Autosomal Recessive Motor and Sensory Neuropathy with Excessive Myelin Outfolding in Two Siblings

F. Barbieri, R. Santangelo, G. Capparelli, A. Ciccarelli and C. Crisci

Abstract: Two siblings, a 35-year-old male and a 37-year-old female, offspring of first cousins, presented with a hereditary motor and sensory neuropathy with type I clinical features which began to manifest at about age 10 years. Nerve biopsy in the proband showed it to be a type characterized by excessive myelin outfolding. Morphometric study revealed hypomyelination with focal thickenings due to outfoldings. Clinical, electrophysiological and morphological findings are virtually identical to those described by Ohnishi et al. The peculiarity of the neuropathological picture suggests a particular form of hereditary motor and sensory neuropathy.

Résumé: Neuropathie sensitivo-motrice autosomale récessive avec plissements excessifs de la myeline chez deux membres d’une même fratrie. Un homme âgé de 35 ans et sa sœur âgée de 37 ans, issus d’une union entre cousins germains, ont consulté pour une neuropathie héréditaire motrice et sensitive, avec des manifestations cliniques du type I, ayant débutée vers l’âge de 10 ans. Une biopsie nerveuse effectuée chez le cas index a montré des plissements excessifs de la myeline. L’étude morphométrique a révélé une hypomyélinaison avec des épaissements en foyers dus aux plissements de la myeline. Les observations cliniques, électrophysiologiques et morphologiques sont virtuellement identiques à celles décrites par Ohnishi et al. Ce tableau neuropathologique est particulier et suggère qu’il s’agit d’une forme distincte de neuropathie héréditaire motrice et sensitive.


Hereditary motor and sensory neuropathies (HMSN) are a group of heterogeneous disorders classifiable into various types according to clinical, genetic, electrophysiological and neuropathological characteristics. Of these forms, HMSN type I and III (heterogeneous groupings themselves) are hypertrophic neuropathies. In type III (Dejerine-Sottas disease; congenital hypomyelination polyneuropathy)1 inheritance is autosomal recessive, while in type I (hypertrophic Charcot-Marie-Tooth disease) it can be autosomal dominant (HMSN types IA and IB) or, rarely, autosomal recessive, X-linked dominant or recessive.2 Molecular genetics showed DNA duplication in 17p 11.2 in HMSN type IA,3,4 while linkage studies allowed gene locus mapping in HMSN type IB5,6 and in X-linked dominant form.7,8 Recently attention has been drawn to sporadic/autosomal recessive cases of congenital demyelinating motor and sensory neuropathy characterized by phenomena of focal myelin thickenings.9-13

In 1989 Ohnishi et al.14 described in two Japanese patients a hypertrophic motor and sensory neuropathy with autosomal recessive transmission, and excessive myelin outfolding. To our knowledge very few other cases of this disorder have been described in the literature since then.15

We present a clinical and electrophysiological study on two siblings, and sural nerve biopsy of the proband only, whose parents were first cousins, and who present characteristics analogous to the cases described by Ohnishi et al.14

CASE REPORTS

Patient 1

S.S., a 35-year-old male, proband, third of ten children, presented with gait difficulty by about age 10 years, followed by marked atrophy and weakness of the hand and forearm muscles. At age 15 he underwent several surgical operations to correct pes cavus and at age 28 he had surgery to correct a right inguinal hernia.

On admission to our department, steppage gait, marked atrophy and weakness of the hand and forearm muscles, atrophy of the leg anterolateral muscles, and pes cavus were present. He was unable to stand or walk on the heels. Tendon reflexes were absent. Vibration sense was distally markedly reduced at the lower limbs and slightly at the upper limbs. Touch and pain sensations were slightly decreased below the middle portion of the leg. Romberg sign was negative. On palpation the nerves were not thickened.

Needle electromyography revealed signs of severe neurogenic muscle damage. Electrophysiological tests (Table I) showed markedly reduced nerve conduction velocity, indicative of a predominantly demyelinating pathology. A sural nerve biopsy was performed.

Neurological examination of the proband’s three offspring (2 male, 1 female) gave normal results. No electrophysiological tests were performed.

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Patient 2

R.S., a 37-year-old female, second born, began to have gait difficulty by about age 12 years; a few years later she noted wasting and weakness of the hands, particularly the right. On admission to our department neurological examination showed: steppage gait, inability to stand or walk on the heels, pes cavus, bilateral atrophy of the leg muscles, marked amyotrophy of the distal third of the forearms and hands, from digit 3 to elbow, in the ulnar nerve from digit 5 to wrist; velocity was studied in the two patients along the median nerve and ulnar nerve; tendon areflexia in all four limbs; distally reduced vibration sense more marked in lower limbs. Romberg sign was negative. On palpation the nerves were not thickened.

Electrophysiological tests gave results virtually identical to those of the proband (Table 1). The patient refused to undergo biopsy of the sural nerve.

The father, deceased at age 58 from intestinal cancer, was not examined by us. He did not present difficulty in gait, pes cavus or evident amyotrophy at the hands. The mother, age 62, was found normal on neurological and electrophysiological examination. None of their remaining 8 children presented clinical signs of peripheral neuropathy.

METHODS

Orthodromic sensory (SCV) and motor conduction (MCV) velocity was studied in the two patients along the median nerve from digit 3 to elbow, in the ulnar nerve from digit 5 to wrist; MCV was also performed in the peroneal nerve, according to the technique described by Buchthal and Rosenfalck16 and by Behse and Buchthal17. The sensory responses were led off via recording electrodes and needle stimulating electrodes. Nerve conduction and sensory and motor potential amplitudes were electronically averaged. Motor conduction velocity was determined analogously according to standard procedures, using surface recording electrodes and needle stimulating electrodes. Nerve conduction and sensory and motor potential amplitudes were compared to the age-matched control groups18 (Table 1). Needle electromyography was performed in the left tibialis anterior and electron microscopy which was further subdivided; both fragments were fixed in 2.5% glutaraldehyde in cacodylate buffer (pH 7.4) at 4°C for 2h, postfixed in 2% osmium tetroxide in the same buffer for 2h, dehydrated in a graded alcohol series, and embedded in Epon 812. One-micrometer sections of the whole nerve were cut transversally and stained with 0.1% toluidine blue for light microscopy. The same sections were used for the morphometric study. The G-ratio (axon O / total fibre O) and the myelin area / axon area ratio were also calculated in 250 fibres (25% of all myelinated fibres) chosen by a systemic random method (every other field); images were analysed by VIDAS-Zeiss equipment. Ultrathin sections were cut on an LKB Nova ultramicrotome, stained with uranyl acetate and lead citrate, and examined with a Philips EM 301 electron microscope. Teased fibre procedure was not performed.

RESULTS

Conduction studies in the median and ulnar nerves are shown in Table 1. In the peroneal nerve, no sensory or motor potential could be detected. Needle electromyography showed in both patients a marked increase in duration of the motor unit potentials (+70%)18 and a normal incidence of polyphasic potentials.

Microscopically, the nerve consisted of 10 hypertrophic fascicles. The epineurium and perineurium were normal. The prominent features were a markedly reduced number of large myelinated nerve fibres, demyelinated and remyelinating axons, numerous onion bulb formations and particularly the presence of a high number of fibres with markedly irregular contour and thickness of the myelin sheaths (Figure 1). On longitudinal section these irregularities were found in both inter- and paranodal regions, and seemed to be linked to the presence of irregular myelin foldings (Figure 2).

Morphometrically, the total number of myelinated fibres was 1858 (normal values in our laboratory, based on eleven normal subjects: 5600-10700) and their density was 1.84 mm 2 (n.v.: 0.7-1.2 mm 2). The histogram of the distribution of myelinated fibre diameters revealed a unimodal curve, peaking at 4.5-5.0 μm (Figure 3). The frequency of fibres with focally irregularly folded myelin was 44%.

The ratio of myelin to axon areas was low, suggesting a general hypomyelination (Figure 4). G-ratio values (axon O / total fibre O) ranged widely: from 0.23 to 0.88 (normal 0.6 - 0.8), indicating a great variability in myelin thickness.

Ultrastructural examination showed that the parts of the myelin sheath with irregular contour and thickness seemed to present excessive outfoldings of the myelin itself toward the Schwann cell cytoplasm, thus not affecting the general form of the axon (Figure 5). However, in some cases this excessive folding was inward, toward the axon (infoldings), and the axon was

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Figure 1: Sural nerve biopsy (patient 1). Transverse section showing marked fibre loss, presence of various onion bulbs, aspects of de- and remyelination and many fibres with myelin sheath characterized by highly irregular contour and thickness. Semithin section. Toluidine blue stain. Original magnification x 640.

Figure 2: Sural nerve biopsy (patient 1). Longitudinal section showing irregular myelin foldings in internodal and paranodal portions of some nerve fibres. Semithin section. Toluidine blue stain. Original magnification x 640.

thus deformed. The general picture of this myelin modification was extremely varied, the myelin often presenting regressive signs in relation to these out- and infoldings. Outside of the regions of excessive foldings, the myelin enclosing the axon was usually disproportionately thin with respect to axon calibre. Some axons were completely demyelinated; many onion bulbs were observed.

DISCUSSION

Autosomal recessive HMSN with excessive myelin outfolding has peculiar pathological aspects. The general picture is that of a hypertrophic neuropathy, with onion bulbs and de- and remyelination phenomena; however, the diffuse presence of marked myelin changes (out- and infoldings) clearly distinguishes it from HMSN type I and III, although out- and infoldings are not specific in themselves, since they can be found in several disorders, albeit infrequently.

These two cases are further examples of this disorder. The most likely transmission modality in these cases is autosomal recessive, given the consanguinity of the parents (first cousins), despite the fact that we could not examine the father, deceased. In keeping with the cases of Ohnishi et al., the presence in ours of severe distal amyotrophy in the upper limbs and marked reduction in muscle strength. Our patients' electrophysiological and pathological pictures were virtually identical to those of Ohnishi et al., besides the ultrastructural images revealing the peculiar findings of excessive myelin outfolding, it is also important to note the results of the morphometric study (G-ratio, myelin / axon areas ratio) showing hypomyelination with foci of hypermyelination linked with myelin outfoldings.

Focal myelin thickenings with or without hypomyelination are a feature of the congenital demyelinating motor and sensory neuropathy. Unlike the cases of Ohnishi et al. and ours, in this sporadic/autosomal recessive affection the onset is before the age of two years. Morphologically, focal myelin thickenings show a more regular profile in this form, with the exception of the cases described by Vallat et al., as compared with the irregular myelin outfoldings of the form of Ohnishi et al. Whether these two neuropathies are distinct or represent two forms of a single disorder distinguishable only by onset and course is, therefore, a matter for discussion.

Focal abnormalities of the myelin sheath are also found in tomaculous neuropathy. However this condition has autosomal dominant transmission and usually manifests before age 20 with pressure palsy of a peripheral nerve; a high number of tomacula are present as myelin thickenings with a regular external surface, unlike thickenings due to outfoldings. Furthermore, ultrastructural studies show that these tomacula consist of an excessive number of normal myelin lamellae, or foldings and
The clinical data, mode of transmission, and especially the morphological findings of HMSN with excessive myelin outfolding suggest a disorder different from the better known forms of HMSN. Only molecular genetic analysis will establish whether it is a variant or a unique form of HMSN.

REFERENCES

