

A Novel Dominant Mutation in SAG, the Arrestin-1 Gene, Is a Common Cause of Retinitis Pigmentosa in Hispanic Families in the Southwestern United States

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PURPOSE. To identify the causes of autosomal dominant retinitis pigmentosa (adRP) in a cohort of families without mutations in known adRP genes and consequently to characterize a novel dominant-acting missense mutation in SAG.

METHODS. Patients underwent ophthalmologic testing and were screened for mutations using targeted-capture and whole-exome next-generation sequencing. Confirmation and additional screening were done by Sanger sequencing. Haplotypes segregating with the mutation were determined using short tandem repeat and single nucleotide variant polymorphisms. Genealogies were established by interviews of family members.

RESULTS. Eight families in a cohort of 300 adRP families, and four additional families, were found to have a novel heterozygous mutation in the SAG gene, c.440G>T; p.Cys147Phe. Patients exhibited symptoms of retinitis pigmentosa and none showed symptoms characteristic of Oguchi disease. All families are of Hispanic descent and most were ascertained in Texas or California. A single haplotype including the SAG mutation was identified in all families. The mutation dramatically alters a conserved amino acid, is extremely rare in global databases, and was not found in 4000+ exomes from Hispanic controls. Molecular modeling based on the crystal structure of bovine arrestin-1 predicts protein misfolding/instability.

CONCLUSIONS. This is the first dominant-acting mutation identified in SAG, a founder mutation possibly originating in Mexico several centuries ago. The phenotype is clearly adRP and is distinct from the previously reported phenotypes of recessive null mutations, that is, Oguchi disease and recessive RP. The mutation accounts for 3% of the 300 families in the adRP Cohort and 36% of Hispanic families in this cohort.

Keywords: retinitis pigmentosa, dominant disease, SAG, Hispanic, arrestin-1

Retinitis pigmentosa (RP) is the most common form of inherited retinal disease (IRD) and has a prevalence of approximately 1 in 4000 worldwide. It is characterized by night blindness and progressive loss of peripheral vision, often leading to complete blindness. The presence of pigmentary deposits, attenuated retinal vessels, and changes to the ERG are typical clinical findings. RP is exceptionally heterogeneous, with more than 100 genetic causes already described (RetNet: https://sph.uth.edu/retnet/, in the public domain). Currently, mutations in 23 genes are known to cause dominant RP, 53 cause recessive RP, and 5 cause X-linked disease. At least 70 syndromic or systemic diseases include RP as a component. Variability in age of onset, secondary symptoms, rate of progression, and penetrance add to the complexity.

Our autosomal dominant RP (adRP) cohort, described previously, currently consists of 300 well-characterized families with evidence of an autosomal dominant form of disease. Likely pathogenic mutations have been identified in 226 of these families (75%). Of the 300 families, 195 (65%) have dominant mutations in known adRP genes, 25 (8%) have X-linked mutations, 3 (1%) have multiple segregating muta-
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METHODS

Patient Ascertainment and Ophthalmologic Testing

This study adhered to the Declaration of Helsinki and was approved by the Committee for the Protection of Human Subjects at the University of Texas Health Science Center at Houston (UTHealth) as well as the institutional review board at each participating institution. Written informed consent was obtained from each participant before examination and genetic testing.

Patients in the adRP cohort have been ascertained over the past 25 years from collaborating clinicians at a variety of institutions. The subset of families described in this report was ascertained primarily at the Retina Foundation of the Southwest (Dallas, TX, USA); the Jule Stein Eye Institute, UCLA School of Medicine (Los Angeles, CA, USA); the Casey Eye Institute, Oregon Health and Science University, (Portland, OR, USA); and the Cizik Eye Clinic, UTHealth (Houston, TX, USA). Patients underwent detailed ophthalmologic examinations including best-corrected visual acuity, static visual fields (Humphrey), kinetic visual fields (Goldmann and Octopus), dark adaptation, dark-adapted full-field ERGs, spectral-domain optical coherence tomography (SD-OCT), and fundus photography.6

Blood and/or saliva samples were collected from affected family members and additional relatives. Genomic DNA was extracted as previously described.7,8

Sanger Sequencing

Fluorescent di-deoxy Sanger sequencing was performed as previously described on a panel of adRP genes and exons containing mutation hotspots.9-14 Additional Sanger sequencing was performed to confirm the results of NGS and to screen additional samples for the SAG mutation. Sanger sequencing was performed on a 3500xl DNA Sequencer (Applied Biosystems, Carlsbad, CA, USA).

Targeted-Capture Retinal Panels

Genomic DNA from each available family member was analyzed for the presence of mutations in retinal disease genes listed in RetNet (https://sph.uth.edu/retnet/, accessed June 1, 2016) using a targeted-capture NGS research protocol. Targeted capture was performed either in the Laboratory for Molecular Diagnosis of Inherited Eye Diseases, UTHealth, or at Baylor College of Medicine, as described previously15 (Bowne SJ, et al. IOVS 2015;56;ARVO E-Abstract 1251).

Whole-Exome NGS Sequencing

Genomic DNA (1 μg) was used to make Illumina paired-end libraries according to the manufacturer’s protocol (Illumina, Inc., San Diego, CA, USA), with modifications as described previously.16,17 Exome capture was performed using a Nimblegen SeqCap EZ Human Exome Library v2.0 (Roche, Madison, WI, USA) according to the manufacturers’ protocols. Illumina paired-end sequencing (2 × 100 bp), alignment, and variant calling were performed as described previously.16,17

Bioinformatics

MendelScan18 was used to rank single nucleotide variants (SNVs) and indels based on a combination of segregation, population frequency, retinal expression, and predicted functional impact. The Exome Aggregation Consortium (ExAC) data set (http://exac.broadinstitute.org/, in the public domain) and 1000 Genomes Database (http://www.1000genomes.org/, in the public domain) (both accessed December 12, 2016) were searched for the presence of the SAG p.Cys147Phe mutation. Variant Effect Predictor (http://www.ensembl.org/info/docs/tools/vep/index.html, in the public domain), which aggregates a number of different pathogenicity prediction algorithms including SIFT, PolyPhen-2, and Condel, predicted the pathogenicity of the SAG p.Cys147Phe mutation. Additional analyses using PROVEAN (http://provean.jcvi.org/, in the public domain) and MutationTaster (http://www.mutationtaster.org/, in the public domain) also were used. Multiple sequences were aligned using Clustal Omega (http://www.ebi.ac.uk/Tools/msa/clustalo/, in the public domain).

Haplotype Reconstruction

Haplotype analysis was done using short tandem repeat (STR) markers and SNVs identified as polymorphic from exome data and subsequent sequencing/typing in all samples. Specific markers included custom STRs and the following SNVs: rs80235307, rs7609436, rs57009436, rs4663420, rs4663420, rs181231851, rs745498, rs13018934, rs11563246, rs950834, rs4003827, and rs4003827. Genomic DNAs from all available family members were amplified, separated, and genotyped on a 3500xl DNA Sequencer (Applied Biosystems) as described previously.16 STR alleles were determined using GeneMapper V3.7 (Applied Biosystems).

Molecular Modeling

The molecular effects of the mutation were predicted based on the 2.8-Å crystal structure of bovine arrestin-1.19 Structures were obtained from the Research Collaboratory for Structural Bioinformatics Protein Data Bank (PDB: www.rcsb.org/pdb/, in the public domain). The structure (PDB ID: 1CF1) was visualized using ViewerPro 6.0 (Acceleyx, San Diego, CA, USA). The effect of mutations on side chain clashes was determined using PyMOL software (available from pmol.org, distributed by Schrödinger, Cambridge, MA, USA).

Genealogy Inferences

Unstructured interviews with informative family members were conducted to determine the earliest known ancestors with vision problems consistent with RP, the general geographic locale of ancestral and living family members, and historical information on country or countries of origin. Pedigrees of contemporary family members and known ancestors were established in genetic counseling sessions (Fig. 1).

RESULTS

Exclusion of Known Dominant Genes

Families in the adRP cohort in the Laboratory for Molecular Diagnosis of Inherited Eye Diseases, UTHealth, were ascer-
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Fig. 1. Pedigrees of families included in this study. Males are represented by squares, females by circles, multiple children of unknown sex by diamonds. Patients affected with RP are represented by filled circles or squares; numbered individuals were typed for the SAG mutation.

Sanger sequencing of known dominant RP genes was negative in the remaining cohort families, including RHO, PRPH2, PRPF31, RPGR, RP2, and mutation hotspots in HK1, IMPDH1, KLHL7, NR2E3, PRPF3, PRPF8, RP1, TOPORS, and SNRNP200. Targeted-capture NGS using a large retinal disease gene panel confirmed these negative findings and extended

tained by clinical collaborators in several institutions. Families included in the cohort have a primary diagnosis of adRP by a knowledgeable retinal specialist and three or more disease generations or male-to-male transmission. Earlier testing led to detection of the disease-causing gene in 75% of families, leaving 74 families without mutations in known genes.
Human S-arrestin (SAG)  

Human β-arrestin 1 (ARRB1)  

Human β-arrestin 2 (ARRB2)  

Human arrestin C (ARR3)  

Mouse S-arrestin  

Cow S-arrestin  

Frog S-arrestin  

Zebrafish S-arrestin  

Sea urchin β-arrestin  

Tunicate β-arrestin  

Roundworm β-arrestin

| SIFT | PolyPhen | Grantham Distance | BLOSUM62 Score | Condel Score | PROVEAN Score | Mutation Taster |
|------|----------|-------------------|---------------|-------------|---------------|----------------|
| 0    | 1        | 205               | -2            | 0.945       | -10.92        | Disease causing |

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| FIGURE 2. Sequence alignment. Clustal alignment of arrestin sequences flanking SAG Cys147 from evolutionarily diverse members of the arrestin gene family. Conserved sequence of the structural 139-loop is shown in green, conservation of Cys147 in yellow. NP_000532.2:β-arrestin (Homo sapiens); NP_004032.2:beta-arrestin 1 isoform A (H. sapiens); NP_004304.1:beta-arrestin 2 isoform 1 (H. sapiens); NP_033144.1:S-arrestin (Mus musculus); NP_851343.1:S-arrestin (Bos taurus); NP_001081898.1:arrestin-C (H. sapiens); NP_956855.1:S-arrestin (Danio rerio); NP_001160226.1:beta-arrestin-1 isoform X1 (Strongylocentrotus purpuratus); NP_001041447:arrestin (Caenorhabditis elegans). A collection of 4000+ genome sequences from Hispanic individuals in Texas was also queried and the variant was not found (Craig L. Hanis, PhD, School of Public Health, UTHealth, written personal communication, 2016).  

Identification of a Novel Dominant SAG Mutation  

Exome sequencing of the unsolved cohort families provided an extensive list of potential disease-causing variants. Comparison across families identified a novel heterozygous variant in SAG (NM_000541.4:c.440G>T; NP_000532.2: p.Cys147Phe), which was originally classified as a variant of unknown significance. Exome testing in six cohort families also did not identify any likely pathogenic changes in known adRP genes nor in genes flanking the SAG mutation in chromosome band 2q37.1.  

**Mutation Analysis**  

The SAG p.Cys147Phe mutation affects an amino acid that is conserved throughout the entire gene family of arrestins (Fig. 2). Alternative bioinformatic methods to predict protein impact yield nearly identical results: p.Cys147Phe is highly likely to be a deleterious mutation (Table 1). In databases of healthy control populations (e.g., ExAC, 1000Genomes), this variant appears once only; at a frequency of 1 per 120,594 alleles overall, and in 1 per 11,566 Hispanic alleles (ExAC database accessed December 14, 2016; http://exac.broadinstitute.org/). A collection of 4000+ genome sequences from Hispanic individuals in Texas was also queried and the variant was not found (Craig L. Hanis, PhD, School of Public Health, UTHealth, written personal communication, 2016).

**Structural Analysis**  

The three-dimensional structure of mammalian arrestin-1 was solved at high resolution (PDB ID: 1CF1 for the basal conformation of the bovine protein19 and 4ZWJ for the rhodopsin-bound mouse arrestin-120). The cysteine at human position 147 (bovine position 143) is located in the center of the molecule, near the rhodopsin-binding finger loop and the 139-loop, a highly conserved feature21 (Fig. 3). The small cysteine side chain faces inward and is in close contact with several neighboring hydrophobic residues (Fig. 3B). The replacement of this cysteine with the much bulkier phenylalanine is likely to affect protein folding: numerous side chain clashes (in particular with Leu132, Val251, Tyr255, and Ile323) are likely to occur. The cysteine side chain faces inward and is in close contact with several neighboring hydrophobic residues (Fig. 3B). The replacement of this cysteine with the much bulkier phenylalanine is likely to affect protein folding: numerous side chain clashes (in particular with Leu132, Val251, Tyr255, and Ile323) are likely to occur. The mutation likely arose. A region of 1.31 Mb, including the SAG mutation, is found in all affected individuals (Fig. 4). In several families, there is insufficient information to determine phase; however, all affected individuals have genotypes consistent with the inferred ancestral haplotype. No potential disease-causing mutations were detected in genes flanking SAG within the shared haplotype and no known genes causing retinal disease, other than SAG, are in this region. Thus, linkage disequilibrium with another potential disease-causing mutation is highly unlikely.

**Haplotype**  

Polyphyletic SNVs and STRs were typed in all available individuals to identify the underlying haplotype on which the mutation likely arose. A region of 1.31 Mb, including the SAG mutation, is found in all affected individuals (Fig. 4). In several families, there is insufficient information to determine phase; however, all affected individuals have genotypes consistent with the inferred ancestral haplotype. No potential disease-causing mutations were detected in genes flanking SAG within the shared haplotype and no known genes causing retinal disease, other than SAG, are in this region. Thus, linkage disequilibrium with another potential disease-causing mutation is highly unlikely.
of field constriction. All patients showed elevations in final dark-adapted visual thresholds following 45 minutes of dark adaptation. Unlike in patients with Oguchi disease, extended (3 hours) dark adaptation did not lead to a further gain in sensitivity. None of the patients with SAG mutations showed the Mizuo-Nakamura phenomenon characteristic of Oguchi disease associated with recessive SAG mutations.4 The fundus typically showed pigmentary changes characteristic of RP. In addition, several patients had multiple hyperreflective spots, which appeared on OCT to be distributed through all retinal layers (Fig. 5).

**Family RFS138.** A 51-year-old male presented with visual acuity loss to 20/200 OU. Rod ERGs were nondetectable and cone ERG responses were reduced greater than 95%. Visual fields measured 4 degrees in each eye. His final dark-adapted threshold with an 11-degree test centrally fixated was elevated by 3.5 log units. His daughter presented at age 28 years reporting peripheral vision loss and nystagmus. Visual acuity was 20/15 OD and 20/50 OS. Her final dark-adapted threshold was elevated 0.9 log unit with an 11-degree test centrally fixated. Rod responses were minimally detectable and cone ERG responses were reduced by 90%. The proband's son was diagnosed with adRP at age 54 years with acuity reduced to 20/40 OU. Visual fields were constricted to 10 degrees. Rod ERG responses were nondetectable. Cone ERG responses were reduced by 85%. His final dark-adapted threshold was elevated by 3.4 log units with an 11-degree test centrally fixated. Bone-spicule pigment was present in the midperiphery and cystoid macular edema (CME) was evident on OCT.

**Family RFS328.** A 42-year-old female reported first noticing peripheral vision loss and night blindness at age 40 years. Visual acuity was decreased to 20/65 OD and 20/32 OS with nondetectable rod ERG responses and cone ERG responses reduced by 97%. Visual fields were constricted to 25 degrees' diameter and the final dark-adapted threshold was elevated 2.5 log units with an 11-degree, centrally fixated test. Severe bone-spicule pigment was present in the midperiphery with vessel attenuation and optic disc pallor.

**Family RFS395.** A 59-year-old male first reported noticing progressive peripheral vision loss, nystagmus, and photophobia at age 53 years. Visual acuity was 20/40 OD and 20/25 OS with nondetectable rod ERG responses and cone responses reduced by 96%. Visual fields were constricted to less than 10 degrees and the final dark-adapted threshold was elevated by 0.8 log unit. The fundus showed peripheral pigmentation in all quadrants but minimal vessel attenuation. The patient’s 45-year-old sister reported noticing symptoms only within the past year. Visual acuity was 20/32 OD and 20/32 OS. Rod ERGs were nondetectable and cone ERGs were reduced by 78%. Visual fields were constricted to 25 degrees. Her final dark-adapted threshold was elevated 0.6 log unit with an 11-degree test.

**Family RFS516.** A 48-year-old male first noticed field loss at age 24 and is aware of progressive field constriction. The patient retained 20/32 acuity in each eye with nondetectable rod ERGs and cone ERG amplitudes reduced by 98%. Visual fields were constricted to 15 degrees and the final dark-adapted threshold was elevated 1.0 log unit with an 11-degree test centrally fixated. Moderate bone-spicule pigment clumping was observed in the midperiphery along with arteriolar attenuation. The patient’s 26-year-old son was also diagnosed with adRP and provided a DNA sample for analysis.

**Family RFS741.** A 59-year-old male first reported progressive peripheral vision loss beginning at age 51 years and severe night blindness since age 44 years. Acuity was reduced to 20/40 OU. Rod ERG responses were nondetectable with cone ERG responses reduced by 93%. Visual fields were constricted to 20 degrees with a temporal island in the periphery. The dark-
adapted threshold was elevated 3.6 log units. Moderate bone-
spicule pigment was observed in the midperiphery along with
atrophic areas in the mid-to-far periphery and optic disc pallor.

**Family UTAD0746.** A 55-year-old female first noted
decreasing night vision and restriction of peripheral vision at
age 42 years. Visual acuity was 20/16 OD and 20/20 OS. Rod
ERGs were nondetectable and cone ERGs were reduced by
62%. Visual fields were constricted to less than 20 degrees. The
final dark-adapted threshold was borderline normal with an 11-
degree test centrally fixated but elevated at least 2.0 log units
elsewhere in the retina. The fundus showed moderate
midperipheral bone spicules with pigmentary mottling in both
eyes, but vessels showed minimal attenuation.

**Family UTAD1112.** A 41-year-old female first reported field
loss at age 34 years and reported progressive peripheral vision
restriction and nyctalopia. The patient retained 20/20 vision
with correction but 20/20 OS. Rod ERGs were nondetectable and cone ERG amplitude
reduced by 96%. Visual fields were constricted to less than
20 degrees centrally, and the final dark-adapted threshold was
elevated 0.5 log unit with an 11-degree test centrally fixated.

Spare bone-spicule pigment was observed in the midperiph-
eery. The patient’s 67-year-old father was also diagnosed with
adRP, and retained 20/25 OD, 20/20 OS vision with non-
detectable rod ERG responses. Cone ERGs to 30-Hz flicker
were reduced in amplitude by 30% and significantly delayed in
b-wave implicit time. Kinetic visual fields to spot size 3 were
full, with a midperipheral scotoma in the superior field.

**Family UTAD0080.** A 45-year-old female with a family
history consistent with dominant RP was first diagnosed at age
36. Visual acuity with correction was 20/40 OU with visual
fields constricted to less than 10 degrees. On fundoscopy,
scattered pigment clumps and a few bone spicules were seen
throughout the retina. Some RPE atrophy in a lobule pattern
was observed, especially in the temporal areas.

**Family ORD4432.** This 57-year-old male has a family
history consistent with autosomal dominant inheritance. His
visual acuity was 20/50 OD and 20/50 OS. Kinetic perimetry
showed intact peripheral fields with midperipheral scotomas.
Rod ERG responses were nondetectable. Cone responses
showed moderate attenuation with delayed b-wave implicit
times. His fundus showed moderate vessel attenuation,

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**Figure 4.** Haplotype surrounding the SAG Cys147Phe mutation. Seventeen single-nucleotide polymorphisms (SNPs) and four STRs were typed in all available members of the 12 families with the SAG Cys147Phe mutation. Genotypes were examined and the disease haplotype determined in as many families as possible. All individuals carried a haplotype or corresponding genotype consistent with a founder mutation occurring several generations ago. The minimal haplotype region was determined to be 1.31 Mb and located between SNPs rs181158151 and rs950834.
### Table 2. Clinical Description of Representative Patients

| Patient ID | Diagnosis  | Age at Examination | BCVA          | VF                  | ffERG                                      | DA Threshold Elevation, Log Units | Notes               |
|------------|------------|--------------------|---------------|---------------------|--------------------------------------------|----------------------------------|---------------------|
| RFS138-3119 | adRP       | 28                 | 20/15; 20/50  | -                   | Rods reduced by 95%; cones reduced by 90% and delayed | 1.0                             |                     |
|            |            | 31                 | 20/30; 20/25  | -                   | Rods reduced by 95%; cones reduced by 90% and delayed | 0.6                             |                     |
|            |            | 40                 | 20/25; 20/60  | HVF 10°             | Rods nondetectable; cones reduced by 98% and delayed | 0.3                             |                     |
| RFS138-11896 | adRP      | 54                 | 20/40; 20/40  | Kinetic 10°         | Rods nondetectable; cones reduced by 85% and delayed | 3.2                             | Diabetic            |
| RFS138-1142 | adRP       | 51                 | 20/200; 20/200| Kinetic 4°          | Rods reduced >95%; cones reduced by >99%       | 5.2                             |                     |
| RFS3288038  | Possible adRP | 42             | 20/63; 20/32  | HVF 25°             | Rods nondetectable; cones reduced by 97% and delayed | 2.5                             |                     |
| RFS3959690  | adRP       | 59                 | 20/40; 20/25  | HVF 10°             | Rods nondetectable; cones reduced by 96% and delayed | 0.6                             |                     |
| RFS3959706  | adRP       | 45                 | 20/32; 20/32  | HVF 25°             | Rods nondetectable; cones reduced by 78% and delayed | 0.4                             |                     |
| RFS51611005 | adRP       | 48                 | 20/32; 20/32  | HVF 15°             | Rods nondetectable; cones reduced by 98% and delayed | 1.1                             | Diabetic            |
|            |            | 51                 | 20/20; 20/25  | Kinetic 20° with peripheral islands | Rods non-detectable; cones reduced by 93% and delayed | 1.3                             |                     |
| RFS74111993 | Possible adRP | 59             | 20/40; 20/40  | Kinetic 20° with peripheral islands | Rods non-detectable; cones reduced by 93% and delayed | 3.6                             | Foveal swelling OU   |
| UTAD0746-8794 | adRP     | 55                 | 20/16; 20/20  | HVF 20°             | Rods non-detectable; cones reduced by 62% and delayed | 0.0                             |                     |
| UTAD1112-10597 | adRP   | 41                 | 20/25; 20/25  | HVF 20°             | Rods non-detectable; cones reduced by 96% and delayed | 0.5                             | Diabetic            |
| UTAD1112-12157 | adRP   | 67                 | 20/25; 20/20  | Full with midperipheral scotoma | Rods non-detectable; cones reduced by 30% | 0.3                             | Diabetic            |
| UTAD0080-01 | adRP       | 42                 | Not done      | HVF 12°             | Nondetectable                             | 1.4                             |                     |
|            |            | 45                 | 20/40; 20/40  | HVF 10°             | Nondetectable                             | 2.0                             |                     |
|            |            | 65                 | 20/200; 20/80 | HVF <10°            | -                                           | -                               |                     |
| ORD 4452   | adRP       | 57                 | 20/50; 20/50  | Full with midperipheral scotoma | Rods non-detectable; cones moderately reduced and delayed |                     |                     |

BCVA, best-corrected visual acuity; DA, dark-adapted; ffERG, full-field electroretinogram; VF, visual field; HVF, Humphrey visual field.
**FIGURE 5.** Fundus montages and OCTs of representative patients with the SAG Cys147Phe mutation. (A, C) Patient RFS516-11005, a 48-year-old male with adRP. Retinal thinning, bone-spicule pigmentation and patches of atrophy are present in a midperipheral ring. A horizontal midline OCT scan (C) shows an intact ellipsoid zone in the central macula but extensive degeneration outside the arcades. The *arrows* indicate hyperreflective spots, which are seen in all retinal layers. (B, D) Patient RFS 138-11896, a 54-year-old male with adRP. Extensive atrophy is present throughout the periphery. A horizontal midline OCT scan (D) shows an intact ellipsoid zone in the central macula. Immediately peripheral to the central macula there is loss of the photoreceptor layer, including the nuclear layer, and extensive disruption of the RPE.
pigment mottling in the macula, and a ring of depigmentation around the arcades.

**DISCUSSION**

The p.Cys147Phe Mutation in SAG Is Common in Hispanic adRP Patients of Mexican Ancestry

The p.Cys147Phe mutation in SAG accounts for 36% of Hispanic families in our adRP cohort (8/22) and 3% of the overall total (8/300). Haplotype analysis is consistent with the hypothesis that the p.Cys147Phe mutation is the result of a founder mutation event and that all of the families are distantly related. Although families could not be linked by pedigree reconstruction, interviews of the oldest individuals indicated that many families had lived in what is now Texas for multiple generations, perhaps since the 1700s when the Spanish colonizers of Mexico established permanent settlements in the region via land grants.

In regions of the United States with large populations of Hispanic individuals of Mexican origin (e.g., Texas and California), it is likely to be a significant cause of disease, whereas in other parts of the country it may be less common among Hispanic individuals and rare otherwise. This is borne out by an earlier study of families with adRP from the Northeastern region of the United States,22 where the Hispanic population is significantly smaller than in Texas or California and is also primarily of Caribbean and South American origin (U.S. Census Bureau Report C2010BR-04: http://www.census.gov/library/publications/2011/dec/c2010br-04.html). In that study, neither the p.Cys147Phe mutation nor any other pathogenic mutation in SAG was found. A recent study of 35 Hispanic adRP probands, ascertained in Miami with origins in the Northeastern region of the United States,22 where the Hispanic population is significantly smaller than in Texas or California and is also primarily of Caribbean and South American origin (U.S. Census Bureau Report C2010BR-04: http://www.census.gov/library/publications/2011/dec/c2010br-04.html). In that study, neither the p.Cys147Phe mutation nor any other pathogenic mutation in SAG was found. A recent study of 35 Hispanic adRP probands, ascertained in Miami with origins in Cuba, Colombia, and Puerto Rico, also did not report the SAG mutation.23

**Dominant Mutations in a Recessive Gene**

Before our finding, mutations in the SAG gene had been identified in only two human diseases: Oguchi disease,4 a form of congenital stationary night blindness, and recessive RP.5 Both diseases are recessive and all mutations are presumed to be functional nulls, resulting in absence of a functional protein. A single example of a patient with both an amino acid substitution and an exonic duplication has been reported, but without evidence of compound heterozygosity:24 All previously reported pathogenic SAG alleles are nonsense or frameshift mutations, and heterozygous carriers are asymptomatic. Additionally, recessive canine retinal atrophy (PRA) in Basenji dogs has been attributed to a homozygous stop-loss mutation in SAG.25

Finding genes that harbor both dominant and recessive mutations is becoming more common. For IRDs, six genes have been identified (seven including SAG) with both dominant and recessive pathogenic mutations. BEST1,26 RH0,27 RPI,28 and RPE6526 cause both dominant and recessive RP; IMPG1 mutations cause dominant and recessive macular degeneration,29 and GNAT1 can cause dominant or recessive congenital stationary night blindness (CSNB).30

**Role of S-Antigen Visual Arrestin-1 in Phototransduction**

S-antigen visual arrestin-1 is the second most abundant protein in rod photoreceptors, present at approximately 0.8:1 molar ratio to rhodopsin.31,32 SAG is part of the highly conserved arrestin gene family and has homologues in species as evolutionarily distant as roundworms and tunicates33 (Fig. 2). In humans (and all other mammals) the arrestin family is composed of four members: SAG (S-arrestin/S-antigen/arrestin-1), ARR3 (cone arrestin/X arrestin), ARRB1 (arrestin beta 1), and ARRB2 (arrestin beta 2). SAG and ARR3 are the visual arrestin genes and their expression is photoreceptor-specific,34 whereas the two beta-arrestins are ubiquitously expressed.35 Crystal structures for all four vertebrate subtypes are solved and they have a common two-domain structure with an extended carboxy-terminal tail35–37 (Fig. 3). Extensive work has been done to map the functional properties of S-antigen visual arrestin-1 onto the three-dimensional structure and to delineate critical characteristics.

The primary function of S-antigen visual arrestin-1 in photoreceptors is well understood. Rods are sensitive to a single photon of light,38 and it is crucial for the signaling cascade driven by rhodopsin, a G protein-coupled receptor, to be under tight control. After being activated by light, rhodopsin amplifies the signal by activating multiple transducin molecules, which in turn activate cyclic guanosine monophosphate (cGMP) phosphodiesterase. As the concentration of cGMP decreases in the rod outer segment, cGMP-gated cation channels close. To be ready for the next incoming photon, the rod cell must reset, and this occurs with a series of inactivation steps. The role of arrestin-1 is to bind to rhodopsin that has both been activated by light and also phosphorylated at specific serine and threonine residues in its C-terminus by rhodopsin kinase (systematic name GRK1). The binding of arrestin-1 to phosphorylated rhodopsin physically blocks transducin molecules from binding39,40 and prevents subsequent steps in phototransduction from taking place, resulting in the timely return of the cell to its photosensitive state.

Analysis of the protein structure supports both pathogenicity and a likely mechanism of disease caused by the SAG Cys147Phe mutation. Replacement of the cysteine with the much larger phenylalanine is likely to perturb the tertiary structure to the point in which misfolding occurs, leading to protein aggregation and the unfolded protein response (UPR). This is analogous to the many dominant rhodopsin mutations that cause RP via protein misfolding.31,41,42 Recessive mutations in SAG also may elicit the UPR by constitutive rhodopsin signaling35 but perhaps at a slower rate, usually resulting in CSNB but sometimes causing progressive degeneration.31,43 The presence of arrestin-1 in cones, as well as rods,44 is likely to explain how cone function is also affected by this mutation.

**CONCLUSIONS**

The history of the SAG p.Cys147Phe mutation is intriguing, and its prevalence in American Hispanic individuals with adRP may be analogous to that of the rhodopsin p.Pro23His mutation in American non-Hispanic individuals.45 If the mutation occurred in Spain, before the Spanish colonization of what is now Mexico and Texas, it may be prevalent in other parts of the world as well. Until now, the p.Cys147Phe mutation in the SAG gene has been consistently overlooked because it does not conform to our expectations: monomeric variants found in the “canonical recessive genes” are routinely classified as variants of unknown significance, if they are annotated at all. Reanalysis of NGS data with fewer preconceived ideas of genotype/phenotype relationships may help in finding additional pathogenic genes and mutations for IRDs.

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