Genomic duplication in Dyggve Melchior Clausen syndrome, a novel mutation mechanism in an autosomal recessive disorder

E Kinning, C Tufarelli, W S Winship, M A Aldred, R C Trembath

Methods

Genotyping and DMC gene mutation analysis

Four subjects from three independently ascertained kindreds were investigated. Multicentre research ethics approval was given for the study (South East MREC, London). Pedigrees are illustrated in fig 1. Genomic DNA was extracted from peripheral blood leucocytes according to standard protocols. Microsatellite markers extending over a 3.1 Mb distance surrounding the DMC locus (fig 1 and supplementary tables 1 and 2) were amplified in multiplex polymerase chain reactions (PCR) and the products separated by electrophoresis on an ABI 377 DNA fragment analyser. Alleles were scored manually using Genescan v3.1 and Genotyper v2.0 software (Applied Biosystems, Foster City, California, USA). All 16 coding exons of the DMC gene, together with the exon–intron boundaries, were amplified as previously described. Products were purified and sequenced on an ABI 377 automatic sequencer using the fluorescent dye terminator method. Primer sequences are available as supplementary material.

Reverse transcriptase PCR and fluorescent dosage PCR

RNA was extracted either from peripheral blood collected into Paxgene® collection tubes (Qiagen, Valencia, California, USA) following the manufacturer’s instructions or from EBV transformed lymphocyte cell lines (subjects 7261 and 7345) into Paxgene® collection tubes (Qiagen, Valencia, California, USA) following the manufacturer’s instructions or from EBV transformed lymphocyte cell lines (subjects 7261 and 7345) using RNeasy kit (Qiagen), Complementary DNA was synthesised by priming with oligo-dT using First Strand cDNA synthesis kit (Roche Diagnostics, Lewes, UK). DMC cDNA was amplified in eight overlapping PCR products and size separated on 2% agarose gel (supplementary table 3). Products of an aberrant size were extracted using a Gel purification kit (Qiagen) and purified DNA sequences as described above.

Fluorescent dosage PCR to quantify locus specific genomic DNA was carried out as a multiplex assay using the primers listed in supplementary table 4 and Qiagen multiplex PCR master mix. DMC exons 2, 14, and 15 were amplified together with ALK1 exon 2 (control), samples run on an ABI 377 DNA fragment analyser using the fluorescent dye terminator method. Primer sequences are available as supplementary material.

Background: Dyggve Melchior Clausen syndrome (DMC) is a severe autosomal recessive skeletal dysplasia associated with mental retardation. Direct sequencing of genomic DNA has identified causative mutations in the gene Dymeclin (chromosome 18q12–21), with the majority predicting the generation of a truncated protein product.

Objective: To carry out molecular genetic studies in three DMC kindreds.

Results: Two novel nonsense mutations and two complex genomic duplication events resulting in exon repetition were identified.

Conclusions: Exon dosage assessment or mRNA analysis, in addition to direct genomic DNA sequencing, should be employed in the investigation of DMC affected individuals. Genomic duplication may be the causative mutation mechanism in other autosomal recessive disorders.

The skeletal dysplasias are a heterogeneous group of at least 200 distinct clinical disorders affecting approximately 1 in 4000 individuals in European populations. Dyggve Melchior Clausen syndrome (DMC, OMIM 223800), an autosomal recessive spondylo-epimetaphysal dysplasia, is characterised by short trunk dwarfism, microcephaly, a coarse facial appearance, and mental retardation. Radiographs show multiple abnormalities including vertebral platyspondyly, lacy iliac wings, laterally displaced irregularly ossified femoral heads, and a hypoplastic odontoid. Aspects of the DMC phenotype mirror the clinical features seen in the mucopolysaccharidosis known as Morquio’s disease, hence chondrocyte dysfunction was postulated to be a result of abnormal metabolite storage. However, biochemical studies have been inconsistent, such that pathogenesis of DMC remains unclear. Autozygosity mapping led to the localisation of a DMC disease gene to chromosome 18q12–21, with no evidence of locus heterogeneity. A few mutations were detected in a gene encoding the predicted transcript termed FLJ 20071. The DMC gene spans approximately 400 kb and is composed of at least 17 exons. Dymeclin, the protein product of the DMC gene, is widely expressed and several splicing isoforms have been predicted (Aceview, NCBI), and differentially sized transcripts seen on northern blot. The longest open reading frame encodes a putative transmembrane protein of 669 amino acids (75 kDa), but to date the exact nature and function of the protein is unknown. The analysis of naturally occurring mutations is instrumental in gaining functional insights and more generally in understanding the molecular mechanisms causing human genetic disease.

For a better understanding of the allelic architecture of DMC mutations, we undertook a molecular screen of affected subjects from three kindreds. We now report the identification and characterisation of four novel disease alleles including complex genomic rearrangements which, when expressed, result in exon duplication or repetition. Recent studies have suggested that regions of the human genome may be subject to regional duplication/deletion on a large scale, which we show may lead to gene disruption and subsequent disease.

Abbreviations: DMC, Dyggve Melchior Clausen syndrome; DQ, dosage quotient; STR, simple tandem repeat

The supplementary tables can be viewed on the journal website (www.jmedgenet.com/supplemental).
377 DNA fragment analyser, and gels analysed using the software packages Genescan v3.1 and Genotyper v2.0. Three independent experiments were done and used for comparison. The mean peak area for each exon was calculated for 10 control samples and used to compare with patient samples. Dosage quotients (DQ) were calculated for the exon of interest in relation to two other DMC exons or ALK1 exon 2 using the following formula, as previously described: \[DQ = \frac{\text{peak area of exon a / peak area of exon b}}{\text{peak area of control exon a / peak area of control exon b}}\]

Southern blot analysis
A total of 10μg of genomic DNA was digested with the restriction enzyme PvuII or AplII and separated on 0.8% agarose gel. After electrophoresis DNA was transferred to Zeta probe membrane (Biorad) by alkaline blotting according to the manufacturer’s instructions. A 25 ng sample of probe DNA was \(^{32}\text{P}\) labelled with 15 μCi \([\alpha-^{32}\text{P}]\text{dCTP (NEN Life Science, Boston, Massachusetts, USA) in a standard random primed reaction (Invitrogen, San Diego, California, USA). Hybridisation was undertaken at 65°C for 16 hours, membranes were washed twice at 65°C in 2×SSC/0.1% SDS (sodium dodecyl sulphate) for 20 minutes, then twice in 0.1×SSC/0.1% SDS at 65°C for 20 minutes. The filters were exposed to a Storage Phosphor screen for 16 hours and analysed on a Typhoon phosphoimager using ImageQuant software (Amersham Biosciences, Amersham, UK). DMC exon 2 and exon 14 probes correspond, respectively, to 298 base pairs (bp) of DMC cDNA including all of exon 2 and 1040 bp of DMC genomic DNA including all of exon 14, as illustrated in fig 3A.

RESULTS
Identification of new mutations in the DMC gene
In kindred DMC01, the proband (7758) was heterozygous for a paternally inherited C→T transition at nucleotide position 208 (exon 4).
Figure 2  (A) Reverse transcriptase polymerase chain reaction (RT-PCR) gel image and schematic of cDNA amplified. Dyggve Melchior Clausen syndrome (DMC) cDNA amplified with forward primer in exon 1 (5' untranslated region) and reverse primer in exon 3 produces product of 574 base pairs (bp) in control compared with approximately 800 bp in 7345. DMC cDNA amplified with forward primer in exon 13 and reverse primer in exon 15 produces product of 594 bp in control compared with approximately 1000 bp in 7263 and 7264. (B) RT-PCR gel image and schematic of aberrant subject cDNA amplified. Forward primer at the end of exon 2 and reverse primer at the beginning of exon 2 do not produce a product in control cDNA as expected. A product of 122 bp is shown in 7345. Forward primer at the end of exon 14 and reverse primer at the beginning of exon 14 do not produce a product in control cDNA but products of 79, 182 and 285 bp are shown in 7261, 7263, and 7264. A BMPR2 amplicon of 191 bp is produced for all samples.
replacing an arginine residue (R70X). The maternal chromosome harboured a mutation at nucleotide position 1363 (exon 12), also a C→T transition, which results in the introduction of a premature stop codon (R455X).

Homozygosity for all eight simple tandem repeat (STR) markers across the DMC locus at chromosome 18 was observed for each of the three affected subjects from kindred DMC02 (7345) and DMC03 (7263, 7264). Family studies identified distinct putative disease bearing haplotypes not shared by either family (fig 1). Direct sequencing of genomic DNA in each of the affected subjects failed to reveal any deleterious sequence variants. Reverse transcriptase polymerase chain reaction (RT-PCR) analysis of lymphocyte derived DMC exon 1 to exon 3 cDNA from subject 7345 (DMC02) revealed an abnormally sized product of approximately 800 bp compared with the 574 bp fragment observed in control subjects. Sequencing of the gel purified product revealed a 193 nucleotide duplication encompassing all of exon 2 including the coding region (140 bp) and the 5’ untranslated region (53 bp) (fig 2A).

Amplification of exons 13 to 15 of DMC cDNA of probands in kindred DMC03 identified approximately 1000 bp fragments, significantly larger than the 594 bp amplification seen in control subjects, and sequence analysis revealed tandem repeats of four copies of exon 14 (fig 2A). To confirm these findings, RT-PCR was carried out using primer pairs, each predicted to generate a product specific to the aberrant repeated cDNA sequences but not in the normal sequence (fig 2B and supplementary table 3). Bands of the sizes expected (122 bp in 7345 and 79, 182, and 285 bp in 7263 and 7264) were seen in affected individuals and parents from

**Figure 3** (A) Schematic illustrating probes used for Southern blotting, and restriction enzyme cleavage sites. Exon 2 probe comprises 298 base pairs (bp) of cDNA sequence including all of exon 2 in addition to partial exon 1 and 3 sequence. Following digestion of wild type sequence with Apal and probing with exon 2 probe, a fragment of 8977 bp will be produced. Exon 14 probe comprises 1040 bp of genomic DNA sequence including all of exon 14 and surrounding intronic sequence. Following digestion of wild type sequence with Apal or PvuII and probing with exon 14 probe, fragments of 3564 and 6985 bp, respectively, will be produced. (B) Southern blot analysis. Following digestion of genomic DNA with Apal and sequential probing with DMC exon 2 and exon 14 probes, fragments of the expected sizes, respectively 8977 and 3564 bp, were seen both in control samples and in DMC02 proband (7345) and father (7344). Densitometry of exon 2 to exon 14 restriction fragments showed a ratio of 2:1 in 7345, 3:2 in 7344, and 1:1 in control. (C) Southern blot analysis. PvuII digestion of genomic DNA in DMC03 subjects (7263 and 7264) and both parents (7260 and 7261) revealed a smaller abnormally sized fragment of approximately 5500 bp in addition to a fragment of the expected size (6985 bp).
whom RNA was available (fig 2B) and were not detected in control (normal) cDNA. Positive control RT-PCR amplification of a 191 bp product of BMPR2 (chromosome 2q36–37) was done and revealed a product in all samples tested.

Analysis by fluorescent dosage PCR revealed DQs for DMC exon 2 compared with either DMC exons 14, 15, or ALK1 exon 6 (chromosome 12q11–14) in individual 7345 (DMC02) of exon 2 compared with either DMC exons 14, 15, or ALK1 exon 6 in affected subjects 7263 and 7264 (DMC03) were ~4 (data not shown).

**Southern blot analysis**

Following digestion of genomic DNA with A sperm and sequential probing with DMC exon 2 and exon 14 probes, bands of the expected sizes (8977 and 3564 bp, respectively) were seen in control samples and DMC02 proband (7345) and father (7344), as illustrated in fig 3B. A maternal DNA sample was not available. ImageQuant software analysis confirmed the ratio of exon 2 and exon 14 band density as 2:1 in 7345 (affected) and 3:2 in 7344 (father) normalised to control samples. Following digestion of high molecular weight DNA with the restriction enzyme PvuII and Southern blot analysis with a DMC exon 14 probe, a band of the expected size (6985 bp) was seen both in control samples and DMC03 affected individuals (7263, 7264) and parents (7260, 7261). However, an additional fragment of approximately 5500 bp was observed in each DMC03 family member tested, as shown in fig 3C. Analysis following digestion with A sperm revealed restriction fragments of the expected size only.

**DISCUSSION**

In a screen of subjects affected with DMC we have identified four novel DMC mutations including complex genomic rearrangements that when transcribed result in exon duplication or repetition. We unexpectedly identified two nonsense mutant alleles inherited as compound heterozygote in the offspring of cousin parents (DMC01). These findings highlight a potential limitation to the method of autozygosity mapping (the method used to identify the DMC disease gene), widely adopted in gene mapping of rare recessive disorders.

Previous reports have identified 16 DMC mutations distributed among 21 kindreds (table 1) by direct sequencing of genomic DNA; hence gross deletions/rearrangements/duplications would not have been identified. These disease alleles were seen to segregate with classical DMC or the allelic disorder Smith McCort dysplasia, distinguished from DMC by the absence of mental retardation. Previous reports have identified 16 DMC mutations distributed among 21 kindreds (table 1) by direct sequencing of genomic DNA; hence gross deletions/rearrangements/duplications would not have been identified. These disease alleles were seen to segregate with classical DMC or the allelic disorder Smith McCort dysplasia, distinguished from DMC by the absence of mental retardation. Reviewing all mutations including those in this report, it is apparent that the majority (14/20 (70%)) of DMC alleles predict the generation of a truncated protein product, some as a result of a frameshift (including by exon skipping) and subsequent premature termination codon. This supports the hypothesis that loss of dymeclin function is the likely mechanism of disease pathogenesis. While the spectrum of mutations now identified provides compelling evidence that the DMC protein product is necessary for normal chondrocyte activity, the precise function of dymeclin remains unclear. The observation of diluted rough endoplasmic reticulum and multiple vacuoles in chondrocytes of affected individuals suggests that lack of dymeclin may lead to abnormal processing or defective synthesis of cartilage proteins.

The exon 2 duplication and exon 14 repetition predict the insertion of 22 and six amino acids, respectively, followed by a premature termination codon. It is expected that the transcript would be destroyed by nonsense mediated decay with subsequent failure to produce a protein product. The ATG initiator codon is located in exon 2; thus if translation was initiated in the second copy of exon 2 a normal transcript would be produced. However, exon 2 contains only 53 bp of untranslated sequence, which may not be sufficient to promote the use of this initiation site and hence translation may preferentially occur at the first ATG.

The existence of exon repetition of mRNA in the absence of genome duplication has been reported both in rat and human genes. However, fluorescent dosage analysis of PCR

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Table 1  Dyygeve Melchior Clausen syndrome (DMC) mutations

<table>
<thead>
<tr>
<th>Nucleotide change</th>
<th>DMC exon</th>
<th>Amino acid change</th>
<th>Consequence on protein</th>
<th>Phenotype</th>
<th>Ethnic origin</th>
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amplified lymphocyte derived genomic DNA provided independent confirmation of the genomic rearrangements. Southern blot analysis in subjects from kindred DMC02 did not reveal any novel restriction fragments, and a doubling of exon 2 band density in the proband (compared with exon 14 and normalised to control) suggests that the duplicated sequence includes the entire sequence between ApaI restriction sites (fig 3, panels A and B). In contrast, hybridisation with the DMC exon 14 probe after PvuII digestion of genomic DNA from DMC03 subjects revealed an aberrantly sized fragment of approximately 5500 bp in addition to a fragment of the expected size (6985 bp), as illustrated in fig 3C. ApaI digestion showed only fragments of the expected size. We interpret the above data to suggest that at least one of the external copies has been unaffected by the duplication, and hence a fragment of the expected size is produced. The exact nature of the duplication is not yet known but our results suggest the duplicated sequence is likely to start between the ApaI and PvuII restriction sites (fig 3A).

We used BLAST analysis of the DMC intronic sequence surrounding exons 2 and 14 to search for regions of homology where non-allelic homologous recombination or replication slippage may have arisen. This revealed multiple Alu repeat sequences, including several in the region between PvuII and ApaI cut sites surrounding exon 14 which Southern blot analysis indicated may be the site of initiation of duplication. Rearrangements arising at Alu elements have been reported in several conditions including Duchenne muscular dystrophy, and are estimated to account for 0.3% of human genetic diseases.16 Recent studies have highlighted the existence of variation in copy number of large segments of the human genome and suggested that this phenomenon may be more common than previously thought. Genomic imbalance was detected in approximately 50% of individuals tested in one study, including regions containing known genes.13 However, the sample sizes were small and using mathematical estimates based on Duchenne muscular dystrophy as a model, others have suggested that one in eight humans will have a segmental deletion and one in 50 a duplication.4 Interestingly, increased gene density has been found in duplons, suggesting that this phenomenon may be an important mechanism underlying mutations causing human diseases.4 It may even account for a proportion of individuals whose phenotype appears to be linked to a disease gene but in whom mutations are not identified by standard mutation screening techniques.

While we were unable to characterise exactly the real extent of genomic duplication, implicit in our observations is the assumption that the duplication events encompass sufficient intrinsic sequence to enable normal splicing. We are now seeking to characterise in more detail the duplicated intrinsic sequence in both cases, which may provide further insight into the underlying mechanism.

Conclusions
This study has identified novel mutations including complex genomic rearrangements of the DMC gene, each likely to disrupt the normal function of the protein product significantly. Genomic duplication mutations have not previously been reported in DMC and this study highlights the merit of using additional methods of mutation analysis in this severe skeletal dysplasia. Genomic duplication leading to gene disruption may be an important mechanism underlying other human genetic disorders.

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Electronic databases
The supplementary tables can be viewed on the journal website (www.jmedgenet.com/supplemental).

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