Linkage refinement localises Sorsby fundus dystrophy between markers D22S275 and D22S278

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Abstract
Sorsby fundus dystrophy is an autosomal dominant disorder which both clinically and histopathologically bears striking similarities to age related macular degeneration, one of the leading causes of blindness in the developed world. Recent studies have suggested a genetic localisation of the disease to chromosome 22q in a large genetic interval of approximately 25 cM. Independent genetic linkage analysis in a six generation British pedigree confirms linkage to the chromosome 22q region. A maximum two point lod score of 7·09 with no recombination was obtained with marker D22S280. Haplotype data positioned the disease between loci D22S275 and D22S278, thus significantly reducing the region on chromosome 22q where the gene is located.


Sorsby fundus dystrophy (SFD) was originally described as an autosomal dominant retinal dystrophy with complete penetrance in five British families.1 Affected subjects are symptomatic by the fourth decade of life, and loss of central vision occurs either through choroidal neovascularisation or by geographic atrophy.2 The disease is characterised by delayed choriocapillaris filling on fluorescein angiography2 with later deposition of confluent subretinal lipid containing material at the level of Bruch's membrane.3 These characteristics have features in common with pathological changes seen in age related macular degeneration,4 the commonest cause of blindness in Westernised society. Thus, elucidation of the precise pathogenic mechanism in SFD may shed light on the pathogenesis of age related macular degeneration.

Recently linkage analysis in a Canadian SFD family of Irish origin has shown localisation of the disease to chromosome 22q.13-qter in a 25 cM region flanked by markers D22S275 and D22S274.5 To determine if one of the original families described by Sorsby et al6 was linked to chromosome 22q, conventional linkage analysis was carried out.

Patients and methods
From a six generation pedigree, 25 family members were studied including 12 affected subjects, 12 unaffected subjects, and two spouses (figure A). Subjects were only confirmed as unaffected if they were over 55 years of age, with normal fundal appearance on clinical examination and fluorescein angiography, and had normal vision. Informed consent explained the nature of the study was obtained from each patient. Seven microsatellite markers6 mapping to the chromosome 22q SFD region7 were used to genotype subjects. Two point lod scores were determined using MLINK7 and multipoint analysis was undertaken using LINKMAP7.

Results
LINKAGE ANALYSIS
Two point lod scores for markers mapping to chromosome 22q in the region previously identified as linked to SFD are presented in the table. A maximum two point lod score of 7·09 (θ = 0·00) with marker D22S280 confirmed tight linkage in our family to the SFD locus on chromosome 22q13.1. In addition, two other markers, D22S273 and D22S281, also showed no recombination with the disease giving two point lod scores of 2·80 and 6·19, respectively.

MULTIPOINT LINKAGE ANALYSIS
A series of four three point analyses were performed with the linked markers to generate a multipoint map of the region and is presented as a single graph (figure B). Two markers and disease were used in each analysis to calculate the likelihood of the disease gene being located between these two markers. The data suggested that the SFD disease locus was most likely to be in the interval between D22S273 and D22S280 with a peak lod score of 8·51, or between D22S280 and D22S281 with a peak lod score of 7·82. The two lod confidence interval suggested that the disease locus was
Discussion
This study has confirmed linkage in our family to chromosome 22q13.1. The differences in clinical presentation of the disease that have been reported may represent phenotypic variation because of different mutations within the same gene, similar to that which has been seen in peripherin/RDS mutations. This hypothesis can be confirmed when the disease gene is identified.

The identification of critical recombinants within the pedigree has significantly improved the localisation of the disease locus to chromosome 22q13.1 within an 8 cM interval in this family. There are two genes in the region which are possible candidates for the disease. The A1 adenosine receptor was localised to chromosome 22q13 and has been shown to be expressed in the retina. The second candidate is TIMP-3 which was recently mapped to chromosome 22q13. TIMP activity has been detected in interphotoreceptor matrix which lies between the neural retina and the retinal pigment epithelium. These two genes are currently being assessed for a role in the molecular aetiology of SFD. If they are not involved in the disease pathogenesis then generation of new markers in the region and positional cloning will be used to identify the disease causing gene for SFD.

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