and the 10 hours containing a microbout were also selected by the participant. The participants were asked to complete 10 bouts and not 9, because preliminary data showed that subjects performed 10% less of the physical activity when instructed to perform microbouts. The goal was therefore o
have them completing 45-min if moderate intensity activity per week, like participants in the ONE group.

To ensure compliance, each subject was asked to wear an activity tracker (Charge 2, FitBit, San Francisco, CA) at all time except during water-based activities. Every day, the subject downloaded the recorded data on their personal account created by the research team. If the subject was not reaching the instructed level of activity required, a member of the research team was contacting the subject to provide advices and help finding strategy to implement the interventions into their daily life. An example of the data retrieved by the FitBit is shown in Figure 2.

![Figure 2](image)

**Figure 2.** Example of the FitBit data obtained during a day with microbouts of activity to monitor compliance in real-time.
2.2.4 Physical Activity and Sedentary Behavior

For 14 days before the start of the intervention and during the last two weeks of the intervention, daily time spent sitting, standing and stepping was monitored via ActivPAL activity monitor (PAL Technologies: Glasgow, Scotland) on the participants right leg, and Actigraph activity monitor (Actigraph GT3X+, Fort Walton Beach, FL) worn around the participant’s waist. Data from the monitors was time stamped and includes time spent sitting/lying, standing, and stepping per day. Using a customized R program\textsuperscript{15}, the data is converted into a second by second file that can be used to determine time stepping, time sitting, time lying, breaks in sedentary time, number of daily steps, and duration of bouts\textsuperscript{16,17}. Time sitting and lying did not include time sleeping, only waking time, by using daily journals participants were given to record bedtime and waketime.

2.2.5 Resting Metabolic Rate, Total and Activity Energy Expenditure

TEE was measured for two weeks prior to the start of the intervention and during the final 2 weeks of the intervention using the doubly-labeled water (DLW) method as previously described\textsuperscript{18}. Subjects ingested a premixed 2g/kg total body water (TBW, estimated as 73.2\% of fat free mass) dose of DLW composed of 0.2 and 0.15 g/kg estimated TBW of H\textsubscript{2}\textsuperscript{18}O and \textsuperscript{2}H\textsubscript{2}O, respectively. Urine samples were collected on days 0 (+4 and +5 hours post dose), 1, 4, 6, 8, 12, 14, 14+1hr. \textsuperscript{2}H and \textsuperscript{18}O enrichments were determined in urine samples by isotope ratio mass spectrometry (IRMS, Delta V Advantage, Thermo, Germany). TEE was calculated in MJ/d using
Schoeller’s equation \(^{19,20}\). The food quotient was assumed to be equal to 24h respiratory quotient (RQ) measured in resting condition in the whole room calorimeter.

AEE was calculated using the following equation:

\[
AEE (MJ/d) = TEE (MJ/d) - TEF (MJ/d) - RMR (MJ/d);
\]

Where RMR is resting metabolic rate and was determined using a metabolic cart (ParvoMedics, for 1hr after an overnight fast)\(^{21}\), and TEF is the thermic effect of food that was estimated to be 10% of TEE (reference needed).

2.2.6 Body Composition

Body composition was measured in pre and post by isotopic dilution as previously described \(^{22}\).

2.2.7 Daily Glycemic Variability

To measure the daily glycemic variability in free living conditions, a constant glucose monitoring system (CGMS) was worn by participants for 14 consecutive days during pre-intervention baseline measurements and during the final two weeks of the intervention. The CGMS (FreeStyle LibrePro, Abbott) is a small adhesive patch containing a micro-needle that is inserted subcutaneously into the triceps. Blood glucose is collected and stored every 15 minutes for the two weeks, and once completed, the monitor is scanned via Bluetooth and data is retrieved and stored electronically. Data was analyzed using a R software version 3.2.1 (www.rproject.org), to calculate various
glycemic variability outcomes\textsuperscript{8,15}. Mean glucose (in mg/dl) is the arithmetic mean of all raw CGM values. Standard deviation (SD) is the standard deviation of all raw CGMS glucose values taken. Mean amplitude of glucose excursions (MAGE) is calculated as the arithmetic average of absolute value differences between adjacent glucose peaks and nadirs, where differences exceed 1 SD from the mean. Continuous overall net glycemic action (CONGA) represents the SD of all valid differences between a current observation and an observation (n) hours earlier\textsuperscript{23}.

2.2.8. Run-in period diet and physical activity prior to the 24h stay in the whole room calorimeter and during the study visit

All the meals were provided by the University of Colorado Anschutz Metabolic Kitchen. A three-day run-in diet was provided to the subjects during their run-in period, 3 days prior to their chamber stay, in both pre and post intervention. The macronutrient composition of the diet was 30\% fat, 55\% carbohydrates, and 15\% protein of total energy intake. Percent energy per meal was as 30\% breakfast, 35\% lunch, 35\% dinner. Total energy was calculated to match their free living energy requirements by using resting metabolic rate (RMR) which was derived from the average of (1) direct measurement via hood indirect calorimetry and (2) the equation: \[ 24hr intake = (23.9 \times FFM \text{ in kg}) + 372, \] dual energy X-ray absorptiometry (DXA), and an activity factor estimated from the free living activity of the participants (ActivPAL data, IPAQ data and pedometer data). During the run-in period, participants were instructed to perform the physical activity regimen they were assigned to including the day prior to the study visit.

The assessment of energy needs in the chamber used the same equation but an activity factor of 1.5 was applied to all our subjects based on data from prior studies including chamber visits\textsuperscript{11,24}.  

10
Macronutrient composition was (65%, 20% and 15% of total energy intake for CHO, fat and protein respectively). Breakfast, lunch, dinner and snack respectively represented, 30%, 30%, 30% and 10% of total energy intake. To avoid deconditioning while confined in the chamber, subjects were asked to perform three bouts of stepping to mimic their baseline physical activity. The duration of the stepping bout was adjusted individually based on their daily steps recorded in free living conditions by the activity monitors. The estimated energy expenditure induced by stepping was added to the total estimated energy intake. Duration of stepping bout as well as total energy intake were similar in pre and post intervention.

2.2.9 Twenty-four hour chamber visit

Participants reported to CTRC at 0700h, voided and were weighed. An IV catheter was placed into the antecubital vein for collecting blood samples, and baseline breath tests were collected. Subjects then entered the whole-room calorimeter at 0800 for a 23-h stay. Participants remained seated and quiet for one hour to achieve steady state within the calorimeter. Breakfast, lunch, dinner, and snack were provided at 0900h, 1300h, 1830h, and 2130h with the same macronutrient composition and energy intake as the run in diet. Four grams of a U-13 stable isotope tracer were given in the breakfast meal to measure exogenous carbohydrate oxidation. CO₂ measurements and blood samples were collected every hour for 14 consecutive hours (0900h-2200h), and again at 0600 the following morning. Urine was collected in one jug for the “waking time” (0900h-2230h), and a different jug for “sleep-time” (2230h-0600h). Participant exited the room at 0700 the following morning. Stepping bouts were performed at 1100h, 1500h, and 1900h using an aerobic step and stepping 72 beats per minute.
2.2.10. Twenty-four total and exogenous substrate oxidation

O₂ consumption (VO₂) and CO₂ production (VCO₂) were determined from the flow rates and differences in gas concentrations between air entering and air exiting the calorimeter as previously described\textsuperscript{25}, providing respiratory quotient (RQ) values. To isolate the non-protein respiratory quotient (npRQ), urinary nitrogen excretion was used.

Participants also collected an hourly breath sample for \(^{13}\)CO₂ by blowing through a tube into two 15 ml vacutainer tubes. Breath CO₂ was sampled directly from the vacutainer with a syringe, and \(^{13}\)CO₂/\(^{12}\)CO₂ was measured with isotopic ratio mass spectrometer (IRMS, Delta V, Thermo Electron, Bremen, Germany). The average baseline enrichment value was subtracted from the enrichment relative to the subject’s own baseline. By using time matched CO₂ production rates from the whole room calorimeter, \(^{13}\)C glucose was calculated as the instantaneous percentage recovery of \(^{13}\)C in expired CO₂ per hour for 14-h and after 24-h.

2.2.11. Insulin Sensitivity

Whole body insulin sensitivity was determined via ‘intravenous glucose tolerance test’ (IVGTT). Participants reported to the CTRC at 0700h after an overnight fast. Participants were instructed to maintain physical activity three days prior to the test. They were also asked to eat a high carb
diet the night prior that contained either pasta, potatoes, rice or white bread (examples of meals were provided to the participants). They were asked to take pictures and write down in a dietary log the exact meal they consumed in pre intervention and to have the exact same meal post-intervention. For the IVGTT, two antecubital catheters were inserted, one on one arm for infusion of exogenous glucose and insulin, and one on the other arm for blood draws. After baseline samples were obtained at minute 0, intravenous glucose (0.3 g/kg) was infused over 1 minute and samples were collected at minutes 2, 3, 4, 5, 6, 8, 10, 12, 14, 16, and 19. After 19 minutes, 0.03U/kg of insulin was administered and blood samples were collected at minutes 22, 23, 24, 25, 27, 30, 40, 50, 70, 90, 100, 120, 140, 160, and 180.

2.2.12. Plasma metabolite and hormones analysis

Whole blood was added to a preservative (3.6 mg EDTA plus 2.4 mg glutathione in distilled water). Plasma and serum were separated after spinning and stored at -80°C until analyzed. EDTA plasma samples were analyzed for glucose and insulin. Insulin concentrations were measured using a standard double antibody radioimmunoassay (EMD Millipore, St Charles, Missouri). Serum glucose concentrations were determined using the hexokinase method (reference).

2.3 Data and Statistical Analysis

2.3.1. Metabolic Flexibility
Metabolic flexibility, i.e. shift from carbohydrate oxidation during the day (while feeding) to fat oxidation at night (while fasting), was assessed as the delta between the waking npRQ and sleeping npRQ values.

\[
\text{Metabolic flexibility} = \Delta \text{npRQ} = \text{waking npRQ} - \text{sleeping npRQ}
\]

A metabolically flexible individual utilizes primarily carbohydrates during the daytime as they are feeding, and oxidize primarily fat during fasting hours at night. Therefore, a metabolically flexible individual who is capable of a large change in substrate utilization based on availability of the course of 24 hours, have a greater arithmetic difference between waking npRQ and fasting npRQ than someone who is metabolically inflexible.

2.3.2. Insulin Sensitivity Analysis

The minimal model analysis by Richard Bergman’s MinMod program (MINMOD 6.02, MinMod, Los Angeles, CA, USA) has been applied to various insulin resistant states, including obesity, gestational diabetes, experimental hyperinsulinemia, and insulin dependent diabetes. It uses a mathematical model to examine the closed system feedback loop of the body in response to an IVGTT (Figure 3). Outcome variables include insulin sensitivity (SI) which represents the effect of insulin to accelerate the ability of glucose to enhance glucose disappearance from extracellular fluid via suppression of endogenous glucose output and increased glucose disposal. The first phase of the body’s response to exogenous glucose can be measured via the Acute
Insulin Response to glucose (AIRg). This is found by calculating the area under the curve from the start of the mixing phase up until the exogenous insulin is pushed, and is indicative and the body’s endogenous response to glucose. Lastly, is glucose effectiveness (Sg), or the ability of glucose to clear itself without the presence of insulin. This is hypothesized to be via uptake in the brain. However, it is known that Beta cells can compensate for lack of insulin sensitivity by upregulating insulin secretion; using the fourth variable called Disposition Index (DI) which is the product of SI x AIRg. If an individual’s endogenous organs become less insulin sensitive pancreatic beta cells will compensate for this by releasing more insulin; this produces a larger AIRg which will compensate for the decrease in SI and allow a healthy individual to maintain their DI. A DI that changes overtime shows the lack of ability to compensate and becomes a risk factor for developing diabetes.

Figure 3. Example of the plasma glucose and insulin concentrations during an IVGTT. Blue represents blood glucose post exogenous bolus is given in minute 1 Green represents blood
insulin (the first peak representing endogenous insulin in response to glucose, the second peak representing exogenous insulin being given at minute 19).

2.3.3. Statistical analysis

Changes (pre vs post) in 24 hour sitting/lying, stepping and standing time, waking sedentary time, 24hr TEE and AEE, RMR, body weight and composition, daily glycemic variability, whole body insulin sensitivity, 24hr exogenous CHO oxidation, 24h npRQ and metabolic flexibility were analyzed using linear mixed model with time as repeated effect, group and group by time interaction as main effect and subjects as random effect with compound symmetry as covariance. Associations between primary outcomes were determined using linear mixed model as well. Mean and standard deviation (SD) were reported unless stated otherwise. Significance was set up at  p<0.05 except for the time-by-group interaction for which it was set up at 0.1. Statistical analysis were performed with SPSS software (SPSS v.22, SAS, IBM).
3. RESULTS

3.1 Participant Characteristics

To date, nineteen sedentary adults (n=7M/12F; 33.0±8.3 years; BMI=29.9±3.0 kg/m²) out of 28 participants completed the study. There were no differences at baseline among groups, and participant’s characteristics are displayed in Table 1.

Table 1. Anthropometric characteristics of study participants

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<th>MICRO 3M/6F</th>
<th>ONE 4M/6F</th>
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<tr>
<td>Sex</td>
<td>3M/6F</td>
<td>4M/6F</td>
</tr>
<tr>
<td>Age (year)</td>
<td>32.8 ± 10.0</td>
<td>33.1 ± 6.5</td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>87.9 ± 12.7</td>
<td>80.2 ± 10.7</td>
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<tr>
<td>Lean body mass (kg)</td>
<td>51.3 ± 8.3</td>
<td>50.1 ± 9.3</td>
</tr>
<tr>
<td>% Fat Mass</td>
<td>41.3 ±7.2</td>
<td>31.1 ±7.4</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>31.1 ± 3.9</td>
<td>28.1 ± 2.1</td>
</tr>
<tr>
<td>Step counts (pedometer)</td>
<td>4782 ± 993</td>
<td>4954 ± 1286</td>
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</table>

Mean (±SD)

3.2 Changes in sedentary behavior, physical activity, energy expenditure and body composition

At baseline, participants in ONE spent more time stepping than those in MICRO (p<0.05). No other baseline difference was observed. Stepping time over 24hr, sitting/lying time over 24hr (sedentary time), and TEE are shown in Figure 2. As designed, both ONE (1.12±0.52 versus 1.77±0.88 hrs) and MICRO (1.91±0.74 versus 2.19±0.86 hrs) increased time spent stepping over 24hours. Neither MICRO nor ONE modified time spent sedentary over waking hours, sitting/lying over 24hr and standing. Similarly, no significant changes were observed in TEE,
AEE and physical activity level (PAL). RMR tended to increase (p=0.07) after four weeks of active intervention but was no longer modified when the slight increase in FFM (p=0.07) was taken into account. Body and fat mass were maintained, indicating that subjects were in stable energy balance. No between-group nor interaction were observed (Figure 4).

Table 2. Effects of 4 weeks of MICRO versus ONE on body composition, RMR, TEE, AEE, PAL measured by doubly labeled water and a metabolic cart

<table>
<thead>
<tr>
<th></th>
<th>MICRO</th>
<th>ONE</th>
<th>Group effect</th>
<th>Intervention effect</th>
<th>Group-by-Intervention effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PRE</td>
<td>POST</td>
<td>PRE</td>
<td>POST</td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td>84.5 ± 13.8</td>
<td>84.6 ± 14.5</td>
<td>78.1 ± 10.0</td>
<td>79.8 ± 10.0</td>
<td>0.4</td>
</tr>
<tr>
<td>Fat mass (kg)</td>
<td>34.1 ± 12.1</td>
<td>33.0 ± 12.7</td>
<td>26.9 ± 5.6</td>
<td>27.0 ± 4.6</td>
<td>0.2</td>
</tr>
<tr>
<td>Fat free mass (kg)</td>
<td>50.4 ± 6.1</td>
<td>51.6 ± 6.4</td>
<td>51.3 ± 10.0</td>
<td>52.8 ± 10.1</td>
<td>0.8</td>
</tr>
<tr>
<td>Percent body fat (%)</td>
<td>39.5 ± 8.6</td>
<td>38.1 ± 8.6</td>
<td>34.7 ± 7.0</td>
<td>34.1 ± 6.4</td>
<td>0.3</td>
</tr>
<tr>
<td>RMR (MJ/d)</td>
<td>6.43 ± 0.7</td>
<td>6.64 ± 0.93</td>
<td>6.59 ± 1.1</td>
<td>6.91 ± 1.1</td>
<td>0.6</td>
</tr>
<tr>
<td>RMR/FFM (MJ/kg)</td>
<td>0.13 ± 0.01</td>
<td>0.13 ± 0.01</td>
<td>0.13 ± 0.01</td>
<td>0.13 ± 0.02</td>
<td>1.0</td>
</tr>
<tr>
<td>TEE (MJ/d)</td>
<td>10.8 ± 2.1</td>
<td>10.7 ± 1.6</td>
<td>11.6 ± 3.3</td>
<td>12.8 ± 3.5</td>
<td>0.2</td>
</tr>
<tr>
<td>TEE/FFM (MJ/kg)</td>
<td>0.21 ± 0.04</td>
<td>0.21 ± 0.02</td>
<td>0.22 ± 0.03</td>
<td>0.24 ± 0.03</td>
<td>0.14</td>
</tr>
<tr>
<td>AEE (MJ/d)</td>
<td>4.16 ± 1.8</td>
<td>4.10 ± 0.82</td>
<td>5.02 ± 2.4</td>
<td>5.95 ± 2.9</td>
<td>0.14</td>
</tr>
<tr>
<td>AEE/FFM (MJ/kg)</td>
<td>0.09 ± 0.04</td>
<td>0.08 ± 0.01</td>
<td>0.09 ± 0.03</td>
<td>0.11 ± 0.04</td>
<td>0.2</td>
</tr>
<tr>
<td>PAL (TEE/RMR)</td>
<td>1.64 ± 0.26</td>
<td>1.62 ± 0.09</td>
<td>1.76 ± 0.27</td>
<td>1.86 ± 0.38</td>
<td>0.09</td>
</tr>
</tbody>
</table>

Mean ±SD
**1 MICRO and 2 ONE group participants were excluded from this data set due to inability to analyze doubly labeled water data**

**Figure 4.** Waking sitting, standing, stepping and doubly-labelled water-derived total energy expenditure (TEE), before (PRE) and during the last two weeks of a 4-week intervention (POST) consisting of either nine 5-min brisk walking bouts per day, 5 d/wk (n=9, MICRO) or of 45-min brisk walking performed once per day, 5 d/wk (n=10, ONE).
### 3.3 Daily Glycemic Variability

Data obtained over 14 days from the CGMS in pre and during the last two weeks of the intervention are provided in Table 3. No between-group differences at baseline were observed. Mean, SD, MAGE, and CONGA did not differ between the groups. Intervention-by-group interactions were observed for SD, MAGE, and CONGA (p<0.1 for all). While MICRO decreased SD from 0.95 ±0.28 to 0.81±0.21 mg/dl, ONE increased SD from 0.78±0.14 to 0.81±0.16 (p=0.06). MICRO decreased MAGE from 2.4±0.9 to 2.0±0.5, ONE increased MAGE from 1.9±0.5 to 2.2±0.5 (p=0.03). MICRO increased CONGA from 3.9±0.4 to 4.3±0.7, ONE decreased CONGA from 4.2±0.6 to 4.0±0.7 (p=0.07). Daily mean glucose was not modified by either of the active interventions.

#### Table 3. Effects of 4 weeks of MICRO versus ONE on 24hr glycemia measured by CGMS

<table>
<thead>
<tr>
<th></th>
<th>MICRO</th>
<th>ONE</th>
<th>Group-by-Intervention effect</th>
<th>Group effect</th>
<th>Interventions effect</th>
<th>Group-by-Intervention effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PRE POST</td>
<td>PRE POST</td>
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<tr>
<td>n</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean glucose (mg/dl)</td>
<td>4.7 ± 0.44</td>
<td>5.0 ± 0.78</td>
<td>4.9 ± 0.60</td>
<td>4.7 ± 0.79</td>
<td>0.9</td>
<td>1</td>
</tr>
<tr>
<td>SD (mg/dl)</td>
<td>0.95 ± 0.28</td>
<td>0.81 ± 0.21</td>
<td>0.78 ± 0.14</td>
<td>0.81 ± 0.16</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>MAGE</td>
<td>2.4 ± 0.89</td>
<td>2.0 ± 0.49</td>
<td>1.9 ± 0.45</td>
<td>2.2 ± 0.51</td>
<td>0.6</td>
<td>0.3</td>
</tr>
<tr>
<td>CONGA</td>
<td>3.9 ± 0.44</td>
<td>4.3 ± 0.69</td>
<td>4.2 ± 0.55</td>
<td>4.0 ± 0.66</td>
<td>0.9</td>
<td>0.8</td>
</tr>
</tbody>
</table>

Mean ±SD, Standard deviation (SD), mean amplitude of glucose excursions (MAGE), continuous overall net glycemic action (CONGA)

**1 ONE group participant was excluded due to loss of CGMS monitor by participant**
3.4 Insulin Sensitivity

Data from the IVGTT are reported in Table 4. No between-group differences at baseline were observed. No group effect was observed in SI, AIRg and DI. Intervention-by-group interaction were observed for SI (p=0.08) and AIRg (p=0.02). While MICRO increased SI from 5.5 (±3.2) [mU/l]^{-1} min^{-1} to 11.3 (±12.5), ONE decreased SI from 6.6 (±5.2) to 5.8 (±4.5). While MICRO increased AIRg from 603.9 (441.7) mU/l × min to 470.3 (±560.1), ONE decreased AIRg from 673.2 (±419.9) to 810.5 (±573.1). None of the physical activity interventions modified DI.

Although the activity interventions did not change Sg, it tended to be lower in MICRO than in ONE (group effect: p=0.08) across the intervention period.

Table 4. Effects of 4 weeks of MICRO versus ONE on whole body insulin sensitivity measured by IVGTT

<table>
<thead>
<tr>
<th></th>
<th>MICRO</th>
<th>ONE</th>
<th>Group</th>
<th>Intervention</th>
<th>Group-by-Intervention</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>PRE</td>
<td>POST</td>
<td>PRE</td>
<td>POST</td>
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<tr>
<td>n</td>
<td>9</td>
<td>9</td>
<td>10</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>SI (mU/l^{-1} min^{-1})</td>
<td>5.5 ± 3.2</td>
<td>11.3 ± 12.5</td>
<td>6.6 ± 5.2</td>
<td>5.8 ± 4.5</td>
<td>0.4</td>
</tr>
<tr>
<td>AIRg (mU/l × min)</td>
<td>603.9 ± 441.7</td>
<td>470.2 ± 560.1</td>
<td>673.2 ± 419.9</td>
<td>810.5 ± 573.1</td>
<td>0.4</td>
</tr>
<tr>
<td>DI</td>
<td>2549.1 ± 1421.8</td>
<td>2307.2 ± 1991.2</td>
<td>3428.2 ± 1656.7</td>
<td>3507.0 ± 2111.9</td>
<td>0.18</td>
</tr>
<tr>
<td>Sg (min^{-1})</td>
<td>0.021 ± 0.012</td>
<td>0.015 ± 0.010</td>
<td>0.029 ± 0.013</td>
<td>0.025 ± 0.012#</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Mean ±SD, #p=0.07 vs. MICRO; IVGTT, Intra-venous glucose tolerance test; MICRO, 4 weeks of microbouts of activity (9 bouts of 5-min brisk walking, 5 d/wk); ONE, 4 weeks of single continuous 45-min of brisk walking (5d/wk)
3.5 Metabolic Flexibility and Substrate Utilization

No between-group difference was observed at baseline. A group-by-intervention interaction was observed for the index of metabolic flexibility, i.e. delta between day npRQ and night npRQ; while it increased in MICRO, it did not change in ONE (p=0.03; Figure 5). While MICRO (post: 59.5±7.0% versus pre: 46.3±8.9% recovery of ingested dose) maintained 24h exogenous carbohydrate oxidation at baseline levels, ONE decreased it (post: 59.9±5.4% versus pre: 50.4±7.8% recovery of ingested dose; group-by-intervention interaction: p=0.007).

**Figure 5.** Effect of 4 weeks of microbouts of activity (MICRO, n=9) vs single continuous bouts of activity (ONE, n=10) matched for total active time (45-min), on 24h exogenous carbohydrate oxidation and metabolic flexibility in overweight/obese adults.
4. DISCUSSION

In this longitudinal randomized intervention study, we showed that four weeks of frequent interruptions of sedentary time with short bouts of brisk walking performed each hour for nine consecutive hours reduce glycemic variability, improve whole body insulin sensitivity, and improve metabolic flexibility to a greater extent than four weeks of intervention consisting of a single bout of brisk walking matched for total active time. Importantly, these differential metabolic responses were observed while both interventions had similar impact on the pattern of physical activity, i.e. increase in time spent stepping and MVPA, decrease in time spent sitting and light-intensity activity, and no change in AEE or TEE. These results indicate that the beneficial effects of breaking up sedentary behaviors are independent of changes in time spent active and sedentary or changes in energy expenditure.

Our study measured free living blood glucose concentration and fluctuations over the course of 14 days using constant blood glucose monitoring systems prior to and in the final two weeks of the exercise intervention. We found that frequent brisk walking bouts spread throughout the day decrease glycemic variability over 24 hours when compared to the performance of a single continuous bout of brisk walking. Blankenship et al. also used constant glucose monitoring systems to study frequent bouts of physical activity in adults with type 2 diabetes. They found that when active time was matched to 30-minutes, breaks from sitting with moderate intensity walking bout were more effective at decreasing glycemic variability than the performance of moderate intensity walking as a continuous bout. One of the benefits to constant glucose monitoring systems is the ability to examine glucose values and changes in free-living conditions, during rest days, and during sleeping hours. Dempsey et al. used this
monitoring system to examine the effects of glucose values in type 2 diabetics after completing a seven hour intervention of either 3 minutes of walking every 30 minutes, 3 minutes of resistance exercise every 30 minutes, or sedentary for 7 hours, and continued to monitor them for a total of 22 hours. They found that the repeated walking bouts and resistance exercises significantly decreased blood glucose during nocturnal hours compared to the sedentary control, showing that this type of intervention has acute lasting effects that limit the episodes of hyperglycemia in type 2 diabetics. Our study however didn’t show any significant decrease in mean blood glucose during wake or sleep times (data not shown) meaning maybe it is a temporary effect that dissipates with an intervention as long as ours. Importantly, our data took into account both the days during which participants performed the intervention but also the days during which they did not. Future analysis will have to only consider only the “active days” to examine the effect of the interventions on mean blood glucose. The decrease in glycemic variability with the microbouts of activity may therefore be even more pronounced when only looking at the active days. This has important clinical implications because high fluctuations of glycemia is a recognized risk factor for type 2 diabetes and cardiovascular disease.

Although no change in mean glycemia was detected over 24h (day or night), we observed a decrease in post prandial glucose concentration (data not shown) in response to standard breakfast, lunch and dinner. This result is in line with an increasing number of intervention that observed a decrease in postprandial glycemia in response to an acute exposure to the breaks. Importantly, our data suggests that breaking up sedentary behaviors improves glucose control not only acutely, but also when participants are resting, after 4 weeks of intervention. We recently showed that the decrease in post-prandial glycemic variability induced by 4 days of breaking was associated with a greater use of carbohydrate oxidation for 24 hours.
not observe an increase in 24h total carbohydrate oxidation (data not shown), but a maintenance of 24h exogenous carbohydrate oxidation, contrary to the performance of 4 weeks of continuous bout of brisk walking that decreased it. De Jong attributed this effect to the fact that bouts of activity are effective at training the body to preferentially use glucose as they believe the reason to be that glycogen at the skeletal muscle was being utilized due to increased translocation of the GLUT4 transporter which triggered uptake and oxidation of glucose. Once again, it is important to note that this greater use of dietary carbohydrate as fuel was observed when subjects were rested, and not exercising, suggesting that other adaptations come at play going beyond the translocation of the GLUT4 transporter to facilitate glucose uptake. In addition, the frequent bout group had an improved metabolic flexibility compared to the single bout. This means that they were essentially utilizing carbohydrates as substrates during the feeding/waking hours (npRQ closer to 1.0) but mainly relied upon fat as fuel during sleeping/fasting hours (npRQ closer to 0.7). In previous study, our group showed that physical inactivity induced by bed rest trigger the development of metabolic inflexibility, which precedes the development of glucose intolerance. Our present data suggest that the lack of frequent muscle contraction, use of energy or use of carbohydrate contribute to this metabolic inflexibility rather than physical inactivity per se, i.e. fact of not reaching the recommended level of physical activity. Future studies will be warranted to confirm this hypothesis.

Understanding effective ways to treat metabolic inflexibility is important as it is a common characteristic of type 2 diabetics, insulin resistance, obesity, and metabolic syndrome. Here we observed a positive correlation between the 4-week changes in whole-body metabolic flexibility and in insulin sensitivity. The directionality between the two however remains unknown, and will need to be investigated. Our study showed that frequent bouts of physical activity improve
whole body insulin sensitivity as well as in the acute insulin response to glucose in overweight/obese adults in a greater extent than what is observed when physical activity is performed as a single continuous bout. This is in line with the observations from other studies of the acute effects of breaking up sedentary behavior on post prandial insulinemia which suggests an improvement in insulin sensitivity with breaking up sedentary behavior. In this line, our study also showed improved insulin sensitivity post prandially (data not shown).

While there are many valid ways measure insulin sensitivity, our study used IVGTTs, however the choice to not use an hyperinsulinemic euglycemic clamp or oral glucose tolerance test (OGTT) did come with downfalls, as each test has it’s own variety of pros and cons. While a hyperinsulinemic euglycemic clamp is technically the “gold standard” for measuring insulin sensitivity, and has the ability to show tissue-specific changes in insulin sensitivity, the IVGTT is a shorter test at around 4 hours total compared to around 7 hours of a clamp. IVGTTs are also less costly were within the scope of funding for this study, and do not require a physician present (making participant scheduling easier and compliance better). They are validated against the hyperinsulinemic clamp when using the Minimal Model program for analysis. Another validated method that could have been used and had outcomes that the IVGTT did not was the OGTT. This test is also less costly than a glucose clamp, and is also more effective at classifying participants into diabetic or normal as the estimated efficiency of glucose disposal rates is better shown with OGTTs with participants with mild glucose tolerance abnormalities when compared to the IVGTT. The IVGTT was chosen over this however as when it is paired with minimal model, it still gives a valid insulin sensitivity value as well as the DI and AIRG which the OGTT does not.
The improvement in AIRg suggest that frequent bouts of physical activity lead to an acute release of insulin in response to exogenous glucose load when compared to the single bout group. The reasons for this remain unknown but there are two hypothesis, either (1) a better pancreatic function leading to a quick insulin secretion or (2) the presence of insulin in the beta cell ready to be quickly released into the blood stream. To determine this mechanism, future studies will need to test the effects of breaking up sedentary behavior on beta cell function by using a stepwise glucose tolerance test in which c-peptide is measured.

A major strength of this study was the randomized controlled trial and the study for the first time of the sustained effects of breaking up sitting time. It also included use of a 24 hr whole room calorimeter and 3 day run in diet, as well as using active-time matched controls. Our limitations include the fact that there was a lack of beneficial changes shown in the single bout group. However, we believe that this is due to the fact that this group did not increase their overall total energy expenditure. We believe that because they adopted exercise into their daily routine, they were inadvertently doing less physical activity during the rest of the waking hours. Other limitations include the fact that some participants either gained or lost weight throughout the study rather than remaining weight stable, which could have affected our main outcomes. With this being said, while there were changes in individuals (weight gain and weight loss), there were no significant differences on average in body weight or fat mass in neither group. While there is not enough statistical power with our 19 participants currently to run post-hoc tests to determine the pre/post changes in each group and the between-group differences in post. However, upon completion of the study, changes in energy balance, body weight, and body composition will be taken into account in the definitive statistical analysis. IVGTTs were also conducted at 0700 for each participant, however recent studies have shown that insulin
sensitivity is associated with circadian rhythms and therefore the tests should have been conducted in relation to each of the participants natural wake time. Finally, when determining 24 hour energy intake to feed the participants in their 3 day run in diet and the chamber, we had to multiply RMR with an activity factor. This activity factor was estimated based on pedometers, ActivPAL, IPAQ, but was not objectively measured. Applying an activity factor of 1.5 or 1.6 automatically induce a 10% difference in energy intake and potentially energy balance, which can also affect the results.

Conclusions and future perspectives

In conclusion, we showed that 4 weeks of microbouts of physical activity result in decreased 24hr glycemic variability, which is associated with improved metabolic flexibility, high levels of exogenous carbohydrate oxidation, and increased whole-body insulin sensitivity, compared to the performance of 4 weeks of single continuous bout of brisk walking matched for total active time. It is important to note however that these findings are specific to sedentary overweight and obese adults and is aiming to find an intervention that can compare to the traditional exercise recommendations of 150 minutes of MVPA, not be overall better. The number one indicator of premature death is lack of cardiorespiratory fitness and our frequent bouts of physical activity does not increase this, however for some of the US population, our intervention could be effective to combat common barriers of to exercise (time, knowledge, enjoyability, etc), and be better than nothing at all. It is also designed specifically with individuals who have high glycemia, and would be used as an intervention specific to this issue rather than a normal or healthy individual. Ideally, we believe that the perfect intervention for health of our study population and the general public would be breaking up sedentary behavior.
with the addition of 150 minutes of MVPA. Future studies will be needed to further our understanding of the health effect of physical activity interventions varying in terms of dose, timing, duration, etc, so that more specific guidelines on physical activity will be developed. In addition, future research should also be conducted to help determine the most effective way to implement frequent bouts of exercise into daily life. It’s been shown that even when reaching 150 minutes of moderate to vigorous physical activity, it does not protect an individual from cardiometabolic risk factors if they are sedentary otherwise\textsuperscript{10}; while we and others have shown that breaking up sedentary behavior is beneficial on these same biomarkers \textsuperscript{6,11,35}. However, the minimum amount of time active in order to offset sedentary behavior still remains unknown. Determining this, or the maximum amount of time that one can remain sedentary until walking without having detrimental health outcomes, would provide extremely valuable knowledge for public health and clinical purposes and the ACSM physical activity guidelines could be made more specific.
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