Haemogram, liver function tests, serum, calcium, phosphorus, copper, caeruloplasmin estimations were normal. Blood sugar levels in the fasting state and during the ictus were normal. Slit lamp examination for K-F ring was negative. Radiological evaluation with CT scan of the head, radiographs of the chest and skull was normal. An interictal EEG whilst awake, showed frontally dominant intermittent rhythmic delta activity. No epileptiform discharges were seen.

Serial trials of phenobarbitone (120 mg/day), phenytoin sodium (300 mg/day) and clonazepam (2 mg/day) for four weeks each did not help. A year later the attacks were unchanged.

Paroxysmal choreoathetosis emerged as a clinical entity in a description of a family by Mount and Reback in 1940. Clinical features of the three subgroups are summarised in the table. Our case belongs to the intermediate type and is the first sporadic case in this group. The history, examination and investigations ruled our underlying conditions, such as, cerebral palsy, multiple sclerosis, Wilson’s disease, idiopathic hypoparathyroidism, hypoglycaemia, encephalitis or space occupying lesion which could produce focal dyskinesias.

The well documented families of Lance and Plant et al, which form the basis of the intermediate type, are characterised by cases whose attacks were induced by prolonged exercise. In our case the attacks were not only induced by prolonged exercise, such as, running, but also by prolonged work with a part of the ipsilateral limb. A similar observation is made in case 1 by Plant et al in which writing to dictation could induce the attacks of involuntary movements.

As a trigger factor cold temperature has not been reported in the intermediate type. Absence of response to warm water suggests that the trigger factor is cold related and not related to the striking force of the water drops to the body surface. Cold breeze also induced the attacks. This is the first case of the intermediate type to show unilaterality of the involuntary movements. In the cases of Lance and Plant et al, the involuntary movements predominantly affected both the lower limbs and only when severe, involved the whole body. The EEG finding of frontally dominant intermittent rhythmic delta activity in the awake state is another new observation although possibly of non-specific significance. Previously reported cases had normal EEGs.

The trigger factors and involuntary movements in our case, suggest a functional disturbance of the contralateral basal ganglia or their connections leading to abnormal responsiveness to external stimuli sensed by kinaesthetic and cold receptors.

I am grateful to the Principal, Jawaharlal Nehru Medical College and the Administrator of the KLE Society's Hospital and Medical Research Centre for granting permission to publish this report.


She improved with simple analgesics and was allowed home after four days. Six weeks later she again presented with a two week history of progressive generalised headaches, vomiting and a left sided weakness. Examination revealed a left sided weakness, the presence of primitive frontal reflexes but no papilloedema. She was pyrexial (temperature 37.5°C). Further CT (figure b) showed a cystic enhancing mass in the right frontal area. A right frontal burr-hole was made and the mass aspirated, 40 ml of pus was obtained. Bacteriological studies showed Gram positive cocci and a growth of Staphylococcus aureus.

She was initially treated with gentamicin, metronidazole and benzyl penicillin. This was later changed to lincomycin, sodium fusidate and metronidazole. Antibiotics were continued for ten weeks—one week intravenously, nine weeks orally. Her symptoms improved as did the left hemiparesis; the

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**Table**

<table>
<thead>
<tr>
<th>Inheritance</th>
<th>Paroxysmal dystonic choreo-athetosis</th>
<th>Paroxysmal kinesigenic choreo-athetosis</th>
<th>Intermediate type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autosomal dominant</td>
<td>Single sporadic Case</td>
<td>Autosomal dominant</td>
<td>Autosomal dominant</td>
</tr>
<tr>
<td>Age of onset</td>
<td>Neonatal/infancy</td>
<td>Infancy to 4th decade</td>
<td>2/month; rarely daily</td>
</tr>
<tr>
<td>Frequency of attacks</td>
<td>1/month to 3/day</td>
<td>Usually 1 minute; always less than 5 minutes</td>
<td>5 to 30 minutes</td>
</tr>
<tr>
<td>Duration of attacks</td>
<td>2 min to 6 hours</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Induced by</td>
<td>Alcohol, other drinks, fatigue, rest after physical stress, excitement</td>
<td>Sudden movements, startle; Continuous exertion, passive movements, local vibration</td>
<td></td>
</tr>
<tr>
<td>Therapy</td>
<td>Clonazepam</td>
<td>Phenobarbital</td>
<td></td>
</tr>
</tbody>
</table>

Modified from Buruma and Roos¹

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Figure 1 An attack of right hemidystonia induced by treadmill exercising.

Figure 2 An attack induced by cold water shower.
primitive frontal reflexes slowly disappeared and she remained appyreal. CT scans at weekly intervals showed no re-accumulation (figure c) and she regained her normal state of health.

The risk of infection after craniotomy is between 1.1–3.5% but after surgery from aneurysm, is less than 1%. Cerebral abscesses are common secondary to otitogenous-laryngological infection. In this case craniotomy did not pass through an infected sinus. Among other causes are haemogenous spread in patients with cyanotic congenital heart disease and following cysophageal dilatation. In our case, the infection was most likely introduced at the time of operation. The mortality from intra-cerebral abscess ranges between 10–70% in untreated patients. The surgical management options include: tube drainage, mastupalisation, Kahn’s migration method, tapping, aspiration and excision. At present only two methods are used, aspiration and excision. Excision of the abscess cavity carries a high risk of mortality and morbidity.

MRI in hemiballism due to Sydenham’s chorea

Parainfectious and immunologically mediat-
ed chorea may be caused by a cytotoxic reaction in the basal ganglia. We report the MRI appearances in a patient with hemiballism due to Sydenham’s chorea.

A 17 year old woman showed right sided hemiballism of subacute onset. She had a history of juvenile rheumatoid arthritis and repeated tonsillitis, but no family history of neurological diseases. Routine laboratory investigations showed normal blood cell counts and blood chemistry. Positive immunological findings were antistreptolysin O (ASO) 181 (< 155), anti-streptokinase (ASK) 256(< 640) and decreased IgA of 32-9 mg/ dl (135–340). There were no abnormal find-
ings in anti-virus titres, hormonal levels, CSF and ECG. Although her involuntary move-
ments were reduced by administration of sodium valproate and phenytoin, phenytoin caused an allergic agranulocytosis which required treatment with prednisolone, 40 mg daily for 14 days. Prednisolone alleviated both the drug allergy and involuntary move-
ments. After withdrawal of prednisolone hemiballism reappeared and was treated with haloperidol 5 mg IM followed by 2.25 mg daily by mouth for 20 days and the hemiballism resolved during this time. CT scan showed left caudate nucleus swelling only, and angiography was normal. T2 weighted MRI 31 days after onset of symptoms dem-\nonstrated hyperintensities in the left caudate nucleus, putamen, lateral pallidum, periru-
bral area and substantia nigra (figure). Hyperintensity in the prerubral area sug-
gested involvement of the subthalamic nucleus or its connecting pathways. Fourteen months later, she had a convulsion with mental confusion. No ballistic movements were observed and a repeat MRI was normal. There were bursts of theta and delta wave activity in the EEG. Serological test showed positive antinuclear antibody (ANA: X320, speckled type) and negative anti-DNA antibody.

Despite the fact that systemic chorea is associated with systemic disease, unilateral involuntary movement are often seen. The lesions in our case seemed to spread during neural connections rather than in a vascular territory. The lesions were oedematous but were unlike lesions seen in vascular accidents or multiple sclerosis plaques. An allergic reaction to phenytoin, amelioration of hemiballism by prednisolone and positive ANA, which appeared later, suggest an immuno-
nologically mediated disorder. The history of repeated tonsillitis and elevated ASO and CRP indicated the diagnosis of Sydenham’s chorea. Sydenham’s chorea is thought to be an immunologically mediated reaction in the basal ganglia and subthalamic nucleus. Increased signal intensity in T2 weighted MRI at an early stage suggests an exudative or inflammatory lesion, which is compatible with the hypothesis of an immunological reaction in the basal ganglia.

Besides lesions of the subthalamic nucleus, hemiballism can be caused by single or combined lesions of the striatum, pallidum, substantia nigra, thalamus and their connec-
tions in each other. Where the pallidum is affected, the lesion is usually in the outer segment. Disinhibition of the medial palli-
dum by the lesion of the inhibitory sub-
thalamopallidal pathway and a subthalamic nucleus itself is thought to cause ballism. MRI of our case showed a widespread lesion in the basal ganglia and related structures on the left. Of these, the perirubral area, where the subthalamopallidal pathway is located, and lateral pallidum were thought to be responsible for hemiballism.

Radiological findings of symptomatic chorea have been poorly documented. MRI of the present case offers the information about distribution and possible nature of the lesion.

Correspondence to: Dr Kagnaya

Letters to the Editor

Figure A) CT without contrast showing low density area in the right frontal region; B) CT without contrast showing cystic mass in the right frontal area which is enhanced; C) CT scan at three months after aspiration showing low density area. There is no abscess and no shift.