Secondary addition of methotrexate to partial responders to etanercept alone is effective in severe rheumatoid arthritis

J D Cohen, S Zaltni, M J Kaiser, M C Bozonnat, C Jorgensen, J P Daurès, J Sany


Etanercept, a soluble tumour necrosis factor α (TNFα) receptor fusion protein, may be combined with methotrexate (MTX) in severe rheumatoid arthritis (RA).1 Etanercept is generally given to patients treated with MTX who have an inadequate response. However, MTX could be introduced to patients who have already been treated with etanercept and who initially had a good response but subsequently a secondary failure. It was decided to evaluate, in an open prospective study, the clinical, biological, and functional benefit of the secondary addition of MTX in patients with RA with an inadequate response to etanercept alone.

PATIENTS, METHODS, AND RESULTS

A cohort of 93 patients with active and severe RA was observed for a mean duration of 8 months (range 1–20). The group comprised 75 women, 18 men, with an average age of 49.5 years (range 17–75), and positive rheumatoid factor in 67 (72%). All had previously been treated with a median of four disease modifying antirheumatic drugs (DMARDs). The mean disease duration was 12.4 years (range 3–42). Patients were treated with subcutaneous injections of etanercept alone, 25 mg twice weekly. Eighteen of the 93 patients had an inadequate response (did not reach the American College of Rheumatology (ACR) 20 criteria, prerequisite for inclusion in this cohort) with etanercept alone; we therefore added MTX to the treatment, no other DMARDs being allowed. The mean dose of added MTX was 15.5 mg a week. The dosage was given orally to 12 patients and intramuscularly to the remaining six. All 18 patients (16 female, two male, average age 51 years, mean disease duration 13.3 years, with a positive rheumatoid factor in 14 (78%) patients) had previously received MTX before etanercept alone, but it had produced an insufficient result. This failure with MTX was a requirement before starting etanercept. A prospective follow up of these patients took place every month for 3 months and from then on every 3 months during a 12 month period, with recording of the usual clinical and biological measures in order to obtain the ACR20, 50, and 70 and the Disease Activity Score 28 (DAS28). This enabled us to assess the efficacy of this addition. The Health Assessment Questionnaire (HAQ) was also used in this study.

Table 1 summarises the results obtained. The data show an improvement in disease activity, which was sustained through the 12 month follow up. A favourable response was obtained at 3 months with 10 (56%) achieving the ACR20, 5 (28%) the ACR50, and 2 (11%) the ACR70. At 12 months of this combination therapy, 12 (67%) had achieved the ACR20, 9 (50%) the ACR50, and 3 (17%) the ACR70. DAS28 decreased from 4.9 to 3.0 at 3 months and to 2.4 at 1 year, suggesting a significant clinical improvement in disability. The HAQ score changed from 1.7 to 1.1 at 1 year.

There was also a rapid therapeutic biological response, with a decrease in the erythrocyte sedimentation rate (ESR) and C reactive protein (CRP) of 50% after 3 months. No serious adverse events were reported and only two patients required antibiotics (bronchitis, urinary infection).

DISCUSSION

In summary, this open study suggests that the secondary addition of MTX to etanercept produced a marked improvement among patients who had had an inadequate response to etanercept alone after a first inadequate response to MTX given alone. To our knowledge, this is the first open study to evaluate the efficacy of MTX in addition to etanercept in a two step strategy. Previous studies have demonstrated the efficacy of etanercept alone compared with MTX,2 its efficacy in early RA3 or in addition to methotrexate,1 and, in particular, in juvenile idiopathic arthritis.4 This suggests that when MTX is added to the etanercept regimen patients with RA improve both clinically and biologically without any increase in side effects. The addition of MTX to the treatment of patients who respond incompletely to etanercept alone may be useful in the management of RA. It will be interesting to evaluate this strategy in further studies. Therefore, the preferred treatment

<table>
<thead>
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<th>Table 1 Results of secondary addition of MTX to etanercept</th>
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<td>MTX duration (months)</td>
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<tr>
<td>------------------------</td>
</tr>
<tr>
<td>ACR20 (%)</td>
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<td>ACR50 (%)</td>
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<td>ACR70 (%)</td>
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<tr>
<td>DAS28, mean</td>
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<td>Score HAQ, mean</td>
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<tr>
<td>ESR (mm/1st h), mean</td>
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<td>CRP (mg/l), mean</td>
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for RA may very well be the combination of MTX plus a TNF antagonist.

**Authors’ affiliations**

A Barsotti, M Cutolo, F Indiveri, A Sulli, M Ghio, G P Bezante, L Deferrari, C Craviotto, V Sebastiani, M Setti, G Filaci, F Puppo, A Barsotti, M Cutolo, F Indiveri

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**Blunted coronary flow reserve in systemic sclerosis: a sign of cardiac involvement in asymptomatic patients**

A Sulli, M Ghio, G P Bezante, L Deferrari, C Craviotto, V Sebastiani, M Setti, G Filaci, F Puppo, A Barsotti, M Cutolo, F Indiveri

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Cardiac disease is often present in systemic sclerosis (SSc), even if rarely of clinical significance. Therefore, we investigated the coronary flow reserve (CFR), by transthoracic contrast enhanced second harmonic Doppler echocardiography, a non-invasive method that might detect early heart dysfunction in patients with SSc even in the absence of clinical signs or symptoms.

Twenty-nine consecutive patients (2 male, 27 female, mean (SD) age 55 (14) years) affected by SSc, not complaining of signs or symptoms of cardiovascular involvement, were recruited. No further serious disease other than SSc was present. Eleven healthy subjects matched for age and sex (mean (SD) age 53 (5) years) were also evaluated as controls. Echocardiography was performed with an ultrasound unit using a broadband transducer with second harmonic capability in both B mode and Doppler modality. Levovist was used as the ultrasound contrast agent. The CFR, expressed as the ratio between hyperaemic and resting, both peak, diastolic velocity (PdvCFR) and velocity time integral (VtiCFR), was non-invasively assessed in the distal left anterior descending coronary. Peripheral microangiopathy was assessed by nailfold videocapillaroscopy (NVC), as previously reported.

All patients were found in sinus rhythm, without any significant ECG alteration. All ECG parameters were normal. The study showed a reduced CFR in 14/29 patients with SSc, when compared with the normal range of healthy subjects matched for age and sex (CFR >2.00). In particular, both PdvCFR and VtiCFR were strongly reduced in patients with SSc (mean (SD) 1.93 (0.56) and 1.81 (0.56), respectively) in comparison with controls (3.11 (0.72) and 2.83 (0.51), respectively) (p<0.0001). Furthermore, both PdvCFR and VtiCFR were significantly lower in patients with diffuse SSc (1.74 (0.46) and 1.59 (0.38), respectively) than in patients with limited SSc (2.39 (0.52) and 2.35 (0.38), respectively) (p<0.004 and p<0.001, respectively) (fig 1).

Nineteen patients (mean (SD) age 52 (13) years) and 10 patients (mean (SD) age 63 (12) years) had diffuse SSc (dSSc) and limited SSc (ISSc), respectively; the patients with dSSc were younger than those with ISSc (p<0.04).

Glucose serum levels were normal in all patients with SSc. No statistically significant correlation was found between CFR and history of smoking and cholesterol or triglyceride serum levels. Moreover, no statistically significant correlation was found between CFR and blood pressure values.

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**REFERENCES**


Incidentally discovered asymptomatic necrotising intra-abdominal vasculitis after peripheral gastric bypass surgery for morbid obesity

A Boungas, M Melachrinou, G Giannopoulos, N Meimaris, P Aroukatos, F Kalfarentzos, A P Andonopoulos

We describe the case of a 35 year old woman, who underwent peripheral gastric bypass type Roux-Y surgery for morbid obesity, and was incidentally discovered to have diffuse abdominal necrotising vasculitis, which has remained silent for at least 15 months.

CASE REPORT
The patient was referred to us because a biopsy of the appendix removed at surgery had disclosed necrotising vasculitis (fig 1), and her serology had shown positive antinuclear antibodies (ANA), 1/320 homogeneous. She had received bronchodilators in the past, amoxicillin clavulanate, oral contraceptives, and fenfluramine for a short time. No recent vaccination had been administered. She denied systemic symptoms, arthralgias or arthritis, skin or mucosal lesions, abdominal pain or rectal bleeding, sicca symptoms, or Raynaud’s phenomenon. After surgery she had lost 45 kg and, as expected, she had been experiencing mild abdominal cramps and diarrhoea shortly after meals. Physical examination was normal.

Routine laboratory investigations, including haematology, biochemistry, and urine analysis, chest x-ray examination, and a purified protein derivative skin test were normal. Besides the positive ANA, complete serology, including antineutrophil cytoplasmic antibodies perinuclear and cyttoplasmic, anti-proteinase 3, and anti-myeloperoxidase antibodies, and tests for hepatitis B and C virus, was unrevealing. Abdominal and cardiac ultrasound, mammography, and PAP test were normal. A biopsy of skin and muscle of the gastrocnemius did not show abnormalities. An abdominal multislice computed tomography-angiography was consistent with vasculitis of the polyarteritis nodosa (PAN) type, showing bead-like, concentric stenoses and dilatations of multiple branches of the superior mesenteric artery (fig 2).

The diagnosis of a medium and small, muscular-type, necrotising arteritis was made, albeit silent, and the patient followed up closely. Today, 15 months after surgery, she has remained clinically asymptomatic, without laboratory abnormalities, enjoying a normal life and approaching the ideal body weight.
DISCUSSION

Vasculitis of the PAN type, localised to the appendix or the gall bladder, has been accidentally found, usually after surgery.1 Its prognosis is almost always good, in contrast with vasculitis that is found when systemic PAN has already been diagnosed.1 Several possibilities can be suggested for the aetiology of the diffuse intra-abdominal vasculitis of our patient. Firstly, a relationship with the operation can be easily excluded because the disease was already present in the appendix before the procedure.

A second possibility may implicate the morbid obesity of the patient. Intra-abdominal processes such as pancreatitis and malignancy, but other systemic diseases also, such as infectious endocarditis, atrial myxoma, and drug abuse, have been reported to mimic the visceral angiographic appearance of vasculitis.2 No such problems were present, and furthermore, morbid obesity has never been reported to cause vasculitis.

The positive ANA may suggest systemic lupus erythematosus as a third possibility, but our patient had no other evidence to support this diagnosis.

Drug induced vasculitis may be a fourth possibility.3–5 In the vast majority of cases it affects the skin, but it may extend to internal organs. The diagnosis is usually one of exclusion and a timely relationship to the offending drug should be documented. Almost any drug can be blamed. However, it is rather unlikely that one of the drugs the patient had taken would have induced abdominal vasculitis, without skin manifestations.

Finally, it is quite likely that our patient had asymptomatic PAN, which will either remain as such, or will become manifest in the future. In such cases, clinicians should be aware of the possibility that a systemic necrotising vasculitis may, for some time, remain asymptomatic. Furthermore, mesenteric angiography, when used to help diagnose PAN in cases of multisystem clinical presentations, is not expected to disclose vasculitis when hepatic enzymes are normal, but even then the procedure may be of diagnostic help.

REFERENCES

Ultrasound detection of knee patellar enthesitis: a comparison with magnetic resonance imaging

M Kamel, H Eid, R Mansour

The diagnosis of enthesitis in clinical practice is difficult and usually based on conventional radiographic findings, which are not helpful in most cases. We previously reported that ultrasound (US) was sensitive in detecting peculiar pathological features of enthesitis around the heel. Furthermore, we have continued to study the efficacy of ultrasonographic diagnosis of enthesitis of other tendon and ligament insertion sites.

METHODS AND RESULTS
Sixteen patients (10 male, six female, mean age 45.6 years) with a diagnosis of seronegative arthropathy were recruited from the population for the study. Their mean disease duration was 6.3 years. They had seronegative arthropathy and knee enthesopathy without typical conventional radiographic evidence. An HDI 3000 ATL US machine (Advanced Technology Laboratories, USA), equipped with a 12 MHz linear transducer, was used to examine the knee patellar enthesis.

The present study produced interesting findings (figs 1A and B). The US images of the knee patellar enthesis showed loss of the normal fibrillar echo texture of the patellar tendon, no homogeneous pattern, blurring of the patellar tendon margins, irregular focal or generalised increased tendon thickness, and focal ill-defined tendon defects, with loss of their tightly packed echogenic dots. The US images clearly showed the definition of the patellar tendon margins, which were more precise and anatomically defined than the magnetic resonance (MR) images (figs 1C and D).

DISCUSSION
The US examination of the knee joint clearly detected the early calcification foci of the patellar tendons. However, the calcification process of the knee patellar ligament developed less often in the patients than the calcification of the Achilles tendon found in a previous report.

The process of fatty degeneration of the patellar tendon was detected early in US images, and appears as hyperechoic intratendinous lesions.

This study detected a significant thickening of the patellar tendon, which can be measured by US. We believe that this US feature is more sensitive and reliable in diagnosing early enthesitis than a classical MR high signal intensity image.

Figure 1 (A) A sagittal US scan shows thickened proximal entheses of the patellar ligament with loss of its fibrillar echo pattern, loss of the sharp definition of its posterior aspect compared with the distal portion (arrow head), calcific foci (arrow). (B) A transverse US scan of the same patient shows the thickened medial part of the patellar ligament with calcific focus (arrow). (C) A sagittal T2 fat suppression image shows the thickened distal part of the patellar tendon with altered signal intensity (arrow head) and prepatellar bursitis (arrow). (D) A sagittal T1 weighted image shows high intensity signals of the proximal patellar tendon.
within the superior medial and central aspects of the patellar tendon at its proximal attachment. This interesting observation has been confirmed in other related studies.

In conclusion, we found several pathological differences between the pattern of patellar enthesitis and that of Achilles tendon and plantar fascia of the heel. The enthesial changes of the patellar tendon occurred at the tibial or patellar insertion either on its medial or lateral aspect, but in the case of the Achilles tendon, the enthesal changes were detected only in the calcaneal insertion. The presence of calcific foci were found more often in the Achilles tendon than in the patellar tendon. US is a valuable and sensitive diagnostic method in patients with spondenegative spondyloarthropathy and knee enthesopathy who have normal findings with conventional radiological images of the knee joint.

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Longlasting effects of immunoadsorption in severe Sjögren’s syndrome
M Böhm, T Dörner, F Knebel, A Bruns, N Jochmann, G Baumann

Despite advances in understanding the immunopathogenesis of Sjögren’s syndrome, successful therapeutic interventions are extremely limited.

CASE REPORT
With this in mind we began a immunoadsorption treatment of a 38 year old woman, diagnosed 1.5 years ago with Sjögren’s syndrome. She reported dry eyes and mouth for several months and intermittent arthralgias, especially of the small finger joints and elbows, as well as swollen joints of the hands, elbows, and ankles for six years. During the six months before treatment her joint symptoms had increased significantly and considerably impaired everyday activities. A Schirmer’s test was positive. The antinuclear antibodies and rheumatoid factor were raised, autoantibodies against Ro/SSA and La/SSB were positive. Thus, the patient fulfilled six of the six revised criteria of primary Sjögren’s syndrome.

Previous corticoid treatment (prednisolone 20 mg a day for four weeks) had led to oropharyngeal candida mycrosis, and methotrexate (25 mg a week) did not have a therapeutic effect. Because chloroquine had side effects in her family, the patient refused to take hydroxychloroquine. Upon presentation in our clinic, the patient took 5 mg a day of prednisolone. Because the severe arthralgias and sicca symptoms did not respond to conventional treatment we started to treat the patient with immunoadsorption, to improve the symptoms by reducing IgG to 10–20% of its initial level.

Approval of the ethics committee and informed consent by the patient were obtained, and we started immunoadsorption therapy according to a previous protocol used for patients with dilated cardiomyopathy.

Immunoadsorption treatment took place in two consecutive cycles with an interval of four weeks. The first cycle comprised three treatment days and the second, two. The plasma filtration was similar to haemodialysis. The IgG-Therasorb Adsorber and a Mirosoorb treatment unit (Plasmaselect, Teterow, Germany) treated 7 litres plasma a day. Plasma IgG, antibody complexes, and fragments of antibodies were bound to the Fc fragment of polyclonal sheep antihuman antibodies, bound in turn to Sepharose.

As a result of the striking and sustained clinical benefit achieved, the treatment was discontinued after the second cycle of immunoadsorption.

DISCUSSION
To prevent a rebound and infection after immunoadsorption, the treatment protocol requires intravenous IgG substitution (0.5 g per kg body weight Venimun) after each cycle, indicating that the patient’s IgG has a significant role in the disease. We cannot exclude the possibility that immunoglobulin infusion may have had some effect on the outcome. This needs to be examined by additional studies.
This case report suggests the potential advantages of immunoadsorption in severe Sjögren’s syndrome which is refractory to conservative treatment. After 16 months the patient remains free of tender and swollen joints, and the sicca symptoms remain stable. A current open trial will provide further data, allowing better assessment of the value of immunoadsorption in patients with Sjögren’s disease.

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REFERENCES

Juvenile temporal arteritis and activated protein C resistance

B Granel, J Serratrice, N Ene, P E Morange, P Disdier, P-J Weiller

Juvenile temporal arteritis (JTA) is a rare benign vascular lesion limited to the branches of the external carotid artery, first recognised by Lie et al. Histological features of JTA include vasculitis (without giant cells and granulomas), lymphocytes infiltrate, and fibrous intimal proliferation. Despite frequent local arterial thrombosis associated with the lesion, no study on the coagulation pathway has been undertaken until now. We present an exceptional case of JTA occurring in a man in association with activated protein C resistance.

CASE REPORT
A 34 year old male smoker (12 cigarettes/day), without past medical history, consulted in January 2001 for right headache associated with swellings of both temples. Temporal regions were painful when he wanted to brush his hair. No visual or systemic symptoms occurred. Physical examination showed no pulse of the right temporal artery and a pulse of the left one; both arteries were thickened, tensed, indurated, and painful. Ultrasound echo Doppler disclosed arterial thrombosis of the right superficial temporal artery and normal aspect of the carotids and subclavicular arteries. Biological evaluation showed no inflammation (erythrocyte sedimentation rate 2 mm/1st h and C reactive protein <4 mg/l) and normal white blood cell count (8 x10^9/L, with 1.4% eosinophils), haemoglobin level, and platelet count. Histopathological analysis of the right temporal artery disclosed an inflammatory infiltrate with mononuclear cells without giant cells or granuloma, and a fibrous intimal proliferation. No eosinophils were seen in the biopsy. An electrocardiogram was normal. The patient was treated with aspirin (100 mg/day) and the temporal pain resolved. He stopped smoking.
Past familial history showed multiple phlebitides and a pulmonary embolism in the grandmother, a phlebitis in the mother, and three phlebitides in the father. Blood coagulation evaluation in the patient showed activated protein C resistance with heterozygous mutation in factor V gene (Arg 506-Gln). There was no mutation in factor II gene, normal levels of antithrombin, protein S, protein C, and protein Z, no lupus anticoagulant, and no anticardiolipin antibody. Homocystine level was normal. A search for antinuclear, antineutrophil cytoplasmic antibodies, and cryoglobulinaemia was negative. Fundoscopic examination was normal. Angiographic magnetic resonance imaging of the cerebral arteries only performed a few months later (when the patient was referred to our medical department) was normal. Control of ultrasound echo Doppler performed 2 years later (when the patient was still receiving aspirin treatment) showed a persistent thrombosis of the right temporal artery associated with a myointimal thickness and a non-occlusive myointimal thickness of the left artery. Carotid, vertebral, and subclavicular arteries appeared to be normal. The patient continued to receive aspirin treatment.

DISCUSSION
JTA is a very rare entity of benign course and fewer than 15 cases have been reported. It is usually unilateral, although bilateral involvement, as seen in our case, has also been reported. JTA occurs in children or adults under 40 years, and is characterised by (a) pain in the temporal region with arterial induration or palpable nodules; (b) no associated inflammatory symptoms; (c) no objective ophthalmic symptoms; (d) no biological inflammation; (e) possible blood eosinophilia with eosinophilic infiltrate in biopsy; and (f) no need for steroid treatment. The reported patients who were treated with steroids had systemic symptoms and an associated systemic vasculitis. More recently, rapid regression of the temporal pain in a patient with JTA treated with non-steroidal anti-inflammatory drugs has been reported. Differential diagnosis includes Kimura’s disease (angiolymphoid proliferative disorder of soft tissues with eosinophilia and raised immunoglobulin E, occurring almost exclusively in oriental men), angiolymphoid hyperplasia with eosinophilia, giant cell arteritis, panarteritis nodosa,
Serum osteoprotegerin but not receptor activator of NF-κB ligand correlates with Larsen score in rheumatoid arthritis

M Skoumal, G Kolarz, W Wołoszczuk, G Hawa, A Klingler

Osteoprotegerin (OPG) is a soluble decoy receptor, produced by osteoblastic cells and in the inflamed synovium of RA by dendritic cells, B cells, and other immunocompetent cells. 1 It inhibits the differentiation of osteoclast precursor cells and the activation of mature osteoclasts by neutralising the receptor activator of NF-κB ligand associates with Larsen score in rheumatoid arthritis. 3 Histopathological analysis often discloses arterial thrombosis associated with non-giant cell, non-destructive, and non-granulomatous vasculitis. A search for blood coagulation disorders has never been performed until now. Factor V Leiden is a well-known risk factor for venous thrombosis but may also play a role in arterial thrombosis. This observation reminds doctors of this rare entity and highlights the need to study blood coagulation in cases of JTA.

Table 1 Correlation of OPG and sRANKL with clinical data

<table>
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<th>Clinical data</th>
<th>Mean</th>
<th>SD</th>
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<th>p Value</th>
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OPG, osteoprotegerin; sRANKL, soluble receptor activator of NF-κB ligand; ESR, erythrocyte sedimentation rate; CRP, C reactive protein; RF, rheumatoid factor; DAS, disease activity score.
Successful management of neonatal cryoglobulinaemia after a gemellar pregnancy in a woman with symptomatic type I cryoglobulinaemia

J Sibilia, O Feugeas, V Laugel, A Dreal, J Messer, J Goetz

CASE REPORT
Since 1996 a 32 year old woman had had cold-induced clinical manifestations: Raynaud’s phenomenon, livedo reticularis, necrotic and purpuric lesions on the legs, and acrocyanosis on the ears and fingers. She had no family history of cold intolerance, autoimmune disease, or recurrent thrombosis. In 1998 these symptoms were traced to a monoclonal IgGλ cryoglobulin (type I) present at 1.5 g/l and precipitating at 27°C. Cold agglutinin and cryofibrinogen were absent while total complement (CHs) and C4 fraction were low. There were no antinuclear, anti-dsDNA, antineutrophil, anticardiolipin, or anti-β2-glycoprotein I antibodies, and viral serology was negative for hepatitis B and C and cytomegalovirus. A skin biopsy showed leukocytoclastic vasculitis and thrombosis of the capillaries related to the cryoglobulin. There were no renal, gastrointestinal, or neurological manifestations, but the patient had intermittent distal polyarthritis and was treated with pentoxifylline accompanied by protective measures against cold.

This mother had born a first healthy child in 1996 and had experienced no previous miscarriage. In 2000, during a second dichorionic gestation of twins, the lesions were progressive while the cryoglobulin persisted at 1.10 g/l. Because the IgGλ chain can cross the placenta and the initial temperature of cryoprecipitation was 27°C, clinical manifestations could be expected in the newborns at room
temperature. Treatment of the mother consisted only of protection against cold. Plasmaphereses were not performed as these do not totally eliminate cryoglobulin and have potential drawbacks, such as inducing premature labour by altering the plasma volume or hormone levels, or decreasing fetal nutrients and maternal and fetal serum immunoglobulin levels. General measures were defined to avoid cryoprecipitation at delivery: a room at 28°C, adapted clothes permitting medical supervision, warming of liquids to 37°C before perfusion to the mother, and placing the newborns in incubators.

The twins (female and male weighing 2440 g and 2690 g, respectively) were delivered after 39 weeks’ gestation and immediately put in incubators at 32°C. One and five minute Apgar scores were 10 for both neonates, and no cutaneous or visceral involvement was detected, despite the presence of cryoglobulin at 1.70 g/l in cord blood. Two days later the twins were placed in a room at 25°C, but after 3 hours small erythrocyanotic maculae appeared on the boy’s fifth finger (left hand) (fig 1) and second and fourth toes (left foot). These lesions were very similar to those seen in the mother, whereas careful examination of the girl disclosed no cutaneous manifestations. The twins were replaced in incubators and the lesions disappeared slowly within one week. After another 2 days they were removed from the incubators and dressed warmly, no further lesions were observed and the dressing precautions could be gradually abandoned before discharge. The monoclonal IgG1 cryoglobulin was detected in the serum of both neonates at comparable levels, but it was unfortunately not possible to obtain follow up of these results. Six months later the twins were healthy while the mother’s status remained stable with an unchanged serum cryoglobulin (1 g/l).

**DISCUSSION**

Maternal IgGs can cross the placenta, as is well illustrated by certain neonatal autoimmune diseases related to maternal autoantibodies. In the present case a monoclonal IgG1 cryoglobulin identified in the mother and her twin children was associated with transient cold-induced lesions in one child. This is the first report of a pathogenic neonatal effect due to the transplacental passage of a type I cryoglobulin. Although the lesions were probably favoured by the high precipitation temperature (27°C), many other physicochemical parameters can influence cryoprecipitation and hence might explain the difference between the twins. Progressive clearance of the maternal IgG1 after its passive transfer may further explain the favourable outcome and absence of recurrence. Such cases are rare, probably on account of the low incidence of type I cryoglobulinaemia in women of child bearing age. Associations between pregnancy and other types of cryoglobulin have been described but without related neonatal lesions. In a patient with mixed type II cryoglobulin and hepatitis C infection, cryoglobulinaemia was responsible for placental lesions inducing pre-eclampsia, but no specific neonatal lesions were seen.

The consequences for the fetus of the transplacental passage of IgG type I cryoglobulin are still unknown and the management of such pregnancies is difficult owing to the lack of a consensual procedure for the prevention of maternofetal complications. Corticosteroids or immunosuppressive drugs have failed to demonstrate any benefit. In our experience, protection against cold seems to be the only way to avoid precipitation of the cryoglobulin in the vessels of the mother and neonates with its pathogenic effects.

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Primary Raynaud’s phenomenon in monzygotic twins

T Oskay, Ü Ölmez

We report two cases of primary Raynaud’s phenomenon in 16-year-old monzygotic female twins.

CASE REPORT

The twins were admitted to the dermatology department with a one-year history of the sudden onset of episodic purple lesions on both hands. Presentation of the lesions in the twins was synchronous. After careful questioning, we determined that it was usually induced by emotion or cold temperature. On dermatological examination, the order of purple discoloration, erythema, and pallor was observed symmetrically in both hands. There was no history of smoking, the use of drugs, occupational hazard, or trauma. They also had sensations of coolness and paraesthesia. Photographs of the hands were obtained during an attack and used to confirm the history. We also performed a provocative test—namely, immersion of the patient’s hand in ice water for three minutes, to confirm the diagnosis. Nailfold capillaroscopy examination also demonstrated normal capillaries. The female twins were born at 38 weeks of gestation by caesarean section. Obstetric anamnesis was normal. The twins’ past medical history was unremarkable. Their parents were relatives, but the family history was negative for cutaneous or collagen vascular disease.

Results of laboratory examinations, including complete blood count, biochemistry, urine analysis, thyroid function tests, serum cryoglobulins, complement levels, immunoglobulins, erythrocyte sedimentation rate, and C reactive protein, were normal. Serological testing for anticardiolipin antibody, antiphospholipid antibodies, and antibody to Ro, La, dsDNA, ribonucleoprotein, and Smith antigen were negative. The twins were diagnosed as monzygotic with almost complete certainty by examining the ABO blood type, HLA typing, fingerprints, and from the physical resemblance between them. Identical HLA antigens were HLA-A2, A24, B17, B57, Cw3, Cw6, DR4, and DR11. Medical evaluation subsequently disclosed no disorder known to cause secondary Raynaud’s phenomenon (RP).

DISCUSSION

RP is an exaggerated vascular response to cold temperature or emotional stress, primarily seen in young women. It is classified into primary or idiopathic without underlying disease and secondary when it is related to various medical problems.1,2 Clinical criteria are used to distinguish between patients with uncomplicated or primary RP, and those with secondary RP. The suggested criteria for primary RP are symmetric attacks; the absence of tissue necrosis, ulceration or gangrene; the absence of a secondary cause on the basis of a patient’s history and general physical examination; normal nailfold capillaries; a negative test for antinuclear antibody; and a normal erythrocyte sedimentation rate.2,3 The most common diseases associated with secondary RP are scleroderma, mixed connective tissue disease, systemic lupus erythematosus, occlusive vascular disease, drug effects, haematological abnormalities, and use of vibrating tools.3,4 Primary RP occasionally evolves into a connective tissue disease, most commonly scleroderma.4,5

Even though the cause of primary RP is not known, increased α2 sympathetic receptor activity on vessels, endothelial dysfunction, deficiency of calcitonin gene related peptide, protein-containing nerves, or some central thermo-regulatory defect have been implicated.5,6 A genetic cause for primary RP was first suggested by Lewis and Pickering, based on an apparent familial aggregation of primary RP.2,7,8 MacGregor et al. first assessed the genetic basis of RP in a population-based twin study and found a substantial genetic contribution to the symptoms of RP.5 Recently, three potential candidate genes have been described in the genetic susceptibility to primary RP.9

To the best of our knowledge, this is the first report documenting the occurrence of primary RP in monzygotic twins. Association with HLA antigens, parental consanguinity, and development in identical twins may indicate that genetic factors have a role in the occurrence of this disease. Larger series of twins and investigation of multicase families are needed to clarify the genetic component in the pathogenesis of primary RP.

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