Pure Motor Stroke Due to Vertebral Artery Dissection

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ABSTRACT: A 39-year-old man presented with a pure motor stroke 9 days after cervical chiropractic manipulation. Computerised tomographic scanning showed a pontine infarct. Cerebral angiography showed changes consistent with the diagnosis of bilateral vertebral artery dissection. It is postulated that the infarct resulted from artery-to-artery embolism.

RESUME: Ictus moteur pur du à la dissection des artères vertébrales Un homme âgé de 39 ans s’est présenté avec un ictus moteur pur survenu 9 jours après des manipulations cerviales chiropractiques. Le CT scan a montré un infarctus au niveau du pont. Une angiographie cérébrale a montré des changements compatibles avec un diagnostic de dissection bilatérale des artères vertébrales. Nous postulons que l’infarctus est le résultat d’une embolie d’artère-à-artère.

Fisher has described more than 20 clinical syndromes of lacunar infarction, i.e. small deep cerebral infarcts due to hypertension-associated arteriolar disease. However, it has become increasingly apparent that clinical findings do not reliably indicate the site of the underlying brain lesion or its pathogenesis. Pure motor stroke (hemiparesis without associated sensory deficit, homonymous hemianopia, aphasia, or agnosia) — one of the commonest lacunar syndromes — has been described in association with: lacunar infarcts in the internal capsule or basis pontis; infarction of the cerebral cortex, centrum semiovale, medullary pyramid, and cerebral peduncle; small hemorrhages involving the internal capsule, putamen, and basis pontis; saccular basilar artery aneurysm; brain abscess; cysticercosis; and primary or metastatic brain tumour. A recently reported study of a new model of stroke in the rat suggests that small deep infarcts may be produced by platelet emboli from the cervical carotid artery. Here we report a patient who had a pure motor stroke due to a pontine infarct caused by a presumed embolus from a vertebral artery dissection.

CASE REPORT

A 39-year-old man experienced the sudden onset of dizziness, speech disturbance, and left-sided weakness while hurrying to work on April 15, 1988. He was transferred from his local hospital to Camp Hill Hospital on April 19.

Nine days before the onset of these symptoms he completed a 10-day “course” of chiropractic treatment for a painful, stiff neck which he had had for about 6 weeks. Otherwise, he had been fit and well. There was no past history of migraine, neurological symptoms, cardiovascular disease, hypertension, diabetes, alcohol abuse, or use of tobacco.

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cocaine or other sympathomimetic drugs. Family history was negative
for premature vascular disease. He was not taking medications.

At the time of admission to our hospital, the general medical exami-
nation was unremarkable. Features of Marfan’s syndrome were absent.
While in hospital, his systolic blood pressure ranged between 110-140
mm Hg, and the diastolic between 70-80 mm Hg. No cardiac abnormal-
ities were detected, and there were no bruises over the head and neck.
Neurological examination showed a left hemiparesis involving face,
arm, and leg. No other cranial nerve signs were detected. Speech and
language were normal. There were no cerebellar or sensory signs. His
neurological deficit was moderately severe; he could not walk, and had
no useful function in the left upper extremity, i.e. grade 6 on the Stroke
Severity Scale.13

The following investigations were normal or negative: hemoglobin,
hematocrit, platelet count, ESR, serum proteins, serum protein elec-
trophoresis, VDRL, chest X-ray, and ECG. Plain films of the cervical
spine showed no evidence of congenital anomaly, subluxation, arthritic
or degenerative changes. An echocardiogram showed prolapse of the
anterior leaflet of the mitral valve.

A computerised tomographic (CT) scan done 6 days after stroke
onset was normal. A CT scan done 18 days after stroke onset showed a
right-sided mid-pontine infarct (Figure 1).

Four-vessel cerebral angiography was performed 13 days after
stroke onset. The extracranial carotid arteries and the intracranial circu-
lation were normal. Both posterior cerebral arteries filled from the
carotid system. The left vertebral artery tapered to an occlusion in the
foramen transversarium at the level of C6 (Figure 2). There was collat-
eral flow to the distal left vertebral artery via branches of the left occip-
tal artery. The right vertebral artery was irregularly stenosed through-
out much of its course in the foramen transversarium (Figure 3). The
distal right vertebral artery and the basilar artery appeared normal. The
angiographic appearances were considered indicative of bilateral verte-
brary artery dissection.

During a 16-day stay in hospital he received daily occupational
therapy and physiotherapy. At the time of discharge he still had a grade
6 neurological deficit on the Stroke Severity Scale,13 but his level of
functioning had improved. He was able to walk with the assistance of a
drop-foot splint and a cane, and opposition movements had begun to
return in the left hand.

**DISCUSSION**

Head and neck trauma is an important cause of cerebral
infarction in young adults.14,15 Cervical chiropractic manipulation
has been linked with dissections of the cervico-cerebral
arteries, particularly the extracranial vertebral artery.16-19
Vertebral artery dissections are typically bilateral in symptomat-
ic cases. They can occur at any point in the course of the verte-
brary artery, but are most frequent at the C1-C2 and C6 levels,
presumably because the artery is most subject to mechanical
stress at these points.19 Nucho-occipital pain and the lateral
medullary syndrome are the commonest clinical
manifestations.18,20,21 The “locked-in” syndrome,18,22 and fatal
brainstem infarction17 have also been reported in association
with vertebral artery dissections. Subarachnoid hemorrhage can result from dissection of the intracranial vertebral artery because this type of dissection tends to occur in the subadventitial plane.\(^23,24\)

We believe that our patient had bilateral vertebral artery dissections induced by cervical chiropractic manipulation. Atypical features were the absence of neck pain at the onset of neurological symptoms, and the 9-day delay between the last chiropractic treatment and the stroke. However, these features do not cast serious doubt on the diagnosis because painless dissections are known to occur, and the interval between trauma and stroke may vary from hours to several days.\(^16\) Angiographic diagnosis of vertebral artery dissection is presumptive, unless a pseudoaneurysm or double-lumen is demonstrated.\(^19\) These abnormalities are most likely to be demonstrated by early angiography; our patient was studied 13 days after stroke-onset.

The long ribbon-like narrowing of the right vertebral artery, and the tapering stenosis and occlusion of the left vertebral artery were considered diagnostic of dissection in this patient because of his age, the history of neck trauma, and the absence of any other cause for the angiographic findings. Echocardiography demonstrated the presence of mitral valve prolapse but an origin and tapers distally, an embolus small enough to traverse the vertebral artery would ordinarily not block the basilar artery. We think this is the first time a pure motor stroke has been associated with vertebral artery dissection. We think that the pontine infarct was probably caused by artery-to-artery embolism. This is a recognized mechanism of brainstem infarction,\(^25,26\) and our investigations failed to provide a viable alternative explanation.

In his description of the "top of the basilar" syndrome, Caplan\(^27\) commented that since the basilar artery is widest at its origin and tapers distally, an embolus small enough to traverse the vertebral artery would ordinarily not block the basilar artery except distally. However, the present case and recent experimental evidence\(^12\) suggest that small-diameter penetrating cerebral arteries can be embolized by platelet thrombi from lesions situated in more proximal segments of the arterial tree. We admit that the evidence on which our conclusions are based is circumstantial, but direct proof of such a stroke-mechanism would be difficult or impossible to obtain, even at autopsy.

Recognition of this mechanism of stroke is important both from the clinical and scientific points of view.\(^28\) Infarcts in the territory of end-arteries are not always due to small-vessel disease.\(^29,30\) The absence of pre-existing hypertension should prompt investigation for an alternative explanation, as it does in the normotensive patient who has a spontaneous intracerebral hemorrhage.\(^31,32\) Precise definition of the mechanism of stroke is necessary for rational therapy,\(^33\) and is especially important in young patients and individuals who have mild strokes. In research studies it is desirable to categorize strokes according to mechanism of causation; patients who have small deep cerebral infarcts do not form a pathogenetically homogeneous group.

The management of dissections is controversial.\(^16\) Antiplatelet agents or anticoagulants are often used, provided that intracranial vertebral artery dissection and subarachnoid hemorrhage have been excluded, but it is not known whether these drugs influence outcome. We did not treat our patient with these drugs because 4 days had elapsed between stroke onset and admission to Camp Hill Hospital, his neurological deficit had already started to improve, the vertebral artery injury was caused by an avoidable form of trauma, and there was no evidence of an underlying arteriopathy that could predispose to recurrent dissection.

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REFERENCES

33. Caplan LR. TIA: we need to return to the question, “What is wrong with Mr. Jones?” Neurology 1988; 38: 791-793.