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The telomere of human chromosome 1p contains at least two independent autosomal dominant congenital cataract genes

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Aims: Multiple genetic causes of congenital cataract have been identified, both as a component of syndromes and in families that present with isolated congenital cataract. Linkage analysis was used to map the genetic locus in a six generation Australian family presenting with total congenital cataract.

Methods: Microsatellite markers located across all known autosomal dominant congenital cataract loci were genotyped in all recruited family members of the Tasmanian family. Both two point and multipoint linkage analysis were used to assess each locus under an autosomal dominant model.

Results: Significant linkage was detected at the telomere of the p arm of chromosome 1, with a maximum two point LOD of 4.21 at marker D1S507, a maximum multipoint exact LOD of 5.44, and an estimated location score of 5.61 at marker D1S507. Haplotype analysis places the gene inside a critical region between D1S228 and D1S199, a distance of approximately 6 megabases. The candidate gene PAX7 residing within the critical interval was excluded by direct sequencing in affected individuals.

Conclusion: This is the third report of congenital cataract linkage to 1ptel. The critical region as defined by the shared haplotype in this family is clearly centromeric from the Volkmann cataract locus identified through study of a Danish family, indicating that two genes causing autosomal dominant congenital cataract map to the telomeric region of chromosome 1p.

Autosomal dominant congenital cataract (ADCC) is one of the most common causes of childhood blindness. The disorder is clinically heterogeneous, with the presentation of cataract varying considerably between families and even between individuals carrying identical mutations.1 Genetically, ADCC is also highly heterogeneous, but within a given family congenital cataract tends to be monogenic with high penetrance. Autosomal dominant inheritance is the most common form, although recessive forms have been described.2 Mutations causing cataracts have been identified in genes from diverse classes including membrane proteins, cytoskeletal proteins, and transcription factors.3–5 Causative genes remain to be identified at further loci.6–8 Cataracts have been identified in genes from diverse classes including membrane proteins, cytoskeletal proteins, and transcription factors.3–5 Causative genes remain to be identified at further loci.6–8

Here, we use linkage analysis to map the ADCC gene segregating in a large family from the Australian island state of Tasmania. Using linkage and haplotype based approaches we have shown that this locus is independent of the described Volkmann-type congenital cataract locus previously mapped to this region.

Methods: We identified a six generation family originating from the Australian island state of Tasmania with ADCC. This is one of several families previously screened for mutations in crystallin genes.9 Family members were recruited with written informed consent obtained from all participating individuals or their guardians. DNA was available from 16 members of the six most recent generations and was extracted from whole blood or buccal mucosal swabs using the PureGene DNA Isolation Kit (Gentra Systems). Ethics approval for this study was obtained from the human research ethics committees of the Royal Children’s Hospital, Melbourne, the Royal Victorian Eye and Ear Hospital, Melbourne, and the University of Tasmania, Hobart, and we adhered to the tenets of the Declaration of Helsinki.

Genotyping: Forward primers for each marker were labelled with fluorescent dyes to allow detection of polymerase chain reaction (PCR) fragments on an ABI PRISM 310 genetic analyser (Applied Biosystems). Primer sequences and PCR conditions are available from the Genome Database (www.gdb.org). Microsatellites were genotyped by the Australian Genome Research Facility in or near all known ADCC loci, including: D1S468-D1S2660 (1ptel) and D1S252-D1S498 (1q21), D2S2358-D2S325 (CRYG), D3S1569-D3S1593 (BESP2), D10S192-D10S597 (PITX3), D11S898-D11S4090 (CRYAB), D12S83-D12S313 (MIP), D13S1236-D13S175 (Cx46), D15S117-D15S1033 (15q21-q22), D16S3066-D16S515 (HSF4), D17S1847-D17S836 (17q24), D17S849-D17S831 (17p13), D20S115-D20S871 (20p12-1q11), D21S1255-D21S266 (CRYAA), D22S315-D22S1154 (CRYB). Additional microsatellites located on the 1p telomere were genotyped: D15S214, D15S450, D15S244, D15S2667, D15S489, D15S228, D15S507, D15S278, D15S436, D15S2644, D15S199.

Linkage analysis: The analysis model assumed a high genetic risk, autosomal dominant model. The disease gene frequency was assumed to be rare at 0.001 and genotype penetrances of aa 0.001 Aa 0.95 AA 0.95. Two point LOD scores were calculated using MLINK. Multipoint LOD scores were calculated using approximate location scores using the SimWalk2 program7 and exact LOD scores using the Vitesse program.8 Analysis using Vitesse was limited to three markers (D1S507-D1S2728-D1S2644).
because of the size of the kindred. The genetic positions of the markers were taken from the Decode17 and Marshfield18 genetic maps and the marker order confirmed using their physical position in the human genome (genome.ucsc.edu/cgi-bin/hgGateway). Allele frequencies were taken from the genome database. Individuals 16.10 and 16.11 are monozygotic twins and hence only patient 16.10 was included in the linkage analysis.

Re-sequencing of \textit{PAX7} in affected patients

The coding region of the candidate gene \textit{PAX7} was resequenced in two affected patients as well as one unaffected individual. Primers were designed to amplify each of the eight identified exons (primers available on request) from genomic DNA. Amplified products were purified using spin columns (MoBio) and sequencing was performed on an ABI PRISM 310 genetic analyser (Applied Biosystems).

RESULTS

Clinical notes from two patients in the Tasmanian pedigree describe the phenotype as a complete cataract (16.05 and 16.14) although most affected members are elderly and aphakic, with no preoperative clinical notes available. The median age of diagnosis was less than 1 year (range birth to 9 years), however most were diagnosed shortly after birth and the mean age of cataract surgery was 3 years 4 months (range 0–9 years). Twelve of the 13 patients had bilateral cataract surgery and were aphakic (eight had acuities <6/60 and four had acuities 6/19–6/60), corrected with spectacles. There is one affected member (16.16, fig 1) who is an obligate carrier but has a relatively mild phenotype, phakic with good vision (6/6–1 6/7.5+2) but not available for slit lamp examination. Eleven of 12 of the aphakic individuals had nystagmus. Two individuals had exotropia (16.02, 16.08) and four esotropia (16.03, 16.06, 16.07, 16.11). Four individuals had bilateral aphakic glaucoma (16.03, 16.05, 16.08, 16.15). No other ocular or systemic abnormalities were noted.

Linkage analysis using the microsatellite markers genotyped across the known cataract loci detected significant linkage at the telomere of chromosome 1p. Two point LOD scores of greater than three were achieved at markers D1S507 and D1S2644, with LODs of 4.21 and 3.23 (\( h = 0 \)) respectively (table 1). Multipoint analysis achieved maximum location scores\(^{19}\) and LOD scores\(^{20}\) of 5.61 and 5.44 respectively (table 1). Combined, these results indicate significant linkage to this region making it highly probable that the gene segregating in this family maps to this locus. All other regions tested failed to reach significance or were inconsistent with linkage (data not shown).
The inferred haplotypes across 1pter are displayed in figure 1. Two informative recombinants, 16.10 and 16.14, define a critical region between centromeric marker D1S199 and telomeric marker D1S228. The physical location of these markers indicated that the region containing the gene spans a physical distance of approximately 6 megabases (Mb).

Eiberg et al.7 studied a large Danish pedigree affected with Volkmann-type congenital cataract to map the gene for this disorder to the interval between the 1p telomere and D1S214. In the Tasmanian family the recombinants 16.14 and descendants of 16.20 all indicate that the gene segregating in this family is located at a position 6 Mb telomeric to D1S214 (fig 2). As there is 6 Mb between the two critical regions defined by these studies, it is likely that the genes in the Tasmanian and Danish pedigrees are not allelic and provides strong evidence that there are at least two genes predisposing to ADCC on the telomeric region of chromosome 1p.

The PAX transcription factors have been implicated in ocular development and congenital cataract.19 The presence of the PAX7 gene inside the critical region defined by our study made it an excellent candidate gene. However, re-sequencing of the coding region of this gene in two of the affected patients from the Tasmanian family did not identify any mutations, apart from a common synonymous single nucleotide polymorphism (SNP) that has already been identified as a part of the human genome project (rs2743201) and therefore was very unlikely to have any influence on disease. While we cannot exclude that a mutation in a regulatory region of this gene may have a role, the absence of a coding region mutation suggests that the disease allele may lie in another gene within the critical region.

**DISCUSSION**

The linkage analysis results from the Tasmanian family clearly indicate the existence of a susceptibility gene at 1p36. This is the third report of significant linkage in congenital cataract to the telomere of 1p. Eiberg et aII studied a large Danish pedigree affected with Volkmann-type cataract and mapped the predisposition gene for this disorder to the interval between the 1p telomere and D1S214. This region is 6 Mb telomeric to the region defined by the recombinant individuals in the Tasmanian family. Both the Danish and our own results have reached highly significant linkage (Danish pedigree LOD = 14.04 Tasmanian pedigree LOD = 5.44) and with the physical positions better determined via the human genome project, we can determine that there is no ambiguity in the marker’s position used by either study. The lack of overlap between the segregating haplotypes from the two studies makes it likely that the genes in the Tasmanian and Danish pedigrees are not allelic and there are two genes predisposing to congenital cataract on the telomere of 1p.

The second report of linkage to this region was in a British family presenting with ADCC.8 Based on the haplotype in this British family, and particularly a homozygous stretch in a critical recombinant, the authors suggest that the gene segregating in this family might be allelic to the Danish Volkmann cataract pedigree. However, in reviewing the haplotype segregating in the British family in light of the region outlined here, we believe it also possible, if not more likely, that the gene in the British family could be allelic to the Tasmanian pedigree.

Interestingly, the critical region defined by the Tasmanian family also overlaps with that of the recessive primary congenital glaucoma locus, GLC3B.20 Four of 13 patients from the Tasmanian family had aphakic glaucoma. This is relatively high for congenital cataracts, suggesting possible involvement of the same gene with both congenital glaucoma...
and cataract. However, the phenotypic differences between congenital cataract and congenital glaucoma, and genetic differences (a dominant versus recessive inheritance), make it equally likely that these disorders are not caused by the same gene. The mapping of the congenital cataract, congenital glaucoma and Volkman-type cataract loci in this region indicates that the telomeric region of the chromosome 1 contains a number of genes that are relevant to both eye development as well as congenital eye disorders.

The physical distance defined as the critical region in the Tasmanian family is about 6 Mb. Inside this interval are 30 named genes, of which 18 are clearly noted to be expressed in the eye (www.ncbi.nlm.nih.gov/unigene). Also in this region are many more undescribed mRNAs and algorithm predicted genes. This makes the positional cloning of the causative gene a formidable task at this point. The identification of additional families linked to this region and containing informative recombinants may further restrict this region so that a positional cloning attempt is realistic. Identification of this gene and others that cause congenital cataract increases our understanding of the molecular events that lead to the development of cataract as well as the development of the eye.

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Competing interests: none declared

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REFERENCES


