Allgrove or 4 “A” syndrome: an autosomal recessive syndrome causing multisystem neurological disease

J Kimber, B N McLean, M Prevett, S R Hammans

As in 1978 by Allgrove, the syndrome may thus manifest itself during the first decade of life with dysphagia or severe (occasionally fatal) hypoglycaemic or hypotensive attacks, related to adrenocortical insufficiency. Onset of adrenal insufficiency or other features may be delayed to adulthood. In contrast with paediatric patients, adult patients with Allgrove’s syndrome may present with multisystem neurological disease; childhood history of achalasia or alacrima may be overlooked. The authors describe two families with two affected siblings and a further unrelated patient with typical clinical features of Allgrove’s syndrome, who exhibit signs of multisystem neurological disease including hyperreflexia, muscle wasting, dysarthria, ataxia, optic atrophy, and intellectual impairment. None of the cases have developed adrenocortical insufficiency but all have progressive neurological disability. Autonomic dysfunction was a significant cause of morbidity in two cases. The three index cases represent the longest described follow up of Allgrove’s syndrome into adulthood. It is speculated that they represent a subgroup of patients who follow an often undiagnosed chronic neurological course. Recognition of the syndrome presenting in adult life permits treatment of unrecognised autonomic dysfunction, adrenocortical insufficiency and dysphagia, and appropriate genetic advice.

Allgrove and colleagues in 1978 first described this syndrome in two pairs of siblings (aged 4–6 years). All four had achalasia and ACTH insensitivity, three had impaired lacrimation, and one had autonomic dysfunction. The onset of adrenocortical impairment is usually before puberty, although preservation of cortisol secretion into the third decade has been reported. The syndrome may thus manifest itself during the first decade of life with severe hypoglycaemic or hypotensive attacks, which may lead to sudden death.

Follow up of the patients originally described by Allgrove, showed all developed variable combinations of sensorimotor polyneuropathy, long tract degeneration, mild dementia, and in one case an akinetic-rigid syndrome by the third decade. A further review of 20 cases aged 2–29 showed additional neurological abnormalities in 17 with onset between 2 and 17 years. These included amyotrophy, dysarthria, ataxia, optic atrophy, intellectual impairment plus autonomic dysfunction.

Previous genetic analysis of eight families with Allgrove’s syndrome demonstrated linkage to markers on 12q13. Recent studies in North African and other kindreds with triple dysphagia or severe (occasionally fatal) hypoglycaemic or hypotensive attacks, related to adrenocortical insufficiency. Onset of adrenal insufficiency or other features may be delayed to adulthood. In contrast with paediatric patients, adult patients with Allgrove’s syndrome may present with multisystem neurological disease; the childhood history of achalasia or alacrima may be overlooked. The authors describe two families with two affected siblings and a further unrelated patient with typical clinical features of Allgrove’s syndrome, who exhibit signs of multisystem neurological disease including hyperreflexia, muscle wasting, dysarthria, ataxia, optic atrophy, and intellectual impairment. None of the cases have developed adrenocortical insufficiency but all have progressive neurological disability. Autonomic dysfunction was a significant cause of morbidity in two cases. The three index cases represent the longest described follow up of Allgrove’s syndrome into adulthood. It is speculated that they represent a subgroup of patients who follow an often undiagnosed chronic neurological course. Recognition of the syndrome presenting in adult life permits treatment of unrecognised autonomic dysfunction, adrenocortical insufficiency and dysphagia, and appropriate genetic advice.

A syndrome has identified mutations in a WD-repeat protein gene, labelled AAAS. This report describes the clinical features in three adult patients with this disorder, with particular attention to the clinical, neurophysiological, and autonomic abnormalities found in adult patients.

**CASE REPORTS**

**Patient 1**

This man aged 40, was born to non-consanguineous parents. At age 5 he was felt to have delayed developmental milestones and subsequently attended a school for learning difficulties. At age 9 he underwent a Heller’s cardiomyotomy for achalasia of the cardia and the next year required achilles tendon lengthening surgery. His brother also had achalasia and died of an oesophageal perforation aged 45. The patient presented aged 34 to neurological services with progressive muscular weakness and gait ataxia. Alacrima, hypotension, and erectile dysfunction were also noted together with regional hyperhidrosis. Currently he remains ambulant but has developed dysarthria and refractory orthostatic hypotension.

On examination he had dysmorphic facial features with dysarthria and nasal speech. He had dilated pupils with an absent direct light response and poor response to accommodation. Fundoscopy revealed bilateral optic atrophy. Bilateral palatal paresis with an absent gag reflex was noted. His tongue appeared small and spastic with normal filiform papillae. There was global wasting of both arms and legs more marked distally but without fasciculations. Weakness was present in the same distribution. Tendon reflexes were generally brisk but both ankle jerks were absent. Both plantar responses were extensor and he had pes cavus. Limb coordination was relatively preserved but he had marked gait ataxia. Cutaneous sensation including pain perception was normal.

**Patient 2**

A woman aged 45, was born of non-consanguineous parents. She was noted to have a nasal voice as a child. She underwent a Heller’s cardiomyotomy for achalasia of the cardia aged 24. She first presented to the neurology service 10 years later with paraesthesiae in the right arm and hand. On examination mild palatal weakness was noted but tendon reflexes were preserved and no firm diagnosis was made. Aged 37 she complained of gait disturbance, posturally induced presyncope, and regional hyperhidrosis and was re-investigated at a neurological centre. Examination findings included reduced direct and consensual pupillary light response with bilateral pallor of the optic discs. Wasting and fasciculation of the tongue with palatal paresis was present. There was distal wasting in the arms and mild pyramidal weakness of the legs. All reflexes were brisk and both plantar responses extensor. She had bilateral pes cavus. Mild limb ataxia was present together with some distal sensory loss in the legs. A marked postural fall in blood pressure was noted. Her diagnosis was made at the age of 43 when she developed frequent blackouts.
and dysphagia. Currently she requires PEG feeding and is treated with midodrine for postural hypotension. Her sister died of carcinoma of the ovary in her fifth decade and had milder but qualitatively similar neurological symptoms. Both siblings also exhibited features of gonadotrophic dysfunction; delayed menarche, early menopause, and reduced fertility.

**Patient 3**

This man aged 60 years, was born to non-consanguineous parents. At age 12 he was noted to have a scoliosis. By the age of 20 his gait was unsteady and was aware of reduced sensation in his feet. Aged 28 he developed dysphagia and subsequently underwent a Heller’s procedure for achalasia of the cardia. Aged 60 he presented to the neurology service with frequent blackouts related to sudden postural change. Erectile dysfunction and regional hyperhidrosis were noted. His walking had deteriorated and had limited mobility with one stick. On examination there was right sided partial ptosis together with miosis. Pursuit eye movements were fragmented. Speech was normal. Wasting of the interossei was present bilaterally, although limb power was relatively preserved. Reflexes were brisk and the left plantar response was extensor. Reduced sensation was evident in the legs to the mid calf. Lower limb and gait ataxia were present.

**DISCUSSION**

In all our cases achalasia preceded the development of progressive neurological disease—typically a combination of sensorimotor polyneuropathy, long tract degeneration, optic neuropathy, and cerebellar features together with impaired parasympathetic and sympathetic autonomic function.

Palatal paresis was a feature in two of our cases and is reported by other authors. It is an uncommon finding in neuropathy and may represent central neuronal degeneration.

There are several reports in the literature of severe neuropathy, ataxia, or alacrima with achalasia and we speculate that these too are forms of Allgrove’s syndrome. The syndrome seems least likely to be recognised if presenting with neurological features in adult life. Intelectual impairment was suggested by the clinical history in case one, although formal neuropsychometry was not performed. Cognitive problems have been recognised in some paediatric patients with Allgrove’s syndrome.

### Table 1A Electrophysiological tests

<table>
<thead>
<tr>
<th>Sensory nerve</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>R sural</td>
<td>Absent</td>
<td>Amp 2.0 µV, Lat 4.8 ms, C V 29 ms</td>
<td>Amp 3.7 µV, Lat 4.0 ms, C V 30 ms</td>
</tr>
<tr>
<td>R digiti I</td>
<td>Absent</td>
<td>Amp 2.0 µV, Lat 2.8 ms, C V 39 ms</td>
<td>Amp 9.9 µV, Lat 2.9 ms, C V 43 ms</td>
</tr>
<tr>
<td>R digiti V</td>
<td>Absent</td>
<td>Amp 2.0 µV, Lat 3.0 ms, C V 33 ms</td>
<td>Amp 4.6 µV, Lat 3.4 ms, C V 35 ms</td>
</tr>
<tr>
<td>R median CAP</td>
<td>Amp 7.7 µV, Lat 4.0 ms, C V 68 ms</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>R ulnar CAP</td>
<td>Amp 7.5 µV, Lat 4.1 ms, C V 68 ms</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>R radial</td>
<td>Amp 6.5 µV, Lat 2.3 ms, C V 49 ms</td>
<td>–</td>
<td>Amp 3.4V, Lat 1.6V, C V 54 ms</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Motor nerve</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>R tibial</td>
<td>Absent</td>
<td>Amp 5.0 µV, Lat 5.4 ms, C V 40 ms</td>
<td>–</td>
</tr>
<tr>
<td>R peroneal EDB</td>
<td>Absent</td>
<td>Amp 3.5 µV, Lat 5.0 ms, C V 39 ms</td>
<td>Amp 7.4 µV, Lat 4.7 ms, C V 43 ms</td>
</tr>
<tr>
<td>R ulnar</td>
<td>Amp 0.2, Lat 3.7 ms, C V 40 ms</td>
<td>Amp 4.0 µV, Lat 3.8 ms, C V 47 ms</td>
<td>Amp 6.7µV, Lat 3.1ms, C V 53 ms</td>
</tr>
<tr>
<td>R median</td>
<td>Absent</td>
<td>Amp 9.0 µV, Lat 4.6 ms, C V 43 ms</td>
<td>Amp 7.6 µV, Lat 4.7 ms, C V 49 ms</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>EMG</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>R EDC</td>
<td>Nil spontaneous, Large polyphasic AP, ↓ interference</td>
<td>Nil spontaneous, ↑ duration polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
<tr>
<td>R Quadiceps</td>
<td>–</td>
<td>Fibrillations, Large polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
<tr>
<td>R Biceps</td>
<td>–</td>
<td>Nil spontaneous, ↑ duration polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
<tr>
<td>R Triceps</td>
<td>–</td>
<td>Fibrillations, Large polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
<tr>
<td>R Tib Ant</td>
<td>–</td>
<td>Nil spontaneous, Large polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
<tr>
<td>R APB</td>
<td>–</td>
<td>Nil spontaneous, Large polyphasic AP, ↓ interference</td>
<td>–</td>
</tr>
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</table>

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<tr>
<th>Evoked potentials</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>VEP</td>
<td>Bilateral delay</td>
<td>Bilateral delay</td>
<td>Normal</td>
</tr>
<tr>
<td>BSEAP</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>SSEP</td>
<td>Bilateral delay</td>
<td>Bilateral delay</td>
<td>Bilateral delay</td>
</tr>
<tr>
<td>Skin sympathetic response</td>
<td>Absent in all four limbs</td>
<td>Absent in all four limbs</td>
<td>–</td>
</tr>
</tbody>
</table>

Amp, amplitude; Lat, distal latency; CV, conduction velocity; CAP, compound action potential; AP, action potential; Tib Ant, tibialis anterior; EDC, extensor digitorum communis; APB, abductor pollicis brevis; VEP, visual evoked potential; BSEAP, brain stem auditory evoked potential; SSEP, somatosensory evoked potentials from the lower limb.
Our cases are remarkable in that adrenocortical function was preserved into the fifth decade confirming that this feature of Allgrove's syndrome may be absent or occur later in the course of the disease. In one patient aldosterone responses alone appeared impaired. This has been reported in younger patients.

Autonomic dysfunction contributed significantly to morbidity in two of our cases and seemed to involve cardiac parasympathetic, cholinergic sympathetic (sudomotor), cranial, and pelvic parasympathetic (pupillomotor, lacrimotor, erectile function). Previous authors have suggested dysfunction is limited to cholinergic neurones in Allgrove's syndrome. Two of our cases also had impaired sympathetic vasoconstrictor (noradrenergic) function, although the possibility of an additional central autonomic defect cannot be ruled out.

Sural nerve biopsies in two patients showed changes consistent with axonal degeneration and loss of both myelinated and unmyelinated fibres with no amyloid deposition. The aetiology of the neuropathy in Allgrove's syndrome is obscure. Previous authors have suggested it may result from a defect of ACTH receptors on neurones/glia with secondary demyelination, but further work has not provided support for this theory.

In two of our cases a second sibling was affected by a familial glucocorticoid deficiency with achalasia of the cardia and deficient tear production. The aetiology of the neuropathy in Allgrove's syndrome is obscure. Previous authors have suggested it may result from a defect of ACTH receptors on neurones/glia with secondary demyelination, but further work has not provided support for this theory.

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