tions is unclear. The groups compared may have been too small to obtain statistical significance. These groups were similar in size to those of others, however, and since the trends were different in men and women this would seem to be an unlikely explanation. Alprenolol-induced increases in urate concentration have been shown to return baseline levels within two years despite continuous treatment. If this decreasing effect of alprenolol on urate is also true for other beta-blockers, no difference in urate values should be expected between patients treated long term with beta-blockers and those not treated with beta-blockers. In fact, no differences were found in this study, where the patients taking beta-blockers had been treated with such agents for an average of more than four years before the study. Moreover, this study confirmed the positive correlation between urate and triglyceride concentrations, and since triglyceride values correlate negatively with HDL-cholesterol concentrations, the negative correlation between urate and HDL-cholesterol concentrations is not surprising, although the underlying mechanism is unknown. Thus the lack of a difference in the plasma lipid and lipoprotein concentrations between patients on long-term beta-blocker treatment and those not on such treatment may be explained by the observed interrelationships between urate and these lipids and by assuming that propranolol, like alprenolol, has a decreasing effect on triglyceride and HDL-cholesterol concentrations, as well as on urate values.


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Arthritis and arthralgia associated with toxocarial infestation

Arthritis and arthralgia have been associated with various worm infestations—for example, filariasis and dracunculiasis. Commonly retinal lesions have been associated with toxocarial infestation, but encephalitis and disease of the liver, lung, and heart may occur. We report a case of arthritis and arthralgia associated with toxocarial infestation.

Case report

In April 1977 an 18-year-old woman presented with a seven-day history of blurring of vision in the left eye. She had kept dogs and rabbits as pets for many years. On examination visual acuity was 6/6 on the right and 6/4 on the left. Examination of the left fundus showed an area of new choroiditis and a pigmented scar near the macula. The vitreous was noted to be cloudy. White cell count was 8.3 x 10⁹/l and erythrocyte sedimentation rate 3 mm in first hour. No eosinophil count was carried out. A toxoplasma dye test gave negative results. Her symptoms settled with oral prednisolone 30 mg daily and betamethasone eye drops. In November 1977 she presented with transient swelling and stiffness of the right elbow and left wrist and ankle. On examination limitation of movement and slight swelling of the affected joints were noted. White cell count, erythrocyte sedimentation rate, liver function tests, antistreptolysin O titre, autoantibodies, rheumatoid factor, Wassermann reaction, and toxoplasma dye and haemagglutination tests were all normal. HLA typing was A2, A26, B12, and B14. Her symptoms improved on treatment with benorylate.

Between January 1978 and January 1979 she suffered repeated episodes of choroiditis in the left eye and arthralgia. A toxocarial fluorescence antibody test was performed, which was positive. The eosinophil count was 2 x 10⁹/l.

She was treated with a 21-day course of diethylcarbamazine citrate (3 mg/kg). After this her ocular and joint symptoms settled rapidly and oral steroids and non-steroidal anti-inflammatory drugs were tailed off.

Comment

We are unaware of any other reports of arthritides and arthralgias associated with toxocarial infestation. In visceral larva migrans larvae of Toxocara canis migrate through lymphatic and vascular channels throughout the body. Commonly the liver, lung, heart, and eyes are affected. Presumably joint symptoms might occur from direct joint disease as in dracunculiasis. In this case joint aspiration was not possible owing to the transient, fitting nature of her arthritis.


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Polycythaemia rubra vera and congenital deafness in monozygotic twins

Polycythaemia rubra vera occurring in families is rare, and we can find no previous report of it in monozygotic twins. In this case we describe provide an interesting parallel with the recent report of primary thrombocythaemia in uniovular twins.

Case reports

CASE 1

A 49-year-old woman who had been deaf since birth presented in 1954 after an episode of haematemesis and melaena. On examination splenomegaly was noted, and investigation showed haemoglobin concentration 14-4 g/dl; white cell count 37.2 x 10⁹/l with a normal differential; platelet count 319 x 10⁹/l, and hypercellular bone marrow. Polycythaemia rubra vera was diagnosed, and she was treated with phosphorus-32 and splenic irradiation.

Over subsequent years she suffered from chronic gastrointestinal blood loss in an undetermined site, with asymptomatic hypochromic microcytic anaemia (haemoglobin concentration 6-4-10·0 g/dl) that showed rapid rises to 17-0 g/dl and 20·4 g/dl after courses of oral iron. In March 1968, with a haemoglobin concentration of 21·3 g/dl, she suffered a fatal mesenteric vein thrombosis. Her white cell count had remained raised during the whole of the illness.

CASE 2

The twin of the first patient, who was physically identical to her and shared the same blood group (B Rh positive), presented in January 1980, aged 75 years, with a rash of some months’ duration. Skin biopsy showed the histology of eczema, which responded satisfactorily to topical steroids. The tip of the spleen was palpable, and splenomegaly was confirmed by technetium scan.

Investigations showed: haemoglobin 19·3 g/dl; red cell count 6780 x 10⁹/l; packed cell volume 0·614; white cell count 21·0 x 10⁹/l (54% neutrophils, 39% lymphocytes, 5% monocytes, 2% eosinophils); platelet count 235 x 10⁹/l; red cell volume 20·2 ml (48 ml/kg); arterial oxygen pressure 13·1 kPa; and neutrophil alkaline phosphatase score 225 (normal range 10-110). Haemoglobin electrophoresis showed a normal pattern and an intravenous pyelogram no abnormality. A bone-marrow aspirate was hypercellular with no abnormality. The bone-marrow karyotype was 46xx with no Ph¹ chromosome.

Like her twin sister, she had been deaf since birth. Audiometry showed high-tone loss of inner-ear origin compatible with congenital deafness. She was treated by regular venesection to maintain a packed cell volume below 0·45. White cell and platelet counts remained unchanged.
Shrapnel presenting with symptoms 62 years after wounding

Several reports exist of penetrating foreign bodies presenting up to 39 years after the initial injury. We describe a patient who first presented 62 years after being wounded with shrapnel.

Case report

An 85-year-old man presented in May 1979 with a four-month history of an expanding painful swelling over the left lower chest. Examination showed a large fluctuant abscess with no axillary lymphadenopathy. Radiography disclosed no underlying effusion or pulmonary change, and the only other finding was mild congestive heart failure partially controlled by diuretic treatment. Investigations showed pronounced neutrophil leucocytosis (26.0 x 10^9/l), erythrocyte sedimentation rate 17 mm in first hour, haemoglobin concentration 9.8 g/dl, and mean corpuscular volume 59%; a blood film showed hypochromia. Surgical drainage of 30 ml of sterile pus was carried out and a piece of shell shrapnel 10 x 9 mm removed. With the benefit of hindsight the shrapnel was visible on the initial chest x-ray film. Further questioning elicited that he had received minor shrapnel wounds in 1917. He had recovered rapidly and returned to the front and had subsequently been captured and spent some months as a prisoner of war. He had had no symptoms after this until he had fallen against his left side in January 1979 and had subsequently complained of increasing pain.

Comment

The longest recorded interval of a foreign body presenting many years after the initial wounding is 39 years. This was in a woman presenting with a swelling of the lip attributed to a piece of glass that had become embedded after a land-mine explosion in 1940. A piece of shrapnel that had lodged in the chest wall during D-Day worked its way to the surface 35 years later in a 62-year-old man. With the removal of the shrapnel the patient lost the ability to forecast rain. Other unusual late presentations include biliary colic caused by a grenade splinter migrating into the common bile duct 34 years after injury, choledochoduodenal fistula produced by migration of a bullet 32 years after injury, and haemorrhagic cyst formation in the region of the thyrohyoid membrane occurring 30 years after a penetrating injury.

We think that the presentation of symptoms 62 years after the original entry in the case described merits recording.


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Toxic optic neuropathy caused by benoxaprofen

Benoxaprofen is a recently introduced non-steroidal anti-inflammatory drug with useful analgesic and anti-inflammatory properties in the treatment of rheumatic diseases. It is a member of the proprionic group of acidic anti-inflammatory drugs but appears to vary from other compounds in this group since it has weak prostaglandin synthetase inhibitory activity but more effect on macrophage function. Phototoxic and photoallergic rashes and onycholysis are well-described adverse reactions, but we report on a patient who developed a partially reversible toxic amblyopia while receiving this drug.

Case report

A 65-year-old woman underwent aortic and mitral valve replacement in 1975 for valve damage secondary to previous rheumatic fever and was subsequently treated with warfarin sodium. In May 1979 she developed an inflammatory polyarthritis with a strongly positive rheumatoid factor (rheumatoid antibody haemaggulination titre 1/320), a raised erythrocyte sedimentation rate, and radiological evidence of joint erosion. In May 1980 she developed an endoscopically proved gastric ulcer and was started on cimetidine in a maintenance dose of 400 mg nightly. In July 1980 benoxaprofen 600 mg nightly was started for her active rheumatoid arthritis, with definite improvement in the degree of pain and synovitis.

On 1 September 1980 she complained of progressive blurring of vision over 10 days unassociated with any ocular discomfort. Acuities were reduced to 6/12 in each eye with a small hypermetropic spectacle correction. She showed a severe red-green colour defect and had bilateral central scotomata to a 15 mm red target (Bjerum’s screen). Electrotoretinography showed normal rod and cone responses and normal dark adaptation pattern. Visual evoked responses to a 50 flash/min pattern reversal stimulus showed low-amplitude potentials that were grossly delayed (P100 = 185 ms right and left; normal 108 ± SD 2 ms). Benoxaprofen was stopped, and over the next month her vision improved.

By March 1981 visual acuities were 6/9 in each eye. The right visual field was full to both at 3 mm white and 15 mm red target, but in the left eye there was still a relative central scotoma to a 15 mm red target. An electroretinogram remained normal and the latency of the P100 component of the visual evoked response had improved to 138 ms right and 140 ms left. The optic discs remained ophthalmoscopically normal.

Comment

A reversible toxic amblyopia has been described in patients taking ibuprofen, the first of the proprionic subseries to be introduced. Larger, prospective studies have failed to confirm this association, suggesting an idiosyncratic response to the drug.

In our patient the clinical and electrophysiological evidence suggest that the visual deterioration was the result of a toxic optic neuropathy due to benoxaprofen, her other treatment having remained unchanged. Confirmation could have been sought by reintroducing the drug, but this was not ethically justifiable. Although this is the first example of...