SHORT REPORT

DNA analysis in Finnish patients with hereditary neuropathy with liability to pressure palsies (HNPP)

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Abstract
Hereditary neuropathy with liability to pressure palsies (HNPP) (MIM number 162 500) is a dominantly inherited disorder that presents as recurrent mononeuropathies precipitated by apparently trivial traumas. The presence of a deletion in 17p11·2 was analysed in 13 Finnish families with HNPP. The deletion was found in all patients who were neurologically and neurophysiologically confirmed to have HNPP. In the problematic cases the detection of the gene defect is the method of choice in the diagnosis of HNPP. Analysis of DNA can also be used to detect clinically unaffected family members.

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Hereditary neuropathy with liability to pressure palsies (HNPP) is a dominantly inherited disorder that presents as recurrent mononeuropathies precipitated by apparently trivial traumas.1 Typically, the onset of symptoms occurs in the second or third decade. Neurological examination shows weakness and sensory loss in the distribution of the affected nerve and many patients also have signs of a more generalised neuropathy.2 The electrophysiological findings in HNPP are characteristic.3 In addition to abnormalities due to mononeuropathies, the nerve conduction velocities are generally mildly reduced and the distal motor latencies are prolonged. Markedly reduced conduction velocities are often found at typical entrapment sites (the median nerve in the carpal tunnel and the ulnar nerve at the elbow). In older patients distal muscles in the lower extremities show mild to moderate neurogenic abnormalities. Pathological changes include segmental demyelination and tomacular swellings.4 Several studies5–8 have confirmed the autosomal dominant mode of inheritance in HNPP. Recently, Chance et al9 reported a large interstitial deletion in 17p11·2 associated with HNPP in three unrelated pedigrees. The deletion spans about 1·5 Mb and includes all the markers previously shown to be duplicated in Charcot-Marie-Tooth disease type IA (CMT1A or HMSN 1A).** The candidate gene for CMT1A, PMP22, encoding a peripheral myelin protein map within the area of duplication. Furthermore, the breakpoints in HNPP and in CMT1A map to the same interval in 17p11·2, suggesting that these two disorders may be due to reciprocal products of unequal crossing over.

Here we report the results of DNA analysis in 13 Finnish families with clinical diagnosis of HNPP.

Family material
Our family material consisted of 13 families with HNPP (fig 1). Fifty one family members had been neurologically and neurophysiologically examined. Forty three of them had clinical evidence of mononeuropathies and many of them had a history of recurrent neuropathies. A minor trauma or compressive labour often preceded nerve lesions. Furthermore, most of them also had reduced tendon reflexes. None of them had systemic diseases; nor was there any evidence of drug or toxic exposure. Reduced nerve conduction velocities were found both in affected and non-affected nerves. The remaining eight family members showed no clinical or neurophysiological evidence of HNPP.

Blood samples were obtained from 33 out of 43 patients who were clinically and neurophysiologically shown to have HNPP, and from two subjects from family 4 who were clinically and neurophysiologically unaffected. Blood samples were also obtained from 18 family members who were not clinically or neurophysiologically examined and from five patients who were clinically affected but not neurophysiologically studied. A nerve biopsy was obtained from two patients.

Methods
Genomic DNA was isolated by a standard phenol-chloroform extraction procedure. The DNA was digested with the restriction enzymes MspI or EcoRI according to the manufacturer’s conditions, separated by 1·0 or 1·2% agarose gel electrophoresis, respectively, and transferred to a nylon membrane. The filters were hybridised with the 32P

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Gene copy number was estimated by visual assessment of polymorphic bands on autoradiograms and corroborated by measurement of band density and comparison with control marker (SF85) using a densitometric analysis (MCID, Image Analysis System).

Nerve biopsies were processed for semithin epoxyle section, teasing, and electron microscopy according to our modification of the previously published method.30

Results
All 33 patients from 13 families (fig 1) who were clinically and neurophysiologically (two of them also neuropathologically) confirmed to have HNPP harboured the deletion in 17p11-2. The deletion was also detected in the five patients who were clinically affected but not neurophysiologically examined. In all affected patients the presence of the deletion was determined by comparing the relative intensity of the 17p11-2 band with the constant band of the reference probe SF85 on EcoRI digestion (fig 2). The intensity differences noticed by visual inspection were confirmed by densitometric analysis. Due to the differences in labelling and hybridisation in each film, the calculated ratio of the signals for pVAW409R3a and SF85 differed significantly between affected and non-affected patients. Thus only the ratios within the same film could be compared. The ratios for the patients were about half of the ratios for the control patients in each film (table).

In all families the segregation of pVAW409R3a/MspI alleles was also followed. Nine families were informative and a loss of a pVAW409R3a allele was found. For example, in family 8 (fig 2) the two affected males (II-1 and II-2) are hemizygous for the pVAW409R3a locus. They have inherited their only 1-9 kb allele (c) from their healthy mother and no pVAW409R3a allele from their affected father. Similarly III-2 has inherited the 2-8 kb allele (a) from her healthy mother and no allele from the affected father. The deletion was also detected in four subjectively healthy subjects (III-1 in family 4, II-2 in family 7, II-2 in family 10, and I-2 in family 12) who had no history of mononeuropathies but who have not been clinically or neurophysiologically examined.

The transmission of the deletion (also male to male) through three generations was seen in two families and in eight families deletion was found in two generations confirming the autosomal dominant inheritance of HNPP.

Discussion
Recently, the HNPP locus was assigned to chromosome 17p11-2 and an interstitial deletion was shown to be associated with the disorder in three pedigrees.7 Similarly, Mariman et al 11 have detected a loss of heterozygosity of the D17S122 (pVAW409R3) locus in one large HNPP family. Here we report the presence of a deletion in 13 unrelated Finnish families with HNPP. All family members,
who were confirmed to have HNPP harbour the deletion whereas neurologically unaffected family members had two copies of the pVAW409R3 locus. The deletion was also detected in four subjectively healthy subjects aged 3, 15, 25, and 46 years, with no history of illness. Unfortunately, these four subjects have not been investigated clinically or neurophysiologically to assess whether the deletion may appear in truly asymptomatic form.

The detection of the deletion in the same chromosomal region in all reported HNPP patients indicates that HNPP is due to the deletion of pVAW409R3 in most if not all of the cases. The possible allelic heterogeneity remains to be shown by further studies, however, with markers of the critical region of 17p11.2.

In most patients with HNPP there are usually no problems with differential diagnosis, but in patients with no evident heredity, establishing a firm diagnosis may sometimes be difficult. Also, in some patients there may be problems in differentiating HNPP from other polyneuropathies, especially HMSN1. In these patients the value of the DNA analysis is obvious. In our data a loss of a pVAW409R3a allele was easily seen on analysing several members of nine informative families out of 13 studied. In these cases the diagnosis of HNPP is conclusive. When DNA analysis is used diagnostically in a single case, however, it is recommended that two 17p11.2 probes are used as in the diagnostic test for CMT1a,12 because the control and patient ranges for one probe may overlap (table).

Although HNPP is a benign disorder, we believe that the gene defect should be identified in clinically unaffected family members at an early age. If the patients are aware of their liability to develop pressure neuropathies they may avoid situations where their nerves may become exposed to chronic pressure either at work or at leisure time.

So far there have been no specific laboratory tests for HNPP. Diagnosis of HNPP has relied on medical history, clinical, neurophysiological, and neuropathological findings, and on heredity. In most of our patients the tentative diagnosis of HNPP was made when a patient with mononeuropathy showed generalised abnormalities of nerve conduction. If the patient had the typical clinical presentation and neurophysiological findings the diagnosis of probable HNPP was made. Subsequently the family members were studied. If another family member showed the typical neurophysiological findings the diagnosis was definite HNPP. Nerve biopsy was done on only two patients. We believe that nerve biopsy, which may cause considerable inconvenience to the patient, is not to be recommended in this benign disorder, all the more now with the DNA diagnostics available. The accuracy of diagnosis based on medical history, heredity, and neurophysiological findings is shown by our study: all the patients diagnosed as having HNPP harboured the 17p11.2 deletion.

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