knowledge only one patient has been reported with an association between obstructive sleep apnoea proven by a sleep study and adult acquired micrognathia due to rheumatoid arthritis. However, in that case there was no upper airway imaging. Our patient showed an extremely reduced upper airway area (21 mm² and 68 mm² at the oropharyngeal and hypopharyngeal levels, respectively) which may have been a major factor in the collapse of the upper airway.

The destruction of the cervical spine by rheumatoid arthritis, leading to potential compression of the cord or medulla, has been advocated as a causative factor for central sleep apnoea. Although our patient had an atlantoaxial dislocation and a subluxation of C3–C4, she had predominantly obstructive apnoeas, a normal ventilatory response to carbon dioxide, and no brainstem compression on the CT scan. This mechanism is therefore unlikely to explain the sleep apnoea syndrome in this case.

Sleep fragmentation in rheumatoid arthritis with marked disruption of sleep continuity can occur without sleep apnoeas as a result of pain, periodic leg movements, depression, and effects of drugs. Sleep fragmentation could favour respiratory instability and sleep apnoea including obstructive events.

In the present study, however, the abnormalities of the upper airway can reasonably be considered to be the main cause of the obstructive sleep apnoeas.

In clinical practice it is important to consider the diagnosis of sleep apnoea in patients with rheumatoid arthritis with temporomandibular joint destruction or cervical spine lesions who present with sleepiness, snoring, and/or cardiovascular disease. Prospective studies to assess the incidence and causative factors of sleep apnoea in large groups of patients with rheumatoid arthritis are needed.


Paradoxical vocal cord adduction in an adolescent with cystic fibrosis

P Shiels, J P Hayes, M X FitzGerald

Abstract

Many patients with cystic fibrosis have symptoms of dyspnoea and wheeze which are responsive to treatment with bronchodilators. An adolescent woman with cystic fibrosis is described who presented with inspiratory stridor and in whom the classical features of paradoxical vocal cord adduction were found.

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Keywords: cystic fibrosis, vocal cord adduction, airways obstruction.

Paradoxical vocal cord adduction is a very rare functional disorder caused by inappropriate adduction of otherwise normal vocal cords. Typically, it affects women under 40 years of age who often have a background of employment in health care. The upper airways obstruction associated with this condition may be mistaken for asthma and symptoms may persist in spite of varying therapeutic interventions. We describe here an adolescent woman with cystic fibrosis with the classical features of paradoxical vocal cord adduction – an association, to our knowledge, not previously described.

Case report

A 17 year old adolescent woman was admitted in July 1993 for assessment of chronic, predominantly dry, daytime and nocturnal cough with mucus clearing. Cystic fibrosis was diagnosed at birth and she attended the Adult Cystic Fibrosis Unit at St Vincent's Hospital, Dublin from 1991. Features of her illness included chronic bronchiectasis, multiple nasal polyps, pancreatic insufficiency, and recurrent meconium ileus equivalent. Her sputum was
Paradoxical vocal cord adduction in cystic fibrosis

Flow-volume loop showing severe inspiratory obstruction.

Paradoxical vocal cord adduction is an unusually rare condition due to inappropriate adduction of the vocal cords. It usually affects young adult women working in health care-related jobs. While functional airways obstruction is most often the result of inappropriate adduction of the vocal cords, it may also be caused by inappropriate movement of the pharyngeal wall muscles. This is considered to be an hysterical conversion reaction which may be responsive to a wide range of psychological therapies. Although initial reports suggested that it was a discrete isolated disorder, we have recently described a number of patients in whom paradoxical vocal cord adduction co-existed with documented asthma, and in whom objective evidence of refractory symptomatic paradoxical vocal cord adduction persisted for over a decade in spite of a wide range of psychotherapeutic interventions. This present report further extends the spectrum of pulmonary disorders in which paradoxical vocal cord adduction may be found.

Many patients with cystic fibrosis have symptoms of wheeze and shortness of breath with a positive response to inhaled bronchodilators which may vary with pulmonary exacerbations of the disease. To our knowledge there have been no previous reports of paradoxical vocal cord adduction in patients with cystic fibrosis. One of the puzzling psychodynamic features of this disease is its virtually exclusive occurrence in young women, the majority of whom have been involved in health care-related occupations. Our patient, like many individuals with cystic fibrosis, had been introduced to the hospital milieu from an early age and had many hospital admissions. Interestingly, the patient reported here had been heavily involved in lifetime courses, had spent considerable time as a beach lifeguard, and had expressed a desire to be a medical laboratory technician. This functional disorder is obviously rare, but it is clear that it could be easily overlooked in patients with cystic fibrosis who may present with various disorders of the upper and lower airways including recurrent respiratory infections, asthma, oropharyngeal candidiasis, and dysphonia due to inhaled steroid. We believe that the diagnosis of paradoxical vocal cord adduction should be considered in patients with cystic fibrosis in whom atypical chest symptoms, hoarseness, or stridor persist, and we suggest that appropriate physiological and endoscopic assessments are carried out.

**Discussion**

Paradoxical vocal cord adduction is an unusually rare condition due to inappropriate adduction of the vocal cords. It usually affects young adult women working in health care-related jobs. While functional airways obstruction is most often the result of inappropriate adduction of the vocal cords, it may also be caused by inappropriate movement of the pharyngeal wall muscles. This is considered to be an hysterical conversion reaction which may be responsive to a wide range of psychological therapies. Although initial reports suggested that it was a discrete isolated disorder, we have recently described a number of patients in whom paradoxical vocal cord adduction co-existed with documented asthma, and in whom objective evidence of refractory symptomatic paradoxical vocal cord adduction persisted for over a decade in spite of a wide range of psychotherapeutic interventions. This present report further extends the spectrum of pulmonary disorders in which paradoxical vocal cord adduction may be found.

Many patients with cystic fibrosis have symptoms of wheeze and shortness of breath with a positive response to inhaled bronchodilators which may vary with pulmonary exacerbations of the disease. To our knowledge there have been no previous reports of paradoxical vocal cord adduction in patients with cystic fibrosis. One of the puzzling psychodynamic features of this disease is its virtually exclusive occurrence in young women, the majority of whom have been involved in health care-related occupations. Our patient, like many individuals with cystic fibrosis, had been introduced to the hospital milieu from an early age and had many hospital admissions. Interestingly, the patient reported here had been heavily involved in lifetime courses, had spent considerable time as a beach lifeguard, and had expressed a desire to be a medical laboratory technician. This functional disorder is obviously rare, but it is clear that it could be easily overlooked in patients with cystic fibrosis who may present with various disorders of the upper and lower airways including recurrent respiratory infections, asthma, oropharyngeal candidiasis, and dysphonia due to inhaled steroid. We believe that the diagnosis of paradoxical vocal cord adduction should be considered in patients with cystic fibrosis in whom atypical chest symptoms, hoarseness, or stridor persist, and we suggest that appropriate physiological and endoscopic assessments are carried out.

Paradoxical vocal cord adduction in an adolescent with cystic fibrosis.

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