δ-Catenin Is Genetically and Biologically Associated with Cortical Cataract and Future Alzheimer-Related Structural and Functional Brain Changes

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Abstract

Multiple lines of evidence suggest that specific subtypes of age-related cataract (ARC) and Alzheimer disease (AD) are related etiologically. To identify shared genetic factors for ARC and AD, we estimated co-heritability of quantitative measures of cataract subtypes with AD-related brain MRI traits among 1,249 members of the Framingham Eye Study who had a brain MRI scan approximately ten years after the eye exam. Cortical cataract (CC) was found to be co-heritable with future development of AD and with several MRI traits, especially temporal horn volume (THV, r = 0.24, P < 10⁻⁴). A genome-wide association study using 187,657 single nucleotide polymorphisms (SNPs) for the bivariate outcome of CC and THV identified genome-wide significant association with CTNND2 SNPs rs17183619, rs13155993 and rs13170756 (P < 2.6 x 10⁻⁷). These SNPs were also significantly associated with bivariate outcomes of CC and scores on several highly heritable neuropsychological tests (5.7 x 10⁻⁶ ≤ P < 3.7 x 10⁻⁶). Statistical interaction was demonstrated between rs17183619 and APP SNP rs2096488 on CC (P = 0.0015) and CC-THV (P = 0.038). A rare CTNND2 missense mutation (G810R) 249 base pairs from rs17183619 altered δ-catenin localization and increased secreted amyloid-β₁₋₄₂ in neuronal cell culture. Immunohisto-pathological analysis of lens tissue obtained from two autopsy-confirmed AD subjects and two non-AD controls revealed elevated expression of δ-catenin in epithelial and cortical regions of lenses from AD subjects compared to controls. Our findings suggest that genetic variation in delta catenin may underlie both cortical lens opacities in mid-life and subsequent MRI and cognitive changes that presage the development of AD.

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Introduction

Alzheimer disease (AD) and age-related cataract (ARC) are common age-related disorders. Approximately 5.4 million Americans have AD including 13% of people ages 65 and older and nearly 43% of people ages 65 and older [1]. An estimated 20.5 million persons aged 40 years and older in the U.S. show some evidence of ARC [2]. Both AD and ARC are highly heritable [3,4]. Numerous rare and common genetic variants have been robustly associated with AD [3,5–8], and to a lesser extent with ARC [9], but much of the genetic risk of these disorders is still unexplained.

Numerous lines of evidence suggest common factors linking AD-associated pathology in the brain and lens. Comparing aged controls with AD patients, Goldstein et al. observed amyloid-β
Methods

Subjects

The Framingham Study is a multi-generation community based study that began as a study of cardiovascular disease in 1948 among 5209 individuals comprising the Original cohort [24]. The Framingham Offspring Study includes 5216 spouses and offspring of the Original cohort members who have been followed since 1971 [25]. Surviving members of this cohort were examined and classified for lens opacities during exams conducted between May 1989 and December 1991 as part of the Framingham Offspring Eye Study (FOES) [26]. Starting in 1999, dementia-free surviving members of both Original and Offspring cohorts were invited to have a magnetic resonance imaging (MRI) examination of the brain. Repeat measurement data were obtained from a second MRI scan performed 2.6 years and 6.2 years later on average in the Original and Offspring cohorts, respectively. For the molecular genetic analysis, we focused on 1249 members of the Offspring cohort who were examined in the 1989 and 1991 exams. CTNND2 Link to Cataract & Alzheimer Brain Changes

Phenotypic Evaluation

Ophthalmic examinations in the FOES were performed by two experienced certified ophthalmologists who evaluated the slit lamp photographs obtained through a dilated pupil using a standardized grading system [27]. Both eyes were examined in all study subjects. Nuclear opacification was classified by comparison to a set of standard photographs (grades 0 to 3) according to a validated ordinal rating system [28]. Cortical opacification was measured using a grid to assess opaque area in one-eighth wedges from the circumference of the lens cortex. An ordinal rating of cortical cataract was assigned based on the total number of one-eighth wedges occupied by cortical cataract. Persons with 2+ positive wedges were assigned a rating of 4 (Fig. S2). Posterior subcapsular cataract (PSC) was assessed by measuring the vertical width of posterior opacification recorded by slit lamp photography (range: 0 to 7 mm). We selected the worse measurement from the right and left eyes. Thirty-three subjects with missing ophthalmic examination data or with a history of cataract surgery were excluded from the analysis.

Details of the MRI scan protocol and procedures for deriving structural volumes and measures of degeneration have been described previously [29]. In brief, volumetric brain MRI measures were obtained using analysis of digital images from the MRI scans. Participants were imaged on a Siemens 1 or 1.5-T MR machine (Siemens Medical, Erlangen, Germany). Digital images were processed by a central laboratory blinded to demographic and clinical identifiers. Quantified MRI measures were derived for frontal lobar volume (FBV), occipital brain volume (OBV), parietal brain volume (PV), temporal brain volume (TV), hippocampal volume (HPV), lateral ventricular volume (LVV), temporal horn volume (THV), and white matter hyperintensity volume (WMHV). HPV information was unavailable for 43 (2%) of the subjects at the time of the study.

Subjects were administered a neuropsychological test battery using standard protocols and trained examiners. Details of the tests administered and normative values for the Framingham Original and Offspring cohorts have been published [30,31]. Cognitive test data were selected from the exam at the time the baseline MRI scan was performed.

Co-heritability Estimation

Heritability (h²) in the narrow sense for a quantitative trait is the ratio of variance contributed by the additive genetic effects to the total phenotypic variance and can be estimated from family data by analysis of intraclast correlations (r). Heritability based on analysis of data obtained among siblings is calculated using the formula: h² = 2r. Co-heritability is the ratio of analogous covariance components between two traits [32]. It can be estimated from analysis of cross-trait sibling correlations. A cross-trait sibling correlation is the average of the correlation of the first trait in sibling A with the second trait in sibling B and the correlation of the second trait in sibling A with the first trait in sibling B. By contrast, self correlation is the correlation between two traits within individuals. The design for estimating self and cross-trait sibling correlations is illustrated in Fig. S3.

Self and cross-trait sibling correlations of volumetric brain MRI and cataract phenotypes with AD were estimated using the FCOR program in the Statistical Analysis for Genetic Epidemiology (S.A.G.E) software (version 6.0) [33]. FCOR calculates correlations for dichotomous and quantitative traits in extended families using the Pearson product-moment estimator with generalized...
weights. In addition, FCOR estimates multivariate familial correlations with their asymptotic standard errors without assuming multivariate normality of the traits by using a second-order Taylor series expansion and replacing all correlation parameters with their respective estimates [33,34]. Raw measurements showing significant self and cross-trait correlations were further analyzed by adjusting for sex and age at exam. Age at blood draw (for DNA extraction) was substituted for age at eye exam because this information was not available for a large proportion of subjects at the time the data were analyzed. This was deemed reasonable since the age- and sex-adjusted residuals for age at blood draw and age at eye exam occurred during the same Framingham Study exam cycle. Correlations were also computed for the annual rate of change in the selected MRI traits which was calculated as the difference in the measures between the baseline and second MRI evaluations divided by the number of years between examinations. Because the distributions of many traits were multi-modal, age and sex-adjusted residuals for each trait were derived and normalized, forcing the marginal distribution of the trait to be approximately normally distributed under the null hypothesis, by taking the inverse standard normal transformation of the empirical quantile that was obtained using the formula \( r/yj - 1/3 \) divided by \( n + 1/3 \), where \( r/yj \) is the rank of residuals, \( y \) [20]. This approach has been shown to be valid in family-based association tests even with rare variants [20]. Normalized residuals were used for subsequent analyses. The significance threshold accounting for 38 independent tests was set at \( P = 0.0013 \).

**Genotyping and Data Cleaning**

All available participants were genotyped at Affymetrix (Santa Clara, CA) using the Affymetrix GeneChip® Human Mapping 500K Array Set and 50K Human Gene Focused Panel®. Phenotype and genome-wide association study (GWAS) data including raw genotyping calls and genotypes imputed using the HapMap 2 and 3 reference populations were obtained from the dbGaP website (http://www.ncbi.nlm.nih.gov/gap. Accessed 2012 July 2).

Prior to analysis, SNPs with a call rate less than 98%, SNPs with a minor allele frequency (MAF) less than 2%, and SNPs not in Hardy-Weinberg equilibrium \((P < 10^{-8})\) were excluded. Individuals with SNP call rates below 98% among the remaining SNPs, or whose gender as determined by analysis of X-chromosome data using PLINK [35] was inconsistent with the self-reported gender were also excluded. The 354,310 SNPs remaining among 7,966 subjects were then pruned to remove one from each pair of SNPs in high linkage disequilibrium \((r^2 > 0.8)\) using PLINK with a window size of 1500 SNPs and a step size of 150. After applying these criteria, 187,657 genotyped autosomal SNPs remained. Familial relationships were verified by examining segregation of marker genotypes within families using the MARKERINFO program in S.A.G.E. After confirming these relationships, marker genotypes showing Mendelian inconsistency were set to missing.

Population substructure was examined among those with self-reported ethnicity of white or Caucasian, first among the complete sample and again within a set of unrelated individuals consisting of one randomly selected individual from each extended pedigree. The complete sample of individuals was used for investigation of population structure using the pruned and cleaned genotypes from the Affymetrix 500K array and the CEU HapMap phase 2 reference data. SNP loadings were prepared using the smartpca script implemented in the EIGENSTRAT package [36]. This initial run was performed to confirm the presence of a single ethnicity-specific cluster delineated based on self-declared ethnicity. For evaluation of within-cluster substructure, a set of SNP loadings were prepared using the pruned and cleaned genotypes from the unrelated sample of individuals with the same smartpca script implemented in the EIGENSTRAT package. The SNP weights for each eigenvector from the unrelated sample were then applied to all remaining family members to compute principal components for all individuals in the sample. We did not find any significant association between principal components and any cataract or MRI phenotypes in univariate analyses.

**GWAS Analysis Methods**

The SNP pruning strategy described above was employed to reduce computational burden associated with bivariate analysis in extended families. SNPs were coded under an additive model as 0, 1, or 2 with respect to the number of the reference alleles. The coded SNP values were included as covariates in regression models along with variables for polygenic effects and random effects to account for the various constellations of relative pairs. Association of each SNP with a bivariate outcome comprising age and sex-adjusted normalized residuals of quantitative measures for one cataract trait \((yc)\) and one MRI trait \((ymri)\) was evaluated using the bivariate extension of the two-level Haseman-Elston (tHE) regression method developed for general pedigree data implemented in the RELPAL program in S.A.G.E [37,38]. The regression model is denoted as \( ycc\text{mri}ik = b\text{X} + b\text{Z} + e\text{k} \) for individual \( i \) in pedigree \( k \), where \( b\text{X} \) and \( b\text{Z} \) represent the SNP, polygenic and random effects, respectively; and \( b\text{X} \) and \( b\text{Z} \) are coefficients. This approach uses an iterative generalized squares algorithm which can accommodate various family structures and incorporate both individual-level and pedigree-level covariates. We demonstrated previously by simulation that this regression model has high power (>99%) for a SNP with a large effect size \((\beta > 1.5)\), even with very rare variant \((\text{MAF} < 0.001)\), and produces few false positives in large extended families [39]. Some of the pedigrees in the Framingham Study have similar pedigree structures to those included in the simulations.

Nominal \( P \) values for association were determined using first-level Wald tests computed by RELPAL. The most significant results (i.e., \( \text{SNPs with} \ P < 10^{-5} \)) in the genome-wide scan were investigated further using data for all known SNPs in the implicated genes including those that were genotyped but had been dropped due to pruning. Additional SNPs in the HapMap 2 and 3 reference panels were imputed using MaCH [40]. All SNPs within 100 kb of top-ranked SNPs not located within genes were analyzed. A quantitative estimate between 0 and 2 representing the dose of the minor allele was used in the analysis instead of imputed genotypes to incorporate the uncertainty of the imputation estimates. We excluded imputed SNPs with low MAF (<2%), not in Hardy-Weinberg equilibrium \((P < 10^{-8})\) and low imputation quality \((\text{RSQ} < 0.8)\). A total of 186,192 genotyped SNPs and 1,465 imputed SNPs for a selected region were analyzed for association. Based on this number of SNPs, the threshold for genome wide significance was determined to be 2.66 \times 10^{-7}.

The possibility of a functional role of genes identified by bivariate GWAS in lens opacity and neurodegeneration was evaluated statistically by testing models including the genome wide significant SNP, a SNP from \( \text{APP} \) or \( \text{PSEN1} \), and an interaction term. The two most significant \( \text{APP} \) genotyped and uncorrelated SNPs (pairwise LD < 0.5), each with MAF > 0.1, from the bivariate GWAS were tested for interaction with the top \( \text{CTNND2} \) SNP rs17183619. No \( \text{PSEN1} \) SNPs met these criteria and hence were not tested for interaction. The proportion of genetic variance in
the bivariate trait explained by rs17183619 was estimated using methods proposed by So and colleagues [41,42].

Functional Analysis of the CTNND2 G810R Mutation

Cell Culture. Human Embryonic Kidney 293 cells stably expressing APP Swedish mutation (K595N/M596L of APP695, HEK293-APPsw) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 500 μg/ml Geneticin (G418, Invitrogen) and maintained at 37°C in a humidified atmosphere (5% CO2/95% air).

Transfection, plasmids and cloning. HEK293-APPsw cells were transfected using Lipofectamine™ 2000 reagent (Invitrogen) according to the manufacturer’s protocol. Briefly, plasmid DNA was pre mixed with Lipofectamine (2 μg; 5 μL ratio) in reduced serum media (Opti-MEM®, Invitrogen). DNA-Lipofectamine complexes were added to 80% confluent cells in growth medium without antibiotics for five hours. After incubation, medium was replaced with fresh medium. All assays were performed 24 hours after transfection. pCDNA3 empty vector and pCDNA3His encoding a wild-type human CTNND2 cDNA were from our plasmid library. Mutant CTNND2 (glycine to arginine substitution at amino acid position 810) was PCR engineered using pCDNA3His/CTNND2 as a template and primer sets designed to introduce a missense mutation at codon 810.

Western blot. Cell monolayers were lysed in STE buffer (50 mM Tris [pH 7.6], 150 mM NaCl, 2 mM EDTA, 0.2% NP-40 and 0.5% Triton) supplemented with Complex™ protease inhibitor cocktail (Roche). Lysates were pre-cleared of insoluble material by centrifugation and total protein was quantified using BCA reagent (Pierce). Proteins were boiled at 95°C for 5 minutes in Laemmli buffer, separated by molecular weight in 4–12% NuPAGE® Bis-Tris gels (Invitrogen) and transferred to nitrocellulose membranes (Millipore). The membranes were probed for 1 hour with rabbit anti-CTNND2 (Abcam; 1:5,000) or mouse anti-tubulin (Santa Cruz; 1:2,000), washed in Tris Buffered Saline with Tween and re-probed with donkey anti-mouse or donkey anti-rabbit antibodies conjugated to horseradish peroxidase (Santa Cruz; 10,000) for one hour. The proteins were visualized using an ECL reagent (GE).

ELISA. The levels of secreted amyloid β (Aβ) peptide were determined by using ELISA Aβ40 and Aβ42 kits (Invitrogen). Briefly, the culture medium was collected with serum protease inhibitor AEBSF (Roche) and immediately assayed or kept at −20°C until tested. Incubation time with Aβ detection antibody was 3 hours. All other steps were performed as described in the manufacturer’s protocol. Aβ levels were normalized to total protein and then normalized to the pCDNA3 vector control. Three independent experiments were performed for each condition tested (empty vector, delta-catenin, G810R). Differences in Aβ levels between conditions were evaluated using a two-tailed Student’s t test.

Microscopy. HEK293-APPsw cells grown on glass coverslips were fixed in 2% paraformaldehyde for 15 minutes, permeabilized in 0.1% saponin for 20 minutes and incubated with rabbit anti-NPRAP antibody (Abcam; 1:1,000 in 2% BSA and 0.1% saponin in PBS) for 1 hour. Cells were then incubated for another hour with Alexa Fluor 488-conjugated goat anti-rabbit antibody (Invitrogen; 1:250 in 0.1% saponin in PBS) followed by 15 minutes with DAPI (Sigma; 100 ng/ml). Cells were washed twice in PBS between every step. Coverslips were mounted on slides using Vectashield® mounting medium (Vectorlabs) and observed using an epifluorescence microscope. Semi quantitative analysis of protrusions was performed manually. Images were captured with the 40× objective based on green fluorescence immunoreactivity. Spine-like, filopodia-like and other protrusions on apical segments (branches) of HEK293-APPsw cells were identified based on standard morphological criteria and quantified in areas of equivalent size and magnification. All experiments were performed blinded to the identity of transfected constructs and repeated three times per condition tested (empty vector, δ-catenin or G810R). P values were obtained by two-tailed Student’s t test.

Immunohistochemical Analysis

Eyes from a 68-year-old male and 71-year-old female with neuropathologically-confirmed AD and from two normal male controls ages 68 and 70 were procured from the Boston University Alzheimer Disease Center and National Disease Research Interchange (Philadelphia, PA). Eyes were fixed in 4% paraformaldehyde, embedded in paraffin, and sectioned at 8 μm. Sections underwent antigen retrieval using citrate buffer pH 6.0 diluted to the manufacturer’s instructions and heated to 90°C for 20 minutes. Tissue sections were immunostained with δ-catenin antibody (SC-81793, Santa Cruz Biotechnology, Inc., Santa Cruz, CA) at 1:100 dilution and processed by conventional immunohistochemistry (LabVision Autostainer 360, Fremont, CA). Negative controls were run for each antibody. Tissue sections were visualized by brightfield photomicroscopy.

Results

Cortical Cataract is Predictive of Future Abnormal Brain Changes

To identify quantitative measures of brain damage that are the best surrogate markers for presymptomatic AD and associated with lens opacity, we examined correlations among these traits within the same individual and between siblings (Table S1, Fig. S3). Although most of the MRI traits were significantly correlated with AD among all Framingham Study subjects who had at least one MRI exam, only TBV, LVV, and THV were significantly co-heritable with future AD after correction for multiple testing (Table S2). Lens opacity information was available for 1249 individuals who participated in the Framingham Eye Offspring Study (FEOS), had at least one brain MRI exam and were on average 51.3 years old at the time of the ophthalmic examination (Table S1, Fig. S1). At the time of the first MRI exam, none of these subjects were demented and only nine (0.7%) had a previous stroke. The mean interval between the eye exam and the first MRI exam was 9.3 years. Our analysis also revealed a significant sib-sib correlation of AD with cortical cataract (CC, ρ=0.12, P<10⁻⁴) and posterior subcapsular cataract (PSC, ρ=0.11, P<10⁻⁴), but not with nuclear cataract (NC) (Table S2). Heritability and co-heritability estimates were much greater for CC and several MRI traits, particularly THV and LVV, after adjustment for age and sex, and variable normalization (Table 1, Table S3). Taken together, these results suggest that CC and several AD-related brain changes have shared genetic liability or are associated with another variable under genetic influence, and that development of CC may foreshadow development of AD-related brain changes in later life.

Genome-Wide Association

Since CC was much more strongly co-heritable with THV than LVV (Table 1), a genome-wide search was conducted for association of common variants with the bivariate outcome of CC and THV. For this analysis, we used the set of 186,192 genotyped SNPs that remained after applying quality control procedures and a pairwise linkage disequilibrium (LD) cutoff of 0.8 (see Methods). The quantile-quantile (QQ) plot showed that the
distribution of P values for the majority of SNPs (i.e., \(P > 10^{-5}\)) met expectation under the null hypothesis (genomic inflation factor \(\lambda = 1.036\), whereas more SNPs than expected under the null had \(P\) values \(< 10^{-5}\) suggesting several true associations within this set of SNPs (Fig. S4). While none of the genotyped SNPs were genome-wide significant (threshold \(P = 2.7 \times 10^{-8}\)), suggestive association \((P < 5 \times 10^{-8}\) was observed for CC-THV with three SNPs in CTNND2 (Table S4). This region was further examined by testing association of CC-THV with 1,465 accurately imputed results are supported by strong associations \((P = 0.0015)\), but not for THV.

interaction revealed evidence of a synergistic effect on the bivariate CTNND2 variant in with a decrease in both CC and THV. These results suggest that a indicative that the minor allele (G) of rs17183619 is associated value sign) was consistent in univariate and bivariate models considered as separate outcomes. The effect direction (indicated by \(\beta\) value sign) is consistent in univariate and bivariate models indicating that the minor allele (G) of rs17183619 is associated with a decrease in both CC and THV. These results suggest that a variant in CTNND2 influences a process that is more precisely correlated with by degeneration in both the lens and brain than in either tissue considered separately. The proportion of the total co-correlated with by degeneration in both the lens and brain than in either tissue considered separately. The proportion of the total co-

## Table 1. Correlations of cortical cataract with temporal horn volume and lateral ventricular volume.

| MRI Trait^1 | Cross-Trait Correlations with Cortical Cataract^1 | Within an Individual | Between Siblings^1 |
|-------------|--------------------------------------------------|----------------------|-------------------|
|             | Subjects | COR | \(P\) | Subjects | COR | \(P\) |
| Temporal Horn Volume (THV) | 1249 | 0.318 | \(<10^{-4}\) | 668 | 0.239 | \(<10^{-4}\) |
| Lateral Ventricular Volume (LVV) | 1249 | 0.127 | \(1 \times 10^{-4}\) | 668 | 0.153 | \(5 \times 10^{-4}\) |
| Annual Change of THV | 880 | -0.128 | 0.001 | 497 | -0.071 | 0.116 |
| Annual Change of LVV | 880 | -0.148 | \(2 \times 10^{-4}\) | 497 | -0.095 | 0.043 |

^1 Trait values transformed to normalized residuals.

^2 Adjusted for age and sex.

^3 Calculated by averaging the correlations of cortical cataract in sib 1 with MRI trait in sib 2, and cortical cataract in sib 2 with MRI trait in sib 1.

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A CTNND2 Functional Variant Alters Amyloid-\(\beta\) (A\(\beta\)) Expression

The top-ranked CTNND2 SNPs \((P < 10^{-5}\) are located in the region encompassing exons 14–16, and the three most significant SNPs flank exon 14 (Fig. 1C). Bioinformatic analysis of this region revealed that the top-ranked SNP, rs17183619, is 249 base pairs upstream of a rare non-synonymous coding SNP (rs1754599) that results in a glycine to arginine missense substitution at residue 810. This missense mutation is predicted to have a deleterious effect on the protein structure (Fig. 2A). The sequence surrounding G810R is highly conserved across several species and bears a signature stretch of basic residues which are required for nuclear function of \(\delta\)-catenin, the protein encoded by CTNND2 (Fig. 2B).

Transfection of the human CTNND2 cDNA with the G810R mutation into HEK293 cells that stably express the human APP Swedish mutation \((APP_{sw}) showed that the distribution of mutant \(\delta\)-catenin is altered predominantly at or near the cell surface (Fig. 3A). Transfected cells expressing the G810R mutation had a significantly more elaborate network of protrusions \((P = 0.006, Fig. 3B)\, suggesting that the mutated CTNND2 gene may alter interactions of the plasma membrane with the underlying actin cytoskeleton. Moreover, transfection with the mutant \(\delta\)-catenin resulted in a significant and specific increase in \(A\beta_{1-42}\) secretion \((P = 0.02, Fig. 3C)\.

\(\delta\)-Catenin is Expressed in Human Lens

We detected \(\delta\)-catenin immunoreactivity in the central and bow regions of the lens epithelium and underlying anterior, subequatorial, and deep supranuclear cortical regions of the lens (Fig. 4) and retina (Fig. S5). The tissue localization pattern in postmortem lens from subjects with neuropathologically confirmed AD, but not lens from non-AD controls, was notable for abnormally increased \(\delta\)-catenin immunoreactivity in the lens epithelium and cortex regions of the lens examined. Moreover, we detected pathological accumulation of \(\delta\)-catenin immunoreactive product in AD lenses that was detectable as compacted cytosolic deposits with a striking basolaminar preponderance in the anterior and equatorial epithelium and as heterogeneously distributed punctate deposits
in the superficial, subequatorial, deep cortical and supranuclear lens subregions. The presence of these pathological $\beta$-catenin immunoreactive accumulations in AD lenses comprises a presumptive biological substrate for locally increased light scattering, lenticular opacification, and frank cataract.

Discussion

We observed in a sample of Framingham Study participants a correlation of a cortical cataract score measured during adulthood with future development of AD, and with multiple measures of AD-related brain degeneration obtained from both MRI scan and

**Table 2.** Interaction of CTNND2 SNP rs17183619 and APP SNP rs2096488 in univariate and bivariate models of cortical cataract (CC) and temporal horn volume (THV).

| SNP (Reference Allele) | Cortical Cataract | Temporal Horn Volume | Bivariate |
|------------------------|-------------------|----------------------|-----------|
|                        | $\beta$          | P        | $\beta$ | P   | $\beta_{cc}$ | $\beta_{THV}$ | P         |
| rs17183619 (G)         | −0.266           | 4.0 $\times$ 10^{-5} | −0.123 | 0.0593 | −0.411    | −0.193     | 1.8 $\times$ 10^{-7} |
| rs2096488 (A)          | 0.046            | 0.33     | 0.113   | 0.0176 | 0.049     | 0.093      | 0.19      |
| Interaction            | −0.372           | 0.0015   | 0.007   | 0.95   | −0.227    | 0.18       | 0.038     |

Effect estimates ($\beta$) are based on the dosage of the rs17183619 minor allele (G) which has a frequency of 0.153.

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cognitive GWAS identified genome-wide significant association of three intronic SNPs in CTNND2 with a bivariate outcome of cortical cataract and volume of the temporal horn, an MRI measurement that strongly correlates with AD progression [43]. This association accounts for 19% of the heritable component of brain changes. Our analyses also revealed a significant interaction of the top-ranked CTNND2 SNP (rs17183619) with APP SNP rs2096488 on the degree of cortical cataract and performance on several neuropsychological tests including the BNT, LMI, LMD and TMTB which are strongly associated with future AD risk [44–46] evaluated at the same time as the MRI scan, suggesting that variants in CTNND2 may influence functional as well as structural brain changes. Our analyses also revealed a significant interaction of the top-ranked CTNND2 SNP (rs17183619) with APP SNP rs2096488 on the degree of cortical cataract and on the bivariate outcome of cortical cataract and THV. Interestingly, the interaction between CTNND2 and APP was strongest in the univariate cortical cataract model suggesting the possibility that this interaction may be stronger or is detectable earlier in the lens. In addition, we demonstrated that a rare missense mutation (G810R) located 249 bp from rs17183619 alters the distribution of δ-catenin and Aβ secretion in neuronal cells. Immunohistochemistry experiments using a δ-catenin antibody revealed punctate staining in the cortical and supranuclear region of the lens from autopsy-confirmed AD cases but not from subjects lacking AD-associated neuropathology.

The rarity of the G810R mutation makes it unlikely that this polymorphism explains the association with CTNND2, despite its proximity to rs17183619 and demonstrated effect on APP processing. It is more likely that common variants or multiple rare variants in tight linkage disequilibrium with rs17183619 alter the function or expression of CTNND2 perhaps by affecting splicing or transcription factor binding.

CTNND2 encodes an adhesive junction-associated protein of the β-catenin superfamily that has been implicated in brain and eye development [47]. The δ-catenin protein is a component of the cadherin-catenin complex that recruits presenilin 1 to cadherins and inhibits Aβ production [48,49]. SNPs rs17183619 and rs13170756 surround exon 14 that contains the G810R mutation and encodes part of the highly conserved set of 10 armadillo repeat domains in δ-catenin [50]. Armadillo repeat domains 2–10 are necessary and sufficient to mediate the binding of δ-catenin to the hydrophilic loop of presenilin 1 in human brain [51]. Presenilin 1-deficient mice show significantly reduced expression of δ-catenin [52] and mice lacking normal δ-catenin display severe impairments in learning and memory tasks and in synaptic plasticity [53]. Emerging evidence supports a critical role for δ-catenin in dendritic spine maturation and maintenance in the cerebral cortex [54]. Recently, it has been suggested that δ-catenin tethers γ-secretase near synaptic membranes [55], supporting the previously proposed concept that γ-secretase activity is required for δ-catenin function via a mechanism involving γ-secretase.

The mechanism underlying the change in Aβ secretion observed in cells transfected with the G810R mutation warrants further investigation. We have previously shown that δ-catenin interacts with the TM6-TM7 hydrophilic loop domain of presenilin 1 [57], thus providing a plausible mechanism for direct modulation of γ-secretase activity. However, we also observed that the mutant δ-catenin has a markedly different subcellular distribution from the wild type protein in HEK293 cells transfected with the mutant δ-catenin construct, an observation consistent with the striking abnormal basolaminar distribution pattern noted in the epithelial layers of human AD lenses. Taken together, these observations suggest that δ-catenin may be aberrantly concentrated under the plasma membrane rather than in the nucleus in AD lens and brain. It is therefore conceivable that G810R alters intracellular trafficking of membrane proteins such as APP, or alters a signaling/transcriptional pathway that influences γ-secretase activity.

Our study showed only modest association of CTNND2 with cortical cataract and no evidence for association with measures of brain degeneration when the lens and brain traits were considered independently in univariate analysis. These observations suggest that CTNND2 accounts for a very small portion of the genetic component of late-onset AD and age-related cortical cataract captured by the phenotype classification system used in the FEOS. Alternatively, AD-linked cortical cataract may be a distinct disorder as suggested by the distinctive subequatorial supranuclear
phenotype observed in late-onset AD and Down syndrome [10,11]. Another explanation is that CTNND2 variation affects a very specific process or pathway that is best represented by the bivariate measures of degeneration in the lens and brain that are mechanistically related to altered binding of δ-catenin to presenilin 1. This hypothesis is supported by our data showing apparent

Figure 3. Effect of the CTNND2 G810R mutation on intracellular distribution of δ-catenin and Aβ secretion in HEK293 cells stably expressing human APP with the Swedish mutation (APPsw). A. HEK293 APPsw cells were transfected with empty vector, wild type delta-catenin or G810R mutant delta-catenin (n = 3 each). Effect of G810R on cell morphology. In contrast to wild type δ-catenin, mutant δ-catenin is predominantly located at/under the cell surface and G810R mutant cells show a more elaborate network of protrusions (white arrows), suggesting that it may be altering interactions of the plasma membrane with the underlying actin cytoskeleton. Lower panels show detail for apical segments (red rectangles) at a higher magnification. B. Semi-quantitative analysis shows significantly more protrusions extending from apical segments in cells with G810R mutation (p = 0.006). Error bars represent the standard deviation. C. Effect of G810R on Aβ secretion. Conditioned media was collected at 16 hours and assayed for Aβ1–40 and Aβ1–42. Aβ concentrations were corrected for total protein levels. Error bars represent the standard deviation. When compared to empty vector controls, wild type δ-catenin had no effect on secreted Aβ levels. In contrast, cells expressing mutant δ-catenin displayed a significant and specific increase in the secretion of Aβ1–42 (p = 0.02).

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increased accumulation and abnormal cellular distribution of δ-catenin in lenses from neuropathologically-confirmed AD patients. The significant interaction of CTNND2 and APP SNPs on the univariate measure of cortical cataract highlights the importance of investigating physiological changes in the lens that may precede neuronal loss.

One of the innovative aspects of this study is the application of a bivariate framework to identify genes for co-heritable traits. This approach detected genome-wide significant association of CTNND2 variants with the bivariate outcomes of cortical cataract and temporal horn volume, whereas association with this gene was not statistically remarkable when these traits were considered separately. Family-based samples are particularly well suited for establishing co-heritability of traits before consideration as a bivariate outcome in genetic association studies. Thus, the Framingham Offspring Study provided a unique opportunity to test our hypotheses because of its family-based design, prospective follow-up of individuals unbiased by selection for ocular or neurological disease, detailed ophthalmological data acquired about ten years prior to the first brain MRI examination, longitudinal monitoring of cognitive status, and availability of GWAS data.

While our results are significant and mechanistically plausible, our study has several limitations. The association findings of CTNND2 SNPs with quantitative outcomes derived from measures of cortical lens opacification and AD-linked neurodegeneration should be replicated in an independent sample even though they met genome-wide significance criteria. However, to our knowledge there are no other large cohorts having eye measurement data obtained many years before acquisition of brain MRI scan data. For this reason, we validated these findings through genetic association analyses with measures of cognitive performance and by experimental approaches. Our association findings for CTNND2 with bivariate outcomes including cognitive function are consistent with results from a recent GWAS in a prospectively followed cohort showing near genome-wide significance for association of rate of cognitive decline with rs2973488 [58], a CTNND2 SNP located 2 kb distal of rs13189742 which was strongly associated with the CC-THV bivariate outcome (Table S5). Another limitation relates to the use of an ordinal variable that precludes inclusion of precataractous lens pathology. Nonetheless, we were able to demonstrate statistically significant association with multiple variables derived from this less precise measure of lens opacity. We also note that meaningful comparisons of co-

Figure 4. Immunohistochemistry of δ-catenin in the human lens in Alzheimer disease (AD) from a 68 year-old male and 71 year-old female with neuropathologically-confirmed AD and from two normal male controls ages 68 and 70. Staining was observed in the anterior (Panels A–D), equatorial (Panels E–H) and supranuclear (Panels I–L) regions. In the anterior region, intense δ-catenin immunoreactivity was observed in the epithelial cell layer with a striking basolaminar distribution in the AD cases (Panels B and D) compared to age-matched controls (Panels A and C). Dense punctate δ-catenin immunoreactive deposits were observed in the epithelium (white arrowheads in Panels B and D) with laminar preponderance in the subjacent cortex and in the subequatorial and deep cortex of AD lenses (black arrowheads in Panels B and D). Dense punctate δ-catenin immunoreactive staining was also observed in the supranuclear region in the AD cases (black arrowheads in Panels J and L). doi:10.1371/journal.pone.0043728.g004
heritability estimates for clinically ascertained AD with specific cataract and MRI phenotypes in the same sample could not be made as only seven incident AD cases currently exist among the 1249 subjects who participated in both the eye and MRI examination arms of the study. However, analysis of an enlarged sample including 139 AD cases who had the eye exam only revealed significant co-heritability of CC and PSC with AD (Table S2). Thus, the co-heritability estimates for lens and MRI traits are especially remarkable since all of these subjects were dementia-free at the time of the ophthalmic examination and ten years following at the time of the initial MRI examination. Future studies of this cohort are likely to provide additional insight into the relationship of AD pathogenesis in the brain and lens.

Taken together, prior observations and the results from our study support the existence of a pathway leading to AD-linked pathology in the brain and lens, a hypothesis that supports a systemic rather than brain-limited focus for age-dependent AD pathogenesis. This hypothesis is indirectly supported by the epithelial origin of the lens and brain (surface ectoderm and neuroectoderm, respectively) and the long-lived nature of the terminal differentiated cell types affected by AD pathology in the lens and brain. Further investigation is needed to determine the specific molecular and cellular mechanisms underpinning presumptive linkage of AD pathology in these two anatomical compartments. The implication that a genetic variant can alter the function of a protein affecting cortical cataract and AD suggests that these two systemically distinct diseases may be related, thus suggesting possible convergent pathogenic mechanisms. Moreover, δ-catenin, and possibly other members of the cadherin-catenin complex, may provide new therapeutic targets for AD and cortical cataracts. Finally, detection of AD-linked lens pathology could serve as a peripherally accessible biomarker to facilitate discovery, development, evaluation, and implementation of emerging AD therapeutics.

Supporting Information

Figure S1 Experimental design for computational studies. A. Dates and mean ages of Framingham Study Original Cohort and Offspring Cohort participants at the time of the eye and brain MRI exams, and mean age at onset of incident AD cases subsequent to these exams. B. Samples included in the co-heritability and bivariate GWAS components of the study. Fewer than 15% of the 5,209 Original Cohort members participated in the eye and MRI exams approximately 43 and 53 years, respectively, after entry into the Framingham Study in 1948. Approximately one-half of the 5,216 Offspring cohort members enrolled in 1971 participated in these exams. Co-heritability analyses included members of both cohorts whereas the GWAS study was limited to Offspring Cohort members since eye, MRI and GWAS data were available for very few individuals from the Original Cohort.

(TIF)

Figure S2 Retroillumination slit lamp photomicrograph of a dilated right and left eyes from a 57-year-old female Framingham Offspring Study participant demonstrating equatorial cortical opacification with cortical spoking and posterior extension.

(TIF)

Figure S3 Heritability and co-heritability estimation. The heritability of a trait is derived from the correlation among siblings, as shown in the diagram by the solid blue and orange arrows. Co-heritability of two traits is derived from the cross-trait sibling correlation which is obtained by averaging the correlation of the first trait in sibling A with the second trait in sibling B and the correlation of the second trait in sibling A with the first trait in sibling B shown by the crossed blue and orange arrows.

(TIF)

Figure S4 Quantile-quantile (Q-Q) plot of observed (y-axis) vs. expected (x-axis) P-values from genome-wide association tests for the bivariate outcome of cortical cataract and temporal horn volume. Black dots represent all genotyped SNPs and red dots denote the imputed SNPs from CTNND2 with P<10^{-5}. The number of SNPs (188,629) includes genotyped SNPs (186,192) and imputed SNPs (2,437) from selected gene regions.

(TIF)

Figure S5 δ-catenin in the human retina. A. δ-catenin immunostaining in a retina from a 70-year-old female with AD. B. same retina in panel A stained with hematoxylin and eosin. C. δ-catenin immunostaining in a retina from a 50-year-old male control. Abbreviations: VIT, vitreous body; ILM, inner limiting membrane; GCL, granule cell layer; INL, inner nuclear layer; OPL, outer plexiform layer; ONL, outer nuclear layer; PRL, photoreceptor layer; RPE, retinal pigment epithelium; CHO, choroid; SCL, sclera.

(TIF)

Table S1 Sample characteristics.

(DOCX)

Table S2 Trait correlations.

(DOCX)

Table S3 Correlations for cortical cataract with selected MRI traits.

(DOCX)

Table S4 Top-ranked GWAS results (P<10^{-5}) with 186,192 genotyped SNPs in bivariate models of cataract and temporal horn volume.

(DOCX)

Table S5 Top-ranked association results (P<10^{-5}) for CTNND2 SNPs with cortical cataract (CC), temporal horn volume (THV), and the bivariate outcome CC-THV.

(DOCX)

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Author Contributions

Conceived and designed the experiments: GJ LEG PStG-H LAF. Performed the experiments: JAM CK. Analyzed the data: GJ JB LAF. Contributed reagents/materials/analysis tools: SS ACM PAW. Wrote the paper: GJ PStG-H LEG LAF. Critical evaluation and edited the manuscript for intellectual content: GJ JAM SS GL PAW PStG-H LEG LAF. Provided study supervision: PStG-H LEG LAF.

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