Regional cerebral blood flow by SPECT imaging in Sturge-Weber disease: an aid for diagnosis

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SUMMARY Regional cerebral blood flow (rCBF) was studied using SPECT (single photon emission computed tomography) with 133-Xenon in 13 patients with confirmed Sturge-Weber disease, aged 9 months to 18 years. CT scan, performed at the same time, showed evident cerebral angioma in 10 but not in three. A marked hypoperfused area was found in all patients, ranging from −32% to −72% and of the same location as the CT signs. The hypoperfusion seems to result from post ictal phenomenon as well as from chronic ischaemia. SPECT imaging is therefore a sensitive method for visualising intracranial angioma in Sturge-Weber disease and it provides an aid for diagnosis when a CT scan is not reliable.

Sturge-Weber (SW) angiomatosis is a non-inherited neurocutaneous syndrome characterised by a congenital trigeminal port-wine stain and a piamater angioma.1 This intracranial angioma, venous and purely meningeal,23 is embryologically derived from the neural crest. Along with the angioma, cerebral lesions, which are ischaemic in nature, are seen in the cortex underlying the angioma.4

The diagnosis of SW disease is based on the association of three characteristics — cutaneous, neurological and radiological. Among the neurological features are epilepsy, hemiplegia and mental retardation. CT is currently the most reliable technique used to confirm the existence of intracranial angioma by visualising calcifications, focal cortical atrophy and opacification after contrast enhancement.5 Diagnosis is easy in severe cases of this disease since all clinical and radiological characteristics are present, but it is difficult in milder cases where one or more of these features is absent, such as an absence of cutaneous angioma, a delayed onset of neurological signs6 or an atypical appearance of the angioma on the CT, on which calcifications, focal atrophy or angioma contrast may be absent, particularly in young children.8 Progressive aggravation of the neurological condition, for which seizures seem to be responsible, occurs in SW disease.4 Therefore, the diagnosis of this disease, that is, the presence of an intracranial angioma, needs to be determined.

Hypometabolism but also hypermetabolism have recently been reported using positron emission tomography (PET).9 We studied regional cerebral blood flow (rCBF) in children with confirmed SW disease to determine whether a characteristic rCBF pattern exists and to test the diagnostic value of this technique with that of CT.

Patients

We studied thirteen patients, eight boys and five girls, aged between 9 months and 18 years (mean 6 years). The examinations were performed with the informed consent of the children’s parents.

All 13 were diagnosed as having SW disease. Diagnostic criteria are summarised in table 1. Facial angioma was found in all patients except one (number 5). In this case, the neurological and radiological characteristics were those of Sturge-Weber disease; arteriography was normal, excluding other types of vascular malformation. Two patients did not have seizures before SPECT imaging (numbers 8, 9); one of these showed neither neurological disorders nor mental retardation (number 8) but there was no doubt about cutaneous and cerebral angioma. All patients showed asymmetrical electroencephalographic features, including depressed background activity on the side of the angioma (fig 1b), interictal focal epileptiform activity (fig 1b), or focal epileptic discharges (fig 1c). In ten patients, the CT revealed a typical pattern of intracranial angioma; for the three others, the CT disclosed only focal atrophy (number 9), only contrast enhancement (number 7), or no abnormality (number 12).
Patient number 7 had a special history. Anticonvulsant treatment had been given since birth for suspected SW disease because of facial angioma and to prevent seizures. Two CTs given when the patient was aged six months and one year were considered normal. Treatment was therefore stopped after the second CT. One month later, partial motor status occurred, confirming the diagnosis of SW disease. One year later, although epilepsy was severe, the patient showed no permanent focal motor deficit and a CT revealed only moderate abnormalities (table 1).

Methods

The rCBF was measured by SPECT with a Tomomatic 564 using 133 Xenon, and injected intravenously in doses ranging from 1.5 to 2.5 mCi/Kg. Data collection lasted 4.5 minutes. Results were obtained on five 20 mm-thick slices, parallel to the orbito-mental (OM) line, at levels OM + 20, OM + 40, OM + 60, OM + 80 mm and OM + 100 mm. The radiation dose to the target organ, the lung, was 250 to 350 mrad. To prevent the head from moving during acquisition, children under 6 years old received 4 mg/Kg of rectal pentobarbital and 0.5 mg/Kg of intramuscular droperidol. All patients were receiving anticonvulsant drugs (table 1) and none of the patients experienced seizures on the day of the study.

The rCBF was measured on 20 cortical regions of interest (ROI) per slice. The ROI were circular and ranged from 1.5 to 2.5 cm² in size. For each patient, the mean rCBF or ‘‘global rCBF’’, was calculated as the mean half-slice rCBF for slices OM + 40, + 60, and + 80, on the undamaged or the less damaged side. ROI rCBF values were expressed as a percentage of the individual mean rCBF and also as a percentage of the contralateral value in the symmetrical ROI. ROI rCBF was classified as ‘‘decreased’’ when the difference with the contralateral value was greater than 20%.

A CT, with contrast injection, was performed on each patient, using the same head position as that required for the SPECT study. Nine mm-thick slices were obtained at the same levels OM + 20, + 40, + 60, + 80, and + 100, as in the SPECT study. Both studies were performed at the same time in all except two patients for whom there was a time interval of 7 and 26 months between the CT and the SPECT (numbers 6 and 7 respectively); no neurological changes were observed during this time interval. The ROI employed on SPECT images were traced on the CT images after the size of the ROI was homothetically reproduced on the CT image. Three types of abnormalities were studied on the CT scan: calcifications, atrophy and contrast enhancement after injection.

Results

In six patients (numbers 1–4, 11, 13), the CT abnormalities were large since contrast enhancement and calcifications covered an entire hemisphere. In all six patients, an rCBF decrease was observed in a large cortical and subcortical area of the same hemisphere. In the cortex, the hypoperfused area was approximately of the same size as the CT abnormal area (fig 1A). In the subcortical regions, the size of the hypoperfused area was easily measured, but could not be compared with that of the CT abnormal area which was poorly defined. The intensity of the rCBF decrease was considerable, with maximal value ranging from 32% to 72% depending on the patients.
Fig 1 Patient number 13. (a) Large abnormal CT area involving the whole left hemisphere (top) with a hypoperfused area (bottom) of approximately the same size. (b) Depressed interictal background activity on the whole left hemisphere. (c) Left temporal epileptic discharge at the same age.
Table 2 Comparative data on SPECT and CT scan

<table>
<thead>
<tr>
<th>Number of patients</th>
<th>Location of CT abnormalities</th>
<th>Location of SPECT hypoperfusion</th>
<th>Number of pathological cortical ROI</th>
<th>Minimal rCBF observed (symmetrical rCBF) ml/100g/mm</th>
<th>Mean rCBF ml/100g/mm</th>
<th>Maximal rCBF decrease*</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>R diffuse</td>
<td>R diffuse</td>
<td>On CT 30</td>
<td>21 (58)</td>
<td>52</td>
<td>-59%</td>
</tr>
<tr>
<td>2</td>
<td>Bilateral atrophy</td>
<td>L central</td>
<td>On SPECT 30</td>
<td>46</td>
<td>47</td>
<td>-45%</td>
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<tr>
<td>3</td>
<td>R diffuse</td>
<td>R diffuse</td>
<td></td>
<td>31 (43)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>L posterior</td>
<td>L posterior</td>
<td></td>
<td>25 (37)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>R diffuse</td>
<td>L central</td>
<td></td>
<td>47</td>
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<tr>
<td>6</td>
<td>L posterior</td>
<td>R posterior</td>
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<td>39 (71)</td>
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<tr>
<td>7</td>
<td>R posterior</td>
<td>L posterior</td>
<td></td>
<td>73</td>
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<tr>
<td>8</td>
<td>R posterior</td>
<td>R anterior</td>
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<td>73</td>
<td></td>
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<tr>
<td>9</td>
<td>R posterior</td>
<td>L posterior</td>
<td></td>
<td>67</td>
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<tr>
<td>10</td>
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<td>R posterior</td>
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<td>63</td>
<td></td>
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<tr>
<td>11</td>
<td>L diffuse</td>
<td>L posterior</td>
<td></td>
<td>72</td>
<td></td>
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<tr>
<td>12</td>
<td>R posterior</td>
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<tr>
<td>13</td>
<td>L diffuse</td>
<td>L diffuse</td>
<td></td>
<td>80</td>
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</tbody>
</table>

*Expressed in percentage of mean rCBF.

In six patients (numbers 5–10), the CT abnormalities covered a small area. In all six, an rCBF decrease was found in an area corresponding to the CT scan abnormal area. In the cortical region, the size of the hypoperfused area was similar to that of the CT abnormal image in three patients (fig 2) and slightly larger in the other three (numbers 7, 9, 10) (figs 3, 4). In these six patients, the rCBF decrease was also marked,
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Fig 3  Patient number 9. CT scan performed after contrast injection (top) shows no enhancement but only focal atrophy (the posterior contrast area corresponds to the superior longitudinal sinus). The hypoperfused area is clearly seen on rCBF images (bottom).

with individual values ranging from 32% to 45%.

In the remaining patient (number 12), SPECT showed a localised rCBF decrease, with maximal decrease of 50%, although the CT was normal (fig 5).

These results show that in all the patients in this series, SPECT clearly defined a hypoperfused area, which was also seen in the patients with or without the typical appearance of angioma on CT.

Discussion

The diagnosis of Sturge-Weber disease is based on the association of clinical and radiological characteristics. Cutaneous angioma is usually present at birth. Epilepsy usually appears in the first years of life and hemiplegia follows the first prolonged seizures although some patients may remain seizure free. Cerebral angioma is constant but radiological characteristics may be delayed, or reduced to non-specific features, or may even be occasionally absent. Sturge-Weber disease is easily diagnosed when all the features are present, but diagnosis is difficult when the neurological characteristics are absent, and especially when a CT provided no additional information. An additional diagnostic method would therefore be useful.

The method used to measure rCBF has been previously described. It is easy to perform, does not require any blood samples, and only takes 5 minutes. The SPECT system used is especially adapted to the brain and is highly sensitive. The validity of this method has been studied for the cortex by comparing rCBF values in the same patients with those obtained by SPECT and PET using continuous inhalation of C15O2. The rCBF values obtained by both methods were significantly correlated. Adapting the SPECT technique for children required two modifications: intravenous (IV) injection of the 133-Xenon instead of inhalation and the use of premedication. Using the IV injection method, rCBF values increased by a mean of 10%, probably due to the shape of the Xenon input curve. The premedication used (pentobarbital and droperidol) did not cause any significant change in rCBF in our research.

A study of patients with confirmed Sturge-Weber disease was necessary to test the ability of SPECT imaging to visualise the intracranial angioma. Our results show that all patients had a marked hypo-
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Fig 4  Patient number 7. (a) CT scan after contrast injection at one year is considered normal (top). SPECT imaging at one year five months shows a well defined posterior hypoperfused area (middle). At two years, CT scan after injection only shows a discrete enhancement in the posterior cortex and a hypertrophy of the left choroid plexus (bottom). (b) Left temporal focus of spikes and right temporal focus of slow waves on the interictal EEG recorded at one year six months.
perfusion in the pathological cerebral regions, suggesting that such hypoperfusion is a reliable sign of Sturge-Weber disease. In our series, the CT scan pattern was incomplete in some patients and diagnosis was established after the neurological features appeared: in one of them CT scan was normal, in another it showed such mild abnormalities that they had been overlooked on first examination. In three patients, the clinical pattern was incomplete. Two of the three had never had any seizures. However, in these patients, a hypoperfused area was also clearly defined. This indicates that CBF decrease may be present even when CT and neurological features are incomplete or absent. SPECT imaging may therefore be considered a complementary method for diagnosing Sturge-Weber disease, which is especially useful in cases with an atypical CT scan.

The hypoperfusion described in our patients should not be confused with the hypoperfusion of immature areas, which is seen before the age of six months in the temporoparieto-occipital cortex and is always symmetrical. Hypoperfusion should not be related to the anticonvulsant drugs taken by our patients because such drugs produce diffuse changes in blood flow and metabolism but not focal defects. The hypoperfusion observed here is therefore probably due to the lesions of SW disease.

Focal hypoperfusion is a non-specific finding that can reflect different mechanisms. Two of them may be taken into account in this disease, ischaemic and epileptic. The anatomical lesions described in Sturge-Weber are mainly angioma and ischaemic lesions. The angioma affects the superficial areas because it is meningeal. Ischaemia progressively develops in the brain underlying the angioma and involves cortical and subcortical regions, due to the obstruction by the angioma of the normal venous return. This produces an abnormal drainage into the deep plexus and hypertrophy of the choroid plexus visible on CT scan. Stasis and slowing in venous cortical circulation have been demonstrated by dynamic angiography, and hypoxia, preoperatively, has been shown. The ischaemic mechanisms alone may explain the hypoperfusion seen in our two non-epileptic patients.

Hypoperfusion may also be a consequence of the seizures in the post-ictal period. Evidence of consistent presence of epileptic foci related to the angioma is well known in SW disease and was also seen in our series. This mechanism alone may be involved in our patient with seizures without CT scan abnormalities, as described in partial epilepsy. Alexander suggests that both ischaemic and epileptic mechanisms may be involved together. The complex circulatory condition in SW disease may explain why seizures are particularly severe and often followed by permanent hemiplegia. In normal conditions, cerebral blood flow locally increases during seizures. In SW disease, the insufficient increase in blood supply during seizures may exacerbate ischaemic brain lesions. But the respective responsibility of ischaemic and epileptic phenomena in lesional mechanisms remains unclear, as well as the relation between hypoperfusion and the reported abnormalities on metabolism.

Our results show that the considerable hypoperfusion observed by SPECT imaging in patients with Sturge-Weber disease is a reliable characteristic of the disease. Since it is consistently observed, even in cases without clear evidence of angioma on CT, SPECT imaging may be considered a useful complement to CTs for diagnosing SW disease in childhood. Further studies are needed to test its diagnostic value in the first year of life.

We thank Doctors J Motte, C Billard, P Pineau and C Bouilloche for their kind assistance.

References

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J Neurol Neurosurg Psychiatry 1989 52: 1402-1409
doi: 10.1136/jnnp.52.12.1402

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