LETTERS

Extended colonic ulcerations in a patient with microscopic polyangiitis

C-N Tsai, C-M Chang, C-H Chuang, Y-T Jin, M-F Liu, C-R Wang

Microscopic polyangiitis is a necrotising vasculitis primarily affecting small vessels, with few or no immune deposits. Patients are characterised by positive antineutrophil cytoplasmic antibodies (ANCA), mainly perinuclear pattern. The spectrum of clinical manifestations is broad, including the kidney, musculoskeletal system, lung, gastrointestinal tract, skin, ear, nose, and throat, and neurological system. Although gastrointestinal disease is noted in half of the patients, the presentation is usually mild. Here, we report a patient with microscopic polyangiitis with initial presentation of extended colonic ulcerations and haemorrhage, characterised by a crypt abscess. To our knowledge, such a finding has not been reported previously.

CASE REPORT

A 69 year old man was admitted to hospital owing to leg oedema and body weight loss in the past 3 months. No systemic disease had previously been noted. He had a poor appetite and abdominal discomfort, but denied bowel habit changes, bloody or tarry stool.

On examination, vital signs were stable. Pale conjunctiva, mild abdominal tenderness, and pitting oedema over the legs were noted. Laboratory tests showed leucocytes 11.9 $\times$ 10$^9$ l$^{-1}$, haemoglobin 67 g/l, albumin 225 $\mu$mol/l, C reactive protein 1157 mg/l, and erythrocyte sedimentation rate more than 150 mm/1st h. Daily urinary protein loss was 0.7 g. Stool examination showed positive occult blood. However, the results of panendoscopy and lower gastrointestinal investigations were negative.

Intermittent high fever was noted several days after admission. No infectious focus was identified by repeated blood and stool cultures. Abdominal computed tomography disclosed mild inflammation over the mesentery. Colonoscopy showed markedly swelling mucosa with haemorrhage and ulcers from rectum to the cecum (fig 1A). A pathology examination reported inflammatory infiltrates in the lamina propria, destruction of the mucosal gland, and a crypt abscess (fig 1B). Ulcerative colitis was diagnosed, and mesalazine and prednisolone 30 mg every day were prescribed.

Sudden onset of haemoptysis occurred 1 week later and pulmonary haemorrhage was diagnosed. Intravenous methylprednisolone 20 mg every 8 hours was given, and the haemoptysis disappeared 1 week later. Repeated urinary analysis showed a daily protein loss of 4 g. An autoantibody examination showed negative antinuclear antibodies and antifibrinoluculent basement membrane antibody. ANCA were positive (perinuclear pattern, $\times$320), and further analysis showed positive antithromboperoxidase 103 U/ml (normal range <20 U/ml). A renal biopsy showed glomerular necrosis, cellular crescent, interlobular arteritis, diffuse tubular atrophy, and few immune deposits. The final diagnosis was microscopic polyangiitis with a rare presentation of extended colonic ulcerations. Pulse intravenous cyclophosphamide 700 mg was given and steroid was gradually tapered. Proteinuria, hypoalbuminaemia, anaemia and inflammatory measures improved with treatment. He was discharged and received monthly pulse cyclophosphamide treatment thereafter. Neither abdominal pain nor diarrhoea was noted during follow up at the outpatient clinic.
DISCUSSION

In microscopic polyangiitis, necrotising glomerulonephritis is the most common abnormality and pulmonary capillaritis often occurs. Perinuclear ANCA are present in more than 60% of patients, and antineutrophil cytoplasmic antibody is closely allied.1 Gastrointestinal involvement is usually mild and there is no report of extended colonic ulcerations and haemorrhage.2 In addition, colonic biopsy disclosed crypt abscess, which can be found in ulcerative colitis, infectious colitis, radiation induced colitis, and graft versus host disease.3 However, such a finding has not been reported previously in patients with microscopic polyangiitis. Ulcerative colitis was diagnosed initially according to the colonicoscopic and pathological findings. Pulmonary vasculitis is very rare in patients with inflammatory bowel diseases.6

Renal disease with necrotising glomerulonephritis is not commonly seen in patients with ulcerative colitis.7 In addition, antineutrophil cytoplasmic antibody is usually not detected in such patients.8 The clinical course of this patient who had lung and renal manifestations did not favour a diagnosis of ulcerative colitis. The final diagnosis was microscopic polyangiitis with gastrointestinal involvement. Extended colonic ulcerations with haemorrhage have not been reported previously in patients with microscopic polyangiitis. Vasculitis should be considered for differential diagnoses of colonic ulcerations, especially when the presentation is atypical, to avoid delay of prompt treatment.

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REFERENCES


Severe digital ischaemia treated with phosphodiesterase inhibitors

C R Kumana, G T Y Cheung, C S Lau

Pulmonary hypertension is associated with autoimmune diseases, giving rise to digital ischaemia, including Raynaud’s phenomenon (RP). As sildenafil, a phosphodiesterase-5 (PDE-5) inhibitor, relieves the pulmonary hypertension,9 we suggested that it might also benefit digital ischaemia and RP symptoms. This report describes the impact of oral sildenafil (Viagra, donated by Pfizer HK) on three women with autoimmune disease and progressively severe digital ischaemia, despite treatment with a variety of drugs, including intravenous iloprost (fig 1). All patients gave written informed consent to try this off-label treatment.

CASE REPORTS

A 42 year old woman with dermatomyositis “sine myositis” and thyrotoxicosis (case 1) taking prednisolone, hydroxychloroquine, and carbimazole, experienced regression of troublesome RP and ischaemic digital ulceration after receiving diltiazem SR 200 mg/day and thrice monthly iloprost infusions. Ten months later, antituberculosis chemotherapy was started and the steroid dosage reduced, after she developed a febrile illness with cough, pleuropneumonial effusions, and sputum culture positive for tuberculosis (TB). Digital gangrene supervised in both hands and both feet despite daily intravenous iloprost infusions for 2 weeks.

A day after starting sildenafil 50 mg three times a day her digital circulation and pain improved markedly; facial flushing was the only adverse reaction. In view of the slightly deteriorating digital ischaemia and experimental evidence,2 we inferred that rifampicin treatment was inducing the metabolism of sildenafil. Its dosage was therefore doubled. Symptomatic improvement continued and ischaemic tissues became demarcated. However, 26 days after starting sildenafil she succumbed to uncontrolled pulmonary TB.

A 28 year old woman with scleroedema/lupus (case 2) who had been receiving diltiazem, domperidone, omeprazole, and penicillamine for about 4 years presented with progressive breathlessness. Computed tomography of the thorax suggested fibrosing alveolitis; prednisolone and azathioprine were started. Thereafter, she incurred occasional chest infections, other complications, and features of autoimmune diseases. She then developed persistent high grade fever, chills, rigors, night sweats, and increasingly severe digital arthralgia, vasculitis, and ischaemia. A polymerase chain reaction (PCR) of blood disclosed disseminated TB. Despite resolution of the sepsis after anti-TB chemotherapy, she developed progressive ischaemia and impending gangrene of her fingers, toes, and feet, despite regular infusions of iloprost for 2 weeks. One day after starting sildenafil 50 mg
three times a day, all four extremities showed marked symptomatic improvement, becoming warm, less painful, and less discoloured. Her digital vasculitis resolved; weeks later necrotic zones became demarcated and autoamputated. Owing to resurgence of active lupus, her prednisolone dosage was increased; her haematological/serological abnormalities regressed. After 48 days of sildenafil treatment, tadalafil (Cialis) 10 mg once daily was substituted for 78 more days, without any adverse reaction. Currently, the patient remains well and can stand.

A 76 year old woman with prior hypertension and ischaemic heart disease (case 3) presented with erythematosus facial and finger rashes, myalgia, finger and toe ischaemic lesions, and proximal muscle weakness. Dermatomyositis and a disseminated malignancy were diagnosed. After mutually agreeing to stop further investigation, prednisolone treatment was started. Her digital ischaemia responded poorly to diltiazem SR and any improvement during iloprost infusions disappeared when they stopped. After 12 days and despite her past medical history, sildenafil treatment was cautiously started, with immediate reduction in peripheral pain and ischaemia and healing of toe ulcers. Although well tolerated, sildenafil was discontinued after 3 weeks, as the patient required treatment with nitrates.

**DISCUSSION**

All three patients had immediate, subjective and objective improvements after initiating sildenafil, which was entirely consistent with Lichtenstein’s observations, except that our patients were very severely afflicted.\(^4\) In two patients, worsening RP and digital ischaemia appeared during active TB, consistent with the effects of sepsis.\(^5\) Rifampicin, a powerful inducer of relevant P450 enzymes (apt to enhance sildenafil elimination\(^2\)), appeared to affect their management. These

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**Figure 1** (A), (B), and (C) refer to cases 1, 2, and 3, respectively. In all three cases, ischaemia of viable tissues resolved after treatment with sildenafil (and tadalafil in case 2). (B) shows that increased finger extension became possible after treatment.
Hereditary C1q deficiency and secondary Sjögren’s syndrome

E P A H Hoppenreijs, P J van Dijken, P J Kabel, J M Th Draaisma

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A 13 year old Turkish boy with a known C1q deficiency and SLE-like disease developed recurrent parotitis. Investigations confirmed a secondary Sjögren’s syndrome (SS). As far as we know, the association between C1q deficiency and SS has not been described before. The potential role of C1q in the pathogenesis of SS and systemic lupus erythematosus (SLE) might stimulate further research in understanding the pathogenesis of these and other autoimmune diseases. Screening patients with SS for complement deficiencies, including C1q deficiency, seems indicated.

Deficiency of the complement component C1q is a rare genetic disorder with susceptibility to recurrent infections with polysaccharide encapsulated micro-organisms and a high prevalence of autoimmune phenomena, most often SLE.

CASE REPORT

At the age of 4 years a boy of consanguineous Turkish descent had meningitis of unknown origin. At the age of 8 years he presented with meningococcal septicaemia and meningitis. In the late convalescent phase of this infection he developed arthritis of his right elbow and pericarditis. At the age of 10 years he was admitted owing to lobular pneumonia. Immunological studies at that time demonstrated no functional or antigenic activity of C1q based on a homozygous Glu-86 stop mutation in the C1qA gene exon 2. Both parents and a sibling sister were found to be asymptomatic heterozygous carriers. His sibling brother, who also developed SLE-like symptoms, was homozygous for this mutation.

At the age of 13 years the patient developed recurrent arthritis of the ankle and elbow. Two years later he presented with six episodes of alternate right and left parotitis. There were no complaints of dry mouth or dry eyes. Ultrasonography of the parotid gland during an episode of parotitis disclosed diffuse swelling. Salivary gland biopsy showed lymphocytic sialoadenitis with a focus score of 4 and a percentage of plasma cells containing IgA of 9%, consistent with SS. Microscopic haematuria and proteinuria were not present. Immunological studies showed positive antinuclear antibodies (ANA), RNP antibodies, anti-Sm antibodies, anti-SSA antibodies, and a positive rheumatoid factor (RF). Anti-SSB antibodies and anti-dsDNA antibodies were not detected. A direct Coombs test and a serological test for syphilis were negative. Serum immunoglobulins were slightly raised. C3 and C4 were normal.

DISCUSSION

In our patient a homozygous point mutation in the C1qA gene was demonstrated that has earlier been described in five families from the Slovak republic and Turkey. With arthritis, a positive ANA test, and anti-Sm antibodies our patient has an SLE-like disease.3 Patients with SLE commonly have sicca symptoms, which may be related to the concomitant occurrence of SS.5 Secondary SS, in which the disease coexists with an autoimmune disease, is defined by the presence of either ocular or oral symptoms and two of four objective classification criteria.4 Our patient with recurrent swollen salivary glands, autoantibodies to ANA, RF, and SSA, a salivary scintigraphy showing delayed uptake, and biopsy findings consistent with SS, meets the full criteria for secondary SS.

These observations suggest that absence or abnormal function of C1q leads to susceptibility for SLE and SS. This may be due to ineffective immune complex clearance that causes tissue injury, expresses autoantigens, and stimulates an autoantibody response.7 In this respect it is significant that our patient had late onset reactive arthritis and pericarditis related to his meningococcal disease, which is also mediated by immune complexes.6 Other data suggest a defective clearance of apoptotic cells, which promotes accumulation of nucleosomes (suggested antigens in SLE), which in turn drives an autoimmune response.7

In conclusion we report a patient with C1q deficiency, SLE-like disease, and secondary SS. This is the first time that an association between C1q deficiency and SS has been reported. It suggests a role for C1q in the pathogenesis of autoimmune diseases, possibly due to ineffective immune complex clearing or defective apoptosis, and warrants further research.

Screening of patients with C1q deficiency for SS, but especially screening of patients with SS for complement deficiencies, including C1q deficiency, seems indicated.

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Acute pneumonitis starting 2 hours after intramuscular gold administration in a patient with rheumatoid arthritis

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Gold salts have been used for the treatment of rheumatoid arthritis (RA) for over 50 years. Their use is limited by the incidence of adverse reactions, including pneumonitis. Most case reports describe pneumonitis developing after a cumulative dose of several hundred milligrams. Only four cases of lung disease have been reported with a cumulative dose of <100 mg.

Here we report a case of gold pneumonitis, starting only 2 hours after the first maintenance dose of 20 mg was administered.

CASE REPORT

A 54 year old woman with RA had been followed up since 1984. Her arthritis was well controlled with chloroquine. In 1997 her disease had flared up, and alternative disease modifying drugs were introduced—sulfasalazine, then methotrexate (both of whom were stopped because of adverse side effects), and finally, leflunomide. The last of these caused leucopenia and hence was discontinued. Subsequently, her disease became more active again with prolonged joint stiffness, hand synovitis, and erythrocyte sedimentation rate (ESR) 102 mm/1st h.

Treatment was started with gold injections, with a test dose of 10 mg (no adverse effects noted), followed a week later by a dose of 20 mg. About 2 hours afterwards she complained of dizziness, nausea, then she developed a cough and breathlessness. She was admitted to hospital later that day. On examination she was apyrexial, tachypnoeic, cyanosed (oxygen saturation 79% on air), pulse 110/minute, blood pressure 80/46, heart sounds normal, and had no signs of heart failure.

Investigations showed haemoglobin of 103 g/l with normal white cell and platelet counts. C reactive protein was 115 mg/l and troponin-I (a marker for myocardial infarction) was normal at 0.1 µg/l. Arterial blood gases on air showed a type 1 respiratory failure pattern. A chest x ray examination showed bilateral lower zone alveolar shadowing. An echocardiogram showed a normal ejection fraction.

After high dose oxygen treatment, intravenous hydrocortisone 100 mg, and intramuscular adrenaline 0.5 mg 1:1000, some improvement was seen clinically, oxygen saturation rising to 90% and blood pressure to 116/69. She was admitted to the high dependency unit and treatment started with pulse methylprednisolone 1 g daily, nasal continuous positive airway pressure ventilation, and required parenteral nutrition. On the day after her admission, she developed a generalised macular rash. Over the next 7 days her clinical condition improved (serial chest x ray examinations over this period showed clearing of the previous alveolar shadowing), and her treatment was changed to a reducing regimen of oral steroids. She was discharged, continuing frumil once daily and prednisolone 5 mg daily. Her arthritis was inactive and blood pressure 110/70. The ESR was 22 mm/1st h.

During her admission, it was considered that owing to the severity of her illness she was not fit enough to have other investigations—for example, computed tomography chest scan, bronchoalveolar lavage, and pulmonary function tests.

The gold was discontinued and her arthritis remains in remission.

DISCUSSION

Gold pneumonitis is a potentially fatal complication of gold salt treatment. This case illustrates two important points in management:

- Some of the early symptoms could have been confused with a nitritoid reaction. Indeed, hypertensive patients with RA who are taking angiotensin converting enzyme inhibitors (in this case ramipril) show a higher frequency of such reactions even if long term gold treatment is established. This combination should be avoided where possible.

- The clinical picture started to develop after only 2 hours of gold administration with just a 30 mg cumulative gold dose. Most case reports have suggested higher cumulative doses.4 5 6 and longer duration of treatment. To our knowledge such a florid onset has not been previously reported.
An unusual complication of appendicitis

S L Mackie, A Keat

32 year old man presented with a 3 day history of fever, central abdominal pain, and frequent loose bowel motions without blood or mucus. The pain had recently shifted to his right iliac fossa. He had previously been well and he was receiving no regular drugs. On examination he was systemically well, but rebound tenderness was noted in the right iliac fossa, and C reactive protein (CRP) was raised (78 mg/l, reference range <10). A diagnosis of acute appendicitis was made and he underwent appendicectomy. Histology disclosed serosal congestion with a predominantly eosinophilic infiltrate in the mucosa and deeper layers.

Postoperatively the fever resolved, but the diarrhoea continued for several weeks. Two weeks later he developed a right knee effusion associated with raised inflammatory markers (CRP 37 mg/l). Synovial fluid from the knee and blood cultures was sterile; the diarrhoea was mild and stool cultures were not performed. Treatment with non-steroidal anti-inflammatory drugs and sulfasalazine was started. The arthritis resolved over the next 3 months, with normalisation of inflammatory markers; at the last review the patient had stopped all treatment and was well.

Serological testing showed raised titres (1/1280) of antibodies to Yersinia enterocolitica O:3 (>1/160 being considered significant), which remained at 1/640 3 months later.

On the basis of the clinical picture and serology a diagnosis of yersiniosis was made.

DISCUSSION

Reactive arthritis following enteric Y enterocolitica infection is well described.1 Y enterocolitica and Y pseudotuberculosis are also well known causes of mesenteric adenitis, ileocolitis, and appendicitis,2 but reports of reactive arthritis following Y enterocolitica appendicitis are surprisingly rare.3,4

Pure cultures of Y enterocolitica have occasionally been isolated from acutely inflamed appendices, suggesting a primary pathogenic role for this species.5 Serological evidence of acute infection by Y enterocolitica was also reported in three of 90 patients with acute appendicitis,6 of these, two had postoperative diarrhoea. In another series7 evidence of Y enterocolitica or Y pseudotuberculosis was found by a polymerase chain reaction technique in 10 of 40 cases of granulomatous appendicitis but in none of 30 cases of non-granulomatous appendicitis.

An unexpected feature of this case was that granulomas were not seen on histology. The significance of the eosinophilia is uncertain, but in other respects the histology was typical of acute appendicitis. The association of appendicitis and yersinia induced reactive arthritis appears to be surprisingly unusual, despite the common occurrence of each condition individually. Nevertheless, it is clearly important for physicians to be aware that abdominal pain in patients with reactive arthritis may signify the potentially lethal complication of appendicitis, and for surgeons to be aware that appendicitis may be complicated by reactive arthritis.

REFERENCES

A 17 year old girl was admitted to our rheumatology clinic with right knee pain and swelling for the past 3 months. A diagnosis of familial Mediterranean fever (FMF) had been made 8 years previously: she had recurrent attacks of fever, abdominal pain, and knee arthritis. She had been using colchicine 1.5 mg/day regularly since then. Although her abdominal attacks and fever had subsided with colchicine treatment, recurrent mild knee attacks occurred almost every month. Two years ago she was found to have the homozygous M694V gene mutation for FMF. Three months ago, she was admitted to hospital owing to monarthrosis in her right knee—but this time with a prolonged and severe episode. She was given antibiotics, and a suspected diagnosis of septic arthritis was made. As the cultures did not yield any bacterial growth and her symptoms persisted, she applied to our department.

A physical examination disclosed painful and limited movement, swelling and warmth in the right knee joint. Laboratory findings were as follows: erythrocyte sedimentation rate: 96 mm/1st h, C reactive protein: 165 mg/l (0–8), fibrinogen: 8.5 g/l (1.4–4.3). Protracted arthritis of FMF was diagnosed and the dose of colchicine was increased to 2 mg/day. Ten days later, she was seen on a control visit with complaints of abdominal pain, diarrhoea, skin rash—probably due to colchicine—and persistent knee arthritis, which did not respond to colchicine treatment. Accordingly, interferon alfa 4.5 million IU (twice a week) was started and the colchicine dose was decreased to 1.5 mg/day.

Thereafter, she has been followed up every 10 days for 3 months and her knee arthritis has disappeared and the laboratory measures have improved (table 1). She is still being routinely followed up and is completely normal with the above mentioned regimen of interferon alfa and colchicine together with a protocol of isometric quadriceps strengthening exercises.

**DISCUSSION**

Arthritis in FMF usually comprises acute attacks, with complete resolution within a few days or 2 weeks. It is more common in patients with the homozygous M694V gene mutation. Occasionally, the attacks are protracted, lasting for several months, and chronic joint disease has been estimated to contribute about 5–10% to the joint manifestations in FMF.1,2 However, once degenerative or necrotic changes ensue, the recurrent attacks of arthritis are no longer punctuated by symptom-free intervals. Additionally, the irreversible morphological changes in the joints become non-responsive to colchicine, corticosteroids, or non-steroidal anti-inflammatory drug treatment. Although ruling out a coexisting chronic inflammatory arthritis should remain a prerequisite, surgery is usually warranted in such cases.1,4,5 To the best of our knowledge, a medical treatment alternative for these patients that would make synovectomy unnecessary has not been reported. Only one report, by Tunca et al, has described favourable effects of interferon alfa on the abdominal attacks of seven patients with FMF.6 As recent studies have disclosed up regulation of the MEFV gene by interferon,7 the mechanism—which could not have been substantiated previously—has become evident. Thus, overall, we advocate the use of interferon alfa as an adjunct for the treatment of colchicine resistant arthritis attacks in FMF.

**Table 1 The changes in the laboratory variables of the patient with interferon alfa treatment**

<table>
<thead>
<tr>
<th>Interferon treatment</th>
<th>ESR (mm/1st h)</th>
<th>CRP (mg/l)</th>
<th>Fibrinogen (g/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immediately before</td>
<td>61</td>
<td>106</td>
<td>8.8</td>
</tr>
<tr>
<td>Two weeks after</td>
<td>35</td>
<td>11</td>
<td>4.9</td>
</tr>
<tr>
<td>Six weeks after</td>
<td>30</td>
<td>4</td>
<td>4.8</td>
</tr>
<tr>
<td>Twelve weeks after</td>
<td>25</td>
<td>4</td>
<td>4.5</td>
</tr>
</tbody>
</table>

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Transient bone marrow oedema in a child
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Transient bone marrow oedema (TBMO) is an uncommon condition associated with joint and bone pain on activity. The most common localisation is the proximal femur, and it mainly affects only one bone.1 Reports of bone marrow oedema in children are scarce.2–4

CASE REPORT
An 8 year old boy was referred to hospital owing to difficulty in walking. He had started ski jumping 2 weeks earlier, but his history showed no injuries in the extremities. He complained of pain in the knees, ankles, and wrists. Mild fever and signs of upper respiratory tract infection were detected and reactive arthritis was suspected. Treatment with naproxen was started and he was discharged. Ultrasound investigation showed only slight symmetrical oedema in the ankle joints. Haemoglobin, white blood cell count, and thrombocytes were all within normal limits.

During the next 2 weeks, the pain was localised in both feet and the patient was hardly able to walk. Palpation and compression of both feet was very painful. An x-ray examination which was carried out 2 weeks after the first visit showed marked osteopenia in the tarsal area and in distal parts of the tibia and fibula (fig 1). A magnetic resonance imaging (MRI) scan was performed 4 weeks after the beginning of the symptoms showed marked bone marrow oedema (fig 1). Transient osteoporosis was suspected. Bone scintigraphy showed areas of hyperperfusion both in the feet and the hands. An MRI scan was subsequently obtained of the hands also, which also showed bone marrow oedema. Axial bone mineral density was measured by dual x-ray absorptiometry and was normal. Serum calcium, phosphate, parathyroid hormone, alkaline phosphatase (AP), and vitamin D metabolites were normal. However, bone markers were raised (table 1).

During follow up the pain became less intensive and after 3 months, the condition had resolved clinically, although an MRI scan was still abnormal. MRI normalised in 8 months. However, serum osteocalcin values remained raised, suggesting increased bone remodelling (table 1).

DISCUSSION
There are no data on the incidence of TBMO in children. However, it seems to be a relatively rare cause of foot and ankle pain also in adults.5 In another study of 1123 patients referred for MRI imaging of the foot, 72 patients with oedema-like bone marrow abnormalities were registered.6

The aetiology of TBMO is unknown but it may be associated with local vascular disturbances, microtrauma, bone contusion, or altered biomechanics.6–7 The x-ray examination is usually either normal or shows localised osteoporosis, as in our case. However, osteoporosis is a rare finding in bone biopsies.8 Histological studies have suggested ischaemic origin.9

The definitive diagnosis of TBMO is made by MRI. Bone contusion and bone bruises normally manifest as focal areas of low signal intensity in T1 weighted images and increased signal intensity in T2 weighted images, whereas bone oedema shows diffuse changes of intensity.1 In our patient marked bone marrow oedema was seen in both feet and hands.

Because of the unknown aetiology of bone changes we measured biochemical markers of bone metabolism. Serum AP values were within the normal range but both osteocalcin and procollagen type I C-peptide were high and remained high, although the clinical condition was relieved. However, except for AP, interpretation of these results is difficult because there are no established normative data for bone markers in children.10

The treatment of TBMO is not well established. Core decompression and vasodilatator treatment with iloprost have been suggested.4,7 Conservative treatment aims at protecting bone from weight bearing, both to prevent collapse of the articular surface and provide relief from pain. In this

Figure 1  A radiograph showing marked osteoporosis in the tarsal and metatarsal bones. In the MRI scan, hyperintesity in the tarsal bones can be seen as a sign of oedema in T2 weighted images.
Joint lavage and pseudogout

P Pasquetti, E Selvi, K Righeschi, M Fabbroni, R De Stefano, E Frati, R Marcelongo

Joint lavage, although its efficacy is still under debate,1–4 seems to be effective, mostly, in the treatment of gonarthrosis associated or not with chondralocalcinoses (CC).2–5 Paradoxically, acute pseudogout is a complication of this technique.6 Our study aimed at evaluating the incidence of pseudogout in 73 patients with gonarthrosis, associated or not with CC, who underwent arthroscopic lavage (AL).

METHODS AND RESULTS

In this retrospective study we assessed the incidence of pseudogout attacks that occurred 24 hours after surgery in 73 consecutive patients with gonarthrosis (52 women (71%), 21 men (29%)), who underwent AL of the knee at our hospital. All the patients had medium-severe symptomatic osteoarthrosis, according to Kellgren and Lawrence’s classification (II-III-IV degrees),9 and were unresponsive to drugs (that is CC, who underwent arthroscopic lavage (AL).

Of the 73 patients (mean (SD) age 59.4 (6.7)), nine (12%) had an arthritic episode within 24 hours after surgery. In each of these cases, wet sinovianalysis confirmed the inflammatory nature of the phenomenon (leucocyte counts between $5 \times 10^9$ and $30 \times 10^9/\text{l}$; polarising microscopy demonstrated the presence of CPPD. In all nine cases, culture and bacterioscopic examination of the synovial fluids were negative. The joint was injected with steroids, and the phenomenon resolved completely within 24 hours. The joint was injected with steroids, and the phenomenon resolved completely within 24 hours.

REFERENCE


The OR of group A compared with group B was estimated to be 5.5 (95% CI 1.24 to 24.60; \( p < 0.05 \)) (table 1).

**DISCUSSION**

The results of this study underline the significant incidence (26%) of pseudogout as a possible complication of AL in patients with CC (table 1).

Among the pathogenic hypotheses, mechanical-traumatic is the most likely; in fact, the lavage fluid could promote “crystal shedding”, due to the release of CPPD embedded in the joint tissues.8

Furthermore, it should be noted that this complication was also documented in three patients with gonarthritis without radiological or laboratory evidence of CC. This might be due to the fact that in a variable percentage of cases, microscopy for crystals in synovial fluid may yield false negative results.3 10 According to our data the age, sex, and radiographic grade of the patients had no significant association with the incidence of pseudogout after lavage.

In conclusion, pseudogout should be considered as a possible complication of AL, mainly in patients with CC, with about a fivefold risk compared with patients with osteoarthritis.

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Hereditary C1q deficiency and secondary Sjögren's syndrome

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