Localization of the Gene for Distal Hereditary Motor Neuronopathy VII (dHMN-VII) to Chromosome 2q14

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Distal hereditary motor neuronopathy type VII (dHMN-VII) is an autosomal dominant disorder characterized by distal muscular atrophy and vocal cord paralysis. We performed a genomewide linkage search in a large Welsh pedigree with dHMN-VII and established linkage to chromosome 2q14. Analyses of a second family with dHMN-VII confirmed the location of the gene and provided evidence for a founder mutation segregating in both pedigrees. The maximum three-point LOD score in the combined pedigree was 7.49 at D2S274. Expansion of a polyalanine tract in *Engrailed-1*, a transcription factor strongly expressed in the spinal cord, was excluded as the cause of dHMN-VII.

The peroneal muscular atrophy syndrome, also known as Charcot-Marie-Tooth disease (CMT [MIM 118200]) is the most common inherited disorder of the peripheral nervous system, affecting 1 in 2,500 people (Keller and Chance 1999). The cardinal clinical features are weakness and wasting of the distal limb muscles, hypo- or areflexia, and pes cavus deformity of the foot. Distal sensory loss is present in some cases. Three distinct groups of peroneal muscular atrophy conditions are recognized: (1) a demyelinating form, hereditary motor and sensory neuropathy type I (HMSN I or CMT1); (2) an axonal form, hereditary motor and sensory neuropathy type II (HMSN II or CMT2); and (3) distal hereditary motor neuronopathy (dHMN, also known as distal spinal muscular atrophy, or spinal CMT) (De Jonghe et al. 1998).

dHMN accounts for ~10% of peroneal muscular atrophy syndrome (Harding and Thomas 1980). Motor and sensory nerve conduction velocities are normal, and

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electromyography typically shows features of neurogenic atrophy, consistent with degeneration of the anterior horn cell of the spinal cord. On the basis of clinical and genetic criteria, dHMN is divided into seven subtypes (Harding 1993). The genes for several dHMN subtypes and related conditions have been localized (table 1). However, no causative dHMN genes have thus far been isolated.

dHMN-VII (MIM 158580) is an autosomal dominant condition characterized by distal muscular atrophy associated with unilateral or bilateral vocal cord paralysis. Young and Harper (1980) first described the condition in a large Welsh kindred. Other families with similar phenotypes were subsequently reported (Serratrice et al. 1984; Boltshauser et al. 1989; Pridmore et al. 1992). We report the localization of the dHMN-VII gene to chromosome 2q14 by linkage and allelic association analyses in two families with dHMN-VII, including the original family, YH1, described by Young and Harper.

Detailed clinical and pathological features of members of family YH1 have been published elsewhere (Young and Harper 1980). Permission for the current study was obtained from the local research-ethics committee, and all individuals analyzed were reevaluated clinically by one of the authors (M.M.). dHMN-VII typically presented in the second decade, with weakness and wasting

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Table 1

Distal Hereditary Motor Neuronopathy Conditions for Which Predisposition Genes Have	e Been Localized
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Disorder	Features	Inheritance ^a	Locus	Reference
dHMN-II	Adult-onset distal wasting and weakness	AD	12q24.3	Timmerman et al. (1996)
dHMN-V	Upper limb predominance;		7p15	Christodoulou et al. (1995)
	occasionally pyramidal features	AD	-	
dHMN-VI	Severe infantile form with respiratory distress	AR	11q13-21	Grohmann et al. (1999)
dHMN-VII	Adult onset with vocal cord paralysis	AD	2q14	Present report
dHMN–Jerash type ^b	Juvenile onset with pyramidal features	AR	9p21.1-p12	Middleton et al. (1999); Christodoulou et al. (2000)
Congenital dSMA ^{b,c}	Congenital nonprogressive dHMN with contractures	AD	12q23-q24	van der Vleuten et al. (1998)

^a AD = autosomal dominant; AR = autosomal recessive.

^b dHMN–Jerash type and congenital dSMA were not included in the original classification of these disorders by Harding (1993).

 $^{\circ}$ dSMA = distal spinal muscular atrophy, interchangeable with dHMN.

of the small muscles of the hand and thenar eminence. Subsequently, weakness and wasting of the distal muscles of the lower limbs occurred. All but one affected individual developed a hoarse voice, which resulted from unilateral or bilateral vocal cord paralysis. The age of subjects at vocal cord involvement varied and preceded the hand changes in some individuals. As muscle wasting progressed, deep-tendon reflexes were lost, but sensation remained intact in all cases. Disease progression was slow and compatible with a normal life span. Motor and sensory nerve conduction velocities were normal, and electromyography was suggestive of chronic partial denervation.

A genomewide linkage search was performed on 10 affected individuals and 3 spouses in family YH1. A total of 230 microsatellite markers, spaced at 20-cM intervals, were selected from the ABI PRISM linkage-mapping set version 2 (Applied Biosystems) and were PCR amplified using standard protocols. Amplified markers were electrophoresed through 4.5% denaturing polyacrylamide gels on an ABI 377 DNA sequencer and were analyzed with GENESCAN and GENOTYPER software (Applied Biosystems).

dHMN-VII was modeled as an autosomal dominant trait with a penetrance of 95% by age 20 years. Unaffected individuals aged <20 years were coded as "unknown." A disease allele frequency of .0001 and equal recombination fractions in males and females were assumed. Two-point LOD scores were calculated using the MLINK program of LINKAGE, and multipoint (threepoint) LOD scores were generated using the VITESSE program. Marker allele frequencies were estimated from unrelated individuals from the pedigrees.

In the linkage search, marker D2S112 generated the highest two-point LOD score, and positive LOD scores at consecutive markers occurred only at D2S160 and D2S112 on chromosome 2q14 (table 2). To confirm and refine the location of the dHMN-VII predisposition

gene, 10 additional markers, spanning a 40-cM region on chromosome 2q14, were analyzed. DNA from an additional six unaffected relatives were included in these analyses. The order and distance from the p-telomere of the chromosome 2q14 markers are shown in table 3. Examination of the marker-allele haplotypes, segregating in affected individuals only, placed the dHMN-VII gene in a 36-cM region flanked by D2S2216 and D2S2288 (fig. 1). Meiotic recombinants in unaffected individuals IV:3 and V:7 refined the dHMN-VII region to a 20-cM interval flanked by D2S2264 and D2S2215 (fig. 1). The maximum two-point LOD score was 4.21 at D2S275, and the maximum multipoint LOD score was 5.01 at D2S100.

A second pedigree with dHMN-VII (family P2) was examined for linkage to chromosome 2q14. The clinical, pathological, and electrophysiological features in members of family P2 were similar to those in members of family YH1 and have been published elsewhere (Pridmore et al. 1992). All individuals analyzed in this study were reevaluated by one of the authors (H.H.). Twelve microsatellite markers, spanning the 20-cM dHMN-VII interval defined in family YH1, were analyzed. The marker haplotypes demonstrated that affected individuals shared a haplotype of marker alleles that was not present in their unaffected relatives (fig. 2). Positive twopoint LOD scores were observed at all markers with a maximum two-point LOD score of 2.18 at D2S100 (table 2). The maximum multipoint LOD score was 2.25 at D2S2265. At a candidate locus, these values provide strong evidence in favor of linkage. The disorder in family P2 is thus highly likely to be due to mutation of the dHMN-VII gene on chromosome 2q14.

Scrutiny of the marker haplotypes revealed the disease-associated alleles in family P2 to be identical to those segregating in family YH1 for 11 of the 12 markers initially analyzed. This was suggestive of a founder mutation predisposing to the disease in both families. To

Table 2

Two-Point LOD Scores at Chromosome 2g14 Markers in dHMN-VII Pedigrees

Marker		Two-Po	int LOD	SCORE	AT $\theta =$	
and Family	0	.01	.05	.10	.20	.30
	0	.01	.03	.10	.20	.50
D2S2216:						
YH1	-3.85	-1.23	56	30	11	04
P2	.95	.93	.83	.70	.47	.26
D2S2209:						
YH1	2.02	2.01	1.92	1.75	1.30	.79
P2	.25	.24	.20	.15	.08	.04
D2S2264:						
YH1	1.83	1.80	1.67	1.48	1.08	.66
P2	2.12	2.08	1.92	1.71	1.27	.79
D2S274:						
YH1	3.66	3.59	3.29	2.91	2.11	1.29
P2	1.79	1.76	1.61	1.42	1.02	.62
D2S293:						
YH1	4.06	3.97	3.64	3.22	2.33	1.42
P2	.38	.37	.33	.28	.20	.13
D2S160:						
YH1	.95	.91	.75	.56	.25	.06
P2	2.09	2.05	1.89	1.67	1.23	.75
D2S100:						
YH1	3.44	3.38	3.11	2.75	2.00	1.21
P2	2.18	2.14	1.98	1.76	1.31	.82
D2S275:						
YH1	4.21	4.12	3.79	3.37	2.47	1.53
P2	1.85	1.82	1.67	1.49	1.09	.67
D2S2215:						
YH1	2.19	2.20	2.14	1.96	1.46	.88
P2	.21	.20	.17	.13	.07	.03
D2S112:						
YH1	2.36	2.37	2.29	2.10	1.56	.94
P2	2.10	2.06	1.90	1.69	1.26	.79
D2S2219:						
YH1	1.10	1.13	1.16	1.09	.82	.49
P2	1.68	1.64	1.50	1.32	.94	.56
D2S2288:						
YH1	-3.85	-1.45	55	16	.10	.12
P2	.51	.50	.47	.43	.35	.25

investigate this further, 30 markers from the dHMN-VII interval were analyzed in both families (table 3). Identical disease-segregating alleles were seen in both families at 22 markers, which supports our conjecture of a common ancestral origin for the two families. Genealogy investigations subsequently confirmed that the families are indeed related, with individual I:1 in family YH1 being the uncle of individual I:2 in family P2. LOD scores calculated with the combined pedigree generated a maximum two-point LOD score of 6.63 at D2S275 and a maximum three-point LOD score of 7.49 at D2S274.

There were two separate regions in the disease-associated haplotypes in families YH1 and P2, at which the segregating alleles were identical in both families (table 3). The centromeric interval, demonstrating allelic association in the two families, was between D2S2216 and D2S160. The telomeric interval, showing allelic association, was between D2S2970 and D2S2288. The multipoint LOD scores were strongly positive within both of these intervals (fig. 3). Between the two regions there was a 7-cM interval, flanked by D2S1890 and D2S100, in which the disease-segregating alleles differed in the two families (table 3) and the multipoint LOD scores were much lower (fig. 3). The location of the markers analyzed within the D2S1890-D2S100 interval could be confidently assigned by reference to the genetic radiation hybrid and physical maps of chromosome 2q14, and they clearly bisect the disease-associated haplotype that

Table 3

Microsatellite Marker Alleles Segregating with dHMN-VII in Families YH1 and P2

Distance from p-tel		Disease- All	Two-Point LOD Score in Combined Pedigree at	
(cM)	MARKER	YH1	P2	$\theta = 0$
104	D2S2333	10	6	-3.01
111	D2S2216	6	3	-2.52
112	D2S2209	1	1	2.60
114	D2S2264ª	3	3	4.51
115	D2S274	6	6	6.04
116	D2S2229	6	6	4.27
117	D2S1897	3	3	2.67
118	D2S293	3	3	4.96
120	D2S1890	4	4	3.56
122	D2S160	5	3	.19
124	D2S308	3	6	1.41
125	D2S437	4	5	.03
127	D2S2970	3	5	2.24
128	D2S100	4	4	6.33
129	D2S2265	4	4	1.12
130	D2S2224	1	1	1.92
131	D2S347	12	12	4.05
132	D2S2339	4	4	5.18
132	D2S275	4	4	6.63
132	D2S1273	8	8	2.03
132	D2S1272	11	11	3.19
133	D2S2271	1	1	3.97
133	D2S95	6	6	6.05
134	D2S2215ª	9	9	2.84
137	D2S1260	9	9	5.36
141	D2S112	3	3	4.95
142	D2S2219	10	10	3.18
145	D2S1334	11	11	4.59
147	D2S2288	1	6	-3.06
152	D2S151	15	11	-6.34

NOTE.- The two intervals in which the disease-linked alleles in families YH1 and P2 are identical are shown in boxes. Genealogy investigations confirmed that the two families are related, and the most likely explanation for the interval between D2S1890 and D2S100, in which the disease-linked alleles differ, is a historic double recombination between the two families.

^a Recombinants at this marker in family YH1 (individuals IV:3 and V:7) define the minimal dHMN-VII interval.

Reports

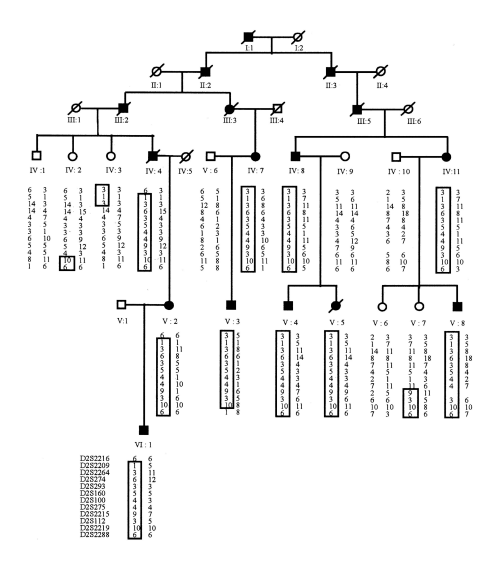


Figure 1 Haplotypes for 12 markers from chromosome 2q14 from members of family YH1. *Blackened symbols*, individuals affected with dHMN-VII. *Unblackened symbols*, unaffected individuals. Boxes around marker alleles indicate disease haplotypes present in the affected individuals. Boxed alleles in unaffected individuals IV:3 and V:7 define the minimum interval encompassing the dHMN-VII gene to the region between D2S2264 and D2S2215.

segregates in the two families. It is therefore most likely that a double recombination event has occurred between the two families, with one recombination event occurring between markers D2S1890 and D2S160 and the other between D2S2970 and D2S100. The data strongly suggest that the dHMN-VII gene does not lie between markers D2S160 and D2S2970.

The combined linkage, allelic association, and recombinant data from families YH1 and P2 localize the dHMN-VII predisposition gene either to the 8-cM region flanked by D2S2264 and D2S160 or the 7-cM region flanked by D2S2970 and D2S2215. The gene has been designated *DHMNVP* (distal hereditary motor neuronopathy with vocal cord paralysis).

No genes for distal hereditary motor neuronopathy

have been identified, and it is therefore difficult to predict what the function of *DHMNVP* might be. However, it is highly likely that the gene is expressed in the spinal cord. Two possible candidates for the dHMN-VII gene are *Engrailed-1* (*EN1*) and the neuronal PAS domain protein-2 gene (*NPAS-2*), both of which encode transcription factors that are strongly expressed in the spinal cord of mice (Davis et al. 1988; Zhou et al. 1997). *EN1* is a homeobox-containing gene that has been implicated in the regulation of axon pathfinding by association interneurons that project to motor neurons in the spinal cord (Saueressig et al. 1999). To confirm that *EN1* is within the dHMN-VII interval(s), a dinucleotide repeat located 3 kb 5' of *EN1* was identified, and amplifying primers were designed and optimized (table 3). The re-

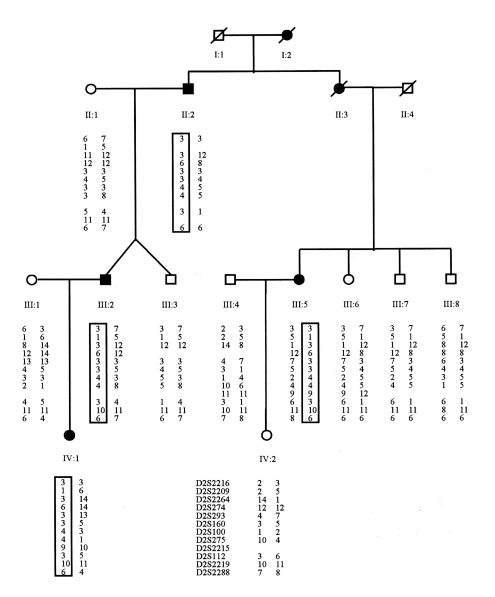


Figure 2 Haplotypes for 12 markers from chromosome 2q14 from members of family P2. Data from at-risk individuals aged <20 years have been omitted. Boxes around marker alleles indicate disease haplotypes present in the affected individuals.

peat was amplified in families YH1 and P2, and a single allele segregated with the disease in both families, generating a two-point LOD score of 3.58 (data not shown).

The first exon of *EN1* contains an imperfect polyalanine repeat of the form alanine(10)-valine-alanine(9). Pathogenic expansions of short polyalanine tracts have been reported in a number of disorders, including oculopharyngeal muscular dystrophy, in which expansions of alanine(6) to alanine(8-13) in the *PABP2* gene have been identified (Brais et al. 1998). Primers amplifying the polyalanine tract in *EN1* were designed (table 4) and analyzed in one affected and one unaffected individual from families YH1 and P2. No alteration in the size of the amplified fragment was identified in the two indi-

Table 4

Primer Pairs for PCR Amplification of EN1 Repeat and Polyalanine Tract

	Primer (5'→3')			
	Forward	Reverse		
Dinucleotide repeat Polyalanine tract	GACTAGTGGGGGCTGACTTGG GCAGGTAGAGACCCTGTCCA			

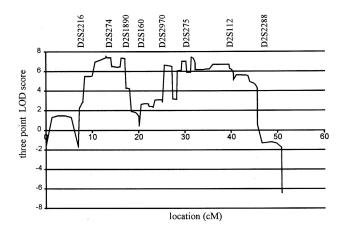


Figure 3 Multipoint LOD score for combined dHMN-VII pedigree.

viduals affected by dHMN-VII compared with the controls (data not shown). Thus, expansion of the *EN1* polyalanine tract is not responsible for dHMN-VII. However, intragenic *EN1* mutations were not investigated and may be responsible for dHMN-VII.

Hereditary motor and sensory neuropathy type IIC (HMSN-IIC [MIM 158580]) shows considerable overlap with dHMN-VII. HMSN-IIC is an autosomal dominant disorder characterized by motor and sensory involvement of the limbs, with progressive weakness of the vocal cords, diaphragm, and intercostal muscles (Dyck et al. 1994; Yoshioka et al. 1996; Donaghy et al. 1999). The distinguishing feature of HMSN-IIC is sensory involvement, which is not seen in dHMN-VII. The gene for HMSN-IIC has not been localized, but it has been suggested that HMSN-IIC and dHMN-VII may be allelic disorders that are caused by mutations of the same gene (Dyck et al. 1994). There is a precedent for such allelism, as dHMN-V and HMSN-IID have both been linked to chromosome 7p15 (Christodoulou et al. 1995; Ionasescu et al. 1996). Moreover, a kindred in which both phenotypes segregate in association with the same chromosome 7p15 haplotype has been reported (Sambuughin et al. 1998). Linkage analyses at chromosome 2q14 in HMSN-IIC pedigrees should determine whether or not the condition is likely to be due to the dHMN-VII gene on chromosome 2q14.

Other conditions with phenotypic overlap with dHMN-VII would also be worth examining for linkage to chromosome 2q14. A family with dHMN, vocal cord paralysis, and sensorineural deafness has been reported; these features may be the result of a *DHMNVP* mutation (Bolthauser et al. 1989). However, a family with some individuals having CMT, sensorineural deafness, and vocal cord paralysis has been described, in which a pathogenic *PMP22* point mutation was identified (Kovach et al. 1999). Thus, genetic heterogeneity in conditions

with peroneal muscular atrophy and vocal cord paralysis is likely to exist.

The identification of a further disease locus for dHMN adds to our growing knowledge of the disease loci involved in this subset of peroneal muscular atrophy. Isolation of genes altered in dHMN will not only improve our understanding of these disorders but will also help to clarify the relationship between dHMN and HMSN.

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Electronic-Database Information

Accession numbers and URL for data in this article are as follows:

Online Mendelian Inheritance in Man (OMIM) http://www .ncbi.nlm.nih.gov/Omim (for Charcot-Marie-Tooth [MIM 118200]; dHMN-VII and HMSN-IIC [MIM 158580]; dHMN-II [MIM 158590]; dHMN-V [MIM 600794] and [MIM 601472]; dHMN-VI [MIM 604320]; congenital distal SMA [MIM 600175])

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