Familial amyloidotic polyneuropathy with muscle, vitreous, leptomeningeal, and cardiac involvement: Phenotypic, pathological, and MRI description

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Abstract

Familial amyloidotic polyneuropathy (FAN type 1) is a rare systemic disease that causes severe and disabling peripheral neuropathy. We describe the phenotypic, radiological, and pathological characteristics of a patient with familial amyloid polyneuropathy type 1 who had evidence of motor-sensory-autonomic neuropathy, ocular vitreous deposits, diffuse leptomeningeal involvement, and hypertrophic cardiomyopathy. Muscle involvement, an infrequently reported feature, was also observed. Early recognition of the disease has significant therapeutic implications.

Key Words

Amyloidosis, cardiomyopathy, familial amyloidotic polyneuropathy, leptomeningitis, myopathy, neuropathy, vitreous deposits

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Serum electrophoresis and immunofixation, and the bone marrow were normal. Nerve conduction study of the median, ulnar, common peroneal, and sural nerves showed unexcitable motor and sensory nerves in all the four extremities. Autonomic function tests showed absent sympathetic skin response (SSR) in all four limbs and postural drop of blood pressure (54 mmHg systolic and 9 mmHg diastolic). The heart rate response to deep breathing and standing was also impaired. Routine ECG was within normal limits and 2D-echocardiogram revealed a nonobstructive cardiomyopathy. The possibilities of mitochondrial neurogastrointestinal encephalomyopathy (MNGIE), familial amyloidotic neuropathy (FAN), and chronic inflammatory demyelinating neuropathy (CIDP) with autonomic involvement was considered.

Sural nerve and biceps muscle biopsies showed multiple amyloid deposits. In the nerve it was abundant in the subperineurial and perivascular areas of the endoneurium [Figure 2a] and, to a lesser extent, in the epineurial vessel walls. Myelinated fiber loss was almost total. In the muscle the amyloid deposits were typically maximal at the periphery of the fascicles surrounding individual myofibers and in the perimysium [Figure 2b and c]. In addition, accumulations were seen within intramuscular nerve twigs [Figure 2d], in consonance with their known predilection for this site, and also in muscle spindles [Figure 2e], probably as an extension of the intraneural deposition. The deposits revealed characteristic apple-green birefringence on Congo red staining and polarization microscopy. Immunostaining for kappa and lambda light chains was negative.

MRI of the brain and spinal cord revealed extensive leptomeningeal enhancement in the post-gadolinium T1-weighted sequences. There was no signal alteration in the brain or spinal cord parenchyma [Figure 1c–f]. The patient underwent vitrectomy, following which there was improvement in his vision. Conjunctival biopsy and vitreous material also showed evidence of amyloidosis [Figure 3a–d]. After 6 months, there was slight improvement in vision without any changes in the other parameters.

Discussion

Primary amyloidosis is due to deposition of immunoglobulin light chain, is associated with paraprotein, and often manifests with the carpal tunnel syndrome rather than symmetrical and extensive sensory-motor-autonomic neuropathy. Our patient had evidence of motor-sensory-autonomic neuropathy, ocular vitreous deposits, diffuse leptomeningeal involvement, asymptomatic myopathy and hypertrophic cardiomyopathy, and normal serum electrophoresis and immunofixation. A diagnosis of FAN could be confirmed by detection of mutation and/or immunohistochemistry of the tissue with anti-human TTR antibody.[1] These investigations could not be carried out in our case due to lack of facilities. TTR is a normal tetrameric
protein that is involved in the transport of thyroxine and retinol-binding protein. It is located in chromosome 18 and is produced mainly in the liver, retinal pigment epithelium, choroid plexus, and yolk sac. It has an autosomal dominant pattern of inheritance, and a family history may not be present in more than 50% of cases.[1] More than 100 causative mutations have been described, with the commonest being FAP ATTR V30M.[3] Familial amyloidosis was first described in 1929 by DeBruyn and Stern and constitutes about 2–4% of all patients of amyloidosis.[10] There are four major types of FAP, depending on the type of aberrant proteins accumulated (e.g., TTR, ApoA1, or gelsolin). Type 1 FAN is the commonest and presents with peripheral and autonomic neuropathy, and vitreous and leptomeningeal involvement. Type 2 is the restricted form of type 1 and manifests with carpal tunnel syndrome, but no autonomic involvement. Type 3 is characterized by duodenal ulcers and early renal involvement. Type 4 manifests with cranial nerve involvement and corneal dystrophy.[4] Our patient had features of type 1 FAN.

TTR-related FAN was first described from Japan and certain European countries. It is an invariably progressive disease, with a mean age of onset of 35.3 years; it reaches the terminal stage in around 10 years.[10] The dominant phenotype shows sensory disturbances (100%), motor weakness and wasting (80%), autonomic dysfunction (>80%), and leptomeningeal and ocular involvement (20%).[3] Cerebral involvement is rare but, when present, manifests with cerebral infarction and hemorrhage, dementia, hydrocephalus, seizures, and pyramidal signs.[5] This might be due to amyloid deposits in the leptomeningeal and cerebral vessels causing ischemia and impaired autoregulation.[4] These patients have evidence of multisystem involvement, including cardiac and renal involvement, which often leads to early death. Survival is for 10–15 years. MRI studies in these patients have shown enhancement of the leptomeninges, labyrinth, dentate ligament, and vitreous body following gadolinium administration.[6,7] The present case did have extensive enhancement of leptomeninges and vitreous. Liver transplantation has been tried to halt the progression of the disease.

Our patient had demonstrable amyloid deposition in nerve, muscle, conjunctiva, and vitreous. Spuler et al. (1998) studied 13 adults with muscle weakness of 3 months’ to 4 years’ duration with a diagnosis of systemic amyloidosis and demonstrated congophilic deposits around blood vessels and muscle fibers in all the muscle specimens. The authors stressed the importance of fluorescent Congo red stain as a routine procedure in the diagnosis of amyloidosis.[8] Muscle involvement was also studied in another series of 35 patients with amyloid neuropathy, and 9 of the 25 patients of FAN had muscle involvement.[9] Amyloid myopathy in a 62-year-old man with progressive proximal weakness has been reported. This diagnosis should always be a consideration in adults with progressive neuromuscular weakness of uncertain cause.[10] Neuropathy is attributed to preferential deposition of amyloid in the sensory and autonomic ganglia, which causes axonal degeneration. Also, there is endoneurial edema due to the amyloid, resulting in ischemia. Nerve biopsy in the early stages may reveal a relatively greater reduction of small myelinated and unmyelinated fibers. In the advanced stages of the illness, large myelinated fibers also decrease. The largest and most numerous amyloid deposits are seen within the nerve fascicles.

This is one of only two case reports of FAN from India with ocular, skeletal muscle, cardiac, and leptomeningeal involvement; the other case, reported from north India, was that of a lady presenting with sensory/motor neuropathy, without any systemic involvement.[11] A high index of suspicion, with careful attention to clinical clues and histological characteristics are essential for accurate diagnosis. Genetic analysis is essential for confirmation. Symptomatic treatment for dysautonomia and visual disturbances, along with rehabilitative measures, can improve the quality of life in these patients.

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