Antioxidant Vitamin C Prevents Decline in Endothelial Function during Sitting

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Background: This study was designed to test the hypothesis that antioxidant Vitamin C prevents the impairment of endothelial function during prolonged sitting.

Material/Methods: Eleven men (24.2±4.4 yrs) participated in 2 randomized 3-h sitting trials. In the sitting without vitamin C (SIT) and the sitting with vitamin C (VIT) trial, participants were seated for 3 h without moving their legs. Additionally, in the VIT trial, participants ingested 2 vitamin C tablets (1 g and 500 mg) at 30 min and 1 h 30 min, respectively. Superficial femoral artery (SFA) flow-mediated dilation (FMD) was measured hourly for 3 h.

Results: By a 1-way ANOVA, there was a significant decline in FMD during 3 h of SIT (p<0.001). Simultaneously, there was a significant decline in antegrade (p=0.04) and mean (0.037) shear rates. For the SIT and VIT trials by a 2-way (trial x time) repeated measures ANOVA, there was a significant interaction (p=0.001). Pairwise testing revealed significant between-SFA FMD in the SIT and VIT trial at each hour after baseline, showing that VIT prevented the decline in FMD 1 h (p=0.009), 2 h (p=0.016), and 3 h (p=0.004). There was no difference in the shear rates between SIT and VIT trials (p>0.05).

Conclusions: Three hours of sitting resulted in impaired SFA FMD. Antioxidant Vitamin C prevented the decline in SFA FMD, suggesting that oxidative stress may contribute to the impairment in endothelial function during sitting.

MeSH Keywords: Antioxidants • Oxidative Stress • Sedentary Lifestyle

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Background

Increased sitting time has been associated with increased risk of cardiovascular disease and mortality [1]. Prolonged sitting can create a unique physiological milieu in the lower extremity vasculature, leading to conditions conducive to the development of atherosclerosis [2]. For instance, 5 h of sitting has been shown to increase calf pooling and decrease thigh blood flow [3,4]. Greater than 3 h of sitting has also been shown to increase diastolic blood pressure (BP), mean arterial pressure (in the arm and leg), and total peripheral resistance [4,5]. Furthermore, only 2 h of sitting significantly increased whole-blood viscosity in the leg [6]. In the first experiment investigating the effects of sitting on vascular shear stress and endothelial function, Padilla et al. [5] discovered that 3 h of sitting significantly attenuated vascular shear in the popliteal artery; however, this was not associated with a concomitant decline in endothelial function measured by FMD. Contrary to this study, we recently discovered that 3 h of sitting results in a decline in mean shear rate in the SFA in addition to a significant decline in endothelial function when all measurements are made in the seated position [7]. Interestingly, in the vascular system, areas of low vascular shear stress have been associated with increased oxidative stress [8] and impaired endothelial function [8]. Additionally, low shear stress may be an etiological mechanism for atherosclerosis [9].

The human vasculature is internally coated with a single layer of cells called the endothelium, which becomes functionally impaired in the process of atherosclerosis [10]. Nitric oxide (NO), which is a critical molecule involved in maintaining the antiatherogenic properties of the endothelium [11], is synthesized from L-arginine by endothelial NO synthase (eNOS) in the presence of laminar shear stress and co-factors such as nicotinamide adenine dinucleotide phosphate (NADPH) and tetrahydrobiopterin (BH4) [12]. Shear stress associated with physical movement augments NO bioavailability [13] and thereby improves endothelial function [12]. Conversely, sitting decreases lower extremity shear rates [5,7], and hence it is possible that sitting may reduce NO bioavailability in the lower extremities. Indeed, low shear stress leads to decreased expression of NOS [8]. Our group has recently proposed that prolonged sitting may lead to low vascular shear-related impairment in endothelial function involving mechanisms related to oxidative stress [2]. We have recently address the first part of this hypothesis and shown that prolonged sitting decreases mean shear rate and impairs endothelial function in SFA [7].

The purpose of the current study was to determine if antioxidant Vitamin C prevents the decline in SFA endothelial function during prolonged sitting. We hypothesized that there would be a significant decline in SFA FMD from baseline during 3 h of sitting. Previous studies from our lab and others have demonstrated vitamin C to be effective in preventing the oxidative stress-mediated decline in endothelial function in a variety of physiological stressors, including, but not limited to, hypertension [14] and increased retrograde shear [15]. Therefore, we hypothesized that vitamin C would prevent the decline in SFA endothelial function during 3 h of sitting.

Material and Methods

Study design (Figure 1)

This study consisted of 2 screening visits and 2 sitting trials conducted in random sequence, the first with 3 h of uninterrupted sitting (SIT) and the second with oral Vitamin C during uninterrupted sitting (VIT). FMD was measured at baseline, 1 h, 2 h, and 3 h in both the trials. All procedures for the study were approved by the Indiana University Institutional Review Board, and participants gave written informed consent for their participation.

Participants

To be included, participants (all men) had to self-report that they were nonsmokers, and not taking any anti-hypertensive, lipid lowering, or anti-diabetic medications. Participants needed to have total cholesterol ≤240 mg·dl$^{-1}$ and triglycerides ≤200 mg·dl$^{-1}$ and fasting blood glucose ≤120 mg·dl$^{-1}$. We recruited individuals who performed <150 min/week$^{-1}$ of moderate-intensity physical activity or <75 min/week$^{-1}$ of vigorous-intensity physical activity [16]. Participants were asked to maintain their regular diet patterns throughout the study duration and to discontinue any over the counter antioxidant supplement at least 7 days prior to the first session of testing. Based on our pilot data, for the primary dependent variable SFA FMD, we estimated that 12 participants would be needed to find a significant difference between trials with statistical power of >0.80 at α=0.05.

Screening visits

At the first visit, all experimental procedures were explained to the participants and they were familiarized with the lab setting.
If the participants volunteered to participate in the study, a written informed consent was obtained. Height, weight, and BP (in triplicate in a seated posture) were measured using standard procedures and a medical health history and habits questionnaire [7] was completed to screen for any preexisting cardiovascular or metabolic condition and physical activity levels. BP (in triplicate) was measured during an additional screening visit at the same time of day, to confirm previous measurements.

**Testing trials**

Participants arrived at the laboratory after an overnight fast of at least 6 h between 07:00 and 09:00 h. The arrival time was matched for both trials within each subject. Participants were asked to refrain from any caffeinated drink for at least 8 h prior to reporting to the lab. Upon their arrival in the lab, all participants self-reported adhering to this dietary schedule. During both trials, the participants remained seated for 3 h without moving their legs (perpendicular to the floor) and feet (flat on the ground). Participants were allowed to move their arms, although not vigorously; for example, to use a computer or do light reading during the non-testing periods of the trials. Arm movement was not quantified. The FMD and other vascular function parameters were measured at baseline, 1 h, 2 h, and 3 h. Baseline measurements were made after ~20 min of seated rest. The 2 trials were performed in random order, separated by a minimum of 2 and a maximum of 7 days. The SIT trial was uninterrupted sitting for 3 h, whereas in the VIT trial participants ingested 2 Vitamin C tablets (with water); 1 g at 30 min and 500 mg at 1 h 30 min during the sitting interval. Doses were determined based on pilot data and were not tailored to body weight.

**SFA FMD**

SFA endothelial function was measured by FMD in accordance with current guidelines [17]. We chose SFA for 2 reasons: (1) It is a readily accessible artery to use for measuring FMD in a sitting position; and (2) SFA FMD has been shown to be largely NO-dependent [18]. Each measurement was performed in a dark, quiet, and climate-controlled room (22–25°C) room. A 5×84 cm automatic BP cuff (E-20 rapid cuff inflator; D.E. Hokanson, Bellevue, WA, USA) was placed on their right thigh about 7 cm above the knee joint, distal to the SFA recording location. Images of the SFA were obtained longitudinally 7–10 cm below the knee joint with a 2-D high-resolution ultrasound system (Terason t3000, Teratech Corp., Burlington, MA, USA), using a 5–12-MHz multifrequency linear-array transducer. Once satisfactory images of near and far arterial walls were obtained, the transducer was secured and stabilized in a stereotactic clamp, and landmarks were made on the subject's skin to ensure similar placement of the transducer for subsequent FMD procedures and shear rate assessments within and between trials. Participants were encouraged to maintain the landmarks between the 2 trials. In addition to imaging the arterial dimensions, Doppler ultrasound was used to concurrently measure SFA blood velocity. Doppler flow signals were corrected at an insonation angle of 60°, and the sample volume was placed in the middle of the artery.

Diameter images and Doppler measurements of blood velocity were continuously recorded for 45 s at baseline prior to cuff inflation. The automatic BP cuff was then rapidly inflated to 250 mmHg and maintained for 5 min until cuff deflation. Diameter and blood velocity recordings resumed prior to cuff deflation and continued for 5 min after deflation. Ultrasound images were continuously recorded at 5 frames/s−1 with Camtasia (TechSmith, Okemos, MI, USA), and stored as .avi files [7]. This procedure was repeated hourly across the sitting intervals. All measurements were made in the seated position.

**Arterial diameters and blood velocities**

Off-line analysis of diameters were performed using automated edge-detection software (Brachial Analyzer, Medical Imaging Applications LLC, Coralville, IA, USA) as previously described [7]. This software allows the technician to determine a region of interest where the near and far vessel walls are most clear. The vessel wall borders are then detected using an optimal graph search-based segmentation that uses a combination of pixel density and image gradient as an objective function. All analyzed images were reviewed by the technician and edited when needed to ensure that diameter measures were always determined from the intima-lumen interface at the near and far vessel wall. Blood velocities were determined using internally validated custom-made software by selecting a region of interest that surrounded the Doppler wave. The velocity-time integral was used to calculate the mean blood velocity. The peak dilation after cuff deflation was determined using the highest 3-s moving average and FMD was presented as a percentage change from baseline diameter (FMD%). SFA shear rate used as an estimate of arterial shear stress and was calculated for each FMD% at baseline and during the post-occlusion period using the following formula: \( V_mD \), where \( V_m \) is mean blood velocity (cm·s\(^{-1}\)) and \( D \) is mean arterial diameter (cm). All measurements and analysis were performed by a single researcher (ST) who was blinded to the participant identity and treatment condition for each image file.

**Statistical analysis**

Descriptive analysis was performed to summarize subject characteristics (Table 1). Within both trials, 1-way ANOVA was conducted on the baseline diameter as the dependent variable. Within the SIT trial, a 1-way ANOVA was conducted on the dependent variables baseline diameter, FMD%, and shear rates (antegrade shear rate, mean shear rate, retrograde shear rate, peak shear rate, and shear rate [area under the curve]) SRauc.
Table 1. Subject demographics.

| Variable            | Value     |
|---------------------|-----------|
| N                   | 11        |
| Age (yrs)           | 24.2±4.4  |
| BMI (kg·m⁻²)        | 23.6±3.4  |
| Systolic blood pressure (mmHg) | 116±10    |
| Diastolic blood pressure (mmHg) | 77.3±6    |
| Total cholesterol (mg/dl) | 171.5±32.7 |
| LDL cholesterol (mg/dl)   | 98±29.6   |
| HDL cholesterol (mg/dl)   | 57.8±13.6 |
| Triglycerides (mg/dl)    | 78±49     |
| Glucose (mg/dl)        | 91±6.5    |

Data represented as mean ±SD.

(area from deflation up until peak dilation)). When an effect was found, pairwise comparisons were used to locate significant differences across time from baseline. Observed effect size was reported for ANOVA interactions as partial eta squared ($\eta^2$). Comparisons across the 2 treatment conditions for FMD and shear rates (antegrade shear rate, mean shear rate, retrograde shear rate, peak shear rate, and shear rate area under the curve) were performed using 2-way repeated measures ANOVA, evaluating the effect of sitting on the vasculature differed between conditions (with and without Vitamin C). We were only interested in the differences between SIT and VIT at each time-point; hence, pairwise comparisons were used to identify significant differences at each time-point if the main effects were positive. Alpha level for statistical significance was set a priori at 0.05. All statistical calculations were performed using IBM SPSS Statistics 22.0 software (IBM SPSS, Inc.).

Results

We recruited and tested 12 healthy, inactive participants (See Table 1 for subject characteristics). We had an incomplete ultrasound image at 1 time point on 1 subject and this subject was omitted from all analysis. As a result, data on 11 participants is presented.

Tests to identify the effects of SIT

SFA FMD% (Figure 2)

By one way ANOVA there was a significant decline in FMD% from baseline during 3 hr of sitting (p=0.001, $\eta^2=0.406$). The variable of interest was FMD% between each time-point. We conducted pairwise comparisons to investigate the differences between SIT and VIT FMD% at each hour. Baseline SFA FMD% did not differ between the 2 trials ($p=0.093$). However, FMD% was significantly different between trials at 1 h ($p=0.009$), 2 h ($p=0.016$), and 3 h ($p=0.004$) with VIT preventing the decline of FMD.

SFA shear rates

There was a significant decline in antegrade shear rate ($p=0.04$, $\eta^2=0.238$) and mean shear rate from baseline to 3 h ($p=0.037$, $\eta^2=0.506$). There were no significant differences in retrograde shear rate, peak shear rate, and shear rate area under the curve (SRauc) from baseline to 3 h ($P>0.05$).

Baseline diameters between SIT and VIT:

Using pairwise comparisons, there was a significant difference between baseline diameters in the SIT and VIT trial ($p=0.019$). There was, however, no significant difference between the baseline diameters at 1 h, 2 h, or 3 h ($p>0.05$).

Tests to identify differences between SIT and VIT

SFA FMD% between SIT and VIT (Figure 2)

There was a significant trial x time interaction for SFA FMD (p=0.001, $\eta^2=0.406$). The variable of interest was FMD% between each time-point. We conducted pairwise comparisons to investigate the differences between SIT and VIT FMD% at each hour. Baseline SFA FMD% did not differ between the 2 trials ($p=0.093$). However, FMD% was significantly different between trials at 1 h ($p=0.009$), 2 h ($p=0.016$), and 3 h ($p=0.004$) with VIT preventing the decline of FMD.
there were multiple participants who did not have an arterial dilatation in response to hyperemia. There were no significant differences between antegrade (p=0.193), mean (p=0.605), peak (p=0.187), or retrograde shear (0.723) rates between SIT and VIT trials (p=0.05). Baseline diameters and all flow data are presented in Table 2.

### Discussion

The purpose of this study was to determine if antioxidant vitamin C prevents impairment of SFA endothelial function during prolonged sitting. This is the first study to our knowledge to experimentally investigate the role of oxidative stress during prolonged sitting. We hypothesized that there would be a significant decline in SFA FMD from baseline during 3 h of sitting and that antioxidant vitamin C would prevent the decline in SFA FMD during 3 h of sitting. Indeed, in support of our hypothesis, we found a significant decline in SFA FMD during 3 h of sitting (SIT trial), which was counteracted by Vitamin C (Figure 2). Our results suggest that endothelial function in the SFA is significantly impaired from baseline during a 3-h session of uninterrupted sitting, beginning sometime before the first hour. We also found that there was an associated decline in antegrade and mean shear rates from baseline to 3 h of sitting in the SIT trial. Based on the results from the VIT trial, the mechanism responsible for the decline in FMD during SIT is suggestive of an oxidative stress mechanism. There is only 1 other study to our knowledge that has directly investigated oxidative stress as a potential mechanism for endothelial dysfunction in a model of physical inactivity. In an animal model for chronic inactivity, Laufs et al. [19] found that 6 weeks of inactivity significantly increased NADPH oxidase, superoxide, and reactive oxygen species (ROS), and resulted in endothelial dysfunction and increased atherosclerosis in sedentary mice as compared to mice who voluntary trained on running wheels. It is well known that exercise-induced shear stress facilitates beneficial endothelial adaptations [20]; hence, it is tempting to speculate that the endothelial dysfunction from oxidative stress in Laufs’ [19] study may be a result of low shear stress in the sedentary mice as compared to the running mice. It is important to note that this study induced chronic inactivity, whereas we were looking at acute inactivity, but there is no other study to our knowledge which has looked at oxidative stress during inactivity. We have recently shown (and data presented in this paper) that there is a significant decline in mean shear rate during 3 h of sitting. Low mean shear rate has been implicated in reduced NO bioavailability and in decreased activity of eNOS [21]. Hence, in the absence of biological markers (see limitations) to test if the impairment in endothelial function is caused by mechanisms related to oxidative stress, we used vitamin C in our research design.

### Vitamin C and endothelial function

Use of vitamin C is a novel and critical aspect of this project because it provides suggestive evidence on the mechanisms for endothelial dysfunction during sitting. Vitamin C is an antioxidant and has been previously used in endothelial function to counteract oxidative stress (and to discover this mechanism responsible for the decline in FMD during SIT). It is well known that exercise-induced shear stress facilitates beneficial endothelial adaptations [20]; hence, it is tempting to speculate that the endothelial dysfunction from oxidative stress in Laufs’ [19] study may be a result of low shear stress in the sedentary mice as compared to the running mice. It is important to note that this study induced chronic inactivity, whereas we were looking at acute inactivity, but there is no other study to our knowledge which has looked at oxidative stress during inactivity. We have recently shown (and data presented in this paper) that there is a significant decline in mean shear rate during 3 h of sitting. Low mean shear rate has been implicated in reduced NO bioavailability and in decreased activity of eNOS [21]. Hence, in the absence of biological markers (see limitations) to test if the impairment in endothelial function is caused by mechanisms related to oxidative stress, we used vitamin C in our research design.

### Table 2. Baseline diameters (mm), Shear rates (s⁻¹), SRauc and normalized FMD in the SFA during the SIT and VIT trial at baseline, 1 hr, 2 hr and 3 hr.

|               | SIT          | VIT          |
|---------------|--------------|--------------|
| Baseline diameters | 6.32±0.66   | 6.36±0.74   |
| Antegrade     | 7.7±1.36     | 5.77±2.39   |
| Mean          | 5.03±1.58    | 2.9±2.94    |
| Retrograde    | 2.65±1.16    | 2.87±1.79   |
| Peak          | 54.21±69.10  | 21.01±13.50 |
| SRauc (u)     | 831.9±228.1  | 439.5±294.2 |
| FMD: SRauc    | 0.005±0.002  | 0.24±0.05   |

SRauc and FMD: SRauc are from 5 participants. Please see limitations for details.

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mechanism) from smoking [22], high-fat meals [23], exercise [24], and increased oscillatory shear [15]. Our results show that vitamin C prevents the decline in SFA FMD during 3 h of sitting, suggesting a possible role of mechanisms involving oxidative stress causing the decline of FMD during sitting. The dose of vitamin C ingested by our participants was higher than the recommended daily allowance and was expected to saturate participants’ plasma [25]. There are several potential pathways by which vitamin C may have prevented the decline in SFA FMD during sitting. Vitamin C has been shown to restore NO activity [14] and decrease the activation of vascular NADPH oxidase (a pro-oxidant) [26]. Vitamin C supplementation has been found to protect BH₄ and thus the activity of eNOS [27,28]. Vitamin C also allows the recycling of BH₄ radicals to a stable BH₄ in the presence of ROS [29], thereby maintaining NO production [30]. Indeed, vitamin C may have directly counteracted oxidative stress by scavenging elevated superoxide radicals [31]. Vitamin C, when ingested at 30 min and 1 h 30 min, prevented the decline in endothelial function during 3 h of sitting. This may suggest that mechanisms involving oxidative stress may be present during prolonged sitting, which are counteracted by vitamin C. Using a placebo in addition to vitamin C was also considered. Placebos are used in trials to conceal whether a treatment is being given or not and hence to control for the psychosomatic effects. It is highly unlikely that the vascular responses including blood flow patterns and diameters can be modulated by psychosomatic effects in a period of 3 h. Vitamin C also did not change blood flow patterns during the trial. Furthermore, use of vitamin C was based on pilot data and was meant only to uncover the oxidative stress mechanism.

**Alternative mechanisms**

There are alternative mechanisms which may attenuate endothelial function during prolonged sitting. For example, viscosity has been shown to be higher in the legs after a session of 2 h of sitting [6], and in the presence of low mean shear rates [32]. Viscosity has been positively associated with increased coagulation and inflammatory markers [33], which in turn have been strongly associated with impaired endothelial function [34]. It is possible that vitamin C counteracted the viscosity and inflammation in our experiment. Indeed, plasma vitamin C has anti-inflammatory effects and is inversely associated with viscosity [35]. Additionally, skeletal muscle inactivity results in an increase in intracellular calcium concentration, which may result in increased NAD(P)H oxidase activity elevating superoxide production [36].

**Limitations**

Our study is novel and the results have public health significance. However, it is not devoid of limitations. For example, not being able to obtain blood samples from the lower extremities is a clear limitation. Blood samples from the lower extremities would have increased our understanding about the physiological milieu in the legs during sitting and would have allowed us to measure viscosity and oxidative stress markers. Viscosity measurement would have allowed us to explore alternative mechanisms for FMD attenuation during sitting, and calculate shear stress rather than using shear rate as a surrogate measure for shear stress. Oxidative stress measurements from the legs would have conclusively supported the oxidative stress mechanism. It has to be understood, however, that obtaining continuous blood samples from a lower extremity vein in this design is difficult, and these investigators tried this technique in 5 people multiple times before it was deemed logistically impossible (technical difficulty in addition to subject discomfort) even with the use of a topical anesthetic. We did collect venous blood samples from the antecubital vein and analyzed the plasma for oxidative stress biomarkers. However, these biomarkers did not change during 3 h of sitting as compared to baseline, similar to the brachial artery FMD, implying that the local lower extremity effect could not be detected using the upper extremity measurements [37]. We did not measure BP during SIT. BP is known to increase during sitting [4,5] and may lead to endothelial dysfunction. Increased BP is also known to impair endothelial function via the oxidative stress mechanism [14]. Some of our participants had no FMD at some time-points, which did not allow us to calculate SRauc until the point of maximum dilation. Since SRauc is the stimulus that causes the dilation, we cannot speculate if it was the stimulus that was attenuated. Our study was conducted in young and inactive men and more research needs to be done to replicate these findings in women and subjects of different age groups. Despite these minor limitations, our study is the first to our knowledge to discover the mechanism behind vascular impairment during prolonged sitting.

**Conclusions**

Three hours of sitting leads to a significant decline in endothelial function, most likely through an oxidative stress mechanism. Sitting is a common activity of modern human beings. Repeated sitting sessions may lead to chronically altered shear patterns, oxidative stress, endothelial dysfunction, and a predisposition to atherosclerosis. With the prevalence of sedentary behavior increasing, vitamin C as a regular supplement taken during the seated sessions may play a role in maintaining vascular health in the inactive population.

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Competing interests

The authors report no competing interests.

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