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Characterization of a variant of gap junction protein α8 identified in a family with hereditary cataract

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Abstract

Purpose
Congenital cataracts occur in isolation in about 70% of cases or are associated with other abnormalities such as anterior segment dysgenesis and microphthalmia. We identified a three-generation family in the University of California San Francisco glaucoma clinic comprising three individuals with congenital cataracts and aphakic glaucoma, one of whom also had microphthalmia. The purpose of this study was to identify a possible causative mutation in this family and to investigate its pathogenesis.

Methods
We performed exome sequencing and identified a putative mutation in gap junction protein α8 (GJA8). We used PCR and DNA sequencing of GJA8 in affected and unaffected members of the pedigree to test segregation of the variant with the phenotype. We tested cellular distribution and function of the variant protein by immunofluorescence and intercellular transfer of Neurobiotin in transiently transfected HeLa cells.

Results
Exome sequencing revealed a variant in GJA8 (c.658A>G) encoding connexin50 (Cx50) that resulted in a missense change (p.N220D) in transmembrane domain 4. The variant was present in all three affected family members, but was also present in the proband’s grandfather who was reported to be unaffected. The mutant protein localized to the plasma membrane and supported intercellular Neurobiotin transfer in HeLa cells.
Conclusions

We identified a variant in transmembrane domain 4 of Cx50 in a family with autosomal dominant congenital cataracts. This variant has been previously identified in other cataract cohorts, but it is also present in unaffected individuals. Our study demonstrates that the mutant protein localized to the plasma membrane and formed functional intercellular channels. These data suggest that \( \text{GJA8 c.658A}>\text{G} \) is most likely a benign rare variant.

Introduction

Cataracts form when crystalline lenses lose transparency and become opaque. Worldwide, cataracts are a major cause of treatable visual impairment in children, occurring in 1.2–6 per 10,000 live births [1]. Hereditary congenital cataracts account for 10–25% of pediatric cataract cases [2]. In these cases, cataracts can occur in isolation or with other ocular and systemic abnormalities. Mutations in lens crystallins and gap junction proteins constitute one of the most common genetic alterations causing hereditary congenital cataract, but there is considerable genetic heterogeneity [3].

Ocular lenses are avascular structures that derive nutrients from the adjacent aqueous humor. Water and solutes enter the lens driven by pumps in the epithelium, move through extracellular spaces, and cross the cell membranes of lens fiber cells before being brought back to the surface through channels contained in gap junctions; this process allows for nutrient and metabolite exchange, maintenance of resting potentials, and lens clarity [4,5]. Gap junction proteins (i.e., connexins) oligomerize to form hexameric hemichannels that traffic to the plasma membrane. Two hemichannels from adjacent cells dock with each other to form a gap junction channel containing a central pore that allows for communication between the cells. Connexins share a similar membrane topology with four transmembrane domains, two extracellular loops, one cytoplasmic loop, and cytoplasmic N- and C-termini [6]. In human lenses, three different gap junction proteins have been identified; mutants of those found in mature lens fiber cells, connexin50 (Cx50) and connexin46 (Cx46) have been associated with congenital cataracts [6].

In this study, we identified a family with congenital cataracts associated with a variant of \( \text{GJA8} \) encoding a substitution in the fourth transmembrane domain of Cx50 (c.658A>G; p. N220D) and examined the biochemical and functional consequences of this variant.

Materials and methods

Subjects

This study was approved by the institutional review board at the University of California, San Francisco (UCSF). Written informed consent was obtained from all participants or their parents (for children under 18 years of age). Medical records were reviewed. Subjects provided peripheral venous blood or saliva samples (Oragene Discover ORG-500, DNA Genotek, Ottawa, Ontario, Canada), and genomic DNA was extracted according to the manufacturer’s instructions (DNeasy Blood & Tissue Kit, Qiagen, Valencia, CA, USA).

Genetic analysis

Exome sequencing. DNA samples were sent to the UCSF Institute for Human Genetics Genome core for library preparation, exome capture, and sequencing. Briefly, genomic DNA
was sheared using a Covaris S2 sonicator (Woburn, Massachusetts, USA) to a target size of 200–300 base pairs, and assembled into a library with TruSeq adapters containing indexes that differentiate different libraries in a capture reaction as well as a sequencing run (KAPA Library Preparation Kit, Kapa Biosystems, Wilmington, MA, USA). Libraries were pooled into a capture reaction that contained biotinylated DNA oligonucleotides ('baits') from Nimblegen, (SeqCap EZ Human Exome Library v3.0; Roche Nimblegen, Madison, WI, USA) for 72 hours. The DNA bait-DNA hybrids were pulled out of the complex mixture by incubation with streptavidin-labeled magnetic beads and captured onto a strong magnet. After washing, the targeted DNA of interest was eluted and subjected to 18 cycles of DNA amplification. The sample was then sequenced using an Illumina HiSeq2000 sequencer (Illumina, San Diego, CA, USA). The data were filtered using GeneTalk (http://www.gene-talk.de) [7]. Variants were analyzed with Polyphen-2 (http://genetics.bwh.harvard.edu/pph2/) and SIFT (Sorting Intolerant From Tolerant Web Server, version 5.1.1, database release February 3, 2015; http://sift.bii.a-star.edu.sg/) to predict pathogenicity) [8,9].

Polymerase Chain Reaction (PCR) genotyping. DNA samples were genotyped using PCR. Taq Polymerase (Thermo Scientific, Waltham, MA, USA) was used to amplify the DNA according to the manufacturer’s directions using forward (5' -AGGCACTAAGAAGTTCCGGC-3') and reverse (5' -CAACCTCGGTCAAGGGGAAA-3') primers. The thermocycler was run at 94˚C for 3 minutes, then cycled 40 times at 94˚C for 30 seconds, 55˚C for 30 seconds, and 72˚C for 60 seconds, before final extension at 72˚C for 5 minutes. The PCR product was purified with ExoSAP-IT (Affymetrix, Santa Clara, CA, USA). Then 10 ng of the PCR product was diluted in sterile water to 15 μl total volume with 8 pmol of primer (forward or reverse) for DNA sequencing (ELIM Biopharmaceuticals, Inc.; Hayward, CA, USA) in both directions.

Molecular analysis

Generation of Cx50 constructs. The coding region of the wild-type human GJA8 gene had been previously subcloned into pcDNA3.1/Hygro(+) (Life Technologies, Waltham, MA, USA) [10]. A corresponding plasmid encoding the GJA8 mutant (c.658A>G; Cx50N220D) was obtained by PCR using High Fidelity Phusion DNA polymerase (New England BioLabs, Ipswich, MA, USA) [11]. Primers were designed in opposite directions to incorporate the 658A>G mutation into the PCR product (sense: 5' -GTGATGGAGTTGGGC CACCTGGGC-3' and antisense: 5' -GTCGAGGAATAGGGACACAGAGGCCAC-3'). The coding region of the construct was fully sequenced at the University of Chicago Comprehensive Cancer Center DNA Sequencing and Genotyping Facility to ensure that PCR amplification did not introduce unwanted mutations.

Cell culture and transfection. Communication-deficient HeLa cells were grown in MEM supplemented with non-essential amino acids, 10% fetal bovine serum, 2 mM glutamine, 100 units/ml penicillin G and 10 μg/ml streptomycin sulfate. Cells at 50% confluence were transiently transfected with wild type Cx50 in pcDNA3.1/Hygro(+) or with Cx50N220D in pcDNA3.1/Hygro(+) using Lipofectin (Life Technologies, Carlsbad, CA, USA).

Immunoblotting and immunofluorescence. Immunoblotting to detect Cx50 and immunofluorescent localization of Cx50 in cultured cells were performed essentially as described previously [10,12]. Immunolocalization of Cx50 was assessed in many microscopic fields in three independent experiments. Photomicrographs were obtained using a Zeiss Axioplan 2 microscope (Carl Zeiss, Munich, Germany) equipped with a mercury lamp and a digital camera.

Intercellular transfer of gap junction tracers. Intercellular communication was assessed by microinjection of low molecular weight tracers as previously described [12]. Briefly, one
cell within a cluster was microinjected for 1 min with a solution containing 5% Lucifer yellow (charge -2; MW: 444.4) and 9% Neurobiotin (charge: +1; MW: 287.2; Vector Laboratories) using a picospritzer (model PLI-188; NikonInstruments Inc., Melville, NY). After 5 min following microinjection of the tracers, cells were fixed in 4% paraformaldehyde, subjected to immunofluorescence (using rabbit anti-Cx50 antibodies and Alexa 488-conjugated goat anti-rabbit IgG antibodies), and incubated with Cy3-streptavidin conjugate to allow detection of the Neurobiotin by fluorescence microscopy. Lucifer yellow was used to identify the injected cell, because Cx50 shows limited permeability to this dye. Incidence of coupling was quantified in clusters of Cx50-immunopositive cells as the number of injections that resulted in Neurobiotin transfer divided by the number of injections.

**Results**

We identified a three-generation Caucasian family comprising three individuals with congenital cataracts and aphakic glaucoma inherited in a pattern consistent with autosomal dominant transmission (Fig 1). The proband (III-1) was born 5 weeks prematurely and had a 14-day stay in the neonatal intensive care unit. She was noted to have partial cataracts at 4 weeks of age, which became very dense nuclear and cortical cataracts by 6 weeks of age. She underwent uncomplicated bilateral lensectomy and anterior vitrectomy at 7.5 weeks of age. She was diagnosed with aphakic glaucoma at 3 months of age and was treated with a glaucoma drainage tube in the right eye and aqueous suppressant medications in the left eye. The proband’s mother (II-2) was diagnosed with bilateral cataracts at a few months of age. She had bilateral cataract extraction at age 5 months and subsequently developed aphakic glaucoma in both eyes. The proband’s maternal aunt (II-3) had bilateral congenital cataracts and microphthalmia. She developed aphakic glaucoma in both eyes following cataract extraction, requiring multiple surgical interventions. The family’s medical history was otherwise unremarkable.

To investigate the causative mutation in this family, we first performed exome sequencing on DNA from subject II-2, which identified 66,151 variants. We filtered the results to allow only certain protein consequences (nonsynonymous, insertion/deletion, splice site affecting), leaving 12,118 variants. We then limited allele frequencies to <1% compared to public databases (1000 Genomes, Exome Sequencing Project, Exome Aggregation Consortium), resulting

![Pedigree of the family with autosomal dominant congenital cataract](https://doi.org/10.1371/journal.pone.0183438.g001)
in 1,223 variants. We searched for mutations within candidate genes (Table 1) that have been associated with autosomal dominant cataract in multiple studies. Only one of the remaining variants was present within a candidate gene: GJA8.c.658A>G, which leads to a missense mutation p.N220D within the fourth transmembrane domain. When we tested for segregation of this variant with the phenotype we found that it was present in each of the individuals with cataracts, but it was also found in the proband’s maternal grandfather (subject I-1) (Fig 2). This individual was reported by the family to have had no congenital cataracts; however, he was not available for examination. The GJA8 c.658A>G variant has a population frequency of 0.005 (1000 Genomes, Exome Sequencing Project). The asparagine residue at position 220 is highly conserved across species and isoforms (Fig 3) and is predicted to be pathogenic by SIFT (score 0.01 out of 1.00, “damaging”) and Polyphen-2 (score 0.999 out of 1.00, “probably damaging”).

We then examined the consequences of this variant using communication-deficient HeLa cells transiently transfected with wild type GJA8 (encoding wild type Cx50) or with the c.658A>G allele (encoding Cx50 N220D). Immunoblots of homogenates from HeLa cells transiently transfected with wild type Cx50 or Cx50 N220D showed an immunoreactive Cx50 band of similar electrophoretic mobility (Fig 4A). To test the ability of the mutant protein to form gap junction plaques, we examined the cellular distribution of the proteins by immunofluorescence. Cells transfected with wild type Cx50 and Cx50 N220D had substantial labeling at

| Gene   | Name                                      | Exons (#) |
|--------|-------------------------------------------|-----------|
| AQP0/MIP | aquaporin/major intrinsic protein of lens fiber | 4         |
| BFSP2 | beaded filament structural protein 2 | 7         |
| CHMP4B | charged multivesicular body protein 4b | 5         |
| CRYAA | crystallin, alpha A | 3         |
| CRYAB | crystallin, alpha B | 3         |
| CRYBA1 | crystallin, beta A1 | 6         |
| CRYBA2 | crystallin, beta A2 | 4         |
| CRYBA3 | crystallin, beta A3 | 6         |
| CRYBA4 | crystallin, beta A4 | 6         |
| CRYBB1 | crystallin, beta B1 | 6         |
| CRYBB2 | crystallin, beta B2 | 6         |
| CRYBB3 | crystallin, beta B3 | 6         |
| CRYGB  | crystallin, gamma B | 3         |
| CRYGC  | crystallin, gamma C | 3         |
| CRYGD  | crystallin, gamma D | 3         |
| CRYGS  | crystallin, gamma S | 3         |
| Cx46/GJA3 | connexin 46/gap junction alpha-3 | 2         |
| Cx50/GJA8 | connexin 50/gap junction alpha-8 | 2         |
| EYA1  | eyes absent homolog 1 (Drosophila) | 18        |
| FOXE3  | forkhead box E3 | 1         |
| FTL   | ferritin, light polypeptide | 4         |
| HSF4  | heat shock transcription factor 4 | 15        |
| MAF   | v-maf avian musculoaponeurotic fibrosarcoma oncogene homolog | 1         |
| PAX6  | paired box gene 6 | 15        |
| PITX3 | pituitary homeobox 3 | 4         |
| WFS1  | Wolframin | 8         |

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appositional membranes (Fig 4B) indicating successful trafficking. To test the ability of Cx50\textsuperscript{N220D} to form functional intercellular channels, we microinjected cells after transfection with the gap junction permeant tracer, Neurobiotin. Both wild type Cx50 and the Cx50\textsuperscript{N220D} variant transferred Neurobiotin to their neighboring cells (Fig 4C); the incidence of coupling in wild type Cx50-transfected cells was 9/9 and in Cx50\textsuperscript{N220D}-transfected cells was 10/11.

**Discussion**

We identified a variant of GJA8 (c.658A>G) in a family with autosomal dominant cataract. This variant results in the substitution of a polar, uncharged asparagine at amino acid position 220 for a negatively charged aspartic acid (N220D). Asparagine at this position in Cx50 is highly conserved across species and across other connexin isoforms. The substitution is predicted to be probably damaging based on bioinformatic algorithms. However, based upon
genetic evidence, the pathogenicity of this variant is equivocal. It was first identified in a subject from a family with congenital cataracts, but was also found in 1 out of 340 alleles in the control population (allelic frequency: 0.003), which led to the conclusion that it was a non-pathogenic polymorphism [13]. The mutation was also found in a subject with congenital cataract and microcornea and in his unaffected father, but not in his affected sibling, indicating that it did not segregate with the disease in the family [14]. In our pedigree, the mutation was present in all affected subjects, but it was also found in the proband’s maternal grandfather who was reported to be unaffected. These observations suggest that the mutation is either an incidental finding or that it is pathogenic with reduced penetrance and variable expressivity.

Fig 3. Protein sequence alignment. (A) Orthologous sequences of Cx50 from human, mouse, rat, chick and sheep show that amino acid N220 is highly conserved across species. (B) Alignment of other human gap junction protein sequences also demonstrated conservation of this residue among isoforms.

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Fig 4. Cx50N220D production, localization and function in transfected cells. (A) Proteins from homogenates of HeLa cells transiently transfected with wild-type Cx50 or Cx50N220D were resolved by SDS-PAGE and subjected to immunoblotting using rabbit polyclonal anti-human Cx50 antibodies. Samples from cells expressing either Cx50 or Cx50N220D contain a single immunoreactive band of similar mobility. Molecular mass standards are indicated on the left. (B) Photomicrographs show the distribution of immunoreactivity to anti-Cx50 antibodies in HeLa cells transfected with wild-type Cx50 (WT) or Cx50N220D (N220D). Cells expressing WT and N220D both show substantial labeling at appositional interfaces between cells consistent with gap junction plaques. (C) Photomicrograph showing intercellular neurobiotin transfer from a microinjected cell (*) to neighboring cells in HeLa cells transiently with CX50N220D. Bar, 16 μm for B and 10 μm for C.

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Environmental factors, stochastic events, epigenetic changes or differences in genetic background (or modifiers) can all contribute to differential effects of a mutation. Notably, in the family reported here all affected members are female, however, in a previous study the affected child and the unaffected father that carried the mutation were male [14].

Ours is the first study to characterize the consequences of the N220D variant on the biochemical and functional properties of Cx50. The most common pathogenic mechanism found in other cataract-associated Cx50 mutants is failure to form gap junctions due to impaired protein folding and/or localization to the plasma membrane [6,15,16]. Unlike most pathogenic mutations, Cx50\textsuperscript{N220D} localized to appositional plasma membranes and formed functional gap junction plaques between adjacent cells. These results demonstrate that this variant does not abolish intercellular channel function (at least not in transfected HeLa cells). Variations of cell density within the same tissue culture dish and of protein expression in transient transfection experiments preclude us from estimating whether the extent of transfer differs between Cx50 and Cx50\textsuperscript{N220D}. The ability of Cx50\textsuperscript{N220D} to support intercellular communication may explain the lack of segregation of the variant with the cataract phenotype in the different families. The results suggest that other factors might contribute to the pathology.

Although Cx50\textsuperscript{N220D} is the only variant in the fourth transmembrane domain of Cx50 identified to date, mutations in the corresponding region of other connexins have been associated with diseases, including autosomal dominant congenital cataract (Cx46\textsuperscript{F206I}) and autosomal dominant and recessive hearing loss (Cx26\textsuperscript{C202F} and Cx26\textsuperscript{N206S}, respectively). Interestingly, the N220D substitution in Cx50 occurs in a homologous position to N206S in Cx26, however, the functional consequences appear to be different. This might relate to the differential permeability of the intercellular channels formed by Cx50 and Cx26. Cx50 channels are permeable to Neurobiotin, but show limited intercellular transfer of Lucifer yellow whereas Cx26 channels are permeable to both cationic and anionic molecules with a cationic preference. This implies that the amino acid residues determining permeation must differ between Cx26 and Cx50 [17–21]. N206 is involved in critical interactions within the Cx26 channel; the crystal structure of Cx26 shows that N206 forms a hydrogen bond with R143, located on the third transmembrane domain [22]. Mutations in this residue are predicted to disrupt this interaction; indeed, Cx26\textsuperscript{N206S} gap junction channels show altered voltage gating properties and decreased permeability to ethidium (charge +1, MW: 314.39), but preserve permeability to anionic Lucifer yellow [18,20]. In Cx50\textsuperscript{N220D}, however, we did not find a difference in intracellular transfer of Neurobiotin compared to wild-type Cx50.

In summary, we identified and characterized a variant occurring in transmembrane domain 4 of Cx50 in a family with autosomal dominant congenital cataracts. Functional studies demonstrated that Cx50\textsuperscript{N220D} forms gap junctions that allow the passage of small, cationic molecules, similar to wild-type Cx50, supporting that this is a rare, benign variant rather than the causative mutation in this family.

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