Next generation sequencing of RB1 gene for the molecular diagnosis of ethnic minority with retinoblastoma in Yunnan

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Abstract

**Background:** Retinoblastoma is a rare intraocular malignancy and typically initiated by inactivating biallelic mutations of *RB1* gene. Each year, ~8,000 children worldwide are diagnosed for retinoblastoma. In high-income countries, patient survival is over 95% while low-income countries is ~30%. If disease is diagnosed early and treated in centers specializing in retinoblastoma, the survival might exceed 95% and many eyes could be safely treated and support a lifetime of good vision. In China, approximate 1,100 newly diagnosed cases are expected annually and 28 hospitals covering 25 provinces established centers classified by expertise and resources for better treatment options and follow-up. Comparing with other province of eastern China, Yunnan province is remote geographically. This might result that healthcare staff have low awareness of the role of genetic testing in management and screening in families.

**Methods:** The patients with retinoblastoma were selected in Yunnan. DNA from blood was used for targeted gene sequencing. Then, an in-house bioinformatics pipeline was done to detect both single nucleotide variants and small insertions/deletions. The pathogenic mutations were identified and further confirmed by conventional methods and cosegregation in families.

**Results:** Using our approach, targeted next generation sequencing was used to detect the mutation of these 12 probands. Bioinformatic predictions showed that nine mutations were found in our study and four were novel pathogenic variants in these nine mutations.

**Conclusions:** It’s the first report to describe *RB1* mutations in Yunnan children with retinoblastoma. This study would improve role of genetic testing for management and family screening.

**Background**
Retinoblastoma (RB, OMIM#180200) is a rare malignant tumor which rapidly develops from the immature cells of a retina and occurs in infancy or in children, usually before the age of five years (about 2/3 children below the age of two years, about 95% children below the age of five years) [1, 2]. The incidence of retinoblastoma is approximately 1 in 16,000–18,000 live births, regardless of sex, race, or geography [3–5]. In the United States and Europe, incidence rate of RB is 2–5 cases per million children according to WHO [6]. In India, rates of RB incidence are 1.9–12.3 and 1.3–6.7 per million in boys and girls, respectively [7]. Dimaras Helen et al. have estimated that approximate 8,000 new cases are predicted each year in worldwide [8]. In China, there are about 1100–1500 new cases each year, but only 50% those children survive [9]. The disease-free survival rate of children with RB in developed countries is over 95% [10] while this is substantially lower, at 10–30% in developing countries [11, 12].

A heritable form and non-heritable form are two forms of the RB disease. Heritable RB accounts for 45% of all cases which 80% bilateral, 15% unilateral and 5% trilateral. Approximately 55% of cases are non-heritable RB that is always unilateral [13]. From presenting signs and clinical examination, diagnosis of RB is usually clear [14]. Leukocoria (white pupil) is the most common sign; Strabismus is the second most common sign when central vision is lost. Due to increased pressure, or non-infective orbital inflammation, advanced disease might present with enlarged cornea and iris colour change [8, 15].

The primary goal of management is to save the child’s life, followed by salvage of the eye and optimization of visual function. Many treatment options of RB are available and depend on the laterality and extent of RB disease [16]. The various treatment modalities for retinoblastoma includes enucleation of the eye, external beam radiotherapy, brachytherapy, thermotherapy, laser photocoagulation,
cryotherapy, systemic chemotherapy, intra-arterial chemotherapy, nanoparticulate chemotherapy and chemoreduction [17]. When children with RB have symptoms, such as leukokoria and strabismus, it is late for these children to have best time for cure. Delayed diagnosis results in incurable invasion of the optic nerve and brain and metastases elsewhere in the body. However, if noticed early, prompt treatment can cure the cancer and save the eye(s) [8]. According to Intraocular International Retinoblastoma Classify (IIRC) which divides retinoblastomas into 5 groups (labeled A through E; Group A means small tumors while group E means large tumor and the eye have no chance to be saved), Children’s Hospital of Fudan University reported that about 75% children with RB were diagnosed at group D and E of IIRC when these children saw doctor for the first time [18] [19].

In the management of RB, genetic testing and counseling are extremely important [20]. Heritability form of RB can be identified by genetic testing of the proband [21]. But, genetic testing has not been carried out in Yunnan. Therefore, how to carry out genetic testing is an urgent problem in the RB clinical in developing countries. In this paper, the genes of RB family in Yunnan were studied in order to obtain genetic forms of RB by genetic testing of the probands and then give counseling to these families. We use targeted next generation sequencing (NGS) to screen probands and then Sanger sequencing was used to identify variants in other members of pedigrees. In our study, nine mutations were found and four were novel mutations.

Methods

Subjects
The non-consanguineous families from the Yun-Gui Plateau was recruited by the Children’s Hospital of Kunming Medical University for genetic diagnosis. In these families, the children suffered from RB, but the parents and other members were normal. This study
was approved by the Ethics Committee of the Children’s Hospital of Kunming Medical University; in accordance with the principles of the Declaration of Helsinki written informed consent was obtained from the participants or their guardians.

Clinical evaluations

All clinical and physical examinations were conducted in the Children’s Hospital of Kunming Medical University in Kunming. CT and ultrasound imaging were performed in these probands.

Targeted NGS and variant analysis

Two-millilitre peripheral blood samples were collected from the probands, siblings and their parents in tubes containing 0.2 M EDTA. Using the QIAamp DNA blood extraction kit (TIANGEN, Beijing, China), DNA was extracted from the venous blood of each subject. According to the manufacturer’s protocol (MyGenostics, Inc., Beijing, China), 3 micrograms of genomic DNA was fragmented by Covaris 32, and the 3’ end of each DNA fragment was A-tailed. Then, Illumina adapters were ligated to these fragments. We aimed for a 350–400 base-pair product, and all samples were checked with Nanodrop 2000 or Qubit systems to determine if they represented a captured library.

Each resulting captured library was loaded on an Illumina MiSeq 2000 sequencing platform, and the sequences were determined to ensure that each sample met the desired average sequencing coverage.

Mutation analysis

Using Bcl2Fastq software (Bcl2Fastq 2.18.0.12, Illumina, Inc.), raw image files were processed for base calling and raw data generation. In addition, to get a quality score ≥20, low-quality variations were filtered out. Then, Short Oligonucleotide Analysis Package (SOAP) aligner software (SOAP2.21, soap.genomics.org.cn/soapsnp.html) was used to align the clean reads to the reference human genome (UCSC hg19,
Polymerase chain reaction (PCR) duplicates were removed by the Picard programme[22, 23]. The single nucleotide polymorphisms (SNPs) were determined by the SOAPsnp programme[24]. The reads were realigned by Burrows-Wheeler Aligner (BWA) software 0.7.15, and the deletions and insertions (InDels) were detected by Genome Analysis Toolkit software 3.7. In addition, the identified indel SNPs were annotated using the Exome-assistant programme (http://122.228.158.106/exomeassistant). To determine their pathogenicity, non-synonymous variants were evaluated by four algorithms, namely, PolyPhen (http://genetics.bwh.harvard.edu/pph2/), Protein Analysis Through Evolutionary Relationships (PANTHER, www.pantherdb.org), Sorting Intolerant from Tolerant [SIFT, (http://sift.jcvi.org/)] and Pathogenic Mutation Prediction (Pmut; http://mmb.pcb.ub.es/PMut/).

**Mutation validation**

PCR and Sanger sequencing with an ABI3500 sequencer were used to confirm potential causative variants in this family. The sites of variation were identified to compare the DNA sequences with the corresponding GenBank (www.ncbi.nlm.nih.gov) reference sequences. The sequences of forward and reverse primers are presented in supplementary table 1 and were used to confirm potential causative variants in this family.

Thermocycling conditions: An initial denaturation of 95°C for 10 min, 35 cycles of denaturation at 94°C for 30 sec., annealing at 64°C for 30 sec, extension at 72°C for 45 sec and a final extension of 72°C for 5 min. The sequences of forward and reverse primers are in supplementary table 1.

**Results**

**Clinical findings**

Twelve patients with RB were selected for this study of targeted *RB1* sequencing from
non-consanguineous Yunnan families. There are no pathogenic or likely pathogenic variants were detected in three probands with our approach while nine variants in nine families.

In these nine families, there are seven boys and two girls (See Fig. 1). Minority nationalities have Dai, Hani, Bai, Hui, Yi and Han. Mean age at diagnosis is 23±9.9 months for unilateral and 11±8.5 months for bilateral retinoblastoma in our study, respectively.

Six probands were diagnosed with bilateral RB while three with the unilateral form. There are common sign leukocoria in probands of families 1, 3, 4, 6, 7; in family 2, left eye of the proband had been removed; the proband of 5 had strange reflection; the proband had blurred vision and shed tears in 8 and 9 (table1).

Targeted NGS
The mean read depth of coverage for each proband sample ranged from 52 to 66X, and the average throughput depth of the target region in each sample ranged from 90 to 99X. NGS produced reads from 32.8 to 56.12 million reads and the read length from 148 to 149 bp. The reads aligned to the human genome and mapped to the target region with a mean coverage from 99.8 to 99.9%. The SNPs and Indels are reported in supplementary table 2.

Identification of SNVs and InDels in children with RB patients
To detect SNVs and InDels, we analyzed blood samples of 6 bilateral patients and 3 familial unilateral patient and identified pathogenic variants in 8 patients and likely pathogenic variants in 1 patient. Four were novel and five were previously reported (Table2). The spectrum of identified mutations includes 5 SNVs (1 nonsynonymous and 4 stop gain), one deletion (1 frameshift), and 3 splice site variants (splice).

Seven of them were shown to be de novo, and remaining two were inherited from one of their mothers.
Identification of pathogenic mutation

Sanger sequencing and cosegregation further confirmed all the pathogenic variants (See Fig.2 and 3). We detected 9 mutations which 4 were novel and 5 were known (Table 2).

Discussion

RB is the most common intraocular malignancy in childhood and presents in one or both eyes. Through presenting signs and clinical examination, diagnosis of RB is usually clear [14]. In our study, five probands had leukocoria; one proband had strange reflection; one proband’s left eye was enucleated; two had blurred vision. Mean age at diagnosis was 23±9.9 months for unilateral and 11±8.5 months for bilateral. According to Gene Reviews and AlAli et al (Mean age at diagnosis is 24 months for unilateral and 15 months for bilateral; mean age at diagnosis to be 27 months for unilateral and 15 months for bilateral, respectively), our mean age at diagnosis is not later than elsewhere in the world [25].

Genetic testing of RB1 is beneficial to provide counselling for families. In patients with RB, identification of gene alterations improves clinical management of patient and relatives at risk[26]. Here, we have used targeted NGS approach for the molecular analysis of Yunnan Children with RB, based on targeted gene enrichment and bioinformatics pipeline. We used in-house pipeline to successfully detect both pathogenic variants in RB patients. The average throughput depth of the target region in each sample ranged from 90–99X. These met the desired average sequencing coverage which was ensured to provide high quality bases for sensitive and efficient variant detection. To detect SNVs and InDels for all the samples, we developed an automatic in-house variant calling pipeline as freely available tools. The pathogenic SNVs and InDels were identified by stringent criteria, and 8 the pathogenic and 1 likely pathogenic variants were further confirmed by conventional methods and cosegregation with phenotype. Targeted mutation analysis is useful to study
mutations in blood and can detect DNA variations. The \textit{RB1} gene of seven families’ member are normal and only in family 3 and 7, the unaffected mother of this family are heterozygous and the same to the proband. Seven proband’s gene are spontaneous mutation. Four mutations were newly discovered mutations and never reported before. With genetic testing, mutation profiles might be created to precisely screen mutations of relatives or subsequent generations in these families. Four families’ other children had no mutations by gene testing while other five families are one-child family. Clearly, study of social determinants of health, such as health seeking behavior, perceptions of medical care, and sociocultural issues related to cancer inheritance would inform counselling approaches that meet the needs of families [27]. This study is helpful for the molecular diagnosis of RB in a comprehensive in Yunnan. These might provide molecular diagnosis to doctor for RB management because lack of genetic testing counseling, poor access to multidisciplinary retinoblastoma-specific health care and socioeconomic factors are one of the factor of higher mortality Globally. However, in this cohort, the small number of patients is not enough to establish a significant frequency reference and functional studies are necessary for assigning pathogenicity to these novel variants.

Conclusion

Here, we reported that this approach with bioinformatics pipeline could detect variants including novel pathogenic variants. This comprehensive approach reduces the time and number of assays required for the detection of pathogenic variants by conventional methods. To the best of our knowledge, this is the first such study using targeted NGS approach to detect pathogenic variants in Yunnan children with RB. Overall, targeted NGS approach is becoming more feasible in clinical settings.
Declarations

Abbreviations

RB: Retinoblastoma
IIRC: Intraocular International Retinoblastoma Classify
NGS: Next generation sequencing
SOAP: Short Oligonucleotide Analysis Package
PCR: Polymerase chain reaction
SNPs: Single nucleotide polymorphisms
BWA: Burrows-Wheeler Aligner
InDels: Deletions and insertions
SIFT: Sorting Intolerant from Tolerant
SNVs: Single nucleotide variants

Ethics approval and consent to participate
The present study was approved by the ethics committee of the Children’s Hospital, Kunming Medical University; written informed consent was obtained from participants or their guardians.

Consent for publication
Written informed consent for publication of their clinical details and data was obtained from their guardians.

Availability of data and materials
The data sets used and analyzed during this study are available from the corresponding author on reasonable request.

Competing interests
The authors declare that they have no competing interests.

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Author contributions
Z Z, Y S X and R S conceived and designed the experiments; Z Z, R S and J M performed the experiments; Z Z, Y S X, J M, HC J, X H Y and L T analyzed the data; Y S X recruited patients and collected clinical information. YS X and H Y G contributed to accumulation and interpretation of clinical data. WJ H and Z Z coordinated the project. All authors read and approved the final manuscript.

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Tables

Due to technical limitations, all table files are only available for download from the Supplementary Files section.

Figures
Figure 1

Pedigree of families with RB. Unaffected subjects are denoted as blank while affected subjects are represented with darkened symbols. The arrow indicates the proband. A. Pedigree of the family 1; B. Pedigree of the family 2; C. Pedigree of the family 3; D. Pedigree of the family 4; E. Pedigree of the family 5; F. Pedigree of the family 6; G. Pedigree of the family 7; H. Pedigree of the family 8; I. Pedigree of the family 9;
Figure 2

The variants of the probands. Arrows denote the mutations (the proband).

Mutations of retinoblastoma identified by Sanger Sequencing in family 1, 4, 7. (A) The mutation of the family 1; (B) The mutation of the family 4; (C) The mutation of the family 7.
Figure 3

The variants of the probands in family 3.

Supplementary Files
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