and the IL1ra have co-evolved as a cytokine control mechanism: agents eliciting IL1 production generally induce IL1ra gene transcription as well, either directly or through a feedback loop triggered by IL1 itself. IL1ra competes with IL1 for binding to the cognate receptor, but produces no known signalling events upon binding. Additionally, one of the two IL1 receptors, type II, also lacks any known signalling capacity and is thought to function as a decoy binding site. If it is assumed that polymorphisms found in various members of the IL1 gene family modify gene function then the implication is that they should be studied collectively, not individually. In support of this argument comes recent experimental evidence showing that the IL1a and IL1b levels are determined from allelic combinations within the IL1 gene family, as well as evidence for linkage disequilibrium between polymorphic markers spread across the IL1 gene family locus. It is fair to say that Crilly and colleagues did allude in their paper to the possible existence of an extended IL1 gene family haplotype. However, there is already sufficient published evidence to justify examination of haplotypic combinations of the investigated alleles in population studies.

On a different note, although the authors’ argument about the validity of surgery as a study end point is convincing, their selection of a 15 year disease duration free from study end point is less convincing. Their argument about the validity of surgery as a control mechanism in population studies would be strengthened by the inclusion of a control group calculated that the median disease duration free from surgery as the cut off point for this study seems dubious. A previous study by the same group had calculated that the median disease duration before surgery was 14.6 years, implying that there is a substantial number of patients in the “no surgery” group who will require surgery shortly after the 15 year time point, thereby potentially confounding any significant statistical findings. One might argue that the aim of this study was not to differentiate between patients that will or will not require surgery but, rather, to differentiate between rapid (<15 years) versus delayed (>15 years to surgery) disease progression. Even so, a more valid statistical approach might have been to analyse all patients as a single group and try to correlate genotype (or haplotype) with disease duration to surgery.

Author’s reply

I thank Dr Vamvakopoulos for his interest in our article1 and would like to make the following response.

We agree that testing for haplotype combinations of the interleukin 1 (IL1) gene family allele would be useful. However, for a large number of alleles to be tested a considerable sample size would be needed to correct for multiple allele testing. This would require a multicentre study; this is limited by funding constraints.

Secondly, for the statistical analysis we predefined early and late surgery. Analysis was undertaken after discussion with a qualified medical statistician. Should the correspondents wish to analyse our data, we would be happy to allow access if guarantees can be provided.

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LETTERS TO THE EDITOR

A case of shingles mimicking carpal tunnel syndrome

A 59 year old woman with an eight year history of seropositive erosive rheumatoid arthritis (RA) receiving sulfasalazine and penicillamine presented with severe onset pain radiating from the left elbow to the left thumb, index and middle fingers. Examination disclosed synovitis of the left wrist, which might have caused median nerve compression. The wrist joint was injected with 20 mg of triamcinolone acetate with 1% lidocaine (lignocaine). She returned the following morning complaining of worsening pain. She was clinically well with no fever. White cell count was normal, but the erythrocyte sedimentation rate (ESR) was raised at 73 mm/1st h. A transcutaneous nerve-stimulating (TENS) machine was applied and she was prescribed amitriptyline 25 mg at night. The following day she had improved significantly but had developed a vesicular rash in the C6 dermatome consistent with herpes zoster infection (fig 1). Viral titres were consistent with current varicella zoster infection.

DISCUSSION

Establishing the cause of pain in patients with RA can be notoriously difficult. In addition to the psychological factors that influence pain perception, wrist and hand pain may result from rheumatoid synovitis, soft tissue inflammation, or mechanical nerve compression at wrist, elbow, and cervical spine. Herpes zoster infection is heralded by burning discomfort in a dermatomal distribution, which may occur for up to five days before the onset of the typical rash. Cervical dermatomes are affected in up to 15% of patients and may result in diaphragmatic paralysis and lower motor neurone paresis. In this case the occurrence of prodromal symptoms of herpes zoster mimicked the symptoms of carpal tunnel syndrome, presumed secondary to RA synovitis. RA increases the risk of herpes zoster infection.1 In one series the use of low dose methotrexate, long duration of disease, and seropositivity were risk factors for subsequent infection.2 Gold treatment may also increase the likelihood of shingles.3

Immunocompromised patients should receive acyclovir early to avoid viral dissemination. In patients with RA the complexity of the differential diagnosis may delay diagnosis unless the possibility of herpes zoster infection is kept in mind.

Conflict of interest: none.

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Azathioprine hypersensitivity

Systemic hypersensitivity is a rare but documented side effect of azathioprine. The common adverse effects of azathioprine include

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fevers, gastrointestinal disturbances of nausea and vomiting, granulocytopenia, and hepato-cellular injury. More rarely, hypersensitivity can present with features of severe systemic infection and end organ dysfunction. Aza-thioprine is a commonly used immunosuppressive drug and is used in many medical specialties. Therefore recognising azathioprine hypersensitivity is important for all doctors. We report a case in which we suspect such an azathioprine hypersensitivity reaction.

A 32 year old Asian woman with mixed connective tissue disorder presented with a one day history of general malaise, fevers, and painful digits. Fourteen days earlier treatment had been started with azathioprine 50 mg daily, and her prednisolone increased from 5 mg to 15 mg daily owing to the development of proteinuria (1.5 g on 24 hour urine collection) and urticarial vasculitis on her lower legs.

On examination she was shocked, with a blood pressure of 70/30 mm Hg and had marked acrocyanosis (fig 1). The differential diagnosis included a lupus crisis or sepsis shock. She was treated in the intensive care unit with intravenous co-trimoxazole, metronidazole, and prostacyclin for digital ischaemia after serial cultures were taken from all major sites. Staphylococcal toxic shock was excluded, with no growth on vaginal swabs and a negative staphylococcal toxin test. After negative cultures 1 g methylprednisolone was given for a possible flare up of her connective tissue disease. The azathioprine was discontinued. She improved despite concomitant intravascular coagulation. Multiple cultures from all sites did not identify any source of sepsis.

We suspected azathioprine hypersensitivity owing to:
- The timing of the patient’s illness in relation to the initiation of azathioprine
- The presence of recognised features of hypersensitivity—fever, chills, diarrhoea, hypotension, and hepatic dysfunction
- The effect of rechallenge. Three weeks after the presenting episode, one dose of azathioprine 25 mg was given for steroid-sparing effect (the initial illness was attributed to disease flare up); the patient had a more severe and rapid hypersensitivity response requiring treatment with intravenous corticosteroids and haemofiltration in intensive care.

She improved after treatment with high dose corticosteroids. No cultures ever isolated an infective source.

This case shows the importance of recognising azathioprine hypersensitivity. Approximately 50 cases have been reported in patients with immune mediated diseases such as inflammatory bowel disease, multiple sclerosis, and immune thrombocytopenias, where the initial illness is often ascribed to sepsis or reactivation of underlying disease. Most reactions occur in the first four weeks of drug initiation.17 Hypersensitivity should always be included in the differential of fever, hypotension, and renal failure. The case was reported to the Committee on Safety of Medicines (United Kingdom).

The mechanism of the reaction is unclear. Azathioprine is composed of a nitroimidazole attached to 6-mercaptopurine. It is proposed that the imidazole component causes hypersensitivity, while the 6-mercaptopurine may cause haematological side effects. However, there are conflicting reports about the component of the drug to which the hypersensitivity reaction can be attributed.18 Fields reviewed 49 cases, where the reaction occurred equally in men and women, aged 16–76 years, and there was a wide variation in azathioprine doses.1 All patients who developed shock were also taking corticosteroids.

Support for an allergic reaction is that it occurs in only a small percentage of patients,1 and the event recurs with drug rechallenge, as occurred inadvertently in our patient. Rechallenge with azathioprine is therefore dangerous and should be done under careful observation. A hapten from the imidazole component may bind to a protein molecule to elicit type 1 hypersensitivity. Reactions mimicking sepsis may result from increased production of mediators such as tumour necrosis factor.

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