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Panagiotou, ES, Sanjurjo Soriano, C, Poulter, JA orcid.org/0000-0003-2048-5693 et al. (15 more authors) (2017) Defects in the Cell Signaling Mediator β-Catenin Cause the Retinal Vascular Condition FEVR. The American Journal of Human Genetics, 100 (6). pp. 960-968. ISSN 0002-9297

https://doi.org/10.1016/j.ajhg.2017.05.001

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Defects in the cell signaling mediator β-catenin cause the retinal vascular condition FEVR

Evangelia S. Panagiotou¹, Carla Sanjurjo Soriano¹, James A. Poulter¹, Emma C. Lord¹,
Denisa Dzulova¹, Hiroyuki Kondo²,³, Atsushi Hiyoshi², Brian Hon-Yin Chung⁴, Yoyo Wing
Yiu Chu⁴, Connie H.Y. Lai⁵, Mark E. Tafoya⁶, Dyah Karjosukarso⁷,⁸, Rob W.J. Collin⁷,⁸,
Joanne Topping¹, Louise M. Downey¹,⁹, Manir Ali¹, Chris F. Inglehearn¹ and Carmel
Toomes¹,*

¹Section of Ophthalmology & Neuroscience, Leeds Institute of Molecular Medicine,
University of Leeds, Leeds LS9 7TF, UK; ²Department of Ophthalmology, Fukuoka
University, Fukuoka 814-0180, Japan; ³Department of Ophthalmology, University of
Occupational and Environmental Health, Kitakyushu 807-8555, Japan; ⁴Department of
Paediatrics and Adolescent Medicine, Centre for Genomic Sciences, The University of Hong
Kong, Queen Mary Hospital, Hong Kong, China; ⁵Department of Ophthalmology, The
University of Hong Kong, Queen Mary Hospital, Hong Kong, China; ⁶Pacific Retina Care,
Waikele, Hawaii 96797, USA; ⁷Department of Human Genetics and Donders Institute for
Brain, Cognition and Behaviour, Radboud University Medical Centre, 6525 GA Nijmegen,
the Netherlands; ⁸Department of Ophthalmology, Hull Royal Infirmary, Hull HU3 2JZ, UK.

* Correspondence: c.toomes@leeds.ac.uk

Running title: Mutations in CTNNB1 cause FEVR.
Abstract

Familial exudative vitreoretinopathy (FEVR) is an inherited blinding disorder characterized by the abnormal development of the retinal vasculature. The majority of mutations identified in FEVR are found within four genes that encode the receptor complex (FZD4, LRP5 and TSPAN12) and ligand (NDP) of a molecular pathway that controls angiogenesis, the Norrin-β-catenin signaling pathway. However, half of all FEVR cases do not harbor mutations in these genes, indicating that further mutated genes remain to be identified. Here we report the identification of mutations in CTNNB1, the gene encoding β-catenin, as a cause of FEVR. We describe heterozygous mutations in two dominant FEVR families (c.2142_2157dup p.(His720*) and c.2128C>T, p.(Arg710Cys)) and a de novo mutation in a simplex case (c.1434_1435insC, p.(Glu479Argfs*18)). Previous studies have reported heterozygous de novo CTNNB1 mutations as a cause of syndromic intellectual disability (ID) and autism spectrum disorder and somatic mutations are linked to many cancers. However, in this study we show that Mendelian inherited CTNNB1 mutations can cause non-syndromic FEVR, and that FEVR can be a part of the syndromic ID phenotype, further establishing the role that β-catenin signaling plays in the development of the retinal vasculature.
Beta-catenin is an essential protein that orchestrates many key processes during human development and throughout adulthood. It fulfills these functions through its roles as both a cell adhesion molecule and a mediator of cell signaling.¹ It is encoded by CTNNB1 (MIM 116806) and somatic mutations in this gene, which predominantly result in the inhibition of the degradation of β-catenin, are well characterized in many different cancers (MIM 114500, 114550, 155255, 167000, 132600).² More recently, large-scale sequencing studies in individuals with syndromic intellectual disability (ID, MIM 615075) or autism spectrum disorder (ASD, MIM 209850) have identified heterozygous de novo mutations in this gene as the cause of these overlapping disorders.³⁻⁴ In this report, we describe inherited Mendelian mutations in CTNNB1 in the blinding disorder familial exudative vitreoretinopathy (FEVR).

FEVR is an inherited developmental disorder caused by incomplete retinal angiogenesis (MIM 133780).⁵ The primary defect, and the hallmark phenotype of this disease, is the absence of vasculature in the peripheral retina but this feature alone often causes no symptoms. In a subset of cases, however, retinal ischemia ensues resulting in a variety of sight-threatening secondary complications including neovascularisation, vitreoretinal traction, exudation, retinal folds and retinal detachment.⁶ The expressivity of FEVR is remarkably variable, even within families, ranging from asymptomatic individuals who are unaware that they have the condition to severely affected individuals who lose their vision in infancy.⁷

In addition to its phenotypic variability, FEVR is also genetically heterogeneous showing dominant, recessive and X-linked modes of inheritance. Mutations in eight genes have so far been found to cause FEVR and a ninth has been mapped but remains to be identified: NDP (MIM 300658), FZD4 (MIM 604579), LRP5 (MIM 603506), TSPAN12 (MIM 613138),
A TOH7 (MIM 609875), ZNF408 (MIM 616454), KIF11 (MIM 148760), RCBTB1 (MIM 607867) and EVR3 on chromosome 11p12-13 (MIM 605750).\(^8,19\) Mutation screening has shown that only \(\sim 50\%\) of FEVR families have mutations in these genes.\(^{20}\) Of these, the vast majority (\(\sim 90\%\) of cases with a molecular diagnosis) harbor mutations in four genes, NDP, FZD4, LRP5 and TSPAN12, all of which encode components of the same molecular pathway, the Norrin-\(\beta\)-catenin signaling pathway.\(^7,20\)

The Norrin-\(\beta\)-catenin pathway (also referred to as the Norrin/Fz4 pathway) is a derivative of the Wnt-\(\beta\)-catenin pathway but it is activated by a non-Wnt ligand and utilizes a specific set of receptors (Figure 1).\(^{21}\) Norrin (encoded by NDP) is the extracellular ligand which binds to a receptor complex composed of Frizzled-4 (FZD4), Low-density lipoprotein receptor-related protein-5 (LRP5) and Tetraspanin-12 (TSPAN12).\(^{22}\) The ligand-receptor signaling complexes are believed to cluster through the multimerization of TSPAN12, and possibly Norrin, to form specialized membrane microdomains. In the absence of Norrin binding, cytoplasmic \(\beta\)-catenin is targeted for ubiquitination and proteasomal degradation through the action of the “destruction complex”. This complex is composed of the scaffolding proteins Axin and Adenomatous polyposis coli (APC) and the Ser/Thr kinases Glycogen synthase kinase 3 (GSK3) and Casein kinase 1 (CK1). CK1 and GSK3 sequentially phosphorylate \(\beta\)-catenin, resulting in \(\beta\)-catenin recognition and ubiquitination by \(\beta\)-transducin repeat containing protein (\(\beta\)-Trcp) and subsequent degradation.\(^{23}\) However, upon Norrin binding, the resulting signal directs the destruction complex to relocate to the receptors at the cell membrane through the action of Dishevelled (DSH). This inhibits the degradation of \(\beta\)-catenin, allowing its cytoplasmic levels to increase. Subsequently, \(\beta\)-catenin translocates into the nucleus and interacts with the T-cell factor (TCF)/Lymphoid enhancing factor (LEF) family of transcription factors in order to control the expression of target genes.\(^{22}\)
The genetic investigation of families with FEVR-related retinopathies has been pivotal in helping to define the components of this unique angiogenesis pathway and it is hoped that the remaining unidentified mutated genes will provide further insight. With this in mind, the aim of this study was to identify new genes mutated in FEVR.

Our experiment focused on an international multi-ethnic cohort of 36 families, with at least one member diagnosed with FEVR, in whom we had previously excluded the presence of mutations in the known genes mutated in FEVR using Sanger sequencing and/or targeted next generation sequencing. The study was approved by the Leeds East (Project number 03/362) Research Ethics Committee and adhered to the tenets of the Declaration of Helsinki. Individuals participated in the study after giving their informed consent. Blood samples were obtained from both affected and unaffected family members and genomic DNA was extracted using standard protocols.

Whole exome sequencing (WES) was performed in one affected individual from 12 of the families in the cohort. The SureSelect All Exon V5 Capture Reagent (Agilent Technologies) was used to enrich exonic regions from genomic DNA and processed according to Agilent’s SureSelect\textsuperscript{QXT} Library Prep protocol. The processed libraries were subsequently pooled and subjected to 150-bp paired-end sequencing on a HiSeq3000 sequencer (Illumina). The resulting reads were aligned to the human reference genome (GRCh37) using Novoalign short-read alignment software (Novocraft Technologies) and processed in the SAM/BAM format using Picard and the Genome Analysis Toolkit (GATK).\textsuperscript{24-26} SNPs and indels were called in the VCF format using the HaplotypeCaller function of GATK, with variants being
filtered on the basis of mapping quality, strand bias and genotype quality. After further filtering to exclude polymorphisms with a minor allele frequency >1% in dbSNP142, ExAC or EVS (see URLs), the remaining variants were annotated and their functional consequences determined with ANNOVAR software (version July 2014). For the full WES data set the average depth of coverage was 60x, with an average 96% of targeted bases covered at a depth of 5x and 90% at 10x. This data confirmed that no pathogenic nucleotide variants or copy number variations (screened for via ExomeDepth software) were present in the genes known to cause FEVR when mutated. The remaining lists of variants were subsequently compared to determine whether any mutations were present within the same gene in two or more families. This analysis led to the discovery of different heterozygous CTNNB1 mutations. We therefore designed primers (primer pairs are shown in Table S1) and used PCR and Sanger sequencing to confirm and segregate mutations, and to screen all 14 coding exons and flanking intronic sequences in one affected member from the 24 remaining families in the cohort. Three CTNNB1 mutations were identified which were not present in the dbSNP, EVS or ExAC databases. These mutations accounted for 8% (3/36) of FEVR cases in this series, although it should be noted that the known mutated genes causing FEVR had previously been excluded in the majority of this cohort.

A heterozygous 16-bp duplication in exon 15 was found in a Hawaiian FEVR family (Family F258) of Japanese origin, c.2142_2157dupTAGCTATCGTTCTTTT, p.(His720*)(Figure 2). Three family members were affected by FEVR and the mutation segregated perfectly with the disease phenotype in the pedigree. The proband (II:2) was assessed in her late teens. Her left eye had a retinal detachment with proliferative vitreoretinopathy and temporal traction resulting in a severe retinal fold through the macula. She had a vitrectomy with silicone oil but her vision remained poor (Hand Movements). Her right eye had temporal macular
dragging but this was stabilized by peripheral laser treatment and she maintained good vision in this eye (20/40). Her father (I:1) had nystagmus and severe retinal scarring and was registered blind. When last assessed, in his fifth decade, his vision was limited to Hand Movements in one eye and No Light Perception in the other. The proband’s affected younger brother (II:3) was diagnosed at age 10 when he started to have active retinal disease in both eyes. This was stabilized with bilateral laser treatment allowing him to maintain good vision. The mother and two unaffected siblings had normal eyes.

In a Japanese family segregating the FEVR phenotype (Family F410), a heterozygous missense change in exon 14 was identified, c.2128C>T, p.(Arg710Cys) (Figure 2). The mutation was present in all affected family members. The proband (III:2) presented at 11 years of age with very low vision. Fundus examination showed retinal avascularity, exudation, retinal holes and bilateral retinal detachment, for which he underwent buckle surgery in both eyes (Figure 3). His affected brother (III:1) had a similar phenotype with retinal avascularity, exudation and bilateral tractional folds, while their father (II:1) showed abnormal retinal vessels and retinal avascularity. The paternal grandmother (I:2) showed no evidence of retinal avascularity upon fluorescein angiography examination but she did have unilateral focal retinal degeneration. As this phenotype was not typical for an FEVR diagnosis but was clearly abnormal, she was given a diagnosis of “query FEVR” but does carry the mutation. The proband’s asymptomatic brother (III:3) was also found to carry the mutation but fundus examination at aged 9 revealed no signs of disease. Analysis of the mutated p.Arg710 amino acid among homologues of CTNNB1 revealed this residue to be conserved down to Danio rerio and Xenopus (Figure S2). Furthermore, this mutation was predicted to be disease causing on various bioinformatic pathogenicity prediction tools (Table S2).
A 1-bp insertion in exon 9 was identified in a simplex Chinese case, a child with congenital FEVR (Family F1321), c.1434_1435insC, p.(Glu479Argfs*18) (Figure 2). He presented at 1-month old and assessment under general anesthesia showed that his left eye had a total retinal detachment with anterior fibrovascular proliferation, vitreous haemorrhage and 360 degree rubeosis iridis. His right eye showed shallow retinal detachment with fibrovascular traction at the optic disc and vitreous haemorrhage (Figure 3 and S1). Neuroimaging and skeletal survey were unremarkable and no facial dysmorphology was noted. Both parents had normal eyes. Segregation analysis revealed that this mutation was de novo, as neither of the unaffected parents carried the mutation but the inheritance of variants in unrelated genes was consistent with the child being related to both parents.

Human β-catenin contains 781 amino acids and is subdivided into three domains; an amino-terminal domain (NTD), a central region containing 12 armadillo (ARM) repeats (residues 138-664) and a carboxy-terminal domain (CTD). The NTD contains the binding site for α-catenin, the phosphorylation sites for GSK3 and CK1 and is the location of the mutations responsible for the oncogenic form of β-catenin. The ARM repeats form the binding sites for the majority of the interacting partners of β-catenin and are crucial for its roles in cell adhesion and signaling. At the start of the CTD is an α-helix motif (residues 667-683), termed Helix-C, that facilitates the binding of proteins to the ARM repeats and is essential for β-catenin signaling. The role of the remaining CTD is less well characterized as many of the experiments on this section of the protein also included the last ARM repeat and Helix-C. However, it has been suggested that this domain influences signaling by preventing the self-aggregation of the ARM repeat domain and increasing the binding specificity of ARM repeat interactions.
The effects of the mutations identified in this study are predicted to produce two different outcomes. The exon 9 frameshift mutation identified in family F1321 is expected to lead to nonsense mediated mRNA decay (NMD) of the mutated transcript and hence to haploinsufficiency of CTNNB1. However, the truncating mutation identified in family F258 is in the last coding exon of CTNNB1, which would indicate that it would not be targeted for NMD and would create a truncated 719 amino acid polypeptide. This truncated form of β-catenin is predicted to be missing the very end of the protein beyond Helix-C. Similarly, the missense mutation identified in family F410 (p.(Arg710Cys)) alters an amino acid residue in this uncharacterized terminal region of the CTD (Figure 2). These findings indicate that FEVR is part of the phenotypic spectrum caused by CTNNB1 haploinsufficiency, and that non-syndromic FEVR, which is at the milder end of this spectrum, is caused by mutations altering the uncharacterized CTD of β-catenin.

De Ligt and colleagues first reported de novo CTNNB1 mutations in three subjects with ID, microcephaly and spasticity. Subsequent reports followed which showed that de novo CTNNB1 mutations lead to a recognizable clinical entity which comprises ID with speech impairment, abnormal muscle tone, ASD, microcephaly, distinctive facial features and brain abnormalities such as corpus callosum hypoplasia. This form of syndromic ID appears to be the result of haploinsufficiency of CTNNB1. While full gene deletions have been reported in two individuals, the majority of intragenic mutations (Figure S3) are predicted to result in a truncated transcript that will undergo NMD and this mechanism has been confirmed experimentally in two subjects. In a parallel study, two de novo CTNNB1 mutations were found in individuals with ASD. At the moment it is not clear if the ASD cases have additional features consistent with them having a diagnosis of syndromic ID as only limited
clinical information was provided. However, these individuals did have low non-verbal IQ scores indicating that they also had ID.\textsuperscript{4} Furthermore, the reported ASD mutations are a nonsense (p.(Trp504*)) change and a missense variant in the crucial ARM repeat domain (p.(Thr551Met)),\textsuperscript{4} suggesting that they will result in null CTNNB1 alleles similar to mutations which cause syndromic ID.

All the affected individuals in the current study had a diagnosis of non-syndromic FEVR, clearly linking mutations in CTNNB1 with this phenotype. However, given that the affected child from family F1321 is predicted to have haploinsufficiency of CTNNB1, and was only 4-weeks old when originally examined, the family was re-contacted following this molecular result. The child is now 3 years old and displays many clinical features associated with syndromic ID: global developmental delay, motor delay (he can stand briefly with support), significant speech delay (he only uses 1-2 single words and most of his communication is at a non-verbal level) and dysmorphic facial features (squint, long face and prominent nasal tip). Interestingly, a recent report has described a 22-month old boy with a history of lipomyelomeningocele, failure to thrive, short stature, developmental delay and ASD along with an FEVR ocular phenotype. This child was reported to have a de novo heterozygous mutation in CTNNB1, c.2112_2116dupAGAAC, p.(Pro706Glnfs*31) but no other genetic information was provided. It is therefore unclear if all the child’s clinical features are attributable to the β-catenin defect but it appears to be another report of FEVR in an infant with a predicted null allele in CTNNB1.\textsuperscript{36} The presence of FEVR in these two cases supports the hypothesis that FEVR is part of the CTNNB1 haploinsufficiency phenotype.

Although mild visual defects have been reported in 19/30 of the CTNNB1-related syndromic ID cases reviewed to date,\textsuperscript{32-35} these were not part of the typical FEVR phenotypic spectrum.
and frequently included either strabismus, hyperopia, myopia or poor eye contact. In addition, a single case of photophobia has been reported and another subject had slight optic atrophy in one eye and moderate optic papilla hypoplasia in the other.\textsuperscript{32-34} Given our findings, we suspect that many of these cases may also have subtle retinal avascularity. In all dominant forms of FEVR the severity of the disease is highly variable and asymptomatic individuals carrying mutations are frequently reported.\textsuperscript{7,10,12,37} The asymptomatic case present in Family F410 in this study indicates that CTNNB1-related FEVR follows the same pattern of variable expression so it would be interesting to perform fluorescein angiography examinations on a cohort of CTNNB1-related syndromic ID and ASD individuals to look for mild FEVR.

The familial mutations identified in this study only caused an FEVR phenotype. It is unlikely that additional defects will have been missed in these individuals given their age but they clearly display the full spectrum of FEVR severity. One could speculate that the mutations are only altering a tissue-restricted isoform of CTNNB1 but there is no evidence to support the existence of such a transcript. There are currently 15 isoforms of CTNNB1 on Ensembl (accessed 03/17) and not all of these contain exons 14 and 15 or encode proteins. However, all four of the annotated RefSeq coding transcripts encode proteins with identical CTDs and are predicted to be altered by the mutations (Genome Browser accessed 03/07). The importance of this part of the transcript is reflected by the lack of variants in the exome databases in this part of the gene. ExAC only shows two loss of function alleles in CTNNB1 and one of these is of low quality while the other is a splice defect which is predicted to alter the exons encoding the uncharacterized CTD. Similarly, there are only 28 variants predicted to cause missense substitutions in the CTD beyond Helix-C and these are all rare, the most common one having an allele frequency of 0.0001.
Undoubtedly, the fact that both of the familial mutations are predicted to produce β-catenin with a defective CTD is the likely explanation why these mutations are sufficiently mild to be passed from one generation to the next and why they cause a restricted ocular phenotype. We did attempt to identify additional familial mutations in this region of the gene by sequencing exons 14 and 15 of CTNNB1 (which encode the CTD beyond Helix-C) in an additional cohort of 36, partially pre-screened, predominantly Northern European FEVR cases. However, no more variants were identified indicating that these mutations are rare.

Previously, Cox and co-workers examined the effects of truncating the Drosophila β-catenin homologue, armadillo, in an in vivo model. One of the mutants investigated was almost identical to the predicted truncated protein present in family F258. Their results showed that the truncation had no effect on cell adhesion but caused a reduction in armadillo (β-catenin) signaling. To investigate if the familial mutations identified in this study caused a similar reduction in signaling, the mutations were modeled using the TOPflash β-catenin transcriptional reporter assay. Untagged expression constructs for wild-type (WT) or mutant β-catenin (p.Arg710Cys or p.His720*) in pDEST40 (Life Technologies) were created using the QuikChange II XL site-directed mutagenesis kit (Agilent Technologies) and the expression capability of the constructs was verified by western blot (Figure S4). The assays were performed in STF cells which are HEK293 cells stably transfected with the TOPflash firefly luciferase construct (a kind gift from Jeremy Nathans). Experiments were performed in triplicate in a 24-well plate and repeated in three independent experiments. 90,000 cells/well were transfected with 399 ng of construct DNA plus 1 ng of Renilla luciferase control plasmid (pRL-TK from Promega) using 1.5 µl of FuGENE 6 transfection reagent (Promega). Luciferase activity was measured after 48 hours using the Dual-Luciferase Reporter assay system (Promega) on a Mithras LB 940 plate reader (Berthold Technologies).
The data were analyzed using GraphPad PRISM 7.0 software (one-way ANOVA and Dunnett’s test). The results showed that both of the variant CTNNB1 constructs produced significantly different TOPflash activity levels when compared to the WT CTNNB1 construct but that they had opposite effects (Figure 4). The p.His720* variant produced significantly reduced levels of TOPflash activation, almost at null allele levels, whereas the p.Arg710Cys variant produced significantly higher levels of TOPflash activation.

It is difficult to envisage a mechanism explaining these contradictory findings. However, contrasting TOPflash results like these have been previously reported for NDP mutations. Similarly, although a reduction in Norrin-β-catenin signaling is usually associated with the angiogenesis defects associated with FEVR, ectopic over-activation of Norrin signaling has also been shown to disrupt embryonic angiogenesis. Nevertheless, this experiment is unlikely to be an accurate biological replicate of the in vivo consequences of these mutations. For example, if the p.(His720*) mutation reduced β-catenin signaling levels to that same extent in family F258, they would be expected to have additional clinical features associated with CTNNB1-related ID syndrome. The molecular mechanisms that control β-catenin signaling are extremely complex and involve a growing number of tissue specific co-factors, inhibitors and activators which vary in a time and context dependent manner and are unquestionably modulated by feedback loops and other signaling pathways. Although a great deal of research into the function of β-catenin has been performed in animal and cell-based models, it is clear that they do not always accurately model the function of β-catenin in humans. A clear example of this is the heterozygous β-catenin knockout mice. These should be a model for CTNNB1-related syndromic ID but are reported to have a wild-type phenotype.
The fact that the FEVR phenotype is also present in individuals who are haploinsufficient for CTNNB1 suggests that these mutations may cause a partial loss-of-function of β-catenin and therefore act in a hypomorphic fashion. However, it’s unclear if the defects alter a retina or Norrin-β-catenin specific function or if they reduce a ubiquitous fundamental function of β-catenin and the retinal vasculature is the tissue most susceptible to this reduction. The location of the mutations and the abnormal TOPflash assay results point to the mutations altering the transcriptional activity of β-catenin. However, in addition to its well-established roles in cell adhesion and transcription, β-catenin has also been reported to play an uncharacterized role at the centrosome and in mitotic spindle formation. The links between asymmetric cell division and angiogenesis, and the fact that the gene encoding the mitotic protein KIF11 is mutated in FEVR, suggest this potential role warrants further investigation.

Conditional knockout studies in mice have shown that β-catenin depletion in endothelial cells leads to angiogenesis defects throughout the central nervous system (CNS) including the retina. Although FEVR is characterized as a disorder which only affects the retinal vasculature, the FEVR knockout mouse models suggests that there may be mild vascular defects in other parts of the CNS. For example, the Ndp and Fzd4 mice have capillary defects in the ear (stria vascularis) and additional vasculature defects have been identified in the cerebellum of the Fzd4 mouse. The retina is the most accessible part of the CNS and this allows detailed examination of its vasculature. It is this accessibility that allows us to diagnose subtle vascular defects in asymptomatic individuals harboring FEVR mutations but defects in the wider CNS could easily be missed. If present in humans, these could potentially contribute to some of the neurological features associated with syndromic ID and ASD.
Consistent with this suggestion, defects in angiogenesis have been reported in ASD, and there are many reports linking vascular defects with brain development, ID and dementia.\textsuperscript{51-54}

In summary, we have shown that heterozygous mutations in CTNNB1 can cause non-syndromic FEVR and that FEVR is part of the CTNNB1 haploinsufficiency phenotype. These findings confirm the importance of β-catenin signaling in the pathogenesis of FEVR, but also raise questions as to how mutations in such a ubiquitously essential protein can lead to such a restricted phenotype.

Accession Numbers

The accession numbers for the variants reported in this paper are ClinVar: SCV000266847-SCV000266849.

Supplemental Data

Supplemental Data includes four figures and two tables.

Acknowledgements

We thank the FEVR families for their participation in this study and Professor Jeremy Nathans (John Hopkins University School of Medicine) for kindly providing the STF cells. This project has received funding from the European Union’s Seventh Framework Programme for research, technological development and demonstration under grant agreement no. 317472 (EyeTN), Fight For Sight (grants 1493/1494 and GR586) and RP Fighting Blindness (grant GR586). We acknowledge the support of the UK Inherited Retinal Disease Consortium. The authors have no conflict of interest.
Web Resources

The URLs for data presented herein are as follows:


Combined Annotation Dependent Depletion (CADD), [http://cadd.gs.washington.edu](http://cadd.gs.washington.edu)


Ensembl, [http://grch37.ensembl.org](http://grch37.ensembl.org)

Exome Aggregation Consortium (ExAC), [http://exac.broadinstitute.org](http://exac.broadinstitute.org)

MutationTaster, [http://www.mutationtaster.org/](http://www.mutationtaster.org/)


Online Mendelian Inheritance in Man (OMIM), [http://www.omim.org/](http://www.omim.org/)

Perl scripts, [https://github.com/gantzgraf/vcfhacks](https://github.com/gantzgraf/vcfhacks)


PolyPhen2, [http://genetics.bwh.harvard.edu/pph2/](http://genetics.bwh.harvard.edu/pph2/)

References


Legends to Figures

Figure 1. Schematic overview of the Norrin-β-catenin pathway. Mutations in the genes encoding the ligand Norrin, the receptors FZD4, LRP5 and TSPAN12, and now the transcriptional activator β-catenin all cause FEVR. For pathway description see main text.

Figure 2. Mutations in CTNNB1 cause FEVR. (A) The pedigrees of the families and CTNNB1 mutation segregation data. The genotypes for all tested family members are shown below each individual, with M representing the mutant allele and + representing the wild-type allele. Affected individuals are shaded black. The grandmother (I:2) in family F410 is shaded grey as fluorescein angiography examination showed no evidence of peripheral retina avascularity which is the hallmark feature of FEVR, although she did show focal retinal degeneration in one eye. The symbol with a dot represents an asymptomatic individual. (B) Schematic representation of the CTNNB1 gene (NCBI: NM_001904.3) showing the location and sequence traces of the three disease-causing variants identified in this study. (C) Schematic representation of the β-catenin protein domains showing the variants identified in this study. The amino-terminal domain (NTD) spans amino acids 1-137, 12 armadillo repeats (ARM) span amino acids 138-664, the carboxy-terminal domain (CTD) spans 665-781 and the Helix-C domain spans amino acids 667-683.

Figure 3. Phenotype of FEVR cases with heterozygous CTNNB1 mutations. Color fundus photograph of the peripheral retina of (A) the right eye and (B) the left eye of the proband (III:2) from family F410 at age 11 years showing retinal holes and tractional retinal detachment. Anterior segment photograph of (C) the right eye and (D) the left eye of the proband from family F1321 aged one month showing the absence of retinal view or red reflex
due to dense vitreous hemorrhage in the right eye and neovascularization of the iris and a retrobulbar fibrovascular plaque obscuring the view of the retina in the left eye.

**Figure 4.** Functional assessment of mutant β-catenin on transcription using the TOPflash luciferase activity assay. STF cells were transiently transfected with expression constructs for wild-type (WT) or mutant β-catenin (p.Arg710Cys or p.His720*) or empty vector (pDEST40) alongside a Renilla luciferase plasmid and luciferase activity was measured 48 hours later. The bars represent the firefly/Renilla luciferase ratios for the different constructs. The results are from three independent experiments performed in triplicate. Error bars depict SEM; **p < 0.01, ****p < 0.0001.