Dynamic Cerebral Autoregulation in Acute Lacunar and Middle Cerebral Artery Territory Ischemic Stroke

Rogier V. Immink, MD; Gert A. van Montfrans, MD, PhD; Jan Stam, MD, PhD; John M. Karemaker, MD, PhD; Michaela Diamant, MD, PhD; Johannes J. van Lieshout, MD, PhD

Background and Purpose—We addressed whether dynamic cerebral autoregulation (dCA) is affected in middle cerebral artery (MCA) territory (MCAS) and lacunar ischemic stroke (LS).

Methods—Blood pressure (MAP) and MCA velocity (V) were measured in 10 patients with large MCAS (National Institutes of Health Stroke score, 17±2; mean±SEM), in 10 with LS (score, 9±1), and in 10 reference subjects. dCA was evaluated in time (delay of the MCA Vmean counter-regulation during changes in MAP) and frequency domains (cross-spectral MCA Vmean−to-MAP phase lead).

Results—In reference subjects, latencies for MAP increments (5.3±0.5 seconds) and decrements (5.6±0.5 seconds) were comparable, and low frequency MCA Vmean−to-MAP phase lead was 56±5° (left and right hemisphere). In MCAS, these latencies were 4.6±0.7 and 5.6±0.5 seconds in the nonischemic hemisphere and not detectable in the ischemic hemisphere. In the unaffected hemisphere, phase lead was 61±6° versus 26±6° on the ischemic side (P<0.05). In LS, no latency and smaller phase lead bilaterally (32±6 and 33±5°) conformed to globally impaired dCA.

Conclusions—In large MCAS infarcts, dynamic cerebral autoregulation was impaired in the affected hemisphere. In LS, dynamic cerebral autoregulation was impaired bilaterally, a finding consistent with the hypothesis of bilateral small vessel disease in patients with lacunar infarcts. (Stroke. 2005;36:2595-2600.)

Key Words: cerebral circulation ■ spectral analysis ■ ultrasonography

Cerebral blood flow (CBF) is maintained by both fast- and slow-acting regulatory mechanisms spanning the prevailing demands of CBF in everyday life. Static cerebral autoregulation (CA) reflects the overall efficiency, whereas dynamic CA (dCA) refers to the ability to restore CBF in the face of a sudden change in perfusion pressure, that is, the delay of CA. Acute ischemic stroke is often associated with a temporally elevated blood pressure (BP) returning to pre-stroke values within days.1 Adequate BP management is of temporary elevated blood pressure (BP) returning to pre-stroke values within days.1 Adequate BP management is of importance, considering CA impairment in penumbral tissue. Thus, in this setting, CBF is likely to depend on cerebral perfusion pressure or BP. With global CBF detected by cerebral 85Kr and 133Xe clearance, static CA in patients with acute ischemic stroke was demonstrated to be impaired.2–4 In patients with large unilateral middle cerebral artery territory stroke (MCAS), a transient BP rise resulted in increased MCA velocity (V) on the affected side only, suggesting 1-sided impaired static CA.5 In minor MCAS, dCA was found not to be relevantly disturbed.6 In contrast, bilateral dCA impairment was reported but without accounting for the type of stroke.7–9 This study determined bilateral dCA in 2 distinct subtypes of acute ischemic stroke, that is, large MCA territory stroke and in lacunar stroke of the basal ganglia (LS) and in reference subjects (RS).

Methods

Subjects
Ten patients with a first occurrence of MCAS and 10 patients with a first occurrence of LS were included in this study. Ischemic stroke was defined as a sudden onset of a nonconvulsive and focal neurological deficit persisting for >24 hours without signs of a cerebral hemorrhage on computed tomography (CT) scan. Patients were excluded from the study in case of a diminished consciousness (Glasgow Coma Scale <10), an inadequate acoustic window, absence or insufficient quality of one of the MCA V signals, atrial fibrillation, or significant carotid stenosis (>70%). After admission to the stroke unit, stroke severity was quantified (National Institutes of Health Stroke Scale; maximal 42-point score) and antihypertensive medication withdrawn (Table 1).

An ischemic MCAS was diagnosed by neurological examination and CT scan. Six of 10 CT scans showed early signs of MCA infarction, and none revealed leukoaraisis or hemorrhage. We diagnosed LS when a LS (pure motor stroke [in 4 patients], pure sensory stroke [in 1 patient], sensorimotor stroke [in 1 patient], ataxic hemiparesis [in 4 patients], or dysarthria [none]) was present. A CT served to exclude a hemorrhagic cause. A unilateral lacunar lesion was present in 6 of 10 CT scans made within 3 hours after

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written explanation of the objectives and techniques of measure-
ments and risks associated with the study. Written informed con-
sent was provided in accordance with the Helsinki Declaration. The study
protocol was approved by the medical ethics committee of the
Academic Medical Center, University of Amsterdam, the Nether-
lands (MEC 00/061).

Protocol and Measurements
Measurements were performed at the stroke unit between 8:00 and
11:00 PM within 72 hours after onset of stroke with the patients in
supine position, in a quiet and dimly lit room to minimize arousal.
RS were studied at the same time of the evening ≥2 hours after a
light meal without caffeine-containing beverages. Recordings were
obtained for ≥15 minutes. Brachial BP was measured before the
measurements. Finger BP was measured (Portapres M2; Netherlands
Organization for Applied Scientific Research, Biomedical Instrument-
ation, TNO-BMI) with the cuff applied to the midphalanx of the
middle finger of the nonparalytic hand at heart level. To detect
changes in BP correctly, a built-in expert system (Physiocal) tracked
the unloaded diameter of the finger artery to establish and to adjust
the arterial unloaded volume.10 The transcranial Doppler-derived
blood V was measured in the proximal segments of the left and right
MCA (DWL Multidop X4). Insonation with a 2.5-MHz probe
located the MCA on the nonaffected side. Then, the patency of the
insonated vessel was evaluated more proximally. Once the internal
carotid artery bifurcation was identified, the velocity signal was
optimized with forward and reversed V set at equal intensity. We
checked the patency of the carotid arteries by Doppler imaging with a
4.5-MHz to 5.5-MHz transducer (Hewlett Packard SONOS 2000).
Next, we located the MCA on the affected side, and waveform and
systolic flow acceleration were compared with those on the nonaf-
fected side. In case of a blunted waveform, patients were excluded
from the study. The probe was secured with a headband (Mark 600,
Spencer Technologies). End-tidal CO tension (Petco2) was monitored by a
sampling infrared capnograph (Datex Normocap 200).

Data Analysis
Finger BP, MCA V spectrum envelope, and PetCO2 were A/D
100-Hz converted and stored. Mean arterial pressure (MAP) and
MCA Vmean were the integral over 1 beat, and heart rate was taken
from the pressure interval. MAP and MCA Vmean data were interpo-
lated and resampled for time-domain dCA analysis. We selected
manually upward and downward BP transients lasting ≥10 seconds
at stable PetCO2. All of the 10-second episodes per 15 minutes of
tracing were isolated, and averaged curves of all of the 10-second
upward and downward BP transients were plotted. dCA is consid-
ered adequate with MCA Vmean remaining stable during a change in
MAP, apart from ≥5-second delay, and impaired when MCA Vmean
follows MAP passively.11 Frequency domain analysis of dCA was
performed as reported.11 In short, for examination of dCA in the
frequency domain, a 2-minute tracing of beat-to-beat data of MAP

### TABLE 1. Subject Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>References, n=10</th>
<th>MCAS, n=10</th>
<th>LS, n=10</th>
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</thead>
<tbody>
<tr>
<td>Male:Female</td>
<td>5:5</td>
<td>5:5</td>
<td>8:2</td>
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<tr>
<td>Age, y</td>
<td>57±2</td>
<td>59±5</td>
<td>63±3</td>
</tr>
<tr>
<td>Weight, kg</td>
<td>72±4</td>
<td>83±7</td>
<td>81±3</td>
</tr>
<tr>
<td>Length, cm</td>
<td>173±3</td>
<td>172±4</td>
<td>176±2</td>
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<td>Infarct side, L/R</td>
<td>6:4</td>
<td>4:6</td>
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<tr>
<td>NIHSS</td>
<td>17±2*</td>
<td>9±1</td>
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</tr>
<tr>
<td>Hypertension</td>
<td>6</td>
<td>7</td>
<td></td>
</tr>
<tr>
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<td>4</td>
<td></td>
</tr>
<tr>
<td>Hyperlipidemia</td>
<td>2</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>History of:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate, min⁻¹</td>
<td>86±9</td>
<td>78±5</td>
<td></td>
</tr>
<tr>
<td>Systolic BP, mm Hg</td>
<td>166±9</td>
<td>178±16</td>
<td></td>
</tr>
<tr>
<td>Mean BP, mm Hg</td>
<td>116±4</td>
<td>121±11</td>
<td></td>
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<tr>
<td>Diastolic BP, mm Hg</td>
<td>86±3</td>
<td>93±9</td>
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<tr>
<td>Glucose, mmol/L⁻¹</td>
<td>5.30±1.20</td>
<td>6.60±0.49</td>
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<tr>
<td>Cholesterol–total, mmol/L⁻¹</td>
<td>5.18±0.38</td>
<td>6.52±0.40</td>
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<td>Cholesterol–HDL, mmol/L⁻¹</td>
<td>1.01±0.10</td>
<td>1.23±0.12</td>
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<tr>
<td>Cholesterol–LDL, mmol/L⁻¹</td>
<td>3.08±0.27</td>
<td>3.41±0.37</td>
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<tr>
<td>Triglycerides, mmol/L⁻¹</td>
<td>2.09±0.36</td>
<td>2.28±0.39</td>
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<tr>
<td>ß blocker</td>
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<td>5</td>
<td></td>
</tr>
<tr>
<td>ACE-inhibitor</td>
<td>2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>ATIIRA</td>
<td>0</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Diuretic</td>
<td>2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Cholesterol-lowering drug</td>
<td>1</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Glucose-lowering drug</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

*LR indicates left vs right sided infarct; NIHSS, National Institutes of Health Stroke Scale; ACE, angiotensin-converting enzyme; ATIIRA, angiotensin II receptor antagonist. *P<0.05 vs lacunar stroke.

### TABLE 2. BP and MCA Blood Velocity

<table>
<thead>
<tr>
<th>Variable</th>
<th>Type</th>
<th>Brachial BP, mm Hg</th>
<th>Finger BP, mm Hg</th>
<th>MCA Vmean, cm/s⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reference</td>
<td>Systolic</td>
<td>127±7</td>
<td>115±6</td>
<td>92±6</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>95±4</td>
<td>78±3</td>
<td>61±3</td>
</tr>
<tr>
<td></td>
<td>Diastolic</td>
<td>79±3</td>
<td>59±2</td>
<td>39±2</td>
</tr>
<tr>
<td>MCAS</td>
<td>Systolic</td>
<td>143±9</td>
<td>125±7</td>
<td>86±9</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>103±5</td>
<td>79±5</td>
<td>51±6</td>
</tr>
<tr>
<td></td>
<td>Diastolic</td>
<td>83±3</td>
<td>59±5</td>
<td>30±3*#</td>
</tr>
<tr>
<td>LS</td>
<td>Systolic</td>
<td>156±10*</td>
<td>145±8*</td>
<td>86±5</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>111±8</td>
<td>89±5</td>
<td>53±4</td>
</tr>
<tr>
<td></td>
<td>Diastolic</td>
<td>88±8</td>
<td>61±5</td>
<td>32±3*</td>
</tr>
</tbody>
</table>

*P<0.05 vs reference; #P<0.05 vs ischemic side.
and MCA Vmean was spline interpolated and resampled at 4 Hz. With discrete Fourier transform, the MAP and MCA Vmean variability was expressed as the integrated areas in the very low frequency (LF; 0.02 to 0.07 Hz), low frequency (LF; 0.07 to 0.15 Hz), and the high frequency (HF; 0.15 to 0.3 Hz) ranges. Blood pressure fluctuations in the HF range are induced mainly by respiration, whereas those in the LF range are independent of such influence and reflect primary baroreflex activity. Furthermore, the very LF range of both flow and

**Figure 1.** MCA velocity during a change in blood pressure in RS and in patients with large MCA territory and lacunar stroke; grey circles (all panels): percentage change in blood pressure (Δ BP); top: percentage changes in left (○) and right (●) mean MCA blood velocity (Δ MCA Vmean); middle and bottom: ischemic (○) and nonischemic (●) hemisphere. Data are mean±SEM.

**TABLE 3. Variability in BP and MCA Blood Velocity**

<table>
<thead>
<tr>
<th>Variable</th>
<th>MAP</th>
<th>MCA Vmean Variability (ΔMCA Vmean/Hz²)</th>
<th>Coherence</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Variability, mm Hg/Hz²</td>
<td>Nonischemic</td>
<td>Ischemic</td>
</tr>
<tr>
<td>Reference, n=10</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>8.5±2.4</td>
<td>6.0±1.3</td>
<td>6.1±1.4</td>
</tr>
<tr>
<td>LF</td>
<td>4.0±0.8</td>
<td>2.1±0.4</td>
<td>2.2±0.5</td>
</tr>
<tr>
<td>HF</td>
<td>0.8±0.2</td>
<td>1.2±0.3</td>
<td>1.3±0.3</td>
</tr>
<tr>
<td>MCAS, n=10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>13.8±3.9</td>
<td>5.5±2.0</td>
<td>6.5±1.6</td>
</tr>
<tr>
<td>LF</td>
<td>5.0±1.7</td>
<td>1.7±0.3</td>
<td>2.3±0.5</td>
</tr>
<tr>
<td>HF</td>
<td>3.5±0.9</td>
<td>2.9±0.8</td>
<td>2.1±0.4</td>
</tr>
<tr>
<td>LS, n=10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>31.1±13.3*</td>
<td>6.9±2.7</td>
<td>9.3±3.9</td>
</tr>
<tr>
<td>LF</td>
<td>4.5±1.1</td>
<td>1.2±0.2</td>
<td>1.6±0.4</td>
</tr>
<tr>
<td>HF</td>
<td>2.0±0.5</td>
<td>1.1±0.3</td>
<td>1.2±0.5</td>
</tr>
</tbody>
</table>

MAP indicates mean arterial pressure; VLF, very low frequency. *P<0.05 vs reference.
pressure variability appears to reflect multiple physiological mechanisms, which confound interpretation. Thus, the LF range was used to identify dCA. From the MAP–MCA V mean cross-spectrum, transfer function gain (cm/s/Hg) and MCA V mean-to-MAP phase lead (degrees) were calculated. The transfer function gain for MAP and MCA V mean (in percentage change in cm/s percentage change in mm Hg) was normalized to account for intersubject variability. The cutoff value for preserved CA depends on the frequency range used. We consider a phase lead ≥50° for the LF range to represent preserved CA.11 With dCA impaired, as in patients with carotid artery obstruction12 or malignant hypertension,11 phase lead decreases to as low as ~30°.

Frequency Domain
In RS, LF MCA V mean-to-MAP phase lead was 56±5° (left hemisphere) and 59±5° (right hemisphere). In MCAS, phase lead was lower at the ischemic (26±6°; range, 10 to 50°; *P<0.05) versus nonischemic side (61±6°; range, 17 to 73°) with phase lead ≥50° in 6 patients and with 50° phase lead in 1 patient (Table 3). In LS, LF phase lead was reduced bilaterally (nonischemic: 32±6°; range, 5 to 49° versus ischemic hemisphere: 33±5°; range, 6 to 54°; #P<0.05), and gain did not differ between the hemispheres (Figure 2).

Discussion
This study determined dynamic cerebral autoregulatory capacity in the time and frequency domain in 2 main subtypes of acute ischemic stroke: large MCAS and lacunar stroke. The main finding is that in acute MCAS, dCA is affected at the ischemic side only, whereas in acute LS, dCA is impaired bilaterally.

CA characterizes cerebral artery ability to modulate arterial wall smooth muscle tone, maintaining CBF. The latency in dCA as derived from spontaneous MAP decreases provides confirmatory information regarding impaired CA, as well as...
produced data akin to rapid thigh-cuff deflation as a stimulus for dCA.\textsuperscript{13}

CA in stroke is controversial. In large MCAS, a transient rise in BP by phenylephrine raises MCA $V_{\text{mean}}$ on the affected side only, indicating unilaterally impaired static CA.\textsuperscript{3} Several\textsuperscript{2}–\textsuperscript{4} but not all\textsuperscript{7} reports suggest global static CA impairment. So far, global bhemispheric dCA impairment in acute and subacute stroke with preserved static CA has been reported without obvious differences between stroke subtypes.\textsuperscript{7}–\textsuperscript{9} In subsequent studies, dCA appeared impaired bilaterally\textsuperscript{8} and remained so for at least 1 to 2 weeks,\textsuperscript{9} although this was not confirmed in minor MCAS.\textsuperscript{6} MCA territory infarcts are likely to result from embolism or large cerebral artery artherothrombosis with a considerable penumbral tissue volume. The present study extends the observations of unilaterally impaired static CA in MCAS\textsuperscript{5} to unilateral impairment of dCA.

In contrast to what was found in large MCAS, dCA was affected bilaterally in 1-sided LS. Intracranial small artery occlusion is the probable cause in the majority of LS.\textsuperscript{14} The penumbral volume in LS is usually small, and we consider dCA impairment in LS by the ischemic event itself unlikely.\textsuperscript{15}

In syndromes with more extensive atherosclerosis of both extracranial and intracranial vessels, such as longstanding diabetes, CA is impaired.\textsuperscript{16} Cerebral small-vessel disease can produce isolated lacunar infarcts or diffuse white matter changes appearing as leukoaraiosis on a CT or MRI,\textsuperscript{17,18} which may explain why dCA in LS was affected on the nonischemic side as well. We examined CA capacity in acute stroke patients. In a follow-up study of patients experiencing minor MCAS, CA was still abnormal on the affected side but preserved on the normal side >2 months after stroke onset.\textsuperscript{19} We did not assess CA capacity before LS, but we speculate that a globally impaired dCA was already present.

In this study, CT scanning was the primary imaging technique used on admission. More ischemic strokes show up on diffusion-weighted imaging than on CT or conventional MRI in the first few hours. However, MRI can be difficult to use routinely in acute stroke and may not identify hyperacute intracerebral hemorrhage correctly, whereas CT scans are superior for differentiating ischemic from hemorrhagic stroke.\textsuperscript{18,20} In addition, lacunar syndromes are highly predictive for small deep infarcts on magnetic resonance. For that reason, brain MRI has even been considered redundant in the setting of a lacunar syndrome if supported by a CT that excludes nonischemic causes of stroke.\textsuperscript{21} An inherent difficulty of both CT and MRI is the ability to differentiate between acute and chronic lesions. In spite of the problem of relating lacunar syndromes to certain locations of lacunae,\textsuperscript{22} our finding that CA capacity was impaired also in the contralateral (“healthy”) hemisphere in LS remains unchallenged.

Critical for the interpretation of the data are to what extent MCA $V_{\text{mean}}$ reflects volume flow. The MCA $V_{\text{mean}}$ is assumed to represent flow velocity in the center of the vessel, and, when flow is laminar, MCA $V_{\text{mean}}$ follows $^{133}$Xe clearance-determined CBF.\textsuperscript{23} Thus, the constancy of vessel diameter links changes in MCA V to those in CBF. When nonlaminar blood flow in the affected cerebral arteries is considered, MCA $V_{\text{mean}}$ may change out of proportion to flow, and, therefore, the analysis was restricted to normally shaped Doppler signals.\textsuperscript{24}

CA was assessed after discontinuation of antihypertensive treatment, but a remaining biological effect cannot be excluded. However, also with β blockade, angiotensin-converting enzyme inhibition or angiotensin-receptor antagonist treatment interplay of CA has been confirmed. We did not create different steady-state BP levels, and quantification of static CA as established by Schwarz et al\textsuperscript{5} was not carried out. Impairment of static CA by pharmacological manipulation also affects dCA,\textsuperscript{25} and we, therefore, consider that the impaired dCA found in this study suggests impairment of static CA as well.\textsuperscript{11}

Conclusions

In large MCAS infarcts, dCA was impaired in the affected hemisphere only. In LS, dCA was impaired bilaterally. The latter finding is consistent with the hypothesis of bilateral small vessel disease in patients with lacunar infarcts.

Acknowledgments

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References

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