Distinct mutations with different inheritance mode caused similar retinal dystrophies in one family: a demonstration of the importance of genetic annotations in complicated pedigrees

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Abstract

Background: Retinitis pigmentosa (RP) is the most common form of inherited retinal dystrophy presenting remarkable genetic heterogeneity. Genetic annotations would help with better clinical assessments and benefit gene therapy, and therefore should be recommended for RP patients. This report reveals the disease causing mutations in two RP pedigrees with confusing inheritance patterns using whole exome sequencing (WES).

Methods: Twenty-five participants including eight patients from two families were recruited and received comprehensive ophthalmic evaluations. WES was applied for mutation identification. Bioinformatics annotations, intrafamilial co-segregation tests, and in silico analyses were subsequently conducted for mutation verification.

Results: All patients were clinically diagnosed with RP. The first family included two siblings born to parents with consanguineous marriage; however, no potential pathogenic variant was found shared by both patients. Further analysis revealed that the female patient carried a recurrent homozygous \texttt{C8orf37 p.W185*}, while the male patient had hemizygous \texttt{OFD1 p.T120A}. The second family was found to segregate mutations in two genes, \texttt{TULP1} and \texttt{RP1}. Two patients born to consanguineous marriage carried homozygous \texttt{TULP1 p.R419W}, while a recurrent heterozygous \texttt{RP1 p.L762YF*17} was found in another four patients presenting an autosomal dominant inheritance pattern. Crystal structural analysis further indicated that the substitution from arginine to tryptophan at the highly conserved residue 419 of \texttt{TULP1} could lead to the elimination of two hydrogen bonds between residue 419 and residues V488 and S534. All four genes, including \texttt{C8orf37}, \texttt{OFD1}, \texttt{TULP1} and \texttt{RP1}, have been previously implicated in RP etiology.

Conclusions: Our study demonstrates the coexistence of diverse inheritance modes and mutations affecting distinct disease causing genes in two RP families with consanguineous marriage. Our data provide novel insights into assessments of complicated pedigrees, reinforce the genetic complexity of RP, and highlight the need for extensive molecular evaluations in such challenging families with diverse inheritance modes and mutations.

Keywords: Retinitis pigmentosa, Genetic heterogeneity, Next generation sequencing, Mutation, \texttt{OFD1}, \texttt{C8orf37}, \texttt{TULP1}, \texttt{RP1}, Consanguinity
Background
Retinitis pigmentosa (RP, MIM: 268000), the most common form of inherited retinal degenerations, affects over one million individuals globally [1, 2]. Night blindness is usually the initial symptom for RP, followed by subsequent visual field constriction, and eventual vision loss. RP is featured by great clinical heterogeneities. Its onset age ranges from early childhood to mid-adulthood. Inter- and intra-familial phenotypic diversities caused by the same RP causing mutations have also been revealed [3–5]. Thus, clinical diagnose for RP patients are sometimes challenged by its wide phenotypic spectrum and under certain conditions, like in a young patient without fully onset RP phenotypes. In such situations, molecular testing could help to address the clinical ambiguity in RP diagnosis. RP also shows high genetic heterogeneity. To date, 83 RP causing genes involving hundreds of mutations have been identified (RetNet). Next-generation sequencing (NGS), enabling simultaneous parallel sequencing of numerous genes with high efficiency, is an efficient tool for molecular diagnosis of RP [2, 4]. Genetic annotations with NGS promote better clinical assessments and gene therapy, and therefore should be recommended for RP patients. However, pedigrees with puzzling inheritance patterns could sometimes confuse the genetic diagnoses. Herein, we described the genotypic and phenotypic findings in two complicated RP pedigrees using NGS. Distinct inheritance patterns and RP causing genes/mutations were found in both families.

Methods
Sample collection and clinical assessments
Our study, conformed to the Declaration of Helsinki, was approved and prospectively reviewed by the local ethics committee of People Hospital of Ningxia Hui Autonomous Region (No. 10 [2017]). Eleven participants from family A (Fig. 1a) and 14 participants from family B (Fig. 1b) were recruited from the People’s Hospital of Ningxia Hui Autonomous Region. Written informed contents were obtained from all participants or their legal guardians before their enrollments. Peripheral blood samples were collected from all 25 participants for genomic DNA extraction. Family history and consanguineous marriages were carefully reviewed. Medical records were obtained from all participants. Each participant received general ophthalmic evaluations, while comprehensive ophthalmic examinations were selectively conducted on the eight included patients. Another 150 Chinese healthy controls free of major ocular problems were recruited with their blood samples donated.

NGS approach and bioinformatics analyses
To reveal the disease causing mutation in the two families, we selectively performed whole exome sequencing (WES) on three participants in family A (A-IV:3, A-VI:2 and A-VI:3) and two patients in family B (B-III:4 and B-IV:1). WES was conducted with the 44.1 megabases SeqCap EZ Human Exome Library v2.0 (Roche NimbleGen, Madison, WI) for enrichment of 23588 genes on patients from family A [6], and with SureSelect Human All Exon V6 60 Mb Kit (Agilent Technologies, Santa Clara, CA) on patients from family B [7]. Briefly, qualified genomic DNA samples were randomly sheared by Covaris into 200–250 base pair (bp) fragments. Fragments were then ligated with adapters to both ends, amplified by ligation-mediated polymerase chain reaction (LM-PCR), purified, and hybridized. Non-hybridized fragments were then washed out. Quantitative PCR was further applied to estimate the magnitude of enrichment of both non-captured and captured LM-PCR products. Each post-capture library was then loaded on an Illumina HiSeq 2000 platform for high-throughput sequencing.

Raw data were initially processed by CASAVA Software 1.7 (Illumina) for image analysis and base calling. Sequences were generated as 90 bp pair-end reads. Reads were aligned to human h19 genome using SOAPaligner (http://www.soap.genomics.org.cn) and Burrows-Wheeler Aligner (BWA; http://www.bio-bwa.sourceforge.net/). Only mapped reads were included for subsequent analysis. Coverage and depth were determined based on all mapped reads and the exome region. Atlas-SNP2 and Atlas-Indel2 were applied for variant calling [8]. Variant frequency data were obtained from the following six single nucleotide polymorphism databases, including
Table 1 Clinical features of attainable patients

| Family member ID | RP causative gene | Age (year)/sex | Onset age (year) | Night blindness | Cataract | BCVA (logMAR) | Fundus appearance | ERG |
|------------------|-------------------|----------------|------------------|----------------|----------|---------------|-------------------|-----|
|                  |                   |                |                  |                |          | O.D.          | O.S.              |     |
|                  |                   |                |                  |                |          | O.D.          | O.S.              |     |
| A-V2*            | –                 | –              | 10               | Yes            | –        | –             | –                 |     |
| A-V12            | C8orf37           | 25/F           | 8                | Yes            | No       | LP            | LP                |     |
| A-V13            | OFD1              | 24/M           | 2                | Yes            | No       | LP            | LP                |     |
| B-II4            | RP1               | 80/F           | 50               | Yes            | Severe   | NLP           | LP                |     |
| B-III3           | RP1               | 59/M           | 30               | Yes            | IOL      | 0.6           | 0.25              |     |
| B-III5           | RP1               | 54/F           | 35               | Yes            | Mild     | 0.3           | 0.3               |     |
| B-IV1            | TULP1             | 27/M           | EC               | Yes            | Moderate | 0.15          | 0.2               |     |
| B-IV2            | TULP1             | 24/F           | EC               | Yes            | Moderate | 0.3           | 0.3               |     |
| B-IV4            | RP1               | 31/F           | –                | Yes            | No       | 0.5           | 0.8               |     |

* Female, M male, EC early childhood, BCVA best corrected visual acuity, logMAR logarithm of the minimum angle of resolution, O.D. right eye, O.S. left eye, IOL intraocular lens, LP light perception, NLP non-light perception, MD macular degeneration, OD optic disk, AA artery attenuation, PD pigment deposits, ERG electroretinography, D diminished, R reduced

* This patient is deceased. His clinical features are obtained based on his medical records.
B. taurus (ENSBTAP00000055698), for all 6 patients, while patient B-II:4 also had chronic nyctalopia since early childhood, while the other four patients showed RP symptoms elder than 30-year-old. RP progression also varied among the 6 patients. RP onset ages ranged from early childhood to 50 years old (Table 1). RP progression also varied among the 6 patients. Patients B-IV:1 and B-IV:2 reported to have onset nyctalopia and rapid disease progress. Best corrected visual acuity was light perception for both patients from family B, B-II:4, B-III:3 and B-III:4. The mutated residue R419 in TULP1 proteins were obtained based on human TUB (FBpp0088961), and C. elegans (F10B5.4). Crystal structural modeling of the wild type and mutant TULP1 proteins were constructed with SWISS-MODEL online server (10, 11), and displayed with PyMol software.

In silico analysis
We applied vector NTI Advance™ 2011 software (Invitrogen, Carlsbad, CA) to analyze the conservation of the mutated residue by aligning protein sequence of human TULP1 (ENSP00000229771) with sequences of the following orthologues proteins: P. troglodytes (ENSEP-TRP000000030898), C. lupus (ENSCAFP0000001922), B. taurus (ENSBTAP00000055698), M. musculus (ENS-MUSP00000049070), G. gallus (ENSGALP0000010281), D. rerio (ENSDARP00000099556), D. melanogaster (FBpp0088961), and C. elegans (F10B5.4). Crystal structural modeling of the wild type and mutant TULP1 proteins were constructed with SWISS-MODEL online server (10, 11), and displayed with PyMol software.

Results
Clinical findings
Two patients from family A, A-VI:2 and A-VI:3, and six patients from family B, B-II:4, B-III:3, B-III:5, B-IV:1, B-IV:2 and B-IV:4, were included in the present study with their clinical details summarized in Table 1. Ophthalmic features of patient A-V:2 were obtained according to his medical records, and were presented in Table 1. All patients from the two families were clinically diagnosed with RP. In family A, all three patients had early onset nyctalopia and rapid disease progress. Best corrected visual acuity was light perception for both patients A-VI:2 and A-VI:3 at their last visit to our hospital at the ages of 25 and 24 respectively. Typical RP presentations and macular degeneration were detected upon their ophthalmic evaluations (Fig. 2A–G and Table 1). In family B, RP onset ages ranged from early childhood to 50 years old (Table 1). RP progression also varied among the 6 patients. Patients B-IV:1 and B-IV:2 reported to have nyctalopia since early childhood, while the other four patients showed RP symptoms elder than 30-year-old. On examination, typical RP presentations were detected for all 6 patients, while patient B-II:4 also had chronic angle closure glaucoma in her right eye (Fig. 2H–S). Noteworthy, all 6 patients presented mild to severe cataracts (Table 1). Patient B-III:3 received bilateral cataract surgeries 2 years ago. No systemic defect was noticed in any of the included patients.

Genetic assessments
To identify the pathogenic mutations, WES with high quality was selectively performed on individuals A-IV:3, A-VI:2, and A-VI:3 from family A (mean coverage: 98.16%; mean depth: 70.89 × ), and patients B-III:5 and B-IV:1 from family B (mean coverage: 98.32%; mean depth: 104.66 × ). NGS data were summarized in Additional file 3: Table S3. Exon-specific coverage report of all known RP genes was presented in Additional file 4: Table S4. For family A, patients A-VI:2 and A-VI:3 were born to parents with consanguineous marriage, supporting potential autosomal recessive inheritance. WES identified 10 homozygous variants and 6 compound heterozygous variants shared by patients A-VI:2 and A-VI:3 (Additional file 1: Table S1). However, Sanger sequencing revealed no variant co-segregated with the disease phenotype. We thus hypothesized that the two patients may have distinct RP causing mutations. Based on WES data, patient A-VI:2 carried a recurrent homozygous C8ORF37 mutation c.555G>A (p.W185*; Fig. 1d and Table 2), while patient A-VI:3 had a novel hemizygous OFD1 mutation c.358A>G (p.T120A; Fig. 1e and Table 2).

As to family B, WES revealed one homozygous variant and 18 compound heterozygous variants shared by patients B-III:4 and IV:2 (Additional file 2: Table S2), while no variant was validated co-segregated with the disease phenotype. According to the family pedigree, patients B-IV:1 and B-IV:2 were born to unaffected parents with consanguineous marriage, indicating a potential autosomal recessive inheritance pattern. However, the RP phenotypes of patients B-III:3 and B-III:4 were likely inherited from the affected mother B-II:4, suggesting a dominant inheritance mode. Upon this hypothesis, a novel homozygous TULP1 mutation c.1255C>T (p.R419W; Fig. 1e and Table 2) was revealed as RP causative for patients B-IV:1 and B-IV:2, and a recurrent heterozygous RPI1 mutation c.2285_2289delTAAT (p.L762Yfs*17; Fig. 1f; Table 2) was found in patients B-II:4, B-III:3 and B-III:4. The mutated residue R419 in TULP1 was highly conserved among all tested species (Fig. 1g). Crystal structures of the wild type and mutant TULP1 proteins were obtained based on human TUB protein (Protein Data Bank ID: 1S31) with a sequence identity of 75.19 and a sequence similarity of 0.54. Our data suggested that the substitution from arginine to tryptophan at residue 419 would lead to the elimination of two hydrogen bonds between residue 419 and residues.
V488 and S534 (Fig. 1h, i), further supporting that this mutation would disturb the tertiary structure of TULP1 and interrupt its function. Residues R419, N463, V488 and S534 were conserved between TULP1 and TUB proteins (Fig. 1j). All four mutations identified in the two families segregated with the disease phenotype (Fig. 1a,
| Gene | Variation | Status | Bioinformatics analysis | Reported or Novel | Population prevalence (allele count) |
|------|-----------|--------|-------------------------|-------------------|-------------------------------------|
|      |           |        |                         |                   | rs no.       | gnomAD     | EXAC     |
| C8orf37 | c.555G>A | Hom    | NA                      | NA                | NA       | Novel  | rs748014296 | 2/246148 | 1/121412 |
| OFD1  | c.358A>G  | Hem    | 0.63 (tolerated)        | 0.006 (benign)    | − 0.616 (neutral) | Novel | rs755625951 | 4/178544 | 1/121388 |
| TULP1 | c.1255C>T | Hom    | 0 (damaging)            | 1 (probably damaging) | 7.976 (deleterious) | Novel | rs775334320 | 12/217192 | 6/121222 |
| RP1   | c.2285_2289delTAAT | Het | NA                      | NA                | NA       | Novel  | NA        | NA       | NA         |

*Hom homzygous, Hem hemizygous, Het heterozygous, NA not available*

SIFT: http://sift.bii.a-star.edu.sg/; PolyPhen: http://genetics.bwh.harvard.edu/pph2/; PROVEN: http://provean.jcvi.org/index.php; gnomAD: http://gnomad.broadinstitute.org/; EXAC: http://exac.broadinstitute.org/
| Gene       | Variation       | Nucleotide | Amino acid | Domain     | Disease     | References |
|------------|-----------------|------------|------------|------------|-------------|------------|
| C8ORF37    | c.155+2T>C      | –          | –          | CRD        | [56]        |
| C8ORF37    | c.156−2A>G      | –          | –          | CRD        | [15, 18]    |
| C8ORF37    | c.243+2T>C      | –          | –          | RP         | [21]        |
| C8ORF37    | c.244−2A>C      | –          | –          | RP         | [17]        |
| C8ORF37    | c.374+2T>C      | –          | –          | EORD       | [20]        |
| C8ORF37    | c.497>A         | p.L166*    | –          | RP         | [15, 18]    |
| C8ORF37    | c.529C>T        | p.R177W    | –          | CRD, BBS   | [15, 18, 19, 22] |
| C8ORF37    | c.545A>G        | p.Q182R    | –          | RP         | [15, 18]    |
| C8ORF37    | c.555G>A        | p.W185*    | –          | RP         | [17]        |
| C8ORF37    | c.575delC       | p.T192Mfs*28 | –          | EORD       | [20]        |
| OFD1       | IVS9+706A>G     | p.N313fs*330 | Coiled coil domain | RP | [13]        |
| TULP1      | c.3G>A          | p.M11      | –          | RP         | [25]        |
| TULP1      | c.99+1G>A       | –          | –          | LCA, RP    | [23, 26]    |
| TULP1      | c.280G>T        | p.D94Y     | –          | LCA        | [27]        |
| TULP1      | c.286, 287delGA | p.E96Gfs*77 | –          | RP         | [57]        |
| TULP1      | c.350−2delAGA   | –          | –          | RP         | [28]        |
| TULP1      | c.394, 417del   | p.E120_D127del | –          | RP         | [20]        |
| TULP1      | c.539G>A        | p.R180H    | –          | LCA        | [30]        |
| TULP1      | c.627delC       | p.S210Qfs*77 | –          | RP         | [32]        |
| TULP1      | c.629C>G        | p.S210*    | –          | RP         | [33]        |
| TULP1      | c.718+2T>C      | –          | –          | LCA, RP    | [33]        |
| TULP1      | c.1275_1328delCCAA | p.P242Qfs*16 | –          | LCA        | [34]        |
| TULP1      | c.901C>T        | p.Q301*    | Tubby domain | LCA, CRD  | [35, 36]    |
| TULP1      | c.937delC       | p.Q301fs*9 | Tubby domain | RP         | [28]        |
| TULP1      | c.932G>A        | p.R311Q    | Tubby domain | RP         | [37]        |
| TULP1      | c.956G>A        | p.G319D    | Tubby domain | RP         | [38]        |
| TULP1      | c.961T>G        | p.Y321D    | Tubby domain | LCA        | [34]        |
| TULP1      | c.999+5G>C      | –          | Tubby domain | LCA, RP    | [33]        |
| TULP1      | c.1025G>A       | p.R342Q    | Tubby domain | RP         | [37]        |
| TULP1      | c.1047T>G       | p.N349K    | Tubby domain | RP         | [39]        |
| TULP1      | c.1064A>T       | p.D355V    | Tubby domain | LCA        | [34]        |
| TULP1      | c.1087G>A       | p.G363R    | Tubby domain | CRD        | [40]        |
| TULP1      | c.1081C>T       | p.R361*    | Tubby domain | LCA        | [41]        |
| TULP1      | c.1102G>T       | p.G368W    | Tubby domain | LCA        | [26]        |
| TULP1      | c.1112+2T>C     | –          | Tubby domain | RP         | [42]        |
| TULP1      | c.1113−2A>C     | –          | Tubby domain | LCA        | [34]        |
| TULP1      | c.1138A>G       | p.T380A    | Tubby domain | LCA, RP    | [43, 45, 46] |
| TULP1      | c.1145T>C       | p.F382S    | Tubby domain | RP         | [47]        |
| TULP1      | c.1198C>T       | p.R400W    | Tubby domain | LCA, RP, CRD | [26, 48, 49] |
| TULP1      | c.1199G>A       | p.A400Q    | Tubby domain | RP         | [50]        |
| TULP1      | c.1204G>T       | p.E402*    | Tubby domain | LCA        | [26]        |
| TULP1      | c.1224+4A>G     | –          | Tubby domain | RP         | [29]        |
| TULP1      | c.1246C>T       | p.R416C    | Tubby domain | RP         | [25]        |
| TULP1      | c.1255C>T       | p.R419W    | Tubby domain | RP         | This study  |
| TULP1      | c.1258C>A       | p.R420S    | Tubby domain | RCD        | [51]        |
| TULP1      | c.1259G>C       | p.R420P    | Tubby domain | RP         | [23]        |
| TULP1      | c.1318C>T       | p.R440*    | Tubby domain | LCA        | [31]        |
b), and were confirmed absent in 150 Chinese controls free of major ocular problems.

**Discussion**

RP is a genetically heterogeneous disease with 83 disease causative genes and hundreds of mutations. In this report, molecular test reveals the coexistence of mutations affecting distinct RP causing genes in two RP families, thus providing novel insights into genetic assessments in complicated pedigrees. Among the four mutations identified in the two families, two were novel (\textit{OFD1} p.T120A and \textit{TULP1} p.R419W) and two were recurrent (\textit{C8ORF37} p.W185* and \textit{RP1} p.L762Yfs*17 [Human Gene Mutation Database ID: CD991855]).

\textit{OFD1} mutations have been reported to cause X-linked recessive Joubert syndrome, orofaciodigital syndrome and isolated RP (Table 3) [12, 13]. \textit{OFD1}, protein encoded by the \textit{OFD1} gene, is a crucial component of the centrioles. \textit{OFD1} is involved in ciliogenesis regulation and exhibits neuroprotective roles [14]. Herein, a hemizygous \textit{OFD1} missense mutation is associated with a severe form of RP presenting early onset age and fast disease progression. \textit{C8ORF37} mutations correlate with a wide spectrum of autosomal recessive retinopathies ranging from RP to Bardet–Biedl syndrome (Table 3) [15–22]. The encoded \textit{C8ORF37} protein is a ciliary protein located at the base of the photoreceptor connecting cilia [16], while its role in modulating retinal function is not fully elucidated. In this study, the patient carrying homozygous nonsense \textit{C8ORF37} mutation presents early onset RP with macular involvement, which is similar to previous reports [15, 17]. \textit{TULP1} mutations are implicated in autosomal recessive RP and LCA etiologies (Table 3) [22–57]. \textit{TULP1} protein plays crucial roles in maintaining retinal homeostasis. According to previous reports, \textit{TULP1} interacts and co-localizes with F-actin in photoreceptor cells of bovine retina [58], and RPE phagocytosis ability was remarkably reduced in \textit{TULP1}−/− mice [59]. Thus, \textit{TULP1} is required for maintaining regular functions of photoreceptors and RPE cells. We herein identified \textit{TULP1} mutations in two siblings demonstrating RP with early onset and quick progression. Further confirmatory functional studies are still needed to better illustrated pathogenesis of the identified novel mutations.

**Conclusions**

In summary, we demonstrate the coexistence of diverse inheritance modes and mutations affecting distinct disease causing genes in two RP families. Our findings reinforce the genetic complexity of RP, provide novel insights into the assessments of complicated pedigrees with consanguinity, and highlight the need for extensive molecular evaluations in such challenging families involving diverse inheritance modes and mutations.
Additional files

Additional file 1: Table S1. Post-filtration variants in family A.

Additional file 2: Table S2. Post-filtration variants in family B.

Additional file 3: Table S3. Overview of data production.

Additional file 4: Table S4. Coverage for all exons in all known RP genes.

Abbreviations

RP: retinitis pigmentosa; NGS: next-generation sequencing; WES: whole exome sequencing; bp: base pair; LM-PCR: ligation-mediated polymerase chain reaction; RPE: retinal pigment epithelium.

Authors’ contributions

XC, XS, YL, and ZL contributed equally to this report. All authors were involved in managing the patients. XC, BY and CZ wrote the report. XC, XS, YL and ZL did the genetic analysis and whole exome sequencing, and CZ reviewed the genetic results. 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

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Consent for publication

Yes.

Ethics approval and consent to participate

Our study, conforming to the Declaration of Helsinki, was approved and prospectively reviewed by the local ethics committee of People Hospital of Ningxia Hui Autonomous Region. Written informed contents were obtained from all participants or their legal guardians before their enrollments.

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