Using multifractal detrended fluctuation analysis to assess sacral skin blood flow oscillations in people with spinal cord injury

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Abstract—The purpose of this study was to investigate whether the multifractal detrended fluctuation analysis (MDFA) of skin blood flow oscillations (BFO) differed between nondisabled controls and people with spinal cord injury (SCI). The study of skin BFO has shown promise for assessing blood flow control mechanisms and risk for pressure ulcers. We recruited 23 subjects, including 11 people with SCI and 12 nondisabled controls. Thermally induced maximal sacral skin BFO were measured by laser Doppler flowmetry. MDFA was used to characterize nonlinear complexity of metabolic (0.0095 to 0.02 Hz), neurogenic (0.02 to 0.05 Hz), and myogenic (0.05 to 0.15 Hz) BFO. We found that maximal vasodilation was significantly smaller in people with SCI than in nondisabled controls ($p < 0.05$). MDFA showed that metabolic BFO exhibited less complexity in people with SCI than in nondisabled controls ($p < 0.05$), neurogenic BFO exhibited less complexity in people with complete SCI ($p < 0.05$), and myogenic BFO did not show significant differences between people with SCI and nondisabled controls. This study demonstrated the feasibility of using the MDFA to characterize nonlinear complexity of BFO, which is related to vasodilatory functions in people with SCI.

Key words: blood flow oscillations, detrended fluctuation analysis, laser Doppler, microvascular function, multifractal analysis, nonlinear analysis, pressure ulcers, spectral analysis, spinal cord injury, vasodilatory function.

INTRODUCTION

Spinal cord injury (SCI) causes interruption of the autonomic pathways from the brain stem and hypothalamus to the intermediolateral cell column of the spinal cord [1]. This interruption results in a loss or attenuation of modulation of the spinal autonomic reflexes in response to various stimuli at the level below spinal injury. SCI also causes immobility with a subsequent consequence of prolonged sitting and bed rest. Assuming these postures for a long time not only causes a reduction of blood flow to the compressed soft tissues but also results in an increase of skin temperature [2]. Such an increase raises the metabolic demands of local cells, which may further aggravate the tissue viability of weight-bearing tissues in people with SCI. In nondisabled individuals, an increased temperature elicits a vasodilatory response to remove excessive heat. People with SCI may not be able to induce effective vasodilation under a thermal stress. The impaired thermoregulatory function contributes to the development of pressure ulcers in people with SCI [3]. In addition to the temperature factor, other causative factors of pressure ulcers include pressure, shear, and moisture [2].

Abbreviations: AV = arteriovenous, BFO = blood flow oscillations, BTI = biphasic thermal index, DFA = detrended fluctuation analysis, eNOS = endothelial nitric oxide synthase, LDF = laser Doppler flowmetry, MDFA = multifractal DFA, NS = nonsignificant, SCI = spinal cord injury.

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Skin blood flow signals contain five periodic components [4–6]. A periodic oscillation with a characteristic frequency around 0.01 Hz is reported to be associated with metabolic activity [5,7]. Other underlying physiological mechanisms include neurogenic (0.02–0.05 Hz), myogenic (0.05–0.15 Hz), respiratory (0.15–0.40 Hz), and cardiac (0.4–2.0 Hz) origins. However, the characteristic frequencies embedded in blood flow oscillations (BFO) show a time-varying feature that cannot be fully characterized by linear methods, such as wavelet and Fourier transforms [8–10]. This time-varying feature of characteristic frequencies may be due to the interactions of each control mechanism [7–9]. Stefanovska et al. proposed a model consisting of linear and nonlinear oscillators to understand this phenomenon [8]. Based on this model, the variations in frequency and amplitude of a periodic component are due to the interactions among the control mechanisms of BFO. The nonlinear properties of skin BFO were also reported by Liao et al. and Humeau et al. [11–12]. These studies showed that nonlinear properties of BFO may be related to vasodilatory function: a lower nonlinear complexity of BFO is associated with a smaller vasodilatory response.

Recently, nonlinear complex properties of BFO were quantified with the use of detrended fluctuation analysis (DFA) [13] by Esen’s group [14–15]. They showed that a distinct scaling region, called local region, of the DFA is directly related to the metabolic, neurogenic, and myogenic mechanisms. Furthermore, the scaling exponents in the local region are from only the local origins and are not influenced by systemic factors [15]. This method also allows researchers to detect microvascular dysfunction based on the baseline blood flow without evoking the microvascular response. (The traditional method needs to induce a stimulus into the microcirculatory system to assess its response for assessing microvascular dysfunction.) However, skin blood flow signals have been shown to have multifractal behavior [16–17]. Whether these nonlinear properties of BFO are associated with specific physiological meanings is unclear.

We have conducted a series of studies correlating pathological conditions to the changes of BFO in order to investigate the role of microvascular dysfunction in pressure ulcer risk [7,9,18]. In our previous studies, we successfully isolated physiological mechanisms by using (linear) wavelet-based spectrum analysis of BFO [6–7]. Our method has shown promise for further quantifying the influences of pathological conditions (e.g., SCI, aging, and diabetes mellitus) on microvascular functions, which may lead to a better understanding of the development of pressure ulcers. To achieve our long-term goal of predicting pressure ulcer risk based on microvascular functions, we examined the effect of SCI on the nonlinear complexity of BFO in this study. The specific aims were to investigate the changes of nonlinear complexity of the metabolic (0.0095–0.02 Hz), neurogenic (0.02–0.05 Hz), and myogenic (0.05–0.15 Hz) BFO by using multifractal DFA (MDFA).

**METHODS**

**Participants**

We recruited 23 subjects, including 11 people with SCI and 12 nondisabled controls. The inclusion criteria for the SCI participants included an injury level between cervical 4 and thoracic 12, nonambulation (wheelchair users), and time after spinal injury of 6 months. None of the participants had cardiorespiratory disease, hypertension, diabetes mellitus, or other pathological conditions or were taking medications that might affect cardiovascular function. People with SCI were recruited from the University of Oklahoma Health Sciences Center and University of Central Oklahoma Wellness Center. The demographic features of the enrolled subjects are shown in the Table.

<table>
<thead>
<tr>
<th>Demographic</th>
<th>Complete SCI</th>
<th>Incomplete SCI</th>
<th>Nondisabled Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of Subjects</td>
<td>4</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>Sex: M/F</td>
<td>3/1</td>
<td>4/3</td>
<td>4/8</td>
</tr>
<tr>
<td>Age (yr)</td>
<td>33.0 (27, 46)</td>
<td>43.0 (34, 49)</td>
<td>24.0 (22, 26)</td>
</tr>
<tr>
<td>Duration of SCI (yr)</td>
<td>7.0 (4.6, 10.5)</td>
<td>5.7 (2.4, 10.6)</td>
<td>—</td>
</tr>
<tr>
<td>Body Mass Index (kg/m²)</td>
<td>24.1 (22.2, 25.2)</td>
<td>24.1 (22.0, 28.1)</td>
<td>23.6 (21.0, 25.8)</td>
</tr>
</tbody>
</table>

F = female, M = male, SCI = spinal cord injury.
Procedures
After at least 30 min of quiet rest to become acclimated to the room temperature (24 °C ± 2 °C), the participant was positioned in a prone posture. Sacral skin BFO were recorded at a sampling rate of 32 Hz with laser Doppler flowmetry (LDF) (PF 5001, Perimed AB; Järfälla, Sweden). A heating probe (Probe 415-242, Perimed AB) was used to heat the skin to 42 °C in 2 min and maintain that temperature for the duration of the 50 min heating period. The protocol included a 10 min preheating period, a 50 min heating period, and a 10 min postheating period [19]. The LDF uses a 780 nm laser diode and a fiber separation (distance between the transmitting and receiving fibers) of 0.25 mm. The configuration leads to a measurement depth on the order of 1 mm [20].

Biphasic Thermal Index
To create a reference benchmark, we used the biphasic thermal index (BTI) [7] to incorporate the information of biphasic vasodilatory response. The index was based on Minson and colleagues’ protocol in which they used a fast local heating procedure to induce a biphasic vasodilatory response [19]. The biphasic response includes two peak values of blood flow under local heating. The first peak is mediated by the activity of sensory nerves and can be modulated by the sympathetic nervous system. The second peak (plateau) is mediated by local release of endothelial nitric oxide. The BTI consists of three ratios: ratio of the first peak to baseline blood flow (P1/Baseline), ratio of the nadir to baseline blood flow (Nadir/Baseline), and ratio of the second peak to baseline blood flow (P2/Baseline). The first peak is mainly mediated by sensory axon reflex and the second peak is primarily mediated by endothelial nitric oxide [19].

Multifractal Detrended Fluctuation Analysis
A monofractal signal is homogeneous, in the sense that it has the same scaling properties throughout the entire signal [17]. A multifractal signal, on the contrary, contains many subsets that have different scaling properties. The multifractality may exhibit in several ways: a type of correlation on small (time) scales and another type of correlation or uncorrelated behavior on larger scales, different scaling behavior in different parts of the time series, and different scaling behaviors in interwoven fractal subsets of the series [17].

MDFA has been shown to reliably determine the multifractal scaling behavior of time series [17]. This method is based on a generalization of the standard DFA. Its procedure can be briefly described as follows. First, for a time series \( x(k) \) of length \( N \) on a compact support, one determines the integrated signal profile

\[
Y(i) = \sum_{k=1}^{i} (x(k) - \langle x \rangle), i = 1, ..., N
\]

where \( \langle x \rangle \) is the mean of the time series. Then, the profile is divided into \( N_s \) nonoverlapping segments of length \( s \) and the same procedure is repeated starting from the opposite end (2\( N_s \) segments total). The local trend for each segment \( v \) is estimated by fitting a \( m \)th order polynomial \( P_v^{(m)} \) and subtracting it from the segment. Next, one determines the variance

\[
F^2(v,s) = \frac{1}{s} \sum_{i=1}^{s} \{Y[(v-1)s+i] - P_v^{(m)}(i)\}^2
\]

for each segment \( v, v = 1, ..., N_s \) and

\[
F^2(v,s) = \frac{1}{s} \sum_{i=1}^{s} \{Y[N - (v-N_s)s+i] - P_v^{(m)}(i)\}^2
\]

for \( v = N_s + 1, ..., 2N_s \). Finally, a \( q \)th order fluctuation function is defined as

\[
F_q(s) = \left\{ \frac{1}{2N_s} \sum_{v=1}^{2N_s} [F^2(v,s)]^{1/2} \right\}^{1/q}
\]

for \( q \neq 0 \) and \( F_0(s) \) is defined as

\[
F_0(s) = \exp\left\{ \frac{1}{4N_s} \sum_{v=1}^{2N_s} \ln[F^2(v,s)] \right\} .
\]

If the series \( x(k) \) is long range power-law correlated, \( F_q(s) \) increases for \( s \) values in a certain range, as a power-law

\[
F_q(s) \sim s^{h(q)} .
\]

For \( q = 2 \), the standard DFA procedure is retrieved. The \( q \)th order fluctuations \( F_q(s) \) plotted versus various observation window sizes (scales) \( s \) form a family of lines. The slopes of these lines estimate the exponents \( h(q) \). Figure 1 shows an example of the relationship between the \( q \) values and observation windows (scales, \( s \).
For a monofractal signal, \( h(q) \) is independent of \( q \). For a multifractal signal, small and large fluctuations scale differently, thus \( h(q) \) is dependent on \( q \). The reason is that for positive values of \( q \), \( h(q) \) describes the scaling behavior of the segments with large fluctuations, while for negative values of \( q \), \( h(q) \) describes the scaling behavior of the segments with small fluctuations. For stationary, normalized series, the exponents \( h(q) \) are directly related to the scaling exponents \( \tau(q) \), and \( \tau(q) \) is defined as \[ \tau(q) = qh(q) - 1 \text{ and} \]

\[ \alpha = h(q) + qh(q) \text{ and } f(\alpha) = q[\alpha - h(q)] + 1 \]  \( (8) \)

where \( \alpha \) is the Hölder exponent and \( f(\alpha) \) is the dimension of the subset of the time series that is characterized by \( \alpha \).

In the standard DFA, the scaling exponent of a skin blood flow signal in the local region originates from only the local factors \([15]\) and a scale \( s \) is related to a frequency \( f \) by \( s = f_s/f \), where \( f_s \) is the sampling frequency \([14]\). Here, we show that for an appropriate \( q \) interval, the exponents \( h(q) \) in the scale intervals \( f_s/0.02 < s < f_s/0.0095 \) and \( f_s/0.05 < s < f_s/0.02 \) reflect the scaling properties of metabolic activity (0.0095–0.02 Hz) and neurogenic activity (0.02–0.05 Hz), respectively. As illustrated in Figure 2, after we remove the components with frequencies 0.0095 < \( f < 0.02 \) Hz, scaling exponents in the region \( f_s/0.02 < s < f_s/0.0095 \) are close to zero for all values of \(-5 \leq q \leq 5\), while the scaling exponents in other scaling regions are identical to that of the original data. The process applies to the neurogenic BFO (0.02 < \( f < 0.05 \) Hz) (Figure 3). This evidence indicates that for \(-5 \leq q \leq 5\), the scaling exponents of a blood flow signal in the interval \( f_s/0.02 < s < f_s/0.0095 \) and \( f_s/0.05 < s < f_s/0.02 \) reflect the scaling behavior of metabolic and neurogenic
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BFO, respectively. The same procedures can be used to process the myogenic BFO.

Previous studies have used the range of Hölder exponent \([21–22]\) or \(h(q_{\text{min}}) – h(q_{\text{max}})\) \([23]\) to quantify the multifractality degree of time series. However, estimates of the range of Hölder exponent often encounter a problem of numerical instability \([24]\). The results of simulation experiments showed that the range of \(h(q)\), \(\Delta h = \max(h(q)) – \min(h(q))\) is more robust than \(h(q_{\text{min}}) – h(q_{\text{max}})\) for typical monofractal signals, e.g., fractional Brownian motions (fBmH, \(H\) is the Hurst exponent, \(0 < H < 1\)). On the other hand, for fBmH with the length of 10 min blood flow data, values of \(\Delta h\) in the scale intervals \(f_s/0.02 < s < f_s/0.0095\) and \(f_s/0.05 < s < f_s/0.02\) are significantly smaller than that of blood flow data, especially in nondisabled people, indicating nondisabled people exhibit multifractality in BFO. Therefore, we used \(\Delta h = \max(h(q)) – \min(h(q))\) in the scale interval \(f_s/0.02 < s < f_s/0.0095\) to quantify the multifractality degree of metabolic BFO, the scale interval \(f_s/0.05 < s < f_s/0.02\) to quantify the multifractality degree of neurogenic BFO, and the scale interval \(f_s/0.15 < s < f_s/0.05\) to quantify the multifractality degree of myogenic BFO.

Statistical Analysis

Nonparametric statistics were used because of the small sample size. Comparisons among groups in the same thermal condition were performed using the Kruskal-Wallis tests. When the factor was significant, we further checked the significance between groups by using the Wilcoxon rank sum tests. Comparisons between two thermal conditions in the same group were performed using the Wilcoxon signed rank tests. The significance level was set at \(p < 0.05\). MATLAB and Signal Processing and Statistical Analysis Toolboxes (R2008b, MathWorks; Natick, Massachusetts) were used to implement MDFA and statistical testing.

RESULTS

People with SCI showed a smaller vasodilatory response than nondisabled controls (Figure 4). During the axon reflex mediated vasodilation (P1/Baseline), a significantly smaller increase in sacral skin perfusion was observed in people with complete \((p < 0.01)\) and incomplete \((p < 0.05)\) SCI than in nondisabled controls. During
the maximal vasodilation (P2/Baseline), a significantly smaller increase in sacral skin perfusion was observed in people with complete (p < 0.01) and incomplete (p < 0.05) SCI compared with nondisabled subjects. During maximal vasodilation (P2/Baseline), significantly smaller increase in sacral skin perfusion is observed in people with incomplete SCI compared with nondisabled subjects (p < 0.05). Values are expressed as mean ± standard error. * indicates p < 0.05 and ** indicates p < 0.01. P1 = first peak of biphasic vasodilation, P2 = second peak of biphasic vasodilation.

Figure 5 shows the multifractality degree (Δh) of metabolic oscillations in nondisabled controls and people with spinal cord injury (SCI) (complete and incomplete). In responding to local heating, multifractality degree of metabolic oscillations significantly increases in nondisabled subjects (p < 0.01) but not in people with complete or incomplete SCI. If multifractality degree of metabolic oscillations is compared during maximal vasodilation, people with complete or incomplete SCI show significantly lower values than nondisabled controls (p < 0.05). No statistically significant difference exists between people with complete and incomplete SCI (p > 0.05). Values are expressed as mean ± standard error. * indicates p < 0.05 and ** indicates p < 0.01.

Figure 6 shows the multifractality degree (Δh) of neurogenic oscillations. In responding to local heating, the multifractality degree of neurogenic oscillations significantly decreased in people with complete SCI (p < 0.05) and people with incomplete SCI (p < 0.05). By comparing the multifractality degree of neurogenic oscillations during the maximal vasodilation, we found that people with complete SCI showed a significantly lower value than nondisabled controls (p < 0.05).

Figure 7 shows the multifractality degree (Δh) of myogenic oscillations. No significant differences were found between people with SCI and nondisabled controls (NS).

DISCUSSION

We have shown that the complexity of metabolic oscillations (0.0095–0.02 Hz) in people with complete and incomplete SCI was significantly lower than in nondisabled controls during the maximal vasodilation. People with complete SCI exhibited a significantly lower complexity of the neurogenic oscillations (0.02–0.05 Hz) as compared with nondisabled controls. Our findings provide new insight into the mechanisms responsible for vasodilatory dysfunction in people with SCI. To the best of our knowledge, this is the first study to characterize the impairment of nonlinear complexity of metabolic and neurogenic oscillations in people with SCI and the results support our hypothesis that SCI causes a decrease of nonlinear complexity in sacral skin BFO. This result has advanced our understanding of physiological meanings of BFO and their roles of vasodilatory dysfunction. The finding may contribute to our long-term goal of using BFO patterns for the early detection of pressure ulcers in people with SCI [7,9,18].
Vasodilatory function could be used to quantify risk for pressure ulcers in people with SCI [25–27]; people with SCI who fail to increase skin blood in response to a causative factor of pressure ulcers (pressure, shear, or heating) are susceptible to pressure ulcers [27–28]. Schubert and Fagrell demonstrated that people with SCI have impaired vasodilatory response to either local heating or local pressure [27]. Hagisawa et al. showed that people with SCI take longer to reach peak hyperemia after removal of loading pressure [29]. Jan et al. showed that pressure loadings (alternating and constant pressures) induce different skin blood flow response between people with SCI and nondisabled controls [28]. Nicotra et al. observed a lower ratio of the first peak to baseline blood flow at the level below the lesion in people with complete high-level SCI (cervical 6 to thoracic 5) as compared with people with low-level SCI (thoracic 6 to 11) [26]. They suggested that this lower ratio is due to the loss of sympathetic innervations in people with high-level SCI. The need for sympathetic innervation for a full vasodilatory response was also indicated by Ping and Johnson [30]. These studies have demonstrated that people with SCI have impaired vasodilatory response to loading pressure and thermal stress. In the literature, the influences of loading pressure have drawn more attention than other factors (temperature, shear, and moisture). We are particularly interested in assessing the thermoregulatory functions and the role of local skin temperature changes due to prolonged seating in the development of pressure ulcers. In this study, we continued our research on sacral skin BFO and their relationship with vasoregulatory functions [6–7].

The LDF is an indirect measure of red blood cell velocity [31]. Previous studies have shown that the LDF signal is independent of the effect of the absorption characteristics of the blood for a given LDF system as the fiber separation and wavelength are fixed [32–34]. Shepherd et al. reported that laser Doppler velocity signals are unaffected by changes in blood oxygenation [32]. Nilsson et al. evaluated skin blood flow measurements based on the laser Doppler principle and found a linear relationship between flowmeter response and flux of red blood cells [33]. Fredriksson et al. evaluated the effects of blood concentration and blood oxygen saturation on the measurement depth and volume in LDF through Monte Carlo simulations of tissue using relevant computer models [34]. The results showed that, for the wavelength of 780 nm, the effect of the oxygenation status of the blood on the measurement depth is very small.

The depth of tissue penetrated by the laser depends on factors such as tissue properties, light source wavelength, and probe configuration [20,35]. The LDF used in this study uses a 780 nm laser diode and the fiber separation is 0.25 mm. This configuration leads to a measurement depth on the order of 1 mm for human skin [20]. The measurement depth of the LDF reaches to the arterioles, capillaries, and postcapillary venules of the upper horizontal plexus but does not extend to the deep horizontal plexus [36]. Thus, skin perfusion as measured in...
this study includes blood flow in the capillaries, arterioles, venules, and arteriovenous (AV) anastomoses (or shunts) [20]. Blood flow in the capillaries provides nutrients and oxygen to local cells, and blood flow in the arterioles, venules, and AV shunts regulate body temperature [20].

BFO have been assessed with use of the amplitudes or power in several frequency bands [5,7,37]. It has been demonstrated that attenuated metabolic and neurogenic oscillations are associated with vasodilatory dysfunction [7,37]. Li et al. observed significantly lower blood flow in people with SCI than in nondisabled controls [37]. They also showed that the amplitudes of metabolic and neurogenic frequencies are smaller in people with SCI than in nondisabled controls. However, the dynamics of skin blood flow or of a frequency component contain not only amplitude information but also structural information (nonlinear complexity). The latter cannot be characterized by the averaged amplitude in a time interval. Although wavelet-based time-frequency-amplitude presentation can describe how frequency and amplitude change with time, difficulties may arise from quantifying the changes [7,9]. MDFA allows us to quantify the dynamics of a frequency component with respect to its complexity. From our results, we may deduce that metabolic oscillations in nondisabled people have richer structures (i.e., more complex behavior) than in people with SCI. Our results support the general concept that the output of a healthy system reveals complex variability and that the complexity decreases with aging and diseases [16,38–39]. Reduced complexity of the physiological system is widely accepted to be associated with an individual’s poor abilities to adapt to environmental stimulus. Such vasodilatory impairment may increase risk for tissue ischemic damage [7].

Variations in the frequency and amplitude of the metabolic oscillations are likely induced by interactions between other control mechanisms (e.g., neurogenic and myogenic) of skin microcirculation. The cardiovascular system has been modeled as coupled linear and nonlinear oscillators to study the time-varying features of characteristic frequencies [8]. Metabolic activity adjusts blood flow to meet the need of cells for oxygen. This activity is dependent on cardiac and respiratory activities, which control the level of oxygenated blood flow. On the other hand, the constrictor activities of myogenic and neurogenic systems diminish blood transport, thus lowering metabolic activity [8]. Ping and Johnson found that the myogenic response could be enhanced by sympathetic tone [30]. According to these studies, we assume that the interactions between other control mechanisms result in the variations in frequency and amplitude of metabolic oscillations. This assumption is supported by our observations, including that metabolic oscillations in nondisabled people are less homogeneous than in people with SCI both in the preheating and maximal vasodilation periods and that both in nondisabled people and people with SCI, metabolic oscillations become less homogeneous in response to local heating. In nondisabled controls, the metabolic oscillator interacts with other oscillators, including the neurogenic oscillator, in an adaptive way. A broad class of outside interactions leads to metabolic oscillations over multiple time scales. These fluctuations may superimpose, counteract, or modulate each other, resulting in complex oscillations. In people with SCI, diminished neurogenic activity may lower the interactions among these control mechanisms. The attenuated interactions result in less complex structures of metabolic oscillations. During local heating, the power of metabolic frequency has been shown to increase at the highest order, while the powers of the other characteristic frequencies also increase but at lower orders [7]. Thus, local heating may lead to changes in the intensities of the interactions among the control mechanisms, resulting in more complex structures of metabolic oscillations.

Our results indicated that people with SCI have a lower complex behavior of metabolic oscillations. The underlying mechanism of the attenuated metabolic endothelial function in people with SCI is unclear. One possible reason is their physically inactive lifestyle. The occurrence of endothelial dysfunction is primarily the result of reduced nitric oxide bioavailability [40]. Nitric oxide is a vasoactive substance that plays a pivotal role in the maintenance of vascular homeostasis. The amount of nitric oxide available depends on several key factors including the activity of the endothelial nitric oxide synthase (eNOS) and nitric oxide degradation. People with SCI are likely inactive relative to nondisabled controls. Physical activity induces increases in shear stress on the endothelium, which can lead to an increase in eNOS expression [40]. Furthermore, exercise increases phosphorylation and can also positively influence the nitric oxide half-life by reducing nitric oxide degradation. According to the literature, physical activity may slow down or even reverse metabolic dysfunction in people with SCI. MDFA of skin BFO, the method proposed in this study, could be used to assess the effectiveness of
exercise interventions on enhancing microvascular function in people with SCI. Our speculation may be validated by examining the influence of exercise on maximal vasodilation and multifractality degree of metabolic oscillations in people with SCI. A longitudinal monitoring of skin perfusion in people with SCI who are enrolled in an exercise program might demonstrate an increase of maximal vasodilatory function and complex behavior of skin BFO. Another reason for metabolic endothelial dysfunction in the SCI population is reduced sympathetic tone. Miranda and Hassouna suggested that the interruption of neurological impulses causes metabolic changes in blood vessels that result in altered venous competence [41]. This hypothesis is supported by the findings of Houtman et al. [42], who found that sympathetic nervous system response was altered in people with SCI in comparison with nondisabled participants. This argument is also supported by the results of BTI. We observed significantly lower values of P1/Baseline and P2/Baseline in people with SCI (Figure 4). Since the first peak is modulated by the activity of sympathetic nerves [43] and the secondary peak is mediated by endothelial nitric oxide [19], we may deduce that in people with SCI, both sympathetic tone and nitric oxide release were reduced. Therefore, the attenuated metabolic endothelial function in people with SCI may be attributed to at least two factors, including reduced sympathetic tone and reduced nitric oxide release.

Using MDFA, we were able to characterize the nonlinear complexity of sacral skin BFO and detect that people with SCI had a lower complexity than nondisabled controls. Our results support the concept that decreased nonlinear behavior of a physiological system is associated with pathological states [16,38–39]. In nondisabled controls, a significant increase of the multifractality degree of metabolic BFO in response to local heating may be attributed to more metabolic activities and the interactions between other mechanisms. On the other hand, in people with SCI, a smaller change in the multifractality degree of metabolic oscillations in response to local heating may suggest the impairment of metabolic endothelial function and/or altered coupling between metabolic control and other regulatory mechanisms of skin microcirculation. More research is needed to investigate the consequence of SCI on the changes of nonlinear properties of BFO.

An important challenge in applying the MDFA to detect the complexity of a frequency component (e.g., metabolic component) of skin BFO is how to accurately determine the frequency band of this component. We found that the frequency band of the metabolic activity slightly varies among the participants and even varies from one time period to another in the same participant. These time-varying features have been indicated in our previous studies [7,9]. Thus, in order to accurately estimate the nonlinear multifractality of the metabolic oscillations or other physiological-related oscillations one must visually inspect, the energy spectrum of blood flow signal and the log ($F_q(s)$) – log ($s$) plot to determine the frequency band.

Previous studies have reported that cigarette smoking influences microvascular function. Warner et al. showed that nicotine caused a decrease in resting and maximal skin blood flow responses to prolonged heating [44]. Argacha et al. demonstrated that passive smoking was associated with a prolonged rise of skin blood flow in response to local heating [45]. We did not exclude smokers from this study. Two smokers were enrolled in this study: one in the complete SCI group and the other in the nondisabled group. The smoker in the SCI group had been smoking about 8 to 10 cigarettes daily for about 10 years. The smoker in the nondisabled control group had been smoking about 15 to 20 cigarettes daily for 3 years. The BTI of the two smokers were (4.3, 4.2, 10.9) and (10.8, 6.6, 21.3), respectively; and the average BTI of the corresponding groups were (4.5, 3.9, 11.6) and (8.5, 6.5, 17.7), respectively. On the basis of the direct comparison, these two smokers had microvascular function consistent with the other subjects in each group. We anticipate that smoking had minimal influence on these two subjects’ skin blood flow responses.

This study had several limitations. First, we only recruited 11 people with SCI into this pilot study. We intended to test the feasibility of using MDFA to assess microvascular dysfunction in people with SCI. Thus, we did not recruit a larger group of research participants. Future studies may consider a larger sample size to validate our results. Second, our design did not match age and sex between the nondisabled controls and people with SCI. Although sex and age might confound our results, the influences of these factors should be minimal [46]. Although the SCI groups were not age-matched to the nondisabled control group, the influence of the age factor should be minimal compared with the SCI factor. Previous studies reported that age is not correlated with skin blood flow in people between ages 17 and 60 [46–47]. This evidence suggests that the effect of age/aging on microvascular function is usually documented in people over age 60 [48]. In this study, the median (lower and
upper quartiles) ages of the complete SCI, incomplete SCI, and nondisabled controls were 33 (27.5, 46), 43 (34, 49), and 24.5 (22, 26), respectively. Skin blood flow of these research subjects was not largely affected by age [46–47]. To assess the effect of age on skin perfusion, we performed a regression analysis between the BTI and multifractality degree and age in three subject groups. The results indicated no relationship between age and the skin blood flow-related parameters (Figure 8). Third, many other nonlinear methods (e.g., entropy, correlation dimension, Lyapunov exponent) may be used to quantify nonlinear properties of BFO in people with SCI. In this study, we only selected the MDFA to quantify complexity of BFO in people with SCI. The rationale for this selection was based on the prior publications of Esen and colleagues [14–15]. In their studies, they demonstrated the feasibility of using DFA to quantify the nonlinear properties of BFO. The feasibility of using other methods to quantify the changes of nonlinear properties of BFO in people with SCI should be examined to better understand the role of SCI in microvascular dysfunction.

CONCLUSIONS

Our results provided initial evidence that the dynamics of metabolic and neurogenic controls of skin BFO in people with SCI changed, exhibiting less complexity than those in nondisabled controls. Unlike traditional methods, in which the normalized amplitude of a frequency component is used as a quantitative measure of the related control mechanism of BFO, MDFA provides information about the nonlinear complexity of a blood flow signal, which may be used to study the interactions among various blood flow control mechanisms and the influences of SCI on microvascular functions. This method may contribute to the understanding of the effects of SCI on microvascular dysfunction and risk for pressure ulcers.

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