Retinal angiomatous proliferation associated with risk alleles of ARMS2/HTRA1 gene polymorphisms in Japanese patients

Yasuhiro Ohkuma
Takaaki Hayashi
Tetsuaki Sakai
Akira Watanabe
Hisashi Yamada
Masakazu Akahori
Takeshi Itabashi
Takeshi Iwata
Toru Noda
Hiroshi Tsuneoka

1Department of Ophthalmology, 2Department of Molecular Genetics, Institute of DNA Medicine, The Jikei University School of Medicine, 3Division of Molecular and Cellular Biology, National Institute of Sensory Organs, 4Division of Ophthalmology, National Hospital Organization Tokyo Medical Center, Tokyo, Japan

Background: The purpose of this study was to investigate the association between ARMS2/ HTRA1, CFH, and C3 gene polymorphisms and retinal angiomatous proliferation (RAP), an infrequent and severe form of exudative age-related macular degeneration, which is characterized by intraretinal neovascularization.

Methods: Diagnosis of RAP was based on fundus photographs, images of fluorescein and indocyanine green angiographies, and optical coherence tomography findings. Six single nucleotide polymorphisms (SNPs), A69S (rs10490924) in ARMS2, rs11200638 in HTRA1, I62V (rs800292) in CFH, Y402H (rs1061170) in CFH, R80G (rs2230199) in C3, and rs2241394 in C3, were genotyped in eight Japanese patients with RAP.

Results: The two SNPs in the ARMS2/HTRA1 were in complete linkage disequilibrium. The frequency of the risk T allele in ARMS2 (the risk A allele in HTRA1) was 93.8% in the RAP patients. The frequency of homozygosity for the risk genotype TT of ARMS2 (the risk genotype AA of HTRA1) was 87.5%. The frequency of the non-risk allele (A) of I62V was 100%. The frequencies of risk alleles of Y402H, R80G, and rs2241394 were 12.5%, 0%, and 18.8%, respectively.

Conclusion: Our results suggest that the risk alleles of the ARMS2/HTRA1 SNPs may be associated with development of RAP and play a major role in the pathogenesis of intraretinal angiogenesis.

Keywords: age-related macular degeneration, retinal angiomatous proliferation, single nucleotide polymorphisms, ARMS2/HTRA1 genes, components of the complement system

Introduction

Age-related macular degeneration (AMD) is the most common cause of legal blindness in the elderly, affecting more than 50 million people worldwide.1 In Japan, the prevalence of AMD has risen from 0.87% in 1988 to 1.4% in 2007.2,3 Maruko et al have classified exudative AMD patients into three subtypes, namely typical wet-type AMD, polypoidal choroidal vasculopathy (PCV), and retinal angiomatous proliferation (RAP).4

AMD is a multifactorial disease with genetic, behavioral, and environmental factors.5 Recently, genetic association studies have revealed that single nucleotide polymorphisms (SNPs) in CFH (1q32), ARMS2/HTRA1 (10q26), and C3 (19p13) have been identified as major contributors to the pathogenesis of AMD.6–17 Among various SNPs of those genes, the Y402H (rs1061170) and I62V (rs800292) variants in the CFH gene and the A69S (rs10490924) variant in the ARMS2 gene have been investigated in detail.6,15,18–27 The differences in genotypes associated with AMD...
have been investigated among various ethnic groups and by subtypes of exudative AMD,\(^{16,18–27}\) showing that the I62V and A69S variants are associated with AMD in both Caucasian and Asian subjects.\(^{18–27}\) The Y402H and R80G (in the C3 gene) variants have been associated with AMD in Caucasians\(^{6–15}\) but not in Asians.\(^{18,19,21–25,28}\) The C allele of the Y402H variant and the G allele of the R80G variant are infrequent in Asians.

The term RAP was first coined by Yannuzzi et al in 2001.\(^{29}\) They suggested the retinal origin of this neovascularization, which proceeds posteriorly and finally forms a retinal-choroidal anastomosis. RAP is sometimes called type 3 neovascularization to distinguish it from type 1 neovascularization (choroidal neovascularization under the retinal pigment epithelium) and type 2 neovascularization (choroidal neovascularization that penetrates the retinal pigment epithelium).\(^{30,31}\) RAP accounts for 4.5% of all exudative AMD in Japanese patients\(^{4}\) and 15% of exudative AMD in Caucasian patients.\(^{32}\) RAP is characterized by bilateral, multiple soft drusen, intraretinal hemorrhages, and intraretinal edema. The natural history of RAP is characterized by a rapid loss of vision.\(^{33}\) RAP resists various treatments and recurs persistently.\(^{34–40}\)

The phenotypic diversity of AMD is thought to be related to differences in genetic backgrounds.\(^{10,24–27}\) Various reports have examined genetic backgrounds in PCV. Lee et al reported that the I62V and A69S variants, but not the Y402H variant, were related to PCV in Chinese patients.\(^{21}\) Hayashi et al reported that all three of these SNPs (I62V, Y402H, and A69S) were related to PCV in Japanese patients.\(^{26}\) Goto et al reported that rs2241394 in the C3 gene was associated with PCV.\(^{25}\)

Wegscheider et al reported that the Y402H polymorphism was associated with RAP in Caucasians.\(^{20}\) However, the genetic association with RAP has not been evaluated sufficiently because of the rarity of RAP in Japan. There are only a few reports about associations between A69S and RAP in the Japanese population.\(^{26–27}\) The purpose of the current study was to investigate the involvement of genetic factors in not only the ARMS2/HTRA1 but also the CFH and C3 genes in Japanese patients with RAP.

**Materials and methods**

The study was approved by the institutional review board of The Jikei Medical University, and all procedures were conducted in accordance with the principles of the Declaration of Helsinki. Eight unrelated Japanese patients with RAP were recruited from the Department of Ophthalmology at the Jikei Medical University and the National Hospital Organization Tokyo Medical Center. Informed consent was obtained from all subjects.

All patients with RAP underwent a full ophthalmic examination, including slit-lamp biomicroscopy, funduscopy, optical coherence tomography, and fluorescein and indocyanine green fundus angiographies. The diagnosis of RAP was based on the criteria of Yannuzzi et al\(^29\) and was classified as a defined anastomosis connecting the retinal circulation to a vascular complex within the retina, usually with surrounding intraretinal blood and intraretinal or cystoid macular edema.

Genomic DNA was extracted from the peripheral blood of each individual. A total of six SNPs consisting of A69S (rs10490924) in ARMS2, rs11200638 in HTRA1, Y402H (rs1061170) in CFH, I62V (rs800292) in CFH, R80G (rs2230199) in C3, and rs2241394 in C3 were genotyped. Polymerase chain reaction amplification was performed using LA Taq polymerase (Takara Bio Inc, Ohtsu, Japan) and primers for ARMS2 (forward primer: 5′-GCCTATACCCAGGACCGATG-3′, reverse primer: 5′-CATGTTCTCAGCATCTCAAGTC-3′), HTRA1 (forward primer: 5′-TCTCTGCGAATACGGACACG-3′, reverse primer: 5′-ACTGTGTCAGCTCTGTCGCG-3′, reverse primer: 5′-GCCTATACCCAGGACCGATG-3′), and rs2241394 (forward primer: 5′-ATGTAACTGTGGTCTGCGC-3′, reverse primer: 5′-GGATTAAGGGCGAACAG-3′, reverse primer: 5′-CATGTTCTCAGCATCTCAAGTC-3′).

Genomic DNA was amplified using an automated sequencer (3730xl DNA analyzer; Applied Biosystems). The two SNPs in the ARMS2/HTRA1 gene was associated with AMD in Caucasian and Asian subjects.\(^{6–15}\)

**Results**

**Genetic analysis**

Five men and three women were analyzed in the study. The mean patient age was 82.6±4.6 years (range 76–91 years). Both eyes were affected in four patients (50.0%). All SNPs were successfully genotyped in all patients (Table 1). The two SNPs in the ARMS2/HTRA1 were in complete linkage disequilibrium. The frequency of the risk T allele in

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**Representative case (patient 1)**

An 83-year-old woman with homozygosity for the risk genotype (TT) of the ARMS2 gene presented with bilateral RAP (Figure 1). She had undergone cataract surgery in both eyes prior to diagnosis of RAP. Her decimal best-corrected visual acuity was 0.3 in the right eye and 0.07 in the left eye. Both eyes were treated by standard-fluence photodynamic therapy with verteporfin (Visudyne®, Novartis Pharma AG, Basel, Switzerland) in combination with 1.25 mg (0.05 mL) of intravitreal bevacizumab (Avastin®, Genentech, San Francisco, CA, USA), and her vision improved to 0.5 in the right eye and 0.15 in the left eye, with a rapid resolution of intraretinal edema. There was no recurrence of intraretinal edema or hemorrhages, and her vision remained stable for 2 years following the combination therapy.

**Discussion**

In this study we genotyped six SNPs in RAP patients that were highly representative of the common genetic variations of exudative AMD. Our results raise the possibility of an association between ARMS2 (A69S)/HTRA1 (rs1120638) variants and RAP, but a weak association for the other SNPs. Hayashi et al recently demonstrated that the A69S, Y402H, and I62V variants are associated with RAP and that the A69S variant has the strongest association for RAP among the three exudative AMD subtypes.26 Tanaka et al also reported that A69S might serve as strong genetic markers of RAP.27 Our findings are consistent with their findings26,27 regarding the A69S variant, but we had negative results for the I62V (CFH) variant (Table 2).

![Figure 1](https://www.dovepress.com/)

**Figure 1** Color fundus photographs (A and D), indocyanine green fundus angiographies (B and E), and optical coherence tomography images (C and F) from the right eye (A–C) and the left eye (D–F) of an 83-year-old woman (patient 1).

**Notes:** We diagnosed her right eye with stage II retinal angiomatous proliferation and her left eye with stage III retinal angiomatous proliferation. (A and D) Fundus image shows intraretinal hemorrhages with a large number of soft drusen and pigment epithelial detachment. (B) Indocyanine green fundus angiographies shows some hotspots. One of them connects retinal vessels (arrow), corresponding to the intraretinal neovascularization. (C) A vertical optical coherence tomography image shows a pigment epithelial detachment, cystoid macular edema, and retinal angiomatous proliferation lesion (arrow). (E) Indocyanine green fundus angiographies shows choroidal neovascularization (arrow) that connects retinal vessels, corresponding to retinal-choroidal anastomosis. (F) A vertical optical coherence tomography image shows a pigment epithelial detachment, cystoid macular edema, and a retinal pigment epithelium line that has ruptured (arrow).

**Table 1** Polymorphisms in ARMS2/HTRA1/CFH/C3 genes: genotypes in Japanese patients with retinal angiomatous proliferation

<table>
<thead>
<tr>
<th>Patient number</th>
<th>Age</th>
<th>Sex</th>
<th>Affected eye</th>
<th>ARMS2 rs10490924 (A69S)</th>
<th>HTRA1 rs1120638</th>
<th>CFH rs800292 (I62V)</th>
<th>rs1061170 (Y402H)</th>
<th>C3 rs2230199 (R80G)</th>
<th>rs2241394</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>83</td>
<td>F</td>
<td>Bilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>TT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>2</td>
<td>79</td>
<td>M</td>
<td>Unilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>CT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>3</td>
<td>84</td>
<td>M</td>
<td>Unilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>TT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>4</td>
<td>87</td>
<td>F</td>
<td>Unilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>CT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>5</td>
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<td>M</td>
<td>Bilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>TT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>6</td>
<td>78</td>
<td>M</td>
<td>Unilateral</td>
<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>TT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
<td>7</td>
<td>76</td>
<td>F</td>
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<td>TT</td>
<td>AA</td>
<td>AA</td>
<td>TT</td>
<td>CC</td>
<td>CC</td>
</tr>
<tr>
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<td>91</td>
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<td>Bilateral</td>
<td>TG</td>
<td>AG</td>
<td>TT</td>
<td>AA</td>
<td>CC</td>
<td>CC</td>
</tr>
</tbody>
</table>

**Note:** Risk alleles are shown in bold.

**Abbreviations:** M, male; F, female.
Components of the complement system have been identified in drusen, indicating a potential role of the complement system in the pathogenesis of AMD.\textsuperscript{31,32} C3 and CFH are key components of the alternative complement pathway. C3 is the most abundant complement component and is synthesized predominantly in the liver. Cleavage of C3 into C3a and C3b is the central step in complement activation and can be initiated by the classic antibody-mediated pathway, the lectin pathway, or the alternative complement pathway. CFH is a critical negative regulator of the alternative pathway of the complement system. It binds to C3b, promotes the decay of C3 convertase, and serves as a cofactor for the factor I-mediated proteolytic inactivation of C3b, resulting in inhibition of the complement cascade.

Because there are bilateral multiple soft drusen in the presence of RAP, we suspected that RAP would be more strongly associated with genetic abnormalities in the complement system than other AMD subtypes. However, we hardly detected the risk alleles of the CFH and C3 genes. One reason for these results is the infrequency of the C allele in Y402H and the G allele in R80G in Asians. Reticular pseudodrusen (RPD) are defined as “drusen that form ill-defined networks of broad interlacing ribbons” in the Wisconsin grading system for maculopathy.\textsuperscript{33} RPD have been recognized as a distinctive morphologic feature observed in exudative AMD.\textsuperscript{41} Recent studies have demonstrated the association between RPD and reduced macular sensitivity.\textsuperscript{45,46} Importantly, it is reported that the prevalence of RPD was high in patients with RAP and the risk genotype (TT) in A69S,\textsuperscript{47} and RPD usually occurs bilaterally,\textsuperscript{25} suggesting the impact of genetic background for RPD.

It was an unexpected result that we did not detect the risk G allele in I62V (Table 2). The I62V variant has been associated with exudative AMD in both Caucasian and Asian patients.\textsuperscript{18,19,21–25,27} In Japanese population samples, it has been demonstrated that the risk genotype (GG) in I62V is significantly associated with RAP (Table 2).\textsuperscript{26,27} However, the risk genotype in I62V was not detected in our RAP patients (Table 2). Our findings suggest that the presence of the risk genotype (I62V) may not be necessarily associated with development of RAP. A larger sample size will be required to determine whether the risk genotype in I62V is eventually associated with RAP.

RAP is characterized by intraretinal neovascularization above the retinal pigment epithelium. Two different origins of this neovascularization have been proposed. Yannuzzi et al suggested that the neovascularization in RAP originates from the neural retina.\textsuperscript{29} On the other hand, Freund et al proposed type 3 neovascularization that originates not only from deep retinal capillaries but also from the choroid.\textsuperscript{31} As for the location of gene expression, the CFH gene is expressed primarily in the retinal pigment epithelium, drusen, and choroidal capillaries;\textsuperscript{4} the C3 gene is expressed in the neural retina, choroid, and retinal pigment epithelium;\textsuperscript{41} and the ARMS2 gene is expressed in the ellipsoid region of the photoreceptor cells.\textsuperscript{43} Since it seems that the location of characteristic neovascularization corresponds to the location of susceptible gene expression in RAP, our results support the hypothesis by Yannuzzi et al\textsuperscript{29} that the origin of neovascularization in RAP is in the neural retina.

In conclusion, our results suggest that the risk alleles/genotypes of the ARMS2/HTRA1 SNPs may be strongly associated with development of RAP and that they play a major role in the pathogenesis of intraretinal angiogenesis.

### Acknowledgments

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### Disclosure

The authors report no conflicts of interest in this work.

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**Table 2 Genotype frequency of A69S (ARMS2), I62V (CFH), and Y402H (CFH) polymorphisms in Japanese controls and Japanese patients with retinal angiomatous proliferation**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>rs10490924 (A69S)</th>
<th>rs800292 (I62V)</th>
<th>rs1061170 (Y402H)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>TT</td>
<td>TG</td>
<td>GA</td>
</tr>
<tr>
<td>Controls (n=1,351)\textsuperscript{26}</td>
<td>196 (14.6%)</td>
<td>638 (47.8%)</td>
<td>502 (37.6%)</td>
</tr>
<tr>
<td>Patients (n=36)\textsuperscript{26}</td>
<td>31 (86.1%)</td>
<td>3 (8.3%)</td>
<td>2 (5.6%)</td>
</tr>
<tr>
<td>Patients (n=51)\textsuperscript{27}</td>
<td>39 (76.5%)</td>
<td>10 (19.6%)</td>
<td>2 (3.9%)</td>
</tr>
<tr>
<td>Patients (n=8) (in present study)</td>
<td>7 (87.5%)</td>
<td>1 (12.5%)</td>
<td>0 (0%)</td>
</tr>
</tbody>
</table>

**Note:** Risk alleles are shown in **bold.**

**Abbreviation:** ND, not described.
References


