**SHORT REPORT**

Postinfectious vasculopathy with evolution to moyamoya syndrome

T Czartoski, D Hallam, J M Lacy, M R Chun, K Becker

Background: Parainfectious vascular events are a known complication of bacterial meningitis, typically occurring within two weeks of disease onset. Delayed vascular complications are rare. We present a case of progressive vasculopathy following bacterial meningitis.

Case description: A 20 year old woman developed progressive vasculopathy after successful treatment of pneumococcal meningitis. Within eight months of her infection, angiography revealed the appearance of moyamoya syndrome. Despite aggressive immunomodulation and anticoagulation, she had multiple strokes. Autopsy confirmed severe narrowing of proximal cerebral vasculature with absence of inflammation or atherosclerosis.

Conclusions: The inflammation and subsequent postinfectious autoimmune response associated with meningitis can lead to a progressive vasculopathy and may represent a pathophysiologic mechanism for the arterial occlusions seen in moyamoya syndrome.

Vascular events related to meningitis typically occur within the first two weeks of disease and are most commonly seen with Streptococcus pneumoniae, Mycobacterium tuberculosis, Neisseria meningitidis, and Haemophilus influenzae. Delayed postinfectious vascular events are less commonly reported in association with flow defects from fixed stenoses. We report a case of postinfectious vasculopathy following pneumococcal meningitis with progression to moyamoya syndrome.

CASE HISTORY

In July 2002, a 20 year old white woman with a history of migraines and bipolar affective disorder presented with 24 hours of fever, headache, and obtundation. Lumbar puncture (LP) was not performed due to marked cerebral oedema. She was treated empirically with vancomycin, ceftriaxone, and aciclovir. S. pneumoniae was cultured from her blood; ceftriaxone alone was sensitive to ceftriaxone. CSF examination was normal. Repeat angiogram on day 150 showed progression of the vascular stenoses (fig 1B). Anti-β2-GP1 G titres had increased to 56 SGU; there was no evidence of a lupus anticoagulant. A trial of intra-arterial papaverine failed to change the calibre of the vessels and improve blood flow. Renal angiography revealed arterial changes within the parenchymal vessels consistent with vasculitis. She was treated with cyclophosphamide on day 165.

Serial transcranial Doppler (TCD) studies were consistent with intracranial stenoses and on day 180 suggested improvement; the anti-β2-GP1 G titres had decreased to 24 SGU. The family declined further therapy with cyclophosphamide due to concerns over side effects. She was discharged on prednisone, 60 mg daily, tapered off over eight weeks; the taper concluded on day 228. Routine outpatient follow up two days later revealed no acute problems. The patient was ambulatory with a mild left hemiparesis and the ability to detect light and motion. On day 237 her carer brought her to the ER because of a change in behaviour and complete blindness. MRI revealed new infarcts in a watershed distribution in the right hemisphere and in the posterior parietal lobe in the left hemisphere (fig 1C). An LP revealed a recrudescence of CSF inflammation with 16 WBCs, predominantly lymphocytes.

The inflammation and subsequent postinfectious autoimmune response associated with meningitis can lead to a progressive vasculopathy and may represent a pathophysiologic mechanism for the arterial occlusions seen in moyamoya syndrome.

**Abbreviations:** β2-GP1, β2-glycoprotein 1; CSF, cerebrospinal fluid; LP, lumbar puncture; TCD, transcranial Doppler; WBC, white blood cell.
Table 1  Course of illness, therapeutic interventions, and response to therapy

<table>
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<th>Day</th>
<th>11</th>
<th>48</th>
<th>55</th>
<th>118</th>
<th>150</th>
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<th>165</th>
<th>180</th>
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<td>–</td>
<td>ST</td>
<td>ST</td>
<td>ST, AC</td>
<td>ST, AC</td>
<td>CY</td>
<td>AC</td>
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<td>Red blood cells (per ml)</td>
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<td>2</td>
<td>2</td>
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<td>55</td>
<td>150</td>
<td>158</td>
<td>165</td>
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<td>237</td>
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<tr>
<td>White blood cells (per ml)</td>
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<td>118</td>
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<td>Monocytes (%)</td>
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<td>15</td>
<td>14</td>
<td>3</td>
<td>100</td>
<td>6</td>
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<td>Protein (mg/dl)</td>
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<td>100</td>
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<td>43</td>
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<tr>
<td>Glucose (mg/dl)</td>
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<td>45</td>
<td>40</td>
<td>56</td>
<td>67</td>
<td>72</td>
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Anti-phospholipid antibodies

| Anti-β2-glycoprotein 1 (IgG; SLE) reference range: 0–9 | 29 | 56 | 24 | 52 |

TCD mean flow velocities (cm/sec)

| Right MCA, reference velocity: 58 | 164 | 312 | 157 | 287 | 195 | 350 |
| Left MCA, reference velocity: 58 | 309 | 334 | 344 | 328 | 328 | 328 |
| Right VA, reference velocity: 45 | 193 | 245 | 105 | 107 | 91  | 64  |
| Left VA, reference velocity: 45 | 231 | 121 | 140 | 108 | 176 | 169 |
| Basilar artery, reference velocity: 38 | NI | 162 | 119 | 193 | 114 | 136 |

AC, anticoagulation; CSF, cerebrospinal fluid; CTX, ceftriaxone; CY, cyclophosphamide; NI, not insonated; ST, steroids; TCD, transcranial Doppler ultrasound.

Figure 1  Imaging demonstrates progressive vasculopathy with corresponding infarcts in multiple vascular distributions. Computed tomography (CT) scan 10 days after presentation with meningitis was unremarkable (not shown). (A) Angiogram performed at day 12. The left vertebral injection demonstrates moderate narrowing of distal bilateral vertebral arteries (white arrowheads) and basilar artery (black arrowhead). The left carotid angiogram shows mild narrowing of the M1 segment of the left middle cerebral artery (MCA) (white arrow). (B) On day 139, fluid attenuation inversion recovery (FLAIR) and diffusion weighted images (DWI) show acute right posterior cerebral artery and left MCA distribution infarcts. On day 150, the angiogram shows severe narrowing in the basilar artery (black arrowhead) and left vertebral artery (white arrowhead). The carotid angiogram shows marked progression of the left M1 segment narrowing (straight white arrow) and new stenosis of the A1 segment of the left anterior cerebral artery (ACA) (curved white arrow). At this point, lenticulostriate collaterals become apparent (black arrow). (C) On day 237, DWI confirms acute infarcts in bilateral cerebral hemispheres. Proliferation of posterior penetrating vessels is shown on lateral vertebral angiogram (black arrowhead). There is further proliferation of lenticulostriate collaterals (black arrow), as well as the previously noted severe narrowing of proximal MCA (straight white arrow) and ACA (curved white arrow).
On day 260, she was brought to the ER by her family with acute obtundation following three days of decreased nutritional intake. The international normalised ratio was 6.3. A head CT scan revealed a large left basal ganglion haemorrhage. She progressed to herniation and expired. An autopsy revealed no systemic vasculitis or arteriopathy. The proximal cerebral vasculature exhibited severe luminal narrowing in the absence of inflammation or atherosclerosis (fig 2).

DISCUSSION
Vascular events are a known complication of bacterial meningitis but are usually limited to the parainfectious period.2-5 This case of postinfectious, progressive vasculopathy following pneumococcal meningitis appears to be the result of a delayed, dynamic process. A recent report by Palacio et al described a patient with late developing intracranial stenoses and benign course following H. influenzae meningitis.6 Their patient developed headache and transtemporal headache and progression to moyamoya syndrome. In our case, early postinfectious vasculopathy was similarly complicated by a delay, progressive intracranial vasculopathy with a more malignant course.

The pathogenesis of this type of postinfectious vasculopathy is unknown, although infection could trigger an autoimmune process towards the cerebral blood vessels. Streptococcal infections are associated with a variety of autoimmune diseases, including glomerulonephritis, chorea, myocarditis, arthritis, tics, and obsessive compulsive disorder.7-9 Triggering of antiphospholipid antibody syndromes (APAS) is also reported following infusions, and antibodies to β2-GP 1 are associated with postinfectious autoimmune APAS.9 In our patient, β2-GP 1 levels were elevated during periods of vascular disease progression, which might indicate an aberrant immune response to infection resulting in increased susceptibility to stenosis and thrombosis.

The pathogenesis of moyamoya disease is not understood. Genetic, autoimmune, and infectious aetiologies have been postulated. Autopsy studies of patients with moyamoya disease classically show intimal thickening with crenulation of the elastic lamina—infarction is absent.10 11 Similar changes were seen in our patient. The absence of inflammation of the cerebral vessels at autopsy implies that the patient either responded to the aggressive immunomodulatory therapy or the autoimmune process spontaneously arrested.

Unfortunately, no vascular studies were done prior to the onset of the patient’s symptoms, so the possibility of pre-existing vascular disease cannot be excluded. Several pieces of evidence, however, suggest that her vascular disease was related to the meningitis. First, she was asymptomatic prior to the event. Second, she developed progressive stenoses with multiple infarcts over the course of her illness. Thirdly, vessels outside the subarachnoid space were spared, suggesting the vasculopathy resulted from intracranial inflammation, as is seen in meningitis.

This case illustrates the occurrence of postinfectious vasculopathy with progression to moyamoya syndrome following S. pneumoniae meningitis. Due to the progressive course, elevated anti-β2-GP 1 titres, and transient response to immunomodulatory therapy, we believe that the vasculopathy was mediated by an autoimmune process and suggests a possible pathogenesis for moyamoya syndrome.

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REFERENCES
Brodmann's cortical maps

Vincent d'Azyr, a physician and artist, described the brain's convolutions in 1786, noting differences in morphology in other animals. Magendie had written similarly. Early attempts to correlate the cerebral anatomy to function by observed neurological deficits began in the 1820s, the result of the work of Franz Gall,1 Bouillaud, Robert Todd, Rolando, and many others (references in).2 Pierre Gratiolet and Francois Leuret mapped the folds and fissures of the mammalian cortex. Vogt suggested to Brodmann that he many concerned with the comparative cytoarchitectonics of mammalian species. However, he faced difficulties, observing: “First and foremost we still lack clear criteria for the recognition of anatomically equivalent cellular elements...There has been occasional talk of ‘sensory cells’ located in particular regions, or of sensorial ‘special cells’. People have invented acoustic or optical special cells and even a ‘memory’ cell, and have not shied away from the fantastic ‘psychic cell’.”

This, he stoutly rejected, concluding, “Functional localisation without the lead of anatomy is utterly impossible.”

Oskar Vogt described Brodmann as having “broad scientific interests, a good gift of observation and great diligence in widening his knowledge”. His interests extended to neurology, psychiatry, physiology, zoology, and anthropology. He was described as “an intense and earnest man, reserved to the point of timidity, but could flare, on occasion, into a tempest”, possibly frustrated by his inability to secure a permanent job. Not until 1916, aged 48, did Brodmann obtain this security. He had left Berlin in 1910 to work with Gaupp at Tübingen, where he was made titular professor in 1913. Finally, in 1918, he accepted an invitation from Munich to take the Chair of the Topographical Histological Department at the research centre for psychiatry. He died in 1918 of septicaemia complicating pneumonia.

In 1919, Cécile Vogt described over 200 cortical areas and six years later von Economo and Koskinas revised the nomenclature.

References
5 Bielschowsky M, Brodmann K. Zur feineren histologie und histopathologie der grosshirnrinde (localisation in the cerebral hemispheres: a comparative study), which described 52 discrete cortical areas. He defined cytoarchitectonics as “The localization of the individual histological elements, their layering, and their parcellation in the adult brain.”

Although there were other investigators of cytoarchitectonics,2 9 Brodmann’s maps and, especially, his numerical system attached to cytoarchitectonics were widely accepted; they were his major contribution. He had studied 64 different mammalian species. However, he faced difficulties, observing:

“...The founders of neurology...”


HISTORICAL NOTE