Dynamic Cerebrovascular Diseases

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Dynamic Cerebral Autoregulation Is Transiently Impaired for One Week after Large-Vessel Acute Ischemic Stroke

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Key Words
Dynamic cerebral autoregulation · Stroke · Transcranial Doppler · Transfer function analysis

Abstract
Background: Dynamic cerebral autoregulation (DCA) is the continuous counterregulation of cerebral blood flow to fluctuations in blood pressure. DCA can become impaired after acute stroke, but it remains unclear to what extent and over what interval this occurs. Methods: We included 28 patients (NIHSS = 12 ± 6.5, age = 68.4 ± 17.1, 16F) with acute large-vessel ischemic stroke in the middle cerebral artery territory and 29 healthy controls (mean age 54.9 ± 9, 16F). DCA was assessed by simultaneous measurement of blood pressure together with blood flow velocities using finger plethysmography/arterial catheter and transcranial Doppler over three 10-minute recordings on days 0–2, 3–6 and ≥7 days after stroke. Transfer function analysis was applied to calculate average phase shift (PS) in the low frequency range (0.06–0.12 Hz). Less PS indicated poorer autoregulation. The affected side was compared with the unaffected side and controls. Univariate comparisons of data were performed using t tests at single time points, and generalized estimating equations with an exchangeable correlation matrix to examine the change in PS over time. Results: At mean 1.3 ± 0.5 days after stroke the average PS in the affected hemisphere was 29.6 ± 10.5 vs. 42.5 ± 13 degrees in the unaffected hemisphere (p = 0.004). At 4.1 ± 1 days, the PS in affected and unaffected hemisphere was 23.2 ± 19.1 vs. 41.7 ± 18.5 degrees, respectively (p = 0.003). At mean 9.75 ± 2.2 days stroke there was no difference between the affected and unaffected hemisphere (53.2 ± 28.2 vs. 50.7 ± 29.2 degrees, p = 0.69). Control subjects had an average PS = 47.9 ± 16.8, significantly different from patients’ affected hemisphere at the first two measurements (p = 0.001), but not the third (p = 0.37). The PS in controls remained unchanged on repeat testing after an average 19.1 days (48.4 ± 17.1, p = 0.61). Using the last recording as the reference, the average PS in the affected hemisphere was –23.54 (–44.1, –3) degrees lower on recording one (p = 0.025), and –31.6 (–56.1, –7.1) degrees lower on recording two (p < 0.011). Changes in the unaffected hemisphere over time were nonsignificant. Discussion: These data suggest that dynamic cerebral autoregulation is impaired in the affected hemisphere throughout the first week after large-vessel ischemic stroke, and then normalizes by week two. These findings may have important implications for acute blood pressure management after stroke.

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Introduction

Under normal circumstances cerebral blood flow is maintained over a wide range of systemic blood pressures, a phenomenon known as cerebral autoregulation (CA). This mechanism ensures that the cerebral blood flow matches the brain’s metabolic demands and protects it from hypoxic or hyperperfusion. The active response of cerebral blood vessels to spontaneous or induced blood pressure fluctuations is also referred to as dynamic cerebral autoregulation (DCA) [1, 2]. DCA may become impaired after ischemic stroke [3–6]; in that setting CBF is likely to depend on cerebral perfusion pressure (CPP), rendering the brain at risk for secondary injury such as further ischemia, hemorrhagic transformation and edema formation. Blood pressure is often elevated early after ischemic stroke [7, 8] and its management in the setting of acute stroke remains unclear. Both high and low blood pressures during the acute stroke period have been associated with poor outcome [9–11]. Given the lack of definitive data, clinical guidelines are based on theoretical assumptions about autoregulation in the ischemic penumbra, and recommend against the administration of antihypertensive agents unless the blood pressure exceeds values >220/120 [12]. However, the temporal course of autoregulatory disturbance remains unknown. Transcranial Doppler ultrasonography (TCD) combined with servo-controlled finger photoplethysmography (Finapres) has allowed continuous, non-invasive assessment of autoregulatory function in the acute stroke period. This approach eliminates the need for potentially harmful induced blood pressure or vasodilatory interventions and allows noninvasive assessment of autoregulatory function in the acute stroke period. This study sets out to describe the specific temporal course of cerebral autoregulation in patients with acute ischemic stroke.

Methods

Subjects and Measurements

Patients with acute ischemic stroke admitted to the Stroke unit or Neurological ICU at Columbia University Medical Center were eligible for the study. DCA measurements were performed on days 0–2, 3–6 and ≥7 days after stroke. Inclusion criteria were MCA territory ischemic stroke, first measurement within 48 h of onset and age greater than 18. Patients were excluded from the study if they had prior clinical stroke, extracranial stenosis greater than 70%, prior SAH or ICH, an inadequate acoustic window, or treatment with vasopressors. Consent was obtained from the patient or the surrogate decision maker prior to enrollment. All patients were provided with detailed written information regarding the intent and methods of the study. Approval for the study was obtained from the Institutional Review Board at Columbia University Medical Center.

Reference autoregulation values were obtained from 29 control subjects without cerebrovascular disease; 8 of them had a repeat study after 20 days.

DCA measurements were performed in the supine position with slight elevation of the upper body while the subjects were breathing spontaneously. Cerebral blood flow velocities were assessed using transcranial Doppler (DWL-Multidop-X, Sipplingen, Germany). Proximal MCA patency was a prerequisite for the Doppler monitoring. The proximal middle cerebral artery was insonated through the temporal window with a 2 MHz probe attached to a head frame, at a depth of insonation of 45–60 mm. Blood pressure was recorded simultaneously using intraarterial catheter or servo-controlled finger plethysmography (Finometer Pro, Amsterdam, the Netherlands). The appropriate finger cuff (size: small, medium or large) was placed on the middle phalanx of the left or right middle finger. After establishing a stable recording, the intermittently occurring calibration procedure was turned off. Data was recorded for 10 min. All analog signals were digitized and stored for editing and offline analysis. Data sampling frequency was 100 Hz.

Data Analysis

The editing process included temporal synchronization of the blood pressure and blood flow velocity wave forms using ICU pilot software (Dipylon Medical, Solna, Sweden). This was followed by visual inspection and removal of all major artifacts. The data was then analyzed using Matlab (MathWorks, Natick, Mass., USA) with an in-house written program. The relationship between changes in arterial pressure and CBFV was assessed with transfer function analysis.

Transfer Function Analysis (TFA)

Because the data being acquired takes the form of 2 repetitive oscillatory patterns – arterial blood pressure (ABP) and cerebral blood flow velocity (CBFV) – their spatial relationship can be best characterized using TFA. This method quantifies the extent to which the input signal (ABP) is reflected in the output signal (CBFV), thus quantifying the efficacy of the autoregulatory system [15].

The transfer function H(f) relating each CBFV to ABP was approximated by assuming linearity and time invariance. Estimation of spectra and transfer function was based on the method described by Welch and has been outlined in more detail previously [16, 17].

In brief, arterial blood pressure ABP and CBFV from the left and right middle cerebral arteries were normalized with respect to the mean prior to frequency spectrum analysis. The Fast Fourier Transform (FFT) algorithm was used to calculate the auto-spectra and the cross spectrum of the two signals (ABP and CBFV). Spectra were calculated using individual segments 1/6th the total length of the time series with 50% overlap. Smoothing was accomplished by employing a Hanning window.

The coherence, gain |H(f)| and phase Φ(f) of the system were calculated. Coherence significance criterion (γmin), above which coherence differs significantly from 0, was derived from the degrees of freedom ν of the spectral estimate at a significance level α of 0.05 [14, 18]. The degrees of freedom are dependent on the de-
tails of the spectral estimation method and in our case resulted in a coherence significance level, \( \gamma_{\text{min}} \) equal to 0.5372. Phase shift and gain were calculated by averaging the values of all valid bins in the low-frequency range (0.06–0.12 Hz; each bin is a frequency value at 0.01 Hz increments) where coherence was >0.53.

The resulting phase shift describes to what extent oscillations in CBFV lead those in ABP and can be interpreted as active early counter-regulation. Less phase shift reflects increased latency in cerebral vasomotion and thus poorer autoregulation [19]. Supplemental figure 1 shows an illustration of the concept of TFA.

### Statistical Analysis

Continuous variables were tested for normality using the Kolmogorov-Smirnov test. Data were reported as means and standard deviation (SD) or median, 25th and 75th percentile. Univariate comparisons at single time points were performed using paired t tests. Generalized estimating equations (GEE) with an exchangeable correlation matrix were used to examine the change in phase shift (PS) over time. GEE is an analysis method for longitudinal data when responses are correlated. It accounts for within-subject or within-cluster correlations with a range of correlations structures, and allows for missing data and irregularly timed or mis-timed measurements. A p value of less than 0.05 was considered statistically significant.

### Results

From May 2011 to March 2012 we recruited 28 patients for the study and performed a total of 56 measurements. Twenty-nine age-matched subjects without cerebrovascular disease served as controls. Demographics and baseline characteristics are given in table 1.

At the first two measurements, recordings from four patients could not be obtained due to technical difficulties or the patient’s inability to cooperate with the exam. On the third measurement, recordings were performed on 8 patients; the remainder had already been discharged from the hospital. Overall, we found a significant decrease in phase shift in the affected hemisphere at the first two measurements followed by normalization at the third measurement. Figure 1a–c shows the average phase shift for the affected and unaffected side at the three time points. The first measurement was obtained 1.3 ± 0.5 days after the stroke. The average PS in the affected hemisphere was 29.6 ± 10.5 degrees vs. 42.5 ± 13 degrees in the unaffected hemisphere (p = 0.004). At the second measurement (4.1 ± 1 days) the PS in the affected and the unaffected hemisphere was 23.2 ± 19.1 vs. 41.7 ± 18.5 degrees, respectively (p = 0.003). At 9.75 ± 2.2 days after the stroke there was no difference between the affected and the unaffected hemisphere (53.2 ± 28.2 vs. 50.7 ± 29.2 degrees, p = 0.69). The results of the hemodynamic and autoregulatory parameters are given in table 2. Control subjects had an average PS = 47.9 ± 16.8, significantly different from the affected hemisphere at the first two measurements (p < 0.001), but not the third (p = 0.37). The PS in controls remained unchanged on repeat testing after an average 19.1 days (48.4 ± 17.1, p = 0.61).

Using GEE we found that time of measurement was a significant predictor of phase shift in the affected hemisphere (\( \chi^2 = 6.4, \ p = 0.04 \) and \( \chi^2 = 11, \ p = 0.004, \) respectively). Using the last recording as the reference, the mean PS in the affected hemisphere was −23.54 (CI −44.1, −3) degrees lower on recording one (p = 0.025), and −31.6 (CI −56.1, −7.1) degrees lower on recording two (p < 0.011). Changes in the unaffected hemisphere over time were non-significant. Figure 2 shows error bars of the phase shift.

### Discussion

This study suggests that DCA is impaired in the affected hemisphere early after moderate-to-severe anterior circulation ischemic stroke. In our cohort autoregulation remained impaired throughout the first week and...
Table 2. Hemodynamic parameters

<table>
<thead>
<tr>
<th></th>
<th>Measurement 1 (n = 24)</th>
<th>Measurement 2 (n = 24)</th>
<th>Measurement 3 (n = 8)</th>
<th>Control subjects (n = 29)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP, mm Hg</td>
<td>107.2±19</td>
<td>105±17.1</td>
<td>104±20.5</td>
<td>87±16.3</td>
</tr>
<tr>
<td>CBFV, cm/s</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Affected</td>
<td>58.5±24.1</td>
<td>51.9±18.1</td>
<td>53.8±19.5</td>
<td>Left 52.8±13.6</td>
</tr>
<tr>
<td>Unaffected</td>
<td>56.5±22.3</td>
<td>54.1±14.5</td>
<td>55.7±14.8</td>
<td>Right 52.6±13.7</td>
</tr>
<tr>
<td>Pulsatility index</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Affected</td>
<td>0.97±0.25</td>
<td>0.97±0.25</td>
<td>0.91±0.17</td>
<td>Left 0.75±0.13</td>
</tr>
<tr>
<td>Unaffected</td>
<td>0.99±0.22</td>
<td>1.02±0.24</td>
<td>0.93±0.16</td>
<td>Right 0.77±0.14</td>
</tr>
<tr>
<td>Phase shift, degrees</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Affected</td>
<td>29.6±10.5</td>
<td>23.2±19.1</td>
<td>53.2±28.2</td>
<td>Left 47±17.5</td>
</tr>
<tr>
<td>Unaffected</td>
<td>42.5±13</td>
<td>41.7±18.5</td>
<td>50.7±29.2</td>
<td>Right 48.7±16.3</td>
</tr>
</tbody>
</table>

Values represent mean ± SD.
then normalized by day ten. Previous studies evaluating the early course of cerebral autoregulation after stroke reported similar results [3–6, 20]. Reinhard et al. reported increasingly disturbed autoregulation in the affected hemisphere within the first days after large anterior circulation ischemic stroke. Their patient selection and methods to measure DCA are comparable to the ones used in our study, and the absolute phase values found in our study are similar to those they reported [3, 4]. However, they focused on the early phase after ischemic stroke, and did not follow their patients beyond the first week. Our study is the first to demonstrate the recovery of autoregulation after acute ischemic stroke.

The pathophysiology for impaired autoregulation is not known. It is believed that tissue lactic acidosis leads to local vasoparalysis in the infarcted area and immediate peri-infarct tissue, inhibiting normal autoregulatory function. Invasive measurements with microdialysis in patients with malignant MCA infarction have shown a close correlation between worsening lactate/pyruvate ratio and impaired autoregulation in the peri-infarct tissue, suggesting that lactate and extracellular fluid spreading along the white matter tracts to the peri-infarct tissue may impair autoregulation in this compartment [21]. This process seems to persist throughout the early post-stroke period, possibly accounting for worsening autoregulation in the first few days after stroke. In this time period, regions of suboptimal flow, which were present within the first hours after stroke, have resolved into an ischemic state with low flow in the infarct, surrounded by histologically normal peri-infarct tissue with normal resting flow. Animal models have confirmed that this peri-infarct tissue nonetheless still exhibits impaired autoregulatory function, at least at 24 h after stroke [22].

DCA is probably normal in the chronic phase after ischemic stroke, although data is limited and methods to assess DCA vary considerably [23–25]. There is even less information at what time recovery of autoregulation occurs. In our study, autoregulation had recovered by day ten after acute anterior circulation ischemic stroke. This has important implications for blood pressure management in the acute stroke period. Instability in blood pressure is very common after ischemic stroke [26] and its management remains unclear. During autoregulatory failure, the brain tissue is particularly vulnerable to blood pressure changes, as they have direct effects on cerebral blood flow. Our data suggests that the period of autoregulatory impairment lasts at least 1 week. During that time careful blood pressure monitoring and tight external regulation of blood pressure may be critical to avoid secondary worsening.

Our findings contrast Dawson et al. who reported persistence of DCA impairment at 7–14 days [27]. However, they used the autoregulatory index (ARI) instead of phase shift to assess DCA. The ARI compares the CBFV response after a rapid step-like decrease in blood pressure with 10 theoretical flow-velocity curves [1, 15]. A single episode of induced hypotension (e.g., thigh cuff deflation) may not measure the same as the response to spontaneous oscillatory changes in blood pressure. The two measurement methods may represent two distinct processes and their recovery after stroke may be dissociated.

In addition, stroke size and etiology may have played a role. Our study population was very uniform and only comprised of patients with partial anterior circulation stroke. The Dawson study also utilized a more heterogeneous study group, including patients with total anterior circulation, posterior circulation and lacunar stroke. Patients with lacunar stroke in particular may have more persistent impairment of DCA, possibly reflecting preexisting endothelial dysfunction due to hypertensive small-vessel disease or metabolic syndrome. This underlying dysfunction may be exacerbated during the acute phase response of an ischemic stroke, when inflammatory

![Fig. 2. Course of DCA in patients with acute MCA Stroke compared to control subjects for phase shift. Bars represent means, whiskers denote standard error. * Indicates significance (p < 0.05) for comparison with control subjects.](image)
changes also affect the cerebral vasculature [4, 28, 29]. This small vessel hypothesis is consistent with the finding of bilateral autoregulatory impairment in patients after lacunar stroke [5]. Thus, the lack of autoregulatory improvement in the Dawson study may be explained at least in part by the inclusion of patients with lacunar stroke who have more persistent impairment of DCA, thereby masking an improvement in autoregulatory function in patients with other stroke subtypes.

This study has limitations including its relatively small sample size in particular at the third measurement, operator-dependence of transcranial Doppler and limited temporal windows for TCD measurement in some patients. In addition, we did not control for white matter hyperintensity or lacunar stroke burden, although the asymmetry between the affected and unaffected hemispheres suggests that the acute infarct itself produced the effect we observed.

In summary, dysautoregulation appears to take place early after moderate-to-severe ischemic stroke and normalize in the subacute period by day ten. Our findings have important implications for blood pressure management in the acute phase of ischemic stroke. Prospective validation of noninvasive measures of cerebral autoregulation should be confirmed in a larger study before individualized blood pressure can be managed routinely based on autoregulatory function.

Disclosure Statement

None.

References


