INTRODUCTION

Age-related macular degeneration (AMD) is a chronic and progressive disease marked by degeneration of the photoreceptors, retinal pigment epithelium (RPE), Bruch’s membrane, and possibly the choriocapillaries in the macula.1-3 AMD is the third leading cause of visual impairment and blindness in the United States and the developed world among people aged 65 years and older.4-5 It has been projected that by the year 2020, approximately 7.3 million people in the United States alone will have developed at minimum the early stages of AMD in at least one eye.6 However, AMD prevalence has been rising across the globe. In 2002, an estimated 37 million people in the world were blind. Among these individuals, more than 82% were aged 50 years or older. Over recent years in the developed countries, the number of people over the age of 50 years has increased by 16%. In the developing countries excluding China, this increase was by 47%. China itself had a marked increase of 27% in their elderly population.7 As the average life span of humans continues to increase, particularly in the developed countries, the incidence of AMD is expected to nearly double within the next 25 years.

Despite remarkable disease prevalence, the etiology and pathogenesis of AMD remain unclear. AMD is a common and multifactorial disease in which both genetic and environmental factors have been implicated.8-10 Complex diseases such as AMD are marked by genetic heterogeneity, a low penetrance, a continuous phenotypic distribution, and a high susceptibility to nongenetic factors.11,12 There have been several controversial reports concerning potential risk factors for AMD development.13 To date, however, only age, smoking, exposure to light, and diet have been successfully identified.13-16

The strongest evidence of a genetic component in AMD development stems from the broad tendency for familial aggregation among cases, with roughly 20% of afflicted individuals reporting a positive family history.17,18 There is also a higher incidence of AMD among monozygotic twins as compared with their spouses or other first-degree relatives.19,20 It is very likely that in common complex diseases such as AMD, variations within several genes, each with a small overall contribution and relative risk, interact to create a genetic background that can be triggered by environmental factors.

Several types of genetic polymorphisms can be found within the human genome, such as repeat polymorphisms, insertions, and deletions. However, most DNA sequence variation in human populations is in the form of single nucleotide polymorphisms (SNPs).21 SNPs can be defined as persistent substitutions of a single base with a frequency of more than 1% in at least one population. Recently, investigators have begun to explore the potential role of SNPs in AMD development. Various SNPs have been correlated, through candidate gene association studies, with age-related diseases, including AMD.22,23

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*Presenter.

Bold type indicates SOE member.
The candidate gene approach is a common method used in association analyses. This approach is based on generating hypotheses about, and selecting candidate genes involved with, plausible pathological pathways. This study investigates the Hemicentin-1 (an extracellular matrix protein identified through a genome-wide scan of extended families with AMD) Q5345R, hOgg1 (which is involved in oxidatively damaged DNA repair) S326C, and E-selectin (an adhesion molecule) S149R SNPs in association with AMD.

METHODS

STUDY SUBJECTS

This protocol was approved by the National Eye Institute Institutional Review Board. Each participant included in this study signed the informed consent prior to participation. This multiple case-control study included an AMD patient group and two normal control groups. The patients and controls included in this study were all white of non-Hispanic descent residing in the surrounding greater Washington, DC, area.

Sporadic patients (n = 89) with advanced AMD and screened normal controls (n = 97) were enrolled in this study. A clinical diagnosis of advanced AMD was defined by geographic atrophy involving the center of the macula and/or choroidal neovascularization in the presence of drusen in at least one eye. Stereoscopic fundus photographs of the optic disc and the macula were taken for all AMD patients. The screened normal controls were spouses and friends of the patients as well as randomly recruited normal volunteers. These subjects were all older than 50 years of age. These clinically screened controls received dilated fundus examinations that showed an absence of drusen or less than five small drusen (<63 µm) in the center of the macula and an absence of all other retinal disease affecting the photoreceptors, outer retinal layers, or both, such as high myopia, retinal dystrophies, central serous retinopathy, vein occlusion, diabetic retinopathy, uveitis, and other retinal diseases.

Healthy blood donors (n = 170), who did not receive a clinical eye examination, served as another control group in order to obtain information on SNP frequencies in a population that has not yet reached the average age for AMD onset. Therefore, the average age of this population is much lower than that of the AMD cases and screened controls. This group is referred to as the random controls in this study. The demographics of the study groups are summarized in Table 1.

DNA EXTRACTION

Venous whole blood (10 mL) was collected from the study subjects. Genomic DNA was extracted and isolated by using a QIAamp DNA Blood Maxi kit (catalog No. 51194; Qiagen, Valencia, California).

SNP TYPING

The SNPs Hemicentin-1 Gln5345Arg, hOgg1 Ser326Cys, and E-selectin Ser149Arg were detected by polymerase chain reaction (PCR) coupled with the restriction fragment length polymorphism (RFLP) method. For each gene, the PCR mixture included 1XJumpStart ReadyMix REDTaq (Sigma, St Louis, Missouri), 25 ng DNA, and 70 pmol of the respective primers. Following RFLP assay, the DNA fragments were separated on 15% TBE polyacrylamide gels and visualized after ethidium bromide staining. A negative (H2O) control and standards for both alleles and a heterozygous DNA control when available were included on each genotyping plate. Representative gel images used for SNP typing are shown in Figures 1 through 3.

![Hemicentin-1 Q5345R](image)

**FIGURE 1**

Autoradiography of Hemicentin-1 Q5345R in patients with age-related macular degeneration and age-matched screened volunteers. Representative gel picture of Hemicentin-1 Q5345R shows the banding patterns that determined the respective single nucleotide polymorphism types. The restriction fragment length polymorphism pattern of Hemicentin-1 Q5345R shows a 159–base pair (bp) pattern representing the wild-type (Q/Q). No variant alleles were detected in either of the case or control populations.
Nucleotide Polymorphisms in Age-Related Macular Degeneration

FIGURE 2
Autoradiography of \textit{hOgg1} S326C in patients with age-related macular degeneration and two control groups (age-matched screened volunteers and random blood bank donors). Representative gel picture of \textit{hOgg1} S326C shows the banding patterns that determined the respective single nucleotide polymorphism types. The restriction fragment length polymorphism pattern of \textit{hOgg1} S326C shows a 200–base pair (bp) pattern representing the wild-type (S/S), 200 and 100 bp representing the heterozygote (S/C), and 100 bp representing the variant homozygote (C/C).

FIGURE 3
Autoradiography of \textit{E-selectin} S149R in patients with age-related macular degeneration and two control groups (age-matched screened volunteers and random blood bank donors). Representative gel pictures of \textit{E-selectin} S149R showing the banding patterns that determined the respective single nucleotide polymorphism types. The restriction fragment length polymorphism pattern of \textit{E-selectin} S149R shows bands at 123 bp and 63 bp in wild-types (S/S); 183 bp, 123bp, and 63 bp in heterozygotes (S/R); and 183 bp in variant homozygotes (R/R).

For \textit{Hemicentin-1}, a 159–base pair (bp) DNA fragment was PCR amplified with use of the following primers: 5’-CAA GTG TAT CTG TCC ACC AGG TC-3’ and 5’-TGT CTG TAA TGC TGT TGA GGT TG-3’. The PCR program was run at 2 minutes at 94°C, followed by 34 cycles of 30 seconds denaturation at 94°C, 40 seconds annealing at 59°C, and 55 seconds extension at 72°C. RFLP analysis was conducted by using the restriction endonuclease SalI. The digestion patterns for each SNP type are as follows: wild-type (Gln/Gln), 159 bp; heterozygous variant (Gln/Arg), 159 bp and 137 bp; and homozygous variant (Arg/Arg), 137 bp.

For \textit{hOgg1}, a 200-bp DNA fragment was PCR amplified by using the following primers: 5’-ACT GTC ACT AGT CTC ACC AG-3’ and 5’-TGA ATT CGG AAG GTG CTT GGG GAA T-3’. The PCR program was run at 2 minutes at 94°C, followed by 34 cycles of 30 seconds denaturation at 94°C, 40 seconds annealing at 61°C, and 55 seconds extension at 72°C. RFLP analysis was conducted by using the restriction endonuclease Fnu4H2. The digestion patterns for each SNP type are as follows: wild-type (Ser/Ser), 200 bp; heterozygous variant (Ser/Cys), 200 bp and 100 bp; and homozygous variant (Cys/Cys), 100 bp.

For \textit{E-selectin}, a 186-bp DNA fragment was PCR amplified with use of the following primers: 5’-AGT AAT AGT CCT CCT CAT G-3’ and 5’-ACC ATC TCA AGT GAA GAA AGA G-3’. The PCR program was run at 2 minutes at 94°C, followed by 34 cycles of 30 seconds denaturation at 94°C, 40 seconds annealing at 59°C, and 55 seconds extension at 72°C. RFLP analysis was conducted by using the restriction endonuclease Pst1. The digestion patterns for each SNP type are as follows: wild-type (Ser/Ser), 123 bp and 63 bp; heterozygous variant (Ser/Arg), 183 bp, 123 bp, and 63 bp; and homozygous variant (Arg/Arg), 183 bp.
STATISTICAL ANALYSIS

The chi-square test was performed in order to compare the carrier and allele frequencies of the cases and controls. Hardy-Weinberg equilibrium was also tested by using the chi-square test within 1 degree of freedom. A \( P \) value of less than .05 was considered to be significant. Odds ratios were calculated and an estimation of confidence intervals was made on the basis of the unmatched case-control design.\(^{24}\)

RESULTS

The 89 AMD patients and 97 screened controls were matched as closely as possible for both gender and race (Table 1). Although the screened patients and controls were roughly matched for gender, the random control population was largely male. Of the 89 AMD patients, 57 had the neovascular, or “wet,” form of the disease.

DNA was successfully extracted from all enrolled patients and controls. The distribution of the Hemicentin-1 Q5345R, hOgg1 S326C, and E-selectin S149R polymorphisms did not deviate from Hardy-Weinberg equilibrium in each group included in this study (all \( P \) values < .05). The distribution of the Hemicentin-1 5345R, hOgg1 326C, and E-selectin 149R variant alleles did not differ significantly (all \( P \) values > .05) between the AMD patients and control populations (Tables 2 through 4). All gels were unable to detect a positive Hemicentin-1 5345R variant band in the screened patients and controls (Figure 1). hOgg1 326C allele frequency was 21.35% (38 of 178) in the AMD group compared with 19.12% (65 of 340) in the random controls and 19.59% (38 of 194) in the age-matched controls (Table 3). Representative autoradiography showed positive hOgg1 SNP banding patterns in several cases (Figure 2). E-selectin 149R allele frequencies were 8.99% (16 of 178) in AMD cases, 9.41% (32 of 340) in random controls, and 10.82% (21 of 194) in age-matched controls (Table 4).

A slightly lowered prevalence of the E-selectin 149R variant was found in the AMD patients as compared with the two control groups. However, this difference was not significant. Figure 3 illustrates a representative gel image of the E-selectin banding patterns. These results do not demonstrate an association between the Hemicentin-1 5345R, hOgg1 326C, and E-selectin 149R alleles and AMD in this small study.

<table>
<thead>
<tr>
<th>GROUP</th>
<th>N</th>
<th>MALE</th>
<th>FEMALE</th>
<th>MEAN AGE (YEARS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screened control</td>
<td>97</td>
<td>48</td>
<td>49</td>
<td>67.5 ± 11.7</td>
</tr>
<tr>
<td>Random control</td>
<td>170</td>
<td>113</td>
<td>5</td>
<td>45.3 ± 14.0</td>
</tr>
<tr>
<td>AMD</td>
<td>89</td>
<td>42</td>
<td>4</td>
<td>79.4 ± 7.0</td>
</tr>
</tbody>
</table>

TABLE 2. ALLELE FREQUENCIES OF HEMICENTIN-1 Q5345R IN PATIENTS WITH AGE-RELATED MACULAR DEGENERATION (AMD) AND AGE-MATCHED SCREENED VOLUNTEERS

<table>
<thead>
<tr>
<th>GROUP</th>
<th>ALLELES N</th>
<th>HEMICENTIN-1 Q5345R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screened control (n = 81)</td>
<td>162</td>
<td>0 (0)</td>
</tr>
<tr>
<td>AMD (n = 88)</td>
<td>176</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

OR = odds ratio; SNP = single nucleotide polymorphism.
TABLE 3. ALLELE FREQUENCIES OF HOGG1 S326C IN PATIENTS WITH AGE-RELATED MACULAR DEGENERATION (AMD) AND TWO CONTROL GROUPS (AGE-MATCHED SCREENED VOLUNTEERS AND RANDOM BLOOD BANK DONORS)

<table>
<thead>
<tr>
<th>GROUP</th>
<th>ALLELES</th>
<th>HOGG1 S326C</th>
<th>S</th>
<th>C (%)</th>
<th>C OR</th>
<th>χ²/P VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screened control (n = 97)</td>
<td>194</td>
<td>156</td>
<td>38</td>
<td>(19.59)</td>
<td>1.114</td>
<td>.18/.67</td>
</tr>
<tr>
<td>Random control (n = 170)</td>
<td>340</td>
<td>275</td>
<td>65</td>
<td>(19.12)</td>
<td>1.148</td>
<td>.36/.55</td>
</tr>
<tr>
<td>AMD (n = 89)</td>
<td>178</td>
<td>140</td>
<td>38</td>
<td>(21.35)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

OR = odds ratio.

TABLE 4. ALLELE FREQUENCIES OF E-SELECTIN S149R IN PATIENTS WITH AGE-RELATED MACULAR DEGENERATION (AMD) AND TWO CONTROL GROUPS (AGE-MATCHED SCREENED VOLUNTEERS AND RANDOM BLOOD BANK DONORS)

<table>
<thead>
<tr>
<th>GROUP</th>
<th>ALLELES</th>
<th>E-SELECTIN S149R</th>
<th>S</th>
<th>R (%)</th>
<th>R OR</th>
<th>χ²/P VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screened control (n = 97)</td>
<td>194</td>
<td>173</td>
<td>21</td>
<td>(10.82)</td>
<td>0.81</td>
<td>.35/.55</td>
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<tr>
<td>Random control (n = 170)</td>
<td>340</td>
<td>308</td>
<td>32</td>
<td>(9.41)</td>
<td>0.95</td>
<td>.02/.87</td>
</tr>
<tr>
<td>AMD (n = 89)</td>
<td>178</td>
<td>162</td>
<td>16</td>
<td>(8.99)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

OR = odds ratio.

DISCUSSION

Statistically significant differences in variant allele frequencies were not obtained between the AMD case and control groups after screening for the Hemicentin-1 Q5345R, hOgg1 S326C, and E-selectin S149R SNPs. Our results suggest that these polymorphisms may not be associated with AMD.

Genome-wide scans and genetic linkage analyses of extended families with AMD are two of the approaches taken to locate candidate AMD loci. ARMD1, an identified disease-related locus, was previously mapped to 1q25-31 in a large family with AMD. Recently, a gene within this region was found to be significantly associated with AMD risk. The Y402H polymorphism within the gene encoding the complement factor H protein has been reported in significant association with AMD by several independent groups. Complement factor H (CFH) is located on chromosome 1q31. It was reported that the 402H variant allele may account for up to one half of all AMD cases and that individuals homozygous for this variant allele have a 7.4-fold increased risk of developing AMD. We also found that a SNP in a CFH intron, which is strongly haplotyped with CFH 402, was significantly higher in our AMD patients compared with the controls (data not shown).

Hemicentin-1 was identified and mapped to 1q25.30-1q31.1 after further paring down of the ARMD1 locus. Several groups have screened this gene for potential AMD associated variants; however, no evidence for any significant allele associations has been generated thus far. Schultz and colleagues previously screened for the Hemicentin-1 5345R variant in sporadic AMD cases and found a very low frequency in both patients and controls. Iyengar and associates recently screened exon 104 where Hemicentin-1 is located and found no evidence of mutations in this region. Abecasis and colleagues were also unable to detect this variation in a total of 620 patients and 237 controls.

In the present study, no variant allele was detected in either the screened patients or controls. It may be possible that this gene’s significant involvement in disease development is specific to the particular multigenerational AMD-affected pedigrees previously studied. In other words, Hemicentin-1 may not be relevant to most isolated or sporadic AMD cases or even necessarily to other family pedigrees that may have more complex underlying genetic factors. Furthermore, successfully identifying an association between a particular allele and a given disease phenotype is greatly influenced by the variant allele frequency in the general population. It is highly possible also that the rarity of the Hemicentin-1 5345R allele in the general population renders the small sample populations
included in this current study incapable of detecting or determining this variation’s involvement in AMD. Lastly, there may be additional variations within or in very close proximity to this gene that may account for the risk associated with this region other than Q5345R. Larger sample sizes are needed to elucidate to what degree and by which mechanisms this gene is involved in AMD development.

Variation within genes involved in the proposed protective pathways against the effects of oxygen toxicity within the eye has been a focus of many AMD association studies. The eye is particularly vulnerable to oxidative damage because oxygen consumption is much higher in the retina than in other human tissues.35,37 Thus, failure of the normal protective enzymes and mechanisms needed to defend the macula against the oxidative stresses imposed by reactive oxygen species may result in the development of AMD and other general ailments of the aging eye. Three SNPs (MnSOD, MEHE, Paraoxonase) related to oxidative stress have previously been reported in association with AMD in Japanese populations.33,34

DNA base modifications resulting in mutations and genetic instability are a major consequence of oxidative stress.35 Efficient DNA repair mechanisms are necessary to counterbalance the damaging effects of oxidizing species. DNA repair mechanisms are also vital in maintaining cellular integrity after oxidative damage has occurred.56 The disruption of DNA repair genes has been associated with the other degenerative and earlier-onset age-related diseases, such as the Werner and Bloom syndromes, as well as with the pathogenesis of aging.37,39

The human Ogg1 (hOgg1) gene encodes a DNA glycosylase that is involved in the base excision repair of 8-hydroxy-2′-deoxyguanain (8-OH-dG) from oxidatively damaged DNA.37 Defects in 8-OH-dG repair have been implicated in the development of several diseases, including cancer.37 8-OH-dG has been found to be highly mutagenic, leading to a mutator phenotype characterized by an increase in GC to TA transversions. These transversions are frequently observed in several oncogenes and tumor suppressor genes.37

Although a statistically significant association was not established between hOgg1 and AMD in this study, this avenue of exploration has yielded promising findings. The Cockayne syndrome B (CSB) gene, also called ERCC6, collaborates with hOgg1 to carry out preferential DNA repair in eukaryotes. This gene also plays a role in the maintenance of efficient hOgg1 expression.29,40,42 Recently, we have found an association between SNPs in the promoter region of this gene and AMD (Tuo J, et al, FASEB meeting, 2005, Abstract). We have found that there is an increased prevalence of variant alleles in AMD cases as compared with the control populations. These findings suggest that an increased risk of AMD development is associated with these polymorphisms.

Investigators have recently begun looking into the potentially critical immunologic mechanisms hypothesized to be involved in the development and pathogenesis of AMD. Macrophage chemoattractant CCL2 (alternate name MCP-1, a CC chemokine) or its cognate receptor-2 (CCR2) knockout mice have been shown to spontaneously develop hallmark features of AMD in their senescent stage.43 Recently, we reported an association between AMD and two SNPs within CX3CR1, a CX3C chemokine receptor of fraktalkine/CX3CL1.44 These SNPs have been associated with a decreased number of CX3CR1/CX3CL1 binding sites as well as with decreased binding affinity.45,46 We have hypothesized that the risk introduced in AMD development is due to inefficient and/or insufficient macrophage recruitment to Bruch’s membrane.44 Efficient macrophage recruitment is needed in order to aid in the clearing of the lipid and protein retinal deposits that later accumulate and develop into soft drusen, the early hallmark of AMD. With such promising initial findings, further investigation of the genes involved in the immunologic mechanisms implicated in AMD development and pathogenesis has become an interesting avenue of research for future association studies.

Adhesion of leukocytes to endothelial cells is an essential step in their migration, rolling, strong adhesion, and diapedesis from the circulation to sites of inflammation.47 E-selectin, a cellular adhesion molecule, mediates leukocyte infiltration and lymphocyte trafficking. Polymorphisms within this gene have been associated with risk of atherosclerosis.37,48 Although we did not establish a significant association between this particular SNP in E-selectin and AMD, we observed a slightly lowered prevalence of the variant allele in our AMD cases compared with our two control populations. This difference is slightly greater when comparing the AMD cases with the screened control group alone. Although not necessarily of high importance in this specific instance, it is worth pointing out that some of the individuals in the random control group with the younger mean age may go on to develop some degree of AMD later on in life. This trend may be attributed simply to the age-related nature of the disease. Therefore, we would expect the AMD cases and the screened controls to have allele frequencies that lie on opposite ends of the spectrum, with the allele frequency of the random control group falling somewhere in between. Again, although these differences are not significant, the allele frequency patterns observed in this study align with this theory. Additional genes involved in chemoattraction, cellular adhesion, and inflammation should be explored in larger sample populations.

Only advanced AMD cases were included in this study in order to minimize the possible bias introduced by subjective and/or poorly defined phenotyping criteria. Association studies are strengthened by the incorporation of larger, more carefully age-matched patient and selected control populations. However, because AMD is an age-dependent disease of high incidence, multiple control groups with different average ages can be used to elucidate a possible quantitative correlation and potential gene-dosage effect of the tested genetic risk. We utilized a study design that aims to compensate for our small sample sizes.

In conclusion, we were not able to demonstrate an association between the Hemicentin-1 Q5345R, hOgg1 S326C, and E-selectin S149R SNPs and advanced AMD. Common multifactorial diseases such as AMD are thought to arise from a complex combination of genetic and environmental risk factors. Therefore, it is highly unlikely that a single gene variant is solely responsible for disease development. Furthermore, it is probable that the contribution of each associated gene is relatively small. Larger sample sizes are needed to adequately study gene-gene and gene-environment interactions as well as to more accurately determine relative risk. We are continuing to recruit larger patient and control populations in order to investigate the potential role of further genetic variants in AMD development.
development and pathogenesis.

ACKNOWLEDGMENTS

We would like to thank Katherine Shimel, RN, and Young Kim, RN, from the National Eye Institute, National Institutes of Health, Bethesda, Maryland, for patient and control recruitment and care.

REFERENCES


PEER DISCUSSION

DR IRENE H. MAUMENEE. Age related macular degeneration (AMD) is the third most common cause of blindness worldwide and hence worth every research effort spent on it. We are experiencing an extraordinary convergence of research results from as diverse fields as epidemiology, histopathology, and molecular genetics. All research directions are striving towards that final goal—knowledge-based treatment—which appears well within grasp for the large subset of patients with an inflammatory component in their disease process. The recently discovered variant in the gene encoding complement factor H (tyrosine-402-histidine) may account for 50% of the attributable risk, but it leaves an equally large subset of patients, who may carry mutations in one or more different genes.

Several risk factors have been linked to AMD; they include age, ethnicity, smoking, hypertension, obesity, diet and a positive family history. It will be challenging to identify the total genetic component of this disease because of its delayed onset and high frequency. The late onset prevents collection of good pedigrees, when handled over the short term. High disease frequency similarly interferes with the quality of pedigrees if many genes are causative and their differences are not recognized, when establishing a pedigree. The high disease frequency is at least in part caused by the absence of selection pressure. A selective advantage may exist in carriers of causative mutations. These different possibilities need to be explored, if one attempts to fully understand gene flow in the attempt to minimize negative research data.

An early approach to understanding AMD consisted of screening patients with severe macular degenerative diseases, such as Best disease, Sorsby pseudoinflammatory disease, Doyne honeycomb degeneration and Malattia Leventinese, and autosomal dominant Stargardt-like and recessive Stargardt disease for milder mutations in the respective genes. These analyses have been negative, except for mutations of the ABR transporter gene, which in the heterozygous state may contribute a small proportion to the genetic risk of AMD.

The genetic analyses proceeded from early candidate gene screening to the use of linkage and sib pair analyses, both followed by candidate gene analysis in the region. Significant loci were identified and the discovery of the polymorphism within Complement Factor H is based on these studies.
The authors describe an association study screening single nucleotide polymorphisms or SNPs in candidate genes in the attempt to detect additional genes and mutations significant for the development of age related macular degeneration.

The investigators have collected DNA from 89 individual patients, 57 of them had hemorrhagic macular changes, 97 age- and sex-matched controls, and 170 random controls. They looked for differences in SNP frequencies in three genes in the three groups of DNA samples. The results were negative.

The genes chosen are Hemicentin-1, hOgg1 and E-selectin. Hemicentin-1 is involved in extracellular matrix biochemistry; the function and localization of hOGG1 suggest that it may play a role in somatic hypermutation of immunoglobulin genes; E-selectin plays a prominent role in allergic disease, such as asthma, and the cell types in which mutations have been identified include cytokine-stimulated endothelial cells. It is thought to be responsible for the accumulation of blood leukocytes at sites of inflammation by mediating the adhesion of cells to the vascular lining. It exhibits structural features homologous to those of LYAM1, including the presence of lectin- and EGF-like domains followed by short consensus repeat (SCR) domains that contain 6 conserved cysteine residues. These proteins are part of the selectin family of cell adhesion molecules.

There are good theoretical reasons to assume that mutations in these genes lead to AMD, since they are part of extracellular matrix biochemistry, DNA repair, and modulation of the inflammatory response respectively.

However, whole genes were not sequenced in the patients, even though genetic diseases are only exceptionally caused by a single mutation in a given gene, but multiple discrete mutations, each leading to disease are typically observed. These different mutations may contribute to the observed phenotypic variability. The single Hemicentin-1 mutation identified in one large and several small pedigrees may be rare, but multiple, at present undetected, additional mutations may exist in this large gene and be causative for additional cases of AMD at locus 1 (where Hemicentin-1 also maps).

Linkage analysis and screening of candidate genes within the locus is less successful in common than in rare diseases, but it may be successful when used in isolated populations. Sequencing of candidate genes is the next most commonly used approach, followed by association studies using SNP analysis. They depend upon linkage disequilibrium for detection and remain time consuming and costly. But I would like to encourage the authors to continue their studies in the attempt to detect the next group of causative genes.

I have several questions to the authors. Have you collected pedigrees, and if not, will you? Have you tried to collect sib pairs? Have your patients been analyzed for the mutation seen in the Complement Factor H? Shall physicians start to screen their patients for the identified Complement Factor H mutation?

DR CHI-CHE CHAN: Age-related macular degeneration has become a serious public health issue and therefore requires serious research into potential prevention and treatment strategies. Upon admitting our case and control subjects, we do obtain information on demographics and other ophthalmic conditions. We also collect behavioral and medical histories. However, we have been unable to collect family pedigrees or sib pairs. This is something that we should consider for the future. The complement factor H (CFH) story is very interesting and perhaps very significant. Although we have not screened our patients (current N = 129) and controls (current N =130) for CFH Y402H specifically, we did screen for a SNP within a CFH intron that is strongly haplotyped with the variation at this position. The allele frequencies that we obtained were very similar to those reported by Edwards and colleagues (37.7% versus 34% in the controls and 51.2% versus 55.3% for the patients, respectively). Although the CFH story is exciting, it is a story that still must continue to unfold. Yes, ideally investigators will one day be able to screen for significant variants in patients in order to predict risk and accordingly introduce preventative strategies. Should the patients presently be screened for this variant? There are a lot of factors that play a role in the answer to that question. We need to consider the cost since screening may not be cost efficient at this point in time. Furthermore, who will screen the samples obtained in the clinics? What screening techniques will be used? You also have to be very careful with the patient identification and confidentiality.