Pulmonary infiltrates and abdominal colic pain in a patient with a connective tissue disorder

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Case report
A 32 year old white women was admitted to our hospital complaining of long term fever and fatigue. She reported recurrent episodes of fever lasting 7 to 10 days accompanied by joint pain and weakness during the past four months. Subsequently she developed a productive cough and pleuritic chest pain on the right side. She had not travelled recently or been exposed to contagious diseases. She had not noticed any skin lesions, but on specific questioning reported photosensitivity and Raynaud’s phenomenon. She had a history of several lung infections and urolithiasis. She had had two normal deliveries and no miscarriages. There was a family history of interstitial lung disease.

On admission she was febrile (37.8°C). Physical examination revealed a conjunctival injection and bilateral submandibular lymph nodes. Wheezes were audible in the right lung base. Cardiac auscultation and abdominal palpation were unremarkable. Joint examination showed tenderness of proximal interphalangeal joints.

Laboratory data included the following: erythrocytes 4.75 x 10^{12}/l, haemoglobin 128 g/l, platelets 200 x 10^{12}/l, leucocytes 6.8 x 10^{9}/l and erythrocyte sedimentation rate (ESR) 70 mm 1st h. Serum biochemistry: creatinine 110 µmol/l (normal range 60–110), calcium 2.42 mmol/l (2.1–2.5) and phosphorous 0.69 mmol/l (0.97–1.45). Urine density was 1010, pH 7 and there were 12–20 erythrocytes/high magnification field and amorphous carbonates in the sediment. Immunoglobulins: IgG 22 g/l (8–16), IgA 2.23 g/l (1–3), IgM 1.24 g/l (0.8–2.5). Results were negative for HIV, Epstein-Barr serology, tuberculin test and direct examination of acid-alcohol resistant mycobacteria in sputum.

A chest radiograph showed bilateral interstitial infiltrates at the lung bases (fig 1) and thoracic computed tomography confirmed the lesions (fig 2). Bronchial lavage cultures demonstrated a growth of an ampicillin resistant *Haemophilus influenzae*. She was given cefotaxime, 2 g three times daily.

During admission she developed an acute episode of abdominal pain accompanied by hypotension and vomiting. Plain radiographs (fig 3) showed multiple calcifications in both renal silhouettes. Uromgrams revealed a functional loss of the left kidney (fig 4) and ultrasonography disclosed a moderate hydronephrosis, which was reversed after extracorporeal lithotrisy. A 24 hour urine analysis demonstrated hyperuricaemia (47.6 mmol/24 h), alkaline pH and hypocitraturia of 72 mg/24 h (normal range > 320 mg/24 h), with normal oxalate and calcium concentration. Plasma bicarbonate was 18 mmol/l.

Despite prolonged antimicrobial treatment with resolution of chest abnormalities the fever persisted and synovitis appeared in proximal interphalangeal joints. Rheumatoid factor as determined by nephelometry was 300 IU/ml (normal below 40), and C reactive protein was 4.8 g/l (normal below 1.0). Serological markers of viral hepatitis were negative. ANA were negative by immunofluorescence in Hep-2 cells and complement proteins were normal: C3 98 g/l (80–140), C4 31 g/l (12–33) and B factor 37 g/l (22–48). Immunodiffusion demonstrated antibodies to the nuclear extractable antigens Ro and La. Schirmer’s test was markedly diminished (1 mm after 10 minutes in both eyes). Thyroid stimulating hormone (200...
µUI/l) and free thyroxine (12 ng/l) were in normal range. A labial salivary gland biopsy showed inflammatory infiltration of the tissue yielding a lymphocytic focus score of > 1.0 per 4 mm² (fig 5).

The final diagnosis was primary Sjögren’s syndrome (SS), interstitial pneumonia attributable to *H influenzae* and urolithiasis favoured by distal renal tubular acidosis. The patient was discharged prescribed non-steroidal anti-inflammatory drugs (NSAIDs) and citrates.

During follow up the patient continued to complain of asthenia and joint pain, with recurrent episodes of arthritis, and maintained an ESR persistently raised. As NSAID treatment failed to obtain symptomatic relief deflazacort was introduced at low doses. At about this time she developed an enlargement of the right parotid gland, but surprisingly did not complain of dryness. Despite continuous treatment with citrate she had several renal colic episodes, ending up in medullospongiosis and nephrocalcinosis, with slight impairment of renal function. She had a pregnancy and delivered a premature baby who had a complete atrioventricular block requiring a pacemaker.

**Discussion**

**DIAGNOSTIC DIFFICULTIES IN SS**

It is not unusual for a rheumatologist to be consulted on a case of fever, joint pain and malaise, while the patient actually has an infectious disease, such as a bronchopneumonia. Besides, patients with connective tissue disorders are predisposed to infections, and this should be borne in mind. The cluster of symptoms presented by our patient suggested initially either a bacterial pneumonia or tuberculosis, the latter being highly prevalent in Spain. However, the onset of symmetric proximal interphalangeal joint arthritis prompted us to search for a connective tissue disorder. Viral arthritis and hypothyroidism were also considered, as they are common causes of mild joint inflammation and fatigue. In fact, thyroid function should be examined in SS patients with extreme tiredness, because autoimmune thyroiditis has been reported in up to 10% of SS patients. The diagnosis of primary SS was established after demonstrating exocrine involvement by Schirmer’s test and salivary gland histopathology together with a suggestive autoimmune profile.

SS is an autoimmune disease characterised by xerostomia and keratoconjunctivitis sicca (KCS). Although it is the most common connective tissue inflammatory disease, patients are correctly diagnosed on average 10 years after the onset of the disease, and usually after several medical consultations.

The typical SS patient is a woman (female/male ratio being 9 to 1) who presents with insidious symptoms and later develops manifestations of exocrine impairment, which make the condition easily recognised. However, there are patients who show features of generalised autoimmune disease, and only further studies reveal a subclinical exocrine involvement. Finally, the most challenging situation in diag-
nostic terms occurs in patients presenting with parenchymal internal organ disease without obvious surface exocrine disease. Indeed, a wide variety of onset forms have been described in SS (table 1). Raynaud’s phenomenon, cytopenic events, pulmonary disease and synovitis are the most common non-diagnostic associated features, while muscular, hepatobiliary or renal involvement are less frequent. Our patient illustrates the latter situation as she presented with sequential lung and renal involvement, which are considered as exocrine internal organ complications of SS, preceding any symptoms of xerostomia and KCS. This particular course of the disease was responsible for a delay in diagnosis.

DIFFERENTIAL DIAGNOSIS

SS and systemic lupus erythematosus (SLE) are diseases that share many pathological and serological features and may therefore be difficult to distinguish. Although photosensitivity is a characteristic of lupus, the absence of other typical manifestations of SLE in addition to the negativity of anti-DNA antibodies excluded the diagnosis of SLE with secondary SS. These two conditions may also overlap resulting in a disease known as the anti-La syndrome, which is identified by this serological marker. The anti-La syndrome could account for the patient’s symptoms, including distal tubular acidosis, but the typical rash and marked hypergammaglobulinaemia of the overlap syndrome were not evident in this case.

Additionally, conditions able to produce exocrine gland enlargement were ruled out by histopathology. These include granulomatous diseases such as sarcoidosis, deposition of amyloid or lipids, necrosis attributable to abnormal proteolytic activation, and neoplastic invasion, currently in cases of mucous associated lymphoma.

PULMONARY DISEASE

The percentage of reported cases of lung involvement in SS varies widely reaching 30% or even 55% in some series. Spirometric alterations of all sorts have been described, although overt clinical manifestations are rare. In general, it seems that hyperinflation—or increase in residual volume—is the most frequent abnormality and it seems to be caused by lymphocyte infiltration of small airway epithelia. It progresses to interstitial lung fibrosis in 4% of patients and, very seldom, to a more specific lymphoid interstitial pneumonitis. The pulmonary radiological pattern observed in our patient could have been interpreted as the start of an interstitial disease, but cultures of bronchial specimens and resolution of the infiltrates with antibiotic treatment favoured the diagnosis of bronchopneumonia. Although H influenzae is not a common infectious agent in healthy adults, hosts need not have a predisposing underlying disease. This microorganism produces acute inflammation of the airway mucosa that can occasionally penetrate the bronchial wall and cause a typical bronchopneumonia, as in this case.

On the other hand, respiratory tract dryness impairs clearance of the thickened secretions from the small airways facilitating lung infections in patients with SS.

RENAL INVOLVEMENT

Renal manifestations of SS can be multiple as shown in table 2. Although membranoproliferative glomerulonephritis has been described and should be considered in the case of rapidly deteriorating renal function—the main targets of CD4+ cells are tubular epithelia and surrounding interstitium. The density of lymphoid infiltration correlates with...
severity of tubular abnormalities, which vary from a loss of urinary concentration and latent tubular acidosis to overt acidosis, Fanconi’s syndrome or even atrophic fibrosis of the epithelium.

The basic molecular mechanism for acidification defects seems to be the absence of H+-ATPase in the intercalated cells and a subsequent impairment of proton secretion. However, anti-intercalated cell autoantibodies have not been demonstrated in SS.23 On the other hand, autoantibodies to renal collecting duct cells have been described in distal renal tubular acidosis complicating SS.26

Distal renal tubular acidosis has been reported in 15–67% of SS patients, while the latent form is demonstrated in another 25% by an overload of ammonium chloride.27 28 Our patient had a urinary pH of 7.0 despite low plasma bicarbonate concentration. In the absence of urinary infections by urea splitting organisms, coexistence of these findings is diagnostic of tubular acidosis. Distal tubular acidosis is accompanied by hypocitraturia and hypercalciuria, both of which are risk factors for calcium stone formation.29 Hypocitraturia is caused by an increase in absorption of citrates in the proximal tubules attributable to intracellular acidosis. Normalisation of the cytoplasmic pH is obtained by alkali supplements.29

An epidemiological study by Eriksson found that up to 16% of a population of women who formed hypocitraturic stones had anti-Ro antibodies30 and six of eight women with distal tubular acidosis and urolithiasis developed SS after an average of 15 years. Therefore, urolithiasis and distal renal tubular acidosis can precede the exocrinopathy for years or even constitute a manifestation of SS in the absence of subjective sicca symptoms.31 The investigation of leucocyte infiltration in salivary glands could be informative in this group of presymptomatic patients. This could help towards establishing an aggressive antirheumatic drug regimen in women with distal renal tubular acidosis and urolithiasis at the pre-sicca syndrome stage to modify the natural evolution of stone forming disease.32

The development of distal tubular acidosis is worsened by hypokalaemia,33 nephrocalcinosis34 and even osteomalacia,34 which can be avoided by preventive treatment with a daily lifelong alkali replacement. In the long term, urolithiasis, along with repetitive urinary infections, can lead to a decrease in glomerular filtration rate and to chronic renal failure.

NEONATAL LUPUS

It is well known that women who have antibodies to Ro or La are at risk of giving birth to children with neonatal lupus, regardless of their clinical disease, with a relative risk of suffering a congenital complete heart block of 500.35 This irreversible manifestation is generally detected between 16 and 24 weeks of gestation, and may be associated with myocarditis.36 37

THERAPEUTIC CONSIDERATIONS

The natural course of SS is poorly understood because of the absence of markers of disease activity.38 The existence of multiple autoantibodies in the sera of the patients is characteristic, but their significance remains unknown.39 Although some of them, such as anti-Ro and anti-La antibodies, are pathogenic for fetus heart, evidence is lacking of their participation in humoral and/or cell mediated immune reactions responsible for tissue damage.40 Some patients show raised levels of acute phase proteins, which could reflect cytokine mediation of tissue damage, but this hypothesis is not completely demonstrated. ANA titres, serum viscosity41 and other serological measurements have been found to correlate with disease activity in several series of patients, but they are not reliable as outcome measures. In fact, in many of these patients lymphoid proliferation remains localised to glands, with sicca syndrome being the sole expression of the disease.32

As a result, SS treatment is empirical and symptomatic. It is focused on the relief of the sicca syndrome and other symptoms that accompany the disease rather than seeking remission of the process (table 3). Immunomodulatory drugs are reserved for patients with severe extra-glandular manifestations as it is not known whether they are capable of modifying lymphocytic activation in this condition.42

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Sjögren’s syndrome is a complex autoimmune disorder characterized by inflammation of the exocrine glands, such as those in the mouth and eyes, leading to symptoms like dryness, fatigue, and joint pain. The syndrome is often associated with other autoimmune conditions, such as rheumatoid arthritis and lupus. It is named after Dr. per Oscar Sjögren, who described it in 1933.

The presence of autoantibodies, particularly against RNA polymerase III, and genetic factors play a role in its development. Risk factors include age, gender (females are more commonly affected), and family history. The disease can present at any age, but it typically appears in middle adulthood.

Diagnosis is often based on clinical symptoms and serological tests for autoantibodies. Treatment is symptomatic, with medication used to manage pain, fatigue, and other symptoms. The goal is to manage the disease while minimizing the risk of complications such as lymphoma.

Prevention strategies are limited, as the cause is not fully known. However, maintaining a healthy lifestyle, avoiding smoking, and managing stress may help reduce the risk of developing autoimmune diseases like Sjögren’s syndrome.