CRB1 is essential for external limiting membrane integrity and photoreceptor morphogenesis in the mammalian retina

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Mutations within the CRB1 gene have been shown to cause human retinal diseases including retinitis pigmentosa and Leber congenital amaurosis. We have recently identified a mouse model, retinal degeneration 8 (rd8) with a single base deletion in the Crb1 gene. This mutation is predicted to cause a frame shift and premature stop codon which truncates the transmembrane and cytoplasmic domain of CRB1. Like in Drosophila crumbs (crb) mutants, staining for adherens junction proteins known to localize to the external limiting membrane, the equivalent of the zonula adherens in the mammalian retina, is discontinuous and fragmented. Shortened photoreceptor inner and outer segments are observed as early as 2 weeks after birth, suggesting a developmental defect in these structures rather than a degenerative process. Photoreceptor degeneration is observed only within regions of retinal spotting, which is seen predominantly in the inferior nasal quadrant of the eye, and is caused by retinal folds and pseudorosettes. Photoreceptor dysplasia and degeneration in Crb1 mutants strongly vary with genetic background, suggesting that the variability in phenotypes of human patients that carry mutations in CRB1 may be due to interactions with background modifiers in addition to allelic variations. The Crb1rd8 mouse model will facilitate the analysis of Crb1 function in the neural retina and the identification of interacting factors as candidate retinal disease genes.

INTRODUCTION

Mutations in the crumbs-like 1 (CRB1) gene lead to various forms of heritable retinal disorders in humans (1–5). CRB1 and related genes in other species encode transmembrane proteins that localize to the apical membrane of epithelial cells. The Drosophila crumbs (crb) gene has been studied extensively. Crb is a key regulator of polarity of many epithelial cells in which it acts as an apical determinant and contributes to the assembly of the zonula adherens (ZA), a belt-like adherens junction that separates apical and basolateral membranes (reviewed in 6).

Recent work has identified multiple roles for Crb in the Drosophila retina. Crb is essential for ZA integrity and for stalk membrane formation during the morphogenesis of photoreceptor cells (7,8). Primary defects in these structures cause a highly abnormal shape of photoreceptor cells, in particular, of the light sensing organelle, the rhabdomere. Moreover, Crb has also been shown to support photoreceptor survival during continuous light exposure (9). These findings suggest that CRB1 may be important for integrity of the ZA of the neural retina, also known as the external limiting membrane (ELM), and for the development of photoreceptor inner and outer segments, which are the subdivisions of the apical membrane

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of photoreceptors that correspond to the fly stalk membrane and rhodopsome, respectively.

Interestingly, CRB1 mutations lead to a variety of retinal diseases including retinitis pigmentosa (RP) characterized by sparing of para-arteriolar retinal pigment epithelium (1), RP with Coats-like exudative vasculopathy (2) and Leber congenital amaurosis (LCA, 2–5). The retinal diseases observed cannot be explained solely by allelic differences as individuals with the same mutation have been reported to have different forms of RP (2). Multiple roles that CRB1 may play in retinal differentiation and maintenance, and its interaction with other cellular components necessary to carry out its function, might in part explain the range of retinal diseases and the variation in phenotype within each retinal disease observed. While much has been learned about crb function in Drosophila, a mammalian model would be extremely useful to begin to elucidate the causes of the broad spectrum of phenotypes observed in individuals with CRB1 mutations.

RESULTS

Focal retinal dysplasia and degeneration with pan-retinal shortening of inner and outer segments is observed in rd8 mice

During the process of introgressing Mfprp16 onto the C57BL/6 (B6) background, a mouse was found with large retinal spots rather than the usual, small, discrete, pan-retinally distributed spots. Through further crosses, we determined that a partial large spots phenotype could be produced in the absence of Mfprp16, suggesting the presence of a second spontaneous mutation. This recessive mutation was named retinal degeneration 8 (rd8).

Clinically, mice homozygous for the rd8 mutation exhibit large, irregularly shaped spots most heavily concentrated in the inferior nasal quadrant of the fundus (Fig. 1). By orienting the eye prior to enucleation, we determined that the clinical spotting corresponds to regions with retinal folds and pseudorosettes that involve the photoreceptors and often distort the inner nuclear layer (Fig. 2B). Areas of retinal thinning that include both inner and outer segments are evident in affected regions. Unlike other models of photoreceptor degeneration that exhibit pan-retinal degeneration, degeneration in rd8 mice is focal in nature. This aspect is particularly striking in older mice, in which the outer nuclear layer may be reduced to a single row of nuclei in a sharply demarcated area, with normally near retina present at the edge of the region of severe degeneration (Fig. 2C and D).

Electron microscopy reveals that the photoreceptor inner segments (PR IS), the equivalent of the Drosophila stalk membrane (7), also lose their orderly arrangement, and at 4 weeks of age are ∼25% shorter than those of wild-type mice (Fig. 2E). This observation is consistent with findings in Drosophila (7), in which the lack of Crb leads to the formation of shortened stalks. Photoreceptor outer segments (PR OS) of rd8 mutant mice are also shortened. By 10 weeks, outer segments begin to fragment, and a breakdown of individual photoreceptor lamellae is observed, resulting in the accumulation of granular debris (Fig. 2F). By 5 months, in the most affected regions, only a few outer segment fragments remain and the inner segments approach the retinal pigment epithelium.

In addition, diffuse swelling of portions of the inner segments is observed (Fig. 2G). Although the inner retina is generally normal, Müller cell processes are unusually prominent both in the inner retina and in the inner nuclear layer. While normally difficult to identify, the Müller cell cytoplasm shows increased electron density and individual cytoplasmic strands that are easily traceable through the inner plexiform layer (Fig. 2H).

rd8 maps to chromosome 1 and is caused by a mutation in Crb1

Since rd8 was first observed, 10 additional mouse strains have been identified that have retinal phenotypes similar to that of rd8. Complementation tests were carried out between B6.C3Ga-rd8/rd8 and five of the strains; all of the retinal mutations were found to be allelic to rd8. Interestingly, all of the strains were incipient congenic strains in which different mutations were being introgressed onto the B6 background in our importation/induced mutant resource facility, suggesting a founder effect arising from a spontaneous mutation in the B6 strain.

To elucidate the molecular basis of rd8 and to obtain a fine structure map for the locus, a mapping cross was produced by outcrossing B6.C3Ga-rd8/rd8 and B6.FVB-rd81J/rd81J to CAST/Ei, a wild-derived strain, and back-crossing the resulting F1 animals to an affected parent of the same strain (herein referred to as rd8 and rd81J/BC, respectively). The rd8 locus was initially mapped by genome scan to chromosome 1, ~73 cM from the centromere, in a cohort of 100 mice. The crosses were expanded to 339 back-cross progeny from the rd8 BC and 267 back-cross progeny from the rd81J BC. All mice were phenotyped by indirect ophthalmoscopy and genotyped, initially with markers D1Mit24 and D1Mit42, and subsequently with the flanking markers D1Mit30 or D1Mit498 proximally, and D1Mit141 or D1Mit423 distally (Fig. 3A). All mice recombinant within the critical interval were progeny tested to confirm whether the rd8 mutation was being transmitted in the recombinant animal. Cumulatively, the critical interval containing rd8 and rd81J, between markers D1Mit538 and D1Mit141, was estimated to be 0.50 ± 0.29 cM.

Using mouse and human genome sequencing data, we assembled a physical contig across the minimal genetic region and found that it contained a mouse orthologue of the Drosophila crb gene (Fig. 3B). Sequence analysis of Crb1 cDNA from rd8 mice showed a one base pair deletion at nt3481, which causes a frameshift and a premature stop codon following amino acid 1207 (Fig. 3C). The entire coding region for rd81J was also tested for mutations by direct sequencing and the same nucleotide deletion was observed at nt3481. Subsequently, DNAs from the remaining strains with retinal spotting similar to that associated with rd8 were tested by allele-specific PCR analysis and all strains harbored the same mutation. This finding supports the notion that the mutations observed in all of the incipient congenic strains were indeed caused by a C57BL/6 founder effect. The predicted mutant protein, if generated, would retain the EGF-like and LG-like domains, contain 47 novel amino acids following amino acid 1160 and end prior to the transmembrane domain of the CRB1 protein (Fig. 3D). From hereon, we will refer to the rd8 mutation as Crb1rd8.
The external limiting membrane is disrupted in Crb1<sup>rd8</sup> mutants

crb loss-of-function mutants in Drosophila do not establish normal ZAs in photoreceptors (7,8). To determine if loss of CRB1 in mouse eyes affects ELM/ZA integrity, a number of proteins that localize to the ELM were examined in Crb1<sup>rd8</sup>/Crb1<sup>rd8</sup> mice. As others have shown, antibodies against ZO-1, β-catenin, p120ctn, and pan-cadherin strongly stain the ELM of the neural retina of WT mice (Fig. 4A and D and data not shown) (10). Fragmented staining of the ELM with all markers was observed in Crb1<sup>rd8</sup>/Crb1<sup>rd8</sup> mice (Fig. 4B, C, E and F and data not shown), even in areas that were not affected by folds and pseudorosettes (Fig. 4C and F).

In order to determine when the abnormalities in the ELM/ZA were first observed, we also examined retinas from 2-week old mice. At this stage of retinal development, all of the retinal layers are present, although some remodeling is still occurring, and the PR IS are formed but the PR OS have yet to reach their adult length (11). Unlike the retinas of 4-week old Crb1<sup>rd8</sup> mice, in which the fragmented ELM was observed pan-retinally, in retinas of 2-week old mice only the posterior ELM was affected (Fig. 4H). A linear, uninterrupted ELM was observed peripherally in rd8 mice (Fig. 4G) and pan-retinally in WT 2-week old mice. Interestingly, the fragmentation and disorganization of the ELM appeared to be more severe in the 2- versus 4-week old retinas.

The aberrant staining pattern of the ELM was confirmed as a loss of adherens junctions by electron microscopy. Adherens junction complexes are considerable distances apart in Crb1<sup>rd8</sup>/Crb1<sup>rd8</sup> mice (Fig. 5B and D) in comparison with wild-type mice (Fig. 5A–C). A focal expansion of the extracellular space, which contains amorphous granular material, or of photoreceptor cell bodies is observed in regions where the ZA is absent in Crb1<sup>rd8</sup>/Crb1<sup>rd8</sup> mice (Fig. 5E). Furthermore, Müller cell apical processes were not observed immediately beneath the regions that did not contain adherens junctions, suggesting a developmental anomaly or retraction of the apical processes of the Müller cells from the ELM region. This correlated with absent or reduced staining of the Müller cell apical processes by antibodies against CD44 (Fig. 6E and F), an adhesion receptor that localizes only within these processes (10).

CRB1 localizes to Müller cells and photoreceptor inner segments

To determine the effect of the mutation of Crb1 on CRB1 protein level and localization in Crb1<sup>rd8</sup>/Crb1<sup>rd8</sup> mice, immunohistochemical studies were carried out. Using polyclonal antibodies raised against synthetic peptides from the extracellular domain of CRB1 (amino acids 423–437 and 586–602), we observed the highest intensity staining of the Müller cell radial processes in the inner nuclear layer (INL) and in apical processes sclerad to the ELM, and of photoreceptor inner segments (Fig. 6A). Our results differ from those of Pellikka et al. (7), who also reported staining in cone photoreceptor outer segments in wild-type mice. That nine of 10 peptide residues used to raise the polyclonal CRB1 antibody used by Pellikka et al. (7) were identical to a sequence from a related protein, CRB2, suggests the potential for cross-reactivity. In mice homozygous for the Crb1<sup>rd8</sup> mutation, CRB1 staining in the apical processes of the Müller cell was absent or diffuse (Fig. 6A and B); however, staining of the inner segment appeared the same in both mutant and wild-type retinas. This suggests that the transmembrane and/or cytosolic domain of Crb1 is important for the localization of CRB1 to the apical processes of the Müller cells. Consistent with this view, MPP5 (also known as PALS1), the mammalian homolog of the Drosophila gene, stardust (12,13), whose product has been shown to interact directly with Crumbs1 (14), was also absent or showed a diffuse staining pattern in the ELM region in rd8 mice (Fig. 6C and D). We note that the outer plexiform layer was brightly stained, and that the apparent localization and intensity of this staining was not affected in rd8 mutants.

Alternative splice variants of Crb1

The absence of CRB1 staining of the Müller cell apical processes in the Crb1<sup>rd8</sup> mutants, and the continued staining of IS, suggested the existence of splice variants that did contain...
the exon 6-encoded epitope for the polyclonal antibodies but not exon 9, which is mutated in Crb1rd8 mice. Database searches with the genomic Crb1 DNA sequence identified the ESTs BB642749 and AW491657, from a retinal and a pineal library, respectively, which were derived from a novel, shorter Crb1 splice variant (B in Fig. 7). This splice variant utilizes a novel 5' exon located in intron 5 which contains an ATG start codon with upstream in-frame stop codons. The putative amino terminal sequence is conserved in a human EST (BM687886, 76% similarity/53% identity over 17 amino acid residues), which also has upstream in-frame stop codons. This splice variant also utilizes an alternate polyadenylation site located in intron 11. The presence of this splice form in retinal mRNA was confirmed by sequencing of PCR products obtained from retinal cDNA using primers derived from the 5' non-coding region of exon 5a and intron 11 within the alternate 3' non-coding region of splice form B. The deduced amino acid sequence lacks the transmembrane domain of the full-length CRB1 splice form A, which is encoded by exon 12. Since the amino terminal sequence of splice form B also lacks a signal sequence, we predict that CRB1 isoform B is localized intracellularly. PCR amplification of retinal cDNA with combinations of primers obtained from all exon sequences yielded additional products that lack individual exons, indicating that additional splice variants may exist (Fig. 7C–G). However, all of the full length splice variants that we were able to amplify contained exon 9.

Figure 2. Retinal dysplasia and degeneration of photoreceptors in wild-type, control (A) and Crb1rd8/Crb1rd8 mice visualized by light microscopy (B–D) and by electron microscopy (E–H). (A) Retina from 4-week-old C57BL/6 mouse. (B) At 4 weeks, most of the retinal architecture is normal. There are occasional retinal folds, involving inner and outer nuclear layers (arrow). There is also focal thinning of the inner and outer segments (arrowhead). In the degenerated region, the outer nuclear layer is reduced to a single row of nuclei and both inner and outer segments are absent. (D) The region of focal degeneration at higher magnification shows a prominent loss of the photoreceptors, which in places are reduced to a single layer of nuclei (arrow). Near this area of severe degeneration, nearly normal retinal architecture is preserved including normal inner and outer segments (arrowhead). Hematoxylin and eosin. Magnification: (A) ×200; (B) ×200; (C) ×100; (D) ×400. (E) By electron microscopy, at 4 weeks of age a shortening of both inner and outer segments is noted. (F) At 10 weeks of age there is inner segment shortening and outer segment disorganization. In both D&E, the inner segments are approximately one-quarter to one-third normal length. (G) At 5 months of age there are only a few short fragments of outer segments (OS). Some inner segments (IS) appear normal while others (asterisk) are swollen. (H) The cytoplasm of Muller cells of the inner retina is more prominent in Crb1rd8/Crb1rd8 mice than normal (specimen from an 8-week-old mutant); two Muller cell processes are indicated by arrows. Inset: a Muller cell body (M) in the inner nuclear layer and its cytoplasmic processes (arrow) are also unusually prominent. (D) and (E) original magnification ×8100; (F) and (G) ×12 000; inset ×8100.
Figure 3. Genetic and molecular analysis of the rd8 mutation. (A) Haplotype analysis. Progeny from the (B6.C3Ga X CAST-rd8/+ )F1 X B6.C3Ga- rd8/rd8 or (B6.FVB X CAST- rd8+/+ )F1 X B6.FVB- rd8/rd8 back-cross were phenotyped for the retinal spotting phenotype and genotyped for the indicated microsatellite markers. Black boxes represent heterozygosity, B6 and CAST-derived alleles and white boxes represent homozygosity for the B6-derived allele. The number of chromosomes sharing the corresponding haplotype is indicated below each column of squares. The order of marker loci was determined by minimizing the numbers of crossovers. The genotype for rd8 was inferred from the phenotype or the results of progeny testing of non-informative recombinants.

(B) Chromosomal map position of rd8. A total of 606 BC progeny was typed with markers listed left of the vertical lines representing the region identified as containing rd8 on chromosome 1. The recombination frequencies are given in centimorgans ± SE. On the right hand of the vertical line is a transcript map of the rd8 minimal region. ‘n’ represents novel uncharacterized genes. (C) A deletion of nt 3481 of the Crb1 cDNA, a cytosine, is observed in rd8 mice but not wild-type controls. The arrow indicates the deleted nucleotide. (D) Schematic of protein truncations. Wild-type CRB1 protein contains a signal peptide (SP), 17 epidermal growth-factor-like domains, three laminin A G-like domains, and a transmembrane (TM) domain. If translated, the mutant mRNA in rd8 mice would produce a protein that does not contain the transmembrane or cytosolic domains of CRB1.
Suppression of retinal spotting and dysplasia observed in Crb1RD8/Crb1RD8 mutants

Observation of Crb1RD8 homozygotes from the mapping cross to the wild-derived CAST/EiJ mice revealed that the degree of the spotting phenotype is highly variable. While parental Crb1RD8/Crb1RD8 mice have a relatively uniform spotting phenotype in the inferior nasal quadrant of the eye, Crb1RD8 homozygotes on the segregating CAST/EiJ background have many more or fewer spots. In fact, we noted 19% of mice that were homozygous for the mutation in Crb1 did not exhibit the retinal spotting phenotype. In addition, during the process of introgressing the Crb1RD8 mutation onto the C57BL/6J background, the retinal spotting was no longer evident by indirect ophthalmoscopy at back-cross generation N7. Therefore, it appears that genetic modifiers exist that are able to suppress as well as enhance the degree of retinal spotting.

Retinal folds or pseudorosettes were not observed in histological sections of retinas from mice that were homozygous for the Crb1 mutation and suppressed for the clinical retinal spotting phenotype. Gross histological examination of retinas from suppressed Crb1RD8/Crb1RD8 mice was essentially normal without shortening and disorganization of the inner and outer segments. However, the anti-ZA markers tested on retinas of suppressed mice revealed a discontinuous, fragmented staining pattern similar to that observed in non-suppressed Crb1RD8/Crb1RD8 mice (Fig. 8), suggesting that retinal folding and photoreceptor inner segment shortening and disorganization are
secondary phenotypic changes that are controlled by genes other than Crb1.

Crossing suppressed B6.Cg-Crb<sup>rd8</sup>/Crb<sup>rd8</sup> mice (N10F3) with non-suppressed B6.C3Ga-Crb<sup>rd8</sup>/Crb<sup>rd8</sup> mice (N3F12), yielded offspring (n = 18) of which 100% were observed to have retinal spots. This suggests that the factor(s) necessary for the retinal dysplasia acts in a dominant fashion and is probably a contribution from C3HfB6/Ga (15).

**DISCUSSION**

The defects observed in Crb<sup>rd8</sup> mutant mice show striking similarities to those caused by crb mutations in the *Drosophila* retina (7,8). As in *Drosophila* crb mutants, the ELM/ZA is fragmented, and the ZA defects are more severe in the developing retina compared to the mature retina in Crb<sup>rd8</sup> mutants (7). In *Crb<sup>frd8</sup>/Crb<sup>frd8</sup> mice, the loss of ELM integrity was more notable at 2 than at 4 weeks, particularly in the posterior retina; this appears to be coincident with the posterior to peripheral maturation and migration of the photoreceptors (16). In the mature Crb<sup>frd8</sup> mutant retina, photoreceptor cell bodies protrude into the inner segment layer. Previous studies have suggested that the ELM provides a barrier function, as mice having a chemically induced deficiency of retinal Müller cells, which form ZAs with photoreceptors, exhibit mislocalized photoreceptor cell bodies in the inner and outer segment layers (17).

The Crb<sup>frd8</sup> mutation truncates the transmembrane and cytoplasmic domain of the CRB1 protein suggesting that it may produce a secreted protein that consists of most of the extracellular domain of CRB1. Overexpression of the extracellular domain of *Drosophila* Crb has a dominant-negative effect on the formation of the stalk membrane (7). We have observed retinal defects only in homozygous mutant Crb<sup>frd8</sup> animals but not in heterozygous animals. This may imply that either the secreted, extracellular domain of CRB1 has no dominant-negative effect, or that the expression from the endogenous CRB1 promoter is not producing sufficient gene product to cause phenotypic consequences. *Drosophila* Crb presumably plays mechanistically distinct roles in the formation of the ZA and the stalk membrane (7,8). Whether both of these roles have been conserved in the mammalian retina remains to be clarified. For example, current evidence is consistent with the view that the primary role of CRB1 is the maintenance of the ELM/ZA and that the shortening of the inner segment is a secondary consequence of the fragmented ELM/ZA observed in Crb<sup>frd8</sup> mutants. To identify conserved and potentially novel roles of CRB1 in the mammalian retina, it will be important to characterize the network of factors that interact with CRB1 in retinal development.

A recent publication, in which retinas of individuals diagnosed with LCA and harboring CRB1 mutations were examined by optical coherence tomography, reports thickened retinas with abnormal lamination and photoreceptor rosettes (18). Although we observed retinal folds and pseudorosettes in B6.C3Ga-Crb<sup>frd8</sup>/Crb<sup>frd8</sup> mice, the retinal architecture was, in general, normal. Only within the dysplastic regions were abnormalities in lamination, involving both the INL and ONL, observed. The discrepancy in phenotypic features observed in the retina between mice and humans could be potentially explained by allelic or by species differences. The former explanation may be less likely given the fact that the thickening and abnormal lamination of the retina were observed in individuals with mutations throughout *Crb1*, including the region predicted to be truncated in the rd8 mutant (Fig. 9) (18). Previous reports, showing species differences in retinal disease manifestation have been reported, for example, in shaker mice, which harbor a mutation within myosin VIIA (19). In humans, mutations within myosin VIIA lead to both hearing and vision loss in patients with Usher Syndrome Type 1B (20), while shaker mice only develop hearing loss. This is presumably due to species differences in the localization of the protein within the retina (21,22). It is also possible that the difference in clinical features found between humans and our mouse model, and the relatively mild retinal disease observed in *Crb<sup>frd8</sup>* mutant mice, may be due to the genetic background in which the *Crb<sup>frd8</sup>* mutation currently resides.
Perhaps the most interesting observation of Crb1rd8/Crb1rd8 mice, which sheds light on the variability in clinical features observed in individuals with CRB1 mutations, is that genetic modifiers are able to determine the severity of the disease phenotype. Den Hollander et al. (2) suggested that the wide range of phenotypic characteristics observed in individuals with CRB1 mutations could be due to environmental or genetic factors. We provide evidence that genetic factors do indeed influence the disease phenotype in the presence of a Crb1rd8 mutation. The fact that retinal degeneration is not observed in Crb1rd8/Crb1rd8 mice suppressed for retinal dysplasia suggests that, in mice, Crb1 may be thought of as a susceptibility locus whose mutant form must interact with mutant/variant alleles at other loci for disease to manifest. Additional studies using different inbred strains may help to identify factors contributing to the retinal dysplasia and to clarify the apparent differences in the presentation of retinal histology in humans and mice carrying Crb1 mutations. Also, such genetic modifier screens may further define the molecules involved in the function of CRB1 and should identify candidate retinal disease genes.

MATERIALS AND METHODS

Mice

Mice were bred and maintained under standard conditions in the Research Animal Facility at The Jackson Laboratory. They were maintained on NIH 4% fat chow and acidified water, with a 12:12 hour dark:light cycle in facilities that are monitored regularly to maintain a specific pathogen-free environment. Procedures used in the experiments were approved by the Institutional Animal Care and Use Committee. B6.C3Ga-rd8/rd8 (N3F12) and B6.FVB-rd8/rd8 (N4F9) were used in the initial mapping crosses with CAST/EiJ mice. B6.Cg-rd7/+;rd8/rd8 (N10F3) and B6.C3Ga-rd8/rd8 (N3F12) were used for the suppressor and phenotypic studies.

Chromosomal localization and fine structure mapping

Tail DNA was isolated from mice generated by backcrosses described above according to Buffone and Darlington (23). DNAs of the 606 BC offspring were genotyped using microsatellite markers to develop a fine structure map of the region. Critical recombinant mice were progeny tested by crossing them to F2(B6.C3GaXCAST)rd8/rd8 mice to confirm informative recombinants or to determine if a presumed uninformative recombinant carried the disease gene. A minimum of 20 offspring from each progeny test were genotyped and phenotyped. For PCR amplification, 25 ng DNA was used in a 10 μl volume containing 50 mM KCl, 10 mM Tris-Cl, pH 8.3, 2.5 mM MgCl2, 0.2 mM dNTP, and 0.02 U AmpliTaq DNA polymerase. The reactions which were initially denatured for 2 min at 95°C were subjected to 49 cycles of 20 s at 94°C, 20 s at 50°C, 30 s at 72°C and a 7 min extension at 72°C. PCR products were separated by electrophoresis on a 4% MetaPhor (FMC, Rockland, ME, USA) agarose gel and visualized under UV light after staining with ethidium bromide.

Clinical examination of eyes

Mice were dark adapted and their pupils dilated with atropine prior to examination by indirect ophthalmoscopy with a 78 or 90 diopter aspheric lens. Fundus photographs were taken with a Kowa fundus camera using a Volk superfield lens held two inches from the eye. The highest flash intensity was used with 400ASA film for the photodocumentation.
Histological analysis

Light microscopy. Eyes were oriented by marking the cornea by cauteryization, after sacrifice but prior to enucleation, and were placed in Bouin’s fixative or acetic acid: methanol (1:3) overnight, embedded in paraffin, and sectioned in a plane that included the ora serrata and optic nerve. Sections were stained with hematoxylin and eosin.

Electron microscopy. The eyes were removed immediately after cardiac tamponade euthanasia of each mouse and fixed for 3 h in a cold, phosphate-buffered, glutaraldehyde–parafomaldehyde solution. After 3 h, the anterior segment was removed and the posterior segment cut into 1×2 mm blocks of retina, choroid and sclera. The additional fixation of the whole eye before section improved adhesion of the retina to the retinal pigment epithelium (RPE) and did not alter the quality of preservation. The dissected tissue was placed in fresh fixative for an additional 2–8 h and was post-fixed in 1% osmium tetroxide, dehydrated and embedded in plastic. Thick sections were cut for orientation and thin sections cut and stained with uranyl acetate and lead citrate and examined using a transmission electron microscope.

Immunohistochemistry. Mice homozygous for the rd8 mutation and wild-type controls were sacrificed by cervical dislocation or carbon dioxide asphyxiation. Enucleated, oriented eyes were fixed with either acetic acid:methanol (1:3) or 4% paraformaldehyde (PFA) in phosphate-buffered saline (PBS) overnight prior to embedding in paraffin. For mutants and for controls, a minimum of three eyes, each from a different animal, were tested with each primary antibody. Sections of 6 μm thickness were cut and mounted on slides pretreated with Vectabond (Vector Laboratories, Burlingame, CA, USA). After blocking with 2% normal horse serum in PBS, sections were incubated overnight with primary polyclonal antibodies for zonula adhaerens-1 (1:200; Sigma-Genosys, The Woodlands, TX, USA). Polyclonal antibodies against PALS1 were provided by Dr. B. Margolis (14) and used at a 1:200 dilution. Secondary antibodies were biotin-conjugated anti-rabbit IgG (1:200, Vector Laboratories) for zonula occludens-1, CRB1, CRB2, and PALS1, biotin-conjugated anti-goat IgG (1:200, Vector Laboratories) for p120 catenin, biotin-conjugated anti-mouse IgG (1:200 Vector Laboratories) for pan-cadherin and β-catenin, and biotin-conjugated anti-rat IgG (1:200, Vector Laboratories) for CD44. Binding was detected using FITC-Avidin D (1:200, Vector Laboratories). Nuclear counterstaining was performed with 4,6 diamidine 2-phenylindole dihydrochloride (DAPI) at a final concentration of 5 μg/ml. Images were collected on a Leica DMRXE fluorescent microscope (Leica, Deerfield, IL, USA) equipped with a SPOT™ CCD camera (Diagnostic Instruments, Sterling Heights, MI, USA) using appropriate band-pass filters for each fluorochrome, or on a Leica TCS NT confocal microscope (Leica, Deerfield, IL, USA) using a 40×0.85 NA lens or a 100×1.25 NA oil immersion lens. Appropriate band-pass filters for each fluorochrome were used and image collection was optimized to fill the 256 gray levels available.

Generation of the CRB1 and CRB2 polyclonal peptide antibodies

Polyclonal CRB1 anti-peptide antibodies were generated in rabbit by Peptide Specialty Laboratories GmbH (Heidelberg, Germany). Peptides corresponding to amino acids 423–437 (PFDDTSRTFYGGENGCP) and 586–602 (CKEKCTTKSSVPV-ENHQ) were synthesized, coupled to KLH and used to immunize two rabbits. Sera were tested by ELISA after three immunizations, with positive results. After collection, aliquots of sera were affinity-purified against both peptides conjugated to SulfoLink Coupling Gel™, according to the manufacturer’s protocols (Pierce Biotechnology, Rockford, IL, USA). Preimmune sera, immune sera, and affinity-purified sera were all tested by immunocytochemistry against target tissue. Preimmune sera showed no staining of tissue.

Anti-CRB2 polyclonal antibody were generated in a rabbit immunized with a CRB2 peptide (aa941-955) coupled to keyhole limpet hemocyanin and was peptide affinity purified (Sigma-Genosys, The Woodlands, TX, USA).

Mutation analysis of Crb1 and identification of alternative splice factors

Total RNA was isolated from whole eyes of B6.C3Ga-rd8/rd8, B6.FVB-rd8/rd8, B6.J-rd8/rd8, and C57BL/6J mice. Tissues were homogenized and RNA was isolated by TRizol (Life Technologies) treatment according to the manufacturer’s protocol. cDNA was generated using the Retroscript kit (Ambion). Primers to detect splice variants of Crb1 were designed from exons or EST sequences obtained from GenBank or Celera databases. PCR assays were carried out using the Expand Template system (Roche) and gel purified amplicons were sequenced with an ABI Prism 3700. Allele-specific PCR (24) was used on genomic DNA to confirm the presence of the rd8 mutation using the primers: mCrb1-mF1 GTGAAGACAGCTACAGTTCTGATC; mCrb1-mF2 GCCCCTGTTTGCATGGAGGAACTTGGA-AGACAGCTACAGTTCTGCGG; and mCrb1-mR GCCCCCATTTGCACTGATGAC.

Figure 8. Genetic modification of the retinal phenotype in mice homozygous for Crb1rd8 and segregating on a C57BL/6 and C3H genetic background. (A) Suppressed Crb1rd8/Crb1rd8 mice do not exhibit retinal folds or pseudosettes. Even at 11.5 months, the general architecture of the suppressed rd8 retina is normal. The shortening of the inner and outer segments observed in Crb1rd8/Crb1rd8 mice is not evident in suppressed mice. Staining of the ELM with anti-adherens junction markers of suppressed Crb1rd8/Crb1rd8 mice (B) and age-matched wild-type controls (C) shows that the adherens junctions are discontinuous and fragmented, but not as severely as observed in non-suppressed Crb1rd8/Crb1rd8 mice.
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REFERENCES


