Task specific focal dystonia: a presentation of spinocerebellar ataxia type 6

Autosomal dominant cerebellar ataxias (ADCA) are characterised by clinical and genetic heterogeneity with a substantial overlap of clinical features and a variable degree of adherence to three distinct phenotypes according to Harding’s clinical classification: ADCA type I, II, and III. The availability of molecular genetic testing has provided increasing appreciation of a wider clinical spectrum than previously thought for each ADCA subgroup. In addition there is an increasing list of genes harbouring disease causing mutations which, to date, include spinocerebellar ataxia (SCA) types 1, 2, 3, 6, 7, 8, 10, 12, and 17. SCA 6 is caused by unstable CAG expansion in the α-1A calcium channel gene (CACNA1A). We present a confirmed case of SCA 6 presenting with a task specific focal dystonia (writer’s cramp) predating the onset of progressive gait ataxia by five years, which further widens the clinical spectrum of SCA 6.

Case report
A 57 year old female pharmacist first presented at the age of 42 with a two month history of progressive difficulty in writing prescriptions as a result of writing induced cramp affecting the right hand. This task specific focal dystonia was aggravated by stress and fatigue. Her father was reported to have had Parkinson’s disease on the basis of developing tremor and gait disturbance; however, none had developed writer’s cramp.

On examination, the patient’s gait was normal and no focal neurological deficits were identified. Her writing, after a normal start, was interrupted by a semiflexed posture of the fingers to maintain the grasp of the pen, with the index finger and thumb appeared hyperextended and the wrist hyperflexed. This was accompanied by an ache in the hand. A working diagnosis of writer’s cramp with a suspicion of familial dystonic disorders was made and she was started on the anticholinergic drug, trihexyphenidyl.

On annual follow up, she reported a satisfactory improvement in her symptoms. During this period and on more than one occasion, no cerebellar signs were identified. Five years after the onset of focal dystonia, she developed an intention tremor in her right hand and upper limb ataxia but with a normal gait. A diagnosis of autoimmune hypothyroidism was made following a confirmatory test results and thyroxine was started concurrently with propranolol to control the tremor. Two years later, at the age of 49, she began to experience difficulties with maintaining her balance when walking downhill, which later progressed to an increasing tendency to sway or trip. The writer’s cramp had persisted with progressive tremor and chuminess in both arms, forcing her to swap hands when writing (fig 1). A more detailed family history revealed additional information about paternal relatives who had a history of tremor and gait disturbance; however, none had developed writer’s cramp.

Neurological examination now showed evidence of gait and limb ataxia, mild intention tremor, and horizontal gaze-evoked nystagmus, with mild dysarthria and no other neurological deficit. Ancillary tests to exclude acquired ataxias were also done, and brain magnetic resonance imaging revealed evidence of marked cerebellar atrophy with sparing of cortical and brain stem structures. With informed consent, genomic DNA was extracted from peripheral blood leucocytes. Screening for trinucleotide repeat expansions for SCA 1, 2, 3, and 6 was done using polymerase chain reaction (PCR) amplifications of the genomic sequences containing each trinucleotide repeat expansion site. The PCR products were size fractionated by electrophoresis in a 6% polyacrylamide gel, which was then dryblotted. After hybridisation with a gamma-32P labelled (CAG)n oligonucleotide probe, allele sizes were visualised by autoradiography. Estimations of trinucleotide repeat size for each gene was made using well characterised positive control samples. A pathological 22 repeat expansion in the CACNA1A gene and another allele of 12 repeats were detected (fig 2). Alleles within normal range were detected in the SCA 1, 2, and 3 genes.

Comment
We describe a case of SCA 6 with a focal dystonia preceding the onset of gait or limb ataxia by a period of at least five years. Recently, there has been a report of similar SCA 6 presentation but with a more progressive, disabling, and treatment resistant shoulder girdle and upper limb dystonia. In the original description of SCA 6, the most common presenting feature was a progressive cerebellar syndrome, accompanied by involuntary movements, dystonic posturing, sensory loss, and changes in the deep tendon reflexes. Other presenting features have been reported as a result of direct molecular testing, including episodic ataxia and positional vertigo associated with downbeat nystagmus. Our report provides further evidence of a wider clinical presentation of SCA 6, but may also be relevant to other ADCA, which were previously considered to be relatively “pure” cerebellar ataxias.

This case also shows that, in the absence of a suggestive family history or presenting cerebellar features, it is unusual for clinicians to include ADCA in the differential diagnosis. Dystonia or extrapyramidal motor signs have often been associated with SCA 3, but are not specific, as overlap exists, though less frequently, with other subtypes in ADCA 1. With disease progression, the focal dystonia in our patient has partially resolved and has been predominantly replaced by a mild pancerebellar ataxia with very few extracerebellar signs and absence of other extrapyramidal features. The evolution of these signs and the partial resolution of the writer’s cramps persuade us that the dystonia and ataxia are part of the same pathological process rather than separate disease entities. Ten years after the onset of gait disturbance, the patient is walking using only one stick and remains employed, consistent with the mild disease course previously reported for SCA 6 phenotype. Increasingly, the availability of molecular investigations is indicating a need to revise the phenotype according to the underlying genotype. It is apparent that the majority of the inherited ataxias have a broader clinical spectrum than has previously been appreciated, particularly with ADCA. Our experience highlights the wide range of syndromic presentations as well as advocating the clinical value of testing SCA 6 alongside SCA 1, 2, and 3 which constitute ADCA 1. In addition, it may be useful to screen SCA loci in the ADCA 1 subgroup in cases of idiopathic focal dystonia who develop features of cerebellar dysfunction or who have a relevant family history.

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Figure 2 Autoradiography showing polymerase chain reaction products containing CAG repeats for SCA 6. Products were blotted and probed with a radiolabelled (CAG)n oligonucleotide. Lanes 2A/2B (loaded twice) showing patients with an allele of 22 CAG repeats. Results were confirmed with controls: lanes 3A/3B (12/22) and lane 5 (14/30). Lanes 1A/1B were another patient with ataxia; lane 4 was an unrelated healthy control, and lane 6 was blank.

Figure 1 The patient’s writing samples showing the use of the dominant right hand (line 2), the eventual use of the left hand (line 3), and the recent use of both hands (line 4), attempting to copy a short sentence (line 1).
Bilateral cerebellar ataxia as the sole manifestation of a unilateral rostral pontine tegmental infarct

It has been reported that a small infarct of the pons can lead to various clinical syndromes such as pure motor hemiparesis, sensorimotor stroke, ataxic sensorimotor stroke, ataxic hemiparesis, dysarthria-clumsy hand syndrome, or ataxic tetraparesis. However, bilateral cerebellar ataxia as the sole manifestation of rostral pontine tegmental infarction has not been described. We report a patient with isolated bilateral cerebellar ataxia as the only sign of a rostral pontine tegmental infarct. This unique presentation may reflect the selective involvement of part of the decussation of the superior cerebellar peduncle.

Case report

A 51-year-old man with hypertension developed acute severe imbalance. On examination, he tended to fall to the right when standing unsupported with eyes open. He did not have dysarthria, limb weakness, vertigo, nystagmus, ophthalmoplegia, diplopia, sensory loss, or Horner’s syndrome. The muscle stretch reflexes were normal and the plantar reflexes were flexor bilaterally. There was severe dysmetria on finger-to-nose and heel-to-shin testing on both sides. Dysmetria was worse on the right. The results of somato-sensory evoked potentials, and pure tone audiography were unremarkable. There were no abnormalities of horizontal saccades, smooth pursuit, optokinetic nystagmus, or caloric responses.

Brain magnetic resonance imaging (MRI) showed a right paramedian infarct of lacunar size situated in the tegmentum of the most rostral pons, corresponding to part of the decussation of the superior cerebellar peduncle (fig 1). Magnetic resonance angiography (MRA) showed no abnormalities. An ECG and transoesophageal echocardiography with a bubble study revealed no abnormalities.

The limb coordination and gait improved steadily over several days, but there was mild dysmetria on the heel-to-shin test on the right side. On discharge, the patient complained only of mild unsteadiness when walking.

Comment

There have been several reports of bilateral cerebellar ataxia caused by a unilateral brain stem stroke. However, previous reports have also described associated neurological symptoms such as mild hemiparesis, dysarthria, sensory change in an extremity, or multiple cranial nerve palsies.

Without pathological confirmation, it is difficult to be certain that the infarct affected only the structure identified (the superior cerebellar peduncle). However, in the rostral pons, the only anatomical structure responsible for bilateral limb ataxia is the superior cerebellar peduncle, which is situated in the dorsolateral side to the fourth ventricle and medial to the lateral lemniscus at the level of the most rostral pons—that is, at isthmus level. From this level, the fibres of the superior cerebellar peduncle move ventromedially towards their decussation. Serial neurological examinations over a period of days did not show any neurological signs except bilateral ataxia. These clinical data, when correlated with the known cross sectional anatomy of the most rostral part of the pons, suggest that the small lesion of our patient on brain MRI corresponded to part of the decussation of the superior cerebellar peduncle. Thus the isolated bilateral cerebellar ataxia in our patient may be explained by ipsilateral involvement of both efferent cerebellar pathways. These which include uncrossed fibres of the superior cerebellar peduncle ipsilateral to the lesion and crossed fibres arising contralateral to the lesion.

From the results of MRA and transoesophageal echocardiography, risk factor analysis, and the size of an infarct on brain MRI, small artery disease (that is, a lacunar stroke) was considered the likely pathogenesis.

In summary, our patient presented with isolated bilateral cerebellar ataxia caused by a small infarct situated in the rostral pontine tegmentum. This unique presentation may result from ipsilateral involvement of both efferent cerebellar pathways, before and after the decussation of the superior cerebellar peduncle. We have previously reported isolated ataxia as the sole manifestation of lateral medullary infarction. Together, these reports highlight the importance of sudden gait disturbance as the sole manifestation of brain stem stroke. Isolated bilateral cerebellar ataxia caused by a unilateral pontine paramedian tegmental infarction should be considered in the differential diagnosis of sudden bilateral cerebellar ataxia, even when classic brain stem signs are absent.
Serial EEG records showed diffuse delta and theta activity with occasional prevalence of this activity in the left sided anterior regions. Cranial CT and MRI revealed diffuse oedema with hydrocephalus and dilatation of the cerebral aqueduct. No focal lesions were observed.

Treatment included dexamethasone (4 mg intravenously twice daily), doxycyclin (100 mg twice daily), rifamycin (600 mg daily), and amphotericin B, starting at doses of 20 mg intravenously per day with a progressive increase to 50 mg daily; simultaneous intrathecal amphotericin B was given at an initial daily dose of 0.012 mg, increasing progressively to a maximum daily dose of 0.250 mg.

Despite this treatment, the clinical situation rapidly worsened and the patient died after 11 days in hospital. A necropsy examination was denied.

Comment

Free living amoebae of the genus *Acanthamoeba* are the causative agents of several infections usually occurring in immunocompromised, debilitated individuals and almost always resulting in death. Most cases, therefore, are diagnosed only at necropsy. Our patient was apparently immunocompetent and without any of the usual predisposing factors, such as a history of aquatic activities, treatment with immunosuppressive, chemotherapeutic, or steroid agents or broad-spectrum antibiotics, and so on. The existence of extraneural infective foci in the skin, paranasal area, or lungs—a possible point of access for amoebae—was also excluded. The most striking feature in our patient was that a firm diagnosis was made only through direct observation of the protozoon in the CSF. To our knowledge, this has not been described before in chronic amoebic meningoencephalitis and underlines the diagnostic value of CSF studies in this type of pathology. The main species reported as causing granulomatous amoebic encephalitis are *A. polyphaga, A. castellanii, A. culbentina, and Balamuthia mandrillaris*.

The diagnosis is usually made by microscopic examination of stained slices of brain specimens obtained at necropsy or biopsy and cultivation of the causal organism in an appropriate medium. This usually consists of non-nutrient agar covered with bacteria for the species determination may prove problematic. In our patient, we were unable to discriminate among the various species of amoebae that could have been involved in the infection. Culture tests on non-nutrient agar covered with bacteria were negative, and other cultures, serological tests, and CSF tests were not done because of the rapid progression of illness. Nonetheless, our case indicates that when an amoebic meningoencephalitis is suspected a careful search for the organisms in the CSF may be a decisive factor in the diagnosis.

In conclusion, this case report emphasises the importance of familiarising ourselves with this form of pathology and provides an example of how the identification of amoebae in the CSF may aid in making a firm diagnosis of this uncommon, undiagnosed, life threatening, and difficult to treat CNS infection. As a successful therapeutic result may sometimes be achieved, a timely diagnosis together with prompt and adequate treatment are essential.

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References


