Case reports

from sulphhaemoglobin. An effective antidote, methylene blue, allows methaemoglobin to be reduced by a NADPH-dependent reductase system. Ascorbic acid and riboflavin can be used to reduce methaemoglobin in the homozygous deficiency but do not act rapidly enough to aid in the emergency management of acquired toxic methaemoglobinemia.

The case reported illustrates a life threatening toxic reaction to a commonly used urinary analgesic. This was the third episode of drug induced cyanosis in the patient. This case provides further evidence that oxidant drugs place heterozygotes (as well as homozygotes) for the deficiency at risk of significant methaemoglobinemia.10

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Involvement of dorsal root ganglia in Fabry’s disease

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SUMMARY Bouts of shooting pain along the extremities are common in the early stages of Fabry’s disease. No pathological explanation has been advanced to clarify the mechanism of such pain. In the present case neuronal storage of glycolipid was confined to dorsal root ganglia neurones only. It is suggested that this may explain the shooting pain in Fabry’s disease. In hereditary sensory radicular neuropathy, familial dysautonomia, and tabes dorsalis, changes in dorsal root ganglia cells cause similar clinical signs and thus it may be concluded that shooting pains in Fabry’s disease may be caused by damage to dorsal root ganglia neurones.

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Fabry’s disease is an X linked recessive, glycolipid storage disease, affecting endothelium, myocardium, fibroblasts, epithelial cells of the glomerulus, smooth muscle, cornea, glial cells, and neurones. Storage of di- and triglycosylceramide in the affected tissues is believed to be caused by a- galactosidase A deficiency.1

Early in the course of the disease the young patient suffers from bouts of severe, spontaneous shooting pain, usually in the distal part of his extremities but sometimes involving teeth, jaws, and abdomen. This characteristic symptom lacks a solid neuropathological correlation. Two possible theories have been advanced, that the pain is caused by damaged peripheral nerve endings or by depletion of small myelinated fibres in peripheral nerves.5 The present report describes a patient with Fabry’s disease in whom, in addition to the usual visceral
involvement, detailed neuropathological evaluation disclosed marked accumulation of storage material within dorsal root ganglia neurones exclusively. We suggest that the involvement of dorsal root ganglia neurones might be responsible for the presence of shooting pains in this disease.

Case report

The male patient was 37 years old at the time of death. He enjoyed normal health up to the age of 15. From then he started to have bouts of unexplained fever, frequently followed by generalised joint tenderness lasting for 2 to 3 weeks. Unrelated to these attacks he had paroxysms of lightning pain, shooting down from the hips to both knees, ankles, and feet. Similar pain in both arms radiated to the tips of his fingers. The occurrence of joint tenderness and fever led to the diagnosis of rheumatic fever by several experienced paediatricians. At the age of 18, proteinuria and hypertension were found. At the age of 25, sudden truncal ataxia and right spastic hemiparesis were found. Two years later, cardiomyopathy was diagnosed, based on the presence of chest pain, palpitations, shortness of breath, and cardiomegaly. The diagnosis of Fabry’s disease was then suspected when, in addition to the above clinical history, angiokeratomas were noticed for the first time over the umbilical area. At that stage he suffered from nephropathy, cardiomyopathy, corneal clouding, and cerebrovascular disease. The painful dysaesthesias had gradually decreased in frequency and slowly disappeared. The clinical diagnosis was confirmed by kidney biopsy in which storage of glycolipid was found. Ceramide trihexoside and trihexoside were found in the urine and a deficiency of ceramide-trihexosidase (α-galactosidase A) was found in leucocytes. The family history was compatible with X linked recessive transmission of lethal kidney disease. The patient had two maternal uncles who died of chronic renal failure. Recently a third maternal uncle was examined by us and was found to have the classical features of Fabry’s disease, which was confirmed enzymatically. The patient’s mother had mild proteinuria and showed relatively low activity of leucocyte α-galactosidase A. During the last 10 years of his life the patient had several cerebrovascular accidents resulting in pseudobulbar palsy, which finally led to his death.

Necropsy findings

The heart, spleen, and kidneys were enlarged. The heart weighed 450 g and the myocardium was pale brown with a yellowish hue. The kidneys weighed 210 g each and were also pale brown to yellow in colour. The spleen weighed 300 g and was firm. In the myocardium, microscopical examination revealed eosinophilic granules measuring up to 0.5 μ in diameter located mainly around the nuclei. These granules stained positively with PAS and Sudan black stains. Electron microscopical examination revealed the presence of osmophilic irregular bodies and a second type of lamellated bodies between the muscle fibres (fig 1). In the kidneys, within the epithelium of both proximal and distal tubules, similar bodies to those described in the myocardium were seen. In the spleen, proliferation of large histiocytes with abundant eosinophilic cytoplasm

![Image 1](http://jmg.bmj.com/10.1136/jmg.20.4.309) Myocardium: lamellar bodies and osmophilic irregular bodies are present between muscle fibres. (Original magnification × 6800.)

![Image 2](http://jmg.bmj.com/10.1136/jmg.20.4.309) Small dermal blood vessel. Endothelial cell and pericyte cytoplasm contains vacuoles and osmophilic and lamellar bodies. (Original magnification × 3300.)
was observed. In these organs, many small arterioles showed a thickened wall which stained heavily with PAS stain. In the dermis the blood vessels contained vacuoles, osmophilic bodies, and lamellar bodies, located in the endothelial cells (fig 2). In the brain, multiple encephalomalacic cysts were found in the pons, cerebellum, and both occipital lobes. Many arterioles in the brain showed extensive irregular thickening of their walls by an eosinophilic homogenous PAS positive material. Extensive study of multiple sections of the brain and spinal cord did not reveal any abnormality in the neurones or glial cells. Examination of spinal ganglia showed extremely large neurones with the nucleus and Nissl substance at the periphery of the neurone and the cytoplasm containing fine eosinophilic granules (fig 3). PAS stain showed that the cytoplasm of the neurones was loaded with PAS positive granules (fig 4) which also stained positively with Sudan black. The somatic peripheral nervous system was intact. However, small blood vessels of peripheral nerves showed extensive thickening by PAS positive material. Electron microscopy showed abundant round or irregular osmophilic bodies dispersed in the cytoplasm of the endothelial cells.

**Discussion**

This patient’s clinical history represents the classical course of Fabry’s disease. During adolescence, lightning pain, burning of hands and feet, and bouts of unexplained fever are characteristic, leading frequently to a presumed diagnosis of rheumatic fever. Later, during early maturity there is diminution and eventual cessation of ‘pain crises’, and during late maturity the patient suffers from small shrunken kidneys, cardiomegaly, hypertension, corneal clouding, retinopathy, cerebrovascular disease, and severe renal failure. Our neuropathological findings are somewhat unusual and may provide an adequate explanation for the bouts of shooting pain so characteristic of this disease. Although widespread neuronal storage of glycolipid was found by several authors, others have failed to find such changes although typical storage was always present in extraneurial tissues. Thus it seems that widespread diffused neuronal storage is not a consistent finding. However, it is generally believed that the autonomic neurones are preferentially affected.

In our patient neuronal storage was localised to sensory root ganglia exclusively.

Kocen and Thomas were the first to suggest that dorsal root ganglia cells might be involved in Fabry’s disease although they were wrong in stating that “no observations appear to have been made on the dorsal root ganglia cells”. Two years before this, Steward and Hitchcock found glycolipid storage in dorsal root ganglia neurones but there was extensive widespread involvement of other neurones as well. Similar findings were reported by Tabira et al, Kahn, and Ohnishi and Dyck. It is obvious that storage in dorsal root ganglia neurones only, as was seen in our patient, can produce bouts of shooting pain when these neurones are still alive and storing glycolipid. Cessation of pain in later stages of the disease may be the result of neuronal death.

Pain of a similar nature is a prominent sign during the course of hereditary sensory radicular neuropathy, familial dysautonomia, and tabes dorsalis. In the first two, a marked reduction of dorsal root ganglia neurones is present, while in tabes dorsalis degeneration of dorsal rootlets and, to a lesser extent, dorsal root ganglion neurones is characteristic. It may be concluded from our study...
that in Fabry's disease bouts of shooting pain can be caused by damage to dorsal root ganglia neurons.

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References

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Correction
In the paper 'Graves' disease and Down's syndrome' by McCulloch et al which was published in Journal of Medical Genetics 1983;20:133-4, 'sputum' should read 'serum' in the last line of the Summary.