Direct Visualization of Arterial Emboli in Moyamoya Syndrome

The Harvard community has made this article openly available. **Please share** how this access benefits you. Your story matters.

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Published Version</td>
<td>doi:10.3389/fneur.2017.00425</td>
</tr>
<tr>
<td>Citable link</td>
<td><a href="http://nrs.harvard.edu/urn-3:HUL.InstRepos:34492326">http://nrs.harvard.edu/urn-3:HUL.InstRepos:34492326</a></td>
</tr>
<tr>
<td>Terms of Use</td>
<td>This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at <a href="http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA">http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA</a></td>
</tr>
</tbody>
</table>
Direct Visualization of Arterial Emboli in Moyamoya Syndrome

Julie G. Shulman1*, Samuel Snider2, Henri Vaitkevicius2, Viken L. Babikian1,3 and Nirav J. Patel4

1Department of Neurology, Boston Medical Center, Boston University School of Medicine, Boston, MA, United States, 2Brigham and Women’s Hospital, Department of Neurology, Harvard Medical School, Boston, MA, United States, 3VA Boston Healthcare System, Department of Neurology, Boston, MA, United States, 4Brigham and Women’s Hospital, Department of Neurosurgery, Harvard Medical School, Boston, MA, United States

Background: Hemodynamic insufficiency is often considered the cause of ischemic stroke in patients with moyamoya syndrome. While high-intensity transient signals (HITS) on transcranial Doppler (TCD) have been reported in this population, the relationship between these signals and ischemic symptoms is not clearly established. Accordingly, current treatment is directed at improving perfusion.

Clinical presentation: We present two patients with symptoms of cerebral ischemia and angiographic findings of moyamoya syndrome. In each case, ischemia may have been caused by either hypoperfusion or embolization. Patient A presented with multifocal right middle cerebral artery (MCA) territory infarctions and angiographic findings consistent with moyamoya disease. She underwent right superficial temporal artery–MCA bypass. Intra-operatively, embolic material was observed and recorded traveling through the recipient MCA branch artery on two occasions. Postoperative TCD demonstrated HITS that resolved with uptitration of antiplatelet therapy. Patient B presented with multifocal, embolic-appearing left MCA infarctions, and unilateral angiographic moyamoya syndrome. She was found to have HITS in the left MCA, which eventually resolved with a combination of antiplatelets and anticoagulation.

Conclusion: Hemodynamic compromise may not be the only cause of brain infarction in patients with moyamoya syndrome. Observations from these two patients provide both direct visualization and TCD evidence of embolization as a potential etiology for brain ischemia. Future investigations into the role of antithrombotic agents should be considered.

Keywords: brain ischemia, cerebral revascularization, embolism, moyamoya disease, stroke

Open Access

Edited by: Andreas Charidimou, Harvard Medical School, United States
Reviewed by: Marco Pasi, Massachusetts General Hospital, United States, Jan F. Schelitz, Charité Universitätsmedizin Berlin, Germany
*Correspondence: Julie G. Shulman, julie.shulman@bmc.org

Specialty section: This article was submitted to Stroke, a section of the journal Frontiers in Neurology

Received: 30 June 2017 Accepted: 07 August 2017 Published: 24 August 2017


For the last 60 years, neurosurgeons and neurologists have described patients with progressive occlusion of the carotid termini with compensatory proliferation of tiny vessels at the base of the brain. These proliferating vessels, which appear as an early capillary blush or “puff of smoke” on conventional angiogram, lend the moyamoya syndrome its name (1). The moyamoya syndrome refers only to these angiographic findings and can be seen as a consequence of systemic illnesses or treatments, such as Down’s syndrome, atherosclerosis, or cranial irradiation (2). Moyamoya disease refers specifically to idiopathic cases and has been histologically characterized by intimal proliferation, irregular lamina,
and medial fibrosis in large arteries, with a marked absence of inflammation. The small proliferative vessels are also atypical and markedly variable in diameter (3).

Patients with moyamoya disease typically present in one of two age groups, either as children or as adults in their 40s, with transient ischemic attack (TIA), stroke, or less frequently, hemorrhage. TIAs and strokes are generally felt to result from proximal stenosis causing hemodynamic insufficiency (4). The possibility of active embolism as a mechanism for cerebral ischemia in moyamoya syndrome has only recently been investigated (5). Understanding the exact mechanism of stroke is important in these patients, as treatment options differ. Hypoperfusion is managed with volume expansion, normocapnia, permissive hypertension, and ultimately surgical bypass. Embolization may be managed with antithrombotic and/or antiplatelet agents (6), which may be risky in fragile moyamoya collaterals.

Transcranial Doppler (TCD) is the primary means of detecting cerebral arterial emboli in vivo. Both gaseous and particulate emboli can be identified as high-intensity transient signals (HITS). These signals have a high in vitro specificity for emboli (as opposed to artifact) (7), and have been identified in patients with moyamoya disease (8–10). In spite of this, there has never been visual or pathological proof of embolization in these patients.

**CASE PRESENTATION**

**Patient A**

A 44-year-old right-handed hypertensive woman presented with left arm weakness. Symptoms had been present intermittently for several months, but had been particularly severe for 3 days. On examination, she had left hemiparesis and left-sided hyperreflexia. Magnetic Resonance Imaging (MRI) showed acute right middle cerebral artery (MCA) territory infarcts in addition to multiple bilateral chronic infarcts (Figure 1A). Conventional angiogram demonstrated diffusely narrowed bilateral paraclinoid and supraclinoid ICAs, ACAs, and MCAs with numerous collaterals (Figure 1B).

Thorough workup for secondary causes of moyamoya was unremarkable. No hematologic, vasculitic, autoimmune, connective tissue, or genetic abnormalities were identified. A transthoracic echocardiogram showed no central embolic source and no cardiac dysrhythmias were seen on inpatient telemetry monitoring. Given her age, bilateral proximal vasculopathy, and negative secondary workup, she was diagnosed with moyamoya disease. Because of her progressive neurologic symptoms, plans were made to proceed with bilateral direct superficial temporal artery (STA) to MCA bypasses, in stages, beginning with the symptomatic right side.

The STA parietal and frontal branches were used for anastomoses to two frontal MCA branches. On two occasions, embolic material was observed passing through the recipient artery in a distal-to-proximal direction (Figure 2; Video S1 in Supplementary Material). A portion was removed and sent for histological analysis, where it was interpreted as non-specific proteinaceous material (Figure S1 in Supplementary Material). The surgery was completed without radiographic or clinical evidence of new ischemia. On post-operative day 1, a surveillance TCD showed 8–10 HITS in the right MCA (Figure 3). While the patient was on aspirin, a platelet aggregation assay was suggestive of incomplete efficacy. Aspirin was increased from 81 mg daily to 325 mg daily, and 4 days later a repeat TCD study detected no HITS and the platelet aggregation study demonstrated improved platelet inhibition.

**Patient B**

A 37-year-old woman with a history of complex migraines and Graves disease status post thyroidectomy presented with three weeks of progressive speech difficulty during the third trimester of pregnancy. On examination, she had a mild mixed aphasia.
Brain MRI demonstrated embolic-appearing infarcts of various ages in her left frontal, temporal, and parietal lobes (Figure 4A). Conventional angiography showed occlusion of the left MCA M1 segment with local proliferation of collaterals and a normal right ICA (Figure 4B). Additional imaging was performed to assess the hemodynamic reserve of the left hemisphere and the patency of the left MCA. TCD vasoreactivity study demonstrated a lesser increase in flow velocity in the left compared with the right MCA at higher end-tidal carbon dioxide concentrations. A single-photon emission computed tomography (SPECT) study performed before and after 1,000 mg of intravenous acetazolamide showed baseline perfusion deficits in the left MCA territory with some areas of improved perfusion after the acetazolamide, suggestive of some degree of preserved autoregulation. Additional workup for etiology of stroke included a transesophageal echocardiogram, telemetry monitoring, and cerebrospinal fluid analysis, all of which were unrevealing. She had no history of radiation, atherosclerosis, or hyperlipidemia, and extensive serologic testing was unrevealing. Given her unilateral vasculopathy, negative secondary workup, and history of hyperthyroidism, she was diagnosed with moyamoya syndrome.

Transcranial Doppler demonstrated 26 HITS in the left MCA during a 30-min period (Figure 4C). When these HITS persisted in spite of therapeutic anticoagulation, aspirin 81 mg daily was added. After 3 months of treatment, no HITS were detected during a 40-min monitoring period. At that point, anticoagulation was discontinued, aspirin was continued, and she underwent a left STA–MCA bypass. She has remained well since the procedure on aspirin monotherapy, with a mild residual aphasia.

**DISCUSSION**

We present two patients with symptoms of cerebral ischemia and angiographic findings of moyamoya syndrome and disease with ultrasonic, visual, and pathological evidence of active brain embolization.

Hemodynamic compromise has long been considered the primary cause of ischemic symptoms in patients with moyamoya and is supported by several lines of evidence. Studies using xenon inhalation or single-photon positive emission computed tomography (SPECT) have shown impaired vasoreactivity distal to the stenosis, indicating that distal arterioles are maximally vasodilated and are unable to further augment flow (8, 11) Additionally, brain infarcts have a tendency to occur in areas of perfusion deficit seen on SPECT, in anatomically watershed distributions (12).

There is also clinical evidence to support the phenomenon of hypoperfusion. In children, there is an association between activities which reduce systemic carbon dioxide (such as crying) and subsequent cerebral vasoconstriction with transient ischemic symptoms (13). Furthermore, direct and indirect bypass operations which augment distal perfusion improve outcomes. The 5-year stroke rate of 12% (14) to 27% (9) in patients treated with any kind of bypass surgery is markedly improved compared with 40% (15) to 82% (16) in patients who did not have surgery. Although there is no randomized trial data to prove efficacy, surgery is considered the best available treatment for symptomatic moyamoya syndrome (17).

The possibility of embolism as an additional mechanism has also been recently investigated. In patients with proximal vascular stenoses, the two mechanisms can be difficult to separate and may coexist. One cannot rely solely on a typical watershed pattern of infarcts on imaging, as such infarcts have been associated with small distal vessels filled with cholesterol emboli (5). Both mechanisms may ultimately contribute to the infarcts, in that a state of chronic hypoperfusion from a proximal stenosis may prevent...
the distal washout of emboli. A TCD study may provide action-
able information, as HITS have been associated with emboli in
patients with proximal MCA stenoses, with number of HITS
linearly correlating with number of infarcts on MRI (18).

In moyamoya disease, HITS are detected in about 12% (8) to
20% (9, 10) of patients. Their presence was predicted by recent
clinical ischemic events and carried a corrected odds ratio of 10.6
for stroke in the next year (10). Furthermore, HITS are equally
prevalent in all stages of the disease (10), a finding that might
not be expected if hemodynamic compromise was the only factor
at play.

The two patients discussed here provide convincing evidence
of the presence of emboli in moyamoya patients. The nature and
origin of this embolic material are more difficult to determine.
While the pathological interpretation was non-specific, the
fact that solid material was surgically extracted eliminates the
possibility of the TCD and surgical findings being artifactual.
It is also very unlikely that the embolus was a result of the
procedure itself as it was seen traveling on two occasions in
a distal-to-proximal direction, from an untouched arterial
territory toward the surgical clamp. Possible sources of emboli
include the severely stenosed proximal intracranial vessels,
with disturbed hemodynamics explaining the reversal of flow
in distal arteries. Alternatively, severe hemodynamic compro-
mise from proximal stenosis could cause stasis in distal vessels
with development of embolic material distally. Given that this
finding was observed in both moyamoya disease (patient A)
and moyamoya syndrome (patient B), we have no evidence to
suggest that the risk of emboli formation would differ between
these two groups.

This is the first report documenting direct visualization of brain
embolism in moyamoya. In both cases, HITS were abolished with
the use of antithrombotic therapy, but our data are insufficient to
establish a cause and effect relationship.

CONCLUSION

Hemodynamic compromise may not be the only cause of brain
infarction in patients with moyamoya. Brain embolism occurs
in this population, as seen here intra-operatively and by TCD,
and may cause brain infarction. This provides a rational basis for
antithrombotic therapy, which should be studied further.

ETHICS STATEMENT

Please note that written informed consent was obtained from
both patients for the publication of this report. As this is a case
report, it was exempt from review by our local Institutional
Review Board.

AUTHOR CONTRIBUTIONS

JS, primary physician for patient A, drafted manuscript, man-
aged revisions, selected figures for patient A, and formatted all
figures. SS, primary physician for patient B, made significant
manuscript contributions, selected figures for patient B. HV, sen-
or physician for patient B, supervised manuscript drafting and
made revisions. VB: senior physician for patient A, supervised
manuscript drafting and made revisions. NP, neurosurgeon for
both patients, creator and owner of intra-operative photos and
videos.

ACKNOWLEDGMENTS

The authors thank Ivana Delalle, MD, Ph.D. for assistance with
interpretation of pathology material.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at
full#supplementary-material.

FIGURE S1 | Supplementary Material figure, in TIFF format: Fragment of
intraluminal cellular debris obtained during Patient A’s right STA-MCA bypass,
noted to be non-specific proteinaceous material.

VIDEO S1 | Supplementary Material video, in mp4 format: Intra-operative video
captured during right STA-MCA bypass of patient A. On two occasions, direct
visualization of emboli formation and distal-to-proximal passage is seen.
REFERENCES

1. Takeuchi K, Shimizu K. Hypoplasia of the bilateral internal carotid arteries. 


5. Momjian-Mayor I, Baron JC. The pathophysiology of watershed infarction 
   in internal carotid artery disease: review of cerebral perfusion studies. Stroke 
   (2005) 36:557–67. doi:10.1161/01.STR.0000155727.82242.e1

6. Kraemer M, Berlit P, Disiner F, Khan N. What is the expert’s opinion on 
   antiplatelet therapy in moyamoya disease? Results of a worldwide survey. 

7. Markus H. Transcranial Doppler detection of circulating cerebral emboli: a 

8. Horn P, Lanczik O, Vajkoczy P, Daffertshofer M, Bueltmann E, Werner A, 
   et al. Hemodynamic reserve and high-intensity transient signals in moyamoya 

   Stroke (2008) 39:1347–51. doi:10.1161/01.STR.29.7.1347

10. Chen J, Duan L, Xu WH, Han YQ, Cui LY, Gao S. Microembolic signals predict 
    cerebral ischaemic events in patients with moyamoya disease. 

11. Tatemichi TK, Prohovnik I, Mohr JP, Correll JW, Jarvis L. Reduced hypercapnic 

12. Iwama T, Hashimoto N, Yonekawa Y. The relevance of hemodynamic factors 
    to perioperative ischemic complications in childhood moyamoya disease. 

13. Tagawa T, Naritomi H, Mimaki T, Yabuuchi H, Sawada T. Regional cerebral 
    blood flow, clinical manifestations, and age in children with moyamoya 

    doi:10.1161/STROKEAHA.111.621300


17. Fung LW, Thompson D, Ganesan V. Revascularization surgery for paediatric 
    doi:10.1007/s00381-004-1118-9

    Mechanisms of acute cerebral infarctions in patients with middle cerebral 
    artery stenosis: a diffusion-weighted imaging and microemboli monitoring 

Conflict of Interest Statement: HV is a paid consultant for Sage Therapeutics. VB 
receives funding from Boston Scientific (CEC member) and from Bayer (BMC-PI 
of a multi-center research study). None of these relationships influenced the work 
presented in this manuscript.

The handling editor declared a shared affiliation, though no other collaboration, 
with the authors.

Copyright © 2017 Shulman, Snider, Vaitkevicius, Babikian and Patel. This is an 
open-access article distributed under the terms of the Creative Commons Attribution 
License (CC BY). The use, distribution or reproduction in other forums is permitted, 
provided the original author(s) or licensor are credited and that the original publica-
tion in this journal is cited, in accordance with accepted academic practice. No use, 
distribution or reproduction is permitted which does not comply with these terms.