The progressive systemic sclerosis/systemic lupus overlap: an unusual clinical progression

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Abstract
Three patients with the unusual combinations of discoid lupus, systemic lupus erythematosus (SLE), and progressive systemic sclerosis (PSS) are reported. The first patient developed PSS eight years after a diagnosis of discoid lupus had been made and this was complicated by myositis six years later. The second patient developed PSS more than 20 years after being diagnosed as having SLE. The third patient developed SLE with predominant features of urticarial vasculitis six years after PSS. Mild myositis also ensued. There were no antibodies to U1RNP demonstrable in any of these patients. The clinical progression of SLE to PSS or vice versa in the absence of features of mixed connective tissue disease is distinctly uncommon.

Although the connective tissue diseases can usually be clinically and serologically defined as distinct and separate entities, such as rheumatoid arthritis, systemic lupus erythematosus, progressive systemic sclerosis, and dermatomyositis, overlap between these conditions may be encountered. A specific overlap termed mixed connective tissue disease is one such condition. Others are simply termed undifferentiated connective tissue disease. Sjögren's syndrome, fibrosing alveolitis, and Raynaud's phenomenon may accompany all of the above but may also exist on their own as 'primary' conditions.

The association of lupus and scleroderma (with or without myositis), one condition merging into the other, without the presence of antibodies to U1RNP or any clinical features of mixed connective tissue disease, is distinctly uncommon. We report three such patients seen over the past five years.

Case reports
PATIENT ONE: DISCOID LUPUS DEVELOPING INTO PROGRESSIVE SYSTEMIC SCLEROSIS WITH LATE COMPLICATING POLYMYOSITIS
A 48 year old white woman of Austrian ancestry had a single discoid lesion aged 21 which was confirmed as being discoid lupus erythematosus in 1975 at the age of 34. Clinical details are sketchy, but skin manifestations had predominated. At this time her antinuclear antibodies were positive (1/320). A biopsy specimen from non-lesional skin showed positive immunofluorescence for IgG and complement, and a lesional biopsy specimen showed typical histological features of discoid lupus erythe-
Figure 2  Patient No. 1.  Radiograph of the hands showing resorption of the terminal phalanges.

was normal but the erythrocyte sedimentation rate (ESR) was raised at 44 mm/h. The antinuclear antibody titre was still positive (1/320), immune complexes were increased (IgG 102 mg/l), she had negative latex and Rose-Waaler tests for rheumatoid factor, but surprisingly antibodies to Sm were present. The serum IgG concentration was raised (17·9 g/l) (normal 5·3–16·5 g/l). She was given diclofenac sodium 100 mg twice a day for the joint pains. Throughout 1988 and 1989 various other non-steroidal anti-inflammatory drugs were prescribed (ketoprofen, indomethacin) with no major clinical effects. A chest radiograph was normal, but radiography of her hands now showed resorption of the terminal phalanges (fig 2). There was no soft tissue calcification evident. In October 1989 she complained of muscle pain and her muscle enzymes were raised. Electromyography showed fibrillation and small polyphasic units characteristic of inflammatory muscle disease. A diagnosis of myositis was made. Her creatine kinase was 608 U/l (normal <250), and hydroxybutyric dehydrogenase 238 U/l (normal <182). The ESR was raised at 82 mm/h and the rheumatoid factor test was also positive. Tissue typing showed her to be A11,28, B14,56, C4,5, Bw6, and DR1 and 6. The muscle symptoms have responded to prednisolone 5 mg twice daily.

was treated with prednisolone (60 mg/day) and cyclophosphamide (125 mg/day orally) with improvement. Cyclophosphamide was stopped after three months because of the development of haemorrhagic cystitis.

He was lost to follow up until 1982 when he was admitted with right middle lobe pneumonia. Extensive skin lesions of chronic discoid lupus affecting the scalp, face, and upper trunk with scarring and alopecia were noted. A loud pericardial friction rub was noted, and echocardiography confirmed a pericardial effusion.

The haemoglobin was reduced at 82 g/l, the total white cell count was 11·3×10^9/l with 17% lymphocytes, and platelets were 729×10^9/l. The ESR was 125 mm/h. The urea was 3·8 mmol/l, creatinine 71 μmol/l, with an uncorrected clearance of 132·3 ml/min and a 24 hour urinary protein excretion of 0·12 g. The antinuclear antibody titre was low positive (1/10), and anti-DNA antibodies raised at 10 mg bound DNA/ml serum (normal 0–5). Radiographs of the hands were normal. He was treated with prednisolone (40 mg/day) and antibiotics for his infection.

In 1985 he presented to the outpatient clinic with pleuritic chest pain and arthralgia. Synovitis of the wrists and the metacarpophalangeal and proximal interphalangeal joints of the hands was noted, and radiographs showed juxta-articular osteoporosis, but no erosions. Sclerodactyly was noted for the first time. The antinuclear antibody titre was positive (1/100; 1/500)(homogeneous), and anti-DNA antibodies were raised at 18 mg bound DNA/ml serum. Treatment with chloroquine phosphate (250 mg/day) was started for control of his arthritis.

Throughout 1986 and 1987 he continued to have pain in the hands and wrists, but no active arthritis was reported. The sclerodactyly became more marked, however, and he developed proximal scleroderma with skin changes extending onto the forearms. Marked resorption of the digits resulted in significant loss of hand function. Radiographs of the hands over a three year period showed progression of erosions and marked resorptive changes at the interphalangeal and metacarpophalangeal joints, as well as the wrists (figs 3A, B, and C).

The patient has never had Raynaud’s phenomenon. The nailfold capillaries could not be clearly seen owing to the pigmentation and thickening of the skin. There was no evidence of skin telangiectasia or calcinosis. He did not have dysphagia, and a barium swallow done in September 1988 was normal. Tolerance of exercise was normal and the chest was clinically and radiologically clear.

There had been no evidence of active lupus for the past two years. The skin lesions were quiescent, he did not have serositis, and the urine was clear. The most recent laboratory investigations showed haemoglobin 154 g/l, white blood cell count 7·6×10^9/l with a normal differential, platelets 275×10^9/l, ESR 12 mm/h, urea 4·6 mmol/l, and creatinine 77 μmol/l. A 24 hour urine collection was negative for protein, and the uncorrected clearance was 131 ml/min. Recent serological tests showed a negative
antinuclear antibody and anti-Ro antibody. Anti-DNA antibodies were normal (≤3 mg bound DNA/ml serum) and other antibodies to extractable nuclear antigens were also negative. The patient was once again lost to follow up and skin biopsy specimens were unobtainable.

PATIENT THREE: PROGRESSIVE SYSTEMIC SCLEROSIS DEVELOPING INTO SYSTEMIC LUPUS ERYTHEMATOSUS

The patient, a 42 year old white woman with no family history of note, had a six year history of polyarthritis and polyarthralgias affecting mainly the hands, thickening of the skin, particularly affecting the face, upper arms, and shoulders, accompanied by Raynaud’s phenomenon. Exertional dyspnoea had recently developed over one year as well as ‘indigestion’, which consisted of substernal burning and discomfort postprandially. Additionally, she was lethargic. She had a longstanding history of dermatographism.

When seen in 1985 she had the typical features of scleroderma—including shiny and taught skin over the digits and loss of digital creasing—accompanied by synovitis, particularly of the metacarpophalangeal joints of both hands. There were no associated pigmented changes, but microstomia was present. She additionally had livedo reticularis.

From 1987 she had an intermittent burning purpuric rash generalised in distribution, which was diagnosed as urticarial vasculitis, for which she was given a course of dapsone in 1986. Hair loss started some three years after the onset of her illness. She was given steroid treatment, but in 1987 developed steroid induced glaucoma. To reduce her steroid requirements, hydroxychloroquine and naproxen were added to her regimen for relief of her arthritic pains. Several ‘flares’ of disease had occurred over the six year period, which were associated with increased shortness of breath but without cough, and the dyspnoea was accompanied by severe arthralgias. These ‘flares’ had previously been treated with temporary increases in steroid dose. She had recently had more frequent episodes of retrosternal chest pains radiating to her neck and occurring particularly at night. She was admitted to St Thomas’s Hospital for assessment in April 1987. Alopecia was noted as well as a widespread fading urticarial rash affecting the limbs and trunk. There was a purpuric element present on the edges of the soles of the feet and toes. Microstomia was obvious but no sclerodactyly was present.

Investigations showed a normal blood count with an ESR of 17 mm/h and a raised platelet count of 519×10⁹/L. Coombs’ test was negative. The Cевичia DNA immunofluorescence test was positive. Antinuclear antibodies were positive at 1/320 and the DNA binding antibody was more than 97 U/ml (normal <25 U/ml). The Venereal Disease Research Laboratory test was negative. Immune complexes were raised at 116 g IgG/l (normal <49). The aspartate transaminase was minimally raised and the hydroxybutyric dehydrogenase activity was also increased. The creatine kinase was persistently normal. Liver function and renal function tests...
were normal. Electromyography showed evidence of a mild myositis with some polyphasicity. Radiographs showed normal heart size and lung fields. Barium swallow and meal examinations were normal. No erosive changes were evident in the hands, and there was no terminal resorption of bone and no calcinosis. There was minor subluxation of the distal interphalangeal joints of both little fingers. Respiratory function tests showed a mild restrictive defect with a significant reduction in gas transfer (transfer factor 5·3) (predicted mean value (SD) 8·5 (2·4)).

Several months later the Coombs’ test became positive. This positivity was accompanied, however, by a normal reticulocyte count and a haemoglobin concentration at the lower end of normal (125 g/l).

A diagnosis of SLE was made with complicating urticarial vasculitis, myositis, and intermittent Coombs’ positivity. Steroid treatment was continued. Postprandial epigastric pains persisted, but an endoscopy was normal. Ranitidine 150 mg twice a day was added and she was also given nifedipine 20 mg twice a day for the Raynaud’s phenomenon. Because of the recurrent chest pains an echocardiogram was performed in 1988. This showed the presence of mitral regurgitation, confirmed on Doppler scanning.

When seen in late 1988 she had developed marked telangiectasias of the face and hands but there was no evidence of a CREST syndrome, such as pigmentation or sclerodactyly, and the telangiectasia were ascribed to her long term steroid treatment. Microstomia was still present. Immunological testing now showed antibodies to extractable nuclear antigens (unidentified), and the rheumatoid factor was positive. The C4 concentration was reduced at 0·11 g/l (normal 0·16–0·45) with a low normal CH100 of 60% (normal 50–125) and C3 at 0·8 g/l (normal 0·7–1·8). Antimitochondrial antibodies were negative and the Coombs’ test was now negative. C1q precipitins were present in serum. A lymphopenia of 0·9×10⁹/l had developed (normal 1·5–3·5×10⁹/l).

The patient continues to have episodes of recurrent urticarial vasculitis unresponsive to most treatments.

Discussion

The three patients described present unusual overlap features of two well defined connective tissue diseases. Patient one originally presented with discoid lupus erythematosus and there was no convincing evidence of systemic disease. Eight years later she had features of PSS predominantly affecting the skin, and severe Raynaud’s phenomenon. Interestingly, in 1987, 12 years after the initial development of discoid lupus erythematosus and still with predominant features of PSS, antibodies to Sm were present, indicating that at least serologically there was some evidence of antibody production against ribonucleoproteins, not usually seen in PSS, indicative and diagnostic of coexisting SLE.

Patient two developed SLE at a young age. Eight years later he presented extensive features of discoid lupus erythematosus and then a clinical picture of PSS seemed to evolve with widespread resorption of digits. An unusual feature was the fact that he never experienced Raynaud’s phenomenon. He lived in a warm climate, however, and the fact that he was naturally racially pigmented might have made any observations of colour changes associated with vasospasm difficult.

Patient three developed typical features of SLE six years after manifesting PSS. The SLE was heralded by the appearance of the typical rash of urticarial vasculitis, which has persisted for three years and has been unresponsive to any form of treatment to date.

In patients one and three a myositis eventually complicated their clinical course.

Mixed connective tissue disease was originally described by Sharp et al5 and by Mattioli and Reichlin in the early 1970s.1 Patients with this disease seem to show features of SLE, polyserositis, and progressive systemic sclerosis, accompanied by a chronic arthritis at one time or another. There is a high incidence of Raynaud’s phenomenon, swollen or ‘sausage’ fingers, and tenosynovitis clinically. All these patients have high titres of antibodies to U1RNP and are negative for antibodies to double stranded DNA (dsDNA). A speckled pattern of antinuclear antibodies is a useful clue to the presence of antibodies to U1RNP. The incidence of renal disease and neuropsychiatric illness in this group seems to be low according to some studies. Younger patients with the condition tend to develop SLE, whereas older patients develop features of progressive systemic sclerosis. Many, however, remain in the category of mixed connective tissue disease.

The familial occurrence of SLE and scleroderma is well described. Flores et al reported eight families containing members with both disorders.4 Relatives with PSS, however, had at least one serological abnormality more characteristic of SLE, such as antibodies to Sm, anti-RNP, anti-dsDNA, haemolytic anaemia, leucopenia, hypocomplementaemia. Two members of different families had enough clinical evidence of both diseases simultaneously to classify them as such.

Other authors have also drawn attention to the coexistence of both PSS and SLE within families,5–7 and these patients are discussed in detail by Flores et al.4

Dubois et al reported PSS and SLE coexisting in the same patients.7 These authors reviewed the incidence of coexistent SLE and scleroderma, as well as cases reported by other authors, and were able to discover only 10 previously published cases. They recorded 12 of 78 patients with PSS from their own clinics, nine with PSS and three with morphea only. Definite skin changes of scleroderma were confirmed by biopsy in nine of the 12. Discoid lupus erythematosus lesions were present in four, pleurisy in six, pleural effusions in five, and pericarditis in five. Seven had a leucopenia (<4·5×10⁹/l) and the diagnosis of the two conditions in single patients was confirmed by necropsy in four.

In an appendix to their paper Dubois et al
then described several other patients in detail who seemed to have varying combinations of the two diseases; either PSS, later developing features of SLE, or the reverse. These patients closely resemble the patients we have reported in this paper.

Chorzelski and Jablonska also described three cases of coexistent PSS and SLE, one with discoid lupus erythematosus and two with SLE. Jablonska et al subsequently reviewed the coexistence of the two conditions and concluded that the association was most uncommon, occurring in less than 1% of cases of PSS.

In summary, therefore, although the association of the two diseases in the same patient may occur rarely and sporadically, the familial occurrence of the two conditions may also be seen, albeit uncommonly.

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