

A novel missense mutation in LIM2 causing isolated autosomal dominant congenital cataract

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	A novel missense mutation in <i>LIM2</i> causing isolated autosomal dominant congenital
,	cataract
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Abstract:

Introduction: Congenital cataract is the most common cause of blindness in the world. Congenital cataracts are clinically and genetically heterogeneous and are mostly inherited in an autosomal dominant fashion. We identified the genetic cause of isolated autosomal dominant cataract in a four-generation British family and a Czech family.

Methods: Whole exome sequencing (WES) was performed on one affected member in the British family and two affected members in the Czech family.

Results: A novel missense variant c.388C>T; p.(R130C) was identified in the Lens integral membrane protein (LIM2) and found to co-segregate with disease in both families.

Conclusions: Here we report the first autosomal dominant congenital cataract variant p.(R130C) in LIM2, causing a non-syndromic pulverulent and nuclear phenotype in European families.

Introduction

Cataract accounts for nearly 50% of worldwide blindness (https://www.who.int/publicationsdetail/world-report-on-vision). Congenital cataracts are phenotypically and genotypically heterogeneous and can occur in isolation or in association with other systemic disorders. Significant progress has been made in identifying the molecular genetic basis of cataract, with 38 genes identified to date associated with isolated cataract.¹ These genes encode many different proteins including, intracellular lens proteins (crystallins), membrane gap junction proteins (connexins), cytoskeletal proteins (*BFSP1, BFSP2* and *VIM2*), transcription factors (*FOXE3, PAX6, PITX3* and *MAFA*), lanosterol synthase (*LSS*), and lens integral proteins (*MIP* and *LIM2*).

LIM2 also known as *MP19*, is the second most abundant integral membrane protein present in the ocular lens fiber cells of vertebrates,² consisted of 173-amino-acid with four transmembrane

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domains, two extracellular loops, a cytoplasmic loop, and cytoplasmic amino and carboxyl termini.³ It localizes to junction regions of the lens fiber cell membrane as well as throughout the fiber cell membrane, suggesting a role in junction communication^{4 5 6} and shown to interact with calmodulin and galectin.⁷

Mutations in *LIM2* have been associated recessive cataracts and age-related cataracts. To date only three mutations have been identified causing age-related and autosomal recessive forms of cataract. Here we report the first autosomal dominant variant in *LIM2* responsible for an isolated pulverulent and nuclear phenotype in Europeans.

Material and Methods:

Phenotyping

The British family studied was identified through the proband attending the Genetic Service at Moorfields Eye Hospital, London, UK. Local ethics committee approval was obtained and all individuals taking part gave written informed consent. Nine family members including 6 affected, 2 unaffected and one married-in, underwent full ophthalmic examination (Figure 1A). The Czech family studied consisted of a mother and child (Figure 1B). This study was managed in accordance with the Declaration of Helsinki and approved by the Ethical committee of the General University Hospital in Prague. Informed consent was obtained by all participants or their legal guardians prior to the start of the study.

Whole exome sequencing (WES) and Bioinformatics

Genomic DNA was extracted from EDTA sequestered blood samples taken with informed consent and local ethical approval using the Nucleon II DNA Extraction Kit (Scotlab Bioscience, Strathclyde, Scotland, UK). Whole exome sequencing (WES) was undertaken in one affected member (III-2) of the family (Figure 1A), and both (II-2 and III-1) affected members in the Czech family (Figure 1B). The DNA was sequenced at Macrogen Europe. Exon capture and target enrichment was performed using the SureSelectXT Human All Exon V6post, (Agilent, Santa Rosa, CA, USA). Paired-end sequencing was performed on an Illumina Hiseg 2500 high-throughput sequencer, generating mean exome coverage of 50x. Raw data in fastg format was analysed using the Phenopolis bioinformatics platform⁸. The short-read sequence data were aligned using novoalign (version 3.02.08). Variants and indels were called according to GATK best practices (joint variant calling followed by variant quality score recalibration). The variants were then annotated using the Variant Effect Predictor (VEP). Variants were then filtered to only contain public control databases Kaviar novel variants, not present in and anomAD (http://gnomad.broadinstitute.org/), in known cataract genes (https://cat-map.wustl.edu/) and predicted to be moderately or highly damaging. Bi-directional direct Sanger sequencing was performed on ABI 3730 to validate the variant p.(R130C) identified by next-generation sequencing.

Structural bioinformatics

The protein structure of LIM2 was analysed using SWISSMODEL (https://swissmodel.expasy.org/repository/uniprot/P55344) (Figure 1). The best PDB match, with a match of 17.83%, was the structure of mouse claudin-3 P134G mutant in complex with C-terminal fragment of Clostridium perfingens enterotoxin, solved with X-ray diffraction by Nakmura et al, in 2019.⁹ All structures were downloaded in PDB format and analysed using Pymol (version 1.8) locally.

Results

 All affected family members in the British family had bilateral pulverulent cataract. Individual II-2 had surgery in infancy; III-2 had aphakia and glaucoma. In the Czech family, information on the cataract type was only available for one individual (III-1), who had a nuclear cataract phenotype, and surgery in infancy. The proband (II-2) also underwent cataract surgery in infancy.

After the Phenopolis genetic variant analysis pipeline by allele frequency, in II-2 of the British family, from a total of 123,692 variants, only 403 variants remained. A list of 115 cataract-associated genes was used for gene panel screening. The variants were then sorted by CADD score. The top scoring variant for CADD (score of 35) was a rare heterozygous damaging variant

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NM_001161748.2 c.388C>T; p.(R130C) in exon 4 of *LIM2*. Direct sequencing confirmed the variant (Figure 2), which co-segregated in all the affected members of the British family (Figure 1A), and in two affected individuals from the Czech family. The R130C substitution is located in the second extracellular loop of the LIM2 protein is likely to perturb membrane trafficking and fiber cell-cell communication. It also modifies the structure of the protein compared to wild-type (RMS=0.286) (Figure 3).

Discussion

In this study we report a novel missense variant, c.388C>T; p.(R130C) in *LIM2* in a fourgeneration British pedigree with non-syndromic AD congenital pulverulent cataract, and a Czech family with a nuclear cataract phenotype. These phenotypes have also been reported in various mouse models with *Lim2* mutations. A *Lim2* missense variant, p.(G15V) has been identified in the To3 mouse and resulted in autosomal semi-dominant congenital cataracts. Heterozygous mice exhibited dense cataracts, whilst homozygous mutant mice suffered microphthalmia, lens rupture, and disorganised primary and secondary fibres¹⁰. *Lim2* homozygous knockout mice have pulverulent nuclear opacities and altered refractive properties of the lens, whereas heterozygotes had normal lenses, suggesting that loss of function of Lim2 is responsible for the phenotype.¹¹

In humans, to date only three recessive variants have been identified in *LIM2*; first in an Iraqi family p.(F105V)¹² associated with mild pulverulent cataract, with a pre-senile or senile onset. The other two, p.(G154E)¹³ and p.(G78D)¹⁴, in families of South Indian and Pakistani origin respectively (Table 1). Here we report the first autosomal dominant congenital cataract variant p.(R130C) in *LIM2*, causing a non-syndromic pulverulent and nuclear phenotype in European families, which we suggest results in aberrant modification of protein structure.

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Conflict of interests: The authors declare that they have no conflict of interest.

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Figure 1: (A) British and (B) Czech pedigrees showing a dominant inheritance pattern. Open and filled symbols indicate unaffected and affected individuals.



B] Structure of LIM2 protein

Figure 2: (A) Sequence analysis of LIM2 showing segregation of the p.(R130C) variant in an affected individual. (B) Structure of the LIM2 protein- c.388C>T; p.(R130C) variant affects the second extracellular domain of LIM2 and modifies the structure of wild-type protein (RMS = 0.286).

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Table 1: Spectrum of *LIM2* Mutations causing cataractogenesis.

Exon	HGVSc	HGVSp	Genomic	CADD	Inheritance	Ethnicity/ Origin	Phenotype	Reference
Ex3/5	c.313 T>G	p. F105V	chr19- 51885684- A>C	23.9	AR	Iraqi- Jewish	Presenile Cortical sutural	Pras et al 2002
Ex3/5	c.233G>A	p. G78D	chr19- 51885764 -C>T	31	AR	Pakistan	Nuclear	Irum et al 2016
EX5/5	c.461G>A	p. G154E	chr19- 51883516 -C>T	32	AR	South India	Congenital	Poonam et al 2008
Ex4/5	c.388C>T	p. R130C	chr19- 51883831 -G>A	35	AD	British and Czech	Pulverulent and Nuclear	Present Study

All mutations are on the transcript ENST00000596399.1 of *LIM2* - produces a 173 aa protein (Uniprot: P55344-1). CADD=Combined Annotation Dependent Depletion; AR=Autosomal Recessive; AD=Autosomal Dominant

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