Clinical and electrophysiological correlates of TTRala71 amyloid neuropathy

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The familial amyloidotic polyneuropathies (FAP) are autosomal dominant, length-dependent, axonal neuropathies first described in Portugal in 1958. They were later shown to be heterogeneous, resulting from tissue deposition of abnormal variants of physiological proteins, including transthyretin (TTR), apolipoprotein-A1 and gelsolin. Mutations in the TTR gene are their most common cause, and the replacement of a valine for methionine at amino-acid position 30 of the TTR protein (TTR Val30Met) is the most frequent mutation, although there are at least 70 different mutations (Human Gene Mutation Database - HGMD) identified. TTR mutations are prevalent in Portugal, Japan, Ireland, Majorca and Sweden. In contrast to Japan, where several mutations coexist, in Portugal the TTRmet30 substitution was the single known pathogenic mutation until the year 2000, when the Val28Met mutation was found. In Portuguese descents, the same mutational prevalence for the TTRmet30 seems to be present. Diagnosis is usually based on clinical manifestations, family history, and identification of the mutation.

The Brazilian patient we describe had no family history of an inherited disease, his neuropathy has unusual clinical and electrophysiological characteristics, and he tested negative for the TTRmet30 in a country of Portuguese descent, resulting in late diagnosis and inappropriate therapy for a long period of time.

CASE
A 37-year-old white male patient, of unknown descent, was referred to our hospital as a case of corticosteroid unresponsive chronic inflammatory demyelinating polyradiculoneuropathy (CIDP). Two years previously he presented burning and “pins and needles” sensations in his feet and hands, that progressed to a state of continuous pain and loss of pain sensation in his feet, legs and hands. Later, the proximal thighs, and the forearms became affected. Fourteen months after onset, he noticed difficulty in running, and some months later walking became abnormal, following trips and falls. More recently, a progressive difficulty in rising and climbing stairs appeared. Bladder and bowel functions were normal, although vomiting, attributed to a gall bladder disease, was frequent. He lost 30 kg since the onset of the disease and recovered almost 10 kg after a gall bladder surgery, which significantly improved his gastrointestinal manifestations. There was no family history of a similar condition. A nerve conduction study (NCS) performed elsewhere (data not available) revealed a demyelinating neuropathy, and he was treated with 60 mg of prednisone for 60 days, without any improvement. On examination, he walked with dropped feet, his sense of position was normal but vibration was absent in the toes, moderately decreased in the ankles, and mildly decreased in up to the knees and fingers. Pain and tactile sensations were absent to the middle third of the legs, and mildly decreased in up to the base of the thighs, to the proximal third of forearms and around the navel. There was hy-
Potrophy of foot and leg muscles, and a very mild loss of bulk of the intrinsic hand muscles. Strength, in the upper limbs, presented according to the modified Medical Research County scale (MRC), was normal proximally (5/5) and very mildly decreased distally (4/5); in the lower limbs, hip flexion and extension (4-/5), and foot flexion and extension were severely impaired (3/5). Tendon jerks were normal in the upper limbs and absent in the lower limbs. Apart a significant weight loss, the remaining clinical and neurological examination was uneventful.

His laboratory investigation was normal, including CSF analysis. Our EMG (Table) was also suggestive of an asymmetrical demyelinating sensorimotor neuropathy, and the sural nerve biopsy revealed a severe chronic axonal neuropathy with amyloid infiltrate. No evidence of demyelination or inflammatory infiltrate was found. After informed consent, testing for the TTRmet30 mutation was negative, but due to the presence of amyloid deposits in the nerve biopsy (Figure), exons 2, 3 and 4 were sequenced, revealing the TTRAla71 substitution (Figure). Immunosuppression and immunomodulatory treatments were contra-indicated and the patient was enrolled in a liver transplant program.

**DISCUSSION**

Although having a worldwide distribution and many identified mutations (>70), TTR amyloidosis is rare, has variable clinical expression, the familiar history may be unclear, and atypical presentations may occur. All these characteristics may hamper an appropriate and fast diagnosis.

Although having a length-dependent, sensory painful neuropathy pattern in the initial period of the disease, the presence of an important proximal weakness in the lower limbs in the absence of a significant distal weakness in the upper limbs, and the non-uniform demyelinating pattern observed on NCS suggested this patient had a non-length-dependent demyelinating neuropathy, compatible to CIDP, that may occasionally be heralded by pain. CSF is highly abnormal in patients with CIDP when several punctures are carried out, but one normal examination does not rule out this disease.
As there was no improvement with steroid treatment, a sural nerve biopsy was carried out, revealing an amyloid infiltrate. Subsequently, the TTR gene was screened, and the TTRala71 substitution identified.

This mutation has been previously reported just three times, none in Portuguese descents. Almeida described a Spanish family whose manifestations began in the third decade of life as a sensorimotor polyneuropathy accompanied by constipation and weight loss. Benson et al. reported a French family with a similar phenotype plus mild autonomic features and vitreous opacity. Haagsma et al. described a Dutch family whose disease began later, in the fifth decade of life, with a phenotype similar to the French family. This patient was submitted to liver transplantation, but died 14 months after, with no significant improvement.

Although FAP is defined as an axonal length-dependent neuropathy; asymmetrical demyelinating patterns on NCS have already been reported with other mutations. As no detailed NCS was provided in the previous papers about the TTRala71 mutation, we cannot associate the electrophysiological pattern found in our patient with this specific mutation.

In opposition to the three previous reports, our patient presented as a sporadic case suggesting either new mutation or low penetrance, mechanisms already known to occur with the TTR gene. Unfortunately, we were not able to test the entire family in order to clarify this point. Recently, Plantée-Bordeneuve et al. demonstrated that sporadic cases are prone to late diagnosis and its deleterious consequences, like inappropriate treatments.

Our data emphasize the importance of performing nerve biopsy in atypical or unresponsive CIDP cases, and of sequencing the TTR gene if an amyloid infiltrate is found or if there are significant small nerve fiber manifestations, even in the absence of a positive family history.

REFERENCES