LETTERS TO
THE EDITOR

Magnetic resonance imaging and vertebral artery dissection

Since the advent of advanced radiological modalities such as MRI and magnetic resonance angiography (MRA), dissections of cervical arteries are increasingly recognised as a common cause of stroke in young adults. Auer et al recently advocated MRA as the initial diagnostic tool for vertebral artery dissection. Conventional angiography might be avoided altogether in subjects with a suspicious history and MRA images suggestive of a dissection (double lumen or mural haematoma).

The sensitivity of MRA for the diagnosis of vertebral artery dissection was only 20% in one study, but the specificity was excellent (100%). The sensitivity was considerably better in the hands of Auer et al, but in this study the specificity (true negative rate in subjects free of disease) was not considered because all patients had vertebral artery dissection. The following case report illustrates that care must be taken to avoid false positive results when using MRA for the diagnosis of vertebral artery dissection.

A 47 year old male pilot suddenly experienced clumsiness and slight loss of strength in the right arm and leg during a long distance flight, while he stooped forward. During the following hours, he developed a global headache without irradiation to the neck, but the other symptoms gradually diminished. Prior history was unremarkable, except for a 3 hour period of horizontal diplopia which suddenly developed 3 months earlier. He had never smoked. Family history was negative for cardiovascular disorders. The patient later confessed that he had recently picked up the habit of gargling his throat with toothpaste twice a day, always with his neck in extreme retroflexion.

General physical examination (8 hours after onset of symptoms) was normal. Neurological examination showed minimal paresis and impaired dexterity of the right hand, mild circumduction of the right leg, and an insecure tandem gait. An MRI (including T1 weighted spin echo images with and without fat suppression, and proton density and T2 weighted fast spin echo sequences, performed on a 1.5 Tesla whole body MRI system) performed several hours later visualised both a fresh and an old right sided cerebellar infarct (figure A). In addition, MRI showed an irregular right vertebral artery in which a patent lumen was partially surrounded by a semilunar area of high signal intensity on T1 and T2 weighted images. On fat suppressed images, this area’s high signal intensity persisted, excluding the possibility that it originated from perivascular fat. This image was suggestive of mural haematoma due to vertebral dissection (figure B). Because we were reluctant to base any treatment decisions (antiocoagulants) merely on MRI findings, digital subtraction angiography was performed on the day of admission. This examination was normal (figure C). Shortly after this procedure, the patient developed vertigo and nystagmus which disappeared after 3 hours. Because we were puzzled by the discrepant findings on conventional angiography and MRI, we performed an MRA 4 days later. At this examination, the semilunar area of high signal intensity was found again (figure D), despite saturation of craniofugal and craniopetal flow respectively, which was applied to exclude the possibility that the high signal originated from flow in the periarterial venous plexus. Therefore, this examination was again suggestive of right vertebral artery dissection. An extensive search for other causes of stroke showed no abnormalities. Hence, due to the continuing discrepancy between conventional angiography and MRA, we had to reject this diagnosis in view of the normal conventional angiography, which remains the gold standard for diagnosing cervical artery dissection. In one series, conventional angiography was never falsely negative in patients with clinical signs or symptoms of vertebral artery dissection. The possibility that conventional angiography had nevertheless yielded a false negative result seems highly unlikely. In dissected arteries, MRI/MRA can detect intimal flaps, mural haematomas, or aneurysmal dilatations that are sometimes missed by conventional angiography, but even in such patients conventional angiography is never completely normal in the acute stage. Follow up examinations of patients with proven vertebral artery dissection indicate that the appearance of a dissected artery on conventional angiography can normalise in a substantial proportion of patients, but always after an interval of at least 1 to (usually) several weeks. Conventional angiography in our patient was performed on the day of admis-

(A) T2 weighted fast spin echo image showing high signal intensity in the right cerebellar hemisphere, indicative for a recent infarct. The older infarct cannot be seen on this section. (B) Axial T1 weighted fast spin echo image with fat saturation at the level of the base of the tongue, showing a semilunar area with high signal intensity around the flow void in the right vertebral artery. (C) Selective contrast injection in the right vertebral artery shows no abnormalities. The remainder of the intra-arterial angiography of the cervical and cranial arteries was also normal. (D) Axial three dimensional time of flight technique, acquired in the axial plane image at the same level showing high signal intensity at the same location as in B.
sion, directly after the “abnormal” MRI and four days prior to the “abnormal” MRA, hence spontaneous resolution of the dissection is very unlikely. Therefore, we consider our MRI/MRA examinations falsely positive, and we hypothesise that the area of semilunar high signal intensity originated from a perivascular venous plexus, in which we were unable to saturate inflow of blood completely, presumably due to extremely slow flow.

Our “pilot study” illustrates the specificity problems of MRI/MRA for the diagnosis of vertebral artery dissection. Two anatomical structures surrounding vertebral arteries contribute to these problems. The first structure is the venous plexus that surrounds vertebral arteries. This structure may have a semilunar appearance, and slow flow in its lumen may give rise to high signal intensity on both MRI and MRA, creating an image suggestive of dissection.\(^1\) It has been suggested that saturation slabs in conjunction with MRA completely suppress flow related high signal, thus distinguishing it from high signal from an intramural haematoma which cannot be suppressed by saturation slabs.\(^2\) The present case report illustrates that flow in this plexus cannot always be suppressed.

The second tissue that may falsely present as a dissection is fat that directly surrounds vertebral arteries, turbulence and magnetic susceptibility near sharp vessel turns can also cause false positive MRA results.\(^1\) In some patients, MRI cannot distinguish between intramural and intramural haematoma, leading to false conclusions.

Decisions based on false positive MRI/MRA results can be hazardous due to the sometimes severe side effects of anticoagulation that is recommended by some to prevent further ischaemic events. Another danger of a false positive diagnosis of vertebral dissection is that it may preclude the search for other causes of stroke that could be amenable to secondary prevention.

MRI/MRA remains important because it helps visualise ischaemic lesions and, in some patients, provides complementary morphological information to cerebral angiography.\(^3\) Furthermore, it is a non-invasive procedure, an important advantage over cerebral angiography which carries a morbidity and mortality risk. Our patient, who developed transient neurological deficits shortly after angiography, underscores this. Therefore, MRA can play a part in the diagnosis of vertebral artery dissection, provided that the pitfalls mentioned above are recognised to avoid false positive results. In case of doubt, cerebral angiography remains the gold standard for vertebral artery dissection.

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Letter to the Editor

Catatonia due to central pontine and extrapontine myelinolysis: case report

Central pontine and extrapontine myelinolysis (CPEM) are recognised complications of hyponaetraemia and its overly rapid correction.\(^1\) CPEM usually presents with spastic tetraparesis and pseudobulbar palsy.\(^2\) We describe a patient with CPEM in whom behavioural manifestations overshadowed corticospinal tract signs.

A 64 year old Chinese speaking woman with a history of episodic psychotic depression that had never required admission to hospital was admitted to a hospital because of vomiting and diarrhea. Her general and neurological examination were normal. On admission she had a sodium concentration of 105 meq/l. An infusion of 3% saline at a rate of 150 ml/hour was given during 6 hours. Ten hours later her sodium was 134 meq/l and she was mute and tetraparetic. She seemed catatonic with motor perseveration. Transfer to our hospital was requested.

On admission her vital signs were normal. She was mute without any spontaneous volitional movements except for visual pursuit. She was tetraparetic and hypertreflexic with increased tone and bilateral Babinski's signs. CPEM was suspected. Admission MRI, EEG, and spinal fluid examination were normal. Over the next 2 days the reflexes normalised and the Babinski's signs disappeared but she continued to have mild diffuse weakness. She had waxy flexibility and assumed bizarre non-physiological postures consistent with catatonia. Psychogenic unresponsiveness was suspected and she was started on risperidone and sertraline. There was no benefit. Electroconvulsive therapy was proposed by a psychiatry consultant but was refused by the patient's family. The clinical picture was dominated by an akinetic mutism with marked catatonia. Catatonia due to CPEM was considered. A repeat MRI 12 days after the onset of symptoms showed high intensity areas in the pons, caudate, and putamen consistent with CPEM (figure A, B). Physical and occupational therapy were instituted and she gradually recovered over the next 2 weeks. She was transferred to a rehabilitation hospital where she recovered completely and returned to live independently. She has been followed up at the neurology clinic and has not shown any residual deficits.

CPEM usually presents with tetraparesis and pseudobulbar palsy. Unusual clinical presentations include extrapyramidal syndromes, ataxia, and neurobehavioural syndromes. Although psychiatric manifestations of CPEM have been recognised they usually manifest as an agitated delirium, or a pseudobulbar state with pathological laughing and crying.\(^2\) When present, neuropsychiatric symptoms are usually overshadowed by florid signs of brainstorm and pyramidal tract dysfunction.\(^3\) Behavioural changes such as inappropriate affect, emotional lability, personality changes, paranoia, poor judgement, emotional incontinence, and disinhibition have been reported.\(^3\) Price and Mesulam described a case of pontine myelinolysis in which transient pyramidal signs were followed by confusion, restless behaviour, pressured tangential speech, and disinhibition.\(^1\) Our patient also had transient long tract signs but they were followed by a catatonic state. The extensive extrapontine myelinolysis present in our patient may explain the behavioural symptoms we encountered.

CPEM may present with unusual behavioural symptoms. At the onset of neurological deterioration MRI may be normal but subsequent imaging studies usually disclose the lesions. CPEM presenting with neuropsychiatric symptoms in patients with normal initial imaging studies might suggest a psychogenic aetiology. Corticospinal tract signs may be temporary. A strong index of suspicion for CPEM is required when patients with recent

(A) Axial T2 weighted image showing prominent high signal intensity within the pons suggestive of central pontine myelinolysis. (B) Axial T2 image showing symmetric bilateral areas of high signal in the caudate and putamen suggestive of extrapontine myelinolysis.
hypotension present with behavioural changes. Kinetic mutism and catatonia may be the dominant clinical features in CPEM.

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Association between butyrylcholinesterase K variant and the Alzheimer type neuropathological changes in apolipoprotein E ε4 carriers older than 75 years

Apolipoprotein E (ApoE) ε4 has a strong influence on the development of sporadic Alzheimer’s disease in many ethnic populations. However, ApoE ε4 is neither necessary nor sufficient for the development of Alzheimer’s disease, suggesting that other genes increase the risk of Alzheimer’s disease. One such new candidate is the butyrylcholinesterase (BChE) gene (BChE). 1 BChE is associated with senile plaques (SPs) and neurofibrillary tangles (NFTs). Lemaître et al recently reported that the K variant of BChE (BChE-K) was associated with the development of Alzheimer’s disease, especially in ApoE ε4 carriers older than 75 years. 2 A possible mechanism as to how BChE-K is related to Alzheimer’s disease under the influence of ApoE ε4 is the acceleration of Alzheimer type neuropathological changes. If BChE-K has an effect on the development of Alzheimer’s disease in ApoE ε4 carriers, the formation of Alzheimer type neuropathological changes may be accelerated by BChE-K in the ApoE ε4 carriers.

We have examined genotypes of BChE and ApoE, and densities of the senile plaques (SPs), with dystrophic neurites (NPs), and neurofibrillary tangles (NFTs) in the brains from 51 patients with Alzheimer’s disease and 90 non-demented subjects from a poor-tem series of Japanese. Clinical and postmortem diagnosis of Alzheimer’s disease was carried out as described previously. 3 The densities of Alzheimer type neuropathological changes were quantified by averaging the counts of those in the hippocampus and superior temporal gyrus. Genotypes of BChE and ApoE in all patients were determined as described elsewhere. 1 Genotypic and allelic distributions of BChE were analysed by χ2 test. The densities of the SPs, NPs, and NFTs, and ages at onset and durations of illness were compared among BChE genotypes with the Kruskal-Wallis test or Mann-Whitney U test in total subjects, those with Alzheimer’s disease, and non-demented subjects. We also examined these relations in the subgroups divided by the ApoE ε4 status or the age of 75 years. Statistical significance was defined as two tailed probabilities of <0.05.

There were no significant differences in the frequency of BChE-K genotypes or alleles between patients with Alzheimer’s disease (0.16 in allele frequency) and non-demented subjects (0.18), and in the total subjects, ApoE ε4 carriers or non-ApoE ε4 carriers, although a strong association of ApoE ε4 alleles with Alzheimer’s disease was found in this population (p=0.004). Genetic association of BChE-K genotypes with sporadic Alzheimer’s disease was non-significant in all subjects older than 75 years, the ApoE ε4 carriers older than 75 years, and non-ApoE ε4 carriers older than 75 years. There was no significant association of BChE-K with the densities of the SPs, NPs, or NFTs in the hippocampus and superior temporal gyrus in the total subjects, in the Alzheimer’s disease or non-demented groups, or with ages at onset or duration of illness in Alzheimer’s disease. However, when we divided total subjects into two subgroups with different ApoE ε4 status, there was significant association between BChE-K and the densities of the SPs and NFTs in the superior temporal gyrus (STG) in the ApoE ε4 carriers (SPs, p=0.04; NPs, p=0.03; data not shown). Further, we analysed the correlation between BChE-K and the densities of the SPs, NPs, and NFTs in the hippocampus and superior temporal gyrus in the ApoE ε4 carriers older than 75 years and non-ApoE ε4 carriers older than 75 years. There was a significant genetic association of BChE-K with the densities of the SPs, NPs, and NFTs in the STG in the ApoE ε4 carriers older than 75 years. There was a decrease of severity of Alzheimer type neuropathological changes with BChE-K in our study was not expected because Lemaître et al showed an increase in frequency of the BChE-K allele in Alzheimer’s disease. Single et al also reported that BChE-K was not associated with the densities of the SPs and NFTs, even in the ApoE ε4 carriers. In addition, BChE-K was not related to the development of Alzheimer’s disease in the ApoE ε4 carriers in our study. Rusu et al also showed a lack of association between BChE-K and the development of Alzheimer’s disease. However, Hiltunen et al showed that BChE-K had a protective effect on the development of Alzheimer’s disease in ApoE ε4 carriers younger than 75 years. The effects of BChE-K on the Alzheimer type neuropathological changes or development of Alzheimer’s disease are different among studies, suggesting that the significant genetic association in the studies by Lemaître et al, Hiltunen et al, and ourselves might be linkage disequilibrium with relevant variability in BChE or other adlacent gene on chromosome 3, and that BChE-K does not play a direct part in the pathogenesis of Alzheimer’s disease.

<table>
<thead>
<tr>
<th>BCHE genotype</th>
<th>ApoE ε4 carriers over 75 years (n=28)</th>
<th>non-ApoE ε4 carriers over 75 years (n=95)</th>
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<tr>
<td></td>
<td>K/K (n=8)</td>
<td>N/N (n=20)</td>
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<tr>
<td>Hippocampus:</td>
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<tr>
<td>SPs</td>
<td>3.0 (0.0, 17.2)</td>
<td>12.2 (4.5, 28.7)</td>
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<tr>
<td>NPs</td>
<td>0.7 (0.0, 11.9)</td>
<td>11.0 (5.9, 25.3)</td>
</tr>
<tr>
<td>NFTs</td>
<td>1.1 (0.4, 23.1)</td>
<td>17.4 (2.5, 59.6)</td>
</tr>
<tr>
<td>Superior temporal gyrus:</td>
<td></td>
<td></td>
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<tr>
<td>SPs</td>
<td>0.2 (0.0, 22.8)</td>
<td>47.9 (12.1, 83.8)</td>
</tr>
<tr>
<td>NPs</td>
<td>0.2 (0.0, 8.8)</td>
<td>10.7 (3.6, 19.0)</td>
</tr>
<tr>
<td>NFTs</td>
<td>0.0 (0.0, 0.2)</td>
<td>0.9 (0.4, 4.9)</td>
</tr>
</tbody>
</table>

Values are medians (25th percentile, 75th percentile). The density represents the average counts in 2.56 mm2 for the SPs and NPs, and in 0.64 mm2 for the NFTs. BChE=butyrylcholinesterase gene; ApoE=apolipoprotein E; K=the K variant allele of butyrylcholinesterase gene; N=the normal allele of butyrylcholinesterase gene; SPs=senile plaques; NPs=senile plaques with dystrophic neurites; NFTs=neurofibrillary tangles.
The patient was a 71 year old, retired physician with a 3 to 4 year history of memory impairment. Neuropsychological evaluation disclosed a high average to superior general intellectual functioning, with mild impairment in naming to confrontation and episodic memory for verbal memory. His visuospatial ability remained relatively unimpaired and was rated as average for his age. His comprehension for verbal and written instructions was intact. At the present time he is still well oriented to time and place, and is somewhat independent in activities of daily living. He is, remarkably, not depressed, but does, repeatedly, raise concern regarding whether he has become to his wife. Moreover, mild hypoprophusia in the frontal temporal lobes bilaterally was seen on SPECT investigation and no evidence of pathognomonic laboratory results were found. Taken together, the pattern of episodic memory and naming impairments and functional imaging findings was thought to be consistent with the early stages of the Alzheimer's type (DAT) in keeping with National Institute of Neurological and Communicative Disorders Association-Alzheimer's Disease and Related Disorders Association criteria. The patient was consequently referred to our department for “prospective memory book training” and follow up assessments to index progression of disease.

During our sessions his wife had stated that the patient could still remain intact. At the time of the interview he complained that her once “blamboyant” and “unblushing” husband could no longer “put any feeling into his lines” when they read play scripts together. He thought that he had “lost his enthusiasm to act” and that he had “lost his ability to “act.”” The patient’s use of prosody during speech conversation. When asked to use prosody to command when reading script, however, this once gallant actor spoke without melody, loudness, stress, or accent, with inappropriate pauses. To quantify this patient’s peculiar deficit, the patient was asked to read and repeat words and sentences to prosodic command and imitation. Observation revealed five single words and one sentence that the patient often and spontaneously uttered with normal prosody, such as “Honey, PLEASE(!).” These 10 items were used to assess the patient’s ability to produce prosody to command and imitation (Table). For example, the patient was told to read the words "HoneyPLEASE!!" with loudness, stress, accentuation of the word “please,” and as if he really meant it. If he failed (the words were read without the acoustical features expected), the patient was asked to imitate the experimenter’s reading of the word(s) or sentence which incorporated the appropriate prosodic elements only after he was asked to describe the affective prosodic quality of the phrase to ensure good comprehension. Five age matched normal healthy controls volunteered to read the items found in the table, and in each case, read spontaneously the word or phrase with appropriate and expected prosody.

The patient was unable to read any words or sentences with normal (appropriate and expected) prosody. Indeed, the patient had lost his ability to “act.” The patient’s use of prosody did, however, improve dramatically with imitation. That is, he was able to repeat eight of the ten items in the table with appropriate and expected prosody. Interestingly, the single item that he continued to have difficulty producing was item three (You know...). There was a time when...). He was able to read the word "Y ou know..." with a pause after the word "you know" suggesting that this patient seems to also have features characteristic of motor aprosodia. Hence, because he had available the knowledge to successfully select and use appropriate prosody, but failed to produce prosodic speech to command, dysfunction of the praxis production system is implicated rather than a conceptual system. Thus, motor aprosodic prosody can be defined as an inability to produce prosody to command during speech. The precise underlying mechanism(s) responsible for producing this deficit is unknown, although Heilman et al. and Tuckers et al. have hypothesised that the right hemisphere is indeed dominant for organising the affective-prosodic components of language and gestural behaviour and that the functional organisation was geared to normal affective language in the right hemisphere was analogous to the organisation of propositional language in the left (non-dominant) hemisphere. Conceptually, the evidence of poor affective prosody to command yet normal spontaneous affective prosody, and good affective prosodic repetition and comprehension would suggest a “transcortical aprosodia.” Note however, that the patient’s spontaneous prosody was unaffected whereas spontaneous speech is affected by a transcortical motor aphasia. Hence, we might place the critical lesion for prosodic apraxia in the right dorsolateral frontal lobe, extending into the deep frontal

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**Ten item prosodic apraxia scale**

<table>
<thead>
<tr>
<th>Item</th>
<th>Type of emphasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Honey PLEASE!</td>
<td>Accented PLEASE!</td>
</tr>
<tr>
<td>2. Are you hungry?</td>
<td>Rise in pitch</td>
</tr>
<tr>
<td>3. You know...there was a time when I could recite all the streets in my neighborhood</td>
<td>Pause after “You know”</td>
</tr>
<tr>
<td>4. Holy COW!</td>
<td>With surprise</td>
</tr>
<tr>
<td>5. YUP, yup, yup, yup, yup...</td>
<td>As if you were distressed with decending intonation and stress</td>
</tr>
<tr>
<td>6. La de da...</td>
<td>With melody</td>
</tr>
<tr>
<td>7. O Canada, our home and native land...</td>
<td>With proper tempo, as if you were singing</td>
</tr>
<tr>
<td>8. SHIT!</td>
<td>As if you were frustrated and upset</td>
</tr>
<tr>
<td>9. Thank you</td>
<td>As if you sincerely meant it</td>
</tr>
<tr>
<td>10. May I go to the bathroom, I really need to go quite badly...</td>
<td>As if you really meant it, accentuating the word “REALLY”</td>
</tr>
</tbody>
</table>

Directions: Read the above word(s) and sentences as if you really mean them. Pretend you are auditioning for a play and you are required to read the lines with the type of emphasis noted beside each line.

*These items were selected based on observation of the patients spontaneous speech. Therefore, they are qualitatively constructed and should not be used as a general measure of prosodic apraxia with all patients.
white matter, in keeping with typical dominant hemispheric lesions producing transcortical motor aphasia. This speculation is supported by the patient’s SPEECD findings of mild hypopufferation in the frontotemporal lobes bilaterally.

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Vocal cord abductor paralysis in spinocerebellar ataxia type I

Vocal cord abductor paralysis (VCAP) is considered a sign of a poor prognosis in neu- rodegenerative diseases, because severe laryngeal dysfunction by VCAP may result in acute airway obstruction and require emergen- cy tracheotomy.1 Although VCAP is a cardinal feature in multiple system atrophy (MSA), it has not been reported in several types of spinocerebellar ataxia with dominant inheritance. We evaluated the movements of the vocal cords of seven patients with SCA1 by laryngofibroscopy.

Seven unrelated patients with SCA1 who had the expanded CAG repeat of ataxin-1 were investigated. There were two men and five women ranging in age from 27 to 67 years old (mean 44.5 years). Spouses and other family members, in addition to the patients, were questioned about events of stridor, dysphagia, and respiratory symptoms. Vocal cord movement was examined by laryngofibroscopy and recorded during inspiration and phonation.

The rating scale used to evaluate maximal abduction of the vocal cords during larygofibroscopy was as follows: (-)=normal; (+)=median position; (++)=paramidline position; (+++)=midline position. For the evaluation of VCAP, we tried the respiratory flow volume loop study as well as VCAP in one patient (patient 2) in whom maximal abduction of the vocal cords was slightly limited (+) on laryngofibroscopy.

The correlations between VCAP and CAG repeat length duration of illness were ana- lyzed with the non-parametric Mann-Whitney U test.

The core clinical features, including the vocal cord findings, are summarised in the table. VCAP was present in five of the seven patients with SCA1. Although it is difficult to know when the VCAP first became manifest in each patient, patient 1 showed VCAP con- firmed by laryngofibroscopy only 2 years after the onset of gait disturbance.

All five patients with VCAP showed mild dysphagia requiring no tube feeding, and four patients had a history of stridor at night. Patient 1 showed VCAP accompanying dysphagia without stridor at night even in an early stage of the disease. The VCAP was found to be severe on laryngofibroscopy in all three patients with breathing difficulty on inspiration. Patient 5, who had the severest VCAP, developed stridor during wakefulness as well. In patients 4 and 5, the breathing dif- ficulty on inspiration was improved by tracheostomy. The respiratory flow volume loop study did not detect abnormality in patient 2.

The CAG repeat number tended to be higher in the patients with VCAP than in the patients without VCAP (p=0.05), but the duration of illness was significantly correlated with the presence of VCAP (p=0.43).

This is the first report that VCAP is often found in patients with SCA1. As VCAP may not usually be a late feature in patients with SCA1, evaluation of VCAP is necessary even in early stages of the disease. It is not surpris- ing to find VCAP in patients with stridor, because stridor is usually caused by airway obstruction of the larynx. However, VCAP was detected by laryngofibroscopy in a patient without stridor who had dysphagia. Furthermore, all patients with VCAP exhib- ited dysphagia. We therefore think that laryn- gofibroscopy should be performed in SCA1 patients with dysphagia as well as stridor.

The mechanism of VCAP may be divided into some types, the paralytic type, the non- paralytic type, and these two combined type.1 The first is possible caused by loss of neurons in the nucleus ambiguous.1,2 The second is considered to be due to over- activity of the intrinsic laryngeal muscles. Stridor due to paralysis has been found to be more prominent in sleep than during wake- fulness; whereas stridor by non-paralytic dysfunction has been found both during the daytime and during sleep.3 We suspect that the VCAP in patients with SCA1 may be dominantly paralytic, because the nucleus ambiguous is sometimes pathologically in- volved in SCA1 and because stridor in our patients with SCA1 was more marked in sleep.

Our laryngofibroscopic findings suggested that severe VCAP caused breathing difficulty on inspiration in the patients with SCA1 by obstructing the airway. Moreover, the stridor during wakefulness as well as sleep indicated it to be very serious. The important question concerns when tracheostomy should be carried out after the diagnosis of VCAP to prevent respiratory abnormalities leading to sudden death. Although we consider tracheo- stomy at the stage when breathing difficulty on inspiration or stridor during wakefulness is noted, it awaits further study with a large number of patients to decide which stage is best for tracheostomy.

Furthermore, we now consider endoscopic cord lateralisation as another possible man- ageement for VCAP.


<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Sex</th>
<th>Onset</th>
<th>CAG repeat</th>
<th>Duration of Illness</th>
<th>Stridor at Night</th>
<th>Stridor during Wakefulness</th>
<th>Dysphagia</th>
<th>Breathing Difficulty</th>
<th>Tracheotomy</th>
<th>(CAG)n of Mutant Allele</th>
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<tbody>
<tr>
<td>1</td>
<td>30M</td>
<td>28</td>
<td>2</td>
<td>(+)</td>
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<td>2</td>
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*(-)=normal; (+)=median position; (++)=paramidline position; (+++)=midline position.
Lateral gaze synkinesis on downward saccade attempts with paramedian thalamic and midbrain infarct

The symptoms of paramedian thalamic and midbrain infarct include ocular motor disturbances mainly in the vertical plane. We here describe a patient with the additional feature of an unusual horizontal eye movement synkinesis.

A 60 year old overweight man, with diabetes and mild hypertension, suddenly fell into a coma that lasted for 4 hours and was followed by slight right hemiparesis, recent memory impairment, hypersomnia, and vertical gaze impairment.

On admission to our centre, about 10 days after symptom onset, the patient still presented fluctuating drowsiness from which he was unable to move his eyes downward. By contrast, the leftward gaze deviation elicited by the attempt to make a downward saccade is at variance with all previous descriptions.

The triggering of a saccade requires not only the activation of the exitory burst neurons (EBNs), but also the deactivation of the omnipause neurons (OPNs), which provide tonic inhibition of both horizontal and vertical EBNs.

Accordingly, any attempt to activate a lesioned riMLF should be associated with maximal OPN inhibition. However, OPNs discharge for saccade in any direction and are not strictly direction selective, as shown by horizontal oscillations during vertical saccades detectable in normal subjects. These oscillations suggest that during vertical saccades the inhibition of OPNs disinhibits both vertical and, to a lesser extent, horizontal EBNs.

In our patient, the horizontal gaze deviation was always directed to the left rather than in both directions as during oscillations. Many ocular motor structures, including those located in the midbrain, trigger a purely vertical (downward) saccade only when stimulated bilaterally, so as to nullify horizontal components with different direction depending on the stimulation side. This probably occurs for the riMLF too, as it shows ipsilateral projections to the abducens nucleus. In our patient, the projections to the left nucleus were probably spared by the fact that the lesion predominantly affected the right side.

Overall, our patient’s horizontal ocular motor synkinesis is unusual, and probably derives from a strong inhibition of OPNs, which in turn frees the horizontal EBNs, and from an unbalanced activation of the left abducens neurons via riMLF projections spared from the lesion, although it is not possible to exclude the possibility that the unbalanced activation of abducens neurons originated from frontal or parietal cortical areas or from the superior colliculus rather than from riMLF projections.

This hypothesis is strengthened by the reinforcement of the leftward eye deviation when the examiner kept the patient’s lids lifted. Since this manoeuvre prevents lid synkinesis, it results in what resembles an attempted forced lid closure which, on the basis of blink induced eye oscillations, is likely to be an additional stimulus for OPN inhibition. Moreover, although they occur in various conditions, saccade oscillations during fixation are in keeping with a reduction of OPN inhibition level.

In conclusion, our patient presented an ocular motor synkinesis that should be listed among those occurring in thalamomesencephalic infarcts. This sign is unusual and it is likely to be overlooked, but it is fully explicable both by neurophysiology and

(A) and (B) show a thalamomesencephalic ischaemic lesion, hyperintense in T2 weighted scans (SE, TR=2300 ms; TE=25 ms). The lesion involves the anteromedial portion of both thalami, but the right one to a larger extent. In the midbrain, the lesion is located around the Sylvian aqueduct, and symmetrically, but prevalently right sided, and involves the area that is located posteromedially with respect to both red nuclei. (C) Recording of the horizontal (upper tracing) and of the vertical (bottom tracing) movement recorded respectively from the left and right eye with the infrared reflection technique (Skalar, IRIS system) during an attempted downward saccade. The vertical tracing is flat, as the patient was unable to move his eyes downward. By contrast, the horizontal tracing shows a concomitant leftrightward saccade. At onset, both tracings show a blink artifact.
by anatomical connections of the saccade system.

Systemic sclerosis (scleroderma) is a multi-system connective tissue disease of unknown aetiology, characterised by progressive fibrosis of the skin and internal organs including the lungs and gastrointestinal tract. Pathological calcification of soft tissues (known as calcinosi) is a common feature in the CREST syndrome of scleroderma (calcinosis, Raynaud’s phenomenon, oesophageal dysmotility, sclerodactyly). By contrast, internal organ calcification is rare, and isolated cases of spinal calcinosi and calcific constrictive pericarditis have been reported. We report here the cases of two patients with systemic sclerosis whose CT examination disclosed extensive brain calcifications.

Case 1, a 48 year old man was referred to hospital because of polyarthralgia involving the wrists and ankles, Raynaud’s phenomenon. Mini mental state examination score was 22/30. The patient was oriented to place, but not to time. Anterograde amnesia was noted. Agnosia, apraxia, and aphaia were absent. There was no muscle weakness and manual tone was normal, tendon reflexes and plantar responses were both flexor. There was no sensory loss or impairment of cranial nerves. Systemic sclerosis signs were unchanged. Routine hematological tests were normal. Results of blood chemical tests were also unremarkable (serum electrolytes, urea, creatinine, iron), including phosphorus and calcium metabolism (serum parathyroid hormone concentration, blood calcium and phosphorus, 25-hydroxyvitamin D, 1,25-dihydroxyvitamin D, calcium, and phosphorus). Serum concentrations of free triodothyronine, free thyroxine, and thyroid stimulating hormone were normal. Serological tests for syphilis, HIV-1, A, and Lyme disease were negative. Brain CT showed bilateral extensive calcification in the dentate nuclei (figure 1A), basal ganglia, and subcortical white matter (figure 1B). On MRI T1 weighted images and T2 weighted images, calcification was visible as a low intensity signal on T1 weighted images and as a high intensity signal on T2 weighted images. The patient was given fluoxetine (20 mg/day) and bremazepam (6 mg/day). At follow up, 1 year later, the patient’s clinical status was unchanged, as was brain CT.

Case 2, a 64 year old right handed woman was admitted to hospital for evaluation of a Raynaud’s phenomenon which had lasted for more than 10 years. At physical examination, sclerodactyly and tightness of the facial skin
were noted. Telangiectasias were present on the face, hands, and palate. The patient complained of pyrosis. Oesophageal manometry showed abnormalities of oesophageal motility. Hand radiography disclosed soft tissue calcifications. Anticentromere antibodies were positive at a 1/1000 dilution. A CREST syndrome was diagnosed and the patient was given butylated hydroxytoluene (600 mg/day) and prednisone (25 mg/day).

One year later she was admitted for the evaluation of recent ischaemic attacks (TIAs). During the previous week she had experienced three bouts of expressive aphasia and right hemiplegia, each lasting about 10 minutes. She never smoked and did not have diabetes, hypertension, or dyslipidaemia. The neurological examination was normal. Routine blood chemical tests were normal (serum electrolytes, urea, creatinine) including phosphorus and calcium metabolism (serum parathyroid hormone concentration, blood calcium and phosphorus, 25-hydroxyvitamin D, 1,25-dihydroxyvitamin D, calciumuria, and phosphaturia). Cranial CT showed bilateral calcifications of the basal ganglia, and faint calcifications of the dentate nuclei (1A and basal ganglia (1B)).

Brain CT of case 1 shows dense symmetric calcifications in the dentate nuclei (1A) and basal ganglia (1B).

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BOOK REVIEWS


If it moves - measure it. Such is the trend in psychiatry and this has led to a proliferation of assessment scales of variable utility; from the esoteric to the ubiquitous. This book has them all, touching on virtually every scale, and one of the most useful volumes I have seen for a very long time. It covers everything from the AMTS through to the MMSE to the Kew cognitive test. Each scale is presented in full together with a short commentary, critical references and, usefully, an estimated time taken to perform the test together with an address to contact the original author.

The scales presented are divided into those covering depression, neuropsychiatric assessments, activities of daily living, global assessments, visible assessments, delirium, caregiver assessments and scales for memory function. This organisation, together with a useful and functional index, will make the task of selecting an appropriate scale much easier in the future. Some of these scales are covered by patent law and my only quibble is that it would have been useful to know which instruments can be reproduced and for what purpose without fear of being billed for the privilege. Maybe highlighting such information would give unpatented authors a stimulus to use their lawyer and therefore increase this deplorable practice.

This aside, I cannot recommend this book highly enough. By contrast with books that ever indicated, and what treatments are clinically effective. My only question is “Why didn’t somebody do this before”.

SIMON LOVESTONE


Professor Franco Postacchini is an orthopaedic surgeon at the University “La Sapienza” in Rome and is a well known widely respected surgical spine. He is to be congratulated on the production of this book which is wide ranging, comprehensive, and beautifully illustrated. The management of lumbar disc disease is fraught with uncertainty and there are many diagnostic and therapeutic pitfalls. The author has succeeded in addressing most of these controversies in a clear and logical fashion. He occasionally blurs the distinction between theory and established practice such as when he Forget the practice of medicine and in particular the management of spinal disorders is full of paradoxes. For example, he states that large extruded disc fragments are unlikely to resolve spontaneously and usually require surgical treatment. This seems a logical proposition but my experience is that many of these large protrusions undergo complete clinical and radiological resolution within 2 or 3 months. Paradoxically, it is often the smaller contained disc prolapses which fail to improve with conservative measures. Like many orthopaedic surgeons he is persuaded by the alluring theories of discogenic low back pain and worships at the altar of segmental microinstability. However, I agree with much that he has written and differences of emphasis are inevitable in a field that is strong on dogma and short of established truths.

I would have no hesitation in recommending the text to treatment as the book is very readable and makes a good introduction to the management of lumbar disc disease. Nearly all aspects of diagnosis and treatment are covered but I was disappointed with the chapter discussing results of surgery. There is no mention of the use of objective validated disability and quality of life instruments in the assessment of outcome. For a text that aims to be comprehensive this constitutes a serious omission. It is because practitioners have failed to use objective outcome measures to establish the natural history of lumbar disc disease and the effects of therapeutic interventions that there remains so much uncertainty about management. These uncertainties cover (among others) physiotherapy, manipulation, timing of radiology, timing of surgery, whether spinal fusion is ever indicated, and what treatments are clinically and cost effective. Despite these drawbacks, surgeons who manage lumbar disc disorders will want to have a copy of this book, either on their own or their departmental library's shelf.

RODNEY LAING
disappointing and somewhat introspective that they may think that subjects such as neurology, microbiology, and immunology may justify equally expert contributors.

Is it an easy read? There is no easy way to write on a subject such as this, bedevilled by lack of objective facts and the writers have chosen a discursive, debating style which when not tightly controlled can slip towards verbosity. This, however, is not a standard medical text book and it would be unfair to make direct comparisons.

Is this a useful book? As a source of references, yes. As a guide to clinicians I am less convinced. The two commonest questions repeatedly mentioned: “How long does the illness last?” and “What are my chances of recovery?” Cognitive behavioural therapy is concisely and usefully summarised. There is a single page on complementary therapy which again is often an area of considerable interest to patients notwithstanding the lack of controlled evidence for or against it.

On that note it is perhaps appropriate to quote one very intelligent patient with chronic fatigue: “You know what I mean. I saw him during his PhD. “I’ve done a lot of reading and internet searching about the causes and possible cures of this, before I came to see you” he said “It seems to me that most people go down this road for a bit like that when you commonly see after glandular fever and nobody seems to think it odd that after glandular fever you can feel unwell for quite a long time” he continued “If most people get better from this” (and many do) and if you try all sorts of other treatments like homeopathy, acupuncture, meditation then the one you were doing when you got better will be the one you call “cured you”. He had of course discovered the maxim of entertaining the patient while nature gets them better. One could do worse perhaps than keep patients with chronic fatigue syndrome occupied, if not necessarily always entertained for quite a while, by recommending this book for them to read. They might end up with a greater understanding of fatigue and they would certainly learn a little really is known and how the search for an instant cure (which drives many of them) is futile.

ANDREW LEVER


New information about how and why migraine happens continues to break on us in a dizzying succession of waves coming from various journals in different disciplines. We need an accessible, understandable, and unitary vehicle to collect, organise, and present this information. Journals, the Internet, and the abstracting services have their place, but for this purpose nothing beats the book. How well does Lars Edvinsson’s Migraine and Headache Pathophysiology meet this need?

This book has several attractive features. Recognising that some of the world’s best science is now being done in the laboratories of industry, it has enlisted as authors several leading researchers from the major pharmaceutical manufacturing companies, in addition to “the usual suspects” from academia. Not only has it introduced some exciting writers to the “review book” audience, but it has provided a particular insight into the science of determining why drugs work, or don’t work, in migraine.

This book is up to date, containing many 1998 references. Another strong point of the book is its comprehensiveness; though only 184 pages long, it covers every major aspect of the pathophysiology of migraine. There are chapters on cranial blood vessels, receptor physiology, neurotransmitters, 5-hydroxytryptamine receptor subtypes, cortical spreading, depression, neurogenic inflammation, systemic shunts, cerebral haemodynamics, and animal modelling. This is achieved at the expense of some pretty terse prose at times, which can make it difficult for the non-expert to follow. It should be noted that, the title notwithstanding, this book deals almost exclusively with the pathophysiology of migraine, and the reader who buys it to get some insight into the mechanisms of tension-type headaches, or other kinds of headache, is going to be disappointed.

Who should read this book? Certainly the migraine researcher should. Though much of it will be familiar to those who have kept up with the literature, it is nevertheless an attractive and handsome volume of current research information. Moreover, the first chapter (by Lars Edvinsson) and the last chapter (by Peter Goadsby) are very pretty syntheses of the field. What about clinicians? Some of it is heavy going for people like me, who are not basic scientists. But I got through it all in about 6 hours, and found I knew more about migraine coming out than I did going in—which makes reading it a very worthwhile exercise.

JOHN EDMEADS


There is something about the anachronistic binding of the Handbook of Clinical Neurology series that is really reassuring. Surely if classic phenomenological neurology is to be found anywhere, it will be between these fake leather embossed covers. This volume, the second of three on the neurology of systemic diseases, does not disappoint. Here, in 450 pages of close type and few illustrations, are covered the neurology of orthopaedic, endocrine, gastrointestinal, genitourinary, and psychiatric disorders. Aminoff and Aminoff, the volume editors, have assembled an authoritative panel of authors that equitably straddle the Atlantic. There are detailed reviews of familiar territory such as diabetes, orthopaedic trauma, thyroid diseases, and porphyria. Cole’s historical survey of B12 deficiency is particularly fine. In addition there are excellent chapters on more arcane topics for instance the neurology of pancreatic transplantation and intestinal pseudo-obstruction. Perhaps the movement disorders associated with coccal disease could have been mentioned and a chapter on the neurology of inflammatory bowel disorders is certainly lacking. But these are trifling complaints against a text that, with its twin volumes, is significantly more comprehensive than any other account of the neurology of systemic diseases.

It is hard to imagine a practicing neurologist requiring (or easily affording) a personal copy of all three volumes, but the local medical library should certainly buy them; both neurologists and general physicians will work the better for having them close to hand.

This is one of a new type of medical textbook written to meet the needs of an increasingly informed patient population. Aimed very much at those with multiple sclerosis, their families, and care-workers, it is simple and clearly written with jargon and technical terms kept to a minimum but without patronising. Chronic diseases, and especially multiple sclerosis, are not always well managed by the physician. Too many of us think that there is no cure and feel helpless in a busy clinic faced with the patient with a long list of complaints. There are too few specialty multiple sclerosis clinics in which neurologists, pain specialists, uroneurologists, physiotherapists etc liaise.

Patients often feel left in the dark, unaware which of their symptoms can be attributed to their multiple sclerosis and whether it is “worth bothering a busy doctor”. Many can cite bad experiences in their past when they have been fobbed off with well meaning reassurance but without practical help. Dysesthesia, sexual problems, and urinary incontinence are only a few of the symptoms that can bring misery to the lives of patients and their families and which are poorly addressed by doctors. This book, in a language accessible to most (and with a glossary to explain some unavoidable jargon), explains multiple sclerosis, its symptoms, and what might realistically be obtained in terms of symptom control. All aspects are covered and nothing considered too trivial; constipation or cold feet might be extremely trying for an individual patient and each is considered.

The old idea that it doesn’t help a patient to know too much about his disease (“it will only make him introspective and hypochondrial”) is outdated. Multiple sclerosis can hit anyone and patients now want, and deserve, to be informed. While doctors find it challenging to be faced with a patient equipped with the latest information down loaded from the internet or well informed having read a book such as this, this is a challenge to which we must be ready to rise. This textbook provides the information patients want and fills the gap left by busy doctors. It should be marketed appropriately and we must be ready to respond to the reaction of patients. Perhaps someone with multiple sclerosis should have been invited to write this review.

GILLIAN HALL

CORRECTION


During printing, the figure in this paper (p 164) was made darker than the original. The correct version appears below.

(A) T1 weighted MRI of the lesion. (B) Anatomical scheme of the centre of the lesion, corresponding to the leftmost image of the bottom row of the MRI. The right side of the figures corresponds to the left side of the brain. GP=globus pallidus; Cd=caudate nucleus; Acb=nucleus accumbens; CI=capsula interna; DB=diagonal band.

GILLIAN HALL
Ideomotor prosodic apraxia

KONSTANTINE K ZAKZANIS

J Neurol Neurosurg Psychiatry 1999 67: 694-695
doi: 10.1136/jnnp.67.5.694

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