SHORT REPORT

Chronic cough in the Holmes-Adie syndrome: association in five cases with autonomic dysfunction

J Kimber, D Mitchell, C J Mathias

Abstract
The Holmes-Adie syndrome consists of unilateral or bilateral tonic pupils with near light dissociation and tendon areflexia. It is associated with autonomic disturbances affecting sudomotor and vasomotor function. Five such patients are reported on who also had a troublesome chronic dry cough, which was of unknown aetiology and was resistant to a range of treatments. The cough may be related to involvement of afferent or efferent pathways in the vagus. Chronic cough may be an accompaniment in the Holmes-Adie syndrome, like other forms of autonomic dysfunction.

Keywords: Holmes-Adie syndrome; autonomic dysfunction; cough

The Holmes-Adie syndrome (tonic pupil, near-light dissociation, and tendon areflexia) has been associated with a range of autonomic disturbance that includes orthostatic hypotension, impairment of cardiovascular reflexes, segmental and generalised hypohydrosis (Ross’s syndrome), carotid gustatory syndrome, and chronic diarrhoea. Limited histological studies in this condition indicate loss of ganglion cells in the parasympathetic ciliary ganglia which correlates with the pupillary abnormalities. Degeneration of the dorsal root ganglia or the fasciculus gracilis and cuneatus may explain the absent tendon reflexes. Autonomic dysfunction has been variously ascribed to lesions of both afferent and efferent sympathetic and parasympathetic neurones.

Although Holmes-Adie syndrome is considered to be benign and of unknown aetiology, troublesome symptoms may result from autonomic dysfunction; these include hyperhydrosis, heat intolerance, and syncope. We report a further feature in five patients with Holmes-Adie syndrome in whom there was a persistent dry cough, of unknown aetiology.

Methods and results
The patients are described individually. None had a history of exposure to irritant chemicals or impairment of exercise tolerance. All had bouts of paroxysmal dry coughing during the day, with lesser frequency at night. Cough was not provoked by lying flat. None was taking angiotensin converting enzyme inhibitors, and all underwent detailed cardiovascular autonomic function tests with thermoregulatory sweat testing (table 1). Pulmonary function testing and other investigations to determine possible causes for their cough are shown in table 2. In all the chest radiograph was normal. Thorax CT was questionably abnormal in patient 4, with peribronchial shadowing in the right upper zone that may have represented localised bronchiectasis, but bronchoscopy was normal. There were no features of gastroesophageal reflux and all were treated with H2 antagonists with no benefit. In the three who underwent cine-video fluoroscopy, there was no evidence of laryngeal aspiration. In three, capsaicin challenge indicated increased cough sensitivity. Laboratory testing to exclude secondary causes of autonomic dysfunction included glucose, erythrocyte sedimentation rate, serum electrophoresis, autoantibody screen, treponemal serology, and brain MRI; and the results were normal in each case. Where Holmes-Adie syndrome was not previously documented, patients also underwent pupillography and EMG (to show absent or reduced H waves) to confirm further the diagnosis.

Case reports
Patient 1 was a 39 year old white man, a non-smoker, who had been investigated when aged 21 for anisocoria and a tonic right pupil that was unresponsive to light. When aged 35 he developed hyperhydrosis affecting the right arm and trunk; a right cervical sympathectomy was performed with only minimal improvement. He then developed symptoms of postural hypotension and was bothered by frequent paroxysms of coughing that resulted in syncope on several occasions. There was no history of lung disease.

Examination indicated a dilated right pupil with near-light dissociation. Other cranial nerves were normal. Tendon reflexes were absent bilaterally. Sweating was marked in the right axilla. There were no respiratory system...
abnormalities. Cardiovascular autonomic testing suggested an afferent baroreceptor lesion. Paroxysmal coughing during head up tilt resulted in a fall in blood pressure from 151/99 to 98/70 mm Hg, inducing presyncope. He was treated with a high salt intake and fludrocortisone to improve orthostatic tolerance. Probanthine (15 mg thrice daily) and clonidine (25 mg thrice daily) were used to reduce hyperhidrosis but were stopped due to a dry mouth. Investigation of cough (table 1) indicated no identifiable aetiology. The cough was refractory to therapeutic trials of inhaled, nasal, and oral steroids, lignocaine spray and nebuliser, local anaesthetic lozenges, and oral H1 antagonists. The patient is currently on proprietary antitussive agents (containing morphine, capsaicin, and ipecacuanha).

Patient 2 was a 58 year old white woman, who had stopped smoking 15 years previously. When 51, she noticed inequality of pupils. Two years later she developed asymmetric facial sweating and paroxysmal coughing attacks, particularly while eating and during prolonged talking. She had occasional dysphonia but no wheezing and on investigation was diagnosed as having late onset asthma. Although bronchodilator treatment improved the dysphonia it did not prevent coughing. She had been hypertensive and noted that treatment with a high salt intake and fludrocortisone to improve orthostatic tolerance. Probanthine (15 mg thrice daily) and clonidine (25 mg thrice daily) were used to reduce hyperhidrosis but were stopped due to a dry mouth. Investigation of cough (table 1) indicated no identifiable aetiology. The cough was refractory to therapeutic trials of inhaled, nasal, and oral steroids, lignocaine spray and nebuliser, local anaesthetic lozenges, and oral H1 antagonists. The patient is currently on proprietary antitussive agents (containing morphine, capsaicin, and ipecacuanha).

**Table 1** Details of investigations of the respiratory and allied systems to determine the cause of cough

<table>
<thead>
<tr>
<th>Patient</th>
<th>Patient 2</th>
<th>Patient 3</th>
<th>Patient 4</th>
<th>Patient 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chest x ray film</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>CT thorax</td>
<td>Normal</td>
<td>—</td>
<td>Normal</td>
<td>Penbribronchial shadowing R upper zone</td>
</tr>
<tr>
<td>Spirometry (observed/expected (n (%)))</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Vital capacity (l)</td>
<td>5.4 (104)</td>
<td>1.9 (89)</td>
<td>2.4 (114)</td>
<td>2.9 (104)</td>
</tr>
<tr>
<td>FEV1 (l)</td>
<td>4.5 (110)</td>
<td>1.6 (90)</td>
<td>1.8 (126)</td>
<td>2.4 (109)</td>
</tr>
<tr>
<td>FEV1/FVC (%)</td>
<td>83 (108)</td>
<td>84 (91)</td>
<td>73 (112)</td>
<td>83 (98)</td>
</tr>
<tr>
<td>Peak expiratory flow rate (l/min)</td>
<td>710 (114)</td>
<td>450 (90)</td>
<td>315 (106)</td>
<td>475 (317)</td>
</tr>
<tr>
<td>Transfer factor (T1CO) (ml/min/mm Hg)</td>
<td>26 (74)</td>
<td>—</td>
<td>6.3 (105)</td>
<td>25 (109)</td>
</tr>
<tr>
<td>Bronchoscopy</td>
<td>Normal</td>
<td>—</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>Nasolaryngoscopy</td>
<td>Small pharyngeal pouch &quot;not functionally significant&quot;</td>
<td>Normal</td>
<td>Normal</td>
<td>Mild rhinitis</td>
</tr>
<tr>
<td>Cine-video fluoresc</td>
<td>Minimal pooling in valleculae and pyriform fossae.</td>
<td>Minimal reduction in tongue base contractions</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>OGD</td>
<td>Reduced tongue base contractions</td>
<td>tongue base monitory</td>
<td>base contractions</td>
<td>—</td>
</tr>
<tr>
<td>Capsaicin challenge</td>
<td>Increased sensitivity</td>
<td>Increased sensitivity</td>
<td>Increased sensitivity</td>
<td>—</td>
</tr>
</tbody>
</table>

ODG=oesophagogastroduodenoscopy; FEV1=forced expiratory flow in 1 minute; —=not performed; T1CO=transfer coefficient for carbon monoxide.

Patient 3 was a 59 year old white woman, a non-smoker with a 20 year history of postural dizziness. Previous investigation at the age of 40 had documented bilateral tonic pupils in association with tendon areflexia. Subsequently she developed facial anhidrosis and a chronic dry cough. Examination confirmed small irregular pupils with near light dissociation and tendon areflexia with no other CNS abnormality. Cardiovascular and respiratory systems were unremarkable. Cardiovascular autonomic testing indicated baroreceptor dysfunction (orthostatic hypotension, blocked blood pressure response to Valsalva’s manoeuvre, and exaggerated pressor responses). Investigations for cough identified no obvious aetiology. Treatment with proprietary antitussive drugs resulted in some improvement of symptoms.

Patient 4 was a 31 year old woman, a non-smoker of mixed (white-AfroCaribbeean) origin. On presentation at the age of 26 she had a 2 year history of posture induced syncope together with paroxysmal throbbing headaches and epistaxis provoked by prolonged recumbency. She had had mild asthma since childhood and used a salbutamol inhaler intermittently. Examination indicated a right tonic pupil with near light dissociation. Tendon reflexes in the lower limbs were reduced. There were no other neurological abnormalities. Cardiovascular and respiratory examination was normal. Cardiovascular autonomic testing confirmed orthostatic hypotension and pressor stimuli raised blood pressure at times to 250/120 mm Hg suggesting impaired baroreceptor afferent activity. Methyldopa (250 mg twice daily) reduced the paroxysmal hypertensive episodes. A prominent symptom was persistent dry coughing that was refractory to increasing bronchodilator treatment and this was investigated further (table 2). Treatment with inhaled and nasal steroids (fluticasone inhaler (200 µg twice daily); fluticasone nasal spray (50 µg/nostril daily)) partially improved her cough. Oral H2 (chlorpheniramine) and H1 antagonists (ranitidine) were of no further benefit. She continued to have an unremitting dry cough.
Orthostatic hypotension indicates a postural fall in systolic blood pressure >20 mm Hg on head up tilt. BP=blood pressure; HR=heart rate; bpm=beats/min.

The increased response may in part have been due to coughing, often induced by hyperventilation.

### Table 2  Details of cardiovascular autonomic function tests in patients

<table>
<thead>
<tr>
<th></th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
<th>Patient 4</th>
<th>Patient 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orthostatic hypotension</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Pressor responses</td>
<td>Exaggerated</td>
<td>Normal</td>
<td>Exaggerated</td>
<td>Exaggerated</td>
<td>Exaggerated</td>
</tr>
<tr>
<td>Hyperventilation (expected increase in heart rate: 15–21 bpm)</td>
<td>↓↑ (+33)*</td>
<td>↓↓ (↓6)</td>
<td>↓↑ (+40)*</td>
<td>↓↑ (+45)*</td>
<td>↓↓ (+2)</td>
</tr>
<tr>
<td>Sinus arrhythmia (expected increase in heart rate: 15–19 bpm)</td>
<td>3</td>
<td>17</td>
<td>&lt;5</td>
<td>10</td>
<td>&lt;5</td>
</tr>
<tr>
<td>Valsalva’s manoeuvre</td>
<td>Abnormal</td>
<td>Normal</td>
<td>Abnormal</td>
<td>Abnormal</td>
<td>Abnormal</td>
</tr>
<tr>
<td>Ratio (normal &gt;1.2)</td>
<td>1.09</td>
<td>1.63</td>
<td>1.07</td>
<td>1.16</td>
<td>1.13</td>
</tr>
</tbody>
</table>

*The increased response may in part have been due to coughing, often induced by hyperventilation. BP=blood pressure; HR=heart rate; bpm=beats/min.

Orthostatic hypotension indicates a postural fall in systolic blood pressure >20 mm Hg on head up tilt.

Patient 5 was a 66 year white woman who was a non-smoker. She had presented 12 years previously with a troublesome dry cough. She had also noticed a unilateral dilated pupil and began having episodic diarrhoea up to 12 times daily. Panendoscopy plus large bowel biopsies failed to identify a cause. She then also noted truncal hyperhydrosis and postural dizziness. She had dry eyes and hypostomia with no history of joint or skin inflammation and no reduction in exercise tolerance. Further serological investigation indicated positive anti-Ro antibodies, suggesting primary Sjögren’s syndrome. Examination showed near light dissociation of the right pupil and tendon areflexia. Truncal sweating was prominent bilaterally. There were no other abnormalities. Cardiovascular autonomic function tests indicated no abnormalities and investigation of her cough identified no obvious cause. She had been treated with numerous courses of antibiotics for presumed upper respiratory tract infections and with antihistin treatment and cough suppressants containing opiates without effect. Proprietary antitussive agents containing capsaicin are currently being assessed.

**Discussion**

Chronic cough is a common symptom and may be caused by various stimuli which range from neoplasms, inflammatory lung disease, and postnasal drip to increased cough sensitivity resulting from respiratory tract infections, asthma, gastro-oesophageal reflux, and drugs such as angiotensin converting enzyme inhibitors. Despite extensive investigation, a proportion of patients have no diagnostic reason to explain their cough. Our findings suggest that the Holmes-Adie syndrome may be a disorder associated with a chronic dry cough.

All our patients had typical ocular findings and tendon areflexia indicative of Holmes-Adie syndrome. They also had associated autonomic impairment, either cardiovascular or sudomotor, and this combination presumably reflects the interests of the referral units. All were investigated in detail for autonomic dysfunction and also for a cause for their coughing. Coughing was provoked in each patient by deep breathing or hyperventilation during autonomic testing, suggestive of increased cough reflex sensitivity. This was formally tested in three. All were investigated for asthma. One patient had had mild asthma as a child and another was a probable late onset asthmatic; asthma may cause chronic cough and increased cough sensitivity, but there was no relation between coughing and indices of airflow limitation and furthermore the cough did not respond to effective asthma treatment. Treatment with H₂ antagonists for occult gastro-oesophageal reflux, and H₁ antagonists and topical steroids for rhinitis, were of no benefit.

Consideration of the cough reflex pathways enables speculation on the mechanism of cough in these patients. Afferent pathways carry stimuli from receptors in the trachea, larynx, and larger bronchi to the dorsal medulla via the vagus nerve. The motor (efferent) reflex arc innervates the muscles of the larynx, diaphragm, and chest wall. Efferent pathways for bronchiolar smooth muscle and mucosal glands are also vagal in origin. In Holmes-Adie syndrome, cardiovascular and peripheral autonomic neuropathy, the cough response to mechanical stimulation and tendo reflex abnormalities suggest an impairment of the afferent reflex arc. This alone is an unlikely explanation as in animals with neuropathy induced by acrylamide (affecting vagal myelinated afferent fibres), and in diabetic patients with peripheral autonomic neuropathy, the cough response to chemical stimulation and citric acid is diminished. Whether impaired afferent activity resulted in a relative increase in vagal efferent traffic cannot be discounted.

An increased sensitivity of airway cough receptors is a possible explanation as cough reflex sensitivity using capsaicin challenge was increased in each of the three patients tested. This may have been due to changes in the local environment of sensory nerves in the airway, particularly bradykinin and prostaglandin production as has been noted in cough induced by angiotensin converting enzyme inhibitor. In the three patients who underwent cine-video fluoroscopic examination of swallowing, there were minor abnormalities of coordination that may have provided an abnormal stimulus for coughing; there was, however, no clear association with swallowing or eating except in patient 2, and in none was there evidence of laryngeal aspiration.

One patient responded to an oral and later an inhaled anticholinergic agent and the mechanism may have been through reduced vagally stimulated (muscarinic) effects on bronchial secretions. In patients with refractory epilepsy fitted with vagal nerve stimulators, a proportion experienced coughing corresponding to the stimulator “on” period. A therapeutic trial of anticholinergic agents, as in patient 2, may help to determine the vagal efferent component to the cough.

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There is uncertainty about the CNS component of the cough reflex. In Holmes-Adie syndrome there is no known evidence of a clear central abnormality, and although it cannot be discounted or tested this seems less likely. Although chronic cough is a common symptom, detailed investigation in each of our patients with Holmes-Adie syndrome showed no obvious aetiology for their cough; in four out of five patients the coughing seemed temporally related to the onset of other autonomic features. In Holmes-Adie syndrome the autonomic features affecting cardiovascular and sudomotor function favour a patchy involvement, with resultant exaggerated activity, such as increased pressor responses in some and compensatory hyperhidrosis in others. Whether a similar process affects the autonomic control of the respiratory tract and involves the pathways involved in the cough reflex in such patients can only be speculated on at this stage. However, in all five patients, chronic cough produced appreciable morbidity. We conclude that these findings indicate yet another troublesome symptom, the mechanism of which needs to be further elucidated, to add to abnormalities of the somatic and autonomic nervous system previously described in the Holmes-Adie syndrome.

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