Photoreceptor Cell Death, Proliferation and Formation of Hybrid Rod/S-Cone Photoreceptors in the Degenerating STK38L Mutant Retina

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Abstract
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Keywords
homozygous mutation, STK38L, dogs, retina, photoreceptor

Disciplines
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**Photoreceptor Cell Death, Proliferation and Formation of Hybrid Rod/S-Cone Photoreceptors in the Degenerating STK38L Mutant Retina**

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**Abstract**

A homozygous mutation in STK38L in dogs impairs the late phase of photoreceptor development, and is followed by photoreceptor cell death (TUNEL) and proliferation (PCNA, PHH3) events that occur independently in different cells between 7–14 weeks of age. During this period, the outer nuclear layer (ONL) cell number is unchanged. The dividing cells are of photoreceptor origin, have rod opsin labeling, and do not label with markers specific for macrophages/microglia (CD18) or Müller cells (glutamine synthetase, PAX6). Nestin labeling is absent from the ONL although it labels the peripheral retina and ciliary marginal zone equally in normals and mutants. Cell proliferation is associated with increased cyclin A1 and LATS1 mRNA expression, but CRX protein expression is unchanged. Coincident with photoreceptor proliferation is a change in the photoreceptor population. Prior to cell death the photoreceptor mosaic is composed of L/M-and S-cones, and rods. After proliferation, both cone types remain, but the majority of rods are now hybrid photoreceptors that express rod opsin and, to a lesser extent, cone S-opsin, and lack NR2E3 expression. The hybrid photoreceptors renew their outer segments diffusely, a characteristic of cones. The results indicate the capacity for terminally differentiated, albeit mutant, photoreceptors to divide with mutations in this novel retinal degeneration gene.

**Introduction**

Mutations in the large repertoire of photoreceptor-specific or enriched genes are causally associated with inherited retinal diseases in both humans (RetNet: http://www.sph.uth.tmc.edu/RetNet/) and animals [1,2]. While the involved genes vary, apoptotic cell death is the final common pathway in retinal diseases [3]. This results from activation of one or several cell death pathways that appear to be mutation/model specific, and results in degeneration and death of photoreceptors with eventual blindness [4,5]. As photoreceptors are terminally differentiated, there is no compensatory neurogenesis to replace the dying cells.

Early retinal degeneration (erd) is an autosomal recessive canine retinal disorder caused by a mutation in STK38L, a novel serine-threonine kinase gene also known as nodal-related protein 2 (NDR2) [6,7]. In erd, an exonic SINE insertion eliminates part of the N-terminal regulatory region that is conserved in the nuclear Dbf2-related (NDR) subclass of AGC protein kinases [8]. These kinases, NDR1, STK38L, LATS1 and LATS2, are involved in the control cell division and morphogenesis in various cell types, including neurons [8,9]. LATS1 and STK38L are expressed in retina, and may function as tumor suppressor (LATS1), or in the control of cell death and proliferation [7,8,10]. The disease is characterized by abnormal retinal function whereby the b-wave of the electroretinogram (ERG) fails to develop, and the ERG remains a-wave dominated, an indication of abnormal synaptic transmission to second order neurons in the ONL [6]. We now report that after photoreceptor differentiation is completed in erd there is a period of sustained photoreceptor proliferation and cell death that occurs independently in different cells, and the newly generated photoreceptors are hybrid rod/S-cones. These results demonstrate that terminally differentiated photoreceptors are able to proliferate and differentiate under the appropriate stimulus, and suggest a possible role for STK38L in the control of retinal cell division.

**Results**

**Early rod defects, and rod opsin delocalization in erd**

In mutants, the early stages of retinal development were normal (Fig. 1A1, B1). Thereafter, rod outer segment (OS) shortening and loss was evident by 12.3 wks even though the outer nuclear layer...
(ONL) remained unchanged (Fig. 2A1,B1,C1). Rod opsin immunolabeling showed distinct abnormalities; OS were irregular and variable in length, and some cells showed opsin mislocalization to the ONL. These changes were present but mild at 4.3 wks (Fig. 1A2,B2), yet more marked later when there was extensive rod opsin mislocalization to the ONL and synaptic terminals, and rod opsin positive neurites sprouted into the inner nuclear layer (Fig. 2B1B2,C1C2; * and oblique arrows).

Concurrent photoreceptor cell death or proliferation in developed mutant retina

TUNEL labeling was used to examine the kinetics of photoreceptor apoptosis/cell death in the disease during (4.3 wks), or after (7.7 wks) the completion of postnatal retinal differentiation [6]. As this could only be done in paraformaldehyde-fixed cryosections, analysis was limited to 4.3–14.1 wks. In 4 wk old normal and mutant, fewer than 13 TUNEL labeled cells/106 μm2 of ONL were present (Fig. 3A), values similar to those reported in a previous study of normal dogs [11]. However, in mutants the number of TUNEL positive cells markedly increased thereafter, and labeled nuclei were distributed uniformly throughout the ONL, including the outermost level where cone somatas were located; similar numbers of TUNEL positive cells were found in central, equatorial and peripheral regions. The high number of apoptotic cells indicated that there was active and sustained cell death occurring in the 7.7–14.1 wk time period that was limited to the ONL.

Despite the ongoing photoreceptor cell death demonstrated by TUNEL labeling, there was no apparent loss of ONL cells (Fig. 3B). The ONL thickness in mutant retinas was constant and equal to that in controls until 14.1 wks of age. Thereafter the ONL decreased dramatically by 40.1 wks, and became even thinner at 62 wks.

To examine why ONL thickness did not change in spite of the high rate of photoreceptor apoptosis, we determined whether cell proliferation was occurring using an antibody directed against proliferative cell nuclear antigen (PCNA). Labeled cells were present in the mutant ONL during the period of sustained cell death, with values ranging from 40–113 PCNA labeled cells/106 μm2 (Fig. 3C). The PCNA labeled nuclei in the ONL were morphologically similar to the TUNEL positive ones. To further characterize the cell proliferation findings in terms of cell location and retinal distribution, selected retinal sections from control (7 and 9 wks) and mutants (7.7 and 9.1 wks) were labeled with KI-67, another marker of cell division, and demonstrated a similar ONL labeling pattern as with PCNA (data not shown).

Phospho-histone H3 (PHH3) labeling is limited to mutant photoreceptor cells

PHH3 labeling was used to differentiate mitotic cells [12,13] from those undergoing DNA repair that labeled with PCNA [14]. In normals, PHH3 labeled nuclei were located adjacent to the external limiting membrane, and, in the immediate postnatal period, limited to the retinal periphery at the time that the outer neuroblastic layer was separating, and the outer plexiform layer (OPL) had just formed (Fig. 4A, arrows). In control dogs 4 wks or older, there were almost no PHH3 labeled cells in ONL. In mutants, on the other hand, PHH3 labeled nuclei only were
present at different levels of the ONL, and these were small, round and distributed uniformly from the center to the periphery; labeled cells were distinct from those undergoing apoptosis (Fig. 4B-E). Rod opsins and PHH3 labeling clearly demonstrated colocalization, and labeled nuclei were enclosed by a rod opsin positive cytoplasmic rim (Fig. 4F,G,G1–4).

We have examined a subset of samples and calculated the number of PHH3 positive cells using the same method used for counting PCNA positive cells (see Materials and Methods). The results indicate that comparable numbers of labeled nuclei in the ONL, expressed as labeled cells/10^6 µm^2, were present with both PCNA (7 wk control = 2 ± 1; errd 7.7 wk = 61 ± 16 and 11.6 wk = 91 ± 30) and PHH3 (7 wk control = 4 ± 2.3; errd 7.7 wk = 153 ± 56 and 11.6 wk = 100 ± 11) labeling.

Müller cells, stem cells or microglia do not contribute to the population of dividing ONL cells

Double labeling with PHH3 and glutamine synthetase (GS) was used to rule out Müller cell contribution to the dividing cell population. Distinct and comparable GS labeling was present in the normal and mutant retinas, and extended from the external to the internal limiting membranes. In errd, GS labeling was not associated with PHH3 labeled nuclei in the ONL (Fig. S1). An antibody against nestin, expressed in neuronal stem cells, was used to label the normal and mutant retinas, and extended from the external to the internal limiting membranes. In errd, GS labeling was not associated with PHH3 labeled nuclei in the ONL (Fig. S1).

The COS-1 and OS-2 antibodies, respectively, labeled the L/M- and S-cone classes comparably in control and mutants, and labeling was restricted to the OS (Fig. 1A3,A4, 5A,B). Mutant cones, on the other hand, appeared shorter in younger animals (Fig. 1B3,B4), and some failed to show hCAR labeling even though the cells were readily identifiable in transmitted light with or without DIC optics. Qualitative assessment of cone numbers based on PNA labeling was normal; however, immunolabeling with antibodies against cone opsin indicated that in the older mutants the OS were shorter and irregular (Fig. 5A,B).

The COS-1 and OS-2 antibodies, respectively, labeled the L/M- and S-cone classes comparably in control and mutants, and labeling was restricted to the OS (Fig. 1A3,A4, 5A,B). To determine qualitative immunolabeling intensity differences between normals and mutants, an antibody dilution series was carried out. The minimal dilution required for labeling S-cones was the same for control and mutants for the 3 antibodies (monoclonal OS-2 (1:400), goat sc-14363 (1:20,000), rabbit AB5407 (1:40,000)). In contrast, polyclonal antibodies sc-22117 and AB5405 against the L/M-cones resulted in distinct labeling at dilutions of 1:20,000 and 1:80,000, respectively in errd, but required double that concentration (1:10,000 and 1:40,000) for comparable labeling in normal cones.

An unexpected finding was the labeling of both S-cones and rod OS with the OS-2 antibody. This was observed in the 7.7–14.1 wk time period, but not at 4.3 wks where OS-2 labeling was restricted to S-cones, and similar to control (Figure 1A4,B4). At 7.7 wks, however, OS-2 labeling of rod OS was faint and...
homogeneous, but clearly localized to the photoreceptor OS layer. By 11.6–14.1 wks, labeling of rods was more intense (Figure 5B). Double labeling with OS-2 and rod opsin antibodies indicated that many of these rods showed colocalization of both proteins in the same OS (Figure 5C). In contrast, labeling with COS-1 or the two polyclonal red/green cone opsin antibodies indicated that label was restricted only to hCAR positive L/M-cones (Figure 5A).

Figure 3. Photoreceptor cell death and proliferation in erd. (A, C) TUNEL and PCNA labeling, respectively, in erd is sustained between 7.7–14.1 wks with many labeled cells in the ONL of the superior (Sup.) and inferior (Inf.) meridians (data from both quadrants combined for normal; data points expressed as mean ± 1 SD). Color insets illustrate the labeled cells (green) from a 7.7 wk old mutant animal for the corresponding assay in sections with hCAR antibody that labels all cones (red). (B) The number of photoreceptors in the outer nuclear layer (ONL), expressed as the mean number of rows of nuclei, remains relatively constant until 14.1 wks, and then decreases. Sup.A1/Inf.A1 = superior/inferior area 1, 2000 ± 500 μm from the optic disc; Sup.A2/Inf.A2 = superior/inferior area 2, mid point ± 500 μm between optic disc and ora serrata. For the ONL, the mean nuclei counts are presented. Between 4.3–14.1 wks of age, all the SD for ONL were ≤15% of the mean, with greater than 60% of the values being less than 10% of the mean.

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To determine if rod OS labeling was dependent on antibody concentration, we carried out a qualitative immunolocalization comparison between control (7 wks) and mutant (7.7 and 11.6 wks) retinas using different OS-2 antibody dilutions. In controls, S-cone OS were intensely labeled at 1:100 and 1:200 dilutions, but weakly at 1:400, and rods were unlabeled. The mutant retina showed labeling of rod or S-cone OS at 1:100 and 1:200, and the intensity, by photoreceptor class, was equal at each dilution. At 1:400 only S-cone OS were labeled with an intensity comparable to lower dilutions.

As the transcription factors NRL and NR2E3 are involved in retinal cell fate specification, we examined the expression of these two gene products in normal and mutants at the time when hybrid rod/S-cones were present. Both show NRL expression in rod and cone inner segments (IS), cone OS and rod nuclei; as well as rodents expressed NRL in presumptive hybrid rod/S-cones as these cells lack a PNA labeled domain (Fig. 5D). In contrast, NR2E3 is mainly expressed in nuclei of rods in normals, although occasional labeling of cone nuclei and IS also was observed. The mutant retina showed absence of NR2E3 labeling in the ONL or photoreceptor layer (Fig. 5D).

Gene expression in normal and mutant retinas

We used qRT-PCR to characterize retinal expression of the mutated gene, STK38L, and another member of the NDR family, LATS1, at different ages during normal development and disease (Fig. 6A). STK38L expression in control retinas was unchanged during development. Although mutant mRNA lacked exon 4, the altered transcript showed a slight increase in expression at the 2 older disease time periods examined when using an exon 6 probe (0.05 ≤ p ≤ 0.1). LATS1 expression was increased at the 3 wk time point in normals, and at the 2 older disease time points. As well, we examined by qRT-PCR the expression pattern of two cell cycle genes. The expression pattern of CCNA1 was similar to LATS1, but CCND1 expression was minimally reduced or unchanged at the three disease stages examined (Fig. 6B).

Immunoblotting demonstrated increased expression of three of four photoreceptor-specific proteins evaluated (S-opsin, L/M-opsin and RDS peripherin) in lysates from 6.4 and 9.9 wk erd-retinas compared to 8 wk normal control (Figure 7A,C). The CRX transcription factor was unchanged, but NR2E3 was significantly increased at 6.4 wks, whereas NRL levels were elevated only at 9.9 wks (Figure 7B,C). The NR2E3 results differed from those obtained by immunocytochemistry, and could be the result of different antibodies used in the blots and tissue sections as well as accessibility of antibody to epitope in the sections.

Cone-like outer segment renewal in erd mutant retina

Autoradiography of control retinas following intravitreal 3H-leucine or 3H-fucose injection showed renewal by band displacement...
in OS characteristic of rods (Figure 8A1–A3, horizontal arrows). Calculated renewal rates were $2.13 \pm 0.08 \text{ mm/day}$ for central and mid peripheral regions. In mutants, a distinct band of radioactivity was not appreciable at any time, and a renewal rate could not be established (Figure 8A4–A6; } illustrates diffuse label). Diffuse label was present at all levels of the OS layer, and the labeling pattern was similar in rods and cones. At the 4 day post injection point, this difference clearly was apparent; in controls, the band of radioactivity, and a trailing label tail, was present distal to the rod OS midpoint, but mutants demonstrated diffuse label and no band.

Normal rod opsin biosynthesis in mutant retina, but abnormal renewal kinetics

Relative incorporation of $^3$H-fucose/$^{14}$C-leucine into normal and $erd$ rod opsin differed (Figure 8B). In normals, opsin labeling with both precursors increased at post injection days 2 and 4 as the labeled precursor pool in the vitreous continued to be incorporated into newly synthesized protein. Regression analysis of combined $^3$H-fucose and $^{14}$C-leucine data from controls yielded a slope (daily increase in label) of $0.225 \pm 0.0746$. In contrast, the mutant retinas demonstrated decreased labeling at 2 and 4 days. Regression analysis on the combined data yielded a slope of $-0.296 \pm 0.0459$, an indication of loss of radioactivity over the time period examined. The difference between control and affected slopes was highly significant ($Z = 5.9401, p = 1.43E-09$).

Discussion

In the $STK38L$ mutant retina, rods and cones begin to differentiate normally, and express polarized molecular markers,
e.g. opsins and cone arrestin, indicating that cell class specification occurred after the final cell division [17]. Subsequently, rods show disparate lengths of the OS, neurite sprouting, and opsin mislocalization. These changes are soon followed by cell death primarily affecting rods and then cones. Apoptosis detected by TUNEL labeling occurs in terminally differentiated photoreceptors after retinal development is completed; it is sustained, and many dying cells are present throughout the ONL between 7.7–14.1 wks of age [6].

An unexpected finding, however, is that the high levels of TUNEL labeling is not accompanied by concomitant photoreceptor loss as is characteristic for inherited retinal degenerations, and the mutant ONL remains at a constant thickness until after 14.1 wks of age [5,11]. This suggests that a concurrent, compensatory proliferative event must be occurring in mutants, and was confirmed by PCNA labeling. The antibody labeled many cells in the erd ONL, but almost none in controls. PHH3 and rod opsin double labeling confirmed that photoreceptor cells were undergoing mitosis rather than DNA repair as the number of labeled cells were comparable, and there was no co-localization of TUNEL and PHH3 labeling [13,14]. Furthermore, as labeling with antibodies against CD18, GS, PAX6 and nestin ruled out the possibility that the ONL mitotic cells are microglia, Müller or stem cells, we conclude that the proliferating cells must be photoreceptors.

We have considered the possibility that the proliferating and dying cells are the same, and that lack of colocalization of TUNEL and PHH3 labeling results from the cells dying at a stage in the cell cycle when the proliferation marker is not expressed. This possibility, however, is not consistent with the results. If cell proliferation were occurring, and TUNEL labeling reflects only those cells that are replicating their DNA, we would have expected to see a marked increase in ONL thickness. We previously have observed such proliferative events in young dogs with an RPGR mutation following intravitreal CNTF administration [18]. Thus between 7.7 and 14.1 wks of age, mutant photoreceptors are undergoing apoptosis or cell division. As the ONL thickness remains unchanged during this time, the magnitude of both events has to be comparable, otherwise decreases or increases in ONL cell number would occur. Because rods are uniformly distributed throughout the ONL and outnumber cones 22:1 outside the area centralis [19], it is likely that rods are the predominant cells that are dying or proliferating. The absence of TUNEL or PCNA labeled cells at the earlier time point suggests that terminally differentiated albeit mutant photoreceptors commit to the cell death or proliferation fate.

Neurogenesis in the developed mammalian retina or CNS is limited. In the brain, the subventricular zone of the lateral ventricle, and the dentate gyrus of the hippocampus show low level of neurogenesis from neuronal progenitors [20,21]. In the retina, and under specific circumstances [22], a limited number of photoreceptors and other neurons are generated from presumably terminally differentiated Müller cells that dedifferentiate, proliferate and express neuronal progenitor markers; this has been reported in the adult rat, mouse, chicken and fish (see [23] for review). In contrast, rod photoreceptor replacement in zebrafish is dependent on the extent of injury. Total rod ablation results in robust Müller cell proliferation and formation of neuronal progenitor clusters, while ablation of a subset of rods results in proliferation of rod precursors [24].

Other than the present study, generation of new photoreceptors in a naturally occurring retinal degeneration has not been reported. A previous study suggested the possibility of photoreceptor proliferation based on PCNA and BrdU labeling in rd1 retinas [25]. However, specific labeling for microglia subsequently showed that all dividing cells in the ONL were microglia rather
than photoreceptors [26]. Thus the STK38L mutant retina appears
to be unique in having terminally differentiated photoreceptors
undergoing cell proliferation.

What is not clear at this time are the signals that commit
terminally differentiated photoreceptors to die or divide. While
apoptosis is the final common pathway in retinal degenerative
diseases, there are multiple potential pathways that link the
mutation to the apoptotic event, and these are yet to be defined for
most species including dogs [3,5]. In ird, cell proliferation likely
results from loss of part of the N-terminal regulatory region that is
highly conserved in all NDR subclass of AGC protein kinases [8].
The exonic SINE insertion removes exon 4 from the mature RNA,
and eliminates the binding sites for S100B and Mob proteins, part
of the protein kinase domain, and a Thr-75 residue critical for
autophosphorylation [7]. We posit that in the normal retina, the
terminally differentiated photoreceptors are kept from dividing by
NDR2-Mob1 interaction. Removing this control in mutants allows
the cell to re-enter the cell cycle and divide, as suggested by
increased cyclin A1 expression [8]. Increased expression of LAT31
could be an attempt to suppress this proliferation. Photoreceptor
cell division, however, is temporally limited as the ONL thins after
14.1 wks. The cell division events are similar to what has been
observed with the transgenic expression of SV40 large tumor
antigen in mouse retina that results in postnatal DNA synthesis
and mitosis. A major difference from ird, however, is that after
mitosis, the transgenic photoreceptors do not progress to a viable
postmitotic stage and die [27].

Coincident with the photoreceptor cell death and proliferation
phase is a change in the visual cell population from the preexisting
normal mosaic of L/M- and S-cones, and rods. Although both
cone types remain, most rod cells that presumably were generated
after proliferation now become hybrid photoreceptors that express
both rod opsin and cone S-opsin. This is supported by absent
NR2E3 expression in mutant photoreceptors, at least by
immunocytochemistry, resulting in rod-like photoreceptors that
express cone genes [28,29]. The discrepancy between the
immunocytochemistry and western blotting results for NR2E3 in
ird is difficult to reconcile at this time. As the results obtained in
control retinas are comparable to what has been reported for other
mammalian retinas, e.g. mouse and Nile rat [30], and monkey
[31], we posit that the antibody is detecting NR2E3, and not an
unrelated antigen. Lack of rod labeling in the mutant retina would
suggest that the NR2E3 epitope is masked by the expression of the
mutant STK38L protein. However, the mechanism for this

Figure 7. Protein expression in ird retina. Expression of (A) CNGA3, L/M-opsin, S-opsin, RDS, and (B) NRL, NR2E3, CRX proteins in cleared whole
retinal lysates. (C) Net intensity densitometric values are expressed as corrected net intensity of the sample normalized to corrected net intensity of
actin loading controls. Actin loading control is the same for S-opsin and RDS, and for NRL, NR2E3 and CRX, respectively. Results are the mean ± SD of
four independent densitometer scans of two individual westerns. Confidence levels: * = 95%, ** = 99%.
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finding is unknown, and we plan to address this question experimentally with GST pull down assays to examine for binding partners of mutant STK38L.

Further evidence of this change in the photoreceptor population from rods to hybrid rod/S-cones comes from the rod OS renewal and opsin biosynthesis studies. A fundamental difference between rods and cones is in the renewal of OS disc proteins [32]. Rod disc membranes are isolated, and autoradiograms following pulse labeling of newly formed discs with radiolabeled precursors show a distinct band of radioactivity that is displaced distally until shed from the OS tip. In contrast, cone OS disc membranes are not isolated, but continuous, and redistribution of labeled proteins by lateral diffusion results in a diffuse pattern of labeling [32,33]. In *erd*, renewal was diffuse, and, unlike normals, a distinct labeled band could not be identified at any time point. Similar results could occur if OS morphogenesis is decreased, but only if the process is asynchronous as a uniform decrease would still result in renewal by band displacement, albeit at a slower rate [34].

The complementary biochemical studies support the interpretation that newly synthesized radiolabeled opsin redistributes rapidly throughout the mutant OS, and loss of label occurs associated with the daily shedding of discs from the distal tips of

Figure 8. Abnormal outer segment renewal in *erd* rods. (A1–3) Rod outer segment renewal examined 4 days following intravitreal injection of \(^{3}H\)-fucose in 10.9 week old normal and (A4–6) *erd*-affected dogs. The images (A1,A2 and A4,A5) are serial sections taken from the posterior pole of the superior retinal quadrant. The transmitted bright-field autoradiogram (A2) shows in the normal a band of labeling whose leading edge extends to 2/3 of the rod OS length (A2, horizontal arrow) with a trailing tail of radioactivity. This is better visualized in the combined transmitted/epipolarizing image (A3, arrow) that also shows diffuse label in the cone OS (oblique arrows). The mutant retina (A4–6) shows diffuse label at all levels of the OS layer (A5,A6; brackets) that is similar in rods and cones. RPE = retinal pigment epithelium; OS = outer segments; IS = inner segments; ELM = external limiting membrane; ONL = outer nuclear layer. (B) Relative incorporation of \(^{3}H\)-fucose/\(^{14}C\)-leucine into opsin at different time points following a single intravitreal injection. All data are normalized to incorporation of label at day 1. The *erd* outer segments initially take up both labels (day 1), but label intensity decreases at later post injection times, suggesting that label is diffusely distributed through the rod outer segment membranes after discs are formed, and outer segment tips are lost to the RPE by daily shedding/phagocytosis events.

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the modified OS [32,35]. This conclusion is based on finding comparable incorporation of both radiolabeled precursors into mutant and control rod opsin initially, but labeling decreased in mutants unlike normals. Decreased OS morphogenesis would result in a lower incorporation of label at the 2 and 4 day post injection time points. However, this lower rate of incorporation still would result in increased labeling over time, and not the decrease found in our study. Thus the change from rod to hybrid rod/S-cone is associated with a change in rod OS structure where the membranous discs now become continuous with the plasma membrane. This would require not only a change in expression of the membrane associated proteins such as rod opsin and S-opsin, but also expression of a new repertoire of genes and proteins that determine photoreceptor OS structure. One such protein, RDS, is involved in cone OS biogenesis and maintenance, and appears to require formation of covalently linked RDS dimers to carry out this function [36]. Although RDS levels were only modestly elevated at 6.4 wks, distinct dimer formation was evident in the western blots.

Our results indicate that in the STK38L mutant retina terminally differentiated photoreceptors undergo cell division and differentiate into hybrid rod/S-cone photoreceptors. Because of the paucity of cones in the canine retina, and the finding that most of the proliferating cells are rods, we assume that reconstitution of the photoreceptor layer is rod-derived, although a cone contribution can not be excluded. The newly generated cells return to either the pool of rod or cone precursors, or upstream to the pool of undifferentiated postmitotic photoreceptor cells [37]. Although we have no direct experimental data to support either alternative, the low levels of CRX (western blots), and absent NR2E3 photoreceptor layer labeling at the time when a new population of photoreceptors is being generated argues for the newly produced cells to come from the rod and possibly the cone precursor pool. However, a cautious interpretation of these findings is important given that expression changes in the eld retina, which is terminally differentiated yet photoreceptors are dying or dividing, may have limited similarities to what happens in normal retinal development with expression of transcription factor genes involved in photoreceptor cell specification [17].

In summary, the STK38L mutant retina is atypical in that terminally differentiated photoreceptors undergo cell death or proliferation, and generate a new class of hybrid rod/S-cone photoreceptors. The results suggest a role for STK38L in the control of cell division and morphogenesis in photoreceptors and possibly other retinal neurons. Future studies will inform on the role of the N-terminal regulatory region in these functions, and suggest approaches that can be manipulated experimentally to reconstitute and preserve a diseased photoreceptor layer.

Materials and Methods

Animals

Dogs were maintained at the Retinal Disease Studies (RDS) facility in Kennett Square, PA, and supported by NEI/NIH (EY-06835) and Foundation Fighting Blindness Center grants. The study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health, and adhered to the ARVO Resolution 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 Protocol #s 801870, 802467, 803269), and all efforts were made to minimize suffering. Intravitreal injections for outer segment renewal and opsin biosynthesis studies were made in dogs anesthetized with thiopental sodium; tissue collection was performed under deep pentobarbital anesthesia, and was followed by euthanasia with an overdose of the same or comparable (Ethanol) agent. The dogs represent an outbred population with a common genetic background segregating eld and other retinal disease alleles [1]. The genotypes, ages studied and procedures performed are detailed in Table 1.

Anatomic/immunochemical studies

For structural, cell counting, immunocytochemical, TUNEL and cell proliferation studies, eyecups were fixed in 4% paraformaldehyde (PF), and central superior and inferior retinal strips (optic disc to ora serrata) were embedded in optimal cutting temperature medium (OCT; Sakura Fientek, Torrance, CA) using standard methods [38]. Archival tissues fixed in Bouin’s solution and stored in 70% ethanol from affected animals were used for structural and cell counting studies. Sections from paraffin embedded tissues were cut in the dorso-ventral plane through the pupil-optic disc axis.

Sections from both the superior and inferior meridians were examined in contiguous fields from the optic disc to the ora serrata; this included evaluation of the rod and cone IS and OS, and the thickness of the ONL. For each dog, a section from the superior and inferior quadrants was used for quantitative evaluation of the ONL cell counts at two specific locations: A1 = 2000±500 μm from the optic nerve edge, and A2 = mid-point (equidistant from optic disc and ora serrata) of the retina ±500 μm. At each of these sites, the number of rows of nuclei in the ONL were counted in three areas of a 40 x field and averaged.

Fluorescence immunohistochemistry was done on 7 μm cryosections taken from the superior and inferior retinal meridians, and incubated overnight at 4°C or at room temperature for 1 hour with the primary antibodies after a blocking step with 1.5% BSA/ PBS, 0.25% Triton X-100 (Sigma-Aldrich, St. Louis, MO). The primary antibodies and cell markers used, cell class specificity, and target protein are detailed in Table S1, and include: cones: human cone arrestin, PNA, CNGA3, L/M- (COS-1) or S- (OS-2) cone opsins [39], red/green cone opsin, blue cone opsin; rods: rod opsin; Müller cells, microglia and macrophages: PAX6, GS, CD18; cell proliferation, retinal stem cells: PCNA, Ki-67, PHH3, Nestin; transcription factors: PAX6, NR2E3, Nrl.

Apoptotic nuclei were visualized by TUNEL (terminal deoxynucleotidyl transferase-mediated biotinylated UTP nick end labeling) with the In Situ Cell Death Detection kit (Roche Applied Science, Indianapolis, IN). Antigen retrieval was performed prior to PCNA and PHH3 labeling by heating in the presence of Antigen Unmasking Solution, High pH (Vector Laboratories, Burlingame, CA) using a microwave oven at 10% power.

Both single and double immunolabeling was used. The primary antibody pairs used for double immunolabeling were combinations of rabbit or goat polyclonal and mouse monoclonal antibodies. The antigen–antibody complexes were visualized with fluochrome-labeled secondary antibodies (Alexa Fluor, 1:200; Invitrogen, Carlsbad, CA), and 4’,6’-diamino-2-phenylindole (DAPI) stain was used to label cell nuclei. Slides were mounted with a medium composed of polyvinyl alcohol and DABCO (1,4 diazobiszydko-[2-2,2]oktan) (Gelvatol; Sigma-Aldrich), and examined with an epifluorescence microscope (Axioplan; Carl Zeiss Meditec, Thornwood, NY). Epifluorescence or transmitted light images were captured with a Spot 4.0 camera (Diagnostic Instruments, Inc., Sterling Heights, MI) and imported into a graphics program (Photoshop and Illustrator; Adobe, San Jose, CA) for display. When precise localization of markers was needed, sections were also imaged by confocal microscopy using a Nikon
### Table 1. Summary of procedures and dogs used in the studies.

| Studies                     | Disease Status | Dogs (or eyes) | Ages-wks (#/time point) | Procedures                                                                 |
|-----------------------------|----------------|----------------|--------------------------|-----------------------------------------------------------------------------|
| **ANATOMY**                 |                |                |                          |                                                                               |
| Control*                    | 11             | Preterm, 1, 1.6, 4, 4.7, 7, 8.7, 9, 12.3 (2), 25.7   | Paraformaldehyde fixation for immunohistochemistry, TUNEL, cell counting, structural assessment |
| Mutant                      | 13             | 4.3, 7.7 (2), 8.3, 8.4, 9.1 (2), 9.3, 10.4, 11.6 (2), 12.3, 14.1 |                                     |
| Mutant                      | 11             | 7.1 (3), 7.3, 9.1 (2), 48.1, 62, 68, 101, 165       | Bouin’s fixation for cell counting, structural assessment                      |
| **GENE/PROTEIN EXPRESSION** |                |                |                          |                                                                               |
| Control                     | 11             | 3 (3), 7 (3), 7.4, 8.2, 16 (3)                       | qRT-PCR, western analysis                                                     |
| Mutant                      | 10             | 6.4 (3), 8.3, 9.6, 9.9 (2), 11.9(2), 14.1            |                                     |
| **OUTER SEGMENT RENEWAL**   |                |                |                          |                                                                               |
| Control**                   | (4)            | 8.7, 10.9, 16.4 (2)                                   | Light microscopic autoradiography with ³H-leucine (6 eyes) or ³H-fucose (6 eyes) |
| Mutant                      | (8)            | 8.3, 8.7, 9 (3), 10.9                                 |                                     |
| **OPSIN SYNTHESIS**         |                |                |                          |                                                                               |
| Control**                   | (9)            | 9.9 (2), 16.4 (4)                                     | Dual labeling with ³H-fucose/¹⁴C-leucine, opsin isolation, quantification of radioactivity |
| Mutant                      | (15)           | 8.3 (6), 9 (4)                                        |                                     |

*one control dog is erd heterozygous.  
*four control dogs are erd heterozygous.  
†-fellow eyes from 5 dogs (2 control; 3 erd affected) were used in the outer segment renewal and opsin synthesis studies.

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A1R Laser Scanning Confocal Microscope with DUS 32 Spectral detector (Nikon Instruments, Melville, NY) through a 63× Plan APO objective lens with 1.2 numerical aperture. The specimens were excited at 488 and 561 nm, respectively, with multi line Argon and DPSS lasers.

TUNEL-, PCNA- or PHH3-labeled cells in the ONL were counted throughout the entire length of the section. In determining the proportion of photoreceptor cells that undergo cell death or proliferation as a function of time, the results were expressed as the number of TUNEL, PCNA or PHH3 labeled photoreceptor cells per 10⁶ m² of ONL [11]. The area of the ONL of each section was obtained by measuring the entire length of the ONL from the optic disc to ora serrata, and multiplying it by the average thickness of the ONL throughout the section (mean value of the thickness measured in ten evenly distributed locations). For each retina examined for TUNEL, PCNA or PHH3 labeling the procedure was performed in triplicate with working concentrations (Table S1). These results were validated in repeat analyses.

Gene expression

Quantitative real-time PCR (qRT-PCR) was used to assess expression of selected genes at different time points of normal development (3 (n = 5) and 7 (n = 5) wks) and disease (6.4 (n = 2), 8.3/9.9 (n = 3) and 11.9/14.1 (n = 2) wks). Retinas from 16 wk (n = 3) normal dogs served as reference control. Total RNA was isolated by Trizol extraction (Invitrogen-Life Technologies, Carlsbad, CA), and concentration measured with a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). RNA samples were treated with RNase-free DNase (Ambion, Austin, Tx), and two µg total RNA from each sample was used for cDNA synthesis using the High Capacity cDNA reverse transcriptase Kit (Applied Biosystems (ABI), Foster City, CA). Quantitative RT-PCR was performed on a 7500 Real Time PCR System and software v2.0 (ABI) using 30 ng cDNA from each sample, and amplified using Taqman assays with canine-specific ABI invented probes for STK38L/NDR2, LATS1 (Cf02627675_m1) and GAPDH (Cf02634245_m1); SYBR green analysis was used for CCND1 (Forward: CATCTACACTGACAACTCCATCC; Reverse: CAGGTTCACGTACCTGTTTGTCC). For analysis of STK38L/NDR2 one probe was located within the exonic deletion (STK38L (exon 4) = Cf02709226_m1), and the second in exon 6, 3’ to the exon 4 splicing defect, and used to exclude alterations in splicing resulting from the SINE element insertion (STK38L (exon 6) = Cf02634613_m1). CT values of each gene were normalized to GAPDH, and comparisons between groups were done with the ΔΔCT method [40]. Statistical significance between different groups in comparisons to the 16 wks old normal control retinas was assessed with an unpaired t-test, and expressed as statistically significant (p≤0.05) or towards statistical significant (0.05≤p≤0.1).

Immunoblotting

Equal amounts of total protein as determined by BCA Protein Assay Kit (former Pierce Biotechnology now Thermo Fisher Scientific, Rockford, IL) were separated by 10% SDS-PAGE under reducing conditions, immunoblotted and probed with antibodies. These are detailed in Table S1 and include antibodies

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We determined qualitative differences in immunolabeling intensity between normal and mutant retinas with antibodies directed against S- and L/M-cones. For this, sections (2) from selected age groups were incubated with different dilutions of the antibodies to determine the lowest concentration needed to provide comparable results to those obtained with the routine working concentrations (Table S1). These results were validated in repeat analyses.

Gene expression

Quantitative real-time PCR (qRT-PCR) was used to assess expression of selected genes at different time points of normal development (3 (n = 5) and 7 (n = 5) wks) and disease (6.4 (n = 2), 8.3/9.9 (n = 3) and 11.9/14.1 (n = 2) wks). Retinas from 16 wk (n = 3) normal dogs served as reference control. Total RNA was isolated by Trizol extraction (Invitrogen-Life Technologies, Carlsbad, CA), and concentration measured with a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). RNA samples were treated with RNase-free DNase (Ambion, Austin, Tx), and two µg total RNA from each sample was used for cDNA synthesis using the High Capacity cDNA reverse transcriptase Kit (Applied Biosystems (ABI), Foster City, CA). Quantitative RT-PCR was performed on a 7500 Real Time PCR System and software v2.0 (ABI) using 30 ng cDNA from each sample, and amplified using Taqman assays with canine-specific ABI invented probes for STK38L/NDR2, LATS1 (Cf02627675_m1) and CCND1 (Cf02634245_m1); SYBR green analysis was used for CCND1 (Forward: CATCTACACTGACAACTCCATCC; Reverse: CAGGTTCACGTACCTGTTTGTCC). For analysis of STK38L/NDR2 one probe was located within the exonic deletion (STK38L (exon 4) = Cf02709226_m1), and the second in exon 6, 3’ to the exon 4 splicing defect, and used to exclude alterations in splicing resulting from the SINE element insertion (STK38L (exon 6) = Cf02634613_m1). CT values of each gene were normalized to GAPDH, and comparisons between groups were done with the ΔΔCT method [40]. Statistical significance between different groups in comparisons to the 16 wks old normal control retinas was assessed with an unpaired t-test, and expressed as statistically significant (p≤0.05) or towards statistical significant (0.05≤p≤0.1).

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Equal amounts of total protein as determined by BCA Protein Assay Kit (former Pierce Biotechnology now Thermo Fisher Scientific, Rockford, IL) were separated by 10% SDS-PAGE under reducing conditions, immunoblotted and probed with antibodies. These are detailed in Table S1 and include antibodies
against red/green opsin, blue opsin, CNGA3, NRL, NR2E3, RDS/peripherin, and CRX. Appropriate HRP-conjugated secondary antibodies were used subsequently, and an antibody against β-actin was used as a loading control. Densitometric analysis of the blots was performed on a Molecular Imaging System (Carestream Health, Rochester, NY). The net intensity was corrected for background intensity observed. Values in figure are expressed as corrected net intensity of the sample normalized to corrected net intensity of actin loading controls. Densitometry data were analyzed using Sigma Stat Version 3.1, and intensities from four independent measurements on two westerns were analyzed by Student t test using 95% or 99% confidence intervals.

Rod outer segment renewal and opsin biosynthesis

Rod OS renewal and opsin biosynthesis were examined at specific time points following the injection of 3H-fucose or 3H-leucine (OS renewal) or a combination of 3H-fucose/14C-leucine (opsin biosynthesis) into the vitreous of anesthetized dogs using previously described methods [34]. Post-injection time points were 1, 2, 3 and 4 days for OS renewal, and 1, 2 and 4 days for opsin biosynthesis (Table 1). In the 4 day interval following injection, the rods have renewed ~50% of their OS due to continuous addition of new discs at the base [34]. For the renewal studies, the eyes were fixed in mixed-aldehyde/osmium solution, embedded in plastic resin, and 1 μm thick sections were coated with a photographic emulsion and maintained at 4°C in the dark until developed [34]. Two series of experiments were conducted to examine opsin biosynthesis, and loss of opsin labeling over time after injection. For these studies, eyes received a combination of 3H-fucose and 14C-leucine intravitreally. Eyes from anesthetized dark adapted dogs were enucleated under dim red light, and crude rod outer segment preparations were made by vortexing and centrifugation in 40% (w/v) sucrose. After sonication, detergent solubilized rod OS proteins were separated in a 10% polyacrylamide gel, stained with Coomassie blue, and gel slices digested and counted in a scintillation counter [34]. Disintegrations per minute (DPM) in the opsin peaks were normalized to the highest protein value for each set of gels, and to the DPM count for the post injection day 1 control eye of the same series.

Supporting Information

Figure S1 Phospho-histone H3 (PHH3, red) and glutamine synthetase (GS, green) double labeling of evd mutant retinas. (A1, B1) Merged images of retinas at 7.7 (A) and 11.6 (B) weeks of age show PHH3 labeled nuclei only in ONL (arrows); (A2, B2) the images of GS labeling shows the lack of label in spaces (arrows) occupied by the PHH3 labeled nuclei. Arrowheads in A1, A2 point to intensely labeled end feet of GS positive cells; * = large retinal ganglion cell surrounded by GS positive processes. Scale bar = 20 μm; DAPI (blue) nuclear staining.

Table S1 Antibodies and reagents used for immunohistochemistry and immunoblotting.

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Author Contributions

Conceived and designed the experiments: GDA WAB AIB. Performed the experiments: AIB KB-B SG OG PJO GMA WAB GDA. Analyzed the data: AIB KB-B SG OG PJO AS GMA WAB GDA. Contributed reagents/materials/analysis tools: AS. Wrote the paper: GDA WAB AIB.

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