Single choroideremia Gene in Nonmammalian Vertebrates Explains Early Embryonic Lethality of the Zebrafish Model of Choroideremia

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PURPOSE. Mutations of the CHM gene underlie the X-linked chorioretinal degeneration choroideremia (CHM). The affected gene product, Rab Escort Protein (REP1), mediates the post-translational prenyl modification of Rab GT9ases. In patients with CHM, the related REP2 partially compensates for the loss of function of REP1. The objective of this investigation was to study the natural history of disease in a zebrafish model of CHM.

METHODS. Zebrafish chm−/− were bred and subjected to extensive histologic analysis and TUNEL assays, and cellular extracts were used for immunoblot and in vitro prenylation assays. A detailed evolutionary analysis was performed on the REP family.

RESULTS. The retina of chm−/− zebrafish develops normally for the first 4 days postfertilization (dpf) but that catastrophic multilayer degeneration synchronous with severe multisystem disease follows. Mean survival time is 4.8 dpf. At the onset of generalized disease, a significant reduction in retinal expression levels and activity, with unprenylated rabs accumulating in the cytosol was demonstrated. Extensive bioinformatic analysis of the REP family of proteins revealed a single rep isoform in fish and other nonmammalian vertebrates and invertebrates that is similar to mammalian REP1.

CONCLUSIONS. REP1 appears to be the ancestral gene in the family, whereas the intronless REP2 gene is restricted to the mammalian lineage. The results of this study propose that in chm−/− zebrafish, maternally derived rep allows initial successful development of the embryo, but its gradual loss leads to multisystem disease and invariably to lethality. In its current form, the chm−/− zebrafish has limited usefulness. (Invest Ophthalmol Vis Sci. 2009;50:3009–3016) DOI:10.1167/iovs.08-2755

CHoroideremia (CHM; Online Mendelian Inheritance in Man [OMIM] 313100) is a progressive chorioretinal degeneration with X-linked inheritance. Its estimated incidence is 1 in 50,000, and this slow-onset disease causes night blindness in affected teenage males that progresses to loss of peripheral vision and complete blindness two to three decades after onset. CHM is characterized by slow progressive loss of the RPE and choroid in scalloped areas and atrophy of the photoreceptor cell layer. Loss of the choriocapillaris and preservation of the large chorioidal vessels result in a characteristic fluorescein angiogram pattern. Heterozygous female carriers do not usually develop blindness but do have patchy areas of chorioretinal atrophy.1,2

The CHM gene was identified as encoding Rab Escort Protein (REP1) with disease caused by loss-of-function mutations leading to a truncated, nonfunctional, or rapidly degraded protein. A homologous protein, REP2, is the product of an intronless gene on chromosome 1. REP2s are necessary for successful prenyl (geranylgeranyl [GG]) modification of the Rab family of small GT9ases. REP5s bind newly synthesized Rabs and present them to Rab GG transferase (RGGT), which catalyzes the transfer of prenyl groups to the Rabs. REP then binds the prenylated Rab and delivers it to its target membrane, to which the Rab is anchored by its prenyl group. REP proteins regulate membrane transport pathways, including vesicle formation, organelle movement, and membrane tethering/fusion, by residing in the membrane and interacting with a range of effector molecules in target vesicle membranes. Other small GT9ases are prenyl lipid modified by GG transferase type I and farnesyl transferase.10

Rat tissue studies have shown ubiquitous expression of both REPs, and the absence of REP1 in CHM may be compensated for by REP2 in most tissues.11,12 However, a subset of Rabs, such as Rab27a in CHM lymphoblasts, remains selectively underprenylated in the absence of REP1.13 The recently created CHM mouse model carries a tissue-specific, tamoxifen-inducible conditional knockout of the Rep1 gene.14 Heterozygous females show early onset and progressive retinal degeneration similar to human CHM. Generation of tissue-specific knockouts of REP1 in the RPE and photoreceptors suggested cell-autonomous degeneration associated with different subsets of underprenylated Rabs in each cell type. Previous attempts to develop a REP knockout mouse were unsuccessful because of embryonic lethality in males and heterozygous females when the gene is inherited from a heterozygous mother.15 This was caused by defects in trophoblast vascularization and development.16

Zebrafish have several advantages over the mouse for use as a model for the vertebrate retina.17 Zebrafish undergo extremely rapid extrauterine embryogenesis, throughout which they remain transparent. Their retinal architecture is similar to that of humans, with a rich cone density and retinal lamination reaching an adultlike state by 72 hours postfertilization (hpf). Additionally, zebrafish have a 3-month generation time; each
breeding pair can produce up to 200 eggs per week, and animal husbandry costs are significantly lower than for mice. A zebrafish strain with a loss-of-function mutation in the orthologous CHM gene was recently described, created by random ENU (N-ethyl-N-nitrosourea) mutagenesis and identified in a screen for balance and hearing defects.18 Positional cloning revealed a nonsense mutation (C→T) in the second exon of rep1, a gene homologous to REP1, at residue 32 of a 566-amino acid protein.19 Mutant fish exhibited a variety of phenotypes culminating in death by 6 days postfertilization (dpf).19 The retinal phenotype was subsequently analyzed by Krock et al.19 in more detail at one developmental time point, 4.5 dpf. At this stage, the mutant fish exhibited degeneration of the RPE and photoreceptors, with complete loss of visual function as measured by electroretinography.19 With the use of transmission electron microscopy, photoreceptor layers were found to be dehisced and degenerating, and melanosomes in the RPE cells were smaller and immature and had reduced density compared with those in the wild-type retina. In addition, photoreceptor outer segment material was not effectively eliminated from the mutant RPE, resulting in the development of large vacuoles in this layer.19 However, the natural history of the disease has not been addressed in previous studies. To understand the processes underlying CHM in the zebrafish and to assess its usefulness as a model for human disease, we sought to further characterize the pattern of retinal degeneration and to correlate this with human pathology. Here we report catastrophic degeneration of the retina after 4.5 dpf. We suggest that the retinal and extraretinal phenotypes are caused by a lack of compensatory REP2, which evolved uniquely in the mammalian lineage. Absence of full-length REP is a lethal failure and death.

**Materials and Methods**

**Aquaculture**

Zebrafish (wild-type AB and chm<sup>mut</sup> embryos) were bred and maintained in the Imperial College animal facility according to standard protocols and the guidelines of the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.20 Embryos were raised at 28.5°C in facility water. The developmental stages are given in hpf or dpf, in accordance with morphologic staging criteria.21

**Histologic and Morphologic Studies**

Zebrafish embryos were fixed in 4% paraformaldehyde/PBS overnight at 4°C. Samples were serially dehydrated in alcohol (5-minute washes in 50%, 70%, 90%, 3 × 100%) before they were embedded in paraffin wax, and 5-μm microtome sections were stained with hematoxylin and eosin. Sections were imaged with a camera (SIS ColorView Illu; Olympus, Tokyo, Japan) mounted on a microscope (CKX41; Olympus). Wholemount zebrafish were imaged with a camera (DFC300 FX; Leica, Wetzlar, Germany) mounted on a fluorescence stereomicroscope (MZ16F; Leica).

**TUNEL Assay**

Wild-type and chm embryos were treated with 0.2 mM phenylthiourea (Sigma, St. Louis, MO) at 10 hpf to prevent pigment formation and were incubated at 28.5°C until 5 dpf. After fixation in 4% paraformaldehyde overnight at 4°C, embryos were washed in PBS three times. Retinal sections were prepared as outlined and were de waxed by two washes in clearing agent (Histoclear; National Diagnostics, Atlanta, GA) followed by two washes in 100% ethanol and one wash in 70% ethanol before they were rinsed in deionized H<sub>2</sub>O. For wholemount TUNEL assays, embryos were dehydrated through a graded methanol series (25%, 50%, 75%, and twice in 100%) and stored in 100% methanol at −20°C. After rehydration, sections and wholemount embryos were digested with proteinase K (10 μg/mL). Embryos were refixed with 4% paraformaldehyde for 20 minutes at room temperature, followed by several washes in PBS. A detection kit (ApopTag Peroxidase In Situ Apoptosis Detection Kit; Millipore, Billerica, MA) was used to detect levels of apoptotic cell death according to the manufacturer's instructions.

**Bioinformatics**

Protein sequences were obtained by querying species-specific databases with human REP1, human REP2, human GDIα, yeast REP, and yeast GDI using the BLAST algorithm.22 Our analysis was complemented by the more sensitive domain-based methods to identify protein families using Pfam23 (GDI domains) and Superfamily24 (FAD/NAD binding domains + FAD linked reductase C-terminal domains). Candidate sequences were visually inspected, and REPs and GDIs were distinguished with the use of phylogenetic methods and by the presence of F279 and F135, respectively.25 Protein sequences were aligned, and Neighbor Joining Trees were calculated, with 1000 bootstraps, with the use of graphics software (ClustalX 1.80: http://www.clustal.org)26 with default parameters. Maximum likelihood phylogenetic inference was performed with Phylib<sup>27</sup> with 100 bootstraps.

**Immunoblot Analysis**

Protein extracts were obtained from 20 wild-type and chm embryos at 6 dpf. Embryos were snap frozen in liquid nitrogen and homogenized in lysis buffer (10 mM Tris, pH 7.5, 10 mM NaCl, 1% SDS, 1× Protease Inhibitor Cocktail [Roche, Basel Switzerland]) by sonication. Protein concentration was measured with the use of a protein assay kit (BCA Protein Assay kit; Pierce, Rockford, IL). For each sample, 30 μg protein was boiled for 5 minutes with an equal volume of SDS sample buffer. Proteins were separated on a 10% SDS-polyacrylamide gel and transferred to a polyvinylidene difluoride membrane (Immu-Blot; Bio-Rad, Hercules, CA). The membrane was blocked for 2 hours in blocking solution (5% dry milk, PBS/0.1% Tween [PBST]), washed three times in PBST, and incubated overnight at 4°C with 1:1000 primary anti-REP J905 antibody.13 After three washes in PBST, the membrane was incubated with a secondary anti-rabbit IgG horseradish peroxidase conjugate (dilution 1:10,000; GE Healthcare Chalfont St. Giles, Buckinghamshire, UK) for 2 hours. The membrane was washed three times in PBST before chemiluminescence detection using the ECL Western blotting detection kit (GE Healthcare). With the use of a stripping buffer for Western blots (Restore Western Blot Stripping Buffer; Pierce), the membrane was stripped and reprobed with polyclonal anti-gapdh antibody (Abcam, Cambridge, MA) as a loading control for each sample. Image data analyzer software was used to determine the relative abundance of rep protein compared with corresponding levels of control gapdh.

**Subcellular Fractionation and In Vitro Prenylation Assays**

Samples were homogenized in hypotonic buffer (50 mM HEPES, pH 7.5, 10 mM NaCl, 1 mM dithiothreitol, 1× protease inhibitor cocktail) by sonication. Nuclei were pelleted by centrifugation at 800g for 10 minutes before the postnuclear supernatant was collected and centrifuged at 100,000g for 1 hour at 4°C with a rotor (TLA45; Beckman-Coulter, Fullerton, CA) to obtain cytosolic protein extracts. In vitro prenylation was performed in reaction buffer containing 50 mM HEPES, pH 7.5, 5 mM MgCl<sub>2</sub>, 1 mM NP40, 1 mM dithiothreitol, 1× protease inhibitors (Roche), and 1× phosphatase inhibitors (Sigma). Each 20-μL reaction contained 4 μM recombinant Rab1a, 100 nM recombinant RGGT, 50 nM 3H-GGPP supplemented with 5 μM unla-
beled GGPP, and 25 μg zebrafish cytosolic extract containing a source of rep. After incubation for 30 minutes at 37°C, reactions were precipitated by the addition of 10% HCl in ethanol and were incubated for 30 minutes at room temperature. As a positive control, zebrafish extracts were used as a source of GGTase-I to prenylate 5 μM recombinant human GST-Rac1a using the same reaction conditions but omitting RGGT. Reactions were filtered on glass microfiber filter discs (Whatman, Maidstone, Kent, UK). Each reaction tube was rinsed once with 2 mL 100% ethanol, and each filter was rinsed three times with 2 mL 100% ethanol and counted in a scintillation counter (LS6000SC; Beckman-Coulter). Gel-based assays used 100 μg extract/25 μL reactions, substituting recombinant Rabs with 2 μM recombinant REP and 1 μM (instead of 5 μM) unlabeled GGPP. Reactions were incubated for 45 minutes at 37°C. Loading buffer (5 μL) was added to each reaction, and aliquots were boiled for 5 minutes at 95°C. Aliquots were separated by SDS-PAGE on a 12.5% gel, and the proteins were fixed by incubation for 1 hour in a solution of 40% ethanol and 10% acetic acid. Gels were incubated for 20 minutes in solution (Amplify; Amersham, Piscataway, NJ) before they were dried in a 60°C oven and were exposed to autoradiography film for 2 weeks.

RESULTS

Phenotypes in Homozygous chm Null Zebrafish

The macroscopic phenotype observed in chm−/− mutants conforms to that previously described by Starr et al.18 Among the gross morphologic abnormalities visible from 3 to 4 dpf onward were edema of the heart and abdomen, absence of a functional swim bladder, reduced body length, curvature of the spine, increased dark pigmentation, and microphthalmia (Fig. 1). Mutants also hatched later than their unaffected siblings and retained noticeably larger yolk sacs, possibly because of the lower nutritional requirements of impaired growth. Invariably, chm−/− fish died early, usually by the fifth day (mean survival, 4.8 dpf; n = 50).

We next examined retinal histology from mutant chm−/− embryos to characterize the development of retinal disease. Compared with other cell types, photoreceptors occupy a large proportion of the eye from 55 hpf, when the formation of outer segments is initiated.28 Hence, we hypothesized that photoreceptor cell death would cause reduced eye size as early as 3 dpf. At this age, wild-type retinas were laminated, and photoreceptor cells had developed their distinctive morphology (Fig. 1A). Surprisingly, no morphologic evidence of photoreceptor cell death was seen in most sections from 3 dpf or even 4 dpf mutants (Figs. 1A, B). The earliest observed abnormality was a patchy loss of photoreceptors in the outer nuclear layer, with the underlying RPE directly abutting the cells from the inner nuclear layer (Fig. 1C). By 5 dpf, multiple cell types throughout the retina were damaged (Figs. 1D, 2). The RPE was hypertrophic and invaded the retina, in some places forming discrete “islands” of pigmented cells; in other areas it was depigmented or absent. Photoreceptors became disorganized and lost their columnar morphology, leaving only regular or pyknotic nuclei in their places, and the outer plexiform layer was almost completely absent. Ordered lamination of amacrines, bipolar, and horizontal cells within the inner nuclear layer was lost, and pyknotic nuclei were present throughout.

**Figure 1.** Morphology and retinal histology of chm mutants from 3 dpf. Progression of systemic and retinal degeneration from 3 dpf (A), 4 dpf (B), 4.5 dpf (C), and 5 dpf (D). Left: comparison of wholemount wild-type (top) and chm (bottom) gross morphology. Middle: coronal retinal sections of wild-type embryos. Right: retinal degeneration of chm mutant embryos with magnified view of retinal slice. (C, D, red arrows) Area of patchy photoreceptor cell loss in chm retina at 4.5 dpf and 5 dpf. (D, white arrow) Area of RPE hypertrophy with invasion of inner nuclear layer. RPE, retinal pigment epithelium; ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer; ON, optic nerve; p, pyknotic nuclei.
the retina, suggesting widespread degeneration (Fig. 2). Severity of the phenotype varied between individuals, suggesting some disparity in the rate of disease progression. For chm−/− mutants that had a milder phenotype at 5 dpf (Fig. 2H), degeneration appeared to manifest first in the peripheral retina while the central retina remained normal and showed no evidence of cell death. The peripheral RPE invaded the dorsal and ventral retina, and photoreceptors receded from the retinal germinal zone. The outer plexiform layer was discontinuous, and pyknotic nuclei were present in the inner nuclear and ganglion cell layers. In all cases, degeneration of the retina in chm−/− embryos was surprisingly rapid, usually occurring in 12 hours, between 4.5 and 5 dpf. The lens displayed various changes, though there was considerable variation between affected animals. When observed, the lens was often smaller, denser, and impacted and had lens opacities on examination (Figs. 1D, 2). Wholemount TUNEL staining in chm−/− mutants at 5 dpf revealed widespread cell death across the entire embryo (Fig. 3). Coronal retinal sections showed high levels of apoptosis within the outer nuclear layer involving photoreceptors, with extension into the inner nuclear layer (Fig. 3). Cell death appeared more prolific at the peripheral retina, in accordance with the histology.
with the histologic findings, and around the exiting optic nerve within the outer nuclear layer. In addition, ganglion cells posterior to the lens capsule were undergoing apoptosis.

**Absence of REP Duplication in Fish**

The rapid, degenerative disease observed in cbm/zebrafish extends beyond the outer retina to cause a more severe retinal, multisystem, and lethal phenotype, more comparable to that seen in yeast with REP mutations (called MRS6) rather than in mice or humans. Furthermore, we had previously established that in model nonvertebrate organisms, only a single REP gene is present. We thus hypothesized that zebrafish, unlike mammals, contain a single REP gene whose deletion accounts for the embryonic lethal phenotype.

To test this hypothesis, we decided to reconstruct the evolutionary history of the REP family in vertebrates. We first conducted extensive database searches and identified members of this family in multiple animals. We found two REP isoforms in every eutherian genome (placental mammals; Fig. 4). The closest extant relatives of eutherians are the marsupials and the monotremes. We found a single REP gene in the early release of the platypus genome, a monotreme, and three REP genes in the latest genome release of the marsupial *Monodelphis domestica*. In amphibians, *Xenopus laevis* exhibits two isoforms, whereas *Xenopus tropicalis* exhibits only a single gene. Further down in the vertebrate tree of life, we found a single REP gene in the completed genomes of the zebrafish and the pufferfishes, Takifugu rubripes and Tetraodon nigroviridis (Fig. 4).

Next, we used phylogenetic reconstruction to identify and order gene duplication events and to group the multiple REP isoforms into subfamilies. Mammalian REP1 and REP2 form two distinct groups, suggesting a single origin for each. Human REP2 is an intronless gene residing in the intron 1 of the gene panopsin. Our analysis further indicated that all mammalian REP genes are intronless.

Opossum REP genes are monophyletic, which indicates two consecutive species-specific duplications. In contrast, the two REP genes in the alloplloid frog *X. laevis* are likely accounted for by the maintenance of genes from both parent species in a pseudotetraploid genome. All other species, including the three fish genome sequences we analyzed, have a single REP gene. This, together with dating of the origin of REP2 to after the separation of the mammalian lineage, supports our hypothesis that a single REP gene in fish accounts for the gene deletion lethality.

**Reduced REP Activity in Choroideremia**

These data established that zebrafish contain a single REP gene, and we know that this is an essential gene in yeast. How can we account for cbm/zebrafish survival for approximately 5 days? This would imply that the fish must have some functional REP activity during that period. We hypothesized that a maternal contribution of functional REP activity allows brief survival of cbm/homozygotes by creating a supply of prenylated Rab proteins in the early embryo. In the embryonic zebrafish, all processes occurring before zygotic gene activation rely on maternal products developed during oogenesis and present in the egg at fertilization. These maternal factors must support all cellular functions. To test this, immunoblot analysis was undertaken on total zebrafish extracts from wild-type and cbm mutants with end-stage generalized disease (in this case, at 6 dpf; Fig. 5). Wild-type embryos expressed 87% rep protein at 6 dpf compared with a loading control, gapdh, whereas cbm mutants exhibited only 15% of control. This source of rep protein is attributed to residual maternal contribution derived from the yolk sac.

To complement this experiment, we sought to obtain direct evidence for Rep activity in the zebrafish embryos. To measure rep activity in zebrafish, we adapted an in vitro prenylation assay measuring the incorporation of radiolabeled GG diphosphate (GGpp) into Rab. In this assay, whole zebrafish protein extracts were used as a source of rep and were incubated with an in vitro reaction mix containing recombinant RGGT, Rab1a substrate, and 3H-GGpp. Given the variability between protein extracts, we introduced an internal control in the assay by measuring GGTase type 1 activity, which does not require REP as a cofactor. When normalized to GGTase-1 activity, 5 dpf cbm/zebrafish extracts exhibited a 2.5-fold reduction in rep activity compared with wild-type fish extracts (1.88 vs. 0.78 compared with normalized GGTase-1 activity).
activity in the same extract; Fig. 6A). Another prediction from our hypothesis was that the whole-zebrafish cytosolic extracts would have accumulated unprenylated rabs. In this assay, the cytosolic extracts were a source of rab substrates incubated with recombinant REP and RGGT in the presence of 3H-GGpp. Under these conditions, we observed the presence of 3H-GG-rabs in \textit{chm}/H11002/zebrafish extracts but not in wild-type extracts (Fig. 6B). As a positive control, we used a mammalian cell line that accumulates unprenylated Rabs on treatment with the mevalonate pathway inhibitor mevastatin.\textsuperscript{13} This experiment demonstrated the presence of unprenylated rabs in \textit{chm}/H11002/zebrafish, which correlated with the rapid onset of generalized disease. Altogether, our results suggest a mechanism for catastrophic multisystem degeneration in \textit{chm}/H11002/zebrafish leading to lethality by 5 to 6 dpf.

**DISCUSSION**

We present here evidence that the REP family of proteins underwent a mammalian-specific gene duplication event that prevented lethality after loss-of-function mutations in humans. When REP1 is affected, the retinal-restricted phenotype manifests as CHM, whereas any disease resulting from the loss of REP2 remains unidentified. In \textit{chm}/H11002/zebrafish, the loss of the single REP gene is lethal by 5 to 6 dpf, and early survival may be attributed to the maternal supply of REP activity.

Human CHM abnormality is limited to the photoreceptors, RPE, and choroid throughout life. In zebrafish, the degenerative process seen in the orthologous gene is more severe. The retina forms relatively normally in the first few days after fertilization; however, after 4.5 dpf, we observed rapid degeneration that included all retinal laminae. Because of the speed of the degeneration, it was difficult to elucidate the sequence of cellular loss in the retina. Indeed, one of our initial goals was to determine whether the RPE or photoreceptors were affected first and whether some tissues underwent secondary damage. We quickly realized that the degenerative process was of a general nature and affected most cell types beyond the outer retina and, indeed, the eye. In addition to inner retinal and lens degeneration, \textit{chm}/H11002/zebrafish exhibited uninflated swim bladders, edema of the heart and abdomen, irregular iridophore distribution, and stunted growth that was often associated with spinal curvature.
While this work was in progress, Krock et al.\textsuperscript{19} reported on the retinal phenotype of \textit{cbm} \textsuperscript{--/} fish and showed evidence of RPE and photoreceptor degeneration. Our observations concur with the reported observations, including normal retinal lamination, and various obvious pathologic changes in the RPE and photoreceptors. However, that work was limited by the absence of time-course studies given that the reported analysis was restricted to 4.5 dpf embryos. This precedes the onset of extensive degeneration of the inner retina (Figs. 1, 2). A mere 12 hours later, the retina looks very different, and generalized degeneration is evident in all cell layers. Therefore, one important issue is whether \textit{cbm} \textsuperscript{--/} fish retinal degeneration bears any resemblance to human CHM (in other words, whether \textit{cbm} \textsuperscript{--/} fish may be used as a faithful model of CHM). Our results indicate that it does not.

The extreme abnormalities of zebrafish \textit{cbm} \textsuperscript{--/} can be explained if this disorder is considered to be a null mutation in an essential gene rather than strictly a retinal degeneration. Embryonic essential genes in zebrafish are classified as those that, when disrupted, result in a phenotype visible by 5 dpf that is invariably lethal.\textsuperscript{35} By this definition, \textit{cbm} is an embryonic lethal gene because REP is essential for embryonic development. Our hypothesis to explain the generalized degenerative process followed by death before 6 dpf in \textit{cbm} \textsuperscript{--/} zebrafish is as follows. The maternal supply of rep allows for enough REP activity to ensure the normal development of the early embryo. Maternal REP activity allows the prenylation of the rab GTPases expressed by these cells. A significant maternally derived complement of mRNA or rep protein must be highly stable to persist for several days in zebrafish \textit{cbm} \textsuperscript{--/} mutants.\textsuperscript{36–39} Indeed, maternal products have been shown to function beyond the midblastula transition, when most maternal mRNAs are degraded.\textsuperscript{54, 56} In this scenario, another variable that could explain variation in disease severity between individuals is the initial dosage of maternal rep received by the embryo.\textsuperscript{58} As the finite supply of maternal mRNA for rep declines, so does the expression of rep protein. Nevertheless, experiments in yeast have shown that reduced levels of REP prenylation activity are sufficient to maintain a functional Rab pool in vivo, and REP function is only transiently required to support normal growth.\textsuperscript{41} An additional specificity of REP function is that it serves to activate Rab, and Rab activity appears to be the required function for cell viability. In yeast, prenylated Rabs are relatively stable, requiring at least a single generation for turnover, and only 10% to 20% of normal levels of prenylated (membrane-bound) Rabs are sufficient to allow cellular growth.\textsuperscript{41} Therefore, declining REP activity would manifest itself phenotypically only after the pool of prenylated Rabs had turned over. At that point, Rab dysfunction would reach a level inconsistent with the maintenance of normal membrane trafficking processes, and generalized degeneration would ensue.

Our analysis revealed that all mammalian REP2 genes are intronless, further supporting a single gene duplication event through reverse transcription. These data strongly suggests that REP1, rather than REP2, is the ancestral form, as proposed previously. Many autosomal genes that arose by retroposition from X-linked genes developed testis expression, presumably to avoid X inactivation during spermatogenesis, when the retrogene may contribute complementary function(s) to the parental parologue.\textsuperscript{43–45} It would be interesting to test whether REP2 follows the same male-specific bias to determine what specific function it may perform during spermatogenesis and whether REP2 malfunction may be associated with infertility or other germline defects in mammals.

It appears then that the absence of a second compensatory REP limits the application of zebrafish as a model for choroideremia. REP2 plays an essential role in disease pathogenesis in humans. Hence, the introduction of a transgenic copy of human REP2 gene into \textit{cbm} \textsuperscript{--/} zebrafish may create a model for the study of human CHM disease. Conversely, a model organism with a single REP will be of use in characterizing therapies that aim to boost REP1 activity, such as novel drug classes that allow read-through of nonsense mutations.\textsuperscript{46–48} Such approaches will afford the zebrafish a continued role in efforts to further understand and seek treatments for CHM.

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\section*{References}


