qualitative, and it is still possible that these pathways have a quantitative influence on the epidermal barrier wound response.

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Materials and Methods Figs. S1 to S3

References and Notes

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Complement Factor H Polymorphism in Age-Related Macular Degeneration

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Age-related macular degeneration (AMD) is a major cause of blindness in the elderly. We report a genome-wide screen of 96 cases and 50 controls for polymorphisms associated with AMD. Among 116,204 single-nucleotide polymorphisms genotyped, an intronic and common variant in the complement factor H gene (CFH) is strongly associated with AMD (nominal P value $<10^{-7}$). In individuals homozygous for the risk allele, the likelihood of AMD is increased by a factor of 7.4 (95% confidence interval 2.9 to 19). Resequencing revealed a polymorphism in linkage disequilibrium with the risk allele representing a tyrosine-histidine change at amino acid 402. This polymorphism is in a region of CFH that binds heparin and C-reactive protein. The CFH gene is located on chromosome 1 in a region repeatedly linked to AMD in family-based studies.

Age-related macular degeneration (AMD) is the leading cause of blindness in the developed world. Its incidence is increasing as the elderly population expands (1). AMD is characterized by progressive destruction of the retina's central region (macula), causing central field visual loss (2). A key feature of AMD is the formation of extracellular deposits called drusen concentrated in and around the macula behind the retina between the retinal pigment epithelium (RPE) and the choroid. To date, no therapy for this disease has proven to be broadly effective. Several risk factors have been linked to AMD, including age, smoking, and family history (3). Candidate-gene studies

have not found any genetic differences that can account for a large proportion of the overall prevalence (2). Family-based whole-genome linkage scans have identified chromosomal regions that show evidence of linkage to AMD (4–8), but the linkage areas have not been resolved to any causative mutations.

Like many other chronic diseases, AMD is caused by a combination of genetic and environmental risk factors. Linkage studies are not as powerful as association studies for the identification of genes contributing to the risk for common, complex diseases (9). However, linkage studies have the advantage of searching the whole genome in an unbiased manner

without presupposing the involvement of particular genes. Searching the whole genome in an association study requires typing 100,000 or more single-nucleotide polymorphisms (SNPs) (10). Because of these technical demands, only one whole-genome association study, on susceptibility to myocardial infarction, has been published to date (11).

Study design. We report a whole-genome case-control association study for genes involved in AMD. To maximize the chance of success, we chose clearly defined phenotypes for cases and controls. Case individuals exhibited at least some large drusen in a quantitative photographic assessment combined with evidence of sight-threatening AMD (geographic atrophy or neovascular AMD). Control individuals had either no or only a few small drusen. We analyzed our data using a statistically conservative approach to correct for the large number of SNPs tested, thereby guaranteeing that the probability of a false positive is no greater than our reported *P* values.

We used a subset of individuals who participated in the Age-Related Eye Disease Study (AREDS) (12). From the AREDS

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sample, we identified 96 case subjects and 50 control subjects as described (13). Because there can be many precursors to the development of either geographic atrophy or choroidal neovascularization, we purposely selected the group of study participants who had both large drusen and sight-threatening AMD as cases. All individuals identified themselves as "white, not of Hispanic origin." To the extent possible, we kept the proportions of males/females and smokers/nonsmokers the same in cases and controls. Controls were purposely chosen to be older than the cases to increase the probability that they would remain without AMD (table S1).

All 146 individuals were genotyped as described (13). A summary of genotyping quality can be found in table S2. Of the 116,204 SNPs genotyped, 105,980 both were informative and passed our quality-control checks. We then proceeded to analyze the 103,611 of these SNPs that lie on the 22 autosomal chromosomes.

Single-marker associations. For each SNP, we tested for allelic association with disease status. To account for multiple testing, we used the Bonferroni correction and considered significant only those SNPs for which $P < 0.05/103,611 = 4.8 \times 10^{-7}$. This correction is known to be conservative and thus "overcorrected" the raw P values (I4). Of the autosomal SNPs, only two, rs380390 and rs10272438, are significantly associated with disease status (Bonferroni-corrected P = 0.0043 and P = 0.0080, respectively) (Fig. 1A).

One criticism of case-control association studies such as ours is that population stratification can result in false-positive results. If the cases and controls are drawn from populations of different ancestry, with different allele frequencies, we might detect these population differences instead of loci associated with the disease. All individuals in this study self-identify their ethnicity as non-Hispanic white, and all of the case and control individuals are drawn from the same AREDS population. There was some differential recruiting of cases from office practices and recruiting of controls from radio and newspaper advertising (3). Finding two SNPs out of >100,000 implied the absence of genetic stratification, but we nonetheless used genomic control methods to control for this possibility (15). We consistently found that the significance of the tests was not inflated and that, therefore, these two SNPs are significantly associated with disease.

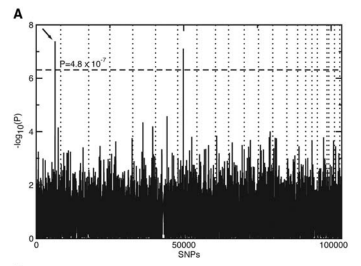
SNP rs380390 was successfully genotyped in all individuals. In 21 individuals, no genotype was determined for SNP rs10272438, and it appears to be excessively out of Hardy-Weinberg equilibrium (HWE $\chi^2 = 36$), indicating possible genotyping errors. Missing genotypes were determined by resequencing (16). After inclusion of these additional geno-

types, the association was no longer significant after Bonferroni correction. Furthermore, the SNP with the third-lowest P value, rs1329428 (Bonferroni-corrected P=0.14), is located 1.8 kb telomeric to rs380390. The genotype frequencies at these two neighboring loci vary between the case and control populations (Fig. 1B). Homozygotes for the C allele of rs380390 and the C allele at rs1329428 have an increased risk of developing AMD (Table 1). The risk conferred by these genotypes accounts for approximately 45% (rs380390) to 61% (rs1329428) of the total population risk (Table 1). We therefore focused on these two SNPs.

Risk haplotype. These two SNPs lie in an intron of the gene for complement factor H (*CFH*), located on chromosome 1q31 (GenBank accession NM_000186). Because both SNPs

are noncoding and neither appears to alter a conserved sequence, we explored whether the two SNPs are in linkage disequilibrium with a functional polymorphism. Analysis of linkage disequilibrium throughout this chromosomal region (Fig. 2A) revealed that the two SNPs lie in a 500-kb region of high linkage disequilibrium. Because this region is longer than typically observed blocks of high linkage disequilibrium (17) and there are long stretches in this region where there are no SNPs in our data set (Fig. 2B), we referred to other data sources with denser SNP coverage to narrow the region.

We used data from the International HapMap project to look at patterns of linkage disequilibrium in a population of residents of Utah with ancestry from northern and western



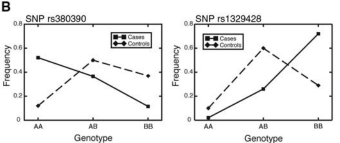


Fig. 1. (A) P values of genome-wide association scan for genes that affect the risk of developing AMD. $-\log_{10}(p)$ is plotted for each SNP in chromosomal order. The spacing between SNPs on the plot is uniform and does not reflect distances between SNPs on the chromosomes The dotted horizontal line shows the cutoff for P = 0.05 after Bonferroni correction. The vertical dotted lines show chromosomal boundaries. The arrow indicates the peak for SNP rs380390, the most significant association, which was studied further. (B) Variation in genotype frequencies between cases and controls.

Table 1. Odds ratios and population attributable risks (PARs) for AMD. The dominant odds ratio and PAR compare the likelihood of AMD in individuals with at least one copy of the risk allele versus individuals with no copy of the risk allele. The recessive odds ratio and PAR compare the likelihood of AMD in individuals with two copies of the risk allele versus individuals with no more than one copy of the risk allele. The population frequencies for the risk genotypes are taken from the CEU HapMap population (CEPH collection of Utah residents of northern and western European ancestry).

Attribute	rs380390 (C/G)	rs1329428 (C/T)
Risk allele	C ,	C
Allelic association χ^2 nominal P value	4.1×10^{-8}	1.4×10^{-6}
Odds ratio (dominant) (95% CI)	4.6 (2.0–11)	4.7 (1.0–22)
PAR (95% CI)	70% (42–84%)	80% (0–96%)
Frequency in HapMap CEU	0.70	0.82
Odds ratio (recessive) (95% CI)	7.4 (2.9–19)	6.2 (2.9–13)
PAR (95% CI)	46% (31–57%)	61% (43–73%)
Frequency in HapMap CEU	0.23	0.41

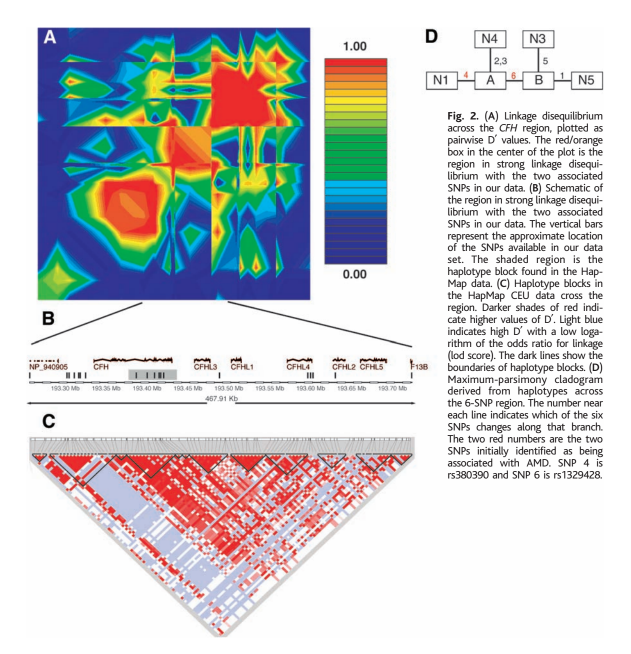
Europe [the Centre d'Étude du Polymorphisme Humain (CEPH) sample] (18). In the 500-kb region of interest, there were only 19 SNPs in our data set as compared with 152 SNPs in the HapMap data set. Using a standard definition of linkage-disequilibrium blocks (17), we found that the two associated SNPs lie in a block that is 41 kb long and entirely contained within the CFH gene (Fig. 2C).

Six SNPs from our data set were in this 41-kb region. These SNPs form four predominant haplotypes, each with a frequency greater than 1% (table S3). Combined, these four haplotypes represent 99% of the chromosomes in this study. Reconstructing inferred haplotypes and building a phylogenetic tree allowed assessment of the evolutionary relationship between haplotypes (Fig. 2D). Using inferred haplotypes for each individual, we computed the

odds ratio of the risk for disease in a nested cladistic framework under both dominant and recessive models (19). The highest risk was conferred by haplotype N1, which is the only haplotype containing the risk allele at SNP rs380390. Being heterozygous for this haplotype increases the likelihood for AMD by a factor of 4.6 [95% confidence interval (CI) 2.0 to 11] in our sample population. Being homozygous for this haplotype increases the likelihood for AMD by a factor of 7.4 (95% CI 3.0 to 19) in our sample population. Therefore, we expected to find the functionally relevant polymorphism in the context of haplotype N1. Most likely, this polymorphism would occur somewhere in the CFH gene, because the 41-kb haplotype block is entirely within CFH.

From markers to candidate functional polymorphism. To identify the polymor-

phism underlying susceptibility to AMD, we chose 96 individuals for exonic resequencing, including the exon/intron junctions. We sequenced all CFH exons, including those outside of the 41-kb block, as well as the region of SNP rs380390 as a control. SNP rs380390 was successfully resequenced in 93 individuals; the genotype derived from resequencing matched the original genotype in all cases. We identified a total of 50 polymorphisms; 17 of these have a minor-allele frequency of at least 5% (table S4). Of these 17, three represent nonsynonymous polymorphisms. We found a polymorphism in exon 9 of CFH (rs1061170) that is located 2 kb upstream of the 41-kb haplotype block, represents a tyrosine-histidine change, and is the polymorphism most strongly associated with AMD among the nonsynonymous SNPs we found. Adding this



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SNP to the haplotype analysis reveals that 97% of the chromosomes with the highest risk haplotype (N1) also have the risk allele (His).

Human complement factor H. Several lines of evidence support the hypothesis that sequence polymorphisms in CFH can lead to AMD. First, the gene for CFH is located on chromosome 1q31, a region that had been implicated in AMD by six independent linkage scans (4-8, 20). Although one study concluded that mutations in a different gene in this region (HEMICENTIN-1) were responsible for AMD (20), mutations in HEMICENTIN-1 have not been found to be generally associated with AMD in three separate, independent studies (7, 21, 22).

CFH is a key regulator of the complement system of innate immunity (23). The complement system protects against infection and attacks diseased and dysplastic cells and normally spares healthy cells. When C3 convertase is activated, it leads to the production of C3a and C3b and then to the terminal C5b-9 complex. CFH on cells and in circulation regulates complement activity by inhibiting the activation of C3 to C3a and C3b and by inactivating existing C3b.

Various components of the complement cascade, including the C5b-9 complex, have been identified in the drusen of patients with AMD (24, 25). We also examined the eyes of four patients with AMD to look for the presence of C5b-9 (fig. S1). Deposition of activated complement C5b-9 was noted in

Bruch's membrane, in the intercapillary pillars, and within drusen. The observation of GC INL ONL RPE Choroid INL ONL Choroid

Fig. 3. Immunofluorescence localization of CFH protein in human retina. Neighboring human retina sections are stained with (A) antibody to CFH or (B) antibody to CFH preabsorbed with CFH as negative control. (C) High-magnification view of the boxed area in (A). For (A), (B), and (C), left panels are the fluorescence images, with CFH labeling in green and DAPI (4',6'-diamidino-2phenylindole)-stained nuclei in blue; right panels are differential interference contrast (DIC) images showing the tissue morphology. In (C), the CFH signal is superimposed onto the DIC image. Labeling of CFH is intense in choroid, including blood vessels and areas bordering RPE [(A)] and (C)this CFH signal is competed away by purified CFH protein (B), which demonstrates the labeling specificity. The fluorescence signal from RPE arises from lipofuscin autofluorescence, which cannot be competed away with CFH protein [(A) and (B)]. The black spots in DIC images correspond to melanin granules in RPE and choroids. The cell layers are indicated: GC, ganglion cells; INL, inner nuclear layer; ONL, outer nuclear layer; RPE, retinal pigment epithelium. Scale bars: 40 μm in (A) and (B), 20 μm in (C).

complement components in drusen in both humans (24, 25) and mice (26) has led to the hypothesis that AMD results from an aberrant inflammatory process that includes inappropriate complement activation (27).

Both age and smoking, two important risk factors for AMD, influence plasma levels of complement factor H (28). CFH sequences have been observed in an expressed sequence tag library derived from human RPE and choroid (29). We confirmed by immunofluorescence experiments that CFH is present in this region of the eye (Fig. 3). Strong staining was observed in choroid vessels (retinal blood vessels) and in an area bordering the RPE. Drusen of similar composition to that found in AMD are found in the eyes of patients with membranoproliferative glomerulonephritis type II (MPGNII), a kidney disease (30); CFH deficiency can cause MPGNII (23). Our immunostaining experiments (Fig. 3 and fig. S1) suggest that in AMD, the risk variant of CFH may give rise to complement deposition in choroidal capillaries (more severe) and choroidal vessels (less severe), with subsequent leakage of plasma proteins into Bruch's membrane. Nutritional supplementation with zinc slows down the progression of AMD; biochemical studies have shown that CFH function is sensitive to zinc concentration (12, 31).

We identified a tyrosine-histidine polymorphism in which the histidine variant almost always occurs in the context of the AMD risk haplotype. This polymorphism is located in a region of CFH that binds to both heparin and C-reactive protein (CRP) (23). It has been previously suggested that this binding could be altered by the replacement of a neutral tyrosine with a positively charged histidine (23). Elevated serum levels of CRP have been shown to be associated with AMD (32). Further work to establish the causal role of the tyrosine-histidine polymorphism in AMD is warranted.

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Fig. S1 Tables S1 to S5 References

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Quantum Phase Transition of a Magnet in a Spin Bath

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The excitation spectrum of a model magnetic system, LiHoF $_4$, was studied with the use of neutron spectroscopy as the system was tuned to its quantum critical point by an applied magnetic field. The electronic mode softening expected for a quantum phase transition was forestalled by hyperfine coupling to the nuclear spins. We found that interactions with the nuclear spin bath controlled the length scale over which the excitations could be entangled. This generic result places a limit on our ability to observe intrinsic electronic quantum criticality.

The preparation and preservation of entangled quantum states is particularly relevant for the development of quantum computers, where interacting quantum bits (qubits) must produce states sufficiently long lived for meaningful manipulation. The state lifetime, typically referred to as decoherence time, is derived from coupling to the background environment. For solid-state quantum computing schemes, the qubits are typically electron spins, and they couple to two generic background environments (1). The oscillator bath—that is, delocal ized environmental modes (2) such as thermal vibrations coupled via magnetoelastic terms to the spins—can be escaped by lowering the temperature to a point where the lattice is essentially

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frozen. Coupling to local degrees of freedom, such as nuclear magnetic moments that form a spin bath, may prove more difficult to avoid, because all spin-based candidate materials for quantum computation have at least one naturally occurring isotope that carries nuclear spin.

Experimental work in this area has been largely restricted to the relaxation of single, weakly interacting magnetic moments such as those on large molecules (3); much less is known about spins as they might interact in a real quantum computer. In this regard, the insight that quantum phase transitions (QPTs) (4) are a good arena for looking at fundamental quantum properties of strongly interacting spins turns out to be valuable, as it has already been for explorations of entanglement. In particular, we show that coupling to a nuclear spin bath limits the distance over which quantum mechanical mixing affects the electron spin dynamics.

QPTs are transitions between different ground states driven not by thermal fluctuations but by quantum fluctuations controlled by a parameter such as doping, pressure, or magnetic field (5, 6). Much of the interest in QPTs stems from their importance for understanding materials with unconventional properties, such as heavy fermion systems and high-temperature

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superconductors. However, these materials are rather complex and do not easily lend themselves to a universal understanding of QPTs. To this end, it is desirable to identify quantum critical systems with a well-defined and solvable Hamiltonian and with a precisely controllable tuning parameter. One very simple model displaying a QPT is the Ising ferromagnet in a transverse magnetic field (5, 7–9) with the Hamiltonian

$$\mathcal{H} = -\sum_{ij} \mathcal{J}_{ij} \ \sigma_i^z \cdot \sigma_j^z - \Gamma \sum_i \sigma_i^x \quad (1)$$

where \mathcal{I}_{ii} is the coupling between the spins on sites i and j represented by the Pauli matrices σ^z with eigenvalues ± 1 . In the absence of a magnetic field, the system orders ferromagnetically below a critical temperature $T_{\rm c}$. The transverse-field Γ mixes the two states and leads to destruction of long-range order in a QPT at a critical field Γ_c , even at zero temperature. In the ferromagnetic state at zero field and temperature, the excitation spectrum is momentum independent and is centered at the energy $4\sum_{i}J_{ij}$ associated with single-spin reversal. Upon application of a magnetic field, however, the excitations acquire a dispersion, softening to zero at the zone center q = 0when the QPT is reached.

We investigated the excitation spectrum around the QPT in LiHoF4, which is an excellent physical realization of the transverse-field Ising model, with an added term accounting for the hyperfine coupling between electronic and nuclear moments (10-12). The dilution series LiHo_xY_{1-x}F₄ is the host for a wide variety of collective quantum effects, ranging from tunneling of single moments and domain walls to quantum annealing, entanglement, and Rabi oscillations (13–17). These intriguing properties rely largely on the ability of a transverse field, whether applied externally or generated internally by the off-diagonal part of the magnetic dipolar interaction, to mix two degenerate crystal field states of each Ho ion.