BEST1 Gene Therapy Corrects a Diffuse Retina-Wide Microdetachment Modulated by Light Exposure

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Guziewicz, K. E., Cideciyan, A. V., Beltran, W. A., Komáromy, A. M., Dufour, V. L., Swider, M., Iwabe, S., Sumaroka, A., Kendrick, B. T., Ruthel, G., Chiodo, V. A., Heon, E., Hauswirth, W. W., Jacobson, S. G., & Aguirre, G. D. (2018). BEST1 Gene Therapy Corrects a Diffuse Retina-Wide Microdetachment Modulated by Light Exposure. *Proceedings of the National Academy of Sciences, 115* (12), E2839-E2848. http://dx.doi.org/10.1073/pnas.1720662115

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Abstract
Mutations in the BEST1 gene cause detachment of the retina and degeneration of photoreceptor (PR) cells due to a primary channelopathy in the neighboring retinal pigment epithelium (RPE) cells. The pathophysiology of the interaction between RPE and PR cells preceding the formation of retinal detachment remains not well-understood. Our studies of molecular pathology in the canine BEST1 disease model revealed retina-wide abnormalities at the RPE-PR interface associated with defects in the RPE microvillar ensheathment and a cone PR-associated insoluble interphotoreceptor matrix. In vivo imaging demonstrated a retina-wide RPE-PR microdetachment, which contracted with dark adaptation and expanded upon exposure to a moderate intensity of light. Subretinal BEST1 gene augmentation therapy using adeno-associated virus 2 reversed not only clinically detectable subretinal lesions but also the diffuse microdetachments. Immunohistochemical analyses showed correction of the structural alterations at the RPE-PR interface in areas with BEST1 transgene expression. Successful treatment effects were demonstrated in three different canine BEST1 genotypes with vector titers in the 0.1-to-5E11 vector genomes per mL range. Patients with biallelic BEST1 mutations exhibited large regions of retinal lamination defects, severe PR sensitivity loss, and slowing of the retinoid cycle. Human translation of canine BEST1 gene therapy success in reversal of macro- and microdetachments through restoration of cytoarchitecture at the RPE-PR interface has promise to result in improved visual function and prevent disease progression in patients affected with bestrophinopathies.

Keywords
AAV BEST1 gene therapy, RPE bestrophinopathy, microvilli, photoreceptors

Disciplines
Medicine and Health Sciences | Veterinary Medicine

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**BEST1** gene therapy corrects a diffuse retina-wide microdetachment modulated by light exposure

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Mutations in the **BEST1** gene cause detachment of the retina and degeneration of photoreceptor (PR) cells due to a primary channellopathy in the neighboring retinal pigment epithelium (RPE) cells. The pathophysiology of the interaction between RPE and PR cells preceding the formation of retinal detachment remains not well-understood. Our studies of molecular pathology in the canine **BEST1** disease model revealed retina-wide abnormalities at the RPE-PR interface associated with defects in the RPE microvillar ensheathment and a cone PR-associated insoluble interphotoreceptor matrix. In vivo imaging demonstrated a retina-wide RPE-PR microdetachment, which contracted with dark adaptation and expanded upon exposure to a moderate intensity of light. Subretinal **BEST1** gene augmentation therapy using adeno-associated virus 2 reversed not only clinically detectable subretinal lesions but also the diffuse microdetachments. Immunohistochemical analyses showed correction of the structural alterations at the RPE-PR interface in areas with **BEST1** transgene expression. Successful treatment effects were demonstrated in three different canine **BEST1** genotypes with vector titers in the 0.1-to-5E11 vector genomes per mL range. Patients with biallelic **BEST1** mutations exhibited large regions of retinal lamination defects, severe PR sensitivity loss, and slowing of the retinoid cycle. Human translation of canine **BEST1** gene therapy success in reversal of macro- and microdetachments through restoration of cytoarchitecture at the RPE-PR interface has promise to result in improved visual function and prevent disease progression in patients affected with bestrophinopathies.

Inherited retinal degenerations (IRDs) encompass a large group of blinding conditions that are molecularly heterogeneous and pathophysiologically distinct (1, 2). The genetic defect often acts primarily on rod or cone photoreceptors (PRs), or both, and the specific defect may involve phototransduction, ciliary transport, morphogenesis, neurotransmission, or others. Less common are primary defects involving the retinal pigment epithelium (RPE), although they have received increased attention due to high-profile clinical trials (3–5).

The most common IRD due to a primary RPE defect is caused by mutations in **BEST1** (6–8), encoding a transmembrane protein associated with the basolateral portion of the RPE (9–11). **BEST1** acts as a multifunctional channel protein responsible for mediating transepithelial ion transport, regulation of intracellular calcium signaling and RPE cell volume, and modulation of the homeostatic milieu in the subretinal space (10–17). Additional insights supporting **BEST1** function as a calcium-modulated anion channel came from recent crystallographic studies, which revealed a similar structural architecture between avian (18) and bacterial (19) homologs. In eukaryotic cells, **BEST1** forms a stable homopentamer with four transmembrane helices, cystosolic N and C termini, and a continuous central pore sensitive to calcium-dependent control of chloride permeation (18). Further analysis of **BEST1** structure with targeted mutations has begun to uncover the specific mechanisms responsible for ion selectivity and calcium-mediated activation of this channel protein (20).

In humans, **BEST1** mutations result in a wide spectrum of IRDs collectively grouped as bestrophinopathies that often involve pathognomonic macular lesions (6, 7, 21). Retinal regions away from the lesions tend to appear grossly normal, despite the existence of a retina-wide electrophysiological defect in the electrooculogram, which reflects an abnormality in the standing potential of the eye (21, 22). Naturally occurring biallelic mutations in the canine **BEST1** gene (c**BEST1**) cause canine IRD with distinct phenotypic similarities to both the dominant and recessive forms of human bestrophinopathies, including the

Significance

One of the most common forms of monogenic macular degeneration worldwide is caused by dominant or recessive bestrophinopathies associated with mutations in the **BEST1** gene. Disease expression is known to start with a retina-wide electrophysiological defect leading to localized vitelliform and atrophic lesions and vision loss. To develop lasting therapies for this incurable disease, there is a need for greater understanding of the early pathophysiology before lesion formation. Here we find that the loss of retinal pigment epithelium apical microvilli and resulting microdetachment of the retina represent the earliest features of canine bestrophinopathies. We show that retinal light exposure expands, and dark adaptation contracts, the microdetachments. Subretinal adeno-associated virus-based gene therapy corrects both the vitelliform lesions and the light-modulated microdetachments.

Author contributions: K.E.G., A.V.C., S.G.J., and G.D.A. designed research; K.E.G., A.V.C., W.A.B., A.M.X., V.L.D., S.J., B.T.K., G.R., V.A.C., E.H., W.W.H., S.G.J., and G.D.A. performed research; A.M.K., A.S., and V.A.C. contributed new reagents/analytic tools; K.E.G., A.V.C., W.A.B., V.L.D., M.S., A.S., G.R., and G.D.A. analyzed data; and K.E.G., A.V.C., S.G.J., and G.D.A. wrote the paper.

Conflict of interest statement: W.W.H. and the University of Florida have a financial interest in the use of AAV therapies, and own equity in a company (AGTC, Inc.) that might, in the future, commercialize some aspects of this work.

This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1720662115/-/DCSupplemental.

Published online March 5, 2018.
salient predilection of subretinal lesions to the canine fovea-like area (17, 23, 24).

Proper anatomical apposition and a sustained interaction between RPE apical microvilli (MV) and PR outer segments (OSs) are considered crucial for normal vision. Both the intensity by position and volume regulation of the subretinal space are essential for maintaining the accurate molecular proximity of this complex and homeostasis of the RPE–PR interface (25, 26). In vitro and ex vivo studies have long shown that genetic mutations, metabolic perturbations, as well as light stimuli alter the ion composition of the subretinal space and physiological responses of the RPE and/or PRs (27–29). More recently, in vivo studies of outer retinal microanatomy in health and disease and its response to light have become increasingly informative with more refined imaging modalities (30–40). Here, we combined ex vivo and in vivo approaches to discover and quantify an abnormal modulation of the RPE–PR distance by light exposure in canine bestrophinopathy (cBest). We demonstrate that this is a retina-wide abnormality, and show that adeno-associated virus (AAV2)-mediated BEST1 gene augmentation corrects this primary subclinical defect as well as the disease.

Results

Early Retina-Wide Pathology at the RPE–PR Interface. To understand the pathophysiology behind the impaired RPE–PR interaction, we first evaluated cBest retinas with clinically obvious disease. The key features of RPE apical membrane responsible for direct interaction with PR OSs were examined by immunohistochemistry (IHC) against EZRIN, a membrane–cytoskeleton linker protein essential for formation of RPE apical MV, and combined with human cone arrestin (hCAR) and peanut agglutinin lectin (PNA) labeling to distinguish the cone PR matrix-specific interface (Fig. 1A). Confocal microscopy and analysis of 3D reconstruction images from the wild-type (WT) retina exposed a complex sheet-like structure of both inherent constituents of RPE apical membrane—cone- and rod-associated MV. Cone-MV (also known as RPE apical cone sheath) (41) were more pronounced than rod-MV, and formed a highly organized wrapping that tethered individual cone outer segments (COSs) to the RPE apical surface (Fig. 1A and Movie S1). In the subretinal space, this intercellular complex was further encased with an equally intricate cone-specific insoluble extracellular matrix sheet (cone-IPM) detected by selective binding of PNA lectin (42) (Fig. 1A and Movie S1). In disease cBest retinas, however, such complex extracellular compartmentalization of COSs was lost, and the cone apical membrane directly contacted the RPE apical surface (Fig. 1A, arrow, and Movie S1). These observations were confirmed in three distinct cBEST1 genotypes (R25*/R25*, P463fs/P463fs, and R25*/P463fs) examined across both the tapetal and nontapetal portions of the retina in 22 eyes after disease onset (age range 45 to 270 wk).

To evaluate the possibility that the structural cone-MV abnormalities are secondary to cone dysfunction and disease, we examined the RPE–COS interaction in a different canine IRD model: a primary cone photoreceptor channelopathy, CNGB3-associated achromatopsia (43). We first examined the RPE–COS complex at 6 wk of age; CNGB3-mutant retinas, harboring either a missense or locus deletion mutation (43), showed no apparent irregularities at the RPE–PR interface, and the proper localization of RPE apical markers was associated with specific anti-BEST1 labeling (Fig. S1A and B). Double immunostaining demonstrated specific distribution of EZRIN along cone-MV interdigitating with hCAR-positive yet CNGB3-negative COSs (Fig. S1A and B). Then we studied the consequences of CNGB3 channel subunit dysfunction in older (ages 57 and 85 wk) mutant retinas, which undergo a gradual cone PR degeneration (44), and found that the microvillar ensheathment of the RPE apical domain still remained largely intact (Fig. S1C and D).

The findings in CNGB3-mutant retinas suggested that the structural alterations associated with cone-MV ensheathment in cBest were not secondary to a cone defect but specific to the RPE channelopathy triggered by mutations in BEST1. We then focused on the 6-wk time point, which is well before clinical disease onset and near the end of postnatal retinal differentiation in dogs (45) (Fig. 1C and D). Hyperscattering in the age-matched WT control, the lack of specific basolateral BEST1 immuno-labeling in the cBest RPE was associated with a rather smooth apical surface and clearly underdeveloped (vestigial) apical microvilli (Fig. 1C, arrowheads). Quantification of the spatial density of cone-MV and the length of cone- and rod-MV was performed on deconvolved 3D z-stack projection images at four retinal locations (Fig. 1D). Significant differences (P < 0.0001) in the mean number of cone-MV were found between cBest and WT in each retinal region examined (Fig. 1D). Even though the cone photoreceptor numbers were comparable to controls, the cone-MV in cBest were much fewer in number, sparsely distributed, and consistently appeared shorter and much finer than those in controls regardless of the topographical location. In control (WT) eyes, the average length of cone-MV was 17.4 (±0.25) μm in the tapetal superotemporal quadrant and 12.3 (±0.23) μm in the inferior nontapetal retina, whereas the length of rod-MV was 6.7 (±0.11) μm and 5.3 (±0.27) μm in the tapetal and nontapetal portions of the retina, respectively. In cBest, however, the average length of rare cone-MV extensions identified was substantially reduced (6.0 ± 0.31 and 6.5 ± 0.74 μm in the central tapetal and nontapetal inferior parts, respectively). A quantitative assessment of the minute rod-MV in cBest was beyond the limits of optical resolution.

cBEST1-Mutant Eyes Have Retina-Wide Microdetalchments That Expand with Light Exposure. To determine in vivo correlates of the early RPE–PR interface abnormalities detected by IHC, we used non-invasive imaging with optical coherence tomography (OCT) to evaluate cBest eyes at young ages, well before ophthalmoscopic lesions are detectable. Qualitatively, central retinas of all evaluated eyes showed an additional hyperscattering layer in the outer retina located distal to the outer nuclear layer (ONL) that was not detectable in WT eyes (Fig. 2A, arrow and double arrow). Unexpectedly, the hyperscattering layer was variable with repeated recordings in the same eye within a single experimental session. Further analyses uncovered that the width of the hyperscattering layer was greater in scans obtained toward the end of an imaging session, when the retina would have been exposed to greater retinal irradiance due to intervening autofluorescence imaging performed with bright short-wavelength lights (Fig. 2A, double arrow, and Fig. S2A). The width of the hyperscattering layer was less in scans obtained early in the imaging session before autofluorescence imaging was performed (Fig. 2A, arrow, less LA). Quantitative studies were performed by obtaining longitudinal reflectivity profiles and making measurements both at nasal and temporal retinal locations. WT eyes (n = 12, ages 15 to 17 wk) showed outer retinal hyperscattering peaks at the outer plexiform layer (OPL) and the external limiting membrane (ELM) defining the intervening hyperscattering layer as the ONL (Fig. 2B). Distal to the ELM was a hyperscattering peak corresponding to the junction between inner and outer segments of photoreceptors (IS/OS), a major peak originating near the RPE–tapetum interface (RPE/T), and an intervening minor hyperscattering peak corresponding to photoreceptor OS tips, which was often difficult to resolve (Fig. 2B). An abnormal hyperscattering layer (Fig. 2B, arrows) was detectable in cBest eyes (n = 6, age 11 wk) with less light exposure. With greater light exposure, the hyperscattering layer became deeper and more distinct (Fig. 2B, double arrows); both nasal and temporal retinal locations showed the same effect. The distance between the IS/OS and RPE/T peaks (Fig. 2A and B, arrowheads) was measured. In WT eyes, the distance was 41.3 (±4.5) μm, whereas in cBest eyes this distance was significantly greater (P < 0.001) at 46.8 (±6.7) μm and 45.2 (±6.8) μm (less light exposure) and 55.5 (±10.5) μm and 53.5 (±6.3) μm (more light exposure) for nasal and temporal retinal regions, respectively (Fig. 2C).
Two types of experiments were designed to better understand the thickness of the hyposcattering layer as a function of light exposure. In the main experiment (WT, n = 12, age 15 to 17 wk; cBest, n = 3, age 13 wk), eyes were dark-adapted overnight and then sequential imaging was performed in the dark over a 2-h period with five intervening brief 488-nm light exposures of incrementally greater intensity ranging from very dim lights to moderate lights produced by standard clinical ophthalmic equipment (Fig. 2D). In a shorter experimental protocol, only the highest two light exposures were used in different eyes (cBest, n = 3, age 13 wk). After overnight dark adaptation, IS/OS–RPE/T distance was 40.0 (±4.5) μm in WT eyes, whereas it was 47.1 (±4.8) μm in cBest (Fig. 2E); the difference was statistically significant (P < 0.001). Increasingly brighter light exposures resulted in monotonic expansion of the IS/OS–RPE/T distance in cBest eyes, reaching an apparent plateau of 59.4 (±8.7) μm (Fig. 2E). In WT eyes, the effect of the light exposure was either negligible or small, with the IS/OS–RPE/T distance reaching a plateau of 40.9 (±4.3) μm. Thus, exposure to light appeared to cause an acute retinal microdetachment of up to 18.4 (±6.3) μm in cBest eyes within minutes at an age preceding any detectable ophthalmoscopic findings. The light-mediated microdetachment disappeared over a time span of less than 24 h (Fig. S2).

In preparation for localized gene therapy, retinotopic distribution of light-driven microdetachments was evaluated in fully light-adapted cBest and WT eyes (Fig. 2F). The mean IS/OS–RPE/T distance across WT eyes (n = 4, age 104 wk) was relatively homogeneous across superior and inferior retinal areas, with a clear boundary corresponding to the transition between tapetal and pigmented (nontapetal) retina. Greater distance in the tapetal retina of WT eyes was likely due to differences in the dominant contributors to the hyposcattering peak (tapetum in the tapetal retina versus pigmented RPE). In a cBest eye (R25*/P463fs) at age 297 wk, there was a relatively diffuse retinal-wide microdetachment in addition to grossly obvious retinal detachment at the fovea-like region (Fig. 2F, demarcated with purple). In a younger cBest eye (P463fs/P463fs) at age 12 wk, there was a distinct band of greater microdetachment along the visual streak and surrounding the optic nerve head even though no ophthalmoscopic abnormalities were evident. Difference maps between mutant eyes and mean WT demonstrated the spatial distribution of the extent of microdetachments (Fig. 2F, Right).

To assess the potential adverse consequences on photoreceptors, ONL thickness was topographically mapped across the retinal areas with microdetachments (Fig. S3). The microdetachments did not result in thinning of the ONL that would be expected from photoreceptor degeneration. Instead, there was a tendency for the ONL in cBest to be homogeneously thicker than WT; hyperthick regions typically included the central-superior tapetal retina, but could also extend into the inferior nontapetal retina (Fig. S3A). Of importance, the hyperthick regions of the ONL, when examined microscopically, had numbers of PR nuclei that were comparable to controls. This suggests an expansion of internuclear spacing as the likely cause of hyperthick ONL observed by imaging.

**Natural History of Canine Bestrophinopathy.** As a prerequisite to assessing gene therapy outcomes, the natural history of cBest was determined from a group of 18 dogs [12 male (M) and 6 female...
subtle focal retinal elevation of the canine fovea-like regions (Fig. S4A). This discrete RPE–PR detachment on cross-sectional imaging, yet unnoticeable on en face imaging, was found to be consistent among cBest eyes examined (n = 34), regardless of genotype. From the subclinical stage, the disease progressed to form a macrodetachment (vitelliform stage) limited to the canine fovea and surrounded by microdetachment (Fig. 3A, Left). The primary lesion gradually evolved to manifest as a characteristic bullous detachment within the area centralis that encompassed the fovea-like region (Fig. 3A, Middle and Right and Fig. S4 B–D). The presence of distinctive hyperautofluorescence was evident in the inferior part of the lesion (Fig. 3A, Middle Insets; pseudohypopyon stage). The advanced disease stages involved a partial resorption and dispersion of the hyperautofluorescent material within the central retina, associated with significant thinning of ONL (Fig. S4 E and F).

In each case followed by serial imaging (Table S1), cBest manifested bilaterally and nearly always presented a remarkable symmetry, albeit with a variable rate of progression (Fig. 3A and Fig. S4). The gross retinal detachments, ophthalmoscopically visible in both eyes, either remained limited to the central retina or became more extensive with extraretinal lesions scattered throughout, still with a strong predilection to the central cone-rich areas and associated with hyperthick ONL.

Subretinal BEST1 Gene Augmentation Therapy Stably Corrects Disease. To evaluate proof of concept for AAV2-mediated subretinal gene augmentation therapy, 22 cBest eyes were injected (vector titers in the 0.1 to 5E11 vector genomes (vg) per mL range or balanced salt solution (BSS) control) at 27 to 69 wk (Table S1) with a canine (cBEST1) or human (hBEST1) transgene driven by the human VMD2 promoter (46). cBest dogs exhibiting different stages of focal or multifocal retinal detachments were either injected unilaterally with AAV leaving the fellow eye un.injected, or with AAV in one eye and BSS in the contralateral eye. Eyes injected at 4 wk formed a partial retinal detachment, while the retinal areas outside of the surgical bleb served as an internal control (Table S1).

A representative result is shown from a compound heterozygous (R25*/P463fs) dog displaying advanced central retinal detachment in the right eye (EM56-OD) (Fig. 3A) that underwent subretinal injection with cBEST1 at the age of 52 wk (Fig. 3B, Left), while the fellow eye was not injected (EM56-OS) (Fig. S4A). Both eyes were examined clinically and by in vivo imaging. Disease reversals were first apparent in the injected eye (excellent effect) postinjection (p.i.) and retained a sustained effect long-term, as illustrated at 43 and 245 wk p.i. (Fig. 3B). In this and other cases with advanced disease that presented extensive accumulation of autofluorescent material within the subretinal macrodetachments (n = 13 eyes), hyperautofluorescent signals were still detectable for several months post AAV injection but gradually faded over time (Fig. 3B, Insets). Based on noninvasive imaging, both the focal as well as extracentral lesions within the AAV-BEST1–treated regions resolved 4 to 12 wk p.i., and localized retinal reattachements remained stable thereafter (Table S1). There was no evidence of inflammatory responses in any of the AAV-treated eyes, and the longitudinal in vivo evaluations did not reveal adverse effects on the RPE or neural retina.

AAV-mediated treatment with the hBEST1 also resulted in lesion reversal and long-term disease correction (n = 13 eyes). Representative in vivo imaging results and IHC evaluations (Fig. 3 C and D) from a cBest dog (R25*/R25*) showed that the early bilateral lesions present before treatment disappeared after the study eye (EMC3-OS) was treated with AAV-hBEST1 (2E11 vg/mL) (Fig. 3D), while the lesion in the contralateral control eye (EMC3-OD) injected with BSS continued to enlarge (Fig. 3C). Based on funduscopic examination, in the illustrated example as well as in all other cases the transient retinal detachment associated with vector or BSS delivery resolved within 24 to 48 h p.i.; however, the retinal lesions injected with BSS reappeared as early

Fig. 2. Light-mediated changes in the outer retinal structure. (A) Cross-sectional imaging along the horizontal meridian through the central area centrals (fovea-like area) in a 15-wk-old normal (WT) dog (Left) and an 11-wk-old cBest (R25*/P463fs) with less (Middle) and more (Right) light adaptation (LA). (Insets) En face reflectance images; the thin white arrows show the superotemporal location of the OCT. The arrowheads and single and double arrows are described in B. (B) Longitudinal reflectivity profiles (LRPs) (average of 85 single LRPs at 3° nasal from the fovea-like area (T, temporal retina) and nasal edge of the optic nerve head (N, nasal retina) in WT dogs (12 eyes, age 15 to 17 wk) compared with cBest (6 eyes, age 11 wk) with less and more LA. Arrowheads mark IS/OS and RPE/T peaks; single and double arrows mark the additional hyperreflective layer in cBest. (C) Distance between IS/OS and RPE/T peaks in WT and cBest eyes under two LA conditions. Symbols with error bars represent mean ± 2 SD distance for each group of eyes at both locations. (D, Upper) Schematic description of the dark and light adaptation protocol. Animals were dark-adapted (D/A) overnight and OCT imaging was performed. Then, five increasing light exposures (L1 through L5) were used. Following each light exposure, OCT imaging was repeated. (D, Lower) Magnified views of the OCT scans in cBest with overlapping LRPs after overnight dark adaptation (Left) and after the maximum light exposure (Right). Arrowheads and single and double arrows are described in B. (E, Left) Distance between IS/OS and RPE/T peaks after overnight dark adaptation and following increasing exposures to light (L1 to L5, 488 nm; vertical lines) in cBest (n = 3; colored traces represent individual eyes) and WT eyes (gray traces). (E, Right) In a different subset of cBest eyes (n = 3; colored traces), results of using an abbreviated protocol involving only L4 and L5 exposures. (F) Spatial topography of IS/OS-to-RPE/T distance in mean WT compared with two representative cBest eyes [pseudocolor panels; EM356-OS: 297-wk-old cmr1cmr3 (R25*/P463fs); LH30-OD: 12-wk-old cmr3 (P463fs/P463fs)]. Purple indicates gross retinal detachment. Grayscale panels show the point-by-point difference between each cBest eye and the mean WT.
as 1 wk p.i., and progressed along the natural disease course (Fig. 3C). This was in sharp contrast to the AAV-treated eyes, where both the early as well as more advanced lesions resolved within the first 6 wk after hBEST1 gene therapy, and the treated areas thereafter remained disease-free (Fig. 3D). Ophthalmological examinations and IHC assessments using RPE- and PR-specific markers showed no adverse effects on the retina up to 207 wk p.i. (Fig. 3 and Figs. S5 and S6). Of particular importance, the assessments of the retinal preservation p.i. revealed a remarkable restoration of retinal architecture at the RPE–PR interface, including extension of cone-MV and actin cytoskeleton rescue, corresponding to the vector-treated bleb area with either the canine or human BEST1 transgenes (Fig. 3 C and D, Lower and Figs. S5 and S6). No differences in the clinical picture or response to the AAV-BEST1 treatment were observed between genders.

Correction of Light-Modulated Microdetachments with Gene Therapy.

To understand the consequences of BEST1 gene augmentation therapy on retinal regions without ophthalmoscopically detectable retinal detachments, IS/OS–RPE/T distance was measured topographically within and outside subretinal blebs (Fig. 4). A representative result with a control subretinal BSS injection at age 69 wk in a cBest (R25*/P463fs) dog showed homogeneous microdetachment covering all the imaged retina at age 87 wk (Fig. 4A). The average microdetachment extent (IS/OS–RPE/T distance of the BSS-injected mutant dog subtracted from colocalized measurements performed in WT eyes) was 11.6 μm in the superior retina and 16.7 μm in the inferior retina (Fig. 4B), consistent with uninjected cBest eyes. Subretinal AAV gene therapy, on the other hand, resulted in substantial reduction of the IS/OS–RPE/T distance in treated regions. EMC3-OS, EML4-OS, and LH21-OS demonstrate results in three genotypes treated with gene therapy using the human BEST1 transgene with titers of ~2E11 vg/mL (Fig. 4A and Table S1). In each case, there was significant reduction of the IS/OS–RPE/T distance returning to WT levels both in superior and inferior retinal regions treated with subretinal gene therapy (Fig. 4B, filled symbols) but not in retinal regions away from the treatment bleb (Fig. 4B, unfilled symbols).

The region of efficacy with subretinal gene therapy is often shown to extend beyond the bleb formed at the time of the surgery to include a penumbral region (47, 48). In cBest dogs with successful gene therapy treatment, there was a defined penumbral region but it appeared to be qualitatively larger than typically encountered previously (Fig. 4A). In some of the most extreme examples, pretreatment maps of retina-wide microdetachment were found to be necessary to demonstrate the extent of penumbral expansion. EML9-OD, for example, at age 29 wk showed a retina-wide microdetachment that was most extreme along the visual streak and included several regions with gross retinal detachments (Fig. 4C). Gene therapy was performed at 69 wk. At 87 wk, microdetachments across the whole retina imaged as well as the majority of the gross retinal detachments had disappeared (Fig. 4C), and quantitative measures showed a normal or thinner IS/OS–RPE/T distance in superior and inferior retinal locations (Fig. 4D). Importantly, IS/OS–RPE/T distance showed substantial improvements at retinal locations corresponding to the bleb formed at the time of the injection as well as in nasal retinal control regions in the same eye. This extreme example of penumbral expansion is likely explained by greater diffusion of the vector via the microdetachment in cBest eyes that resulted in the transduction of the RPE at sites substantially more distant than the initial bleb. A more typical example with a delimited penumbral expansion is illustrated for comparison. EML13-OS at age 37 wk showed microdetachments retina-wide that were especially prominent in the temporal retina and along the visual streak; there were also several gross retinal detachments along the visual streak (Fig. 4E). Gene therapy was performed at 45 wk. At 81 wk

**Fig. 3.** BEST1 gene augmentation therapy results in sustained reversal of foveomacular lesions and restoration of RPE–PR interface structure. (A) Natural history of the central subretinal detachment documented by in vivo imaging in the right eye of cBest dog (EM356-OD; R25*/P463fs) at three time points. A series of representative near-infrared reflectance (NIR) images demonstrate the evolution of the focal lesion (green arrowhead) in the canine macula from the vitelliform stage (Left) through early pseudohyppyony (Middle) to an advanced pseudohyppyony stage (Right). (A, Insets) Autofluorescence and OCT images. (B) Subretinal injection with AAV2–cBEST1 (1.5E10 vg/mL) was performed in the eye shown in A at 52 wk of age, entirely encompassing the lesion depicted in the fundus photograph before injection (b.i.) (Left, Upper Inset). The subretinal bleb area is denoted by the dashed circle. The treated eye showed ophthalmoscopic resolution of lesion 1 wk postinjection (p.i.) (Left, Lower Inset); however, confirmatory OCT scans were not available at this age. Images acquired at 43 (Middle) and 245 wk (Right) p.i. document sustained reversal of the central lesion and fully reattached retina within the treated area. (B, Middle and Right, Insets) Autofluorescence and OCT images. EM356-OD and similar labels designate the individual animal and eye. (Scale bars, 200 μm.) (C and D) Restoration of RPE–photoreceptor interface structure post AAV–hBEST1 treatment in the cBest (R25*/R25*) model in comparison with control. (C and D, Upper) NIR images from EM3 eyes exhibiting early vitelliform lesions shown at ages 19 (C) and 25 wk (D). Both eyes were injected at 27 wk of age with either BSS (C, EM3-OD control eye) or AAV2-hBEST1 (2E11 vg/mL) (D; EM3-OS study eye). Bleb boundaries are marked by dashed circles; the locations of corresponding OCT scans cut through the subretinal lesions before injection or through the matching locations mapped p.i. are marked by green horizontal lines; retinotomy sites are indicated by blue arrowheads. (Scale bars, 200 μm.) (C and D, Lower) Confocal fluorescence images of IHC assessment of AAV–hBEST1– or BSS-injected eyes; in contrast to the BSS control eye (C), cytoskeleton rescue and restoration of RPE–PR interface structure are shown in AAV-treated retina 103 wk p.i. (D). (Scale bars, 10 μm.)
of age, both superior and inferior retina temporal to the optic nerve was lacking micro- and macrodetachments, whereas the untreated nasal retina had retained the microdetachments as well as formed a large number of macrodetachments (Fig. 4E). Quantitative results confirmed the treatment effect (Fig. 4F), which did not reach the nasal retina, unlike EML9-OD.

To understand the potential consequences of gene therapy on retinal degeneration, ONL thickness was mapped across treated eyes (Fig. S3B). Treated retinal regions showing disappearance of microdetachments tended to also correspond to normal ONL thickness, whereas untreated regions retaining microdetachments tended to show hyperthick or normal, or in some regions, thinned ONL (Fig. S3B). In summary, AAV-mediated gene augmentation therapy in canine bestrophinopathies appears to promote a sustained reversal of gross retinal detachments, reestablishment of a close contact between RPE and PRs, and return of ONL thickness to normal values.

**Human Autosomal Recessive Bestrophinopathies: Structure and Function.** To facilitate clinical translation of successful gene therapy in BEST1-mutant dogs, studies were performed to better understand the human pathophysiology of autosomal recessive bestrophinopathies (ARBs) and to gain insight into the distribution of retina-wide disease beyond the gross ophthalmoscopically detectable lesions previously described. Data are shown (Fig. 5) from two patients: P1 was a 39-y-old woman with a best corrected visual acuity of 20/100 carrying biallelic BEST1 mutations (c.341T>C/c.490C>G), whereas P2 was a 36-y-old man with 20/60 acuity also carrying biallelic mutations (c.95T>C/c.102C>T) in BEST1. In both patients, mutant alleles segregated with clinically unaffected parents. Ultrawide imaging of RPE health taking advantage of the natural autofluorescence of lipofuscin granules they contain showed widespread and extensive abnormalities consisting of regions of relative hyper- or hypoaufotuorescence and local heterogeneity. Of note, there was a distinct transition zone (Fig. 5A) at the fovea, where the healthier nasal peripheral retina.

Rod and cone function was sampled at high density along the horizontal meridian to better understand the topography of vision loss and its correspondence to retinal structural abnormalities. Both patients demonstrated a deep (>3 log) loss of rod-mediated sensitivity centrally in long-term dark-adapted eyes; there was relative preservation of rod function in the temporal field (nasal retina) in both patients and the parapapillary area in one patient (Fig. 5B, Upper). Surprisingly, cone-mediated function in light-adapted eyes demonstrated only a moderate loss (<1 log) or normal or near-normal results (Fig. 5B, Lower). Rod and cone function sampled across the full extent of the visual field corroborated and extended these findings and showed strong interocular symmetry (Fig. S7).

Cross-sectional imaging with OCT was performed to evaluate the retinal laminar abnormalities along the horizontal meridian
and the detachment of the central retina from the RPE, likely due to accumulation of subretinal fluid. The retinal lamination showed relative normalization in the parapapillary region (Fig. 5C, red rectangles) and beyond the nasal midperipheral transition (Fig. 5C, green rectangles). Analyses of the two regions of interest showed detectable but abnormally thinned ONL, and detectable IS/OS and cone outer segment tip (COST) with low peak signal in both patients (Fig. 5D and E). In P1, the distances from the ELM to IS/OS and IS/OS to COST were comparable to normal. There appeared to be a hyposcattering layer distal to COST, and the RPE appeared hyperthick (Fig. 5D, Middle). In P2, ELM to IS/OS appeared shorter than normal, whereas IS/OS-to-COST distance was comparable to normal. COST-to-ROST/RPE distance appeared greater than normal with an intervening indistinct hyposcattering layer; the RPE appeared to be comparable in thickness to normal (Fig. 5D, Right). Analysis of the outer retina in the nasal midperipheral region in P1 showed ELM-to-IS/OS, IS/OS-to-COST, and COST-to-ROST/RPE distances greater than normal and RPE thickness which was comparable to normal (Fig. 5E, Middle). P2 features appeared intermediate between P1 and normal (Fig. 5E, Right).

To understand the implications of structural abnormalities at the level of the outer retina and RPE for the kinetics of retinoid transfer between these cellular layers, dark-adaptation testing was performed. At the parapapillary location shown in Fig. 5D, dark-adapted thresholds of P1 were rod-mediated but 1.3 log unit-elevated (Fig. 5F). By 22.5 min following a light exposure, the P1 results had remained cone-mediated on a plateau whereas normal recovery was complete (Fig. 5F). At the midperipheral nasal retinal location shown in Fig. 5E, dark-adapted thresholds of P2 were rod-mediated and ~0.5 log unit-elevated compared with normal (Fig. 5G). By 14.5 min following a light exposure, there was first evidence of rod function which was only incrementally slower than the 11-min cone–rod break in normal. The rate of rod recovery was similar to normal (Fig. 5G). In summary, rod dark-adaptation kinetics of P1 at the parapapillary locus showed an extremely slow time course, whereas dark-adaptation kinetics of rod function of P2 at the midperipheral locus was closer to normal (Fig. 5F and G).

**Discussion**

The RPE has a key role in maintaining the metabolically active environment of the subretinal space (25, 26, 49). Due to the dynamic relationship with adjacent retinal layers, RPE-specific genes often adversely affect the neighboring sensory neurons, leading to loss of visual function and PR degeneration (50–52). Mutations in *BEST1* are known to disrupt transepithelial ion and fluid transport in response to abnormal levels of intracellular calcium (11–13). Abnormal RPE calcium signaling is also thought to lead to dysregulation of other pathways through altered expression and interactions of Ca²⁺-sensitive proteins (20, 53). Based on our findings in cBest, one such protein is EZRIN, a membrane–cytoskeleton linker essential for the formation and proper maturation of RPE apical MV (54–57). It has been demonstrated that the activation of EZRIN’s membrane–F-actin cross-linking function occurs directly in response to Ca²⁺ transients (55), and Ezrin–KO mice exhibit a substantial decrease in elaboration of RPE MV (57). The apparent underdevelopment of RPE apical MV found in the *BEST1*-mutant RPE is consistent with these reports. Furthermore, our comparative IHC assessments with other IRD models demonstrated that these major structural alterations associated with microvillar ensheathment are specific to the primary RPE channelopathy triggered by *BEST1* mutations, and not secondary to cone dysfunction and degeneration.

The structural components of RPE apical processes are very different from those of nonmotile intestinal microvilli (57–59). The presence of contractile proteins (such as myosin) in the RPE apical microvilli, and also molecules typically found at the sites of

**Fig. 5.** Retinotopic phenotype in patients with ARB. (A) RPE health across the retinas of two ARB patients, P1 and P2, imaged with short-wavelength reduced-illumination autofluorescence imaging (SW-RAFI), taking advantage of the natural RPE fluorophore lipofuscin. White arrows depict the location of the parapapillary perimetric profiles and OCT scans; red and green rectangles show the regions of interest shown in other panels; and black arrowheads demarcate disease-to-health transition in the nasal midperipheral retina. (A, Inset) Normal SW-RAFI for reference. (B) Perimetric light sensitivity of rods in dark-adapted (Upper) and cones in light-adapted (Lower) eyes measured across the horizontal meridian. Gray regions represent normal sensitivity except for the physiological blindspot corresponding to the optic nerve (ONH). (C) Retinal cross-section with OCT along the horizontal meridian crossing the fovea. (C, Inset) Normal OCT for reference. (D and E) Detail of outer retinal lamination in patients compared with normal at the two regions of interest at the parapapillary retina (D, red) and midperipheral nasal retina (E, green). Layers and peaks are highlighted for visibility. Orange indicates interface near COS tips, blue indicates interface near ROS tips and RPE apical processes, brick indicates interface near the RPE and Bruch’s membrane. (F and G) Dark-adaptation kinetics measured in P1 at the parapapillary locus (F) and in P2 at the nasal midperipheral locus (G). Time 0 refers to the end of adaptation light. Baseline thresholds obtained after extended dark adaptation are shown before time 0, crossing the fovea (Fig. 5C). There was not a consistent light-exposure history at the time of OCT imaging. Both patients showed a significant loss of ONL and abnormalities at the level of photoreceptor IS/OS in the outer retina across most of the central retina. P2, in addition, showed intraretinal cystic spaces.
cell attachments, suggests that the RPE actively adheres to, and exerts tension on, the neural retina (58, 59). The dearth of a proper microvillar ensheathment at the RPE–PR interface in cBest, and thus an absence of physical and electrostatic support by these projections to the PR OS, would be expected to weaken the adhesive forces and lead to separation of the RPE–PR complex retina-wide. The microdetachment of the PR layer from the underlying RPE found in cBest at the earliest stages of disease would be consistent with this process. Moreover, the presence of contractile elements in the RPE apical projections and the fact that they have evolved from cells in which pigment migration occurred (58) indicate that MV are capable of active contraction while interdigitating with PR OS, and are destined to facilitate circadian phagocytic activity. A single RPE cell can accommodate areas of ∼50 to 500 RPE, depending on the retinal location and packing density (60, 61), the elaborate network of microvilli allows each RPE cell to handle such a high metabolic load on a daily basis. Insights from proteomic profiling support this argument. There is an enriched fraction of retinoid-processing proteins expressed along the RPE apical MV, together with a number of channel proteins and transporters (i.e., Na+/K+/ATPase) central to the efficient transport of water, ions, and metabolites between the RPE and PR OS (26, 59). Considering the topographic differences in the size of RPE cells and taking into account the density and length of MV quantified in this study, MV extensions expressed the functional surface of a single RPE cell by 20- to 30-fold in the central retina, which is consistent with earlier estimates (62). This number is even higher (50-fold) for the small RPE cells in the macular region (26, 63) that adapted to a higher turnover rate of shed POS while facing the most densely packed PRs. Such dramatic reduction in a total apical surface area in BEST1-mutant RPE will lead to a chronic delay in processing of metabolites, and arrest the abilities of the RPE to maintain both the proper cell volume as well as chemical composition and physiological pH levels in the subretinal space (26, 50). Retinal detachments in these eyes are due to the loss of adhesive forces (26), any limitation in the RPE transporting system will alter the balance of hydrostatic forces and result in decreased osmoelastic properties of the RPE–PR complex with subsequent separation from the neuroretina. Indeed, the primary serous detachment in human and canine bestrophinopathy is first evident in the fovea, the central area of highest metabolic activity. The absence of high-reaching RPE apical processes, which in the structurally intact retina tightly wrap COS up to the ellipsoids (64), would exploit the predictability of this cone-rich structure for rapid photoreceptor metabolism. The RPE detachment would be almost exclusive reliance on the frictional interactions with the MV. This is consistent with our observations in cBest, documenting formation of the focal previtelliform lesions within the canine fovea-like area of the area centralis, and also the susceptibility of other central cone-rich areas (like visual streak) to subretinal detachment.

Less expected was the major expansion of microdetachments in cBest upon exposure to dim and moderate light intensities. In normal eyes, light exposure is known to change molecular composition of the subretinal space (65). There is also evidence that measurable structural changes occur in the normal outer retina with light exposure, such as changes in outer segment length (39, 40, 66, 67), hydration of the subretinal space (27–29, 35, 68), increased actin staining along RPE apical MV (37), and phototropism of outer segments (69). However, all of the normal changes are substantially smaller than those measured in cBest. For example, normal human eyes showed changes of ∼1 μm (40), and normal mouse eyes showed changes of ∼4 μm in the outer retina (37), compared with ∼15-μm expansion of the subretinal space (35) in cBest. Hyperthick ONL in patients with BVMD has only recently been recognized and the literature on the earliest disease stages is limited. Specifically, we are not aware of any ARB literature that has performed quantitative measurements of the outer retina. The recessive cBest disease appears to have phenotypic similarities to both dominant and recessive bestrophinopathies in humans. In patients with dominant BEST1 disease (BVMD), there has been some controversy regarding the structural features of retinal regions surrounding vitelliform or later-stage lesions, or retinas in the provitelliform stage of disease. Some studies have demonstrated minor abnormalities at the level of the RPE–PR interface (34, 70, 71), whereas results from others support no detectable structural defects (72–74). Contributing to this controversy could be genotype (75), the resolution of different methodological approaches used, or light history preceding the imaging (36). Indeed, light-dependent outer retinal changes have been described in BVMD using methods like those reported here (36); still, the magnitude of the changes reported in patients was smaller (∼2 μm) than in cBest. In general, however, the abnormal response of the affected retina to light stimuli to a level related to the markedly reduced light peak/dark trough ratio in the electrooculogram, a finding consistent in all, even presymptomatic, BEST1 patients.

Of importance, both the micro- and macrodetachments in cBest had adverse effects on photoreceptor health: Regions of microdetachment tended to correspond to hyperthick ONL, whereas large lesions with gross macrodetachment showed thinning of ONL. Smaller lesions with macrodetachment could not be assessed with the sampling methods used here. ONL contains the nuclei of all rods and cones, and classic studies in animal models and human eye donors have generally shown photoreceptor thinning of the ONL with disease progression (76, 77). Less well known are some of the earliest stages of retinal disease showing ONL thickening, which has only become measurable with the advancement of in vivo imaging methods. Human studies have previously demonstrated such ONL thickening in early stages of retinal diseases (78–81). There has also been evidence in animal studies of ONL thickening associated with retinal stress (82). The hyperthick regions of ONL mapped in cBest when examined microscopically showed the number of PR nuclei to be comparable to control, suggesting a greater intracellular spacing within the ONL, likely corresponding to a level related to the below the apoptosis threshold. Gross retinal detachments, on the other hand, may cause greater retinal stress and progressive degeneration (17, 83).

To prevent the photoreceptor and vision loss associated with BEST1 mutations, we performed subretinal gene augmentation therapy directed to retinal areas with macro- and microdetachments. Our results showed that AAV-mediated BEST1 gene augmentation is safe, reverses the clinically obvious lesions, ameliorates the diffuse microdetachments, and results in normalization of hyperthick ONL. Furthermore, we demonstrated gene therapy success in three distinct BVMD patients, with both focal and multifocal presentations, and confirmed long-term durability of the treatment effect. At the molecular level, we confirmed that both the canine as well as the human BEST1 transgene can correct the apposition of the RPE–PR complex and restore the cytarchitecture of this critical interface. Our study suggests that early as well as more advanced stages of autosomal recessive disease are sensible to approach with this therapy. Further studies utilizing human inducible pluripotent stem cell (hiPSC)-derived RPE models derived from patients harboring autosomal dominant BEST1 mutations (17, 53) will determine whether the gene augmentation approach would also be beneficial for BVMD patients.

To facilitate the clinical translation of successful gene augmentation therapy, ARB patients were studied to gain insight into their retina-wide disease. Consistent with most, but not all, previous descriptions (84–88), retinal disease in our ARB patients extended well beyond the macula into the midperiphery. Our retinotopic mapping of en face and cross-sectional imaging and rod and cone function demonstrated the existence of a distinct tranBest from disease to health in the midperipheral retina, a feature not previously emphasized. Within the diseased region, severe abnormalities in retinal structure were associated with severe loss of rod function; unexpectedly, cone function was relatively retained. Rod dysfunction within the central retina was also associated with extreme slowing of the retinotopic cycle,
whereas the healthier periphery showed near-normal recycling of the retinoids. There are at least two retinoid cycles that provide the 11-cis-retinol chromophore to photoreceptor pigments. The canonical retinoid cycle functions in the RPE to produce chromophore for visual and cone PRs (99). The retinyl ester cycle on the other hand, is thought to regenerate chromophore within the retina for the specific use of cones (90, 91). The abnormal RPE–PR interface in BESTi disease would most likely affect the chromophore delivery from the canonical RPE retinoid cycle; the retinal retinoid cycle may be relatively unaffected, thus explaining the greater retention of cone function.

In summary, our study uncovered new molecular contributors to the pathophysiology of bestrophinopathies at the RPE–PR interface. We discovered the earliest expression of disease-related diffuse microdetachment potentiated by light exposure that was easily detectable by in vivo imaging. We showed that AAV-mediated BESTi augmentation gene therapy reverses both the grossly obvious lesions and microdetachments, and restores the cytoarchitecture of the RPE–PR interface. Evaluation of ARB patients showed retinotopic distribution and properties of structural and functional defects beyond that expected from PR degeneration. Such visual dysfunction may be expected to improve upon successful application of BESTi gene augmentation therapy to patients affected with bestrophinopathies.

Materials and Methods

Human Subjects. Two unrelated patients with ARB were included. Procedures followed the Declaration of Helsinki guidelines. Informed consent was obtained, and all procedures were approved by the Institutional Review Board of the University of Pennsylvania. Patients underwent a complete eye examination and ancillary tests as previously described (92–95). Further experimental details are described in SI Materials and Methods.

Canine BEST1 Models and in Vivo Retinal Imaging. Best-mutant dogs (n = 18) of both sexes (12 M and 6 F) harboring either homozygous (c.737C>T (p.R25*), R25* or c.1388delC (p.P463fs*1) or leucine insertion (c.737C>T-1388delC) (p.R25*P463fs) mutations in c.BESTI (GB*NM_001097545) were included. For ease of annotating the multipanel figures, the three genotypes, respectively, are referred to as cmr1, cmr3, and cmr1/cm3. The study was conducted in comparison with control-crossbred dogs (n = 12; 7 M and 5 F) (Table S1). All animals were bred and maintained at the Retinal Disease Studies Facility (RDSF). The studies were carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the NIH (96) and in compliance with the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research. The protocols were approved by the Institutional Animal Care and Use Committee of the University of Pennsylvania (IACUC nos. 804956 and 803422). En face and retinal cross-sectional imaging was performed with the dogs under general anesthesia as previously described (24, 97). Further methodological details are provided in SI Materials and Methods.

ACKNOWLEDGMENTS. The authors thank E. V. Dutrow, K. Zorych, A. McTish, and A. Dihingra for excellent technical assistance; Theresa Jordan and RDSF staff for animal care; Drs. J. M. Guzman, F. L. Marinho, and J. C. Ch for assistance in subretinal surgeries; M. Leonard for help in graphic design; and L. Mlynk for research coordination. This work was supported by the Foundation Fighting Blindness: FFB-TRAP and FFB-Grant; Macula Vision Research Foundation: Donald and Dorothy Hall LAH VMD2 (33:3385–3393); Li JD, Gavardovskii D, Shibayama H (94) Light-dependent hydration of the space surrounding photoreceptors in the cat retina. Vis Neurosci 11:743–752.

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