CONCISE REPORTS

Respiratory abnormalities due to craniocervical junction compression in rheumatoid disease

R S Howard, F Henderson, N P Hirsch, J M Stevens, B E Kendall, H A Crockard

Abstract
Objectives—To assess the extent and severity of respiratory insufficiency associated with severe rheumatoid atlantoaxial dislocation and its relation to compression of the neuraxis.

Methods—Twelve patients with severe atlantoaxial dislocation due to rheumatoid disease were studied. Detailed clinical, CT myelography and respiratory assessment including nocturnal oximetry, were performed on all patients.

Results—All patients were severely disabled by their underlying disease but none had symptoms of hypventilation. All the patients with C1 compression had myelopathic features. Those with medullary deformation (moulding and/or stretch) had abnormal nocturnal oximetry whilst no significant desaturations were seen in the remaining patients. Postoperative studies showed resolution of nocturnal desaturations.

Conclusion—This study suggests that clinically unsuspected respiratory insufficiency may be common in patients with severe medullary compression associated with rheumatoid atlantoaxial dislocation. It emphasises the importance of careful respiratory monitoring including nocturnal oxygen saturation in patients with major atlantoaxial dislocation due to rheumatoid disease.


The atlantoaxial joint is frequently involved in patients with rheumatoid disease. The combination of ligamentous laxity, bone erosion and hypertrophic synovitis leads to both backwards movement of the odontoid process into the spinal canal and inflammatory soft tissue masses which may contribute to neural compression at the craniocervical junction, however, the extent of atlantoaxial subluxation as assessed radiologically, does not correlate with symptoms or signs of neural compression. The development of progressive cervical myelopathy, upward translocation of the odontoid process, severe cord compression and intermittent neurological disturbances are all poor prognostic features. The poor prognosis of conservatively treated patients with craniocervical compression had led to the development of various operative strategies. It is well recognised that apnoea and sudden death may occur as a consequence of medullary compression disorders in the region of the foramen magnum. The extent to which hypoventilation and apnoea contribute to the high mortality of rheumatoid atlantoaxial dislocation with neural involvement remains unknown.

Methods

Twelve consecutive patients with atlantoaxial dislocation due to rheumatoid disease, not necessarily with respiratory symptoms, were studied as part of an ongoing study of 255 patients undergoing odontoidectomy and posterior fixation. Detailed clinical, CT myelography and respiratory assessment including nocturnal oximetry, were performed on all patients. Six patients were studied postoperatively. Oxygenation was measured continuously using an Ohmeda 3700e pulse oximeter and transferred for subsequent computer analysis. The overnight baseline oxygen saturation (SpO₂) and the number of dips in SpO₂ >4% lasting longer than ten seconds were calculated and expressed per hour of recording. The presence of sleep was documented by the nurse observing the patient but polysomnography was not performed because of the severity of the underlying disability.

Eleven patients were investigated by plain radiography and water soluble CT myelography and one with MRI. The CT technique was as follows: slice thickness of 1-5 mm performed sequentially across the entire craniocervical junction from external auditory meatus to C3 with both flexion and extension of the head, and 5 mm thick slices were made with 2 mm overlap to examine the entire cervical spine from C2 to T1. Multiple sagittal, coronal and axial reformatted images were prepared from this data. The following measurements were made: minimum mid-sagittal diameter of the cord and medulla oblongata when either or both appeared compressed (cord cross sectional area, CSA), the atlanto-dental distance (ADD), and the degree of ventral migration of the dens (VMD) using the method described by Redlund-Johnell. The clivoaxial angle (CAA) was measured, defined as the obtuse angle made between a line drawn from the posterior...
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surface of the clivus intersecting with the upward projection of a line drawn along the posterior surface of the body of the axis.

Finally the degree of kyphosis of the lower brainstem or cervical cord was estimated by assessing the cervical flexure of the neural axis. Precise measurement of this angle was not possible in most studies because not enough of the upper rhombencephalon was visible. An angle of greater than 135° was defined as normal. These distances and angles are illustrated in figure 1. Moulding of the medulla oblongata by the vertically migrated dens was assessed visually and scored as present or absent.

Results

There were 11 women and one man, the mean age of presentation was 53.7 (50–78) years and the mean duration of RA was 22.9 (7–50) years. Eleven patients presented with neck pain, three with limb weakness and one with dysphagia. One patient was breathless at presentation but no respiratory symptoms were present in the remaining patients. There were signs of cord compression in four patients and bulbar weakness was present in a further patient with coexisting downward hindbrain herniation. There was a past history of mild obstructive airways disease in two patients and bronchiectasis in one. Ten patients were taking disease modifying drugs (Gold and penicillamine) and seven corticosteroids, all took non-steroidal anti-inflammatory drugs. Nine took analgesics containing codeine or dextropropoxyphene and six were taking hypnotics; none took opiates. Patients remained on the same medication during pre- and postoperative studies. None of the patients smoked. CT myographic findings are described in the table. Three patients had evidence of underlying pulmonary disease, such as, bronchiectasis, or chronic obstructive airways disease but in most of the other patients spirometric indices were significantly below predicted values. Nocturnal oximetry showed >5 hypoxic dips/hour in four patients. In two other patients there were intermittent prolonged periods of sustained desaturations lasting up to 20 minutes, suggestive of alveolar hypoventilation. The patient with bronchiectasis also had frequent and severe dips below a low baseline. Five patients (including the patient with chronic obstructive airways disease) had normal overnight oximetry. In general, the minimum value of SpO₂ was lowest in those patients with the most frequent dips and the longest duration 4% below baseline.

There was clinical evidence of myelopathy in the four patients with the most severe cervical cord compression and only one of these had moulding or abnormal kyphos of the medulla oblongata. None of the patients with predominant medullary involvement had myelopathy, regardless of severity, and only the patient with coexisting hindbrain descent had cranial nerve involvement. Three patients with myelopathy had mild abnormalities of nocturnal

Radiological features

<table>
<thead>
<tr>
<th>Patient</th>
<th>Site of maximal compression</th>
<th>Other sites of compression</th>
<th>Abnormal cervical flexure</th>
<th>Pannus</th>
<th>Molding of medulla</th>
<th>cord CSA (mm²)</th>
<th>VMD (mm²)</th>
<th>CCA (°)</th>
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<tbody>
<tr>
<td>1</td>
<td>Medulla</td>
<td>C1 (ant and post)</td>
<td>+</td>
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<td>122</td>
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<td>C1 (coexisting ACM)</td>
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<td>12</td>
<td>135</td>
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<tr>
<td>7</td>
<td>C1 (on flexion)</td>
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<td>+</td>
<td>70</td>
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Figure 1 Diagrams to show some of the measurements referred to in the text. Palato-occipital line (a–b); vertical migration of the dens (VMD) (c–d); the clivo-axial angle (CAA) was the obtuse angle between lines e–f and g–h; the cervical flexure of the neural axis was the obtuse angle between lines i–h and h–n.

Figure 2 Nocturnal oximetry tracing (A) pre- and (B) postoperatively for the first two hours of sleep. Trace (A) shows frequent desaturations of variable duration and severity. These have resolved completely postoperatively.
oximetry although none had more than five hypoxic dips/hour.

All patients with medullary deformation (moulding and/or abnormal kyphos) had abnormal nocturnal oximetry. No significant desaturations were seen in the remaining patients. Similarly the presence of marked VMD <15 mm and CAA <125° was associated with abnormalities of nocturnal oximetry. There was no significant association with CSA.

Postoperative studies were carried out on six patients. In three, previously normal nocturnal oximetry remained unchanged. In two patients frequent but mild dips in oximetry were abolished following transoral removal of the anterior compressive mass and posterior occipitocervical fusion. In one patient preoperative study showed frequent and severe nocturnal desaturations (29 desaturations per hour), decompression resulted in complete resolution of nocturnal desaturations (fig 2).

Discussion
Respiratory involvement in rheumatoid disease is commonly due to pulmonary fibrosis or pleural disease and there is also an increased incidence of pulmonary infections, bronchiectasis,1 micrognathia, temporomandibular and cricoarytenoid joint disease, causing upper airway limitation and obstruction sleep apnoea (OSA).13 Disordered breathing during sleep also may be related to lesions of the cranio-cervical junction such as atlantoaxial dislocation.12 13 The patients in the present series were elderly and had severe disabilities related to rheumatoid disease which prevented more detailed study of respiratory function.

In the present series nocturnal respiratory insufficiency was associated with an increased cervical flexure of the neural axis, vertical migration of the dens and the ciboaxial angle, but not with atlantoaxial dislocation. Stevens et al showed that sensorimotor long tract signs correlated with combined anterior and posterior compression of the first segment of the cord. Rogers et al showed that, in patients with vertical translocation of the odontoid process, long tract signs are common and loss of joint position sense is particularly associated with severe compression of the posterior aspect of the spinal cord at the craniocervical junction. In this series only three patients had major compression of the spinal cord. These data suggest that detectable respiratory abnormalities may be associated with deformation of the medulla leading to traumatic neuronal disruption whilst myelopathy is associated with cord compression. Thus, as in the present study, considerable respiratory impairment may occur without clinical evidence of cord compression.

Our study suggests that in rheumatoid disease, clinically unsuspected disturbance of nocturnal respiration can occur even in the absence of respiratory symptoms, focal brainstem signs or myelopathy. Furthermore in patients studied postoperatively there was resolution of pre-existing nocturnal desaturations following decompression, suggesting a causal relationship between compression at the craniocervical junction and nocturnal desaturations. In conjunction with routine tests of respiratory function nocturnal oximetry may be a valuable investigation in identifying patients with rheumatoid atlantoaxial disease who are at risk of sudden apnoeic death. Serious nocturnal hypventilation may also be an important indication for surgical intervention.

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