SHORT REPORT

Cerebellar border zone infarcts are often associated with presumed cardiac sources of ischaemic stroke

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Abstract
It has been suggested that most border zone cerebellar infarcts are embolic infarcts or infarcts due to hypercoagulable states. The aim of this study was to test this hypothesis.

Risk factors for the presumed mechanism of stroke (TOAST criteria) were studied in 14 consecutive patients (nine men, five women; age range 29–84 years) with a total of 17 border zone cerebellar infaracts.

The presumed cause of stroke was "cardioembolism" in nine patients. Three patients had a dissection of the vertebral artery. Two patients had a negative diagnostic investigation, and one had a cardiac arrest.

These findings support the hypothesis that cardioembolism is a frequent mechanism of border zone cerebellar infarcts.

Keywords: cerebellar infarct; cardioembolism; stroke; cerebrovascular disease

Cerebellar infarcts account for about 2% of cerebral infarcts.1 Whereas previous studies took into account territorial cerebellar infarcts,1–4 Amarenco et al5 identified a subgroup of border zone cerebellar infarcts. Their most striking clinical characteristic is the long duration of positional symptoms.5 Their cause remains uncertain.5 Supra-territorial border zone infarcts are usually regarded as low flow infarcts8 but the recently categorised cerebellar border zone infarcts might be embolic infarcts or associated with a hypercoagulable state.5

The aim of our study was to describe the presumed mechanism of stroke in 14 patients with border zone cerebellar infarcts by means of a systematic search for large vessel atherosclerosis and cardioembolism.

Patients and methods
We conducted this study over a 19 month period (1 June 1992—31 December 1993) in 300 emergency admissions for ischaemic stroke or transient ischaemic attacks. During the study period, 11 patients (3·6%) had one border zone cerebellar infarct or more. During the same period, three other patients referred from other hospitals for vertebrobasilar transient ischaemia attacks had a border zone cerebellar infarct identified on MRI.

The study population consisted of nine men and five women, aged from 29 to 84 years. None of these patients had a history of neurosurgery, severe head trauma, or cerebral arterial or arteriovenous malformation. All patients underwent standard blood tests (including sedimentation rate, fibrinogen, packed cell volume, platelet count, and coagulation tests), 12 lead electrocardiography, and 24 hour continuous electrocardiography monitoring. We performed bimdimensional transtroch cerebrovascular echocardiography or transoesophageal echocardiography in all patients. All patients had an MRI except one who had two CT scans. The cervical arteries were studied by doppler ultrasonography and B mode echotomography in all patients, and complete cerebral angiography, including the subclavian and vertebral arteries, was performed in five.

Computed tomography was done without contrast with 5 mm contiguous slices using a Siemens Somatom II machine (Siemens, Germany).
Patients 1, 4, 12, and 14 presented with small cerebellar infarcts.

The presumed causes of stroke were defined according to the criteria used by Adams et al. in the trial of Org 10172 in acute stroke treatment (TOAST).  

**Results**

Table 1 presents the results of the study.

The 14 patients shared a total of 17 border zone cerebellar infarcts: eight in group 1, four in group 2, one in group 3, two in group 4, and two in group 5. The presumed mechanism was cardioembolism (high risk) in four patients: mitral and aortic prosthetic valves in one, left atrial thrombus in one, and septal post myocardial infarction akinesia in two. A medium risk source of cardioembolism was found in five patients: lone atrial fibrillation in two, atrial septal aneurysm associated with a patent foramen ovale in two, and isolated patent foramen ovale in one. Three patients had a dissection of one vertebral artery. One patient with a negative diagnostic assessment and one with a severe cardiogenic shock were classified in the group of undetermined cause.

No patient had a large vessel atherosclerosis as the presumed mechanism of stroke and the erythrocyte sedimentation rate, platelet count, packed cell volume, coagulation test, and serum fibrinogen concentration were normal in all patients.

The figure shows the locations of the infarcts in the 14 patients.

**Discussion**

Our study showed that, of 14 patients with one or more border zone cerebellar infarcts, nine had a potential cardiac source of stroke. Cardiac sources of emboli were more frequent in our study than in previous reports on border zone cerebellar infarcts. Severe vertebrobasilar atherosclerosis was the presumed cause in the eight patients of Barth et al. A focal cerebellar hypoperfusion was present in 27 patients from the 47 of the first study of Amarenco et al. In this series large artery disease was the principal cause (16 patients) of stroke and only seven patients had a cardiac source of embolism. In a more recent study, performed on a smaller group (36 patients), the cause of the border zone cerebellar infarct was cardioembolism in 42%. In our study three patients had large artery disease, and the nine patients with cardiac disease did not have large artery occlusion.

As three patients out of nine with potential cardiac sources of stroke had arterial hypertension, we cannot exclude small vessel
Cerebellar border zone infarcts are often associated with presumed cardiac sources of ischaemic stroke. The TOAST criteria may lead to an underestimation of the prevalence of small vessel occlusion.

Except for the patient with a cardiac arrest all patients had a complete diagnostic assessment. Only one patient was classified in the undetermined cause group by the TOAST criteria; this was less than in the studies of Amarenco et al. Surprisingly no patient had a hypercoagulable state whereas it was the third most common aetiology of border zone cerebellar infarcts in the study of Amarenco et al.

As in the previous studies haemodynamic causes of stroke were rare, suggesting that low flow infarcts are less frequent in border zone cerebellar infarcts than in supratentorial border zone infarcts. The exact cause of stroke in patient with heart disease and no evidence of focal hypoperfusion, however, remains uncertain. A haemodynamic mechanism cannot be excluded. Moreover the role of cardioembolism in the pathogenesis of supratentorial border zone infarcts could have been underestimated, especially in internal junctional infarcts.

Cardioembolism may be a frequent cause of border zone cerebellar infarcts; hence, appropriate cardiac investigations are necessary in these patients.

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