Acute rotatory vertigo caused by a small haemorrhage of the vestibular cortex

Central rotatory vertigo is in most cases caused by a lesion of the cerebellum or brain stem. We describe a patient with acute rotatory vertigo following a small haemorrhage in the left medial temporal gyrus, which probably injured the vestibular cortex.

Case history

A 53 year old woman suddenly experienced leftwards directed rotatory vertigo in the yaw plane and nausea without vomiting. She felt unsteady and had short lasting slurring of her speech. She had no hearing loss or tinnitus. On examination, she could stand unaided but tended to fall after a short while, without a directional preponderance. Gait was severely unsteady and she could not walk unaided. The rotatory vertigo was worse when she was sitting upright than when lying down in bed. Vertigo was also increased by head movements.

Examination of the cranial nerves showed no abnormalities; specifically there was no nystagmus or hearing loss and the eye movements were normal. Neurological examination showed no signs of a stroke (motor and sensory function, coordination, and reflexes) was normal. Electroencephalography showed no abnormalities, supporting a non-epileptic cause of the vertigo.

Magnetic resonance imaging (MRI) on sagittal T1 weighted and transverse T2 weighted spin echo and FLAIR images showed a small (2.0 × 1.5 cm) haemorrhage in the left medial temporal gyrus, adjacent to the superior temporal sulcus (fig 1). There were no lesions of the brain stem or cerebellum. The appearance of the temporal lesion was consistent with haemorrhage from a small cavernous haemangioma.

Functional brain studies have shown that the human vestibular cortical system is located in the superior temporal region posterior to the superior temporal sulcus (fig 1). There were no lesions of the brain stem or cerebellum. The appearance of the temporal lesion was consistent with haemorrhage from a small cavernous haemangioma.

Vertebrobasilar insufficiency is not a common cause of acute vertigo. We describe a patient with acute vertigo, whose vertebral artery was occluded by a small haemorrhage, which was confirmed on MRI. The vertebrobasilar insufficiency was confirmed by transcranial Doppler ultrasonography, which showed a substantial reduction in blood flow in the left posterior cerebral artery.

Vertebrobasilar insufficiency is not a common cause of acute vertigo. We describe a patient with acute vertigo, whose vertebral artery was occluded by a small haemorrhage, which was confirmed on MRI. The vertebrobasilar insufficiency was confirmed by transcranial Doppler ultrasonography, which showed a substantial reduction in blood flow in the left posterior cerebral artery.

References


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Figure 1 Transverse T2 weighted, fluid attenuated inversion recovery (FLAIR) image (left) and sagittal T1 weighted spin echo image (right). Both show a small popcorn shaped area of increased signal intensity demarcated by a rim of decreased signal intensity (haemosiderin), located in the left medial temporal gyrus. The combination of recent blood products and older haemorrhagic residues is consistent with the diagnosis of a cavernous haemangioma.
Coexistent Lewy body disease in a case of “visual variant of Alzheimer’s disease”

Posterior cortical atrophy or the “visual variant” of Alzheimer’s disease is a clinical syndrome with visual agnosia, some or all of the components of Balint’s syndrome, transpatial sensory aphasia, and Gerstmann’s syndrome.1 Although pathologically heterogeneous, several necropsy studies on patients with posterior cortical atrophy have shown Alzheimer’s disease pathology.2 We report a patient who presented with the features of posterior cortical atrophy who later developed mild parkinsonism, visual hallucinations, and dementia. Neuropathological evaluation revealed coexistent Alzheimer’s disease and Lewy body disease.

Case report

A right handed retired diesel mechanic, with 12 years of formal education, was referred for evaluation of an “unusual dementia.” His difficulties started at the age of 58 with the initiation of visuospatial dysfunction. Initially he was not able to fill out bank deposit slips or write numbers correctly. He had been an avid reader but had to re-read material in order to comprehend it, and unsuccessfully used a card to keep his eyes focused when reading. He was not able to locate the refrigerator door handle until he groped over the surface to find it. He revealed that when he was 61 he was having difficulties working as a mechanic. Also, he could not see other cars and obstacles while driving, and he stopped driving at 63 after being involved in two motor vehicle accidents. He developed progressive difficulties with performing calculations, writing, receptive language, and recent memory. Despite the cognitive difficulties, he retained insight in his disorder.

When he was 62, his wife noted that he moved in a stiff manner, did not swing his left arm, and acted “like a little old man.” At age 67, he developed well formed visual hallucinations (he would see bugs, spiders, and people) and paranoid delusions (he expressed concern that people were tearing away his home). He developed personality changes and at times was ophthalmological. He became entirely dependent on his wife for all of his activities of daily living. No features of REM sleep behaviour disorder were ever noted by the family.

The initial neurological evaluation at the Mayo Clinic when he was 67 revealed a complete Balint’s syndrome, a partial Gerstmann’s syndrome, and impairment on visuospatial tasks and recall. On language examination he had paraphasic errors and neologisms. He also showed bradykinesia, a slow wide based gait with reduced arm swing bilaterally, mild generalised rigidity, postural but not resting tremor, and rigidity of the limb apaxes. He had limited upgaze but preserved downward and horizontal gaze. Visual acuity was 20/80 and 20/100 in the right and left eye, respectively. There was no alien limb phenomenon, dystonia, or myoclonus.

Neuropsychological testing showed impairment in verbal skills and verbal memory and the inability to complete the visual tasks. Magnetic resonance imaging and single photon emission computed tomography of the brain showed, respectively, marked asymmetrical (left more than right) parietal-occipital cortical atrophy and hyperperfusion.

Towards the end of his life, he became wheelchair bound and was transferred to a chronic care facility. He developed more behavioural problems, declining vision, and persistent visual hallucinations and delusions. He was unable to recognise family members by sight or sound. He died at 71 years of age.

His past medical history was only significant for a total thyroidectomy for cancer, for which he was on thyroid replacement. There was no family history of any neurodegenerative disorder.

At necropsy examination, standard brain fixation and dissection was undertaken. Tissue sections were cut and stained with haematoxylin and eosin, Bielchowsky silver stain, and immunohistochemically with antibodies to tau (Endogen-AT8), amyloid protein, and synuclein (Zined-LB509).

The brain weighed 1136 g. Focal, asymmetrical (left greater than right) parieto-occipital cortical atrophy and mild pallor of the substantia nigra were observed. The basal ganglia, thalamus, and cerebellum appeared normal. Microscopically, moderate to frequent diffuse and neurtic plaques and frequent neurofibrillary tangles were seen in limbic structures. Accented neuronal loss and increased neurofibrillary tangle density were noted in the limbic lobes. The findings satisfied criteria for Alzheimer’s disease by Braak and Braak staging (stage V/VI)3 and by the National Institute on Aging and Reagan Institute working group diagnostic criteria for the neuropathological assessment of Alzheimer’s disease (high likelihood).4 In addition, synuclein positive Lewy bodies, pale bodies, and Lewy neurites were found in the substantia nigra, amygdala, entorhinal cortex, and cingulate gyrus; however, the substantia nigra was less affected than the limbic structures, where synuclein pathology was severe. These findings are consistent with a diagnosis of transitional Lewy body disease.5

Comment

The clinical syndrome of posterior cortical atrophy is characterised by prominent dysfunction of the neuronal networks in the biparietal and occipital cortices and does not imply an underlying pathology; neuropathological examination in most cases shows neurofibrillary tangles and neuritic plaques characteristic of Alzheimer’s disease, but with a higher concentration of the pathology located in the parieto-occipital cortices and does not characterise the function of the neuronal networks in the parieto-occipital cortices.6 The predominant features of posterior cortical atrophy are followed by dementia more typical of Alzheimer’s disease, but with a higher concentration of the pathology located in the parietal lobe. This suggests that visual hallucinations and parkinsonism in the setting of cognitive impairment reflect underlying Lewy body disease rather than corticobasal degeneration.7 Our case supports this contention. We suggest that underlying Lewy body disease should be considered in any case of posterior cortical atrophy associated with parkinsonism and particularly visual hallucinations.

Acknowledgements

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Reversible collasal disconnection syndrome in internal hydrocephalus

A 74 year old woman was referred to the neurological department for evaluation of progressive gait disturbance. On admission she complained of alienation of her left arm for six months. There were no spontaneous movements without the patient’s intention, but she had always to rely on visual cues. For example, when she was cooking, eating, or doing exercises with her home trainer she had to watch her left arm to
be sure of its movement. On examination she was alert, fully oriented, and cooperative. Snout and palmaromental reflexes were positive. There was no visual, somatosensory, or auditory extinction. Motor examination revealed a mild left sided facial droop and a left sided pronator drift but strength was full and symmetric and there was no lack of spontaneous movement in the left upper limb. The plantar response was extensor on the left. Gait was slow, unsteady, and wide based. The steps were short with reduced step height. Neuropsychological assessment showed fluent speech without dysarthria. Comprehension and reading were intact. Performance in verbal and non-verbal fluency tasks was diminished, and colour-word interference was slightly increased. Long term memory was slightly deficient for verbal and non-verbal material. Visuo-constructive abilities were normal and there was no spatial neglect. There was no apraxia of the left hand for gestures neither on command nor for imitation. Also, there was no apragia or tactile anoma of the left hand. She could perform bimanual tasks without evidence of intermanual conflict. She did not exhibit grasp reflex in either upper limb, and there was no compulsive manipulation of objects.

There was, however, an inability of one hand to imitate the posture of the opposite hand when visual cues were removed. Furthermore, there was an inability to distinguish the left hand from an examiner’s hand when these were placed in the patient’s right hand behind the back, which is known as “strange hand” sign (or “signe de la main étrangère”). Additionally, an inability to cross locate touch of the fingers was found: the patient was blindfolded and touched by the examiner on one finger either of the left or the right hand. Then she was asked to point to the location of touch with the contralateral hand. The accuracy was impaired for both directions but especially for right to left pointing. However, she was correct when asked to point to the location of touch on the face or trunk.

Magnetic resonance imaging showed internal hydrocephalus (fig 1A) and an old lacunar ischaemic lesion in the right anterior limb of the internal capsule. Transcallosal inhibition was assessed by transcranial magnetic stimulation (TMS) as described previously and showed a deficient inhibition particularly from left to right (upper panels “left A” in fig 1B). Cerebrospinal fluid (CSF) pressure was normal during lumbar puncture. After removal of 50 cc CSF the alienation of the left arm, the “signe de la main étrangère” and the impaired cross replication of hand postures

![Figure 1](http://jnnp.bmj.com/)

**Figure 1**  
(A) Magnetic resonance image (T2 weighted) showing internal hydrocephalus with thinning of the corpus callosum.  
(B) Recordings of the rectified tonic electromyographic (EMG) activity of the first dorsal interosseous muscle (ID1) after ipsilateral focal transcranial magnetic stimulation (TMS) are shown. Traces of three trials are superimposed over each other. The recording and stimulus sides as well as the latencies and durations of the transcallosal inhibitory responses (TI) are indicated. Upper panels A: the findings before lumbar puncture are shown. TI is normal after TMS of the right hemisphere (normal values [mean (SD)]; latency 35.8 (7.2); duration 24.8 (5.4)), but missing on the left (arrow). Lower panels B: results after withdrawal of 50 cc cerebrospinal fluid (CSF). TI could be revealed on both sides. The latencies after the lumbar puncture differed clearly and were prolonged after TMS of the left hemisphere.
References


Trigeminal autonomic cephalalgia-tic-like syndrome associated with a pontine tumour in a one year old girl

The so called trigeminal autonomic cephalalgia-tic (TAC) include episodic and chronic paroxysmal hemicrania (CPH), short lasting unilateral neuralgiform headache with conjunctival injection and tearing (SUNCT), and cluster headache (CH). Combinations of cluster headache and chronic paroxysmal hemicrania with trigeminal neuralgia have also been described and have been called cluster-tic syndrome or CPH-tic syndrome.

In order to diagnose TACs, it is essential to record the case history carefully. Only rarely have intracranial lesions such as aneurysms or tumours been observed in association with TACs. In the majority of cases, no brain abnormalities are found using conventional imaging.

We describe a three year old girl who suffered attacks of severe right sided temporal pain and autonomic disturbances and in addition neuralgic shooting pains associated with a pilocytic astrocytoma in the pons and medulla oblongata, extending to the upper cervical cord. The attacks disappeared once the tumour had been debulked.

Case report

A three year old girl presented with a history of extremely painful right sided temporal headache attacks since the age of one year. During these attacks she would grab her right ear and cry intensely. Her eyelids were slightly swollen, with rhinorrhoea on the right side. It seemed that she was in continuous intense pain, lasting seconds, during which she rocked back and forth (the video recording is available at the JNPN website: www.jnnp.com). These attacks occurred spontaneously at any time of day or night with no particular regularity or trigger points. Magnetic resonance imaging (MRI) of the brain revealed a tumour in the pons, extending to the medulla oblongata and cervical myelum (fig 1A), with a syrinx in the cervical myelum. On the transverse slide, the tumour extended to the cerebropontine cistern on the right side (fig 1B). Repeated physical examinations after six weeks showed hyperreflexia of the right arm and leg and positive Babinski reflexes on both sides. After debulking the tumour, the attacks resolved completely and neurological examination normalised.

Comment

The diagnosis of primary headaches associated with autonomic symptoms, such as cluster headache or chronic paroxysmal hemicrania, is based on the patients’ history, because diagnostic tests are not available. As shown here, a video recording of the attacks may be very helpful when patients are unable to describe the attacks in detail themselves. In this case the autonomic symptoms during the headache and the sudden additional shooting pains were recognised by the parents and the physician after studying these video recordings. Although the attacks lasted 12 to 24 hours, the combined headaches best resembled a combination of a TAC-like syndrome in association with trigeminal neuralgia or idiopathic stabbing headache (ISH). It is difficult to distinguish trigeminal neuralgia from ISH in this case, as both headaches only last seconds, may occur many times per day, and not effective. There was no family history of migraine or other severe headaches.

A video recording of several attacks made by her parents showed the child lying still while crying intensely. Her right upper and lower eyelids were swollen and tear production and rhinorrhoea were seen on the right side. It seemed that she was in continuous pain, with superimposed paroxysms of very intense pain, lasting seconds, during which she rocked back and forth (the video recording is available at the JNPN website: www.jnnp.com). These attacks occurred spontaneously at any time of day or night with no particular regularity or trigger points. Magnetic resonance imaging (MRI) of the brain revealed a tumour in the pons, extending to the medulla oblongata and cervical myelum (fig 1A), with a syrinx in the cervical myelum. On the transverse slide, the tumour extended to the cerebropontine cistern on the right side (fig 1B). Repeated physical examinations after six weeks showed hyperreflexia of the right arm and leg and positive Babinski reflexes on both sides. After debulking the tumour, the attacks resolved completely and neurological examination normalised.
Autoimmune myasthenia gravis after cardiac surgery

Autoimmune myasthenia gravis (MG) is a heterogeneous disorder. In young women, the thymus gland is often hyperplastic, and the patients respond well to thymectomy. However, in the increasing number of patients over the age of 40 years, predominantly men, thymic hyperplasia is uncommon, and there are no clear aetiological clues.

We diagnosed MG in three male patients who had undergone cardiac surgery between three and ten weeks before developing symptoms. Table 1 summarises their main features. The patients presented with ocular, bulbar, and generalised weakness (Oksanen classification grade I to IIb). None of them was taking antiarrhythmic agents or other drugs impairing neuromuscular transmission and none of them had had any thyroid or other autoimmune disease. Two showed typical decremental responses to repetitive stimulation on ENG, and all three had positive levels of acetylcholine receptor (AChR) antibodies. None had had postoperative difficulties, which are common encountered in undiagnosed MG, and presented de novo to the immune system; the antithymocyte antibdy response, or allow AChR to be released from the thymus, the presence of other autoantibodies or signs of other autoimmune diseases should be sought.

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Table 1
Clinical and laboratory features of three patients with autoimmune myasthenia

<table>
<thead>
<tr>
<th>Sex, age at the onset of MG</th>
<th>Heart surgery</th>
<th>Time since surgery to onset of MG</th>
<th>Presenting features; (Oksanen group at onset)</th>
<th>Clinical diagnosis</th>
<th>AChR antibodies; normal value &lt;0.8 nmol</th>
<th>Treatment and response</th>
<th>Follow up and evolution AChR-ab present titre</th>
</tr>
</thead>
<tbody>
<tr>
<td>M, 57</td>
<td>Trigeminal aorto-coronary bypass</td>
<td>3 weeks</td>
<td>Ptosis, double vision; (I)</td>
<td>Response to im neostigmine and oral pyridostigmine</td>
<td>1.9 nmol</td>
<td>Cholinesterase inhibitors: good</td>
<td>6 years, fluctuating</td>
</tr>
<tr>
<td>M, 58</td>
<td>Double aorto-coronary bypass</td>
<td>10 weeks</td>
<td>Dysarthria, dysphagia and arm weakness; (IIb)</td>
<td>Decrease on repetitive stimulation, response to iv edrophonium</td>
<td>8.0 nmol</td>
<td>Cholinesterase inhibitors: mild prednisone and plasma exchange: good</td>
<td>3 years, fluctuating</td>
</tr>
<tr>
<td>M, 65</td>
<td>Double aorto-coronary bypass</td>
<td>10 weeks</td>
<td>Double vision, dysphagia and dysarthria; (IIb)</td>
<td>Decrease on repetitive stimulation, response to im neostigmine and oral pyridostigmine</td>
<td>12.0 nmol</td>
<td>Cholinesterase inhibitors: mild prednisone: good</td>
<td>8 years, mild generalised</td>
</tr>
</tbody>
</table>
Chiari I malformation mimicking myasthenia gravis

Chiari I malformation is accompanied by a variety of symptoms and signs suggesting brain stem, cerebellar, or cervical spinal cord lesions. The most common symptoms include headache, neck pain, sensory loss, and ataxia.1 Dysphagia occurs in 5–15% of the patients and it may be the only presenting symptom.2 Progressive dysphagia caused by Chiari I malformation, mimicking myasthenia gravis, has been reported in this journal in 1996 and 2002.2,3 Dysphonia may occur rarely, but it has not been described as an early symptom.4 Pain and stiffness in the posterior neck is a common feature, but severe neck tenderness leading to dropped head syndrome has not so far been reported in Chiari I malformation.

Case report

A 13 year old girl was admitted to our department of neurology four weeks after adenoidectomy under general anaesthesia, because of progressive difficulty in lifting her chin off her chest, together with dysphagia and dysphonia. There was no pain or stiffness in the posterior neck. Computed tomography of the head was reported as normal. There was mild fluctuation of the dysphagia and dysphonia during the day, with worsening of the dysphonia after prolonged conversation or after reading in a loud voice. There was no sleep disturbance.

Neurological examination revealed an increase in deep tendon reflexes in all four limbs. Routine serum biochemistry and blood count were unremarkable.

On the basis of the history and clinical data, myasthenia gravis was suspected, so electrodiagnostic testing was undertaken to investigate the neuromuscular junction. Pencillan 3 Hz repetitive nerve stimulation of the right accessory nerve along with recordings obtained from surface electrodes over the trapezius muscle did not show significant variations in compound muscle action potential amplitude under baseline conditions, or when the test was repeated three minutes after maximal voluntary effort for 30 seconds. Single fibre electromyography of the right extensor digitorum communis muscle during voluntary activity failed to show any abnormal single fibre action potentials.

A response to anticholinesterase agents observed clinically or recorded electrically has been reported in a variety of disorders, including Eaton–Lambert syndrome, botulism, and transverse myelitis, and even in patients with intracranial mass lesions.5 The partial response to anticholinesterase drugs in our case reinforces the view that it is unwise to base the diagnosis of myasthenia gravis purely on a positive pharmacological test.

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Expanding cerebral cysts (lacunae): a treatable cause of progressive midbrain syndrome

A progressive motor deficit presenting in adulthood is an ominous sign, being often associated with either neoplasia or neurodegenerative diseases. Notable if very rare exceptions to this poor prognosis are cerebral expanding lacunae or, as they are sometimes called, benign intraparenchymal brain cysts.1 These are intraparenchymal cavities without an epithelial lining, filled with cerebrospinal fluid (CSF), located in the thalamo-mesencephalic arterial territory.1 They their expanding nature is demonstrated by their progressive clinical course and by the frequent complication of aqueduct stenosis and triventricular hydrocephalus.2

We present a case of progressive midbrain syndrome associated with expanding cysts, which was successfully treated by neuroendoscopy.

Case report

A 43 year old woman with an unremarkable clinical history presented in 1996 with progressive resting tremor and weakness of the left arm. The tremor persisted during posture maintenance and action. Within a year the motor problems extended to the left leg. Brain magnetic resonance imaging (MRI) showed large bilateral thalamic lacunae (10 to 20 mm) well defined lesions with signal intensities identical to CSF occupying most of the right thalamo-mesencephalic region. There was no contrast enhancement either in the lesions or in the surrounding tissue. The ventricular spaces were only mildly

Figure 1 Brain and cervical spine magnetic resonance imaging: herniation of the cerebellar tonsils through the foramen magnum.
enlarged. A search for cystic lesions elsewhere in the body was negative.

In the following year disturbance in posture and diplopia in the right lateral gaze became apparent. The tremor resolved spontaneously, but the left hemiparesis worsened. She was referred for neurological evaluation.

At admission, objective findings were a left hemiparesis (leg worse than arm), hemirigidity, severely reduced automatic movements and left bradykinesia, brisk tendon jerks on the left, diplopia on rightward gaze, and Pari- naud syndrome. The patient was alert and oriented, with intact gross cognition.

A preoperative brain MRI showed multiple large cystic lesions occupying the right pons-mesencephalic region and smaller lesions in the right thalamus (fig 1, panels A and B). CSF flow sequences revealed aqueductal stenosis and slight triventricular hydrocephalus.

Surgical procedure
The patient underwent a surgical endoscopic procedure. A flexible 2.5 mm neuroendoscope (Storz) was inserted through a burr hole, and the third ventricle was cannulated with a 3.9 peel away. The floor of the ventricle posterior to the mammillary bodies appeared severely deformed by a large cystic mass that did not allow access to the aqueduct. The cystic mass was coagulated and opened into the third ventricle. A fragment of the cyst wall was taken for pathology, which showed normal neuroglia with few amilaceaus bodies, no epithelial lining, and no signs of old or recent haemorrhage.

Once opened, the inside of the cyst revealed a multilobular structure. The flux of fluid towards cystic lesions occupying the right paramedian ponto-mesencephalic region and smaller lesions in the right thalamus (fig 1, panels C and D). CSF flow sequences revealed aqueductal stenosis and slight triventricular hydrocephalus.

Figure 1 Preoperative sagittal (A) and coronal (B) T1-weighted magnetic resonance (MR) images (gadolinium enhanced). Large multilobulated cystic lesions occupy the right ponto-mesencephalic region, squeezing the aqueduct and causing mild triventricular hydrocephalus. Postoperative sagittal (C) and coronal (D) T1-weighted MR images. There is a slight reduction in volume of the lesions, with partial resolution of the hydrocephalus.

Follow up
At three months the patient showed a remarkable improvement in motor performance but there was reappearance of a modest resting tremor in the left hand. MRI documented a mild reduction in cyst volume and moderate reduction in ventricular size (fig 1, panels C and D). At 18 months the patient was neurologically normal except for the mild resting tremor of the left hand. She had resumed all her premorbid activities, including dancing.

Comment
A progressive disorder of cognition and hydrocephalus caused by expanding cerebral lacunae in the thalamus and midbrain was first described in 1983.1 These lesions consist of multiple grape-like CSF-filled cavities, usually located bilaterally in the rostral brain stem. Their incidence is extremely rare (seven cases reported thus far).1,2 and apparently not related to any risk factor. Differential diagnosis includes parasitic and neoplastic cystic lesions. However, their pathogenesis remains obscure. Clinical presentation is characterised by signs of triventricular hydrocephalus from aqueduct obstruction, and by various extrapyramidal signs, ataxia, and abnormalities of oculomotion. In our patient the tremor disappeared when the motor defects worsened, and reappeared after their resolution. This observation could be explained by the progression of the cystic lesions towards the ventral thalamus causing the functional equivalent of a reversible thalamotomy, the “benefit” of which reversed upon decompression of the cysts.

The expanding nature of the lesions and the progressive clinical worsening justified surgical management. Treatment of the hydrocephalus (shunting, cisternotomy) has seldom been rewarding.3,4 Opening and draining the cysts, while carrying a higher morbidity risk, seems to give a better clinical outcome, although the cyst volume is not significantly modified by the procedures.3,5 In our case, the use of an endoscopic approach to both the hydrocephalus and the opening of the cysts minimised operative risks and led to an excellent clinical result.

The values of endoscopic neurosurgery in expanding cerebral lacunae has been emphasised by others.7 While the neuropathology of the lesions and their location supports the interpretation that the cystic spaces are dilated Virchow–Robin spaces, the precise mechanism leading to the dilatation remains unknown. The absence of vasculitis or systemic hypertension in all reported cases reinforces the hypothesis of a localised disturbance in vascular permeability and interstitial fluid reabsorption.3 In conclusion we draw attention to this very unusual neuropathological entity. An endoscopic microneurosurgical approach to this type of lesion has the advantage of a good risk to benefit ratio. As the term “lacuna” is usually associated with a small static vascular lesion, and the term “benign cyst” overlooks the expanding nature of the lesion, we suggest that these lesions should be called “benign expanding cerebral cysts.”

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Reversible callosal disconnection syndrome in internal hydrocephalus

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