Fine localisation of the gene for central areolar choroidal dystrophy on chromosome 17p

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Abstract
Central areolar choroidal dystrophy (CACD) is a retinal disease which causes progressive profound loss of vision in patients during middle age. The disease is inherited as an autosomal dominant trait and shows genetic heterogeneity. Mutations in the peripherin-RDS gene on chromosome 6 have been reported in affected members of families transmitting the disease. A new locus at chromosome 17p13 was identified recently by a genome wide linkage search in members of a large Northern Irish family. We now report the refinement of the critical region for this gene to an interval of approximately 5 cM flanked by polymorphic markers D17S804 and CHLC GATA7B03.

Keywords: Central areolar choroidal dystrophy; genetic linkage; chromosome 17p

Central areolar choroidal dystrophy (CACD) is a rare inherited retinal dystrophy which causes progressive profound loss of vision in patients during their fourth decade. A pathognomonic zone of atrophy affecting retina, retinal pigment epithelium, and choriocapillaris develops in the macular area of the eye. Patients develop absolute central visual scotomas leading to severe visual disability by the seventh decade. The gene causing CACD in a large Northern Irish family was mapped in 1996 by linkage to a cluster of markers at 17p13. The maximum two point lod score of 4.7 was found at θ=0.00 with D17S796, with a maximum multipoint lod score of 5.65 at D17S938. Analysis of recombination suggested a location in the interval of about 16 cM between D17S5 and D17S804.

Patients and methods
We have now typed several new markers in this family and have refined the critical CACD gene region. The MLINK and LINKMAP programs of FASTLINK (version 3.0) were used for computerised two point and multipoint analysis of data, assuming autosomal dominant inheritance with a disease allele frequency of 0.0001 and complete penetrance in members of generations I, II, and III. Members of generation IV were assigned as affected or normal on the basis of the presence or absence of electrophysiological abnormalities thought to be indicative of early disease to one of four age related liability classes ranging from 80-95% likelihood of the assigned disease status. This was why markers D17S829, D17S844, and D17S938 were chosen. The critical CACD region is in young family members before the onset of pathognomonic features in the fourth decade.

Subject IV.13 was the youngest family member assessed (aged 14) and had some abnormal signs of possible disease; however, similar signs were also found recently in her unaffected mother. This girl has inherited the normal haplotype from her affected father at all marker loci in the gene region and it is therefore highly probable that she does not carry the disease allele. She was previously assigned an 80% risk of carrying the CACD gene, but has now been excluded from statistical analysis.

Results and discussion
Two point lod scores between the disease locus and markers (table 1) show a maximum of 5.65 at D17S1832. Haplotypes for markers D17S1876, D17S1810, D17S1832, D17S76, GATA7B03, D17S1844, and D17S191, spanning an interval of 8 cM of chromosome 17p13, are shown in fig 1. The CACD gene region is defined by the position of recombination in subjects III.3 and II.6. III.3 is non-recombiant at D17S1832, but recombinant at D17S1810, D17S1876, and more distal markers. The absence of DNA from the parents of II.6 makes his haplotypes more difficult to analyse. He was previously known to have inherited an allele that was not associated with the disease haplotype at D17S756, and was non-recombiant or uninformative at all other markers tested. The possibility of a mutation in his D17S756 allele was considered; however, he is now found to be recombinant also at GATA7B03, D17S1844, and D17S191 which flank D17S76. This subject was not fully informative for D17S1353 and the microsatellite at the TP53 locus. The disease gene is therefore restricted to the interval between D17S1810 and GATA7B03, a distance of less than 5 cM.

Several inherited eye disorders have recently been mapped to chromosome 17p (fig 2). They include the autosomal dominant progressive
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Figure 1 Pedigree of the CACD family showing types for markers D17S1876, D17S1810, D17S1832, D17S796, GATA7B03, D17S1844, and D17S1791 in order from top to bottom. Alleles transmitted by the affected parent are shown on the left of each pair. The haplotype linked to the disease gene is shaded. I.1 and I.2 are of unknown status.

Figure 2 Physical and linkage map of markers on chromosome 17p. Sex averaged distances between markers are shown in cM. The locations of candidate genes and linked eye disorders are also indicated.

corne dystrophies CORD5 and CORD6, autosomal dominant retinitis pigmentosa (RP13), autosomal dominant cone degeneration (ADCD), Leber's congenital amaurosis (LCA), and anterior polar cataract. It is interesting to speculate whether the same gene could be involved in CACD, CORD5, and CORD6. The CORD5 family described shows
central chorioidal atrophy which is also a typical feature of CACD and these disorders have a similar age of onset. Affected members of the CORD6 family have an earlier age of onset, but full clinical details have yet to be reported. If mutations within a single gene are responsible for these retinal dystrophies, the search for the gene can be focused on the 1.7 cM interval between D17S938/D17S796 and the TP53/D17S1353/GATA7B03 cluster.

LCA maps in many families to the interval between D17S938 and D17S1353 and in 1996 mutations in the retinal guanylate cyclase (GC1) gene were identified in affected subjects. GC1 lies within the CACD region. The entire coding region of the gene was sequenced in affected and normal members of the CACD family, but no mutations were found (data not shown). This was not unexpected because both LCA and the autosomal dominant cone degeneration which also maps to 17p have more diverse fundal appearances and are therefore unlikely to be caused by the gene which is responsible for CACD.

Several other candidate genes for retinal disorders map to 17p (fig 2). These include phosphatidylinositol transfer protein (PTITPN), pigment epithelium derived factor (PEDF), recoverin (RCV1), and β-arrestin 2 (ARRB2). The PTITPN gene has homology with the Drosohila retinal degeneration B gene (rdgB) which is responsible for light enhanced retinal degeneration. The PEDF gene product possesses both neurotrophic and neuronal survival activities and recoverin mediates the recovery of the dark current after photoactivation in the retina. These genes, however, map outside the CACD region. The ARRB2 gene is expressed in the retina and is thought to be involved in regulation of receptor binding to a G protein. It maps to 17p13, but has not been finely localized. Regarding the physical position of ARRB2, the Genebridge 4 radiation hybrid map suggests that the gene probably lies distal to the CACD region, but it has not been placed precisely with respect to the polymorphic genetic mapping markers.

The molecular basis of most of the retinal disease genes which have been mapped recently to chromosome 17p remains to be established. Several interesting candidate genes are expressed and have important roles in the retina, but are not yet associated with any retinal phenotypes. Advances in molecular genetics have provided the tools which should allow the genes responsible for these diseases to be found. At present it is unclear whether CACD and the cone rod dystrophies of CORD5 and CORD6 are caused by mutations in one or a cluster of genes. Reducing the region of the CACD gene will assist its identification and will also enable the genetic influence of this locus to be assessed in age related macular degeneration and related macular dystrophies.
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