Short report

Familial cerebellar ataxia with cerebrovascular amyloid

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SUMMARY We report a rare association of familial cerebellar ataxia (without dementia) and cerebrovascular amyloid. Postmortem neuropathological examination of one member of the family showed amyloid angiopathy of the central nervous system with heavy infiltration of capillaries in the hippocampus and cerebellum.

Cerebrovascular amyloid occurs in a number of clinical contexts and may be an asymptomatic finding in the elderly. Patients may develop dementia and the risk of cerebrovascular accidents is increased. There are known associations of amyloid angiopathy with Alzheimer's disease and spongiform encephalopathy. To our knowledge, cerebellar ataxia with cerebrovascular amyloid has previously been reported in only one family in which affected members became demented early in the course of the illness. The present report is of cerebrovascular amyloid affecting one member of a family in which four members in three successive generations developed cerebellar ataxia without dementia.

Case report

Mrs IB, a 49-year-old housewife, presented with slight unsteadiness of gait. She gave no history of alcohol ingestion, exposure to carbon monoxide, previous head injury or encephalitis, and was taking no medication. Her maternal grandfather had been bedridden with "unsteady limbs" for 9 years prior to his death. One maternal uncle died following a cerebral haemorrhage, and two maternal uncles suffered from cerebellar ataxia with nystagmus. Eleven other maternal aunts and uncles as well as the patient's mother died from unrelated illnesses. Two siblings are believed to be healthy.

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Examination disclosed mild, symmetrical cerebellar ataxia of the limbs and trunk with a slurring dysarthria and nystagmus on lateral gaze. The blood pressure was 120/70 mm Hg while lying supine. Haematological and biochemical investigations, including thyroid function tests, were normal. No abnormalities were noted on chest, skull and cervical spine radiographs. A Wassermann test was negative. The ataxia worsened progressively so that within two years she could not walk without support. Three years after her first presentation she developed sudden weakness and stiffness of the left lower limb. Tone was increased in that limb, with left ankle clonus and bilateral extensor plantar responses. Sensation was intact. An electroencephalogram was diffusely abnormal; irregular, intermittent 11 Hz alpha rhythm was present with a left-sided emphasis and irregular 15 Hz activity occurred with a right-sided emphasis. A brain scan was normal. Progressive incapacitation led to her admission to a home for the physically handicapped. During her last few months of life she became incontinent of urine and, for the first time, impairment of memory for recent events was noted, although she remained orientated and lucid. She died ten years after the onset of her illness, at the age of 59 years.

Neuropathological findings Consent was obtained for a post mortem examination limited to the central nervous system. The external appearance of the brain and spinal cord was normal. The unfixed brain weighed 1240 g. After fixation the cerebrum was sectioned coronally. The lateral and third ventricles were slightly dilated. Two slit-like, small infarcts were present in the white matter of the right frontal and temporal lobes. No abnormalities were seen in the brain stem, cerebellum or spinal cord. Blocks were taken from many areas of the cerebral hemispheres, brain stem, cerebellum and spinal cord. In the cerebral hemispheres and leptomeninges, arteries...
and arterioles were diffusely thickened by eosinophilic, acellular material with affinity for Congo red and showing yellow-green birefringence under polarised light thus fulfilling the criteria for amyloid.7 8 In many areas the lumen was narrowed and in places amorphous deposits of amyloid extended for short distances into the surrounding brain parenchyma. Scanty deposits were present in the tunica media of some veins. Involvement of capillaries was virtually restricted to the hippocampal formation where amyloid infiltrated capillary walls, occasionally occluding the lumen, and radiating spicules of amyloid extended into the surrounding parenchyma. These deposits were not argyrophilic with either the Glees and Marsland or Palmgren method of silver impregnation. In addition, numerous neuritic (senile-type) plaques were present in the dentate fascia, hippocampal formation and subiculum but nowhere else in the cortex, and many hippocampal pyramidal cells contained neurofibrillary tangles or granulovascular inclusions. The presence of old infarcts in the white matter was confirmed. The basal ganglia and brain stem showed widespread arterial amyloid, capillaries being spared.

The cerebellum was extensively infiltrated by large deposits of amyloid, many of which were clearly related to capillaries (fig 1). These deposits were numerous in the molecular and granule cell layers and a few were present in the white matter. Generalised arterial amyloid was present in the cerebellum and overlying leptomeninges. There was severe depletion of Purkinje cells and many "torpedoes" were noted on axons of those surviving (fig 2). The granule cells were relatively well preserved. The molecular layer showed Bergmann cell hyperplasia and there was nerve fibre loss and gliosis in the white matter. Arterial amyloid was present throughout the spinal cord and in the spinal leptomeninges with only occasional capillary involvement in the white matter. There was a rim of nerve fibre loss in the subpial white matter of the cervical region. The nerve roots showed no vascular amyloid and no other pathological abnormalities.

Discussion

Worster-Drought, Greenfield and McMenemey described a family with presenile dementia, spastic paraplegia, ataxia of gait and nystagmus.5 6 Post-mortem examination of two members showed generalised "hyaline" arterial wall thickening and argyrophilic pericapillary deposits (later confirmed to be amyloid) most numerous in the hippocampal formation and cerebellum, but also involving the pons and medulla. Neuritic plaques and neurofibrillary tangles were present in the hippocampal cortex, and the cerebellum showed some loss of Purkinje cells with "torpedoes" on axons of surviving cells. Their case, unlike the present one, showed widespread white matter degeneration and gliosis. Although atypical "kuru" plaques have been described in other progressive neurological diseases with cerebellar ataxia,9 10 vascular amyloid has not been present.

In the family described by Worster-Drought et al, dementia was a prominent feature. This may reflect the extensive white matter changes and, possibly, more severe hippocampal damage. The patient reported here remained lucid and alert until her death, and no other family members became demented. However, ataxia of gait and nystagmus were common to both families, with almost identical appearance and distribution of the cerebrovascular amyloid. Familial cerebellar ataxia, with or without dementia, appears, therefore, to be a further manifestation of cerebrovascular amyloid, which is associated with microvascular amyloid in the hippocampus and cerebellum.
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References


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