ABSTRACT: Fifty patients with ruptured intracranial aneurysms and 8 patients with elective clipping of unruptured aneurysms had daily transcranial Doppler (TCD) measurements performed. The highest mean middle cerebral artery velocity (MCA-Vel) was considered to be the best single parameter for judging a patient’s susceptibility to clinically significant vasospasm (VSP). Surgery for the clipping of unruptured aneurysms by itself does not lead to an increase in MCA-Vel. There is a progressive increase in MCA-Vel after subarachnoid hemorrhage (SAH) from aneurysms which peaks between 7 and 10 days. The MCA-Vel is higher on the side of the ruptured aneurysm and the degree of rise is greater if blood is seen on the initial CT scan. It is highly unlikely that a patient whose MCA-Vel remains under 100 cm/sec has a degree of angiographic VSP which causes clinical symptomatology. Patients whose MCA-Vel is > 200 cm/sec are at great risk of developing clinical symptomatology of VSP and are very likely to have significant angiographic VSP. There is a transitional zone in between these two levels.

METHODS

Patient Population

Fifty patients with ruptured intracranial aneurysms admitted to hospital acutely following their SAH, and 8 patients having elective clipping of unruptured aneurysms, had daily TCD examinations performed. Examinations were carried out for 21 days or until discharge, whichever came first. Forty-five of the ruptured aneurysm cases were operated in the first day or two following their admission, 3 were operated after a few days with 2 cases admitted in very poor condition who were operated some months later. The percentage of cases in each grade on
admission were: 0 — 14%, I and II — 47%, III and IV — 34%, and V — 5%. None of the 8 elective cases developed a neurological deficit or died. Of the 27 cases admitted in grade I or II following SAH, 63% had a good outcome, 33% had a deficit and 4% died. The death was in a patient with early massive rebleeding. Of the patients admitted grade III or IV 10% had a good outcome, 70% had a deficit and 20% died. Of patients grade V on admission, 0% had a good outcome, 66% had a deficit and 33% died. The postoperative mortality rate at 30 days was 2%, there was 1 additional late postoperative death at 4 months to make a total postoperative mortality rate of 4%. All cases received prophylactic albumen to maintain a high normal blood volume and about half the SAH cases received a calcium channel blocker in a study which is still blinded. Symptomatic VSP was treated with hypertension/hypervolemia. Clinical VSP was diagnosed from reviewing the charts retrospectively for delayed clinical deterioration (after day 4) not evidently due to some other cause.

**Transcranial Doppler Examination**

A 2 MHz pulsed Doppler unit (TC 2-64, Teca Corp., Pleasantville, New York) was used to insonate the middle, anterior and posterior cerebral and the internal carotid arteries bilaterally through the temporal areas. The usual recording depths from the temporal skin surface were 45 and 55 mm for distal and proximal middle cerebral, 65 mm for anterior cerebral, 75-80 mm for posterior cerebral and 60 mm for the internal carotid arteries. To locate arteries in addition to the depth change the probe angle was changed slightly, forward for anterior or cerebral, backward for posterior cerebral and downward for internal carotid arteries. Slight variation in depth was necessary in some patients. The TCD values were recorded independently of the radiologists who interpreted the angiograms.

In appropriate cases velocity recordings were also made from the basilar artery at depths of 75 and 85 mm through the foramen magnum area. In all patients velocity recordings from the extracranial carotid arteries were made at an estimated angle of 60° with the 2 MHz probe at a usual depth of 50 mm.

Mean MCA-Vel only are presented (the mean of the selected waveform not the mean of multiple separate observations).

**Statistical Methods**

Data was entered into a computer and the means and standard error of the means were calculated. Data comparison used two-tailed unpaired (pooled variance) t-tests. The total number of patients in each group is indicated in the diagrams, however, the ‘n’ on individual days was not constant due to the variation in patient admission to and transfer from the neurosurgical intensive care unit.

**Clinical and Radiological Evaluation**

The diagnosis of “clinical VSP” was made on the basis of clinical deterioration more than 4 days from the SAH which could not be explained on some other basis. Angiographic VSP was diagnosed by the radiologist on the basis of the usual radiographic appearance of diffusely narrowed vessels on angiograms which were repeated after the fourth day following SAH. This diagnosis was made without reference to this subsequent study. There was no routine time for performance of repeat angiograms as some were conducted primarily to check on the efficacy of the operative procedure and only a few were performed specifically to evaluate the degree of VSP.

**Results**

While on occasions the velocities from the other arteries mirrored the changes in the middle cerebral no additional information was gained from these measurements. Ratio of middle cerebral mean velocity to internal carotid arterial mean velocity was calculated. The variation in this parameter mirrored that of the MCA-Vel but was less stable. The wider fluctuations in the ratio compared to the absolute velocity appeared to be due to variation in apparent internal carotid artery velocity. This was probably due to a difficulty in reproducibly insonnating the artery in the neck at an angle of 60°.

Patients having elective clipping of unruptured aneurysms showed no significant elevation in mean MCA-Vel. These remained in the area of 30-60 cm/sec despite the performance of craniotomy and aneurysmal clipping. In contrast, patients after SAH developed a progressive rise in MCA-Vel over the first 7 or 8 days following SAH. By day 8 the mean of all patients had exceeded 100 cm/sec (Figure 1).

Patients having blood clot in the subarachnoid space observed on the initial CT had higher MCA-Vel than those whose initial CT did not show blood. However there were no differences between those whose initial CT subarachnoid clot was considered to be “thick” rather than “thin” (Figure 2).

Lateralized aneurysms (excluding anterior communicating or basilar apex locations) showed higher MCA-Vel on the side ipsilateral to the ruptured aneurysm. This difference became apparent after the 4th day. It achieved statistical significance on the 8th and 9th days (Figure 3).

Twenty-seven of the ruptured aneurysm cases had angiograms performed in the period in which VSP occurs. Twenty-one showed severe VSP, 5 showed mild VSP and 1 showed none. The MCA-Vel for the severe cases was higher than the mild or none cases after day 5 post-SAH. Because of

![Figure 1 — Mean MCA-Vel plotted against days post-SAH or post-op in the case of unruptured aneurysms. The average numbers of daily observations ≤ d5 were 25, d6-10 were 41, d11-15 were 39, d16-20 were 25.](https://www.cambridge.org/core)
large individual variation this only achieved statistical significance on day 10. When the highest MCA-Vel for the entire hospital course was plotted for the group with "severe angiographic VSP" versus "mild or no VSP", the mean and standard error of the mean for the "severe" group was 184 ± 16 cm/sec and for the "mild or no" group it was 124 ± 19 cm/sec (not significant). The lowest maximum MCA-Vel in the severe angiographic VSP group was 80 cm/sec and the highest MCA-Vel for the mild angiographic VSP group was 205 cm/sec (Figure 4). The case with an apparently severe angiographic VSP but low Doppler velocity was of special interest. While the radiological report was "severe, diffuse angiographic spasm" the angiogram on day 13 post-SAH showed a reduction in MCA diameter of only 33% while the intracranial internal carotid was more reduced at 41%.

The actual size of the MCA (which was probably normally on the large size) still looked quite capable of maintaining normal flow (Figure 5), even when florid proximal spasm was evident.

Cases with the diagnosis of "clinical VSP" had higher MCA-Vel than those who did not but the shape of the curve was similar. After 19 days due to very small numbers the relationship was reversed. The shape of rise in the MCA-Vel curve was similar in both groups. The maximum MCA-Vel was achieved on day 9 in the "clinical VSP" group and on day 10 in the "no clinical VSP" group. The mean of the highest MCA-Vel during the hospital course for each patient with a diagnosis of clinical VSP was plotted. The mean and standard error of the group with "clinical VSP" was 176 ± 12 cm/sec. Corresponding figures for the "no clinical VSP" group were 102 ± 9 cm/sec (p<.001). The lowest MCA-Vel in the "clinical VSP" group was 80 cm/sec (Figure 6).

The last Glasgow Coma Score (GCS) recorded for the patient immediately prior to the TCD measurements were recorded. The individual curves varied considerably between patients. There was a tendency for patients with a "flat pattern" for the mean MCA-Vel curves to have high concurrent GCSs. There was also a trend for patients with very high velocity curves to have low GCSs. Some other patients showed a "peak pattern" during days 4-12 and there was a slight tendency for the GCS to be lower during the peak than subsequently. There were, however, numerous exceptions to these generalizations.

DISCUSSION

The results confirm that TCD is an extremely useful technique for monitoring the course of patients with ruptured intracranial aneurysms. Our opinion is that mean MCA-Vel is
Figure 5 — A — anteroposterior view of the right carotid circulation, day 1 post-SAH. B — same view of the right carotid circulation, day 13 post-SAH. The contemporary radiological report was “severe, diffuse vasospasm” but the MCA-Vel remained <100 cm/sec throughout. In retrospect the degree of VSP of the MCA did not appear too threatening. This was the only case of angiographic VSP (independently diagnosed by a neuroradiologist) with a low MCA-Vel.

Figure 6 — The highest daily mean MCA-Vel for each case with diagnosis of clinical VSP or no clinical VSP. The difference was highly significant. Many of these symptomatic episodes were transient.

<table>
<thead>
<tr>
<th>Mean MCA VELOCITY cm/sec</th>
<th>Individual Values</th>
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The most useful single parameter in this type of analysis for patients with ruptured supratentorial aneurysms.

Aaslid et al² studied 18 cases of SAH from aneurysm. The mean MCA-Vel for studies done in the first 4 days following SAH was 84 ± 18 cm/sec in 8 patients. The corresponding diameters of the MCA in concurrently performed angiographic studies were 2.7 ± 0.3 mm. Ten patients had studies performed between days 5 and 12 and the MCA-Vel were 109 ± 45 and the concurrent MCA diameters 2.1 ± 0.6 mm. Angiographically diagnosed VSP in the MCA was associated with MCA-Vel >120-130 cm/sec. Seiler and Aaslid³ presented data on 39 patients with SAH from ruptured aneurysm. In the 20 patients with normal outcomes, 75% had MCA-Vel <140 cm/sec; 10% 140-200 cm/sec and 15% >200 cm/sec. For 11 patients with reversible delayed ischemic deficits (DID) the corresponding values were 27% <140 cm/sec; 36% 140-200 cm/sec and 36% >200 cm/sec. None of the 8 patients with infarction associated with the delayed ischemic deficit had MCA-Vel <140 cm/sec; 13% had velocities 140-200 cm/sec and fully 88% had velocities exceeding 200 cm/sec. In the patients with symptomatic VSP who were operated on after day 10, the maximum velocities were reached between days 7-12. There were examples of asymptomatic patients with velocities >200 cm/sec. Velocities in the range of 120-140 cm/sec were never associated with infarction.

Twelve cases of SAH from ruptured aneurysms were studied by Compton et al⁴ using TCD and Xe133 cerebral blood flow.
In their material MCA-Vel >100 cm/sec were associated with angiographic VSP in 80% of cases. For patients with velocities <100 cm/sec, only 10% had angiographic VSP. The quotient of the cerebral blood flow (Xe133) over the MCA-Vel (cm/sec) correlated better with the clinical grade of the patients than either the cerebral blood flow alone or the MCA-Vel alone.

Fifty cases of ruptured aneurysms were studied by Harders and Gilsbach. They established that the MCA-Vel increased to approximately day 7. The maximum velocities were present between days 11 and 20. Perhaps the apparent continued high velocities to a time when patients usually appear to have recovered from VSP, which was also evident in our data, is due to the selective sampling of the most sick patients while TCD is not as likely to be done on those who have recovered and gone home. There was an inverse relationship between velocity and angiographic diameter of the MCA and intracranial internal carotid but not the proximal anterior cerebral. MCA-Vel varied with the source of the hemorrhage (site of the ruptured aneurysm), side of the operation and use of intravenous nimodipine (velocity was lowered). Fourteen percent of their patients developed reversible DID but only 2% died from VSP. DID occurred in the interval between the 6th and 12th days at which time the MCA-Vel were >137 cm/sec. All patients with DID had MCA-Vel >117 cm/sec. This value was exceeded on day 3 or 4. In their opinion an increase of >20 cm/sec between days 3 and 7 was indicative of impending DID.

Sekhar et al. had 21 cases. Eight of these developed a DID and had reduction in cerebral blood flow demonstrated with the stable Xe-CT method. MCA-Vel were significantly elevated in their cases between days 4 and 12. The VSP group (8 patients) had mean MCA-Vel of 199 ± 27 cm/sec while the 13 cases without VSP had significantly lower velocities (121 ± 5 cm/sec). When the cerebral blood flow was <20 ml/100 gm/min the basal vessels all showed extreme levels of increased velocities by TCD examination.

Seiler et al. analyzed data from 118 patients with ruptured aneurysms having TCD examinations during periods of transient, rapid blood flow variations in 7 patients having extracranial carotid surgery. The extracranial internal carotid blood flow was measured by flow meters at the time of external carotid occlusion and Doppler recording. The mean internal carotid artery flow varied from 167-399 ml/min and the mean MCA-Vel varied from 32-78 cm/sec. The relation between flow volume and blood velocity were nearly linear under the experimental conditions. Seventy-one patients suffering from intracranial hypertension with subsequent brain death were studied using TCD by Hassler et al. Twenty-nine of these patients were assessed by arterial and intracranial pressure measurements. 33 also had four vessel angiograms. With increasing intracranial pressure, the Doppler wave forms exhibited characteristic changes with a progressive lowering of the diastolic part of the velocity curve which ultimately became 0 and then reversed as the intracranial pressure progressively climbed. Following a point of diastolic reversal in the Doppler wave form, all flow except for a small systolic peak ceased and ultimately these also disappeared as intracranial pressure exceeded arterial pressure. DeWitt and Wechsler suggested several pitfalls in the diagnosis of VSP by TCD. They considered that narrowing of branches distal to the proximal segments of the anterior cerebral artery or posterior cerebral artery will usually be beyond the focal range of the Doppler probe and difficult to detect. They observed that changes in the shape of the trace and proximal arteries occurred only if the distal VSP is extensive. Severe VSP of the internal carotid artery may be of sufficient degree to diminish distal flow and thereby decrease velocity in the basal cerebral arteries. This could result in falsely low velocities when VSP coexists in the proximal intracranial or cranial and extracranial carotid basal vessels as well as in the subarachnoid space. In the author’s experience sufficiently severe narrowing of distal branches (such as the M2 or M3 portions of the MCA) in the absence of proximal VSP is exceedingly rare so this is unlikely to be a common problem.

Our own data is consistent with what has previously been noted in the literature. The wide variation in values between different patients probably reflects a variety of anatomical, hemodynamic and rheological influences in addition to a diffuse narrowing of the major conducting vessels when VSP is present. The gradual rise in velocity in the first few days probably reflects the time course of the development of VSP as more spasmogens are released from the lysis of trapped erythrocytes. Probably after 7-10 days the level of vasoconstrictors is stable or falling.

The initial world experience suggests that angiographic VSP which is of a severe degree and involves the middle cerebral arteries is almost certain to be associated with increased MCA-Vel. From a practical point of view if the MCA-Vel is <100 cm/sec then the chance of clinically significant angiographic VSP being present is minimal. At rates above 120 cm/sec very close observation is required. If the MCA-Vel is above 200 cm/sec, severe VSP is almost certainly present and it is very likely to be clinically significant. At that point, if not before, vigorous steps should be taken to prevent ischemia subsequent to the VSP. To some extent TCD may replace angiography.
phy as the diagnostic technique of choice for postoperative VSP in centers having experienced and readily available ultrasonographers.

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