Characterization of Arteriovenous Malformation Feeding Vessels by Carbon Dioxide Reactivity

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PURPOSE: To characterize cerebral hemodynamics in patients immediately before microsurgical resection of moderate to large arteriovenous malformations during isoflurane anesthesia. METHODS: In angiographically defined arteriovenous malformation feeding and nonfeeding arteries, transcranial Doppler studies were performed in 25 surgeries on 22 patients. The mean blood flow velocity and pulsatility index were recorded in the middle, anterior, and posterior cerebral arteries. Transcranial Doppler velocities were measured at end-tidal carbon dioxide tensions (PetCO₂) of about 25 and 35 mm Hg. Carbon dioxide reactivity was calculated as percentage mean blood flow velocity change per mm Hg PetCO₂ change. RESULTS: Patient demographic and clinical data for the arteriovenous malformation group followed the expected strata of a large arteriovenous malformation population. All patients were neurologically stable before surgery. A total of 43 feeding arteries and 55 nonfeeding arteries were studied. Compared with nonfeeders, feeders exhibited higher mean blood flow velocity (68 ± 5 vs 31 ± 3 cm/sec, \( P < 0.0001 \)) and lower pulsatility index (0.64 ± 0.03 vs 0.88 ± 0.04, \( P < 0.001 \)); anterior and middle cerebral artery velocities at normo- and hypocapnia were significantly higher than posterior cerebral arteries for both feeders and nonfeeders (\( P < 0.001 \)). Carbon dioxide reactivity was 0.2 ± 0.2 %/mm Hg in feeders and 2.1 ± 0.2 %/mm Hg in nonfeeders, with no significant difference between arteries. In four of eight patients with lesions fed by the anterior circulation (middle cerebral artery with or without anterior cerebral artery feeders), posterior cerebral artery nonfeeders exhibited low reactivity. In 2 of 5 patients with ipsilateral posterior cerebral artery feeders, contralateral posterior cerebral artery nonfeeders exhibited impaired reactivity. CONCLUSIONS: Quantitative transcranial Doppler studies are technically feasible in the operating room or interventional suite during anesthesia. Hemodynamic assessment using physiologic challenges of arteriovenous malformation feeders as well as angiographically uninvolved vessels may be useful as criteria in the assessment of malformations and arteriovenous malformation patients may exhibit abnormal vasoreactivity in distant uninvolved perfusion territories, suggesting a deranged neural control mechanism.

Index terms: Arteriovenous malformations, cerebral; Blood vessels, flow dynamics; Ultrasound, Doppler; Cerebral blood, flow


Cerebral arteriovenous malformations (AVMs) consist of three main vascular components: the nidus, the feeding arteries supplying the fistula, and the venous drainage. Morphologically, the nidus consists of abnormal arteries and veins that lack muscular resistance vessels and a true capillary bed (1). Feeding arteries may exhibit a lower-than-systemic intravascular pressure with a corresponding hypertension in the draining venous structures (2-4). The combination of arterial hypotension and venous hypertension, if transmitted to contiguous normal circulatory beds, can result in a decrease in cerebral perfusion pressure (2). The abrupt increase in cerebral perfusion pressure with treatment (either embolization or surgery) may be involved in the pathogenesis of certain complications such as brain swelling or
hemorrhage (2, 5). The ability to identify severe physiologic disturbances in cerebral hemodynamics that might lead to postoperative complications would be beneficial.

Standard angiography can identify feeding arteries and their parent vessels but cannot provide quantitative information about shunt flow or the physiologic effects of the fistula on normal circulatory beds. The hemodynamic effects of shunt flow through AVMs on adjacent and distant normal tissue beds have been studied with xenon-133 clearance (6), single-photon emission computed tomography (7), and positron emission tomography (8). Transcranial Doppler studies have also contributed insights into the hemodynamic consequences of the fistula; feeding arteries are characterized by high velocity and low pulsatility, which revert to normal values after obliteration of the AVM shunt (3, 9–12).

Provocative physiologic challenges have been used to assess autoregulatory function in the brain. Carbon dioxide (CO\(_2\)) or its functional analog, acetazolamide, have been applied widely for several cerebral blood flow methodologies in patients with cerebrovascular disease (7, 13, 14). Carbon dioxide (CO\(_2\)) reactivity in normal arteries measured with transcranial Doppler is similar to values reported for hemispheric CBF (tissue perfusion) techniques (15, 16). Lindegaard et al (11) studied middle cerebral artery (MCA) CO\(_2\) reactivity in AVM patients and reported a diminished reactivity to changes in carbon dioxide tension.

The purpose of this investigation was to determine the range of CO\(_2\) reactivity in angiographically defined feeding and nonfeeding arteries in the MCA, anterior cerebral artery (ACA), and posterior cerebral artery (PCA) in the intraoperative setting and attempt to classify AVM feeding and nonfeeding vessels from a physiologic perspective.

**Methods**

All patients underwent elective microsurgical resection of moderate to large AVMs. Preoperative standard four-vessel angiography was used to identify feeding and nonfeeding arteries (feeders and nonfeeders). AVM volume was estimated as described by Pasqualin et al (17).

Intraoperative transcranial Doppler studies were performed during general anesthesia with 0.75% isoflurane and 60% nitrous oxide in oxygen, after induction but before incision. A 2-MHz pulsed Doppler probe (Transpect, Medasonics, Freemont, CA) was placed over the temporal bone to insonate MCAs, ACAs, and PCAs as previously described (10). All vessels were insonated in the proximal portions in approximately the same location and the depth of maximum signal was recorded. Mean blood-flow velocity (MV) and pulsatility index were registered for each artery. The pulsatility index was calculated as: systolic velocity — diastolic velocity)/MV.

After induction of anesthesia, mechanical ventilation was managed by titration of end-tidal CO\(_2\) tension (PetCO\(_2\)) to moderate hypocapnia (about 25 mm Hg). PetCO\(_2\) was assessed using capnography (RGM 5250, Ohmeda, Madison, WI) as an index of PaCO\(_2\) (18). To assess transcranial Doppler velocities at normocapnia (about 35 mm Hg), CO\(_2\) was added to the fresh gas mixture. The MV and pulsatility index were then measured bilaterally in the ACA, MCA, and PCA (for technical reasons it was not possible to obtain transcranial Doppler velocities in all arteries in all patients).

Patients were assigned randomly to begin the sequence of examinations with either hypocapnia or normocapnia.

CO\(_2\) reactivity (%/mm Hg) was calculated as:

\[
\frac{100[(MV_{NC} - MV_{HC})/MV_{NC}]}{\Delta PetCO_2}
\]

The slope of the CO\(_2\) response (cm/sec/mm Hg) was calculated as:

\[
\frac{MV_{HC} - MV_{NC}}{\Delta PetCO_2}
\]

Although the protocol called for maintenance of stable blood pressure between normocapnia and hypocapnia, there were random variations in mean arterial pressure (MAP) between measurements and this change (\(\Delta MAP\)) was introduced as another covariate for statistical analysis. Data were analyzed by analysis of variance with the MV, pulsatility index, and CO\(_2\) reactivity as dependent variables and feeder classification (feeder vs nonfeeder) and artery (ACA, MCA, PCA) as grouping factors. \(\chi^2\) analysis was used for nonparametric data. Linear regression and analysis of covariance was used to determine whether the \(\Delta MAP\) between measurements influenced the CO\(_2\) reactivity between feeders and nonfeeders. A value of \(P < 0.05\) was taken as the threshold of significance. Data are expressed as mean \pm standard error.

**Results**

There were 25 surgeries on 22 patients; three patients underwent staged resection. Their mean ages were 35 ± 2 years (range, 18 to 54) and included 14 men and eight women with 11 left- and 11 right-sided lesions. Five of the 22 patients underwent preoperative embolization. The mean AVM size (cm) was 4.7 ± 0.4 by 3.9 ± 0.3 by 3.5 ± 0.4 and volume was 38 ± 10 cm\(^3\).

The feeding arterial complex was MCA/ACA in eight, MCA/PCA in two, MCA/PCA/ACA in five, MCA-only in one, and PCA-only in six. The predominant location of the AVM was frontal...
(eight), parietal (three), temporal (three), occipital (seven) and dural-cerebellar (one). Patients presented with headache (six), hemorrhage (nine) and seizures (five); two were asymptomatic. All patients were neurologically stable at the time of surgery and no patient had clinical or radiologic evidence of vasospasm or increased intracranial pressure.

AVM volume was smaller in patients who presented with hemorrhage than those with other presentations (15 ± 2 vs 60 ± 13 cm³, P < 0.05). Otherwise, the above demographic data and anatomic considerations were not related to MV, pulsatility index, or CO₂ reactivity by subsequent statistical analysis. The depth of insonation was 65 ± 1 cm for ACA and PCA and 50 ± 1 cm for MCA. A total of 43 feeding arteries and 55 non-feeding arteries were studied.

All MV and pulsatility index mean values are presented in Figure 1A and 1B. Hypocapnic (22 ± 0.2 mm Hg) and normocapnic (36 ± 0.3 mm Hg) PetCO₂ levels were significantly different, with no effect of artery or feeder versus nonfeeder classification. Compared with nonfeeders, feeders exhibited higher mean MV (68 ± 5 vs 31 ± 3 cm/sec, P < 0.001) and lower pulsatility index (0.64 ± 0.03 vs 0.88 ± 0.04, P = 0.001); ACA and MCA velocities at normo- and hypocapnia were significantly higher than PCA for both feeders and nonfeeders (P < 0.001). There was no significant difference in pulsatility index between ACA, MCA, and PCA in both feeders and nonfeeders.

CO₂ reactivity values were significantly different between feeders and nonfeeders and data are shown in Figure 2. CO₂ reactivity was 0.2 ± 0.2%/mm Hg in feeders and 2.1 ± 0.2%/mm Hg in nonfeeders. The slope of the CO₂ response was 0.16 ± 0.17 cm/sec/mm Hg in feeders and 0.99 ± 0.11 cm/sec/mm Hg in nonfeeders. Although feeders had a significantly lower CO₂ reactivity than nonfeeders, there were no significant differences between ACA, MCA, and PCA in either feeders or nonfeeders. For feeders, however, MCA CO₂ reactivity tended to be the lowest.
By analysis of covariance of feeders vs nonfeeders, there was a significant effect of ΔMAP on CO₂ reactivity (P = 0.0001). By linear regression, there was a weak but significant correlation between the ΔMAP and the slope of the CO₂ response, as shown in Figure 3. When feeders and nonfeeders were analyzed separately, the effect of ΔMAP on the CO₂ response was significant for feeders only.

Where applicable, a comparison was made between matched sets of ipsilateral feeders and their corresponding contralateral nonfeeders. Four sets of ACAs, 15 MCAs, and five PCAs were examined. Flow velocities, pulsatility index, and CO₂ reactivity were compared by analysis of variance for ACA, MCA, and PCA. Similar to the analysis of all arteries studied, ipsilateral feeders were significantly less reactive than contralateral nonfeeders, with no significant difference between arteries. Although PCA and MCA pulsatility index values were significantly different between feeders and nonfeeders during hypocapnia, as seen in Figure 4, contralateral nonfeeding ACAs had pulsatility index values that were low and not significantly different from ipsilateral feeders. This was surprising because there was a marked difference in CO₂ reactivity between ACA ipsilateral feeders and contralateral nonfeeders.

In two patients with large frontal AVMs (in whom blood pressure did not change between hypo- and normocapnic measurements), feeding arteries demonstrated a strongly negative reactivity to CO₂ (−3% to −4%/mm Hg); that is, lowering PetCO₂ increased flow through the fistula. There were also several nonfeeders with apparently impaired (<1% change/mm Hg) CO₂ reactivity. In four of eight patients with lesions fed by the anterior circulation (MCA with or without ACA feeders), PCA nonfeeders exhibited low reactivity. In two of five patients with ipsilateral PCA feeders, contralateral nonfeeders exhibited low (about 1% change/mm Hg) reactivity. This is illustrated in Figure 5. The PCA nonfeeders with low reactivity were anatomically distant from the AVM in all cases.

Discussion

Cerebral AVMs are characterized by feeding vessels with significantly higher MV and lower pulsatility index than angiographically defined nonfeeding vessels and the results presented here agree with previous observations made in the
observed fraction of the intracranial-bound cardiac output among the ACA, PCA, and MCA feeders, our patients, a marked redistribution of flow of normal resistive beds and a net increase in the is not clear what factors will determine when the that passes through the fistula was observed during hypocapnia, similar to previous reports (19). The influence of systemic arterial pressure on CO$_2$ reactivity suggests two important considerations for future investigations: 1) Blood pressure changes between transcranial Doppler measurements of CO$_2$ reactivity should be strictly controlled for accurate quantitative interpretation and 2) this modulating influence may be employed as another means to assess hemodynamic function.

An area that deserves further study is the hemodynamic effect of the AVM across Willisian collateral pathways, such as the anterior communicating artery. Because of the physical proximity of the two proximal ACAs, one might expect any decreased pressure in a feeding ACA to be transmitted easily to its contralateral nonfeeding counterpart. We observed a significant lowering of pulsatility index in nonfeeding contralateral ACA segments, which might suggest such a phenomenon. However, the CO$_2$ reactivity did not seem to be affected. Intuitively, one would expect vessels with a low pulsatility index to have similarly decreased CO$_2$ reactivity. This observation awaits corroboration with a larger sample and, ideally, correlation with intravascular pressure measurements to determine which parameter (pulsatility index or CO$_2$ reactivity) is more closely related to intravascular pressure reductions.

The four cases of diminished CO$_2$ reactivity in the PCA with lesions fed by the ACA or MCA and the two patients with ipsilateral PCA feeders and decreased reactivity in a contralateral PCA suggest a distant effect of the AVM fistula. There has been suggestion of distant derangements in cerebral perfusion patterns using other CBF methods (6, 7, 8, 25) and complications are not limited to the ipsilateral hemisphere (26). What awake state (9–11, 19). The PCA tended to have the lowest MV, but the pulsatility index was similar to the MCA and ACA. This was true for both feeders and nonfeeders, reflecting the relative differences in flow velocities seen in healthy subjects (20). The MV values for nonfeeders were about 20% lower than reports in healthy awake subjects (20), similar to other observations in the isoflurane-anesthetized state (21, 22). Our CO$_2$ reactivity values agree with previous studies for MCA feeding vessels and their contralateral nonfeeding MCAs (11). This report extends these previous observations to the ACA and PCA, where there were no differences in CO$_2$ reactivity among the ACA, PCA, and MCA feeders, although there was a tendency for MCA feeders to be the most severely affected. In at least two of our patients, a marked redistribution of flow through the fistula was observed during hypocapnia, similar to previous reports (19). This redistribution is probably explained by vasoconstriction of normal resistive beds and a net increase in the fraction of the intracranial-bound cardiac output that passes through the fistula (12). However, it is not clear what factors will determine when the observed CO$_2$ reactivity will remain near zero and when it will become negative.

Our data support the previous observation (11) that systemic arterial pressure can modulate vascular reactivity to CO$_2$ in feeders. Flow through the fistula is directly pressure-passive because the AVM nidus lacks muscular resistance arterioles. Assuming that autoregulatory vasomotion does not affect the proximal conductance vessels being insonated with transcranial Doppler, one would expect that a normal artery flow velocity would remain constant in the normal autoregulatory range with a change in systemic pressure. Hypothetically, however, a nonfeeding artery may be subjected to lower than normal pressure transmitted across Willisian pathways. Such relative hypotension, even if it did not result in frank ischemia, could result in a state of maximal or near-maximal vasodilation (impairing CO$_2$ reactivity) and would render that territory pressure-passive as well. This remains to be demonstrated in future studies combining other imaging modalities or, ideally, direct intracerebral pressure measurements (23, 24). The influence of systemic arterial pressure on CO$_2$ reactivity suggests two important considerations for future investigations: 1) Blood pressure changes between transcranial Doppler measurements of CO$_2$ reactivity should be strictly controlled for accurate quantitative interpretation and 2) this modulating influence may be employed as another means to assess hemodynamic function.

![Fig. 5. CO$_2$ reactivity in ipsilateral (closed symbols) and contralateral (open symbols) PCA in five patients with lesions fed by the ipsilateral PCA, with no angiographic involvement of the contralateral PCA. Two patients exhibited normal contralateral and decreased ipsilateral CO$_2$ reactivity (A and B), one patient had intermediate reactivity in both vessels (C), and two patients had decreases in both PCAs.](image-url)
remains unclear at present is whether these distant effects are caused by a hemodynamic mechanism (lowered cerebral arterial pressures or a phenomenon functionally analogous to diaschisis seen in the setting of cerebral ischemia) (27). Evidence is accumulating that AVM patients exhibit unusual cerebral blood flow responses to pharmacologic challenges (28, 29). It is tempting to speculate that there may be a derangement in neural modulation or regulation of cerebral blood flow in certain AVM patients; for example, both PCAs may be severely affected by some control defect, even though the contralateral nonfeeding PCA is not involved in feeding the fistula. In fact, this lack of appropriate neurovascular control may very well have importance in the pathogenesis of cerebral AVMs; that is, a localized abnormal vasomotoric response to appropriate metabolic or hemodynamic stimuli may lead to gradual enlargement of what might initially be a trivial injury or abnormal area in such a vulnerable zone.

In summary, quantitative transcranial Doppler studies are technically feasible in the operating room or interventional suite during anesthesia; the values reported here agree with previous observations in the awake state. The behavior of the proximal Willisian vessels appears to be similar, but MCA and ACA feeding artery velocities tend to be higher and CO2 reactivity lower than PCA feeding vessels. AVM patients may exhibit abnormal vasoreactivity in distant uninvolved perfusion territories, suggestive of a deranged neural control mechanism. Hemodynamic assessment using blood pressure or CO2 challenges of AVM feeders as well as angiographically uninvolved vessels may be used as physiologic criteria in the assessment of AVMs. It may be possible to characterize a range of abnormal responses that suggest high risk for development of certain complications after treatment (brain swelling or hemorrhage) and are of assistance in making posttreatment management decisions, such as which patients may benefit from aggressive blood pressure control after embolization or surgery to avoid circulatory breakthrough complications.

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References


