"Dropped head syndrome" in syringomyelia: report of two cases

"Dropped head syndrome" is characterised by weakness of the extensor muscles of the neck, with or without involvement of the neck flexors, and is commonly caused by a variety of neuromuscular disorders, including myasthenia gravis, polymyositis, amyotrophic lateral sclerosis, facio–scapulo–humeral dystrophy, nemaline myopathy, carnitine deficiency, spinal muscular atrophy, and isolated neck extensor myopathy. There are isolated reports of dropped head syndrome occurring in cervical spondylitis and ankylosing spondylitis. In this report, we describe the clinical and imaging findings of two patients who had dropped head syndrome as a rare neurological sign secondary to syringomyelia.

Case report 1

A 46 year old right handed man presented during May 2002 with insidious onset, gradually progressive weakness and wasting of the small muscles of the left hand since August 1998. This deficit stabilised after a period of one year. At the onset of illness, he had also developed mild difficulty in using his right hand for performing fine work but this symptom remained stable. He noticed occasional fasciculations over the arms for one year. After two years and nine months he developed rapidly progressive head drop and required assistance of the hand to maintain the head in an erect posture, and he also had mild difficulty in lifting his head off the bed while rising from the supine position. There was no history of impaired sensation to touch, pain, or temperature, or inadvertent burns over the hands or shoulders. There were no features to suggest Horner’s syndrome or symptoms referring to the cranial nerves, lower limbs, cerebellar system, or sphincters. There was no nuchal pain, restricted neck movements, or symptoms of raised intracranial pressure.

Examination revealed mild cerebellar dysarthria and fasciculations over the tongue. There were no features of Horner’s syndrome and the eye movements, facial sensations, palatal reflexes, and pharyngeal reflexes were normal. Occasional fasciculations were present over the arms. There was Medical Research Council grade 2 weakness of the neck extensor muscles with head drop and grade 4 weakness of the neck flexors. There was mild wasting of the erector spinae muscles in the neck and asymmetric wasting of the small muscles of the hand, including the thenar, hypothenar, and interossei muscles bilaterally, with the left side being affected more severely. Power in the lower limbs was normal. Deep tendon reflexes were sluggish in the upper limbs and exaggerated in the lower limbs, with a bilateral flexor plantar response. The sensory system revealed bilateral C2 hypesthesia, with impaired pain and temperature sensations in both upper limbs and in the shoulder girdle region. Magnetic resonance imaging of the spine and brain revealed a septate intrinsic cord hypointensity in T1 weighted images, becoming hypointense in T2 weighted images extending from the C2 to C7 (fig 1) levels, associated with low lying cerebellar tonsils and brainstem reaching the lower border of C1. Brain images revealed evidence of hydrocephalus. There was no evidence of myelomeningocele. Electromyography of the distal muscles in the upper limbs showed fibrillation and fasciculation potentials. Motor and sensory conduction studies were normal. Investigations including serum chemistry and haemogram were normal.

Case report 2

A 30 year old man presented in December 2003 with progressive weakness and atrophy of the right shoulder girdle muscles with a duration of one year, followed by similar symptoms on the left side for 10 months; at presentation he was unable to raise his arms. In the last six months he had developed head drop, with difficulty in maintaining the head in the erect posture, and since this time he had noticed weakness and atrophy of the hand muscles. There were no symptoms of pain or restricted neck movements, and there was no sensory impairment, sphincter disturbance, or cerebellar ataxia. Neurological deficits were present in the form of thoracic kyphoscoliosis and wasting of the extensor muscles of the neck, resulting in severe weakness and dropped head (fig 2). Power in the neck flexors was grade 4. Severe asymmetric wasting of the shoulder girdle muscles, in addition to moderate wasting of the arm, forearm, and hand muscles was noted, with bilateral claving. There was grade 2 power in the proximal muscles, including the arm, and grade 3 in the small muscles. Deep tendon reflexes were absent in the upper limbs and exaggerated in the lower limbs, with a bilateral extensor plantar response. Temperature sensation was impaired in the entire body except for the face. Posterior column and spinothalamic sensations were normal. Contrast magnetic resonance imaging of the spine, including T1 weighted and T2 weighted images, revealed a cerebrospinal fluid signal intensity lesion involving the entire cord, with expansion of the cord, and no evidence of abnormal contrast enhancement, suggestive of holo-cord syringomyelia.

Discussion

Dropped head syndrome is a well known feature in a variety of neuromuscular disorders, and is also described in diseases of the bone and joints, such as ankylosing spondylitis and cervical spondylitis. Neck flexion weakness is typical in most conditions, but prominent neck extensor weakness has been described in several reports. In the recent review article on dropped head syndrome in amyotrophic lateral sclerosis, 19 different conditions are listed as causes for dropped head syndrome, but syringomyelia is not mentioned as a cause. Our patients presented with classic features of cervical intrinsic cord lesion suggestive of...
syringomyelia and had the typical features of dropped head syndrome.

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The patient gave consent for reproduction of the photograph.

References

Vitamin B-12, serum folate, and cognitive change between 11 and 79 years

A recent Cochrane review reported that although vitamin B-12 deficiency is known to be associated with cognitive impairment in old age, benefits of supplementation on mental ability are unclear.1 The situation is similar to that for blood pressure, in which hypertension is associated with neuropsychological deficits in adulthood but cognitive outcomes of lowering blood pressure in randomised controlled trials are equivocal.2

Lower mid-life blood pressures.

Mean (SD) serum vitamin B-12 was 390 (161) ng/l (n = 422) and mean serum folate was 337 (155) ng/l (n = 391). Pearson correlation coefficients with age 11 IQ were r = 0.04 (p = 0.42) for B-12 and r = 0.13 (p = 0.010) for folate; and with age 79 IQ, r = 0.12 (p = 0.018) for B-12 and r = 0.12 (p = 0.016) for folate. Linear regression of age 79 IQ controlling for age 11 IQ confirmed a significant effect on age 79 IQ for B-12 (β = 0.092, p = 0.016, R² improvement = 0.008) but not for folate (β = 0.038, p = 0.33). Only two participants had folate levels below the normal range (<5 μg/l) and omitting these did not affect correlation coefficients with age 11 and 79 IQ scores. Twenty five participants had vitamin B-12 levels below the normal range (<200 ng/l) and there was a stronger correlation with age 79 IQ in this group (r(25) = 0.57, p<0.001) than in those well within the normal range >250 ng/l (r = 0.10, p = 0.031). The difference between these two correlation coefficients was significant (p = 0.016). After adjusting for age 11 IQ and sex, a significant association was observed to be associated with lifetime change in IQ (sex, APOE 64 status, cigarette smoking, statin use, and number of drugs prescribed),3 vitamin B-12 remained a significant contributor (β = 0.099, p = 0.011). Together, these variables explained 4.5% of total variance in age 79 IQ scores after adjusting for IQ at age 11. The number of units of alcohol consumed per week was also positively correlated with age 79 IQ score (Spearman r = 0.10, p = 0.026), but was no longer significantly associated (β = 0.01, p = 0.73) once age 11 IQ and vitamin B-12 were adjusted for.

COMMENT
Both vitamin B-12 and folate correlate with IQ in old age in a non-demented population. Lower serum B-12 at age 79 is associated with cognitive decline between age 11 and age 79. By contrast, serum folate at age 79 correlates with age 11 IQ, and controlling for this reduces the correlation with IQ in old age to almost zero. Hence, in this sample the relation between serum folate and old age mental ability can be fully explained by its correlation with IQ scores on the same test given 68 years previously. This is a similar situation to that with blood pressure.4 This further emphasises the importance of interpreting associations between cognition and other variables in older people in the context of “pre-morbid” mental ability. The effect size of vitamin B-12 is smaller than those found with more domain specific cognitive tests,5 contributing less than 1% of total variance in age 79 IQ. It is unlikely to be clinically apparent. However, the effect was more significant in the small subsample with laboratory defined deficiency.

Inspection of the relation between vitamin B-12 and the standardised residual score of age 79 IQ regressed on age 11 IQ (fig 1) suggests that the overall correlation is accounted for by a subgroup that is cognitively vulnerable to vitamin B-12 deficiency. Moreover, this vulnerability occurs at levels within the normal laboratory range. An IQ decline of >1 SD occurred in seven of 18 participants (39%) with a serum vitamin B-12 of <200 ng/l, 19 of 84 (23%) with levels of 200–299 ng/l, 21 of 103 (20%) with levels of 300–399 ng/l, and 21 of 149 (14%) with levels of >399 ng/l. Further work is required to confirm this and ascertain what makes these individuals vulnerable. One explanation is that this is a group having “metabolically significant” vitamin B-12 deficiency within the normal range with raised homocysteine or methylmalonic acid levels.6 This may help target B-12 therapy, but at present identifying this group remains challenging. Our data suggest that in a non-demented, relatively healthy population, serum folate concentrations were not related to IQ in old age after controlling for childhood mental ability. However, only two participants in our sample had folate values below the normal range, so other studies are needed to assess the cognitive effects of folate deficiency and treatment with folic acid.

ACKNOWLEDGEMENTS
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Figure 1 Vitamin B-12 levels and standardised residual scores of age 79 IQ regressed on age 11 IQ in a surviving cohort of the Scottish Mental Survey 1932 (n = 422).

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Stent assisted endovascular thrombolysis of internal carotid artery dissection

Spontaneous dissection of the extracranial internal carotid artery (ICA) is a major cause of stroke with severe residual handicap in young adults. Recently, stent supported angioplasty has been used to treat intimal dissection in case of neurological symptoms while on anticoagulation or as an alternative to the traditionally accepted use of anticoagulation. We report a case of internal carotid artery dissection causing hemiplegia successfully treated with emergent endovascular stenting followed by intra-arterial thrombolysis.

Case report

A 44 year old right handed man was admitted to the emergency room after an acute episode of left side weakness which resolved within three hours. The patient had been well until the onset of symptoms. There was no history of trauma, strenuous exercise, hypertension, or other medical problems. In the week prior to admission, he reported intermittent headaches and right side neck pain after four days of diving. A cranial computed tomography (CT) scan was normal. Diffusion weighted imaging (DWI) of the brain performed six hours after the onset of symptoms while the patient was asymptomatic showed no evidence of infarction. T2-weighted magnetic resonance images and fat suppressed images showed a subarachnoid hypersignal of a mural haematoma in the infrapetrous segment of the right ICA suggestive of a dissection. Low molecular weight heparin treatment (enoxaparin sodium 1 mg/kg subcutaneously every 12 hours) was started to obtain true anticoagulation.

Three hours later, the patient had sudden left sided hemiplegia, hypaesthesia, hemianopia, and hemineglect. The National Institutes of Health Stroke Scale (NIHSS) score was 12. Transcranial Doppler ultrasound demonstrated prior to stenting using transcranial Doppler ultrasound. Another alternative to intra-arterial thrombolysis could be the use of mechanical thrombectomy devices to remove the clot from the dissected siphon and MCA.

This case report is to our knowledge the first example of the potential use of stenting followed by intra-arterial thrombolysis to treat and cure symptomatic carotid artery dissection. However, no conclusions can be drawn about the safety of endovascular approach in this clinical setting. Further evaluation is needed to address its risk–benefit ratio.

Discussion

Ischaemic stroke in patients with ICA dissection mainly results from thromboembolic, or, less frequently, haemodynamic mechanisms. Formation of a false channel in the vessel wall or endothelial damage may favour formation of a local thrombus, which becomes less adherent and prone to embolise distally. Although no general agreement exists on the best management of extracranial carotid artery dissection, and because of the threat of an embolic complication, anticoagulation with heparin followed by oral warfarin is used in most institutions. In case of an embolic complication in a patient with known carotid artery dissection revealed by local signs or a transient ischaemic attack, as in our patient, no recommendation exists about emergent therapy. Some authors suggest that intravenous thrombolysis might be safe and effective when given within three hours of onset of stroke without worsening the arterial wall tearing. However, in case of severe stenosis, near occlusion, or even occlusion of the ICA, regardless of the cause, intravenous or intra-arterial thrombolysis has had a poor recanalisation rate, less than 15%.

Endovascular stenting in patients with carotid artery dissection has been successfully used in either selected cases with haemodynamically significant stenosis or when anticoagulation failed to prevent embolic stroke. It permitted resolution of the stenosis and the intraprocedural complications and the patient's symptoms improved gradually after the procedure. On day 7, he was discharged on aspirin and clopidogrel with no residual symptoms.

Intravenous thrombolysis given within three hours of onset of symptoms might be an alternative treatment, but it would have likely not been effective since ICA near occlusion would have been the thrombotic or systemic thrombolysis was contraindicated because of the anticoagulation therapy.

Although so far no clinical trial has documented the efficacy of emergent revascularisation in the setting of acute stroke, accumulated anecdotal data show that endovascular mechanical recanalisation is likely to become an important alternative therapeutic approach in properly selected stroke patients. A potential disadvantage of mechanical reopening is the production of embolic debris. With stent deploying in dissected carotid artery, there is theoretically a risk of the intramural clot contained within the dissected segment breaking into the cerebral circulation leading to embolisation distally. In our case, the MCA emboli were demonstrated prior to stenting using transcranial Doppler ultrasound.

Intra-arterial tissue plasminogen activator (tPA) was infused directly into the thrombus four hours after the onset of the new symptoms (40 mg total). There was good proximal recanalisation with residual filling defects in some branches of the MCA (fig 1C) and immediate clinical improvement. The patient received a 5000 U heparin bolus during the procedure and then continuous 500 U/h infusion for 24 hours. Heparin therapy was then replaced by clopidogrel and aspirin, both 75 mg daily. On day 2, MRI showed brain infarct in the deep MCA territories with asymmetric haemorrhagic transformation. The carotid artery and the MCA were fully patent at ultrasound examination with no evidence of restenosis. There were no periprocedural complications and the patient's symptoms improved gradually after the procedure. On day 7, he was discharged on aspirin and clopidogrel with no residual symptoms.

Intra-arterial mechanical revascularisation is a therapeutic approach in properly selected cases of acute stroke managed to minimise the risk of haemorrhagic complications. By taking advantage of the immediate recanalisation of completely or almost occluded vessels it permits delivery of thrombolytic agents directly in the clot, maximising the chance of total distal recanalisation.
**Bell’s palsy: a study of the treatment advice given by Neurologists**

Bell’s palsy is defined as an isolated unilateral lower motor neuron facial weakness of no obvious cause. The incidence has been estimated at around 23 to 25 cases per 100,000 population annually. Although the prognosis is generally good, around 16% are left with varying degrees of permanent disability.1

The use of steroids and acyclovir in the treatment of Bell’s palsy has been addressed in two recent Cochrane reviews.2 3 These found no benefit from either but concluded that available studies were insufficiently powered to detect a treatment effect. Neurologists are often asked by primary care physicians for treatment advice and in view of this uncertainty we were interested in studying the recommendations given. A questionnaire (appendix) was emailed to all consultant neurologists (n = 35) and specialist registrars (n = 21) in Scotland. Responses were collated at six weeks following an interim reminder. Fisher’s exact test was used to compare groups; odds ratios with 95% confidence intervals and significance were calculated (table 1).

Replies were received from 27 consultants and 17 registrars, response rates of 77% and 81%, respectively. In all, there had been 69 requests received for treatment advice in the preceding three months. Referral for guidance from neurologists amounted to 26% of the total number of cases predicted by incidence studies.1

Only 5% of neurologists said they would always see the patient, with further 29% if atypical features were present. The use of steroids depended strongly on the stage of presentation, 76% giving steroids within 24 hours of onset, 62% within three days, and only 28% up to seven days. Fewer gave steroids in certain subcategories (12% in pregnancy, 19% in Ramsay Hunt syndrome, 62% in a complete syndrome, and 45% in a partial syndrome).

The steroid regimen advised was variable, with most advocating 40 to 60 mg of prednisolone, with or without a tapering dose. Only 20% of neurologists gave acyclovir in every instance; a further 20% gave it if there was evidence of Ramsay Hunt syndrome.

On the whole the responses from consultants and registrars were similar. However, while both advised steroids early on, consultants still recommended steroids up to seven days (42%) compared with only 6% of specialist registrars (p = 0.009). Geographical variability was evident; Glasgow neurologists advised steroids more readily, with 95%, 74%, and 21% giving them at 24 hours, three days, and seven days, respectively. This compared with 42% (p = 0.002), 42% (p = 0.085), and 17% (p = 0.34) at Edinburgh. There was also a trend for Glasgow physicians to prescribe more acyclovir (21% v 11% (p = 0.37)).

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**Table 1. Treatment of Bell’s palsy by subcategory**

<table>
<thead>
<tr>
<th>Subcategory</th>
<th>“Yes” responses by consultants</th>
<th>“Yes” responses by registrars</th>
<th>“No” responses by consultants</th>
<th>“No” responses by registrars</th>
<th>OR (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do you see the patient?</td>
<td>1</td>
<td>1</td>
<td>14</td>
<td>9</td>
<td>0.642 (0.355 to 1.163)</td>
<td>0.5</td>
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<tr>
<td>Siroesteroids within 24 h?</td>
<td>20</td>
<td>12</td>
<td>6</td>
<td>5</td>
<td>1.389 (0.347 to 5.55)</td>
<td>0.25</td>
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<tr>
<td>Siroesteroids within 3 d?</td>
<td>17</td>
<td>9</td>
<td>9</td>
<td>8</td>
<td>1.68 (0.482 to 5.85)</td>
<td>0.18</td>
</tr>
<tr>
<td>Siroesteroids within 7 d?</td>
<td>11</td>
<td>1</td>
<td>15</td>
<td>16</td>
<td>11.7 (1.35 to 102)</td>
<td>0.009</td>
</tr>
<tr>
<td>Siroesteroids in pregnancy?</td>
<td>2</td>
<td>3</td>
<td>19</td>
<td>13</td>
<td>0.456 (0.067 to 3.12)</td>
<td>0.27</td>
</tr>
<tr>
<td>Siroesteroids in Ramsay Hunt?</td>
<td>5</td>
<td>5</td>
<td>20</td>
<td>11</td>
<td>0.55 (0.13 to 2.32)</td>
<td>0.21</td>
</tr>
<tr>
<td>Siroesteroids in complete syndrome?</td>
<td>18</td>
<td>10</td>
<td>8</td>
<td>7</td>
<td>1.58 (0.44 to 5.64)</td>
<td>0.20</td>
</tr>
<tr>
<td>Siroesteroids in partial syndrome?</td>
<td>11</td>
<td>7</td>
<td>14</td>
<td>8</td>
<td>0.9 (0.25 to 3.25)</td>
<td>0.25</td>
</tr>
<tr>
<td>Do you give acyclovir?</td>
<td>6</td>
<td>3</td>
<td>14</td>
<td>9</td>
<td>1.29 (0.255 to 6.49)</td>
<td>0.30</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Subcategory</th>
<th>“Yes” responses by Edinburgh neurologists</th>
<th>“Yes” responses by Glasgow neurologists</th>
<th>“No” responses by Edinburgh neurologists</th>
<th>“No” responses by Glasgow neurologists</th>
<th>OR (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do you see the patient?</td>
<td>2</td>
<td>0</td>
<td>5</td>
<td>14</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Siroesteroids within 24 h?</td>
<td>5</td>
<td>18</td>
<td>7</td>
<td>1</td>
<td>0.4 (0.004 to 0.403)</td>
<td>0.002</td>
</tr>
<tr>
<td>Siroesteroids within 3 d?</td>
<td>2</td>
<td>14</td>
<td>7</td>
<td>5</td>
<td>0.255 (0.05 to 1.19)</td>
<td>0.07</td>
</tr>
<tr>
<td>Siroesteroids within 7 d?</td>
<td>2</td>
<td>10</td>
<td>15</td>
<td>0.75 (0.11 to 4.91)</td>
<td>0.35</td>
<td></td>
</tr>
<tr>
<td>Siroesteroids in pregnancy?</td>
<td>1</td>
<td>3</td>
<td>9</td>
<td>14</td>
<td>0.52 (0.05 to 5.8)</td>
<td>0.39</td>
</tr>
<tr>
<td>Siroesteroids in Ramsay Hunt?</td>
<td>2</td>
<td>5</td>
<td>12</td>
<td>0.48 (0.08 to 3.03)</td>
<td>0.26</td>
<td></td>
</tr>
<tr>
<td>Siroesteroids in complete syndrome?</td>
<td>3</td>
<td>14</td>
<td>9</td>
<td>0.12 (0.02 to 0.63)</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Siroesteroids in partial syndrome?</td>
<td>3</td>
<td>9</td>
<td>8</td>
<td>0.38 (0.08 to 1.89)</td>
<td>0.15</td>
<td></td>
</tr>
<tr>
<td>Do you give acyclovir?</td>
<td>1</td>
<td>3</td>
<td>8</td>
<td>11</td>
<td>0.46 (0.04 to 5.3)</td>
<td>0.37</td>
</tr>
</tbody>
</table>

CI, confidence interval; OR, odds ratio.
COMMENT
Although many neurologists advise steroids and some would recommend acyclovir, the uncertainty regarding the treatment of Bell’s palsy is reflected in our questionnaire responses. The majority of responders indicated that their advice was not based on local guidelines and many commented on the lack of evidence. Some felt it was imperative to discuss the uncertainty with the patient; others that better randomised controlled trials are needed.
A new Scotland based randomised controlled trial will start later this year (Morrison J, personal communication). This study, coordinated by primary care physicians and ear, nose and throat surgeons, will compare four treatment arms (comprising steroids, acyclovir, placebo) within 48 to 72 hours of onset. Assessment of treatment effect will include photographs and questionnaires about objective and subjective outcomes. The aim is to recruit up to 720 patients, of whom 480 will have begun treatment within 48 hours of onset. Given the annual incidence of Bell’s palsy in Scotland, the researchers estimate that this will take up to 18 months. Uncertainty in managing this condition can only be resolved by well conducted randomised controlled trials.

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APPENDIX
Bell’s palsy questionnaire
1. How frequently in the last 3 months have you received a query from a GP about treatment of a Bell’s palsy? Yes/No
2. Do you arrange to see the patient before giving advice? Yes/No
If you are satisfied with the clinical diagnosis:
3. Would you advise steroids
   • within 24 hours? Yes/No
   • within 3 days? Yes/No
   • within 7 days? Yes/No
   • in pregnancy? Yes/No
   • with Ramsay Hunt syndrome? Yes/No
   • with complete facial palsy (loss of taste/hyperacusis)? Yes/No
   • with partial facial palsy? Yes/No
4. If steroids are advised, what regime would you suggest?
5. Is the advice you give on steroids based on local guidelines? Yes/No
6. Would you advise acyclovir? Yes/No
   If yes:
7. Is the advice you give on acyclovir based on local guidelines? Yes/No
8. Any additional comments:

References

Ondine’s curse during pregnancy
We report a case of a 34 year old right handed woman seen at 29 weeks’ gestation who suffered from apnoea of unknown aetiology. This pregnancy, as well as her first gestation, was complicated by generalised oedema and high blood pressure. Starting at week 25, her husband noticed she had developed intermittent brief periods of apnoea only while sleeping, which lasted as long as one minute but of variable duration. Her husband awakened her each time she had a protracted episode of apnoea. She was asymptomatic while awake. In the 29th week (that she suffered a more severe apnoea. She was intubated in the field and taken to the hospital for an emergency caesarean section. There was no spontaneous labour. Inability to breathe spontaneously persisted for two weeks postpartum and a neurological consultation was requested.
On initial evaluation blood pressure was 140/90 and the heart rate was 90 beats/min. The neurological examination revealed upbeat nystagmus of small amplitude in the primary position which did not change with upward or downward gaze. She had lack of spontaneous breathing. She was fully awake and cooperative, sitting up in bed with no assistance. While intubated she had an obvious cough reflex but the gag reflex was not formally tested. Tongue examination showed normal movement and power with no evidence of atrophy or fasciculation. Otherwise, cranial nerve and sensorimotor examinations were entirely normal. There was normal tone, with downgoing plantar reflexes and no evidence of other pyramidal findings. There was no record of arrhythmia. No yawning, vomiting, or hiccups were present during the examination and they were not seen by nursing staff.
It was of interest that this patient had suffered from apnoea presenting immediately after her first vaginal delivery two years previously. This was treated with intubation and resolved spontaneously with successful extubation approximately four hours later. Magnetic resonance imaging of the brain, brain stem, and cervical spinal cord was carried out, and showed a Chiari malformation with tonsillar herniation at C2 level and a cervical syrinx.

Figure 1 Magnetic resonance imaging of the brain (sagittal section) showing a Chiari malformation with tonsillar herniation at C2 level and a cervical syrinx.

Gender influence on the progression of HTLV-I associated myelopathy/tropical spastic paraparesis
HTLV-I associated myelopathy/tropical spastic paraparesis (HAM/TSP) is a chronic and disabling disease caused by the human T-lymphotrophic virus type I. Onset of the disease is insidious and the disease usually progresses slowly over years. However, there have been reports of the rapid evolution of HAM/TSP over months or even weeks.
basis for these different progression patterns is poorly understood and only a few studies have dealt with this matter. The present study aimed at evaluating a Brazilian HAM/TSP population for possible factors implicated in the progression of the disease.

**Methods**

We reviewed the files of 338 HTLV-I infected patients evaluated at the outpatient clinic of the Reference Centers for Neurological Infections and HTLV-I, Evandro Chagas Clinical Research Institute (IPEC), FIOCRUZ, Rio de Janeiro, Brazil. Patients were included in the study if they fulfilled the World Health Organization criteria for HAM/TSP, but were excluded if they had concurrent infections or other disabling diseases that could interfere with clinical progression. The eligible patients were submitted to a clinical history questionnaire and physical examination between June 2002 and February 2003. Clinical severity was evaluated using the IPEC disability scale (table 1), which was developed exclusively for the prospective assessment of HAM/TSP. We evaluated clinical progression using a disease progression index (DPI) defined as the IPEC disability final score divided by the duration of the disease, from onset of symptoms, in years. We used this value to divide our sample into quartiles. Patients whose values were under the 25th percentile were called slow progressors and patients whose values were above the 75th percentile were called fast progressors. Both groups were compared for their demographic and clinical characteristics using Fisher’s test. DPI values were compared using the Mann-Whitney U test. All p values were two sided and an α = 0.05 was employed.

**Results**

A total of 250 individuals were excluded due to lack of neurological disease or the presence of concurrent infections. The mean age of the remaining 88 individuals was 53.1 years, and there were more women (68.2%) than men. The mean age at onset was 40.7 years and the mean duration of disease was 12.5 years. Comparison between the fast (n = 22) and the slow (n = 22) progression groups showed a significantly higher prevalence of women in the former group (p = 0.02). Statistical analysis of other variables failed to show significant differences.

To evaluate the possible role of sex hormones in this difference, because the mean age of menopause is around 50 years, we compared men and women according to age at onset of the disease (early onset: <50 years; late onset: ≥50 years). The mean DPI of women and men were, respectively, 1.79 and 1.17 (p = 0.009) in the early onset group (n = 66; 66% women) and 2.59 and 1.78 (p = 0.731) in the late onset group (n = 22; 68% women), suggesting that women have a faster progression than men if the disease starts before the menopause.

**Discussion**

This is the first study to suggest that HAM/TSP progresses faster in women than in men. This difference seems to be particularly important in women whose disease started before the menopause. Although gender differences regarding the clinical evolution of HAM/TSP have not been reported before, there has been evidence of a disproportional number of women with this disease. Firstly, there is a worldwide female to male preponderance of HAM/TSP patients ranging from 1.5:1 to 3:5:1. Secondly, Nagai et al. analysed the proviral load of 202 HAM/TSP patients and 243 asymptomatic HTLV-I carriers and found a significantly higher proviral load in female patients when compared to males. It is well known that a higher proviral load is associated with the development of clinical disease.

The reason for the gender difference in our study is unknown, but it is possible that sex hormones play a role in the evolution of the disease. To test this hypothesis, we compared the DPI of patients whose disease had started before and after the age of 50, the mean age of the menopause. The finding of a significantly worse evolution in the female group with early onset of disease, coupled with no significant gender difference being observed in the late onset group, suggests that female hormones may be implicated in HAM/TSP pathogenesis and that their presence at higher levels may be associated with a faster clinical progression. Further support for this idea is provided by the beneficial effects of danazol, an androgenic drug, in the treatment of some HAM/TSP cases.

In summary, we found evidence of worse clinical progression in women with HAM/TSP compared to men. We hypothesise that sex hormones may account for this difference. If confirmed by further studies, this information may lead to a better understanding of the mechanisms involved in HAM/TSP pathogenesis and suggest different treatment options.

**Table 1** The IPEC disability scale

<table>
<thead>
<tr>
<th>Motor score: Gait</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Normal</td>
</tr>
<tr>
<td>1. Abnormal but can walk independently</td>
</tr>
<tr>
<td>2. Abnormal and dependent on eventual unilateral support</td>
</tr>
<tr>
<td>3. Abnormal and dependent on permanent unilateral support</td>
</tr>
<tr>
<td>4. Abnormal and dependent on eventual bilateral support</td>
</tr>
<tr>
<td>5. Abnormal and dependent on permanent bilateral support</td>
</tr>
<tr>
<td>6. Abnormal, dependent on permanent bilateral support, and occasional use of a wheelchair (WC)</td>
</tr>
<tr>
<td>7. Permanent use of a WC, stands up, and remains upright without support</td>
</tr>
<tr>
<td>8. Permanent use of a WC, uses arms to stand up, and remains upright without support</td>
</tr>
<tr>
<td>9. Permanent use of a WC, needs assistance from others to stand up and remain upright with support</td>
</tr>
<tr>
<td>10. Permanent use of a WC, unable to stand up, exhibits voluntary movements of the lower limbs when seated</td>
</tr>
<tr>
<td>11. Permanent use of WC, unable to stand up, and does not have any voluntary movements of the lower limbs</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Motor score: Running</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Runs</td>
</tr>
<tr>
<td>1. Unable to run</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Motor score: Climbing stairs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Climbs</td>
</tr>
<tr>
<td>1. Climbs only when holding the handrail</td>
</tr>
<tr>
<td>2. Unable to climb</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Motor score: Jumping</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Jumps on two feet</td>
</tr>
<tr>
<td>1. Jumps on two feet, but not with only one</td>
</tr>
<tr>
<td>2. Jumps on two feet only with hand support</td>
</tr>
<tr>
<td>3. Unable to jump</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Spasticity score: Clonus</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Absent</td>
</tr>
<tr>
<td>1. Only induced by the examiner</td>
</tr>
<tr>
<td>2. Spontaneous</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Spasticity score: Flexor/extensor spasms</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Absent</td>
</tr>
<tr>
<td>1. Present</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sensory score: Paresthesias</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Absent</td>
</tr>
<tr>
<td>1. Present, eventually</td>
</tr>
<tr>
<td>2. Present, permanently</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sensory score: Lumbar and/or lower limb pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Absent</td>
</tr>
<tr>
<td>1. Present, eventually</td>
</tr>
<tr>
<td>2. Present, permanently</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sphincter score: Bowel control</th>
</tr>
</thead>
<tbody>
<tr>
<td>0. Normal</td>
</tr>
<tr>
<td>1. Constipation</td>
</tr>
<tr>
<td>2. Incontinence or total retention, needs manual extraction or enemas</td>
</tr>
</tbody>
</table>

| Total: 0–29 |

---

**References**

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Plasma VEGF as a marker for the diagnosis and treatment of vasculitic neuropathy

Vasculitic neuropathy is treatable with immunotherapy. However, historical evidence of vasculitis is not always obtained from nerve and muscle biopsies. In particular, in cases of non-systemic vasculitic neuropathy showing no or minimum abnormal findings in serological tests, negative biopsy results cause considerable difficulty in the diagnosis.1 Vascular endothelial growth factor (VEGF) is a potent, multifactorial cytokine.2 VEGF is derived from endothelial cells and pericytes in response to hypoxia, and induces angiogenesis and microvascular hyperpermeability through its binding to VEGF receptors.2 Vascular endothelial growth factor (VEGF) levels were found to be raised in dermato mesiostis with peripheral neuropathy.3 These findings suggest that VEGF levels may be increased in patients with vasculitic neuropathy. Although an increase in plasma or serum VEGF concentrations has been reported, we could not find evidence of raised VEGF levels in patients with systemic vasculitis. There have been no studies to evaluate plasma VEGF in a series of patients with vasculitic neuropathy. With respect to VEGF levels in neuropathies, a marked increase in serum levels was reported in the Crow-Fukase (POEMS) syndrome and one with non-systemic vasculitic neuropathy. None of the patients were on drug treatment at the time of sampling. After disease remission was achieved by treatment with corticosteroids or other immunosuppressants, we analysed plasma VEGF again in three of the patients, including two with polyarteritis nodosa and one with SJögren syndrome. As a control group, we used plasma from 18 age-matched healthy volunteers, eight patients with Guillain–Barre syndrome, five with chronic inflammatory demyelinating polyradiculoneuropathy (CIDP), and seven with amyotrophic lateral sclerosis, after obtaining informed consent. Patients with diabetes mellitus or cancer were not included in the study.

Venous blood was sampled into an EDTA tube with minimal stasis. The sample was centrifuged and the plasma VEGF concentration was determined by a quantitative sandwich enzyme immunoassay technique using a Quantikine kit (R&D Systems, Minneapolis, Minnesota, USA). As VEGF is secreted by platelets in the clotting process, we measured plasma samples, not sera, to evaluate the circulating VEGF level precisely.

Differences between the groups were tested by the Kruskal–Wallis test and the Mann–Whitney U test. Differences were considered significant when the probability (p) value was <0.05. Significance tests for group differences were computed with StatView v5.0 (SAS Institute, Cary, North Carolina, USA).

The mean (SD) plasma VEGF concentrations in patients with vasculitic neuropathy (303 (182) pg/ml) were significantly higher than in the healthy controls (30.9 (31.7) pg/ml) (p<0.01) as well as in patients with Guillain–Barre syndrome (85.7 (57.3) pg/ml) (p<0.05), CIDP (49.9 (48.3) pg/ml) (p<0.05), and amyotrophic lateral sclerosis (88.1 (55.7) pg/ml) (p<0.05) (fig 1). There was no statistical difference in plasma VEGF concentrations between healthy controls and patients with CIDP, Guillain–Barre syndrome, or amyotrophic lateral sclerosis. The plasma VEGF concentrations in patients with vasculitic neuropathy before treatment (423 (97.1) pg/ml) decreased significantly after successful treatment with corticosteroids or other immunosuppressants, to 150 (114) pg/ml (p<0.05). One case with polyarteritis nodosa and the patients with vasculitic neuropathy associated with SJögren syndrome had a marked decrease in plasma VEGF after treatment (from 461 to 91.3 pg/ml and from 496 to 77.8 pg/ml, respectively). In the other patient with polyarteritis nodosa, the plasma VEGF levels decreased mildly, from 313 to 281 pg/ml.

**COMMENT**

Our results indicated that increased plasma VEGF could be a useful marker for the diagnosis of vasculitic neuropathy and for monitoring a therapeutic effect. This is the first report to show a significant increase in plasma VEGF levels in patients with vasculitic neuropathy compared with other neuropathies. As our patients with vasculitic neuropathy did not have cancer or diabetes mellitus, and as the plasma VEGF concentrations were significantly decreased after treatment, we consider that VEGF would be secreted into blood by the vasculitic lesions in this condition. We could find no significant increase in plasma VEGF levels in CIDP, Guillain–Barre syndrome, or amyotrophic lateral sclerosis. Vasculitic neuropathy may present with clinical manifestations similar to CIDP or other peripheral neuropathies.1 The increase in plasma VEGF could be a helpful marker to distinguish vasculitic neuropathy from CIDP and other peripheral neuropathies in such patients.

Although our results indicate the potential value of plasma VEGF as a marker in the diagnosis and treatment of vasculitic neuropathy, the significance of the results is limited by the relatively small number of patients. Further studies with a larger study population are necessary to confirm our results.

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doi: 10.1136/jnnp.2004.047571
Competing interests: none declared

**References**


Acute aspiration pneumonia due to bulbar palsy: an initial manifestation of posterior fossa convexity meningioma

False localising signs of intracranial lesions are defined as signs not generally associated with disturbances of function at the site of
An intracranial tumour which has not metastasised may give rise to focal signs of disordered nervous function at a distance from itself in a number of ways. Even though these neurological signs are labelled as false localising signs, it is important to be aware that such signs are in no way “false”. Various cranial nerve palsies have been reported as false localising signs, with the sixth cranial nerve being the most common. According to Gassel, ninth to 12th cranial nerve palsies never provide false localisation. Since Dodge reported the first case of false localising sign involving the lower cranial nerve, only two cases have been reported in the literature. We report a third case of false localising sign involving the left ninth and 10th cranial nerves.

A 29 year old man presented to the medical department of our hospital with history of hoarseness of voice of 15 days duration, dysphagia of 1 week duration, and cough with expectoration and respiratory distress of 2 days duration together with history of fever. On examination, he was febrile, with a pulse of 100 bpm and blood pressure of 120/80 mm Hg. Respiratory examination revealed bilateral coarse crepitations. Neurological examination revealed absent gag reflex on the left side with deviation of the palate to the right side without any other neurological deficit. Indirect laryngoscopic examination revealed paralysis of left vocal cord.

Haematological examination revealed haemoglobin (Hb) 13.6%, a WBC count of 16,800/mm, and an erythrocyte sedimentation rate (ESR) of 120 mm/h. Chest x ray of the patient revealed bilateral pleural pneumonitis. He was treated with antibiotics according to culture sensitivity. He progressively improved and was discharged. At discharge, he had persistent hoarseness of voice and vocal cord palsy on the left side. About 4 weeks later he presented with a history of bifrontal headache and was referred to our department. Neurological examination revealed bilateral papilloedema and left palatal palsy with absent gag reflex. Other cranial nerves were normal. Motor and sensory system examination was normal. Occasional swaying to the left side on tandem walking suggested involvement of the cerebellar system. In view of these findings, a left posterior fossa mass lesion involving the lower cranial nerves such as a schwannoma was suspected. However, magnetic resonance imaging (MRI) of the brain revealed a large isointense homogenously enhancing mass lesion attached to the convexity dura (fig 1). It also revealed evidence of herniation of the cerebellar tonsils below the margin of the foramen of magnum and anterior displacement of the cerebellum causing stretching of the lower cranial nerves on the left side. The patient underwent midline suboccipital craniectomy and total excision of the lesion. The cerebellum was found compressed and deeply indented by the tumour. Postoperatively he improved neurologically. His gag reflex and palatal movements progressively improved and he was asymptomatic at 2 month follow up.

False localising signs are unexpected neurological deficits and reflect pathology distant from the expected anatomical locus. Prominent false localising signs are less common today, as diagnosis is usually made at an early stage. Cranial nerve involvement as a false localising sign is found in 12.5% of cerebral tumours. According to Gassel, false localising signs are more common in patients with signs of raised intracranial pressure. Due to the long intracranial course, sixth cranial nerve palsy is commonly associated with supratentorial mass lesions as a false localising sign. Most reports described single cranial nerve disturbance as a false localising sign. Rarely have multiple cranial nerve palsies been reported as false localising signs.

Ehni proposed various mechanisms responsible for false localising signs. These include: (i) general compression of a nerve having a long course; (ii) meningitis; (iii) oedema and gliosis; (iv) metastatic deposits; (v) infarctions at a distance from the primary lesion. False localising signs are commonly associated with supratentorial mass lesions.

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Ehni proposed various mechanisms responsible for false localising signs. These include: (i) general compression of a nerve having a long course; (ii) meningitis; (iii) oedema and gliosis; (iv) metastatic deposits; (v) infarctions at a distance from the primary lesion. False localising signs are commonly associated with supratentorial mass lesions. Most cranial nerve dysfunctions presenting as false localising signs appear as anoxic dysfunctions. Rarely has a hyperactive dysfunction syndrome involving the cranial nerves such as trigeminal neuralgia and hemifacial spasm, been reported in the literature. A false localising sign involving lower cranial nerves is extremely rare with just two such cases reported to date. Maurice-Williams proposed two mechanisms causing lower cranial nerve palsy: firstly, cerebellar hemisphere impacting the foramen magnum, causing reaction and oedema and thereby compression of the lower cranial nerves; and secondly, displacement of the cerebellum to the contralateral side, forcing the brainstem to the ipsilateral side and thus exerting traction on the contralateral lower cranial nerve. The case reported here involved a large cerebellar convexity meningioma causing cerebellar herniation downwards into the foramen magnum and anteriorly into the lateral cerebello-medullary cistern which resulted in stretching of the lower cranial nerves. The cranial nerve function improved following excision of the tumour. Awareness of the possibility of false localising signs and the conditions in which they are most likely to occur is very important as they may be indicative of serious life threatening pathology within the neural pathway.

**References**

1. Gassel MM. False localizing signs: a review of the concept and analysis of the occurrence in 250 cases reported in the literature. PostScript 297

**Figure 1** MRI of the brain. (A) Sagittal section shows herniation of the tonsils below the foramen magnum due to an isointense mass lesion (arrow). (B) Axial T1 weighted image showing isointense mass lesion, enhanced homogenously with gadolinium, causing compression of the cerebellum and anterior displacement of the cerebellum into the lateral cerebello-medullary cistern resulting in stretching of the lower cranial nerves (arrow).
Acquired Chiari 1 malformation and syringomyelia following lumboperitoneal shunting for pseudotumour cerebri

An important but not widely recognised complication of lumboperitoneal shunting is the development of a Chiari 1 deformity and syringomyelia. We present a case of a patient who developed symptomatic cerebellar tonsillar descent and syrinx formation following treatment of pseudotumour cerebri with lumboperitoneal shunting.

Case report
A 31 year old woman was diagnosed with pseudotumour cerebri following development of headaches, loss of vision, and papilloedema, in association with a cerebrosplinal fluid (CSF) opening pressure of 36 cm H2O. Cranial imaging showed an attenuated ventricular system and no other abnormality. In particular, the posterior fossa was satisfactory in appearance. She was treated with lumboperitoneal shunt insertion, with resolution of her symptoms.

Twelve months later, the patient reported a 6 month history of left hemisensory loss, left hemisensory loss, and numbness. Neurological examination revealed wasting and reduced power of the intrinsic muscles of the left hand, and left-sided hyperaesthesia to pin-prick. Magnetic resonance (MR) imaging showed the development of cerebellar tonsillar descent and syringomyelia throughout the cervico-thoracic spinal cord. The patient underwent insertion of a low pressure ventriculoperitoneal shunt and removal of the lumboperitoneal shunt, with subsequent resolution of her symptoms. However, there was, however, no resolution of the syrinx on follow up MR imaging.

Discussion
The development of cerebellar tonsillar descent is a recognised but rarely reported complication following lumboperitoneal shunting, in the treatment of communicating hydrocephalus.\(^1\)\(^2\) It has been reported to occur in a large proportion of paediatric patients undergoing this procedure, with Chumas et al reporting a 70% incidence in this age group,\(^3\) but its incidence in the adult population is undefined. The development of secondary syringomyelia appears to be much less common, with the above paediatric patients reporting an incidence of syrinx formation of 4%. The development of Chiari 1 and syringomyelia formation following lumboperitoneal shunting for the treatment of pseudotumour cerebri is recognised but has been less commonly reported.\(^4\)\(^5\)

There is a small number of papers reporting chiari development following lumbar shunting for communicating hydrocephalus in children, but only two case reports of syringomyelia formation. The association of syrinx formation and cerebellar tonsillar descent through the foramen magnum is well described, and is postulated to occur as a consequence of a cranial-spinal CSF pressure gradient and diversion of CSF down the central spinal canal rather than over the cerebral convexities.\(^6\)\(^7\) It would seem remarkable that this complication is not seen more commonly in the treatment of pseudotumour cerebri.

The non-resolution of the syrinx, in our case following lumboperitoneal shunt removal, is consistent with other workers’ experiences, although resolution has been reported in one instance.\(^8\)\(^9\)

In conclusion, we describe the development of Chiari 1 deformity and syrinx formation as an important but otherwise poorly recognised complication of lumboperitoneal shunting in patients with pseudotumour cerebri.

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doi: 10.1136/jnnp.2004.045126

Competing interests: none declared

References
6 Williams B. The distending force in the production of communicating syringomyelia. Lancet 1969;4:189-93
We found that eight patients (three vegetative and five minimally conscious) showed consistent improvements in the highest ranked behaviours (table 1; p = 0.008) and total number of behaviours (p = 0.013) observed in the standing position (fig 1). Three patients (two vegetative and one minimally conscious) showed no change and one minimally conscious patient showed only an increase in the highest ranked behaviour observed. Although WHIM scores in three vegetative patients increased during standing, the behaviours observed did not reach a level suggesting awareness of self and/or environment. After standing the WHIM scores in the supine position were equal to or below those acquired before standing. No change in blood pressure was observed (p = 0.3).

Our preliminary results suggest that positional changes may have a significant impact on behaviours in vegetative and minimally conscious patients (although the benefit of this phenomenon in rehabilitation remains unproved, these findings have clear implications for the assessment and categorisation of patients). Neurological assessments used to classify patients according to international guidelines relating to the vegetative and minimally conscious states typically take place with the patient lying in bed. Where physical constraints permit, it may be important to also observe patients in the standing position.

Table 1

<table>
<thead>
<tr>
<th>Patient</th>
<th>VS/MCS*</th>
<th>Supine score</th>
<th>Behaviour observed</th>
<th>Standing score</th>
<th>Behaviour observed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>VS</td>
<td>43</td>
<td>Smiled</td>
<td>43</td>
<td>Smiled spontaneously</td>
</tr>
<tr>
<td>2</td>
<td>VS</td>
<td>4</td>
<td>Eyes held by painful stimulus &lt; 2 s</td>
<td>4</td>
<td>Eyes held by painful stimulus &lt; 2 s</td>
</tr>
<tr>
<td>3</td>
<td>VS</td>
<td>5</td>
<td>Looked at person briefly</td>
<td>26</td>
<td>Frowning/grimacing during physio</td>
</tr>
<tr>
<td>4</td>
<td>VS</td>
<td>1</td>
<td>Eyes opened briefly</td>
<td>49</td>
<td>Vocalised in response to pain</td>
</tr>
<tr>
<td>5</td>
<td>VS</td>
<td>14</td>
<td>Yawned, sighed</td>
<td>26</td>
<td>Frowning/grimacing during physio</td>
</tr>
<tr>
<td>6</td>
<td>MCS</td>
<td>13</td>
<td>Looked at person moving limbs &lt; 3</td>
<td>16</td>
<td>Turned eyes to look at person talking</td>
</tr>
<tr>
<td>7</td>
<td>MCS</td>
<td>20</td>
<td>Vocalised during physio</td>
<td>36</td>
<td>Switched gaze from one person to another</td>
</tr>
<tr>
<td>8</td>
<td>MCS</td>
<td>26</td>
<td>Frowning/grimacing to pain</td>
<td>34</td>
<td>Manosyaptic response to questions</td>
</tr>
<tr>
<td>9</td>
<td>MCS</td>
<td>14</td>
<td>Yawned, sighed</td>
<td>14</td>
<td>Yawned, sighed</td>
</tr>
<tr>
<td>10</td>
<td>MCS</td>
<td>18</td>
<td>Tracked for 3–5 seconds</td>
<td>28</td>
<td>Looked at object when requested</td>
</tr>
<tr>
<td>11</td>
<td>MCS</td>
<td>8</td>
<td>Made eye contact</td>
<td>23</td>
<td>Showed selective response to preferred people</td>
</tr>
<tr>
<td>12</td>
<td>MCS</td>
<td>42</td>
<td>Could find a card from four</td>
<td>43</td>
<td>Smiled spontaneously</td>
</tr>
</tbody>
</table>

*Patient classification at the time of recruitment is denoted VS (vegetative state) or MCS (minimally conscious state).

Acknowledgement

The authors are grateful to Dr R Barker for the neurological assessment.

References


BOOK REVIEWS

Neurological disorders in pregnancy
Editted by Jacqueline M Washington. Published by the Parthenon Publishing Group, 2004, £87.00 (hardcover), pp 150. ISBN 1-84214-189-9

Many neurological disorders occur in women of childbearing age. This small book is designed to bridge the wide gap existing between the disciplines of neurology and obstetrics. It provides a concise overview of the most common neurological disorders that may be seen during pregnancy. Three categories of problems are encountered by obstetricians and neurologists: management of neurological pre-existing disorders during pregnancy, neurological disorders directly due to pregnancy, and neurological affections that require treatment considerations during pregnancy. The book covers these three categories with eight chapters devoted to migraine, cerebrovascular disease, epilepsy, back pain, multiple sclerosis, peripheral neuropathies, myasthenia gravis, and central nervous infections. Each chapter includes consideration of the influence of pregnancy on the disorder, the effect of the disorder on pregnancy, and potential effects of proposed therapies on the developing foetus—all concerns shared by every clinician who care for pregnant women. The different chapters provide a useful resource with lists of dosages, contraindications, monitoring guidelines, and side effects of drugs in pregnancy.

Two other chapters, covering muscle diseases (in particular myotonic dystrophy) and brain tumours, could have been useful. One also could regret the nearly complete omission of figures or diagrams for a book intended not only for neurologists but also for obstetricians. In contrast, most chapters contain many useful tables.

A few remarks are also worth mentioning. For example, the section covering the course of migraine during and after pregnancy is sometimes redundant and could have been summarised. Post partum angiopathy should also be added to the aetologies of postpartum headaches. In the chapter on cerebrovascular disease, eclampsia and hemorrhage sections could have been better detailed.

On the whole, this book represents a useful concise text (more than an in-depth literature summary or detailed analysis of complex issues) written in a balanced, practical, and informative way. It can be used by a wide audience and will facilitate understanding and treatment of neurologic problems in pregnant women.

The neuropahtology of dementia, 2nd edition

Inspecting the hardcover graphics of this new edition your reviewer was startled to find blazoned his comments on the previous, and first, edition to encourage your purchase. It is therefore clear that I am a supporter of this enterprise in principle—though modesty will prevail and I will not flaunt my presence in

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Inspecting the hardcover graphics of this new edition your reviewer was startled to find blazoned his comments on the previous, and first, edition to encourage your purchase. It is therefore clear that I am a supporter of this enterprise in principle—though modesty will prevail and I will not flaunt my presence in
remarking on the desirability of this latest, and any subsequent, reediting.

This review will not take the form of a direct comparison between the present and previous editions since my recollection of the former is based on a fading and rose-tinted affection, rather than direct comparison with the source material. Obsessionality is a professional hazard in neuropathology—either acquired or innate—but in my case it does not extend to detailed record keeping of book loans. I hope the trainee who chose to keep the book has had much joy of it. No doubt his extended loan reflects the esteem he/she felt for the educational value of the first edition.

There is no doubt that the present book is considerably more up-to-date, reflecting significant increased content. However there is no flab, and the overall size and scope are, respectively, manageable and focussed. The book has acquired distinguished American co-editors in place of Dr James Morris (whose career trajectory has taken him deep into health service management) but much of James’s contribution remains, suitably updated, as a core of practical advice related to the diagnostic process in dementia neuropathology, and the particular pathologies associated with Alzheimer’s disease and Vascular dementia. The latter section partly reflects his welcome and homespun wisdom enriched by an experience fraught with unresolved problems of clinicopathological correlation. I am glad they have retained it. Similarly the contribution to the text by Professor Esiri shares this feeling of direct personal tutoring from an expert. The editors have retained the previous structure, roughly summarisable as: what dementia is, where in the brain might be affected, how to go about a pathological survey of a dementia brain, and, finally, what you might find related to specific diagnostic categories. This comprehensive approach is now fleshed out by the introduction of more authors to bring expertise related to individuals’ conditions, additional “introductory” material about the clinical genetics of dementia (styled “molecular diagnosis” for some reason) and neuroimaging in dementia, and increased content reflecting on pathogenesis and research into related conditions. The book is now an edited multi-author compilation rather than a more personal distillation from a small group. Looking at the arithmetic there are 27 USA authors, 14 UK, and four others from Australia, and Scandinavia—the latter empowered only to pronounce on alcohol and dementia and CADASIL. Bar this small non-cross Atlantic contingent the chapters work out at 11 USA and 10 UK, with two more from an unrecognised American propensity for job sharing.

The content is uniformly well presented and informative. Referencing largely dates out in 2000, indicating the long lead time for this type of book but this is not a bad thing. The modern tendency to rush in print with one’s latest minor observation on, for example, yet another apoplectic or oxidative modifier is a relatively insignificant part of the overall progress of neuropathological research in neurodegeneration. The purpose of a book like this is to record those aspects of the understanding of dementia disorders that withstand time and become part of the accepted wisdom rather than twitching with every modish straw in the wind.

The imaging chapter is an especially fine thing with dazzling illustration and some alarming jargon. Continuum-mechanical warping using calculations based on Cauchy-Navier equations with variable Lamé elasticity coefficients, and purple brains, add a new and distracting element to a neuropathology book. However the integration of neuropathology and neuroradiography is a highly desirable goal in clinical dementia research and it would be a pity to kick it out of the book. This content illustrates just how widely a “neuropathology” text needs to cast its net to retain its value in such an interdisciplinary world as neurodegeneration research and the clinical neuroscience of dementia.

In summary, another triumph and an indispensable addition to this field. If I had any hope of getting the book back I would automatically loan it to any new neuro-pathology trainee, but its appeal is far broader and it should be studied by anyone entering dementia research from a tissue-based angle. For a quotation to plug to buy over the sleeve of the third edition I offer the publisher: “buy one, get one free”.

Cordonnier

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Cerebrovascular disease, cognitive impairment, and dementia


The proportion of patients who will be victim of stroke or dementia is terrifying: after the age of 80 years, 1 in 5 people is affected by dementia, and 1 in 10 have had a stroke or transient ischaemic attack. Both the burden of stroke and dementia will continue to increase during the next 20 years in Western countries, owing to increasing life expectancy. Therefore, the economic burden of both disorders will also become a major public health issue. Stroke is an important cause of cognitive impairment and dementia. Stroke prevention, the only way to prevent vascular dementia, may also be an effective way to “prevent” Alzheimer’s disease—or at least to prevent the anticipation of its clinical onset, possibly due to the summation of vascular and Alzheimer lesions. Although the term “vascular dementia” appears in several chapters, the editors discuss two other important concepts. The first is the wide notion of “cerebrovascular impairment”, which includes a large range of severity of cognitive impairments associated with vascular lesions, behind this term is the hope of an effective prevention. The second is the interaction between Alzheimer lesions and stroke, explaining that many patients already have some degree of cognitive impairment before stroke, which may be degenerative in origin in many cases.

The book is divided into 26 chapters, including classification and diagnosis, epidemiology and risk factors, pathophysiology, clinical features, assessment, and management. The organization of the book proceeds logically. All chapters end with the most important references. The information is made clear and is accurate. The target audience consists of all care providers who treat patients with dementia or cerebrovascular disorders. Its length and its level of details make it appropriate for residents looking for a practical knowledge, and also for trained specialists. This book will be of major interest for all those who treat patients with cognitive decline or patients at risk.

D Leys

D Leys has been paid or received funds for research during the last 5 years by Sanofi-Synthelabo, AstraZeneca, Takeda, Lilly, and Servier for educational programmes, speaking, and consulting.