Azathioprine in chronic relapsing idiopathic polyneuropathy

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Summary
A 33-year-old housewife with a 14-year history of relapsing polyneuropathy of unknown cause who has apparently responded to azathioprine therapy is described. The place of this form of treatment in idiopathic polyneuropathy is discussed.

Introduction
A chronic relapsing polyneuropathy similar in clinical and laboratory findings to the acute polyneuropathy of Guillain-Barré has been increasingly recognized in recent years (Thomas et al., 1969; Ashworth and Smyth, 1969; Dyck et al., 1975). The efficacy of steroid therapy in the acute syndrome has been questioned recently (Hughes et al., 1978) but it does appear to have a place in the treatment of some chronic cases (Austin, 1958; Thomas, 1975). Other forms of immunotherapy are being evaluated including azathioprine therapy.

A patient who appears to have responded to azathioprine therapy is described and the literature on its use in chronic idiopathic neuropathy is reviewed.

Case report
The patient, then an 18-year-old shop assistant, first presented in December 1964 with a 6-week history of weakness of grip, clumsiness of hands, and an intermittent weakness of the legs. Examination revealed sluggish tendon jerks and variable limb weakness. She had increasing difficulty climbing stairs over the next 6 months, and CSF protein was then 0.82 g/l with no increase in cells. Without therapy she gradually regained full function over the following few months. In February 1967, however, she was referred with a 2-week history of tired and ‘heavy’ limbs on the right side, and depression. Antidepressant therapy appeared effective, she became pregnant and by May was much improved. Four months later, in the third trimester of her pregnancy, she developed generalized weakness of all limbs, dysphagia, exertional dyspnoea and paraesthesiae in the extremities. There was bilateral facial palsy, palatal palsy, profound weakness of all 4 limbs more marked proximally, impaired sensation in the right hand and areflexia. She was admitted and within 2 days she was bed-bound requiring help with feeding although her respiratory function remained adequate. Her CSF protein was 1.19 g/l with no increase in cells.

ACTH was started in a dosage of 120 mg daily and within 2 days there was marked improvement which was maintained and she proceeded to a successful delivery, leaving hospital with only slight general weakness and able to nurse her child, and walk without support. A relapse occurred in March 1968, with a less impressive response to a 9-week course of ACTH. She was maintained satisfactorily on oral prednisolone until the following January, during which time she was able to do light housework and to walk unsupported. Four months later, she deteriorated. Response to a further course of ACTH was poor. Towards the end of 1969, without treatment she gradually improved and during the following year her only complaint was of intermittent paraesthesiae. She became pregnant again and in March 1971 she had a bout of generalized weakness lasting about 2 months, which resolved without treatment. Her disability was slight and static for the remainder of that year, but in early 1972 there was a gradual increase in weakness and paraesthesiae with some dysphagia. She responded rapidly to a 5-day course of dexamethasone only to relapse when it was stopped.

In June 1972 azathioprine was given for exactly 4 weeks in daily doses of 100 to 200 mg. There was no significant response although 2 months after treatment was stopped she was doing light housework and walking unaided. She relapsed rapidly one month later and showed little benefit from a further course of ACTH.

Over the next 5 years her condition followed a pattern of relapses lasting from one to 6 months with partial remissions of one to 4 months. During this time she received courses of ACTH, tetracosactrin, dexamethasone and prednisolone without lasting benefit. Short courses of high dose i.v. methylprednisolone on 3 occasions seemed drama-
tically effective, but she rapidly relapsed on its withdrawal. In late 1975 weekly BCG immunostimulation appeared successful initially, but was withdrawn after 5 months when she showed deterioration despite it. Transfer factor injections from a matched patient who had successfully recovered from Guillain-Barré syndrome were unsuccessful in 1976.

During the late summer and autumn of 1977 she had shown a slow downward trend and in November was urgently admitted with a profound relapse. She could barely move her limbs, had diplopia in all directions of gaze, facial and palatal palsy with marked dysphagia and increasing breathlessness. Steroid therapy was instituted and she required ventilation. Once spontaneous respiration and oral medication was possible, azathioprine 200 mg daily was started. Within 3 weeks, however, she had deteriorated again and required repeat ventilation. In late December it was possible to start the azathioprine again and gradually withdraw steroids. There was no dramatic response, but she did show a slow upward trend and left hospital using a walking frame. By the following June her improvement was marked and she was walking unaided, doing more and more housework. She has since been maintained on azathioprine with minor alterations in dosage in response to her blood and platelet count. At the time of writing, December 1979, her power in all 4 limbs is almost normal and all her reflexes have returned. Her only symptoms are slight intermittent paraesthesiae in the extremities.

She has been extensively investigated for known causes of peripheral neuropathy including repeated checks for porphyria, vitamin deficiency, metabolic and autoimmune disorders. Although her clinical response is clear and her FEV and FVC are now normal, her motor nerve conduction velocities are still markedly slow (distal velocities for left ulnar and right lateral popliteal nerves, respectively, 13 m/sec and 27 m/sec in 1967; 15 m/sec and 26 m/sec in 1972; 26 m/sec and 25 m/sec in 1979).

Discussion

Three cases of chronic idiopathic polyneuropathy treated with azathioprine have been described. Yuill and colleagues described 5 acute polyneuropathies who improved with azathioprine and one chronic case who did not (Yuill, Swinburn and Liversedge, 1970). The latter case was a 55-year-old man with a 3-year history of polyneuropathy who was unable to tolerate the drug because of dyspepsia and it was stopped within a few days with no benefit. A 58-year-old woman who suffered from relapsing polyneuropathy for more than 3 years repeatedly responded to corticotrophin injections; she was given azathioprine for 6 months without improvement although a subsequent course of corticotrophin again produced a rapid response (Heathfield and Dallos, 1970). More recently a 54-year-old female patient with a 2-year history of polyneuropathy was described in Poland (Prusinski, Szulc-Kuberska and Zawadski, 1978). This woman was unable to walk without support, had difficulty with manual tasks and marked glove and stocking anaesthesia. Steroid therapy was ineffective but she improved within one month of starting azathioprine 100 mg. daily and was walking unaided after 2 months. Her improvement was maintained during the ensuing 9 months of therapy.

The patient described here had a relapsing chronic polyneuropathy for 14 years. For the first 6 years of her illness she had occasional remissions lasting from 2 to 12 months. During the next 6 years there were only short periods of improvement of minor degree until the institution of azathioprine therapy since when she has shown a substantial improvement for 2 years. The delay of 6 months after starting the drug before there was unequivocal improvement is compatible with the action of azathioprine and in keeping with experience of its use in other conditions. It is possible that the treatment merely coincided with a spontaneous recovery from the disease. The use of azathioprine in chronic idiopathic polyneuropathy needs further appraisal, including multicentre controlled trials.

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References


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