Inner retinal change in a novel rd1-FTL mouse model of retinal degeneration

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While photoreceptor loss is the most devastating result of inherited retinal degenerations such as retinitis pigmentosa, inner retinal neurons also undergo significant alteration. Detailing these changes has become important as many vision restorative therapies target the remaining neurons. In this study, the rd1-Fos-Tau-LacZ (rd1-FTL) mouse model was used to explore inner retinal change at a late stage of retinal degeneration, after the loss of photoreceptor nuclei. The rd1-FTL model carries a mutation in the phosphodiesterase gene, Pde6b, and an axonally targeted transgenic beta galactosidase reporter system under the control of the c-fos promoter. Retinae of transgenic rd1-FTL mice and control FTL animals aged 2–12 months were processed for indirect fluorescence immunocytochemistry. At 2 months of age, a time when the majority of photoreceptor nuclei are lost, there was negligible c-fos reporter (FTL) expression, however, from 4 months, reporter expression was observed to increase within subpopulations of amacrine and ganglion cells within the central retina. These areas of inner retinal FTL expression coincided with regions that contained aberrant Müller cells. Specifically, these cells exhibited reduced glutamine synthetase and Kir4.1 immunolabelling, whilst showing evidence of proliferative gliosis (increased cyclinD1 and glial fibrillary acidic protein expression). These changes were limited to distinct regions where cone photoreceptor terminals were absent. Overall, these results highlight that distinct areas of the rd1-FTL central retina undergo significant glial alterations after cone photoreceptor loss. These areas coincide with up-regulation of the c-fos reporter in the inner retina, which may represent a change in neuronal function/plasticity. The rd1-FTL mouse is a useful model system to probe changes that occur in the inner retina at later stages of retinal degeneration.

Keywords: retinitis pigmentosa, Müller cell, c-fos, ganglion cell, cone photoreceptor, plasticity

Introduction

Over recent years there has been a surge of interest in vision restoring therapies that replace the function of dead or dying photoreceptors using a variety of genetic, electronic or transplantation methods (Busskamp et al., 2008; O’Brien et al., 2012). For any of these treatments to succeed, a basic requirement is that the circuitry of the inner retina remains functionally intact and capable of passing visual information to higher visual centers. Yet, over recent years, the notion that the
inner retina remains intact following complete loss of photoreceptors has been questioned (Jones et al., 2003; Marc et al., 2003, 2007; Strettoi et al., 2003; Jones and Marc, 2005; Chua et al., 2009). A range of changes occur in both neurons and glial cells in animal models of inherited retinal degeneration, as well as humans with retinitis pigmentosa (RP). These changes are classified into three phases based on retinal anatomy (Jones and Marc, 2005); Phases 1 and 2 refer to the periods of rod photoreceptor and cone photoreceptor death, respectively, while phase 3 is characterized by major anatomical changes in the inner retina including re-wiring of neural processes and the formation of a glial seal that encases the remaining retina (Marc et al., 2003). As most vision restorative therapies will be employed after photoreceptor death, a greater understanding of the events that occur in phase 3 degeneration will assist in optimizing these therapeutic approaches.

The rd1 mouse is the most widely studied mouse model of retinal degeneration (Farber, 1995; Fletcher et al., 2011). It carries a gene mutation in the β subunit of phosphodiesterase 6 (Pde6b), which is a similar genetic defect to patients with autosomal recessive RP (Pittler and Baehr, 1991; McLaughlin et al., 1993). Studies have documented both functional and anatomical changes in this mouse model of degeneration (Strettoi and Pignatelli, 2000; Strettoi et al., 2002; Jones et al., 2003; Margolis et al., 2008). Apart from the obvious structural changes in the outer retina (the loss of photoreceptors), there are also functional changes in bipolar and ganglion cells within this early period of degeneration (Stasheff, 2008; Chua et al., 2009; Stasheff et al., 2011; Gibson et al., 2013). However, at later stages of degeneration after photoreceptor loss, neurons of the inner retina also show significant anatomical alterations, including migration of somata to ectopic sites, the establishment of aberrant synapses, neuronal death, and glial remodeling (Jones et al., 2003; Marc et al., 2003; Chua et al., 2013). Finally, while neurochemical changes have been described both at early and late stages of degeneration in the rd1 mouse, as well as in other models of retinal degeneration (Fletcher and Kalloniatis, 1996, 1997; Marc et al., 2003; Jones and Marc, 2005; Gibson et al., 2013), the underlying cause and relationship between the different forms of inner retinal change is not clear. For that reason a model system probing these inner retinal changes would be of use.

The Fos-Tau-LacZ (FTL) mouse has an axonally targeted reporter system under the regulation of the c-fos promoter (Wilson et al., 2002). C-fos is an intermediate early gene that is known to be up-regulated in neurons in response to neural activity, cell death, and/or plasticity (Marti et al., 1994; Rich et al., 1997; Wilson et al., 2002; Greferath et al., 2004, 2009). FTL mice have been useful for visualizing neural circuits activated by a range of stimuli, including those involved in regulating water balance (Wilson et al., 2002), learning and memory (Ali et al., 2009, 2012) and respiratory control within the nuclei of the brainstem (Niblock et al., 2012). When FTL mice are bred with rd1 mice, photoreceptors display increased c-fos reporter labeling from an early age that follows closely the time course of photoreceptor death (Greferath et al., 2009). However, nothing is known about c-fos reporter (FTL) regulation at later stages of degeneration in the rd1-FTL mouse and whether this relates to inner retinal neural change.

In view of the increase in aberrant neural activity, plasticity changes and possible cell death that takes place within the inner retina at late stages of degeneration, the rd1-FTL mouse could provide a useful tool for studying the cellular changes that occur in the inner retina well after photoreceptor cell death. In this study we examined the time course of c-fos reporter expression in the rd1-FTL mouse and characterized the accompanying glial and neuronal alterations. Our results show that c-fos reporter (FTL) expression is upregulated in inner retinal neurons within discrete patches in the central retina from 4 months of age. Moreover, increased FTL expression coincides with regions displaying considerable Müller cell change and loss of cone photoreceptor terminals. These areas may represent regions of the retina from which the glial seal may form. Targeting restorative therapies distant to these retinal areas may improve outcomes.

**Materials and Methods**

**Animals**

All animal procedures were performed in accordance with the University of Melbourne Animal Experimentation Ethics Committee and with guidelines outlined by the National Health and Medical Research Council and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Rd1-FTL double-mutant mice were created by breeding rd1 homozygous mice with heterozygous FTL transgenic animals (Wilson et al., 2002). Rd1 mice were originally obtained from Professor Debora Farber, University of California, Los Angeles, CA, USA and backcrossed more than 10 times with the C57Bl6JARC obtained from the Animal Resource Centre (Western Australia). FTL mice were originally generated at the University of Melbourne (Parkville, Australia; Wilson et al., 2002). The rd1-FTL mouse expresses a mutation in the β subunit of Pde6b gene, resulting in retinal degeneration, together with an axonally targeted transgenic marker gene (β-galactosidase) that labels all neurons that have c-fos promoter activation (Greferath et al., 2009). Mice examined in this study were aged from 2 to 12 months and control heterozygous FTL mice were age matched for comparison (at least n > 7 rd1-FTL and FTL at each age group per outcome measure). All mice used in this study were confirmed to be free of the rd8 mutation in the Crb1 gene (Mattapallil et al., 2012).

**Tissue Collection and Fixation**

Mice were killed by an overdose of sodium pentobarbital (Merial Australia, Parramatta, Australia: 120 mg/kg, intraperitoneal). Eyes were then immediately dissected, the anterior segment and lens discarded and the posterior eyecup fixed in one of two ways depending on the type of immunofluorescence to be performed. Posterior eye cups that were processed for indirect immunofluorescence were fixed in 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer, pH 7.4 (phosphate buffered, PB) for 30 min, washed in PB and subsequently processed through...
graded sucrose solutions (10, 20, and 30% w/v sucrose in PB). Samples were stored at –80°C until use.

A subset of eyes from rd1-FTL and FTL mice (n > 8 eyes) were processed for β-galactosidase histochemistry, whereby the mouse was transcardially perfused with 12 ml of 5% sucrose in dH2O, followed by 12 ml of PFA with 0.005% glutaraldehyde in PB. Eyes were immediately dissected and a hole pierced in the eyecup at the corneal margin with a 30G needle. Eyes were then placed in 4% PFA and the anterior eyecup and lens were removed. The posterior eyecup was postfixed in 4% PFA for 10 min and then placed in phosphate buffered saline (PBS). All eyes were immediately processed for enzyme histochemistry.

β-Galactosidase Histochemistry on Wholemount Retinae

Retinae were dissected from the posterior eyecup and placed in the staining buffer (5 mM magnesium chloride, 5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, and 0.4 mg/ml 5-Bromo-4-chloro-3-indolyl-B-D-galactopyranoside) for 16 h under rotation at room temperature (Greferath et al., 2004, 2009). To stop the reaction, an equivalent volume of 4% PFA was added to the solution and the retina was washed in PBS and mounted onto 0.5% gelatine/5% chromium potassium sulfate-coated slides. Wholemounts were air-dried on slides and dehydrated through graded alcohols to histolene and mounted in PBS and 0.5% gelatine/5% chromium potassium sulfate-coated slides. Wholemounts were air-dried on slides and dehydrated through graded alcohols to histolene and mounted in Safety Mount (Frontine, Riverstone, Australia). Wholemount retinae processed for β-galactosidase histochemistry were viewed using a light microscope with either a 4× air objective or 40× oil objective and tissue was imaged using an ImagePoint cooled CCD camera (Photometrics LTD, Tucson, AZ, USA) and assessed using Windows imaging software (Digital Optics Ltd., Auckland, NZ). The extent of β-galactosidase expression (FTL labeling) was quantified as a percentage of total retinal area from n ≥ 8 retinae (mm² FTL labeling/mm² retinal area).

Immunofluorescence

Changes in cells in FTL and rd1-FTL mice were evaluated on vertical sections of retina using indirect immunofluorescence as previously described (Fletcher et al., 2000; Downie et al., 2010; Vessey and Fletcher, 2012). Briefly, sections (14 μm in thickness) were cut from OCT (Tissue Tek, Sakura Finetek, Tokyo, Japan) embedded eyecup blocks and mounted onto 0.5% gelatine/5% chromium potassium sulfate coated slides that were then air-dried. Slides were washed in PB and primary antibody was applied overnight. Following incubation, sections were again washed in PB and secondary antibody was applied for 40 min. Table 1 contains a list of the primary and secondary antibodies that were used. All antibodies were diluted in buffer consisting of 3% normal goat serum, 1% bovine serum albumin and 0.5% Triton X-100 in PB. Cell death was measured using a commercially available fluorometric terminal dUTP nick-end labeling (TUNEL) kit (DeadEnd Fluoro metric TUNEL system, Promega, Madison, WI, USA). Slides were then washed in PB, and mounted and cover-slipped with fluorescent mounting medium (DAKO, Carpinteria, CA, USA).

Retinal sections processed for immunofluorescence and TUNEL were viewed on a confocal laser scanning microscope (LSM 510 PASCAL, Zeiss, Oberkochen, Germany) using either a 20× air objective or a 40× oil objective and images were captured at a resolution of 512 × 512 or 1024 × 1024 pixels using Zeiss LSM image browser software and an appropriate fluorescence filter (Alexa TM 594/CY3: excitation 568 nm, emission filter 605/32; Alexa TM 488/FITC: excitation 488 nm, emission filter 522/32; Alexa TM 647/CY5: excitation 647 nm, emission filter 680/32). Images were collated using Adobe Photoshop CS5 (Adobe Systems, San Jose, CA, USA).

**Histological Analysis**

The immunolabeling of each antibody was assessed across at least n ≥ 6 eyes, with three separate sections imaged per eye. The co-localisation of FTL labeling with specific cell markers was quantified and expressed as a percentage of the total number of FTL positive cells [% (co-label of FTL and cell marker/total cells stained for FTL) × 100]. The markers of retinal Müller cells were quantified in control (FTL) and rd1-FTL retinae (areas exhibiting FTL labeling and neighboring areas lacking expression of FTL) as cell number per mm of retina (n ≥ 6). Total and inner retinal thickness measures were calculated from central retinal sections (n = 6 eyes, at least three sections per eye) that showed FTL labeling and neighboring regions that lacked FTL expression (assessed after indirect immunofluorescence, processed with ImageJ 1.43 freeware, NIH).

**Statistical Analysis**

A one-way analysis of variance (ANOVA) with a Tukey's multiple comparison post hoc test was used to statistically compare the change in c-fos reporter labeling across the rd1 retina. The quantification of glutamine synthetase (GS), Sox9, and cyclin D1 immunolabeling was analyzed using a two-way ANOVA, with Tukey's multiple comparison post hoc test to evaluate the differences. GraphPad Prism (San Diego, CA, USA) was used to graph data and to perform the above statistical analyses.

**Results**

This study examined the retinal changes that take place at late stages of degeneration (6–7 month-old animals) in the rd1-FTL mouse model of retinal degeneration. The rd1-FTL mouse has an axonally targeted reporter system under the control of the c-fos promoter (Wilson et al., 2002), as well as carrying a homozygous mutation in Pde6b. In the results described below, an increase in the c-fos reporter is referred to as “FTL labeling.”

**FTL Labeling is Increased in Inner Retinal Neurons Well after Photoreceptor Degeneration**

Using β-galactosidase histochemistry, the time course of FTL labeling was assessed in rd1-FTL mice up until 1 year of age and compared to age-matched control FTL mice. In the adult (postnatal day 135, P135) control retina (Figure 1A), FTL labeling covered the entire retina and as described previously, was found in a range of inner retinal neurons (Greferath et al., 2004). By contrast, FTL labeling was virtually absent in the rd1-FTL
TABLE 1 | List of the primary and secondary antisera used in this study.

| Antiserum                               | Dilution/Species       | Source/Cat. No.                                                                 |
|----------------------------------------|------------------------|-------------------------------------------------------------------------------|
| β-galactosidase                        | 1:10,000 Rabbit polyclonal | MP Biomedicals; Cat#08559761                                                   |
| β-galactosidase                        | 1:10,000 Chicken polyclonal | Abcam; Cat#ab9361                                                             |
| Caretin                                | 1:500 Mouse monoclonal  | Swant; clone 6B3                                                               |
| CyclinD1                                | 1:1 Rabbit polyclonal   | Merck Millipore; Cat#04-1151                                                  |
| GABA                                   | 1:500 Guinea pig polyclonal | Merck Millipore; Cat#A175                                                     |
| Glutamate                              | 1:45,000 Rabbit polyclonal | Gift from Dr Robert E Marc, available through Chemicon International; Cat #AB133 |
| Glial fibrillary acid protein          | 1:20,000 Rabbit polyclonal | DAKO; Cat# Z0334                                                              |
| GABA                                   | 1:50,000 Rabbit polyclonal | Gift from Prof David Pow, University of Queensland, Brisbane, QLD, Australia |
| GLAST (EAAT1)                           | 1:1,000 Rabbit polyclonal | Abcam; Cat# A841751                                                            |
| Vesicular glutamate transporter 1      | 1:20,000 Guinea pig, polyclonal | Merck Millipore; Cat#AB5905                                                   |
| Glutamine synthetase                   | 1:1,000 Mouse monoclonal | Merck Millipore; Cat# MAB302                                                   |
| Tyrosine hydroxylase                   | 1:1,000 Mouse monoclonal | Chemicon International; Cat #MAB318                                             |
| Kir4.1                                  | 1:1,000 Rabbit polyclonal | Merck Millipore; Cat#AB8818                                                   |
| Neuron specific nuclear protein        | 1:500 Mouse monoclonal  | Merck Millipore; Cat# MAB377                                                  |
| Peanut Agglutinin-Rhodamine            | 1:250 Fluorescent Lectin | Vector Labs; Cat #FL-1072                                                     |
| Sox9                                    | 1:500 Rabbit polyclonal  | Merck Millipore; Cat# AB5535                                                  |
| Anti-Rabbit Alexa594                   | 1:800 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A11037                                   |
| Anti-Mouse Alexa488                     | 1:500 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A11001                                   |
| Anti-Guinea pig Alexa594               | 1:500 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A11076                                   |
| Anti-Guinea pig Alexa488               | 1:500 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A11073                                   |
| Anti-Chicken Alexa594                   | 1:500 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A11042                                   |
| Anti-Mouse Alexa647                    | 1:500 Goat polyclonal   | Life Technologies; Alexa Fluor® Cat# A21235                                   |
| BisBenzimide H                          | 0.1 mg/mL               | Sigma Aldrich; Cat#14530                                                      |

**FIGURE 1** | Fos-Tau-LacZ (FTL) reporter expression increases in the rd1-FTL retina as a function of age. Flatmount retinae from control FTL (A, P135), rd1-FTL (B, P105; C, P135; D, P240; E, dark adapted rd1-FTL aged P210) mice were processed for FTL expression using β-galactosidase histochemistry. In contrast to the wild-type retina that showed FTL labeling throughout the retina (A), the rd1-FTL aged P105 (B) showed no FTL labeling. However, at P135 (C) rd1-FTL retina showed distinct labeling near the optic nerve, that subsequently expanded across the central retina (D,E). The expression of the FTL reporter as a function of retinal area was quantified at different ages from P105 to P365 (F). Data presented as mean ± SEM, n ≥ 8 per group, ****<0.0001 (one-way ANOVA). Scale bar for (A–E) = 1 mm.

retina at a similar age (P105, **Figure 1B**). By P135 (**Figure 1C**), there were discrete regions of FTL labeling lateral to the optic disk in the rd1-FTL retinae. Over time, this labeling spread to cover the central retina (**Figure 1D**), although the peripheral retina never showed labeling, even after 12 months (data not shown).
Previous studies in young rd1-FTL animals (≤P30) have shown that FTL labeling in the retina is influenced by light activation (Greferath et al., 2004). In order to determine whether this increase in FTL labeling observed at this late stage of degeneration (≥P135) was also dependent on light activation, 7 month-old rd1-FTL animals were dark-adapted prior to enucleation and β-galactosidase histochemistry. No difference in the labeling pattern between the light (Figure 1D) and dark-adapted (Figure 1E) rd1-FTL retinae was observed, indicating this increase in FTL labeling is independent of light-induced activation. Quantifying the extent of FTL labeling in the retina showed an age-dependent increase in expression after 105 days post-natal, with 29 ± 3% of the total retinal area exhibiting FTL labeling out to 1 year of age (Figure 1F, one-way ANOVA, p < 0.0001, n ≥ 8 per group).

The retinal cell types that were responsible for this increase in FTL expression were evaluated using immunofluorescence in 6 month-old rd1-FTL retinae. As shown in Figures 2A,C and E, cells in the inner retina were positive for FTL labeling (anti-β-Gal, red). In order to determine the types of cells expressing FTL immunoreactivity, double labeling experiments were performed with markers of amacrine cells including, GABA, glycine, and tyrosine hydroxylase. FTL was observed to co-localize with GABA (Figure 2B, arrows) and glycine (Figure 2D, arrows), but not tyrosine hydroxylase (Figure 2F), suggesting that in late stage degeneration, GABAergic and glycinergic amacrine cells express the c-fos reporter, while dopaminergic amacrine cells do not. This is unlike the inner retinal labeling observed in this model during the early stages of photoreceptor degeneration (<P26) where dopaminergic amacrine cells also express the c-fos reporter (Greferath et al., 2009).

β-galactosidase histochemical labeling of the rd1-FTL retina showed evidence of reporter expression within ganglion cell axons (Figure 2G, black arrows). In order to assess ganglion cell expression of the c-fos reporter, NeuN was used to label all neurons in the ganglion cell layer (GCL), while GABA immunolabeling was used to exclude displaced amacrine cells. Triple labeling with antibodies to β-Gal, NeuN, and GABA identified numerous ganglion cells that were labeled for FTL (NeuN positive, GABA negative, Figure 2H, see arrows). Quantification of FTL positive cells in the inner retina revealed that 28 ± 8% of FTL immunoreactive cells were strongly GABA immunoreactive, 41 ± 20% were strongly glycine immunoreactive, while within the GCL, 16 ± 5% were NeuN+/GABA− neurons (n ≥ 6 animals). These data suggest that specific subpopulations of conventional amacrine cells and ganglion cells express the c-fos reporter during the later stages of retinal degeneration.

**FTL expression during Late-Stage Degeneration Does Not Reflect Inner Retinal Thinning, Nor Exclusively Correlate with Remodeling**

c-fos is known to be upregulated during cell death (Marti et al., 1994). Indeed, our previous study of rd1-FTL mice showed FTL labeling in photoreceptor nuclei during the period of rod loss between post-natal day 10 and 30 (Greferath et al., 2009).

In order to determine whether the amacrine and ganglion cell labeling observed at late stages of retinal degeneration was associated with cell death, inner nuclear layer (INL) thickness was quantified in FTL immunolabeled regions and non-labeled areas that were immediately adjacent. The INLs of the FTL positive regions were not significantly different to the adjacent negative regions (Figure 3A, FTLa 30 ± 2, FTLa 31 ± 3 μm, p > 0.05, n ≥ 6 animals), while total retinal thickness was also unchanged (Figure 3A, FTLa 84 ± 8, FTLa 84 ± 8 μm, p > 0.05, n ≥ 6 animals). To confirm the lack of cell death involvement in FTL labelling, TUNEL labeling was performed within the inner retina of 4 and 6 month-old rd1-FTL mice. Limited staining was detected, similar to that observed in age-matched FTL controls.
Greferath et al. rd1-FTL inner retinal remodeling

FIGURE 3 | Fos-Tau-LacZ reporter expression in the rd1-FTL retina is not due to cell death nor correlated with remodeled inner retina. Total and inner nuclear layer thickness measures were measured from vertical retinal sections in areas exhibiting FTL expression, as well as neighboring areas showing no FTL expression (A). Additionally, vertical sections of rd1-FTL retina were labeled for FTL (anti-β-galactosidase, green) and DAPI (blue). Inner retinal layer and total retinal thickness were similar between FTL+ and FTL− regions (A). Increased FTL reporter expression was observed in inner retinal regions where the structure remained intact (B), as well as in regions where there was evidence of anatomical remodeling (arrows, C). Furthermore, other regions that exhibited DAPI labeled nuclei which were mislocated within the IPL, were FTL negative (arrows, D). Data presented as mean ± SEM, n ≥ 6 per group, p > 0.05 (two-way ANOVA). Abbreviations: INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer. Scale bar: 50 μm.

(data not shown). These findings suggest that at this late stage of retinal degeneration, the FTL labeling observed in the amacrine and ganglion cells of the rd1-FTL retina is unlikely to be the result of cell death.

As c-fos expression has been associated with neural plasticity, retinal areas exhibiting anatomical remodeling were correlated to FTL labeling. As shown in Figure 3B, FTL labeling occurred in areas with normal inner retinal morphology, as well as in regions with abnormally localized nuclei in the IPL, indicative of anatomical remodeling (arrows in Figure 3C). However, other regions that exhibited mislocalized DAPI-labeled nuclei within the IPL, were FTL negative (arrows in Figure 3D). Thus, FTL labeling in the inner retina did not exclusively coincide with areas that showed evidence of remodeling, suggesting that FTL up-regulation does not solely reflect neuronal plasticity changes in the rd1-FTL retina.

FTL Labeling of Inner Retinal Neurons is Localized to Regions of Aberrant Glia

The labeling within the rd1-FTL retina was patchy, with some regions showing FTL in cell somata of the INL (Figure 4A arrow, Figure 4D highlighted region), whereas neighboring areas showed no cell soma labeling. There was, however, a relatively consistent labeling within the GCL likely due to c-fos reporter expression within axons (Figure 4D arrow, Figure 2G). Müller cell labeling with GS (green; Bringmann et al., 2009) also showed a variability in staining (Figure 4B arrow, Figure 4E highlighted region) and revealed an intriguing inverse correlation, with FTL expression observed in retinal regions that were either devoid of, or showed a major reduction in, GS labeling (Figures 4C,F).

Müller cells are integral to maintenance of retinal structure and function, being involved in potassium siphoning, glutamate and GABA turnover, chromophore recycling, energy metabolism and maintenance of the blood retinal barrier (Bringmann et al., 2006). To examine whether these Müller cell metabolic processes might be compromised, rd1-FTL retinas were labeled for Kir4.1 the predominant glial inwardly rectifying potassium channel (Kofuji et al., 2000), GLAST, the principal glutamate transporter expressed by Müller cells (Lehre et al., 1997; Rauen et al., 1998), and glial fibrillary acidic protein (GFAP), a marker of Müller cell gliosis. As shown in Figure 5, regions of the rd1-FTL retina that showed reduced GS immunolabeling also showed reduced Kir4.1 staining (Figures 5A–C). However, while GLAST-labeling was still detected within these regions (Figures 5D–F), it appeared abnormal when compared to neighboring GS-expressing regions. These altered regions showed extensive GFAP labeling (Figures 5G–I), which has been reported previously in the rd1 mouse retina (Stasheff, 2008; Chua et al., 2009; Stasheff et al., 2011; Gibson et al., 2013). These data suggest that while GS and Kir4.1 expression is significantly reduced, gliotic Müller cells with the potential for glutamate uptake remained within these aberrant regions.

These results imply that after photoreceptor degeneration in the rd1-FTL retina, localized subpopulations of amacrine and ganglion cells show increased FTL expression in regions where Müller cells are dysfunctional or absent. In order to determine if Müller cell loss occurred in these FTL-expressing regions, retinal sections were co-labeled for GS and Sox9, a Müller cell nuclear marker. As shown in Figure 6, Sox 9 labeling was 100% co-localized with GS positive Müller cells in
FIGURE 4 | Fos-Tau-LacZ positive regions within the rd1-FTL retina show Müller cell alterations. Vertical sections of the rd1-FTL mouse retina (6 month-old) were labeled for FTL (A,D; anti-β-galactosidase, red) and glutamine synthetase (GS; B,E; green) and areas of co-localisation assessed (C,F). Regions that are intensely labeled for FTL correspond to regions that show low GS labeling (white arrow; B, C). High magnification of the FTL expressing region within the rd1-FTL retina is indicated by the white bracket and corresponds to an area of low GS labeling. Abbreviations: INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer. Scale bar: 250 μm (A–C), 50 μm (D–F).

FIGURE 5 | Müller cells are altered in the FTL positive regions within the rd1-FTL retina. Vertical sections of rd1-FTL retinae (7 month-old) were labeled for glutamine synthetase (A,D,G), Kir4.1 (B), GLAST (E), or GFAP (H), with the double label shown in (C,F,I). Areas that showed little GS labeling (in between arrow heads in C and F) corresponded to Müller cells that lacked Kir4.1. Despite this, these areas still showed GLAST immunolabelling. These altered regions were also positive for the gliosis marker GFAP. Abbreviations: INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer. Scale bar: 50 μm.

control retinae (Figure 6A), and in areas of the rd1-FTL retina that were GS positive (Figure 6B). By contrast, FTL positive areas that almost completely lacked GS, still showed some cells that were Sox9 immunoreactive (Figure 6B, highlighted region). Quantifying the number of GS and Sox9 immunoreactive Müller cells in control FTL and rd1-FTL (FTL⁺ or FTL⁻ regions) retinae (Figure 6E), showed a significant reduction in immunoreactive somata between FTL positive areas and control retinae (GS, –87 ± 2%; Sox9 –56 ± 7%; two-way ANOVA, p < 0.0001, n ≥ 6) as well as FTL positive and negative regions (GS, –86 ± 2%; Sox9 –53 ± 7%; two-way ANOVA, p < 0.0001, n ≥ 6). There were no significant differences between control
retinae and FTL negative regions of the rd1-FTL retinae (GS, –7 ± 6%; Sox9, –6 ± 5%; two-way ANOVA, p > 0.05, n ≥ 6).

As these FTL-positive areas exhibited several markers of proliferative gliosis (reduced GS, and Kir4.1 expression and increased GFAP), cyclinD1 was used to determine whether Müller cells in these affected regions had altered cell cycle regulation. Increased cyclinD1 expression in Müller cells has been previously reported during late stage retinal degeneration (Vessey et al., 2014). As shown in Figure 6C, no cells in the age-matched control retinae showed cyclinD1 expression. However, within FTL+/GS− regions of the rd1-FTL retina, numerous Müller cells showed cyclinD1 expression (Figure 6D, highlighted region). When quantified (Figure 6E), a significantly higher number of cyclinD1 immunoreactive Müller cells were found in regions of FTL labeling compared to control (FTL+ regions 53 ± 5 cells/mm retina; control FTL 0.6 ± 0.4 cells/mm retina; two-way ANOVA, p < 0.0001, n ≥ 6), or neighboring FTL negative regions (FTL+ regions 53 ± 5 cells/mm retina; FTL− regions 4 ± 2 cells/mm retina; two-way ANOVA, p < 0.0001, n ≥ 6). These results highlight that well after photoreceptor degeneration, inner retinal FTL expression coincides with distinct regions that show reduced Müller cell number (reduced Sox9), with the remaining cells exhibiting aberrant expression of functional markers, including Kir4.1, GS, and cyclin D1.

**Cone Loss Occurs in Regions of Glial Loss and Abnormality**

At late stages of retinal degeneration, phase 3 remodeling is characterized by marked anatomical changes that are thought to be precipitated by the loss of cone photoreceptor terminals. The association of cone-mediated remodeling on the patches of glial abnormality and FTL expression was assessed in the rd1-FTL (n ≥ 6 animals). Figure 7 shows representative vertical sections taken from 6 month-old animals that were double labeled with GS and vesicular glutamate transporter (VGLUT1) a marker for the glutamate transporter within glutamatergic terminals (Sherry et al., 2003). Regions that showed GS labeling (Figure 7A) also displayed VGLUT1 labeling (Figure 7B), whereas areas that had low levels of GS immunolabelling, corresponded to regions that showed a complete loss of photoreceptor terminals as indicated by an absence of VGLUT1 labeling (Figure 7C). This finding was confirmed with labeling for the cone marker, peanut agglutinin, with normal GS regions exhibiting remnants of cone terminals, whereas GS depleted regions showed no labeling of synaptic terminals (data not shown). Overall, the results of this study highlight that during late stage retinal degeneration there are localized changes in inner retinal neurons (amacrine and ganglion cells) that are coincident with aberrant Müller cells and that these regions correspond to regions where cone terminals are lost.

**Discussion**

c-fos is an intermediate early gene that is known to be regulated during cell death, altered neural function and neural plasticity. From approximately 4 months of age, a time where most photoreceptor nuclei have been lost, c-fos reporter (FTL)
expression was increased in subsets of amacrine and ganglion cells across the central rd1-FTL retina. These regions of FTL expression were not correlated with cell death or inner retinal remodeling. These areas did correspond to discrete regions of abnormal Müller cells which showed reduced GS, Kir4.1, and Sox9 immunolabelling, in addition to aberrant cell cycle regulation. Furthermore, these aberrant Müller cells were present in regions that showed a complete loss of cone photoreceptor terminals. These results highlight that inner retinal changes occur in the rd1-FTL retina in a localized fashion and are associated with the loss of cone photoreceptor terminals and Müller cell anomalies.

**FTL Expression as a Marker of Inner Retinal Change**

The results of this study show that c-fos reporter (FTL) expression changes in the rd1-FTL with age. Consistent with our previous findings (Greferath et al., 2009), retinal FTL expression was low at 3 months of age in the rd1-FTL compared to control FTL mice, most likely reflecting the lack of neural activity at this age. This is supported by previous work that indicates virtually all photoreceptor nuclei are lost in the central rd1 retina by 2 months of age (Jimenez et al., 1996). While FTL expression was initially low, it re-emerged within subpopulations of both amacrine and ganglion cells by 4 months of age (from P120). This increase in expression was not due to light-induced neuronal activation as we have previously reported in young (> P30) rd1-FTL animals (Greferath et al., 2004). Over the next 8 months (P120–P365) FTL expression increased within the central retina, yet never extended out into the peripheral retina. It must be highlighted that the oldest animals used in this study were 1 year of age and so peripheral FTL expression after this time cannot be discounted. The reason for this central localisation of c-fos reporter expression is unclear, however, a central to peripheral gradient in retinal degeneration has been reported in both the rd1 and rd1-FTL retinae (Carter-Dawson et al., 1978; Greferath et al., 2009).

The FTL transgene was used as a reporter for c-fos, an early intermediate gene that is known to be upregulated during cell death, neural function and neural plasticity. The increase in FTL expression in the amacrine and ganglion cells from 4 months of age was not associated with an increase in inner retinal cell death, as our results showed no difference in thickness of the INL, in either FTL-immunolabeled regions or neighboring FTL-negative regions. There was also no appreciable level of TUNEL labeling and certainly no specific localisation to FTL+ regions. Furthermore, the data did not support a role for c-fos reporter expression as a biomarker of inner retinal remodeling since anatomical remodeling was observed in FTL+ regions, as well as FTL− regions. With previous work on the rd1 mouse model showing evidence of altered inner retinal neurochemistry, as well as aberrant amacrine and ganglion cell function (Stasheff, 2008; Chua et al., 2009; Choi et al., 2014), it is possible that these functional alterations are related to the observed FTL expression.

Our data showed that anomalies in glial cells, specifically Müller cells, coincide with the localized regions of FTL up-regulation. Based on the Sox9 labeling, Müller cells were reduced in number within FTL+ regions of retina. In addition, the Müller cells that remained in these regions were gliotic and showed reduced glutamate synthetase and Kir4.1 immunolabelling. In view of the importance of both glutamate turnover and potassium siphoning in maintenance of neuronal function (Bringmann et al., 2006), we propose that the Müller cell dysfunction observed in localized regions could play a role in the up-regulation of FTL expression within the inner retinal amacrine and ganglion cells. In support of this, Müller cells are known to directly modulate ganglion and amacrine cell function (Newman and Zahs, 1998; Miller, 2004). In addition, these aberrant Müller cells within the FTL expressing regions may be sites of future glial scar/seal formation which is known to occur in phase 2/3 remodeling
et al., 2002) as well as sprouting thin processes from their somata (Strettoi and Pignatelli, 2000). Ganglion cells have shown changes in their oscillatory activity (Margolis et al., 2008; Stasheff, 2008; Stasheff et al., 2011) and ion permeability (Marc et al., 2007; Chua et al., 2009). A caveat of many of these studies is that while they use animals at a stage when the majority of photoreceptors have died, they are in the early stages of the remodeling process (generally < P100). Evidence from this study suggests that the FTL model may provide an early biomarker for glial scar formation and the loss of remnant cones at later stages in the retinal degeneration process, thus allowing these processes to be explored. In particular, it may be useful in identifying and targeting retinal regions that are more amenable to the implantation of vision restorative therapies.

In summary, the results of this study highlight that the rd1-FTL mouse may be a useful and unique tool for analysing retinal integrity during late stages of retinal degeneration. The strong up-regulation of c-fos reporter (FTL) that occurred well after photoreceptor loss and remained in the central retina throughout later stages of retinal degeneration, were independent of cell death, structural remodeling and any residual light response. Rather, it was associated with areas of dysfunctional glial cells and complete cone photoreceptor loss. Inhibiting these glial changes or preserving cone photoreceptors may assist with maintaining inner retinal integrity.

### Significance of Findings for Photoreceptor Restorative Therapies

Many studies of the rd1 mouse retina have shown that there are structural and functional changes in inner retinal cells following photoreceptor loss (Strettoi and Pignatelli, 2000; Strettoi et al., 2002; Marc et al., 2007; Margolis et al., 2008; Sekirnjak et al., 2011). Bipolar and horizontal cells change morphology and retract their dendrites (Strettoi and Pignatelli, 2000; Strettoi et al., 2002) as well as sprouting thin processes from their somata (Strettoi and Pignatelli, 2000). Ganglion cells have shown changes in their oscillatory activity (Margolis et al., 2008; Stasheff, 2008; Stasheff et al., 2011) and ion permeability (Marc et al., 2007; Chua et al., 2009). A caveat of many of these studies is that while they use animals at a stage when the majority of photoreceptors have died, they are in the early stages of the remodeling process (generally < P100). Evidence from this study suggests that the FTL model may provide an early biomarker for glial scar formation and the loss of remnant cones at later stages in the retinal degeneration process, thus allowing these processes to be explored. In particular, it may be useful in identifying and targeting retinal regions that are more amenable to the implantation of vision restorative therapies.

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