Abstract. Autosomal recessive cornea plana is a very rare hereditary ocular disease, characterized by a flattened corneal curvature, marked hyperopia due to low refractive power and frequently consequent accommodative esotropia. Other features include various cornea anterior segment abnormalities, without systemic problems. The purpose of the present study was to investigate the clinical and molecular alterations in a Chinese family with cornea plana. Full ophthalmic examinations of the patients were performed, including slit-lamp examination, fundus examination and ocular ultrasound. Whole-exome sequencing data were screened for pathological variants in the proband, which were confirmed by Sanger sequencing. One novel missense mutation, c.242A>G (p.N81S) and another novel 7 base-pair deletion mutation, c.772_779del (p.G258Cfs*30), were detected in the keratocan (KERA) gene; two affected siblings inherited these variations in a compound heterozygous state, which were derived from the clinically unaffected heterozygous father (c.772_779del) and mother (c.242A>G), respectively. Neither mutation was observed in unrelated healthy controls (n=200). Multiple computer software predictions supported the pathogenicity of the two variants. Furthermore, protein modeling prediction was performed to better understand the molecular basis of cornea plana, particularly the importance of the leucine-rich repeat domain. This study presents the 14th pathogenic KERA mutations identified worldwide and the first in East Asia so far, to the best of our knowledge. These findings guided prenatal diagnosis for the family in question and expand on the variant spectrum of KERA, therefore facilitating genetic counseling.

Introduction

Cornea plana is a rare disorder characterized by small flat corneas, marked hypermetropia, reduced visual activity, iris abnormalities, a hazy corneal limbus and corneal clouding, with or without associated accommodative esotropia. It is inherited as either an autosomal dominant (CNA1; OMIM121400) or autosomal recessive trait (CNA2; OMIM217300); The defective gene in CNA1 has not been identified. CNA2 are associated with the more severe phenotype and additional ocular manifestations, including the presence of a degree of iris malformation and adhesions between the cornea and iris (1-8). Corneal ectasia, corneal hydrops and idiopathic corneal decompensation are very rare findings (2,9-11).

In humans, bi-allelic mutations in keratocan (KERA) cause CNA2. The gene, which has been localized to chromosome 12q22, contains two introns and three exons; a total of 352 amino acids are translated from the majority of exon 2 and a small portion of exon 3 to form a precursor KERA protein, and exon 1 remains untranslated (12). The gene encodes KERA, an evolutionarily-conserved small leucine-rich proteoglycan (SLRP). The SLRPs are a family of highly conserved extracellular matrix proteoglycans with core proteins composed of leucine-rich repeats (LRRs). LRRs are composed of a series of parallel β-strands, which stack into an arched β-sheet array. The leucine-rich region LXXLXLXXNL is highly conserved: N represents cysteine or asparagine; L represents a hydrophobic amino acid or leucine; and X represents any amino acid. The interior of the LRR domain is composed of hydrophobic residues, with the exception of the conserved asparagines (Asn, N) which create an Asn-ladder stabilizing the structure, as reported...
in previous studies (13-15). In addition, the KERA protein is highly expressed in the cornea and is considered to be a key regulator of collagen fibrin production, which regulates the diameter and spacing of fibrils and determines corneal structure and transparency (16). Mouse studies indicate that this gene is exclusively expressed in the cornea and serves an important role during eye development (17). Narrow anterior chamber angles, thin corneal stroma and large-diameter corneal fibers are identified in KERA-knockout mice and no other systemic abnormality (18). To date, 13 different disease-associated KERA mutations have been documented in various populations, all of which manifest as various bilateral anterior segment lesions, including seven missense mutations, three nonsense mutations, one frame shift mutation, one splice-site mutation and one 3 base-pair deletion mutation. Notably, mutations in KERA have not been reported in an Asian population to date (1-4,6-10,19-24).

The present study reports two novel KERA mutations in a Chinese family. The family history was consistent with an autosomal recessive mode of inheritance. Clinical detection and genetic analysis was conducted on five family members.

Materials and methods

Subject recruitment and clinical examination. A family with CNA2 was recruited and identified at the Hunan Jiahui Genetics Hospital (Changsha, China) in October 2017. A pedigree of the family is illustrated in Fig. 1A. The proband, III:2, developed visual disorders shortly following birth, but hadn't been diagnosed or treated. She visited the doctor at the age of 26 years due to a considerable decrease in binocular visual acuity and accompanied by photophobia. Clinical and eye examinations demonstrated temporal iridocorneal synchiae, corneal clouding (Fig. 1B) and an unclear fundus. In addition, she was pregnant at the same time. A similar phenotype was observed in the younger brother of the proband, III:3.

Best-corrected visual acuity (BCVA) was established using the Acta Ophthalmologica 2017 LogMAR charts. Visual acuity was examined using the ETDRS chart. Intraocular pressure was measured by Goldmann applanation tonometry. Anterior chamber depth and axial length were measured with ocular A ultrasound. Motility, pupillary reaction, ocular fundus, slit lamp examinations and keratometry (Zeiss keratometer; Zeiss AG) were performed and documented. The growth and development of III:2 and III:3 were in the normal range. Physical examinations were performed to exclude systemic diseases.

Ethical approval. The present study was performed according to the guidelines approved by the Ethics Committee of Hunan Jiahui Genetics Hospital and adhered to the tenets set out in the Helsinki Declaration. Informed consent was obtained from all subjects.

Genomic DNA extraction and variant screening. Venous blood samples were collected from the study subjects (III:2 and III:3), their family members (II:1, II:2 and III:1) and 200 randomly selected normal individuals without any history of ophthalmic disease. In addition, amniotic fluid was collected from III:2 to reflect the condition of the fetus (IV:1). Genomic DNA from all subjects was extracted via the phenol/chloroform method. Whole-exome sequencing data were screened for pathological variants in the patients (III:2 and III:3), and sanger sequencing of the candidate variants in KERA was performed on all subjects.

The extracted DNA was fragmented with DNase and purifi ed via the magnetic bead method, followed by ligation of the adaptor sequence and polymerase chain reaction (PCR) amplification. PCR was conducted using a Phusion® High-Fidelity PCR Master Mix (New England BioLabs, Inc.) and the follows primers: 5’-AATTGATACGCGGACACCGAGATCTACACT CTTTCCTACACGACGCTTCTCCGATCT-3’ and 5’-CAA GCAGAAGACGCGATACGAGATCCGATCT-3’. PCR was performed under the following conditions: 98˚C for 30 sec; then 12 cycles of 98˚C for 10 sec, 65˚C for 30 sec and 72˚C for 30 sec; and finally 72˚C for 5 min. Library preparation was performed as previously described (25). The whole-exome was captured by NimbleSen SeqCap EZ Exome Probes v3.0 (Roche Diagnostics, Basel, Switzerland), extended, purified and finally sequenced on an Illumina HiSeq sequencer (Illumina, Inc.). The coverage of the target sequence was not less than 97% and the average depth of the target area was 130.95.

The original data were obtained following sequencing of all exons. The Burrows Wheeler Aligner algorithm was used to compare the reference sequence (UCSChg19), GATK version 3.7.0 (https://software.broadinstitute.org/gatk/) and VarScan version 2 software (http://dkoboldt.github.io/varscan/) were used to identify mutations, e.g. single nucleotide polymorphisms (SNPs) and InDels, including detection, annotation and statistical analysis. Annovar (http://www.openbioinformatics.org/annovar/) was utilized to catalog the detected variants. The list of mutations was tabulated, with the variants being described according to the HGVS guidelines. Combining this analysis with the public variation database ExAc, mutations with frequencies <0.1% in the population were identified. Mutations were scored for disease effect using a range of predictive tools [e.g. SIFT (http://sift.bii.a-star.edu.sg/), PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2), Provean (http://provean.jcvi.org/) and MutationTaster (http://www.mutationtaster.org/)]. Based on the results of the data analysis, the mutations were identified with reference to the 1000 Genomes Project (http://browser.i1000genomes.org), the Exome Variant Server (http://evs.gs.washington.edu/EVS/), dbSNP database (https://www.ncbi.nlm.nih.gov/projects/SNP/index_old.html) and the Human Gene Mutation Database (http://www.hgmd.cf.ac.uk/ac/index.php). HomoloGene (https://www.ncbi.nlm.nih.gov/homologene) and T-Coffee multiple sequence alignment (http://www.tcoffee.org) were utilized to determine the conservation of the mutated amino acid residues between different species. A three-dimensional structure of keratocan was modeled using SWISS-MODEL (https://swissmodel.expasy.org). The analysis was performed in combination with assessment of the clinical manifestations and the mode of inheritance. Finally, the nature of the mutation sites were interpreted according to the guideline by the American College of Medical Genetics and Genomics/the Association for Molecular Pathology (ACMG) (26).
The two novel candidate variations were validated using direct bidirectional sequencing methods. Based on the relevant gene standard sequences listed at the NCBI, primers were designed using Primer Premier 5 software (Premier Biosoft International). Primers forward (5'-3'): ccc taAGAGGATTTGATGTAT, primers reverse (5'-3'): GCACTGGTGGGGAACCTT. PCR amplification was performed on the target DNA fragment (700 bp product size, 20 µl reaction system, annealing temperature 62°C) and the product was verified by 1% agarose gel electrophoresis. Direct sequencing was performed on an ABI 3130 DNA Sequencer (Applied Biosystems; Thermo Fisher Scientific, Inc.). The sequenced results were aligned with the reference sequence using SeqMan at the NCBI (no. NM_007035).
Results

Clinical findings. III:2 exhibited a BCVA of 0.2 oculus dextrus (OD; right eye) and 0.3 oculus sinister (OS; left eye) with hyperopic refractive values. Additionally, she exhibited esotropia (right eye), microcornea, corneal opacity, short axial length, shallow anterior chamber, iridocorneal adhesion, pupillary irregularities (1x2 mm irregular right pupil shifting to the nasal side; needle-like changes on the left pupil) and slow light reflection. Extraocular motility, confrontation fields and intraocular pressure were normal, and the lens was transparent, although the fundus was difficult to see clearly (Table I). The younger brother of the proband (III:3) presented with similar clinical symptoms, however the parents had not identified any eye problems.

Mutation screening and bioinformatics analysis. The family of interest, originating from China, displayed an autosomal recessive pattern of inheritance. A compound heterozygous mutation c.242>A>G (p.N81S) and c.772_779delGGATTGGA(P.G258Cfs*30) in exon 2 of the KERA gene was identified in III:2 and III:3, respectively derived from the heterozygous father (c.772_779delGGATTGGA) and mother (c.242>A>G; Fig. 1C and D), although neither of the two mutations were present in the normal controls. These variants have not been previously reported in the 1000 Genomes Project, the Human Gene Mutation Database, dbSNP or the Exome Variant Server. The first variation, an N81S amino acid substitution in the Kera protein, was predicted to be likely pathogenic by the in-silico prediction programs, Mutation Taster, Provean, PolyPhen (27) and SIFT (28) (Table II). The second mutation was a 7 base-pair deletion (G258Cfs*30), leading to the loss of LRR8-11 and producing a strongly truncated protein, which may trigger nonsense-mediated mRNA decay (66 amino acid/10% missing); the prediction algorithms supported its pathogenic nature. Multiple sequence alignment indicated that the residues at position 81 of KERA are highly conserved across species (Fig. 1E) and the lost LRR8-11 regions due to 258 residual changes are also highly conserved. According to ACMG, p.N81S was judged likely to be damaging (PM1-3 and PP1-4) and P.G258Cfs*30 was also judged as damaging (PV1, PM2, PP1, PP3 and PP4) (28).

Primary sequence analysis indicated that the N81S substitution occurs at an LRR motif, the first of 11 such motifs present in the sequence (Fig. 2C) (29). G258Cfs*30 indicates a 7 base-pair deletion leading to a frameshift and premature termination codon (PTC), followed by the loss of LRR8-11. The predicted three-dimensional structure of the KERA protein using the SWISS-MODEL online software suggested that the current mutant protein (PG258Cfs*30) is significantly different from the wild type and these changes may lead to abnormal protein function (Fig. 2D) (30). The architecture of other known LRR proteins reveals that the highly conserved LXXLXXNXXL repetitive motifs form a series of parallel β-strands and these β-strands form a horseshoe-like concave shape. The asparagines (Asn, N) develop a stable Asn-ladder structure. The present study identified that the p.(Asn81Ser) mutation affects the conserved Asn81 residue of LRR1. This substitution to an aspartic acid residue will alter a hydrogen...
bond donor to make it either an acceptor or neutral, possibly resulting in the instability of the Asn-ladder structure or abnormal interactions between the mutant KERA and other proteins, including collagen. The second mutation, G258Cfs*30, that is reported in the present study results in the loss of LRR8-11. This deletion may form an abnormal truncated protein, thereby causing disease.

Discussion

With all the previously reported KERA mutations CNA2 displays a relatively homogeneous phenotype and genetic heterogeneity. Although certain previous studies have documented an atypical phenotype, including superior pellucid corneal marginal degeneration, hypotrichosis and corneal edema (2,9,11,21), they retain a classically identifiable phenotype of CNA2. Any alteration of a highly conserved encoded protein is likely to result in CNA2 with specific characteristics.

The present study reports on the phenotype and identification of suspected pathogenic variations in a Chinese family with CNA2. The proband and her brother, with two novel KERA mutations in a compound heterozygous form, exhibited the typical corneal phenotype, including marked hypermetropia, flat corneas with opacification and iridocorneal synechiae. In addition, the proband exhibited a short axis length. This finding supports the hypothesis proposed in previous studies that the KERA mutation may lead to microphthalmia, although this has not been proven (6,22). At present, there is no known effect on endothelial cell function and no specific studies have been conducted internationally. The unaffected parents in the present case were mutation carriers and did not exhibit corneal abnormalities, suggesting that in the presence of mutant gene products there remains some normal Kera proteins, which allows the cornea to develop normally.

Nearly all the conservative LRRs in the KERA gene are encoded in the larger second exon. Original descriptions of the gene mentioned 11 LRRs. A similar LRR motif exists in the structure of a porcine RNase inhibitor, the crystal structure of which has been elucidated (31). The structure of other proteins containing similar LRR motifs also mimic a β-sheet array and the KERA structural model presented in the present study supports this. These LRRs are flanked by cysteine clusters, a sulfur-containing amino acid that allows for disulfide bonds and the maintenance of the three-dimensional conformation that serves a key role in keratanan functioning.

The most reported mutations in KERA affect the LRR structure/conformation. The present study uncovered a novel missense mutation in KERA exon 2; an Asn81Ser replacement occurs within first LRR motif and disrupts it, potentially altering the stacking or spacing of the β-strand. Another novel deletion mutation, p.G258Cfs*30, in exon 2 leads to a frame shift and PTC, followed by the loss of LRR8-11. The protein is prematurely truncated due to the destruction of 7th LRR and the deficiency of the 8 to 11th LRR motifs, which contain 287 amino acids instead of 353. Conventional LRR motif alignment is believed to be important in the spacing of collagen fibrils on which corneal transparency is dependent. Aberrant KERA in the corneal stroma may not bind to collagen fibrils, therefore damaging the regulatory effect of KERA during the development of the normal corneal structure.

| Mutation at nucleotide level | Prediction | ACMG/AMP classification | Frameshift and PTC | Likely Pathogenic damaging |
|-----------------------------|------------|--------------------------|--------------------|---------------------------|
| c.242A>G                   | p.N81S     | Highly conserved          | LRRs               | Pathogenic               |
| c.772_779delGGaTTTGa       | p.G258Cfs*30 | Highly conserved          | LRRs               | Pathogenic               |

Table II. Predicted effect of 2 novel mutations on function of KERA protein.
Reviewing the literature, it was identified that CNA2 is a rare disorder with a worldwide distribution and a high prevalence in particular populations, e.g., those of Finnish (19), Saudi (3) and Czech descent (23). Almost no mutations have been identified in the Asian population, except for a Chinese American individual who was mentioned in a previous study, although this case was not documented in detail (8). In historically isolated populations, including Saudi and Finnish populations, the occurrence of CNA2 is not only due to a single common founder effect; certain cultural preferences, including intermarriage or endogamy, may also increase the incidence of recessively inherited diseases (8).

In conclusion, the present study identified two novel KERA variants in a Chinese family with cornea plana. These findings,
together with the previously reported mutations, demonstrate the occurrence of CNA2 in different populations and expand the variant spectrum of the KERA gene; therefore, the results of the present study may be valuable for the diagnosis of and genetic counseling for cornea plana.

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Availability of data and materials

All data generated and analyzed during the present study are included in this published article.

Authors’ contributions

CZH, XGL, HT collected clinical data from the CNA2 family. CZH, XGL, LQW and PSYL performed molecular genetics experiments on the family and the controls. CZH, XGL, HUTAN and CP conducted the bioinformatics analysis. CZH, CP and LQW designed and supervised the study. CZH drafted the manuscript. WGL contributed in data analysis. CP and PSYL, WGL and LQW revised the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Informed consent for this investigation was obtained from all participating CNA2 patients and parents, and the principles outlined in the Declaration of Helsinki were followed. The study was conducted in agreement with the Ethical Committee of the Center for Medical Genetics, School of Life Sciences, Central South University (Changsha, China).

Patient consent for publication

The publication of images from the CNA2 patient has the support and informed consent of the patient.

Competing interests

The authors declare that they have no competing interests.

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