A Common Locus for Late-Onset Fuchs Corneal Dystrophy Maps to 18q21.2-q21.32

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PURPOSE. To identify the genetic basis of late-onset Fuchs corneal dystrophy (FCD).

METHODS. Phenotypes and genotypes at 1107 short tandem repeat polymorphism markers were obtained for 43 affected and 33 unaffected individuals in three large families. Two-point genetic linkage analysis was performed with MLINK and multipoint linkage with SimWalk 2.89.

RESULTS. In each family, the most significant cluster of two-point lod scores mapped to chromosome 18, at 18q21.2-q21.3. The highest two-point lod score for each family was at D18S1129, with scores of 3.41, 2.89, and 2.45, with a combined two-point lod score of 7.70. Multipoint analysis yielded a maximum score of 5.94 at D18S1129 for a dominant Mendelian trait exhibiting 85% penetrance and 15% phenocopy rate. Disease interval haplotypes for each family are different.

CONCLUSIONS. FCD2, at 18q21, is the second genetic locus identified for late-onset FCD. Presence of this same locus in all three families may indicate its widespread involvement in late-onset FCD. Allelic differences between disease-associated haplotypes in the families leave open the possibility of independent mutations in the same gene. The incomplete penetrance and high phenocopy rate observed at FCD2 suggest that the origin of FCD in these three families is complex and also depends on other genetic loci or environmental factors. (Invest Ophthalmol Vis Sci. 2006;47:3919–3926) DOI:10.1167/iovs.05-1619

Fuchs corneal dystrophy (FCD), first described in the early 20th century,1,2 is a common age-related disorder that affects 4% of those older than 40 years.3 In its early stages, FCD is characterized by the formation of guttae, which are microscopic excrescences of Descemet’s membrane, the basal lamina that underlies the corneal endothelium.4,5 FCD typically begins in the fifth decade of life and progresses slowly over two to three decades.6,7 End-stage disease is characterized by degeneration of the corneal endothelium,8–10 with the consequent failure of its ion transport and solute barrier functions.11,12

Although examples of familial FCD are well known,12–15 attempts to identify loci and genes have only recently achieved results.7,16,17 A rare, atypical form of inherited FCD that is characterized by unusual corneal histopathology and onset in early childhood18 has been mapped to the FECD locus at 1p34 and associated with point mutations in the COL8A2 gene.7,16 This gene encodes the α2 subtype of collagen VIII, a major component of Descemet’s membrane, the thick basal lamina underlying the corneal endothelium.19–21 Even though this early-onset disorder has provided important insights into the etiology of FCD,7,16,18 we still have no clear evidence that mutations of COL8A2 are involved in the common late-onset form of the disease.22

We have recently identified FCD1, a disease locus for late-onset FCD that maps to 13pter-13q21.13,23 The FCD1 kindred exhibited single-locus Mendelian inheritance with high penetrance, despite the prevailing view that multiple low-penetrance genetic traits modified by environmental factors are generally thought to underlie common human diseases.24,25 It is still unknown whether single-locus or more complex inheritance is typical of this disease. Although relatively few large families have been reported,13,14,26 familial FCD appears to be common, as 50% of clinical patients with FCD are found to have siblings, parents, or offspring with the disease.6,7 To address the genetic basis of FCD more broadly, we now report the results of genetic linkage analysis for three large families showing dominant inheritance of late-onset FCD.

METHODS

Patients and Determination of Phenotype

Participants in this study were recruited, examined, and their written consent obtained under authorization of the Institutional Review Board for Human Subjects Research at the Johns Hopkins University School of Medicine, according to principles of the Declaration of Helsinki. Diagnosis of FCD was by slit lamp biomicroscopy27 and was graded according to the scale of Krachmer et al.28

Genotyping and Linkage Analysis

Whole blood was obtained from family members and stored at ~20°C. DNA from 3.5 mL blood was extracted with a genomic DNA kit (Wizard; Epicenter, Madison, WI) and genotyped by deCode Genetics (Reykjavik, Iceland) at 1107 polymorphic tandem repeat linkage markers distributed at ~4 cM spacing over all autosomes and the X chromosome.29 Pedigree integrity was checked with Relcheck.29 Paternity for all family members was confirmed, and Mendelian consistency of all genotypes were tested by Pedcheck.30 Only 149 genotypes, representing 0.15% of total genotypes, showed Mendelian inconsistency due to
genotyping error and were removed. Two-point linkage mapping was performed with MLINK, using a model that assumed fully penetrant Mendelian inheritance with disease allele frequency 0.0001. Multipoint linkage and haplotypes were obtained using SimWalk 2.89 (http://www.simwalk@genetics.ucla.edu/ provided in the public domain by the University of California Los Angeles). Multipoint analysis was performed in a reduced-penetrance model, having penetrances of \( f_0, f_1, \) and \( f_2 = 0.15, 0.85, \) and 0.85, where \( f_i \) is the chance that an individual carrying \( i \) disease alleles will be affected. The disease allele frequency was set to 0.0001. Marker allele frequencies used in the calculations were estimated directly from the linkage population. Simulated linkage results were obtained by averaging 100 independent runs of MLINK with randomly generated alleles for genotyped individuals.

**RESULTS**

The three largest late-onset FCD pedigrees available for study were ascertained as completely as possible, in preparation for linkage analysis (Fig. 1). Selected individuals were examined fully by means of slit lamp and high-resolution confocal specular microscopy and were found to have corneal guttae with a morphology typical of classic late-onset FCD. Together, the three families had 27 affected women, and 19 affected men. Among those affected, the youngest was age 32, the oldest 97. The presence of FCD patients in multiple generations, and a high percentage of affected siblings was consistent with dominant inheritance, the model implemented with two-point parametric MLINK linkage analysis, and subsequent multipoint link-
FIGURE 2. Two-point lod score genome scans of each kindred. Lod scores of the pedigrees in Figure 1 were obtained for 1056 autosomal markers by using MLINK. The mutant was assumed to be a fully penetrant Mendelian dominant trait with an allele frequency of 0.0001. The recombination fraction $\theta$ was allowed to vary for each point, to achieve a maximum lod score. Markers are organized in chromosomal linkage groups, beginning with chromosome 1 on the left and proceeding from the short arm (p) toward the long arm (q) on each chromosome. Brackets indicate selected chromosomes. Markers corresponding to the two known FCD loci for COLA2 (FECD, 1p34) and FECD1 (1pter-q12.13) are indicated by vertical bars beneath the graph. The marker D18S1129 repeatedly gave the single highest lod score within each of the three kindreds.

age. Initial results of whole-genome scans of the three families by two-point linkage analysis, assuming fully penetrant dominant single-locus Mendelian inheritance, are shown in Figure 2. Note that these lod scores are calculated at the maximum-likelihood estimate of the recombination fraction $\theta$ between each marker and the putative disease gene, with $\theta$ construed to be the same in all families. In all three of the families, a major peak was found on chromosome 18, with the top lod score at D18S1129 in each case. In kindreds 1, 2, and 3, this peak had maximum lod values of 3.41, 2.89, and 2.45, respectively (Table 1). Other two-point linkage peaks were also observed, although none were found for polymorphic markers near the COL8A2 gene at 1p34 or within the FECD1 locus at 13pter-q12 (Fig. 2).

The combined linkage statistics for all three families was estimated by summation of the two-point lod scores for each marker calculated using a single value of $\theta$, then maximizing this combined score across different values of $\theta$ (Fig. 3). The 18q21 locus showed a cluster of very high cumulative lod scores, with a two-point lod score of 7.70 for D18S1129. Secondary peaks on other chromosomes were also observed, but the significance of these is unclear, as they differed among the families. None of the secondary peaks corresponded to the two previously characterized loci for this disease, FECD (COL8A2) at 1p34.16 or FECD1 at 13pter-q12.13 (see Fig. 2). These results were extremely robust over different estimates of the disease allele frequency, which altered the peak D18S1129 lod score from 7.70 at a frequency of 0.0001 to 7.75 at a frequency of 0.01.

Multipoint lod scores were calculated for all 35 markers on chromosome 18 in a dominant model with penetrance of 85% and a phenocopy rate of 15% (Fig. 4A), as described in the Methods section. These traces revealed maxima at D18S1129 for all three families, whereas the combined scores produced a highly significant lod score peak of 5.94 at this marker. Because multipoint analysis is relatively insensitive to misspecification of marker allele frequencies, this peak provides strong confirmation of the significance of the cluster of high two-point lod scores seen in Figure 3. The observed phenocopy rate in these families is 22%, much higher than the 4% prevalence conservatively estimated for the general population. The multipoint linkage is determined with a 5% phenocopy parameter (Fig. 5B), the results are very similar, with a peak combined lod score of 5.51, and the background level for unlinked markers is considerably lower.

Multipoint analysis also yielded inferred haplotypes for each of the three families. These are shown in Figures 5, 6, and 7 along with pedigree structure and disease phenotype. In kindred 1, the 18q21 locus disease haplotype (boxed) showed strong correlation with the disease, although three phenocopies were observed in which the disease was not accompanied by the disease-associated haplotype (Fig. 5). Over all three families there were eight obligatory phenocopies in 36 individuals with no disease (Table 2). An example of nonpenetrance is illustrated by unaffected family member 3 (Fig. 5). Kindred 2 also showed high penetrance (Fig. 6), whereas kindred 3 (Fig. 7) revealed weaker but still substantial association with the disease. Over all three families, there were 41 disease haplotypes, and only four of the individuals carrying the disease haplotype were unaffected, for an observed penetrance rate of 90% for the disease trait, assuming a dominant model of inheritance. When the disease haplotypes of each family were compared by amplimer length (Table 3), they revealed that D18S1152, D18S64, and D18S1134 had identical alleles in kindreds 1 and 2, but that kindred 3 shared disease haplotype alleles with the other two families at only D18S1134.

DISCUSSION

A single locus centered at 18q21.2-q21.32 appears to be the major determining genetic factor for FCD in the three large kindreds analyzed in this study. This was unexpected, because the only genetic prescreening done before genome scan anal-
ysis was performed to rule out known loci, the late-onset FCD1 locus at 13pter-q12.23 and the COL8A2 gene at 1p34.24 These three families were selected purely on the basis of their size and a pronounced clustering of FCD cases. This finding suggests that 18q21 is a common locus in families showing dominant inheritance of late-onset FCD. Heterogeneity of the disease haplotypes between the three families leaves open the possibility of independent mutations at this locus. We cannot yet rule out that all three kindreds share a single ancient mutation that has undergone extensive recombination with its flanking markers.

Two-point linkage of the combined families yielded surprisingly large maximum two-point lod scores at D18S1129. In this case, lod scores were calculated for isolated markers in a dominant, completely penetrant model, with the recombination fraction \( \theta \) between the marker and the disease locus allowed to vary. Two-point linkage analysis is generally robust to misspecification of the genetic model (the penetrances and disease allele frequency), though the estimated recombination fraction between the marker and the disease locus will generally be biased. For a discussion of the effects of model misspecification in linkage analysis the reader is referred to Sham 1998 (Ref. 33, pp 95–97). Thus, the fact that the estimated \( \theta \) were larger than the actual distances between the markers (Table 1) suggests that the data are inconsistent with fully penetrant inheritance of the trait. This map-related inconsistency was confirmed by multipoint analysis of the same data, which yielded no significant linkage for chromosome 18 when calculated using a fully penetrant dominant model of inheritance (not shown). However, multipoint linkage yielded highly significant linkage to D18S1129 when 85% penetrance and a 15% phenocopy rate were incorporated into the dominant inheritance model (Fig. 4A). When the phenocopy rate was set close to the observed sporadic incidence of FCD in the general population (reference level), there was only a minor decrease in the peak score from 5.94 to 5.51. Although multiple linkage analysis is not robust to misspecification of the genetic model, neither two-point nor multipoint linkage analysis give increased false-positive rates in the presence of model misspecification, and so the linkage evidence in these kindreds provides extremely strong support for the contribution of the chromosome 18 locus to Fuchs susceptibility.

<table>
<thead>
<tr>
<th>Marker Name</th>
<th>Loc. (cM)</th>
<th>Maximum Lod Scores for Each Family</th>
<th>Maximum Sum of Lod Scores</th>
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<td></td>
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<td>+2.89</td>
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<td>D18S1154</td>
<td>88.62</td>
<td>+0.05</td>
<td>+1.50</td>
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</table>

Columns contain the marker name, location of each marker on the chromosome 18 linkage group in centimorgans, two-point lod scores for each of the three kindreds and combined lod score for all three kindreds, with the value of \( \theta \) at the maximum combined lod score. That \( \theta \) is in each case greater than the spacing between the markers suggests that a fully penetrant model is not the best fit for the data.
When haplotypes and disease diagnoses were examined on an individual-by-individual basis, deviations from ideal Mendelian inheritance were apparent. Of the 41 individuals with the full three-marker disease haplotype, four were clearly not affected by FCD (Table 2), an observed 90% penetrance of the FCD2 mutation. Lack of penetrance of the mutation in individual 3 of kindred 1, who was 74 years old, is not readily explained as an age-of-onset effect, and the number of cases in this study was too small to allow us to draw any conclusions. As a common disease, we would expect FCD to show complex inheritance, and it is something of a surprise that the observed penetrance rate is as high as 90%. This suggests that environmental or other genetic factors are relatively minor modifiers of the inherited disease trait in these particular families. However,

![Figure 4](image1.png)

**Figure 4.** Multipoint linkage to chromosome 18. Multipoint linkage using all 35 markers on chromosome 18, calculated with SimWalk 2.89. The lod score was plotted against the Marshfield map position of markers in centimorgans (p to q). Lines: individual results for each family, according to the key. **Bold solid curve:** combined multipoint lod score for all three families. All calculations were based on an assumed mutant allele frequency of 0.0001, Mendelian dominant inheritance with 85% penetrance (mutant genotype with normal phenotype) with a phenocopy rate of (A) 15% or (B) 5%. The maximum combined lod score is 5.94 for a 15% phenocopy rate and 5.48 for a 5% phenocopy rate. Each peak is at D18S1129, indicated by a vertical dotted line.

![Figure 5](image2.png)

**Figure 5.** Haplotype analysis of kindred 1. Genotypes are indicated for markers around the chromosome 18 major linkage peak at D18S1129. Alleles are encoded in order of decreasing amplimer size in this family, with 0 signifying unknown or undeterminable genotype. **Boxes** enclose the haplotype of the FCD-associated chromosome, assuming a dominant, incompletely penetrant model of inheritance, with a high phenocopy rate. **Gray bar:** the disease-associated markers for all three families. Individuals representing phenocopies and those showing nonpenetration of the disease trait are indicated at top right.
FIGURE 6. Haplotype of kindred 2. Haplotypes are displayed as in Figure 5. Individual 13 showed impenetrance at only \textit{D18S64}, but was not counted as an obligatory case of impenetrance, as the disease locus boundary could be above this marker.

FIGURE 7. Haplotype of kindred 3. Haplotypes are displayed as in Figures 5 and 6. Individual 21 showed nonpenetrance at \textit{D18S64}, but was not counted as an obligatory case of impenetrance. Note that the offspring of phenocopy 6 included two more phenocopies, and the 29 to 33 sibship showed no evidence of association between 18q21 and FCD.
it remains unknown whether FCD2 alleles found in the general, unselected FCD patient population have the same degree of penetrance.

Another feature of these families is the overall high penetrance of 22%, which is approximately five times higher than that observed in the general population over 40. This high penetrance could be the result of other genetic factors segregating independently in these families, or an unknown environmental component. The penetrance rate seems to be higher for kindreds 1 and 3, and so we cannot rule out variability within this study group. Because kindreds 1, 2, and 3 were selected for genetic analysis on the basis of their striking concentration of disease cases, we may have selected for the presence of those FCD-causing mutations that happen to exhibit high penetrance. The selection for familial clustering might also have independently enriched for families with an unusually high penetrance rate. Thus, the observed penetrance and penetrance rates in these families should be viewed with great caution, as the estimates are subject to ascertainment bias.

Although both penetrance and penetrance rates may be higher than in the general population, it is important to realize that the pathology and late age of onset in these families is indistinguishable from the common clinical form of FCD. These three kindreds were recruited in a single clinic from a total set of 52 families. On this basis, we might expect that at least 5% of familial FCD involves the 18q21 locus. Further studies of many smaller families, and identification of the underlying gene mutation, will be necessary to determine how widely the 18q21 locus contributes to familial and sporadic FCD.

Finally, the disease-associated haplotype of kindred 1 (Fig. 5) defines a 7-Mb genomic interval from D18S487 to D18S1134 as the most likely site of mutations underlying FCD2. Because of the incomplete penetrance observed in these families, the boundaries of this interval cannot be defined as conclusively as in the case of a fully penetrant trait. Given this, the provisional interval shown herein includes 28 annotated genes that are now under investigation.

### Table 2

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<th></th>
<th>PC</th>
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<th>NP</th>
<th>mutH</th>
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<td>9</td>
<td>18</td>
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<tr>
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<td>35</td>
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<td>Total</td>
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<td>36</td>
<td>4</td>
<td>41</td>
<td>77</td>
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</table>

PC, phenocopies (FCD affected with normal haplotypes); nH, number of individuals with normal haplotypes; NP, nonpenetrant (unaffected with mutant haplotype); mutH, number carrying the mutant haplotype; total H, total number (nH+mutH).

### Table 3

<table>
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<th>Position (Mb)</th>
<th>Marker</th>
<th>Kin 1 Allele (bp)</th>
<th>Kin 2 Allele Length (bp)</th>
<th>Kin 3 Allele Length (bp)</th>
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<td>218</td>
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</table>

Columns contain marker position in megabases from the p-end, marker name, and amplimer lengths of the disease-associated allele for each kindred, in base pairs.

### Figure 8

Gene interval of the chromosome 18 FCD2 locus based on haplotypes of kindred 1 (Fig. 5). Gene nomenclature and nucleotide positions relative to the p telomere are based on the current map (May 2004 human genome assembly, University of California, Santa Cruz Genome Browser at http://genome.ucsc.edu).

### References