Cerebrovascular Collaterals Correlate with Disease Severity in Adult North American Patients with Moyamoya Disease


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ABSTRACT

BACKGROUND AND PURPOSE: Cerebrovascular collaterals have been increasingly recognized as predictive of clinical outcomes in Moyamoya disease in Asia. The aim of this study was to characterize collaterals in North American adult patients with Moyamoya disease and to assess whether similar correlations are valid.

MATERIALS AND METHODS: Patients with Moyamoya disease (n = 39; mean age, 43.5 ± 10.6 years) and age- and sex-matched control subjects (n = 33; mean age, 44.3 ± 12.0 years) were graded via angiography. Clinical symptoms of stroke or hemorrhage were graded separately by imaging. Correlations between collateralization and disease severity, measured by the modified Suzuki score, were evaluated in patients with Moyamoya disease by fitting a regression model with clustered ordinal multinomial responses.

RESULTS: The presence of leptomeningeal collaterals ( \( P = .008 \) ), dilation of the anterior choroidal artery ( \( P = .01 \) ), and the posterior communicating artery/ICA ratio ( \( P = .004 \) ) all correlated significantly with disease severity. The presence of infarct or hemorrhage and posterior steno-occlusive disease did not correlate significantly with the modified Suzuki score ( \( P = .1 \) ). Anterior choroidal artery changes were not specific for hemorrhage. Patients with Moyamoya disease were statistically more likely than controls to have higher posterior communicating artery/ICA ratios and a greater incidence of leptomeningeal collaterals.

CONCLUSIONS: As with Moyamoya disease in Asian patients, the presence of cerebrovascular collaterals correlated with the modified Suzuki score for disease severity in North American patients with Moyamoya disease. However, anterior choroidal artery changes, which correlated with increased rates of hemorrhage in Asian studies, were not specific to hemorrhage in North Americans.

ABBREVIATIONS: AchoA = anterior choroidal artery; LMC = leptomeningeal collaterals; MMD = Moyamoya disease; mSS = modified Suzuki score; P1 = proximal segment of the posterior cerebral artery; PCA = posterior cerebral artery; PcomA = posterior communicating artery

Moyamoya disease (MMD) has been largely described by the Asian experience, yet varying epidemiologic and clinical features in North Americans with MMD suggest that pathophysiologic differences exist. MMD in North Americans and Europeans most commonly affects young women in the third-to-fourth decades of life, whereas MMD in Asians typically begins in childhood. North American and European patients present with ischemic stroke or TIA as the most frequent manifestation, whereas hemorrhage is more frequent in Asian cohorts. Additional, gene associations found in Asian cohorts have not been replicated by European investigators.

Shared features for all MMD include predominantly anterior circulation involvement with progressive arteriopathy involving the ICA and proliferation of distinctive basal vessels—changes that have been well-correlated with disease severity. Other collaterals, including dilated anterior choroidal arteries (AchoAs) and posterior communicating arteries (PcomAs), are less well-studied, particularly in North Americans. Histologic analysis of these collaterals has demonstrated evidence of stress related to increased flow, which may predispose patients to hemorrhage. Dilated anterior choroidal and posterior communicating arteries have been shown to be strongly predictive of hemorrhage in Asian MMD. The purpose of this study was to assess correlations between disease severity with less well-characterized collaterals, in-
including the PcomA/ICA ratio, leptomeningeal collaterals (LMC), and AchoA changes, in a cohort of adult North Americans with MMD. Our primary hypothesis is that these collaterals correlate variably with disease severity, as measured by the modified Suzuki score (mSS) and clinical findings of stroke or hemorrhage. Our secondary hypothesis is that these collaterals correlate with disease compared with control subjects.

### MATERIALS AND METHODS

This retrospective review was approved by the local institutional review board. Subjects were identified by an electronic medical record search for all adult patients with MMD who underwent DSA from 2002 to 2012 at our institution. Idiopathic MMD was defined as mSS I–IV in at least 1 cerebral hemisphere without associated predisposing disease. Control subjects were selected sequentially from angiograms obtained for any non-Moyamoya indication. Two certified neuroradiologists, blinded to clinical and imaging findings, graded disease severity separately and resolved disagreement by consensus. Modifications have been made to the Suzuki classification so that the score can be applied to individual cases, rather than longitudinally. The mSS used for this study (Table 1) includes 5 stages of disease severity, examples of which are shown in Fig 1. In separate sessions divided by >2 weeks, the following were graded from DSA:

1) LMC circulation to the anterior circulation was classified into 2 stages: Stage 1, leptomeningeal cortical branches were found coursing from the posterior cerebral artery (PCA) to the frontal, temporal, or parietal Lobes; Stage 2, there was no leptomeningeal collateral circulation. 

2) Ratio of the PcomA lumen diameter to the ipsilateral pre-cavernous ICA lumen diameter is shown in Fig 2. Because the distal ICA is stenosed in MMD, the ICA measurement was performed at the precavernous portion with the widest, parallel lumen. If a PcomA infundibulum was present, the diameter was measured distal to infundibulum.

3) Proximal segment of the posterior cerebral artery (P1) steno-occlusive changes were noted (1, no stenosis or occlusion; 2, stenosis or occlusion). An endovascular neurosurgeon (with 21 years of experience), blinded to clinical and imaging findings, separately graded the AchoA (zero, normal; 1, dilated with distal branching; and 2, dilated with extensive branching).

### Table 1: Modified Suzuki scoring

<table>
<thead>
<tr>
<th>Score</th>
<th>Description of classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No evidence of disease</td>
</tr>
<tr>
<td>I</td>
<td>Mild-to-moderate stenosis around ICA bifurcation with absent or slightly developed ICA MMD*</td>
</tr>
<tr>
<td>II</td>
<td>Severe stenosis around the ICA bifurcation or occlusion of either proximal anterior or MCA branches with well-developed ICA MMD</td>
</tr>
<tr>
<td>III</td>
<td>Occlusion of both anterior and MCA branches with well-developed ICA MMD (only a few of anterior or MCA branches or both are faintly opacified in antegrade fashion through meshwork of ICA MMD)</td>
</tr>
<tr>
<td>IV</td>
<td>Complete occlusion of both anterior and MCA branches with absent or small amount of ICA MMD (without opacification of either anterior or MCA branches in antegrade fashion)</td>
</tr>
</tbody>
</table>

* ICA Moyamoya disease indicates perforating collateral vessels at or around the terminal ICA.
Table 2: Collateral and clinical characteristics by modified Suzuki score in subjects with Moyamoya disease*

<table>
<thead>
<tr>
<th>Metric [P Value]</th>
<th>N</th>
<th>0 (n = 6)</th>
<th>I (n = 7)</th>
<th>II (n = 30)</th>
<th>III (n = 20)</th>
<th>IV (n = 15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PcomA/ICA (0.024)</td>
<td>64</td>
<td>0.125</td>
<td>0.205</td>
<td>0.295</td>
<td>0.330</td>
<td>0.520</td>
</tr>
<tr>
<td>LMC (.001)</td>
<td>78</td>
<td>17% (1)</td>
<td>29% (2)</td>
<td>63% (19)</td>
<td>80% (16)</td>
<td>93% (14)</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>83% (5)</td>
<td>71% (5)</td>
<td>37% (11)</td>
<td>20% (4)</td>
<td>7% (1)</td>
</tr>
<tr>
<td>2</td>
<td>3</td>
<td>83% (5)</td>
<td>57% (4)</td>
<td>37% (11)</td>
<td>15% (3)</td>
<td>33% (5)</td>
</tr>
<tr>
<td>CT/MR (.06c)</td>
<td>78</td>
<td>17% (1)</td>
<td>42% (3)</td>
<td>60% (18)</td>
<td>70% (14)</td>
<td>67% (10)</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>0% (0)</td>
<td>0% (0)</td>
<td>3% (1)</td>
<td>15% (3)</td>
<td>0% (0)</td>
</tr>
<tr>
<td>3</td>
<td>0% (0)</td>
<td>14% (1)</td>
<td>50% (15)</td>
<td>85% (17)</td>
<td>33% (5)</td>
<td></td>
</tr>
<tr>
<td>AchoA (&lt;.001)</td>
<td>78</td>
<td>83% (5)</td>
<td>43% (3)</td>
<td>13% (4)</td>
<td>5% (1)</td>
<td>7% (1)</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>17% (1)</td>
<td>43% (3)</td>
<td>37% (11)</td>
<td>10% (2)</td>
<td>0% (0)</td>
</tr>
<tr>
<td>2</td>
<td>0% (0)</td>
<td>94% (1)</td>
<td>50% (15)</td>
<td>85% (17)</td>
<td>33% (5)</td>
<td></td>
</tr>
<tr>
<td>NV</td>
<td>0% (0)</td>
<td>0% (0)</td>
<td>0% (0)</td>
<td>0% (0)</td>
<td>60% (9)</td>
<td></td>
</tr>
<tr>
<td>PI (.14)</td>
<td>78</td>
<td>100% (6)</td>
<td>86% (6)</td>
<td>90% (27)</td>
<td>95% (19)</td>
<td>67% (10)</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>100% (6)</td>
<td>86% (6)</td>
<td>90% (27)</td>
<td>95% (19)</td>
<td>67% (10)</td>
</tr>
<tr>
<td>2</td>
<td>0% (0)</td>
<td>14% (1)</td>
<td>10% (3)</td>
<td>5% (1)</td>
<td>33% (5)</td>
<td></td>
</tr>
</tbody>
</table>

Note: N indicates the number of nonmissing values; NV, not visualized.

* Numbers after percentages are frequencies.

* PcomA/ICA ratio provided is median.

* CT/MR 1—no infarct or hemorrhage, 2—infarct, 3—hemorrhage.

FIG 3. PcomA/ICA ratio in subjects with MMD versus control subjects by hemisphere. The PcomA/ICA ratio in a patient with MMD (n = 1) was significantly higher (P < .001) compared with control subjects (n = 0). Orange dots are observations for the left cerebral hemisphere, and blue dots are for the right cerebral hemisphere. The dark band represents the median in this boxplot.

The site of ICA occlusion was identified in subjects when the AchoA was not visualized. Hemorrhage or infarct or both were graded on CT or MR imaging performed within 30 days of DSA, and the location of the infarct was classified by territory. Exclusion criteria were the following: 1) For PcomA/ICA ratios, the hemisphere was excluded if the PcomA was not visualized on DSA; 2) both hemispheres were excluded for surgical revascularization before DSA; and 3) for AchoA grading, the hemisphere was excluded if the AchoA was not visualized.

Statistical Analysis

For univariate analysis for the association of the disease severity (mSS) with other collateral and clinical characteristics, the Kruskal-Wallis or Fisher exact test (2-sided) was used. For the comparison of the PcomA/ICA ratio between subjects with MMD and controls, the Wilcoxon rank sum test was used. Because hemispheres were considered separately, a multivariate regression model based on generalized estimating equations23 for clustered ordinal responses with uniform local odds ratio structure was fit to evaluate the correlations between collateralization and disease severity among subjects with MMD. Hemispheres with an mSS of zero were included in the analysis because the typical progression of MMD is bilateral involvement.24 Due to the small number of observations for mSS stages zero and I (6 and 7, respectively), we combined them as stage 1 in the regression model. Asian and African American races were also combined as nonwhite. For multivariate analysis, the CT/MR imaging findings of infarct and hemorrhage were combined. The Fisher exact test was used to correlate the incidence of hemorrhage in patients of Asian descent with non-Asian patients. All MMD statistics were adjusted by age, sex, race (white or nonwhite), LMC, the presence of infarct or hemorrhage, PI and AchoA classification, and PcomA/ICA ratio.

RESULTS

We examined angiograms from 39 subjects with MMD (78 cerebral hemispheres). The mean age was 43.5 ± 10.6 years (range, 26–64 years). The median age was 44 years. Twenty-eight patients were women. Age- and sex-matched controls (n = 33; mean age, 44.3 ± 12.0 years) were included. Subjects with MMD were white (n = 28), African American (n = 7), and Asian (n = 4); control subjects were white (n = 27), African American (n = 5), and Hispanic (n = 1).

Fourteen hemispheres (9 left, 5 right) were excluded from measurement of the PcomA/ICA ratio secondary to lack of ipsilateral PcomA (n = 2), lack of DSA lateral projection (n = 1), prior aneurysm coiling (n = 1), and ipsilateral ICA occlusion (n = 10). All hemispheres excluded due to ICA occlusion had mSS grades of IV. Six subjects with MMD (15%) had unilateral involvement. Collateral characteristics by mSS are summarized in Table 2. Interobserver agreement for mSS rating meets acceptable statistical criteria with a Fleiss-Cohen κ statistic of 0.845 (95% CI, 0.785–0.904).
tion between mSS and the presence of LMC ($P = .008$) for subjects. The OR of having a more severe mSS classification was 4.79 times higher (95% CI, 1.51–15.21) for MMD hemispheres with LMC, compared with those without LMC. Only 2 of 66 hemispheres in control subjects had LMC (1 with a history of seizures and 1 with previously coiled aneurysms, but neither with vascular stenosis). Figure 5 demonstrates the appearance of LMC in 1 subject with MMD. All hemispheres with P1 steno-occlusive involvement had LMC. However, P1 steno-occlusive change was not significantly associated with mSS ($P = .485$).

AchoA grades are shown in Fig 6. Table 3 shows the distribution of AchoA grades by imaging findings. There was a significant association between the AchoA classification and mSS ($P = .02$). The OR of having a more severe mSS classification was 2.76 times higher (95% CI, 0.57–13.24) for hemispheres with grade I AchoA versus control subjects ($P = .21$), and the OR increased to 17.2 (95% CI, 2.26–131.1) when comparing grade II AchoA with control subjects ($P = .01$). In 9 hemispheres, the AchoA was occluded due to ICA occlusion proximal to the AchoA origin and lack of collateral AchoA filling via posterior collaterals. All such hemispheres were mSS IV; none had hemorrhage and 5 of 9 had infarcts. All hemispheres with hemorrhage (4 of 78) had AchoA grade 2, and none had P1 steno-occlusive findings.

There was no statistically significant association between mSS and imaging findings of infarct or hemorrhage ($P = .11$). Forty-six of 78 MMD hemispheres (59%) had infarcts. Of 15 mSS hemispheres, 5—including the only mSS IV hemisphere without LMC—had no infarcts, 2 had infarcts involving the ipsilateral basal ganglia, and all remaining mSS hemispheres had a watershed pattern of infarcts. No patient with mSS IV had posterior circulation or cortical MCA territory infarcts.

Two of 4 subjects with MMD with hemorrhage were of Asian descent. The Fisher exact test gave a 2-sided $P = .045$ for the correlation between the incidence of hemorrhage in patients of Asian descent with non-Asian patients, though findings were limited by the low number of hemispheres with hemorrhage. Only 1/78 MMD hemispheres had both hemorrhage and infarct on imaging. This hemisphere had no P1 steno-occlusive changes or LMC and had grade 2 AchoA changes.

Median follow-up time for subjects with angiography (19 of 39 subjects with MMD) was 463 days (minimum, 105 days; maximum, 1740 days).

**DISCUSSION**

The primary findings of this work are the following: 1) As with prior studies on adult MMD, we observed a strong correlation between vascular collaterals, including the presence of LMC, dilated AchoAs, and larger PcomA/ICA ratios, and mSS in adult MMD. Despite these collateral networks, most subjects with MMD in our study had infarcts, and the presence of infarct and/or hemorrhage did not correlate with disease severity, as reflected by the mSS, due to the high frequency of infarcts in subjects with MMD with less severe mSSs (eg, 60% of mSSs II subjects). 2) In contrast to Asian studies, AchoA changes were sensitive but not specific for intracranial hemorrhage, and P1 steno-occlusive involvement did not appear to correlate with disease severity. Taken together, these findings suggest that mSS may not provide a comprehensive predictive model for the complex hemodynamic stress, which causes infarcts and hemorrhages in subjects with MMD. Additionally, as more nuanced predictive models arise, such as AchoA changes, our findings suggest the models may not generalize across ethnic cohorts.

The ischemic stress in MMD is imparted by progressive anterior circulation stenosis, which may include vascular constrictive changes, which narrows the terminal ICA and, in patients with mSS III–IV, occludes both the anterior and middle cerebral arteries. When ICA stenosis occurs distal to the AchoA, both the AchoA and PcomA may be subjected to increased collateral flow and
The correlation of the PcomA:ICA ratio in our study with disease severity may reflect a compensation mechanism for the hemodynamic stress imposed by progressive ICA arteriopathy. PcomA collateral flow has been shown to be protective against watershed infarcts in patients with atherosclerotic ICA occlusion. Although ICA stenosis in MMD can occlude the PcomA, stenosis typically occurred distal to the PcomA origin in our cohort (87%). The high rate of mSS IV in subjects with MMD in our study with a watershed pattern of infarcts (85% of mSS IV hemispheres with infarcts) suggests inadequate flow despite increased collaterals.

As in other North American studies, our cohort was much more likely to present with ischemia (59%) than hemorrhage (5%). In contrast, Asian studies reported higher rates of hemorrhage, ranging from 25% to 62% (Japan, China, and Korea) compared with 10%–29% (Iowa and Hawaii). Hemorrhage has been shown to be the most significant factor affecting poor outcome in Asian cohorts, and ruptured thin-walled basal perforators have been cited as the source of high rates of hemorrhage. However, studies have not shown a correlation between the reduction in perforators following surgery and rates of hemorrhage. More recently, Asian studies have implicated abnormal branching of the AchoA and dilation of the PcomA as a hemorrhagic source, particularly in subjects with intraventricular hemorrhage.

Grade 1 or 2 AchoA changes were found in 89% of hemispheres with hemorrhage compared with only 44% of ischemic hemispheres in Asian MMD. However, grade 1 or 2 AchoA changes were present in most (80%) subjects with MMD in our study and did not differentiate hemorrhage and infarct. Despite this high rate of AchoA changes, only 4 hemispheres presented with hemorrhage, compared with 27% in the Japanese cohort. These findings suggest that AchoA changes in North Americans may not correlate with increased risk of hemorrhage to the extent that has been found in Asian cohorts. The impact of AchoA occlusion on future hemorrhage and ischemia risk is not clear from our study due to the small sample size and relatively short median follow-up. Only 1 MMD hemisphere had both hemorrhage and infarct, which is in line with published studies, suggesting the etiology for infarct and hemorrhage may be distinct. The incidence of P1 steno-occlusive changes in our cohort (12%) was lower than that reported in Asian studies (20%–43%).

In our study, LMC correlated significantly with disease severity. The hemodynamic importance of LMC to the anterior circulation in MMD has recently been suggested in a Japanese cohort with P1 disease, in which P1 lesions led to MCA territory infarcts much more frequently than posterior infarcts. P1 stenosis in patients with MMD following revascularization was associated with decreased LMC, which may increase ischemic symptoms. LMC have been shown to be independent predictors of an increased oxygen extraction fraction in patients with atherosclerotic ICA occlusion.

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Certainly, we could have used other published grading scales. Togao et al described a 4-stage system in their study, building a multivariate model for the angiographic findings in subjects with MMD, ranging from no PCA occlusive change in stage 1 to PCA occlusion with almost no visualization of distal branches in stage 4. Using this system, Togao et al found 32 stage 1 PCA arteries, 1 stage 2 artery, 5 stage 3 arteries, and 2 stage 4 arteries. Our study had an even larger percentage of subjects with MMD (87%) without PCA steno-occlusive disease. Because only 10 hemispheres...
had PCA steno-occlusive disease, subcategorization of PCA by stenosis degree was not thought to be statistically robust.

The findings of this work should be considered in the context of 4 limitations. First, as with most MMD studies, this study was limited by sample size ($n = 39$), owing to the relatively small prevalence of MMD in the general population (3 cases per 100,000). This led to a relatively small number of hemispheres with hemorrhage. However, this is one of the larger studies of North American MMD; therefore, we believe the results presented should be used as an exemplar for motivating larger studies. Second, constraints are imposed by the retrospective design, which precluded direct PcomA luminal diameter measurement due to differences in magnification on DSA. The PcomA/ICA ratio was, therefore, used as a surrogate. The distal ICA diameter has been shown to decrease in patients with MMD, either due to vascular constrictive changes or as a flow-related phenomenon due to more distal disease; thus, the precavernous ICA was measured. Third, our controls included subjects with intracranial pathology, including aneurysms and vascular stenosis, a limitation imposed by the standard for use of DSA at our institution, where DSA is reserved for subjects with suspected intracranial pathology, due to its invasive nature. Finally, our study represents a static view of what is known to be a dynamic disease. The 20 subjects with MMD without follow-up data, and the relatively short follow-up for several other subjects with MMD preclude drawing conclusions on whether the variables measured have prognostic implications. Longitudinal studies of these collateral pathways are necessary to understand the interplay between ischemic and hemorrhage risks.9,13,42

CONCLUSIONS

MMD in North Americans remains poorly characterized: The etiology is unknown; the natural history, unclear; and optimal treatment, untested. The varied clinical course for MMD may reflect the effectiveness of collaterals in compensating for the hemodynamic stress imposed by progressive ICA arteriopathy without imparting increased risk of hemorrhage. In our analysis, the presence of LMC, dilation of the AchoA, and the PcomA/ICA ratio were independent predictors of mSS severity for North Americans with MMD. However, AchoA changes did not correlate with increased risk of hemorrhage to the extent that has been found in Asian cohorts.

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REFERENCES