**ELECTRONIC LETTER**

**POMGnT1 mutation and phenotypic spectrum in muscle-eye-brain disease**

C Diesen, A Saarinen, H Pihko, C Rosenlew, B Cormand, W B Dobyns, J Dieguez, L Valanne, T Joensuu, A-E Lehesjoki


Muscle-eye-brain disease (MEB; OMIM 253280) was first described in 1977 in Finland,1 where it is enriched because of founder effect and genetic isolation.2 MEB is now known to occur throughout the world, but Finland remains the country with the largest group of MEB patients.

MEB patients present as floppy infants with visual problems and severe mental retardation. The hypotonia is partly caused by muscular dystrophy and partly by cerebral dysfunction. Hypotonia is replaced by spasticity and contractures with increasing age.1 4 Visual failure is the result of progressive myopia, retinal degeneration, and congenital glaucoma. Juvenile cataracts develop by the age of 10 years. The presence of giant visual evoked potentials is an important diagnostic feature.4 The typical central nervous system malformation revealed by magnetic resonance imaging (MRI), referred to as ‘cobblestone complex’,5 consists of cobblestone cortex, midline deformities, flat brain stem, mild cerebellar hypoplasia, and cerebellar cortical cysts.6 Microscopically the cortex is disorganised, with an overgrowth of glia forming a thick membrane on the brain surface.7

The combination of muscular dystrophy and a severe neuronal migration defect is not exclusive for MEB, but is also seen in Walker–Warburg syndrome (WWS; OMIM 236670) and Fukuyama congenital muscular dystrophy (FCMD; OMIM 253800). The recent molecular genetic findings have provided an explanation as to why the distinct clinical features are partially shared in these three diseases. The MEB gene encodes a protein O-mannose b-1, 2-N-acetylglucosaminyltransferase (POMGnT1).10 Mutations in another enzyme involved in O-mannosylation, the O-mannosyltransferase (POMT1), were recently found in a group of WWS patients.11 Fukutin, encoded by the FCMD gene,12 is strongly suspected to play a role in glycosylation.13 The unifying feature in all these disorders is deficient post-translational glycosylation of α-dystroglycan,11 14–16 suggesting that impaired function of α-dystroglycan plays a critical role in their pathogenesis. Recently, mutations in the LARGE gene were reported in a patient with congenital muscular dystrophy, profound mental retardation, white matter changes, and subtle structural abnormalities on brain MRI, suggesting abnormal neuronal migration.17 LARGE is the human homologue of mouse Large, which is mutated in the myodystrophy mouse.18 Compatible with findings in the mouse, the human patient showed reduced immunolabelling of α-dystroglycan, adding a new member to the group of disorders characterised by congenital muscular dystrophy and a neuronal migration defect caused by deficient glycosylation of α-dystroglycan.

Thirteen MEB causing mutations covering the whole POMGnT1 gene have previously been reported.10 19 Ten of these predict protein truncation, while three are missense mutations. Expression studies of mutant POMGnT1 proteins harbouring patient mutations suggest that MEB is inherited as a loss of function of POMGnT1.10 20 We screened the exons and exon–intron boundaries of POMGnT1 in 16 MEB patients from 14 families. We identified nine new mutations and found that the mutation affecting the splice donor site in intron 17, reported previously in three patients, is the prevalent mutation in the Finnish MEB patients.

**METHODS**

**Patients and families**

Fourteen Finnish families with a total of 19 patients, 25 parents, and four healthy siblings were included in the study. Of these families, one was consanguineous, the parents being second cousins. In addition, 12 non-Finnish families with a total of 14 patients, 20 parents, and six healthy siblings were studied. These patients originated from the USA (six patients from five families), Sweden (two patients from two families), Estonia (one patient), Norway (one patient), Italy (one patient), Israel (two patients, Christian Arab siblings15), and Spain (one patient).

**Key points**

- Mutations located throughout the POMGnT1 gene encoding protein O-mannose b-1, 2-N-acetylglucosaminyltransferase underlie muscle-eye-brain disease (MEB), an autosomal recessive disorder characterised by brain malformation, congenital muscular dystrophy, and ocular abnormalities.
- MEB is enriched in the Finnish population, where a previously described mutation, c.1539+1G→A, accounts for 99% of the MEB chromosomes.
- Nine new POMGnT1 mutations in 10 patients of various ethnic origins are described, adding the number of reported MEB associated POMGnT1 mutations to 22.
- The clinical phenotypes of the non-Finnish patients in this study and published previously fall within the variation observed in the Finnish patients who are homozygous for the founder mutation, indicating that in addition to mutations in the POMGnT1 gene, other genetic and environmental factors influence the MEB phenotype.

**Abbreviations:** CDG, congenital disorder of glycosylation; CEPH, Centre d’Etude du Polymorphisme Humaine; FCMD, Fukuyama congenital muscular dystrophy; MEB, muscle-eye-brain disease; PCR, polymerase chain reaction; RT-PCR, reverse transcriptase polymerase chain reaction; SSCP, single strand conformation polymorphism; WWS, Walker–Warburg syndrome.
Finnish controls included 154 individuals, of whom 98 represented regions where the incidence of MEB is higher than the average 1:50 000 in Finland. In addition, DNA samples from 96 unrelated CEPH grandparents (CEPH; Centre d’Etude du Polymorphisme Humaine) of family members were used as controls. All the identified changes were also analysed in a panel of 40 patients with a clinical diagnosis of WWS/MEB.

The study was approved by the ethics committee of the Helsinki University Central Hospital.

**RESULTS**

We screened the whole 1983 base pair (bp) coding region and the exon–intron boundaries of POMGnTI in two Finnish patients representing different haplotypes, as well as in 14 non-Finnish patients from 12 families with a clinical diagnosis of MEB. The identified mutations are summarised in table 1 and fig 1. In addition, a previously identified coding SNP (rs2292487) and several flanking intronic polymorphisms (data not shown) in POMGnTI were identified. Haplotype analysis in Finnish patients (data not shown) suggested the existence of two mutations. A substitution of the highly conserved nucleotide G with nucleotide A at the splice donor site of intron 17 (c.1539+1G→A; reported by Yoshida et al10 and Taniguchi et al10 as IVS17+1G→A) was found to be associated with the founder haplotype (22 of 23 haplotyped chromosomes) in the Finnish patients. The mutation was found to result in skipping of the upstream exon 17 (data not shown), predicting an in-frame deletion of 42 amino acids (aa 472–513 in NP_060209). The mutation co-segregated with the disease phenotype in the Finnish MEB families, where 18 patients (1–17, 33) from 13 families had it in homozygous form, while one patient (18) had it in heterozygous form. The other mutation in patient 18 was a G to T substitution in intron 9, 5 bp from the exon–intron boundary (c.879+5G→T). RT-PCR analysis of lymphoblastoid cell RNA from this patient (data not shown) showed that the mutation abolishes the donor splice site in intron 9 and a cryptic donor site is used instead, resulting in the retention of the first 25 nucleotides from intron 9 in the mature mRNA. The mutation produces an insertion of seven new amino acids after proline 293, followed by a stop codon (Human Genome Variation Society recommended us to use (p.Pro293_Leu294ins7X8) instead of (p.Pro293_Leu294ins7;
These two mutations account for all diagnosed MEB cases in Finland. Screening of 154 Finnish control samples revealed three carriers of the c.1593+1G→A founder mutation. All three originated from western Finland, where the incidence of MEB is highest. No carriers were found for the c.879+5G→T mutation among 75 control samples studied.

Among the 14 non-Finnish MEB patients, nine different mutations in POMGnT1 were identified (table 1). Five patients had homozygous and five compound heterozygous mutations, while in four patients only one mutation was identified. One of the mutations was the c.1593+1G→A mutation common in the Finnish patients (see above) and described previously, and was found in six patients (patients 19–23 and 26) from four different countries. Eight were novel mutations that were not identified in at least 125 controls studied.

Two missense changes (c.1274G→A, p.Trp590X), and five nonsense mutations affected donor or acceptor splice sites, of which at least one introduces a premature stop codon. In patient 20, a c.879+5G→A change was identified. Interestingly, the nucleotide affected is the same as in the rare Finnish c.1593+1G→A mutation (see above), but the change is different (G→A instead of G→T). Therefore it is likely to affect splicing, even though this could not be verified, as no RNA was available from this patient. Patient 24 was compound heterozygous for two different mutations affecting the highly conserved G nucleotide of the splice donor site of intron 21 (c.1895+1G→T and c.1895+1G→A). As no RNA was available from this patient we were not able to investigate the predicted altered splicing on cDNA level. Moreover, the c.1285-2A→G change in patient 32 affecting the highly conserved A nucleotide of the splice acceptor site in intron 15 is likely to influence splicing, although this was not experimentally verified. In patient 31, we identified a 4 bp deletion of GTGA, normally existing in two copies in the immediate sequence downstream of exon 21. This change was shown to result in lack of splicing at c.1895+1 and retention of the whole mutated intron (data not shown), predicting immediate truncation of the protein (p.Val633X).

To illustrate the extent of clinical variation among patients homozygous for the Finnish founder mutation, we describe two patients with different severity of the clinical symptoms.

### Patient 1
The patient was first examined at the age of two months because of hypotonia and poor vision. Myopia of – 6 D was found and she was given spectacles, which improved her visual contact. She learned to roll over at the age of one year and to walk without support at four years. At her present age of 12 years she walks several hundred metres without support, she can feed herself, can use the toilet, and communicates with sounds and a communicator. MRI at the age of three years showed a flat pons, polymicrogyria, pachygyria, and slightly enlarged lateral ventricles (identical to fig 2B). Her muscle biopsy at the age of one year showed a few regenerating fibres and slightly increased variation in fibre size. Her creatine kinase (CK) was 781 U/l (upper limit of normal 170 U/l) at the age of one year. She has epilepsy since the age of nine years and has short absence seizures in spite of drug treatment. She has myopia of – 12 D, and a cataract has been operated on.

### Patient 2
This 37 year old man was born after an uneventful pregnancy. He was floppy and severely neurologically abnormal with poor contact from birth. His head circumference was at 0 SD while his height followed the –1.5 SD curve. A pneumonoecephalogram at the age of six months revealed enlarged lateral ventricles. Treatment with acetazolamide was started. His motor development has been minimal; by the age of three he had learned to roll from side to side and grasp with his hands. He has been treated for seizures since the age eight months. He could vocalise but contact with the surrounding world was minimal. He was placed in an institution for the mentally retarded at the age of four years. An ophthalmological examination at the age of 17 years revealed cataract in his right and myopia of –25 D in his left eye. He was examined by us at the age of 28 years. He was bedridden, with spasticity and contractures in the limbs, and he could roll his head from side to side. His CK was 1322 U/l and his brain MRI is shown in fig 2A. He had cataracts in both eyes, and on MRI his left optic bulb was markedly enlarged. He could not see with his left eye because of myopia of – 30 D, and to walk without support at four years. At her present age of 12 years she walks several hundred metres without support, she can feed herself, can use the toilet, and communicates with sounds and a communicator. MRI at the age of three years showed a flat pons, polymicrogyria, pachygyria, and slightly enlarged lateral ventricles (identical to fig 2B). Her muscle biopsy at the age of one year showed a few regenerating fibres and slightly increased variation in fibre size. Her creatine kinase (CK) was 781 U/l (upper limit of normal 170 U/l) at the age of one year. She has epilepsy since the age of nine years and has short absence seizures in spite of drug treatment. She has myopia of – 12 D, and a cataract has been operated on.

### Table 1

<table>
<thead>
<tr>
<th>Patient/origin</th>
<th>Nucleotide change</th>
<th>Exon/intron</th>
<th>Predicted effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>1–17, 33/Finland</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
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<tr>
<td>10/Finland</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>19/Sweden</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>20/Sweden</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>21/Norway</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>22/Estonia</td>
<td>c.1539+1G→A</td>
<td>IVS17</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>23/USA</td>
<td>c.1539+1G→A</td>
<td>IVS7</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>24/USA</td>
<td>c.1895+1G→T</td>
<td>IVS21</td>
<td>p.Trp409Ser</td>
</tr>
<tr>
<td>25/USA</td>
<td>c.1469G→A</td>
<td>E17</td>
<td>p.Cys490Tyr</td>
</tr>
<tr>
<td>26/USA</td>
<td>c.1539+1G→A</td>
<td>IVS7</td>
<td>p.Leu472_His513del</td>
</tr>
<tr>
<td>27, 28/USA</td>
<td>c.1469G→A</td>
<td>E17</td>
<td>p.Cys490Tyr</td>
</tr>
<tr>
<td>29, 30/Israel</td>
<td>c.1469G→A</td>
<td>E17</td>
<td>p.Cys490Tyr</td>
</tr>
<tr>
<td>31/Spain</td>
<td>c.1274G→C</td>
<td>E15</td>
<td>p.Trp425Ser</td>
</tr>
<tr>
<td>32/Italy</td>
<td>c.1285-2A→G</td>
<td>IVS15</td>
<td>p.Leu472_His513del</td>
</tr>
</tbody>
</table>

**DISCUSSION**

The clinical spectrum of diseases caused by defective protein glycosylation has rapidly increased since the description of the CDG syndrome. The diseases involving N-glycosylation have a wide spectrum of clinical phenotypes—for example,
mental retardation and multiple organ involvement. After the discovery of an O-mannosylation defect behind MEB, several other muscular dystrophies, including WWS and two new types of limb girdle dystrophy, were found to belong to the same category. Fukutin, the protein defective in FCMD, is also postulated to play a part in glycosylation.

We have studied several MEB patients of various ethnic origins, and here describe the Finnish founder POMGnT1 mutation as well as nine novel mutations underlying MEB in patients from different countries, increasing the number of published POMGnT1 mutations in MEB to 22 (fig 1). The Finnish founder mutation, c.1539+1G→A, has previously been reported in patients from Turkey, Belgium, and the USA. We also detected it in six patients of north European or north American origin, in addition to the 19 Finnish patients. This is the most common single POMGnT1 mutation underlying MEB. All other mutations reported to date, with the exception of the p.Cys490Tyr missense mutation identified in three families in this study, have been encountered in single families. Haplotype analysis suggests a single origin for the c.1539+1G→A mutation in the Finnish population. As the majority of patients with this mutation originate from Scandinavia or northern Europe, it is likely that these patients share a common origin for the mutation. Haplotype data available in seven of nine non-Finnish chromosomes bearing the c.1539+1G→A mutation support a common origin with the Finnish mutation in four chromosomes (data not shown). In three Swedish chromosomes an analysis with markers closer to POMGnT1 would be needed to find out whether their “Finnish” haplotype is restricted to a very short segment by historical recombinations or whether these mutations have independent origins. In four patients only one disease allele was identified. As no RNA was available from these patients, mutations in introns or regulatory regions affecting splicing or transcript levels may have remained undetected. Moreover, our mutation screening protocol does not allow detection of heterozygous genomic rearrangements unless they result in aberrant PCR fragments. The 22 POMGnT1 mutations in MEB described to date are evenly located throughout the gene, with no obvious mutational hotspots. In the light of previous studies, it is likely that the mutations identified by us result in loss of function of POMGnT1, as they either affect amino acids in the catalytic domain, or predict truncated proteins.

Every successful genetic mapping has to solve the question of clinical delineation of patients selected for the study. This question has continuously shadowed the molecular genetic studies of WWS and MEB, because both diseases are rare worldwide, and the clinical similarities and differences are confusing. Now that the clinical and genetic identity of these syndromes has been clarified, the question remains as to how the different mutations within each group affect the clinical phenotype. Based on a mutation study of patients of different ethnic origins, Taniguchi and co-workers suggested that the mutations near the 5' terminus cause more severe clinical symptoms than the ones near the 3' terminus. However, based on our data on the Finnish MEB patients, all except one of whom are homozygous for a single mutation, this is not the case. As exemplified with the two short case reports presented in this paper, Finnish patients homozygous for the founder mutation show wide variation in their phenotypic spectrum, with patient 1 having the mildest clinical manifestations and patient 2 representing the severe end of the clinical spectrum. The variation seen in the Finnish patients equals that seen in our molecularly defined non-Finnish MEB patients and in those reported earlier. All Finnish patients were severely handicapped, but some learned to take steps and say a few words, while others remained bed ridden with no communication or contact with the outside world. Some of the patients who have learned to walk have lost this ability.

While comparing clinical phenotypes of MEB patients, two aspects should be kept in mind. First, some features like the ocular symptoms and dystrophic changes in the muscle are progressive. Second, the treatment of severely handicapped persons has improved during the past 30 years. Children, in whom hydrocephalus has been shunted early, myopia corrected, and epilepsy treated, and who are actively rehabilitated have a better chance of using their potential. Therefore one should be cautious while comparing patients from whom limited clinical data are available. We identified no consistent genotype-phenotype correlations in the

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Figure 2. T2 weighted images of three patients with muscle-eye-brain disease (MEB) homozygous for the Finnish founder mutation. (A) In an axial image of a clinically severe case, the lateral ventricles are extremely large and the interventricular septum is missing. There is hyperintensity and volume loss of the entire white matter. The abnormal cortex is clearly visible, especially in the frontal lobes. (B) In a clinically intermediate patient with a ventriculoperitoneal shunt, an axial image shows white matter hyperintensity around the ventricles in the frontal and peritrigonal area. The rest of the white matter shows normal intensity. Note the cobblestone pattern of the frontal cortex. (C) In a clinically mild case, a coronal image shows mild dilatation of the ventricles with absent interventricular septum. The white matter intensity is normal. The abnormal cortex is again seen in the frontal lobes.
non-Finnish patients studied, which is not unexpected in the case of loss of function mutations. Based on the clinical spectrum in the Finnish MEB patients, the largest cohort of patients homozygous for a single POMGnT1 mutation, it is likely that in addition to mutations in this gene, other factors—both genetic and non-genetic—contribute to the observed phenotypic variation.

Data access
The POMGnT1 mutations described have been deposited in Human Gene Mutation Database (http://archive.uwcm.ac.uk/ucwm/mg/hgm04.html).

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Conflicts of interest: none declared

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REFERENCES
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