Oculocutaneous albinism (OCA) describes a group of autosomal recessive disorders featuring hypopigmentation of the hair, skin, and eyes due to an abnormality in the production of melanin. Ocular signs that reflect the abnormal development of the visual system in albinism include reduced visual acuity, nystagmus, strabismus, poor stereopsis, and foveal hypoplasia. In albinism, more fibers from the temporal retina project to the contralateral hemisphere than normal. It can be detected by recording the visual evoked potentials (VEPs). The VEP recordings of patients with albinism show a contralateral predominance in response to monocular stimulation.

Seven types of human OCA (OCA 1–7) have been defined, based on their causative genetic mutations: tyrosinase (TYR), P protein, tyrosinase-related protein 1 (TYRP1), the solute carrier family 45, membrane 2 (SLC45A2), OCA5 (mapped to the 4q24), OCA6 (SLC24A5), and OCA7 (C10orf11). A survey of Japanese OCA patients revealed that OCA1 was the most frequent type (34%), while OCA4, which is rare in other countries, is the second most frequent type (27%). The clinical phenotypes of OCA4 vary from complete depigmentation with brown hair and iris; some patients show improvement in their appearance during the first decade of life. It has been hypothesized that these phenotypic variations of OCA4 patients are dependent on mutations in the SLC45A2 gene. A dominant OCA mutant strain has been isolated in a few animal species. Thus far, however, to our knowledge autosomal dominant albinism has not yet been described in humans.

We report a family with foveal hypoplasia and generalized hypopigmentation, features consistent with OCA. A mutation screening analysis of the affected members showed a likely pathogenic variant in the SLC45A2 gene, confirming a diagnosis of OCA4. To the best of our knowledge, this is the first report on a family with autosomal dominant OCA in which the diagnosis was confirmed by a genetic analysis. We described the clinical characteristics of the dominant OCA family, including the findings from the structural and functional examinations of the fovea in the patients without remarkable nystagmus.

**Patients and Methods**

This study adhered to the tenets of the Declaration of Helsinki and was approved by the Ethics Committee of Oita University.

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**Purpose.** We report the clinical characteristics of a Japanese family with autosomal dominant oculocutaneous albinism and a SLC45A2 gene mutation.

**Methods.** A total of 16 members of a Japanese family with general hypopigmentation and foveal hypoplasia underwent detailed clinical examinations. We evaluated the severity of foveal hypoplasia using spectral-domain optical coherence tomography (SD-OCT) and graded it according to the criteria of Thomas et al. DNA was extracted from 17 family members and used for genome-wide single nucleotide polymorphism genotyping and linkage analysis. Mutational search was performed for the SLC45A2 gene responsible for oculocutaneous albinism type 4 (OCA4).

**Results.** All 16 patients exhibited hypopigmentation of their hair and/or iris. They showed foveal hypoplasia, including 3 patients with grade 1 foveal hypoplasia, 7 with grade 2, and 6 with grade 3. No patient had grade 4 foveal hypoplasia. Optical coherence tomography showed macular ganglion cell complex thinning in the temporal area, and a slight reduction of visual field sensitivity in the centrotemporal area. A maximum multipoint parametric logarithm of the odds (LOD) score of approximately 2.00 to 3.56 was obtained on chromosome 5, spanning approximately 7.2 Mb between rs13187570 and rs395967 that included the SLC45A2 gene. All affected members showed a novel heterozygous variant, c.208T>C (p.Y70H), in the SLC45A2 gene, which supported a diagnosis of OCA4.

**Conclusions.** The present study reports a very rare family with autosomal dominant OCA4 whose diagnosis was confirmed by a mutational analysis. Most family members exhibited mild general hypopigmentation and low-grade foveal hypoplasia.

Keywords: albinism, foveal hypoplasia, autosomal dominant
Faculty of Medicine. Informed consent for the study was obtained from each patient. The proband was a 14-year-old female (V:10; Fig. 1) with amblyopia, congenital nystagmus, and hypopigmentation of the hair and iris. The family history revealed that many family members had amblyopia, nystagmus, and hypopigmented hair. The great-great-grandfather (I:1) and great-grandfather (II:1) of the proband exhibited similar features while they were alive. There was no history of consanguinity in this family. We examined 20 members in three generations of her family. Three members had normal foveal morphology (IV:2, V:4, V:5) and 16 had foveal hypoplasia (7 males, 9 females; age, 6–73 years). One member (III:2) showed foveal schisis; thus, we could not confirm the presence of hypoplasia. However, the patient was strongly suspected to have foveal hypoplasia due to the presence of amblyopia, nystagmus, and esotropia.

All 16 patients underwent a full ophthalmologic examination, which included assessments of decimal best corrected visual acuity (BCVA), refractive errors, type of nystagmus, ocular alignment and stereopsis, slit-lamp biomicroscopy, a fundus examination, and spectral-domain OCT (SD-OCT) imaging. Hypopigmentation of the irides, hair, and fundus was evaluated by a pediatric ophthalmology specialist (RO). Fundus photographs were taken using a digital fundus camera (VX-10; Kowa, Hamamatsu, Japan) to evaluate macular transparency according to a grading system previously described by Summers et al.17 The grading of macular transparency was as follows: Grade 1, the choroidal vessels are easily visible in the macula; Grade 2, the choroidal vessels are visible in the macula but indistinct because of the translucent appearance of the retinal pigment epithelium; and Grade 3, the choroidal vessels are not visible in the macula because of the opaque quality of the pigment epithelium. Stereopsis was evaluated with a Titmus stereo test and a TNO stereo test. Nystagmus was determined by clinical observation and considered to be present even if it was latent or intermittent.

Spectral-domain OCT images of the subjects were acquired using a Spectralis OCT (Heidelberg Engineering, Heidelberg, Germany), which provides 40,000 A-scans per second with 7-μm optical and 3.5-μm digital axial resolution. Vertical line scans through the fovea were obtained for each patient. To minimize test errors caused by missing the fovea due to nystagmus or poor fixation, horizontal and vertical line scans through the fovea center were obtained at a 30° angle, followed by serial vertical scans with an examination field size of 30 × 10°. At each location of interest on the retina, 100 SD-OCT images were acquired and averaged to reduce speckle noise. Measurements were replicated by a skilled investigator (KY). We evaluated the severity of foveal hypoplasia from the OCT images. Foveal hypoplasia was graded according to the criteria of Thomas et al.18 The grades were defined as follows: grade 1, a shallow foveal pit, presence of outer nuclear layer (ONL) widening, and presence of outer segment (OS) lengthening; grade 2, similar to grade 1 but with the absence of the foveal pit; grade 3, similar to grade 2 but with the absence of OS lengthening; grade 4, similar to grade 3 but with the absence of ONL widening.

Pattern VEPs were recorded in 3 patients (V:6, V:8, V:10) and a normal member (V:5) under the condition of monocular stimulation. The distribution of the potentials across the hemispheres was assessed by analyzing the amplitudes of the positive CI components,6 which occurred at approximately 100 ms. Visual evoked potential asymmetry also was assessed by interocular polarity at the L-R trace, which indicates a difference in the potentials in the left and right hemispheres. Four patients with foveal hypoplasia who had stable fixation, good visual acuity, and spherical refraction within
6.3 diopters (D) underwent additional examinations including the measurement of the ganglion cell complex (GCC) by OCT and a visual field examination. Macular GCC was evaluated with a 3D OCT-2000 (Topcon, Inc., Tokyo, Japan), using the 3D macular vertical scan protocols, which consist of a 7 × 7 mm² or 6 × 6 mm² analysis area containing 512 A scans/B scan with 128 B scans per volume. Ganglion cell complex thickness represents the distance from the internal limiting membrane to the outer boundary of the inner plexiform layer. The significance map is color-coded: yellow indicates border-
line results \((P < 0.05)\), and red represents results that are outside the normal limits \((P < 0.01)\). A visual field examination was performed with a Humphrey Field Analyzer with the SITA 10-2 or 30-2 (Carl Zeiss Meditec, Inc., Jena, Germany).

Genomic DNA was extracted from leukocytes from peripheral blood samples. The Illumina HumanOmniExpress-24 v1.1 Array and the Illumina GenomeStudio v2011.1 single nucleotide polymorphism (SNP) genotyping system (Illumina, San Diego, CA, USA) were employed for genome-wide SNP typing according to the manufacturer’s protocols. All SNP markers were tested for Mendelian error before linkage analysis so they could be excluded from further investigation. Multi-point parametric logarithm of the odds (LOD) scores were calculated using GeneHunter v2.1r5 (with easyLINKAGEPlus v5.08). A total of 1748 and 68 SNPs whose pairwise \(r^2\) was 0.01 were automatically selected by easyLINKAGE for linkage analysis to the whole genome (2.0 cM spacing) and to 70 cM candidate region of chromosome 5 (1.0 cM spacing), respectively. The model used in the parametric analyses assumed a dominant model of inheritance, a disease allele frequency of 0.001 and complete penetrance. The marker genetic position was based on the Marshfield linkage map, and the sex-averaged position was applied.

To screen mutations in the coding exons (exons 1–7) of the \(SLC45A2\) gene, oligonucleotide primers on the flanking intron and untranslated region sequences were designed (available on request). Polymerase chain reaction and Sanger sequencing were conducted for the affected proband (V:10) and an unaffected family member (IV:2). To investigate whether the \(c.208T>C\) change in the \(SLC45A2\) gene cosegregates with the disease, variation-specific PCR primers were designed: 5’-AAGAGAGTTCTGCTACGGG-3’ and 5’-CCTGCTACACACAGTAGCCCC-3’. Polymerase chain reaction and direct Sanger sequencing were performed on 14 family members (Fig. 1). Sequences of the membrane-associated transporter protein (MATP), which is coded by the \(SLC45A2\) gene in humans and other vertebrates, were obtained from the University of California, Santa Cruz (UCSC; Santa Cruz, CA, USA) Genome Browser (available in the public domain at http://genome.ucsc.edu/) and aligned against one another using the Clustal W software program, which is provided by the European Bioinformatics Institute (available in the public domain at http://www.ebi.ac.uk/). The pathogenicity of the amino acid changes was evaluated by four computational programs: Polyphen-2 (available in the public domain at http://genetics.bwh.harvard.edu/pph2/), SIFT (available in the public domain at http://sift.jcvi.org), PROVEAN (available in the public domain at http://provean.jcvi.org), and Mutation Taster (available in the public domain at http://www.mutationtaster.org).

**RESULTS**

The clinical characteristics of OCA in these individuals are described in the Table. The spherical equivalent refractive error varied \((-10.5 to +4.88\) D). The binocular decimal BCVA ranged from 0.3 to 1.5 (median, 0.8). Seven patients had a decimal BCVA value of better than 1.0. Four patients had manifest esotropia; 6 had exophoria or intermittent esotropia. Of the 15 patients in whom stereo acuity was examined, 8 had poor stereopsis (240–1980 arc seconds) and 7 had no stereopsis. Four patients had no nystagmus, whereas 12 had horizontal nystagmus (5 of whom had nystagmus on lateral gaze but not in the primary position). Full ophthalmologic examinations and the OCT findings revealed that all 16 patients...
had typical foveal hypoplasia: 3 had grade 1 (BCVA range, 1.0–1.5), 7 had grade 2 (BCVA range, 0.4–1.5), and 6 had grade 3 (BCVA range, 0.3–0.7) foveal hypoplasia. No patient had grade 4 foveal hypoplasia (Table, Fig. 2). Patients with more severe foveal hypoplasia had poorer visual acuity. In addition, patients with grades 1 and 2 foveal hypoplasia had no or subtle nystagmus, while patients with grade 3 foveal hypoplasia had severe nystagmus.

Pattern VEPs from three patients (V:6, V:8, and V:10) showed a well-defined CI component over the right hemisphere for the left eye, and over the left hemisphere for the right eye. The interhemispheric potential differences for the left and right eyes were opposite in polarity, reflecting the hemispheric asymmetry typically observed in patients with albinism. In a normal member (V:5), the pattern VEP shows no significant interhemispheric asymmetries (Fig. 3).

A macular analysis was performed in 4 patients with grade 1 or 2 foveal hypoplasia using SD-OCT. A GCC thickness significance map revealed GCC thinning at the temporal region of the fovea (Fig. 4). This thinning area was noticeably larger in patients with grade 2 than in those with grade 1 foveal hypoplasia. The 10-2/30-2 Humphrey visual fields showed a very slight reduction of sensitivity in the nasal visual field (Fig. 4).

Iris hypopigmentation was evaluated by a slit-lamp examination. All 16 patients with foveal hypoplasia showed mild hypopigmentation of the irides or hair. Among the 16 patients, 13 had brown hair, which darkened with age, and 3 had black

**FIGURE 4.** The GCC thickness significance map and the total deviation of the Humphrey visual fields. The GCC thickness significance map obtained from the 3D-OCT 2000 system and the total deviation of the Humphrey visual fields in a patient with grade 2 foveal hypoplasia and autosomal dominant OCA4. Three-dimensional OCT-2000 imaging with a GCC thickness significance map revealed GCC thinning in the temporal region of the fovea of both eyes. The total deviation of the Humphrey visual fields showed a slight reduction of sensitivity in the nasal visual field in a patient with grade 2 foveal hypoplasia.

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**FIGURE 5.** The variable clinical phenotypes of the family members with autosomal dominant OCA4. Patient V:6, with grade 2 foveal hypoplasia, has black hair, which is similar to the unaffected member (V:5). The others (V:8, V:10, and V:11) have light brown hair that darkened with age.
hair at presentation (Fig. 5). Three patients had discernible iris hypopigmentation compared to normal Japanese individuals. Ten patients exhibited mild peripheral iris hypopigmentation and had no abnormalities under a slit-lamp examination (Fig. 6). Iris transillumination was not performed, as it is generally undetectable in Japanese patients. Most patients showed skin pigmentation similar to that in normal individuals. Their past histories revealed that their skin was slightly creamy in complexion compared to normal Japanese individuals during the first decade of life. The patients were classified according to their macular transparency grade, as follows: grade 1, n = 4 (V:8, IV:6, III:3, V:11); grade 2, n = 3 (V:10, IV:3, IV:5); and grade 3, n = 9 (IV:4, V:2, IV:1, IV:7, V:1, V:6, III:1, V:9, V:7; Fig. 6).

Parametric linkage analysis with genome-wide SNP markers at a 2 cM interval showed a single candidate region at chromosome 5 (Fig. 7A, upper panel). Further linkage analysis using SNP markers at a 1 cM interval revealed the maximum multipoint parametric LOD score of approximately 3.56 on chromosome 5p13.3-p13.1 with an interval delineated by markers rs13187570 and rs395967 (Fig. 7A, lower panel). The region encompassed 7.2 Mb of length that included the SLC45A2 gene (See Supplementary Fig.). A novel nonsynonymous sequence change in the coding sequence of the SLC45A2 gene was found in the proband (V:10):c.208T>C, which predicted a p.Y70H change (Fig. 7B). Codon 70 is located in the second transmembrane domain in the highly conserved MFS/sugar transport protein region spanning codons 35 to 524 (db_xref, CDD: 257676; Fig. 7C). This heterozygous variant was found in all of the affected patients, indicating that the variant cosegregated with the disease. The p.Y70H variant was not found in the Human Gene Mutation Database (available in the public domain at http://www.hgmd.cf.ac.uk), the dbSNP database (available in the public domain at http://www.ncbi.nlm.nih.gov/projects/SNP/), or in the NHLBI Exome Sequencing Project (available in the public domain at http://evs.gs.washington.edu/EVS/). We used the Human Genome Variation Database (available in the public domain at http://www.hgvd.genome.med.kyoto-u.ac.jp/), which contains the genetic variations determined by exome sequencing of 1208 Japanese individuals, instead of performing sequencing in human controls; however, the variant was not found. Computational analyses indicated that p.Y70H was “possibly damaging,” with a score of 0.950 by PolyPhen-2; “damaging,” with a score of 0.013 by SIFT; “deleterious,” with a score of –2.93 by PROVEAN; and “disease-causing,” with a score of 83 by MutationTaster. The c.208T>C change is considered to be a “likely pathogenic variant with moderate evidence of pathogenicity” according to the latest American College of Medical Genetics (ACMG) standards and guidelines for the interpretation of sequence variants (see Supplementary Table for the pathogenicity based on the ACMG guidelines).22

**DISCUSSION**

Oculocutaneous albinism generally is a group of autosomal recessive disorders which feature hypopigmentation of the hair, skin and eyes.13 The clinical characteristics of OCA4 are established by the presence of hypopigmentation of the skin and hair, which ranges from complete depigmentation to partial depigmentation with brown hair, and the characteristic ocular changes.14 We reported a Japanese family in which most members showed foveal hypoplasia and generalized hypopigmentation consistent with OCA4. Mutation screening of the affected members showed a heterozygous missense mutation c.208T>C in the SLC45A2 gene. To our knowledge, this is the first report to describe autosomal dominant OCA4 in a family in which the diagnosis was confirmed by a mutation analysis. The phenotype was highly variable among subjects with the variant. In general, variable expression is seen more frequently in dominant, rather than recessive, conditions.23 According to the linkage analysis, a cosegregated gene is less likely to be expected at other regions.

The SLC45A2 gene is a human orthologue of the mouse underwhite gene (uw). Several mutations in the uw gene are known to cause melanosome anomalies of varying severity in retinal and coat color.16 The uw series contains a semidominant allele, Uw<sup>dir</sup>, which is caused by a p.D153N change. When heterozygous, this change reduces melanin production; when homozygous it results in the loss of nearly all pigmentation.10 A dominant albino locus also is known in rainbow trout.15 Contrary, known human mutations in the SLC45A2 gene are in general believed to be recessive; however, Inagaki et al.13 suggested that a heterozygous mutation p.D157N can have a dominant-negative effect. The dominant-negative effects occur when a mutated gene product interferes with the function of the normal product (the residual function will be <50%). Dominant-negative mutations often cause recessive phenotypes as exemplified by the mutations in the rhodopsin gene that cause retinitis pigmentosa.24 A potential diagnosis of the autosomal dominant form of OCA might be overlooked.

Membrane-associated transporter protein, the gene product of SLC45A2, is required for normal melanin synthesis, but the molecular mechanism of action remains to be elucidated.10,21,22 Membrane-associated transporter protein has the highest degree of homology with sucrose/proton symporter proteins, and these transporters in plants and animals have been investigated as the models for MATP.10,25,26 Notably, mutagenesis studies of plants symporter proteins at a highly conserved residue, located at the end of the first extracellular...
loop close to the start of the second transmembrane domain (e.g., H65 of Arabidopsis protein, AtSUC1), have been shown to cause a wide range of abnormal transport activities. The diverse activities are attributed to the mutations correlated with the different chemical nature of the residues at this site. These studies imply a functional relevance of the human p.Y70H change since the corresponding codon can be aligned in the immediate vicinity of the H65 codon in AtSUC1 (data not shown). Further functional studies should be performed to determine the extent to which the variant described impairs the function of the protein.

This genetic study has several limitations. Basic research using p.Y70H change is not performed. We did not test an involvement of other 49 genes in the candidate region of the positive LOD score (see Supplementary Figure). However, they were not known to be OCA genes, or to be expressed specifically in the skin and oculcar tissues, or function as transport activity according to the UCSC genome browser, available in the public domain at http://genome.ucsc.edu/.

Therefore, the SLC45A2 gene is highly suggestive for the involvement. We assumed that the high rate of inheritance of the variant occurred by chance, however, unusual families like this are more likely to be studied.

Because the present family with OCA4 exhibited a mild phenotype, it was difficult to distinguish some of the affected members from normally pigmented Japanese individuals. In these subclinical cases, an OCT examination, which sensitively and noninvasively identified foveal hypoplasia, could be helpful for the clinical diagnosis of OCA. The grading of foveal hypoplasia using OCT images was strongly correlated with visual acuity. Our patients with low-grade foveal hypoplasia (grade 1 or 2) showed good visual acuity (better than 0.8). Moreover, 9 of 10 patients with low-grade hypoplasia had no nystagmus at the primary position. This finding may represent that the absence of nystagmus is a concomitant feature in patients with albinism with low-grade foveal hypoplasia.

Although the mechanisms of reduced acuity in patients with albinism are multifactorial, foveal hypoplasia and nystagmus are the main prognostic indicators for visual acuity, and the severity of both features were well correlated in the present study.

Several members of the family showed strabismus, reduced stereoscopic vision, and interhemispheric asymmetry on VEP recordings, which suggested the misrouting of the optic nerves. Many other morphologic anomalies have been reported in the visual pathway of patients with albinism, such as reduced central retinal ganglion cell (RGC) numbers, optic nerves and chiasms of smaller diameter, and decreased lateral geniculate nuclei numbers and gray matter volume at the occipital cortex. These anomalies are likely to be the direct result of decreased ganglion cell numbers in the central retina in albino patients. In the present study, the OCT findings showed a significant reduction in the GCC thickness in the...
temporal parafovea in patients with foveal hypoplasia. In general, the development of the foveal pit occurs from 24 weeks of gestation and reaches adult conformation by 15 months postpartum. Changes that occur later in gestation, including the centrifugal migration of the inner retinal layers, result in the progressive reduction of the ganglion and bipolar cell layers at the fovea. Therefore, our GCC significance map data might indicate the incomplete migration of RGCs or the degree of foveal hypoplasia.

Another interesting result in our study is that static perimetry showed a slight reduction of sensitivity in the cenronasal visual field. A few studies have assessed visual fields of various forms in human albinism and various findings have been reported, including contracted fields, reduced sensitivity of the centronasal visual field, and normal patterns. In a previous study which compared the contrast sensitivity of the abnormally projecting part of the temporal retina was not selectively reduced, which indicates the mechanisms of cortical self-organization result in the temporal retina to the cortex. Further studies are needed to confirm whether the reduced sensitivity of the centronasal visual field is a distinctive finding in human patients with albinism.

The clinical characteristics of a family with foveal hypoplasia and generalized hypopigmentation who were diagnosed with autosomal dominant OCA4 were presented. Most family members exhibited mild general hypopigmentation and low-grade foveal hypoplasia. All of the affected members showed a heterozygous variant, c.208T>C (p.Y70H), in the SLC45A2 gene, which supported the diagnosis.

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References


