Retrospective analysis in oculocutaneous albinism patients for the 2.7 kb deletion in the OCA2 gene revealed a co-segregation of the controversial variant, p.R305W

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Abstract
Background: Oculocutaneous albinism (OCA) is an autosomal recessive disorder. A significant portion of OCA patients has been found with a single pathogenic variant either in the TYR or the OCA2 gene. Diagnostic sequencing of the TYR and OCA2 genes is routinely used for molecular diagnosis of OCA subtypes. To study the possibility that genomic abnormalities with single or multiple exon involvement may account for a portion of the potential missing pathogenic variants (the second), we retrospectively analyzed the TYR gene by long range PCR and analyzed the target 2.7 kb deletion in the OCA2 gene spanning exon 7 in OCA patients with a single pathogenic variant in the target genes.

Results: In the 108 patients analyzed, we found that one patient was heterozygous for the 2.7 kb OCA2 gene deletion and this patient was positive with one pathogenic variant and one possibly pathogenic variant [c.1103C>T (p.Ala368Val) + c.913C>T (p.R305W)]. Further analysis of maternal DNA, and two additional OCA DNA homozygous for the 2.7 kb deletion, revealed that the phenotypically normal mother is heterozygous of the 2.7 kb deletion and homozygous of the p.R305W. The two previously reported patients with homozygous of the 2.7 kb deletion are also homozygous of p.R305W.

Conclusions: Among the reported pathogenic variants, the pathogenicity of the p.R305W has been discussed intensively in literature. Our results indicate that p.R305W is unlikely a pathogenic variant. The possibility of linkage disequilibrium between p.R305W with the 2.7 kb deletion in OCA2 gene is also suggested.

Keywords: Albinism, TYR gene, OCA2 gene, p.R305W, Deletion, Linkage disequilibrium

Background
Albinism is a disease caused by a lack of or reduction in the production of melanin pigment. There are many genes involved in the complex production of melanin, known as melanogenesis. Albinism affects about 1 in 17,000 individuals worldwide, with varied subtypes and prevalence among different ethnic populations [1]. Oculocutaneous albinism (OCA) is associated with a hypopigmented state of melanin in skin, hair, and eyes. In addition, OCA may manifest with eye symptoms such as photophobia, strabismus, nystagmus, and reduced visual acuity. Nonsyndromic OCA describes the disorders in which the manifestations are apparently limited to skin and eye functions. Nonsyndromic OCA is an autosomal recessive condition with genetic heterogeneity. Six

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genes and one additional locus have been identified as responsible for seven nonsyndromic OCA subtypes [2]. They are OCA1 (TYR gene), OCA2 (OCA2 gene), OCA3 (TYRP1 gene), OCA4 (SLC45A2 gene), OCA6 (SLC24A5 gene), and OCA7 (C10orf11 gene). OCA5 was identified by linkage analysis to a genomic locus at 4q24 [3]. OCA1 is the most common subtype found in Caucasians and accounts for approximately 50% of cases worldwide [4, 5]. OCA-2 accounts for 30% of cases worldwide and is most prevalent among African and African-American populations. OCA2 is estimated to affect one in 10,000 individuals of African background, while it affects 1 in 36,000 individuals of Caucasian background [6].

The OCA2 gene (also known as Pink-Eyed Dilution, P gene) codes for a membrane protein localized at melanosomes in the melanocytes. Its function may involve transport of tyrosine and some other small molecules [7]. The OCA2 protein may also play a role in stabilizing some melanosomal proteins such as tyrosinase and initiating melanin synthesis in melanocytes. Mutations in the OCA2 gene (cause the OCA2 phenotype (MIM 203200). The OCA2 gene consists of 24 exons (23 coding), spanning almost 345 kb of genomic DNA in the region of 15q11.2-q12. The OCA2 genomic location is also highly polymorphic.

In this study, we evaluated 108 clinically diagnosed OCA patients by sequencing their TYR and OCA2 genes. Then, we retrospectively analyzed the previously reported 2.7 kb OCA2 gene deletion using long-range PCR. We also confirmed the co-segregation of the deletion and the known controversial variant, p.R305W. We suggested that the p.R305W was the more ancestral allele and the 2.7 kb deletion was introduced into this allele more recently. This association will have a clinical impact on the pathogenicity assessment of the p.R305W variant and genetic consultation.

### Results

In our past clinical genetic tests, OCA patients were sequenced for pathogenic variants in the TYR gene at the Molecular Diagnostics Laboratory in the University of Minnesota Medical Center, Fairview (Table 1). In the 68 OCA patients tested for the TYR gene mutations, single pathogenic variant was found in 23 patients and 2 pathogenic variants were found in 45 patients. Long range PCR was performed on these patients with a series of overlapped primer sets and with OCA patients with two pathogenic variants including one equivocal mutation of the single mutation samples as PCR controls (Additional file 1: Table S1). No abnormal PCR fragment was identified (data not shown). OCA2 gene sequencing test was developed later and used as a reflex clinical test for the patients without two pathogenic variants in the TYR

### Table 1 TYR gene genotypes of OCA patients tested at University of Minnesota

| Patient ID | Allele 1 | Allele 2 |
|------------|----------|----------|
| MN1        | c.1118C>A, p.T373K |          |
| MN2        | c.1118C>A, p.T373K |          |
| MN3        | c.1118C>A, p.T373K |          |
| MN4        | c.665T>C, p.I222T |          |
| MN5        | c.1118C>A, p.T373K |          |
| MN6        | c.446A>G, p.Y1149C |          |
| MN7        | c.242C>T, p.P81L  |          |
| MN8        | c.140G>A, p.G47D   |          |
| MN9        | c.346C>T, p.R116*  |          |
| MN10       | c.832C>T, p.R278*  |          |
| MN11       | c.650G>A, p.R217Q  |          |
| MN12       | c.286dupA, p.M96Nfs*73 |          |
| MN13       | c.71G>A, p.C24Y    |          |
| MN14       | c.1037-7T>A, p.I52-7 |          |
| MN15       | c.820-2A>G, IVS1-2 |          |
| MN16       | c.572delG, p.G191Dfs*35 |          |
| MN17       | c.1467dupT, p.A490Cfs*20 |          |
| MN18       | c.242C>T, p.P81L   |          |
| MN19       | c.1118C>A, p.T373K |          |
| MN20       | c.823G>T, p.V275F  |          |
| MN21       | c.1366-4A>G, IVS4+4 |          |
| MN22       | c.1366-4A>G, IVS4+4 |          |
| MN23       | c.242C>T, p.P81L   | c.649C>T, p.R217W |
| MN24       | c.929dupC, p.R311Kfs*7 | c.929dupC, p.R311Kfs*7 |
| MN25       | c.140G>A, p.G47D   | c.140G>A, p.G47D |
| MN26*      | c.820-7T>A, IVS1-7 | c.1090A>C, p.N364H |
| MN27*      | c.732_733delTG, p.Cys244* | c.823G>T, p.V275F |
| MN28       | c.1111A>T, p.N371Y  | c.1118C>A, p.T373K |
| MN29*      | c.1184+1G>A, IVS3+1 | c.1184+1G>A, IVS3+1 |
| MN30       | c.140G>A, p.G47D   | c.140G>A, p.G47D |
| MN31       | c.1118C>A, p.T373K | c.1467dupT, p.A490Cfs*20 |
| MN32       | c.661G>A, p.E221K  | c.1118C>A, p.T373K |
| MN33       | c.1A>G, p.M17      | c.1217C>T, p.P406L |
| MN34       | c.242C>T, p.P81L   | c.1204C>T, p.R402* |
| MN35       | c.896G>A, p.R299H  | c.1199G>T, p.W400L |
| MN36       | c.242C>T, p.P81L   | c.1036+2T>G, IVS4+2 |
| MN37       | c.1217C>T, p.R406L | c.1392dup, p.Lys465* (novel) |
| MN38       | c.832C>T, p.R278*  | c.1264C>T, p.P422W |
| MN39*      | c.32G>A, p.W11*    | c.895C>A, p.R299S |
| MN40       | c.606T>G, p.H202Q  | c.1036+2T>G, IVS4+2 |
| MN41       | c.242C>T, p.P81L   | c.1118C>A, p.T373K |
| MN42*      | c.572delG, p.G191Dfs*35 | c.1150C>G, p.T384A |
| MN43       | c.1037-7T>A, p.I52-7 | c.1037-1G>A, IVS3+1 |
| MN44       | c.242C>T, p.P81L   | c.1204C>T, p.R402* |
| MN45       | c.1118C>A, p.T373K | c.1147G>A, p.D383N |
| MN46       | c.242C>T, p.P81L   | c.1037-7T>A, IVS4+2 |
| MN47       | c.242C>T, p.P81L   | c.1217C>T, p.P406L |
| MN48*      | c.149C>T, p.S50L   | c.892C>T, p.R298W |
| MN49       | c.678_680delAGG, p.G227del | c.1118C>A, p.T373K |

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gene sequencing at the National Eye Institute (Table 2). For the patients without pathogenic variant or with single pathogenic variant or variants with unknown significance in either TYR gene or OCA2 gene, we analyzed the OCA2 gene 2.7 kb exon 7 deletion. We found that none of these patients had the common exon 7 deletion. We then extended our exon 7 deletion analysis to the rest of the samples in the Tables 1 and 2. We discovered that one patient (NE27) was indeed heterozygous for the exon 7 deletion (Fig. 2). Interestingly, this patient NE27 was also positive for one known OCA2 gene pathogenic variant, c.1103G>T, p.Ala368Val [8], and one variant with uncertain significance but likely pathogenic, c.913C>T, p.Ala305Val [8], and found that they were all homozygous for p.R305W too. One additional patient who was heterozygous for the deletion was also heterozygous for the p.R305W variant. It clearly indicated the phase that the 2.7 kb deletion spanning the exon 7 of OCA2 gene was indeed in cis with the p.R305W variant in the exon 9 of OCA2 gene.

**Discussion**

In our study, we sequenced DNA samples from 108 OCA patients for TYR or OCA2 gene pathogenic variants. Our results consistently supported the published allele distributions [4]. We also performed long range PCR to search for potential large genomic rearrangements such as deletion or duplication in the TYR locus and for the 2.7 kb deletion in OCA2 gene. We did not identify any abnormality at the TYR intronic regions using the long range PCR methodology, suggesting that other mechanisms, such as splicing error variants, might be involved. We cannot rule out that this approach was not sensitive enough for smaller sizes of deletions/insertion or even larger massive genomic rearrangements [10]. In addition, we also noticed high frequent of the Alu repetitive sequences in the TYR intronic regions.

The three primer PCR assay system that we adapted from Durham et al. [9] enabled us to determine the genotype of the individuals in a single multiplex PCR with an internal wild type intron sequence control. Observation of the 802 bp deletion-specific junction fragment, the 3.5 kb fragment for the intact genomic region, and the 237 bp fragment spanning one of the breaking points in OCA2 heterozygotes provided a convenient carrier status confirmation analysis for routine clinical diagnostics (Fig. 2). In a comparison of gender, ethnic background, genotype of the individuals in a single multiplex PCR with an internal wild type intron sequence control. Observation of the 802 bp deletion-specific junction fragment, the 3.5 kb fragment for the intact genomic region, and the 237 bp fragment spanning one of the breaking points in OCA2 heterozygotes provided a convenient carrier status confirmation analysis for routine clinical diagnostics (Fig. 2).

In a comparison of gender, ethnic background, genotypes from sequencing results and the deletion status in five samples that were positive for the p.R305W variant (Table 3), besides the patient NE27, we found that the OCA2 gene in patient NE12 contains two heterozygous pathogenic variants: an A–G change at nucleotide 1465 (c.1465A>G) in exon 14, and a T–G change at nucleotide 2310 (c.2310T>G) in exon 22. These pathogenic variants change the amino acid Asparagine (N) to aspartic acid (D) at residue 489 (p.N489D) and amino acid Tyrosine...
(Y) to a termination codon (*) at residue 770 (p.Y770*), respectively, in the P protein. The pathogenic variant p.N489D has been identified in unrelated patients diagnosed with Oculocutaneous Albinism Type 2. The pathogenic variant p.Y770* has not been identified in patients diagnosed with Oculocutaneous Albinism Type 2 to our knowledge. However, this nonsense variant is predicted to cause a premature termination in translation and lead to protein product truncation and expected to be pathogenic. The p.R305W was also present in this patient with

| Patient ID | TYR | OCA2 |
|------------|-----|------|
| NE1 | c.1118C>A (p.Thr373Lys) | c.1441G>A (p.Ala481Thr) |
| NE2 | Negative | c.2228C>T (p.Pro743Leu) |
| NE3 | Negative | c.1842+1G>T (IVS17+1G>T) |
| NE4 | c.346C>T (p.Arg116*) | c.896G>A (p.Arg299His) |
| NE5* | Negative | Not tested |
| NE6 | c.816G>C (p.Trp272Cys) | c.1118C>A (p.Thr373Lys) |
| NE7* | Negative | Not tested |
| NE8 | Negative | c.1327G>A (p.Val443Ile) |
| NE9 | c.1217T>C (p.Pro406Leu) | c.1291C>A (p.Pro431Thr) |
| NE10 | c.1217T>C (p.Pro406Leu) | Not tested |
| NE11 | c.238T>C (p.Trp80Arg) | c.1037-7T>A (IVS2-7T>A) |
| NE12 | Negative | Not tested |
| NE13 | Negative | c.1103C>T (p.Ala368Val) |
| NE14 | c.929dupC (p.Arg311Lysfs*7) | c.929dupC (p.Arg311Lysfs*7) |
| NE15 | c.1118C>A (p.Thr373Lys) | Not tested |
| NE16 | Negative | c.157delA (p.Arg53Glyfs*49) |
| NE17 | Negative | c.632C>T (p.Pro211Leu) |
| NE18 | Negative | c.1240-1G>C (IVS12-1G>C) |
| NE19 | Negative | c.1372G>A (p.Val443Ile) |
| NE20 | Negative | c.913C>T (p.Arg305Trp) |
| NE21 | c.1118C>A (p.Thr373Lys) | Not tested |
| NE22 | c.242C>T (p.Pro81Leu) | c.913C>T (p.Arg305Trp) |
| NE23 | Negative | c.913C>T (p.R305W) |
| NE24 | c.1037-1G>A (IVS2-1G>A) | c.1118C>A (p.Thr373Lys) |
| NE25 | c.61C>T (p.Pro21Ser) | Not tested |
| NE26 | c.1037-7T>A (IVS2-7T>A) | Not tested |
| NE27 | Negative | c.1103C>T (p.Ala368Val) |
| NE28 | Negative | c.913C>T (p.R305W) |
| NE29 | Negative | c.1118C>A (p.Thr373Lys) |
| NE30 | c.229_230insAGG (p.Arg77delinsGlnGly) | Not tested |
| NE31 | c.1366+4A>G (IVS4+4A>G) | Not tested |
| NE32 | c.346C>T (p.Arg116*) | c.929dupC (p.Arg311Lysfs*7) |
| NE33 | Negative | Not tested |
| NE34 | c.242C>T (p.Pro81Leu) | Not tested |
| NE35 | c.1037-7T>A (IVS2-7T>A) | Not tested |
| NE36 | Negative | c.1465A>G (p.Asn489Asp) |
| NE37 | Negative | c.2228C>T (p.Pro743Leu) |
| NE38 | Negative | c.1327G>A (p.Val443Ile) |
| NE39 | c.242C>T (p.Pro81Leu) | c.1217C>T (p.Pro406Leu) |

*NE13-NE25, NE27-NE29 have been submit to BRICS for research (Additional file 2: Table S2)*

*b These two samples were found positive for pathogenic alleles in other OCA related genes later*
unknown phase. In three other patients (NE21, NE22, NE30), it was interesting to note that they were all positive for a single pathogenic variant in the \textit{TYR} gene. It was more likely that there would be an unknown secondary pathogenic variant at their \textit{TYR} locus on the other chromosome.

The \textit{OCA2} (\textit{P}) gene consists of 24 exons (23 coding), spanning almost 345 kb of genomic DNA in the region of 15q11.2-q12. The 2.7 kb deletion was expected to result in a frameshift in the first luminal loop, predicting a truncation of the polypeptide and a non-functional gene product. Regarding the protein structure and predicted function, the p.R305 locates in the first luminal loop. Kamaraj and Purohit provided a computational model to depict the functional consequence of p.R305W change [2]. It was predicted that the p.R305W leads to more flexible in conformation, which might alter the catalytic function in melanin biosynthesis. We propose that such functional alteration is not sufficient to lead

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**Fig. 1** Analysis of the 2.7 kb deletion through PCR and Sanger sequencing. Genomic region around the exon 7 of the \textit{OCA2} gene showed the primer locations, exon 7 and identified deleted region. The expected PCR products were illustrated. The breakpoints of the deletion was confirmed in the Sanger sequencing.

**Fig. 2** PCR determination of genotypes of the 2.7 kb deletion in OCA patients. Invitrogen 1 Kb Plus DNA ladder were on both sides. Lane 1 is from a patient who was homozygous of the 2.7 kb deletion [7]. Lane 2 was NE27. Lane 3 was the mother of NE27. Lane 4 was patient NE28. Lane 5 was no DNA control.
to OCA. Therefore, homozygous genotype itself would not present OCA and it is demonstrated by unaffected individuals in the population. But in a subpopulation of p.R305W, co-segregated 2.7 kb exon 7 deletion was the real disease-causing mutation. Our previously identified exon 7 deletion positive samples in homozygous and heterozygous status were consistently associated with the p.R305W variant. It suggests that the p.R305W variant was more likely an associated variant as a more ancient allele in linkage disequilibrium. The pathogenicity of the p.R305W has been discussed intensively in literature [2, 3, 11, 12]. Until this study, we were still interpreting that the p.R305W was a VUS with likely pathogenic. Based on the analysis on allele origination, our results are not consistent with the p.R305W variant functioning as a pathogenic variant in an autosomal recessive inheritance pattern. We speculate that the dispute for the pathogenicity of p.R305W was due to lack of analysis on this exon 7 related deletion in the past, which presented as disease associated marker. It should be brought to an attention for our clinical molecular diagnosis on OCA2. More attention should be given to the patient ethnic background and the possibility of a 2.7 kb exon 7 deletion associated with the p.R305W variant in sequencing tests. The p.R305W variant itself is likely benign variant in cis with the exon 7 deletion and likely misleading for OCA accurate molecular diagnosis.

### Conclusion

In summary, we herein evaluated 108 clinically diagnosed OCA patients by sequencing their TYR and OCA2 genes, and alkalized the 2.7 kb OCA2 gene deletion using long-range PCR. We confirmed the co-segregation of the deletion and the known controversial variant, p.R305W. Our results suggest that the p.R305W is the more ancestral allele and the 2.7 kb deletion is introduced into this allele more recently. These results provided evidence that do not support a pathogenic classification for the p.R305W variant. This association will have a clinical impact genetic consultation for OCA patients.

### Methods

Patients: OCA patients were clinically diagnosed in the United States. Peripheral blood was collected from each patient and DNA was extracted in the Molecular Diagnostics Laboratory in the University of Minnesota Medical Center, Fairview in Minneapolis, Minnesota from 2003 to 2008 and in the NEI DNA Diagnostic Lab at the National Eye Institute (NEI) in Bethesda, Maryland from 2009 to 2016. DNA samples were sequenced for TYR and OCA2 gene pathogenic variants in the Molecular Diagnostics Laboratory in the University of Minnesota Medical Center, Fairview or sequenced in the NEI DNA Diagnostic Laboratory at the National Eye Institute (NEI) in Bethesda, Maryland. This study was reviewed and approved by the University of Minnesota University IRB and the Combined Neuroscience Institutional Review Board of the National Institutes of Health. Informed consent was obtained from each participant as adhering to tenets of the Declarations of Helsinki.

Genomic DNA was extracted using the Puregene genomic DNA extraction kit (Qiagen), and quantified with the ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE). Genomic DNA samples from patients were sequenced using protocol described previously [13].

Long range PCR was performed using the LA PCR kit 2.1 from Takara (Clontech Laboratories, Mountain View, CA). PCR primers for the TYR gene intron regions were designed using GeneRunner 3.01 software [13] and PCR condition were optimized by annealing temperatures, magnesium concentrations, and DMSO concentrations (Additional file 1: Table S1). PCR primers for the OCA2 2.7 kb deletion and PCR conditions were adapted from

### Table 3 Comparison of OCA patients bearing the p.R305W variant

| IDs | Gender | Ethnic background | TYR Allele 1 | TYR Allele 2 | OCA2 Allele 1 | OCA2 Allele 2 |
|-----|--------|-------------------|--------------|--------------|---------------|---------------|
| NE27 | Male   | American Indian or Alaska Native, Black or African American, White | c.1103C>T (p.Ala368Val) | c.913C>T (p.Arg305Trp) VUS | 2.7 kb Exon 7 deletion |
| NE12 | Female | White             | c.1465A>G (p.Asn489Asp) c.2310T>G (p.Tyr770*) Phase unknown: c.913C>T (p.Arg305Trp) VUS |
| NE21 | Male   | White             | c.1118C>A (p.Thr373Lys) | c.913C>T (p.Arg305Trp) VUS |
| NE22 | Female | White             | c.242C>T (p.Pro81Leu) | c.913C>T (p.Arg305Trp) VUS |
| NE30 | Male   | Asian             | c.229_230insAGG (p.Arg77delinsGlnGly) | c.913C>T (p.Arg305Trp) VUS |

Asterisk represents termination codon
Durham-Pierre et al. [9]. For this modified PCR protocol, we used a 2 primer set (MHB51F + MHB71R in a 1:1 ratio) or a 3 primer set (MHB51F + MHB72F + MHB71R in a 4:1:2 ratio) combinations (Fig. 1). The PCR condition was modified as: Denaturation at 94 °C for 1 min, followed by 30 cycles of denaturation at 94 °C for 30 s and annealing at 69 °C for 4 min, followed by final elongation at 72 °C for 10 min. All PCR products were analyzed using 0.8 or 2 % agarose gel electrophoresis.

Additional files

- **Additional file 1:** Table S1. Primers for TYR intronic regions.
- **Additional file 2:** Table S2. Data accessibility.

Authors’ contributions

XW conceived and designed the experiments; JG and KW performed the experiments; XW, JG, LD and KW analyzed the data; CA, MR, DA and ST contributed sample/materials/analysis; XW and JG wrote the paper. All authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

All data generated or analyzed during this study are included in this published article and its supplementary information files. Samples were collected and studied in the Molecular Diagnostic Laboratory of University of Minnesota Medical Center, Fairview, the eyeGENE Program, the NEI Clinic, and NHGRI OCA Natural History Study. The access to information of samples from the eyeGENE Program, the NEI Clinic, and NHGRI OCA Natural History Study. The access to information of samples from the eyeGENE Program can be found at eyeGENE.nih.gov.

Ethics approval and consent to participate

This study was reviewed and approved by the University of Minnesota University IRB and the Combined Neuroscience Institutional Review Board of the National Institutes of Health. Informed consent was obtained from each participant as adhering to tenets of the Declarations of Helsinki.

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