# Membrane-type frizzled-related protein regulates lipidome and transcription for photoreceptor function 

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#### Abstract

Molecular decision-makers of photoreceptor (PRC) membrane organization and gene regulation are critical to understanding sight and retinal degenerations that lead to blindness. Using Mfrp ${ }^{r d 6}$ mice, which develop PRC degeneration, we uncovered that membrane-type frizzled-related protein (MFRP) participates in docosahexaenoic acid (DHA, 22:6) enrichment in a manner similar to adiponectin receptor 1 (AdipoR1). Untargeted imaging mass spectrometry demonstrates cell-specific reduction of phospholipids containing 22:6 and very long-chain polyunsaturated fatty acids (VLCPUFAs) in Adipor1 $1^{-/-}$and $\mathrm{Mfrp}^{r d 6}$ retinas. Gene expression of pro-inflammatory signaling pathways is increased and gene-encoding proteins for PRC function decrease in both mutants. Thus, we propose that both proteins are necessary for retinal lipidome membrane organization, visual function, and to the understanding of the early pathology of retinal degenerative diseases.


## KEYWORDS

Adipor1, inflammatory signaling, matrix-assisted laser desorption/ionization imaging mass spectrometry (MALDI IMS), retinal degenerations, RPE cell, VLC-PUFAs

## 1 | INTRODUCTION

MFRP is a glycosylated transmembrane protein with an extracellular frizzled-related cysteine-rich domain ${ }^{1,2}$ that recognizes frizzled Wnt receptors, ${ }^{2,3}$ and participates in cell fate
and development. ${ }^{1,3} M F R P$ is expressed in the retinal pigment epithelium (RPE) and ciliary bodies, ${ }^{2}$ and gene mutations encoding this protein are associated with photoreceptor cell (PRC) degeneration, nanophthalmos, posterior microphthalmia, retinitis pigmentosa, foveoschisis, and optic disc

[^0]drusen. ${ }^{4}$ The $r d 6$ mice displays a mutated MFRP and also leads to retinal degeneration. ${ }^{2}$ Fundus analysis of these mice reveals discrete dots across the retina due to macrophage infiltration in the subretinal space ${ }^{5-7}$ as in flecked retinal diseases ${ }^{8}$ with similarities to human retinitis punctata albescens. ${ }^{7,9} \mathrm{Mfrp}{ }^{r d 6}$ has a 4 bp deletion in a splice donor sequence resulting in exon 4 being skipped, and in a truncated protein. ${ }^{2}$

Although MFRP mutations are linked to PRC degeneration, there is an incomplete understanding of MFRP protein function ${ }^{10,11}$ and its potential association with ADIPOR1 (Adiporl) including a shared pathogenic mechanism due to phenotypic similarities of the mutants of these two proteins. ${ }^{10}$ Moreover, comparisons of Mfrp ${ }^{r d 6}$ mice with wild-type (WT) littermates revealed the absence of ADIPOR1 in the RPE, ${ }^{10}$ but retention in the retina, which suggested that an ADIPOR1 deficiency in RPE may initiate retinal degeneration in $r d 6$ mice. ${ }^{10}$

ADIPOR1 is expressed in the retina and RPE, and the mutant of the Adiporl gene in these cells results in inability to take up and incorporate docosahexaenoic acid (DHA, 22:6,n-3), loss of PRC, and retinal degeneration. ${ }^{10,12}$ A single amino acid mutation of Adiporl occurs in different forms of retinitis pigmentosa. ${ }^{13,14}$ Polymorphisms of this receptor have been found in age-related macular degeneration (AMD) ${ }^{15}$ as well as in other forms of retinal degenerations. 22:6 is an omega-3 (n-3) essential fatty acid highly enriched in and required for the biogenesis of PRC membranes ${ }^{16}$ and is supplied either directly by diet or synthesized from dietary linolenic acid in the liver (18:3,n-3). ${ }^{17}$ 22:6 supplied by the liver ${ }^{18}$ is taken up from the choriocapillaris by the RPE and then delivered to the PRC inner segments. ${ }^{16,19}$ Here, it is incorporated at the sn-2 position of the glycerol backbone, especially in phosphatidylcholine (PC), which comprises more than $50 \%$ of PRC phospholipids ${ }^{20}$ mainly used for the biogenesis of PRC outer segments. ${ }^{21,22}$ 22:6-containing phospholipids modulate lipid rafts, ${ }^{23}$ become associated with RHODOPSIN, ${ }^{24}$ and permit efficient conformational changes of RHODOPSIN and its associated G proteins as photons are absorbed. ${ }^{25-27}$ The inability of the retina to take up and incorporate 22:6 leads to homeostatic compromise and PRC death as in retinal degenerative diseases. ${ }^{12,22,28}$ PRC outer segment tips containing 22:6-rich disks are phagocytized daily by the RPE, and the 22:6 is recycled back to PRC. ${ }^{16,19}$ PRC also contain very long-chain polyunsaturated fatty acids (VLC-PUFAs; $\geq 28$ carbons) formed by ELOVL4 (ELOngation of Very Long chain fatty acids-4) ${ }^{29,30}$ that elongates $26: 6, n-3$ derived from 22:6 or 22:5,n-3 (eicosapentaenoic acid, EPA). ${ }^{31,32}$ Despite the low abundance of retinal 22:5 compared to $22: 6,22: 5$ is the favored substrate for VLC-PUFA production. ${ }^{32}$ Generated by retro-conversion from 22:6 in peroxisomes, $22: 5$ provides the $22: 6$ substrate for ELOVL4. ${ }^{32}$ ELOVL4 synthesizes VLCPUFAs in retina ${ }^{21,33}$ and testes, ${ }^{34}$ and generates VLC-saturated fatty acids (VLC-SFAs) in skin and brain. ${ }^{35,36}$ VLC-PUFAs in PRC then become acyl chains of PC, incorporating at the sn-1 position. ${ }^{37}$ Under uncompensated oxidative stress (UOS), 32C
and 34C VLC-PUFAs are released and converted to elovanoids (ELVs) through enzymatic lipoxygenation as a PRC protective response. ${ }^{37}$ Because UOS also triggers 22:6 release and conversion to neuroprotectin D1 (NPD1), ${ }^{28} 22: 6$ availability is part of a dual protective signaling which sustains PRC integrity. ${ }^{28}$

To maintain fundamental PRC functions, membrane phospholipids provide an environment for phototransduction to proceed, and precursors of biologically active lipid mediators in the inner segment of PRC, as well as in the RPE. ${ }^{28,37}$ Therefore, a tight balance on retinal lipidome homeostasis must be maintained because UOS affects highly unsaturated fatty acids, forming $22: 6$ protein adducts that contribute to retinal degeneration. ${ }^{38}$ Overall, a healthy retinal lipidome is essential for PRC function within a highly stressful environment of bright light, high-oxygen demands, and an abundance of polyunsaturated fatty acids which are highly susceptible to oxidative damage. ${ }^{19,28}$

Here, utilizing the $M f r p^{r d 6}$ mouse, we have characterized key events that lead to PRC degeneration by mass spectrometry imaging, LC-MS/MS, physiology, and transcriptomic approaches. We have shown phenotypical, histological, and physiological similarities between Adipor $I^{-/-}$and Mfrpr ${ }^{r d 6}$ mouse retinas with comparable onset of PRC degeneration. Retinas and RPE in both mice presented a dramatic and selective deficit of 22:6,n-3-containing PC phospholipids, but not of arachidonic acid (20:4,n-6)-containing molecules, and an absence of VLC-PUFAs in the retina. We compared gene expression profiles of Adipor1 ${ }^{-/-}$and Mfrp ${ }^{r d 6}$ mice and found Casp1 and Pycard, part of the pyroptotic cell death pathway, upregulated in both mutants. Further analysis of target genes in enriched pathways indicated that the mutation of ADIPOR1 in the retina resulted in upregulation of genes belonging to JNK phosphorylation and activation mediated by the activated human TAK1 pathway, while the mutation of MFRP was linked to activation of the NOD-like receptor signaling pathway. Overall, our results indicate that the inability to enrich the retina with 22:6 results in the absence of VLC-PUFAs from the lipidomes of Adiporl ${ }^{-/-}$and $M f r p^{r d 6}$ mice, and, consequently, the inability to synthesize the neuroprotective elovanoids, which correlates with the activation of inflammatory signaling pathways and the onset of retinal degeneration.

## 2 | MATERIALS AND METHODS

## 2.1 | Animals

Throughout this study, C57BL/6J (wild-type; WT, The Jackson Laboratory, stock \# 000 664), B6.129P2-Adipor $1^{\text {tm1Dgen }} /$ Mmnc (Adiporl ${ }^{-/-}$; MMRRC, stock \# 011599-UNC), and B6.C3Ga-Mfrp ${ }^{r d 6} / \mathrm{J}$ ( $M f r p^{r d 6}$; The Jackson Laboratory, stock \# 003684) male and female mice were used. Mice were maintained in the LSU animal colony on a $12 \mathrm{~h}: 12 \mathrm{~h}$ light cycle
at $\sim 30$ lux (light onset was at 0600 h ), and given water and food ad libitum. Adiporl ${ }^{-/-}$and Mfrp ${ }^{r d 6}$ were screened for Crbl (rd8) mutation by PCR according to Mattapallil et al., 2012, ${ }^{39}$ and we confirmed that they do not contain an inherent Crbl mutation; hence, the retinal phenotypes observed in these mice derive from the mutations of Adiporl and Mfrp. Summary of the Mouse Universal Genotyping Array (MUGA) and MMRRC computational tools used to assess the genetic background of the Adiporl ${ }^{-/-}$mice can be found at: http:// www.csbio.unc.edu/MMRRC/index.py?run=StatsTable. viewSample\&sample=3826\&mm=0. All experiments were conducted in accordance with the Association for Research of Vision and Ophthalmology (ARVO) statement for the use of animals in ophthalmic and vision research, and the protocols were approved by the Institutional Animal Care and Use Committee (IACUC) for the LSU Health Sciences Center.

## 2.2 | Spectral domain-optical coherence tomography (SD-OCT)

Fundus images of 3-month old and SD-OCT images of WT, Adiporl ${ }^{-/-}$, and Mfrp ${ }^{r d 6}$ retinas of 1,3, and 5 months of age were obtained (Heidelberg Spectralis HRA OCT system; Heidelberg Engineering, Heidelberg, Germany) as previously described, ${ }^{12}$ and viewed with ImageJ (http//imagej.nih. gov/ij). The thickness of the outer retina (PRC + RPE) was defined as the distance from the proximal edge of the outer plexiform layer nuclear layer to the tips of the photoreceptor outer segments; the inner retina thickness was defined as the distance from the distal edge of the inner nuclear layer to the proximal edge of the ganglion cell/nerve fiber layer. ${ }^{40}$ Three animals/genotype/time point were analyzed.

## 2.3 | Electroretinograms

Conventional electroretinograms (ERGs) techniques were employed to determine general health and to follow physiological changes within the retinas of the experimental animals. Animals were dark adapted overnight (12-16 hours), anesthetized, and pupils dilated with topical $1.0 \%$ atropine (Mydriacyl 1\%, Alcon). Body temperature was maintained at $38^{\circ} \mathrm{C}$ throughout by the recording platform of the Espion ERG apparatus (Diagnosys, Lowell, MA). A drop of $1 \%$ methylcellulose placed on the cornea prevented corneal desiccation and provided improved electrical contact. Fiber optics contacted the corneas and delivered the light flashes generated by the Espion system. Scotopic ERG responses were elicited with short duration LED flashes delivered from the Espion ERG apparatus, with interstimulus intervals of 0.5-2 minutes, depending on intensity. Flash intensities ranged from 0.0001 to $10 \mathrm{~cd}-\mathrm{s} / \mathrm{m}^{2}$. Three to four responses were averaged for each
step, depending on the stimulus intensity (the flash interval was adjusted to assure dark adaptation following the previous flash). ERG responses were filtered (low-pass 0.125 Hz , highpass 300 Hz ) and archived for later analysis. Intensity-response amplitude data were displayed on log-linear coordinates.

## 2.4 | Matrix-assisted laser desorption/ ionization imaging mass spectrometry (MALDI IMS)

Whole eyes of 2-month-old mice were embedded in gelatin, frozen, cryosectioned ( $20 \mu \mathrm{~m}$ thickness), and then collected on alternating glass slides (for staining with hematoxylin and eosin) and coverslips (for MALDI IMS). Coverslips were attached to MALDI stainless steel plates using thermally conductive dual tape. Plates were then attached to a magnet (thermally conductive) glued to the bottom of a sublimation chamber. 2,5-dihydroxybenzoic acid (DHB) (Fisher Scientific, Pittsburg, PA) formed the matrix for the positive ion mode. A Synapt G2-Si (Waters, Milford, MA), equipped with a MALDI source that uses a solid-state laser ( 355 nm ) at a firing rate of 2000 Hz for positive ion mode data collection, was employed. HDImaing software (Waters, Milford, MA) was used to design the pattern of tissue scanning ( $30 \mu \mathrm{~m}$ spatial resolution for both horizontal and vertical movement) and data analysis. Each image spot consisted of a collection of 1 seconds data acquisition. Ions created by the MALDI source were further separated by ion-mobility-separation with helium gas in the TriWave region of the instrument, with an ion-mobility-separation wave velocity of $600 \mathrm{~m} / \mathrm{s}$ and height of 40.0 V . Data processed with HDImaging was converted with an in-house program, and BioMap software (Novartis) was used to generate images.

## 2.5 | Phospholipids molecular species as regional markers

Prior to analysis, specific markers for retinal layers/cell types were determined. To ensure good separation of retina and RPE-eyecup data, anterior segments were removed and retinas and RPE-eyecups were isolated separately on nitrocellulose filters, frozen, cryosectioned, and analyzed by MALDI IMS. PRCs were highly enriched with $\mathrm{m} / \mathrm{z} 756.6 \mathrm{PC}(16: 0 / 16: 0), \mathrm{m} / \mathrm{z} 760.6 \mathrm{PC}(16: 0 / 18: 1)$ labeled the inner retina, $m / z 1046.8 \mathrm{PC}(34: 6 / 22: 6)$ denoted an $n-3$ derived VLC-PUFA, and $m / \mathrm{z} 738$ (not identified) demarked the uvea/RPE complex. The shorthand notation used for lipid species here follows those outlined by Liebisch. ${ }^{41}$ The PC lipids observed (positive ion mode) were identified to lipid species level by collision-induced dissociation (CID) and are indicated by the number of acyl carbons (C) and the number of double bonds, for example $\mathrm{PC}(56: 12)$. Lipids were identified to the
fatty acyl level by CID and are indicated by the fatty acyl groups esterified to the sn-1 and sn-2 position, such as $\mathrm{PC}(34: 6 / 22: 6)$. To determine which phospholipids were most abundant between retinas, difference spectra were generated. Equal areas were selected from identical regions of retinal sections and total spectra obtained from 700 to $1100 \mathrm{~m} / \mathrm{z}$. One spectrum was then subtracted from a spectrum of another animal to produce a difference spectrum, which revealed molecular abundance within the two retinas. If the remaining peaks extended upward, there was more of that molecule within retina 1 , while peaks extending downward indicated more within retina 2 . Possible identification was then assigned to these PCs, and corresponding MALDI IMS retinal images obtained. WT and mutant retinas were compared in this manner. Since spectrum amplitudes differ among retinas, even if normalized, this method only revealed very large differences that were considered interesting; quantitation using this method is inaccurate.

## 2.6 | Lipid extraction and mass spectrometry for analysis by XevoTQ-S

Lipid extraction was performed similarly to our previous work. ${ }^{12}$ Briefly, each sample was homogenized in MeOH $(3 \mathrm{~mL})$ followed by addition of $\mathrm{CHCl}_{3}(6 \mathrm{~mL})$ and the internal standard (PC 28:0). After sonication in a water bath, samples were centrifuged, and the supernatant was added with $\mathrm{pH} 3.5 \mathrm{H}_{2} \mathrm{O}$ for phase separation. The bottom phase (organic phase) was dried down under $\mathrm{N}_{2}$ and reconstituted in an $\mathrm{AcN}: \mathrm{MeOH}: \mathrm{CHCl}_{3}$ (90:5:5) solution. A Xevo TQ-S equipped with Acquity UPLC BEH HILIC $1.7 \mu \mathrm{~m} 2.1 \times 100 \mathrm{~mm}$ column was used with solvent A (acetonitrile:water, $1: 1 ; 10 \mathrm{mM}$ ammonium acetate pH 8.3 ) and solvent B (acetonitrile:water, 95:5; 10 mM ammonium acetate pH 8.3 ) as the mobile phase. Solvent B (100\%) ran for the first 5 minutes isocratically was graduated to $20 \%$ solvent A for 8 minutes, and then ran at $65 \%$ of A for 0.5 minutes. It ran isocratically at $65 \%$ of A for 3 minutes, and then returned to $100 \%$ of B for 3.5 minutes for equilibration. The capillary voltage was 2.5 kV , desolvation temperature was set at $550^{\circ} \mathrm{C}$, the desolvation gas flow rate was $800 \mathrm{~L} / \mathrm{h}$, cone gas was $150 \mathrm{~L} / \mathrm{h}$, and nebulizer pressure was 7.0 Bars with the source temperature at $120^{\circ} \mathrm{C}$. Mutant and WT samples for males and females ( $\mathrm{n}=3$ each) were treated and analyzed separately. There was no difference observed between the two sexes; therefore, we combined the data for each type of mouse for $n=6$.

## 2.7 | LC-MS/MS data analysis

Raw mass spectrometry data were based on percent composition of relative molecular species per biological replicate.

Determination of differentially occurring molecular species was carried out using the loading score for the top ten PC molecular species for principle component 1, Principal component analysis (PCA), hierarchical clustering, and visualization of significant overlapping PCs (Venn diagram). Visualization of selected 22:6-containing and 20:4-containing PCs was obtained using BioVinci software (v1.1.3, r20180606), © 2017 BioTuring Inc, San Diego, CA, USA.

## 2.8 | RNA isolation and cDNA preparation

Retinas and RPE-eyecups from 1-month-old WT, Adiporl $^{-/-}$, and Mfrp ${ }^{r d 6}$ animals ( $\mathrm{n}=6$ each) were isolated. Mutant and WT samples for males and females ( $\mathrm{n}=3$ each) were treated and analyzed separately. There was no difference observed between the two genders; therefore, we combined the data for each type of mouse for $n=6$. Total RNA was extracted using RNeasy Plus Mini Kit (Qiagen, Germantown, MD) according to the manufacturer's instructions. One microgram of RNA was reverse transcribed using an iScript cDNA Synthesis Kit (BioRad, Hercules, CA). Diluted cDNA ( $100 \mathrm{ng}, 1.25 \mu \mathrm{~L}$ ) was used for the preamplification reaction by adding $1 \mu \mathrm{~L}$ of PreAmp Master Mix (Fluidigm, San Francisco, CA), $0.5 \mu \mathrm{~L}$ of a mix of all primers, and water to a final volume of $5 \mu \mathrm{~L}$. The temperature profile was $95^{\circ} \mathrm{C}$ for 2 minutes followed by 12 cycles of amplification $\left(95^{\circ} \mathrm{C}\right.$ for 15 s , and $60^{\circ} \mathrm{C}$ for 4 min using the Bio-Rad CFX96 thermocycler). Each preamplification reaction of cDNA was then subjected to Exonuclease I treatment, to remove unincorporated primers. The resulting preamplified and treated cDNA was then diluted 5 times in TE Buffer.

### 2.9 High-throughput qPCR

The qPCR reaction mixture had a volume of $5 \mu \mathrm{~L}$ and contained $2.25 \mu \mathrm{~L}$ of diluted preamplified cDNA, $0.25 \mu \mathrm{~L}$ of DNA Binding Dye (Fluidigm), and $2.5 \mu \mathrm{~L}$ SsoFast EvaGreen Supermix with low ROX (Bio-Rad). The primer reaction mixture had a final volume of $5 \mu \mathrm{~L}$ and contained $2.5 \mu \mathrm{~L}$ Assay Loading Reagent (Fluidigm) and $0.25 \mu \mathrm{~L}$ of a mix of all reverse and forward primers, corresponding to a final concentration of 500 nM in the reaction. The Biomark 96.96 IFC $^{\mathrm{TM}}$ (Integrated Fluidic Circuit) was first primed with an oil solution in the Juno ${ }^{\mathrm{TM}}$ Controller (Fluidigm) to fill the fluidic circuit. In all, 96 sample reactions ( $5 \mu \mathrm{~L}$ each) were loaded into individual sample wells, and 96 forward and reverse primer mixtures were loaded into each assay wells ( $5 \mu \mathrm{~L}$ each). The IFC was then placed in the Juno ${ }^{\mathrm{TM}}$ Controller for automatic loading and mixing. After 90 minutes, the IFC was then transferred to the Biomark ${ }^{\mathrm{TM}} \mathrm{HD}$ qPCR platform (Fluidigm). The cycling program consisted of Thermal Mix at $70^{\circ} \mathrm{C}$ for 40 minutes followed by $60^{\circ} \mathrm{C}$ for 30 seconds. Hot Start was 1 minutes at $95^{\circ} \mathrm{C}$,
followed by 30 cycles of denaturation at $96^{\circ} \mathrm{C}$ for 5 seconds, and annealing at $60^{\circ} \mathrm{C}$ for 20 seconds. Melting curves were collected between $60^{\circ} \mathrm{C}$ and $95^{\circ} \mathrm{C}$ with $1^{\circ} \mathrm{C}$ increments $/ 3$ seconds.

### 2.10 | qPCR data processing and analysis

The Ct value of target genes was normalized to the housekeeping genes Actb, Gapdh, Tbp, and Tfrc. For hierarchical clustering, the gene expressions were calculated as number of transcripts with the housekeeping gene set as 10000 transcripts. Mouse groups (columns) and genes (rows) were clustered using one minus Pearson correlation metric. For differential gene expression analysis, the fold-change value was calculated using the comparative threshold cycle method $(\Delta \Delta \mathrm{Ct})$ with the WT mouse sample as control. Calculated data were presented as box-plots with averaged Ct from six animals as a data point. Student's t test analysis was used to compare the differences between mutants and WT. All significant genes that were upregulated or downregulated to the WT were plotted as Venn diagrams. To interpret the distinct and shared transcriptional modulation between Adiporl ${ }^{-/-}$and Mfrp ${ }^{r d 6}$ mice, we used ConsensusPathDB ${ }^{42}$ (http://cpdb.molgen.mpg.de/MCPDB) and uploaded the lists of genes that are increased or decreased in the mutant datasets (Supplemental Table 1). We picked the top five pathways (only four pathways found with the common downregulated genes) and plotted the percentage of genes within the total pathway population. All graphs were made using the Bio Vinci program (Bioturing Inc, San Diego, CA).

### 2.11 | Statistics

The Kolmogorov-Smirnov test was employed to determine the assumption of normality for the SD-OCT measurements; an AR (1) as covariance structure was used for ERG measurements. A MANOVA test showed that differences existed among the retinal phosphatidylcholine (PC) data from the WT, Adiporl ${ }^{-/-}$, and $M f r r^{r d 6}$ animals; all $P$ values $<0.0001$ from Wilk's lambda, Pillai's Trace, Hotelling-Lawley Trace, and Roy's Greatest Root. Tukey-Kramer's multiple comparisons gave the final results (see Supplemental Table 2 for statistical details). $* P \leq .05$, ${ }^{* *} P \leq .01,{ }^{* * *} P \leq .001, * * * * P \leq .0001$ throughout.

## 3 | RESULTS

## 3.1 | PRC degeneration in Mfrp and Adipor1 mutant mice

Fundus imaging of 3-month-old $M f r p^{r d 6}$ mice demonstrated a uniformly flecked retina similar to that found in the Adiporl $1^{-/-}$retina ${ }^{12}$ and in human fundus albipunctatus. These
white spots of uniform size were evenly distributed within the fundus and exhibited similar density in the $M f r p^{r d 6}$ and Adiporl ${ }^{-/-}$retinas (Figure 1A). Optical coherence tomography (OCT) revealed a reduction in outer retinal thickness; by three months of age, about $40 \%$ of PRC nuclei had been lost, and at five months only about $40 \%$ remained (Figure 1B and C). Interestingly, the inner retina of the $M f r p^{r d 6}$ appeared thicker (Figure 1B and C). Overall, the $M f r p^{r d 6}$ phenotype closely resembles the Adiporl ${ }^{-/-} .{ }^{12}$

## 3.2 | Retinal physiology is similarly impaired in Mfrp and Adipor1 mutants

ERGs were recorded from 2-month-old WT, Mfrp ${ }^{r d 6}$, and Adipor1 ${ }^{-/-}$during the onset of PRC death when the PRC layer began to decrease (Figure 1B-D). ERGs obtained from light flashes ranging in intensity from 0.0001 to $10 \mathrm{~cd} \cdot \mathrm{~s} / \mathrm{m}^{2}$ revealed a $50 \%-75 \%$ decreased response in a-waves, for both the $M f r p^{r d 6}$ and Adiporl ${ }^{-/-}$mice vs WT, indicating a reduction of PRC response. The b-wave was reduced by about $50 \%$ throughout, displaying an inner retinal effect (largely bipolar cells). Interestingly, the dimmer light rod response in all mice plateaued through $0.005 \mathrm{~cd} \cdot \mathrm{~s} / \mathrm{m}^{2}$ to $0.075 \mathrm{~cd} \cdot \mathrm{~s} / \mathrm{m}^{2}$, followed by a similar rise in the brighter light cone response for all mice, indicating that reduced 22:6 affected both rods and cones (Figure 1E). Since there is no apparent loss of inner retinal cells in these mutations, this suggests that the reduced signal originated in functionally impaired PRC.

## 3.3 | Perturbations in the retinal lipidome containing highly unsaturated molecular species of phospholipids precedes PRC death in the Mfrp ${ }^{\text {rd6 }}$ retina

22:6 is important for visual function and retinal homeostasis. However, this essential fatty acid must be obtained by diet, collected and packaged by the liver, and released into the circulation for targeted delivery to the RPE and PRC. Failure to achieve adequate delivery results in impaired photoreceptor function and eventual cell death.

MALDI IMS revealed relative molecular abundance (spectra from 400 to $1200 \mathrm{~m} / \mathrm{z}$ ) from scanned histological sections (Figure 2A). MALDI analysis by positive ion mode of Adiporl ${ }^{-/-}, M f r p^{r d 6}$, and WT retinas demonstrated that the same phospholipids were present in the three genotypes, although the abundance of specific species within the mutant retinas was different from those of the WT. Analysis of MALDI sections revealed reliable identifying markers for the retinal layers (Figure 2C-G): m/z 738 Uvea and RPE, m/z 756 PC(32:0) photoreceptor layer, m/z $760 \mathrm{PC}(34: 1)$ inner retina. $\mathrm{m} / \mathrm{z} 900$ $\mathrm{PC}(22: 6 / 22: 6)$, an $n-3$ phospholipid, labeled the PRCs of the

WT, but was absent in both mutants; $\mathrm{m} / \mathrm{z} 832 \mathrm{PC}(20: 4 / 18: 0)$, an n-6 phospholipid, was present in all three animals, but was absent (greatly reduced) in the PRCs of the WT, indicating an $\mathrm{n}-3$
phospholipid preference in WT PRCs. m/z 1046 PC(34:6/22:6), an n-3 VLC-PUFA-containing molecule, was prevalent at the outer edge of the PRC layer in the WT, but was absent in both


FIGURE 1 Progressive PRC degeneration is similar in $M f r p^{r d 6}$ and Adiporl mutants. A, Fundus imaging revealed dense, evenly distributed white, punctate spots deep within 3-month-old mouse retinas, as observed in the Adipor $1^{-/-}$mouse. Images are representative of four mice. B, Optical coherence tomography (OCT) for 1-, 3-, and 5-month-old Mfrp ${ }^{\text {rd6 }}$ and Adipor $1^{-/-}$retinas demonstrated a gradual reduction in the photoreceptor layer thickness when compared to wild-type (WT) mice. C, The outer retina (OR, photoreceptors + RPE) thickness in both mutants at 1 month of age is reduced compared to the WT retina; the Mfrp ${ }^{\text {rd6 }}$ decreases slightly by 5 months, while the Adipor ${ }^{-/-}$OR continually decreases. The inner retinas (IR) remain unchanged for the WT and the Adipor1 ${ }^{-/-}$; however, the Mfrp ${ }^{r d 6}$ IR is thicker at 3 and 5 months. Error bars represent SEM of three individuals. D, Histological sections illustrating cellular morphology and changes in thickness of the Mfrp ${ }^{\text {rd6 }}$ and Adipor1 ${ }^{-/-}$photoreceptor layers, relative to WT mice, especially evident in the reduction of photoreceptors nuclei within the ONL. Each image is representative of three mice. E, a- and b-wave electroretinographic (ERG) responses of 2-month-old animals, to increasing intensities of light (cdes/ $\mathrm{m}^{2}$ ). Mfrp ${ }^{r d 6}$ and Adipor1 ${ }^{-/-}$retinal responses were reduced by approximately $75 \%$ and $50 \%$, respectively, from those of WT mice. Generally, both mutant mice displayed similar reductions. Error bars represent SEM of three individuals. GCL, ganglion cell layer; IPL, Inner Plexiform Layer; INL, Inner Nuclear Layer; OPL, Outer Plexiform Layer; ONL; Outer Nuclear Layer; ELM, External Limiting Membrane; IS/OS, Inner/Outer Segments; RPE, Retinal Pigment Epithelium; IR, inner retina; OR, outer retina; PR, photoreceptors. Magnification bars: (A) $200 \mu \mathrm{~m}$; (B) $100 \mu \mathrm{~m}$; (D) $100 \mu \mathrm{~m} . * P \leq .05,{ }^{* *} P \leq .01,{ }^{* * * P \leq .001, * * * * P \leq .0001}$
mutants, indicating that the lack of 22:6 impaired the synthesis of the VLC-PUFA phospholipids.

LC-MS/MS-based retinal lipidomic analysis displays the quantitative PC molecular species distributions in bar graphs for the retina (top) and the RPE (bottom) (Figure 2I). Clearly, 22:6-containing PCs are reduced in both Adiporl ${ }^{-/-}$and Mfrp ${ }^{r d 6}$ retinas and RPE, especially $\operatorname{PC}(44: 12)$ and $\mathrm{PC}(56: 12)$, which are abundant in PRC outer segments (Figure 2I, insets). The composition of the mass numbers used for retinal layer specification was determined by full fragmentation in negative ion mode (Figure 2 K and Supplemental Figure 1). Matching of the peaks in positive and negative ion modes confirmed the identity of the molecules selected for retina specification (Figure 2J). In our study, m/z 1104.6 (negative ion) and 1046.7 (positive ion) was identified as the $\mathrm{m} / \mathrm{z}$ of VLC-PUFAs (Figure 2 F , J and K) that was not detectable in Adiporl ${ }^{-/-}$and Mfrp ${ }^{\text {rd6 }}$. These spectra indicate that a dysfunctional MFRP results in a remarkable altered retinal lipidome, and, based on the importance of 22:6 in retinal maintenance and function, suggests that reduction of 22:6 contributes to eventual PRC degeneration in the $M f r p^{r d 6}$ mouse.

To further define differences in the retinal lipidome of the WT, Mfrp ${ }^{\text {rd6 }}$, and Adiporl ${ }^{-/-}$, retinal sections were obtained and imaged by MALDI IMS. Total spectra from m/z 700 to m/z 1200 were collected from small retina regions for comparison (Figure 3). A difference spectrum was then constructed by subtracting the Adiporl ${ }^{-/-}$or Mfrp ${ }^{\text {rd6 }}$ profile from the WT profile (Figure 3) A and B. The resulting plots, while not quantitative, emphasized lipid abundance, with the prevalent WT lipids (pointing up) and Mfrp ${ }^{r d 6}$, and Adipor1 ${ }^{-/-}$lipids indicated as downward peaks. 22:6-containing PCs (m/z 828.61 PC(16:0/22:6) and m/z 856.64 $\mathrm{PC}(18: 0 / 22: 6))$ were more prevalent in the WT retina, whereas the 20:4-containing PCs ( $\mathrm{m} / \mathrm{z} 804.6 \mathrm{PC}(16: 0 / 20: 4$ ) and $\mathrm{m} / \mathrm{z} 832.64 \mathrm{PC}(18: 0 / 20: 4))$ were enhanced in mutant retinas. Analysis of VLC-PUFAs spectra (from m/z 1000 to m/z 1200) revealed them to be almost non-existent in both mutant retinas. Supplemental Table 3 depicts the more abundant
phospholipid species revealed by the differential spectra of the WT, the $M f r p^{r d 6}$, and the Adipor $1^{-/-}$retinas, emphasizing the lack of VLC-PUFA species in the mutants. To further highlight the retinal lipidome differences between Adiporl ${ }^{-/-}$ and $M f r p^{r d 6}$, we generated differential spectrum of the two mutants (Figure 3C). The relative amount of 22:6-containig PCs, although severely depleted in $M f r r^{r d 6}$, was still detectable compared to Adiporl ${ }^{-/-}$. Supplemental Table 3 depicts the distribution of the more abundant phospholipid species revealed by the differential spectra of the $M f r p^{r d 6}$ and the Adipor1 $1^{-/-}$retinas. Overall, this comparison suggests that the 22:6-containing molecules are reduced in the $M f r p^{r d 6}$ and Adiporl ${ }^{-/-}$retinas, whereas the 20:4-containing species are either retained or enhanced, indicating that products of $n-3$ fatty acid synthesis are reduced in the Mfrp ${ }^{r d 6}$, while the n-6 fatty acid pathway may be conserved.

## 3.4 | VLC-PUFA-containing phosphatidylcholines are negligible in Mfrp ${ }^{\text {rd6 }}$ mouse eyes

MALDI IMS suggests that abundance of 22:6-containing PCs in the $M f r p^{r d 6}$ is drastically decreased (Figures 2 F and 3 B ). Inability to take up and incorporate 22:6 in retina and RPE was discovered in the Adiporl ${ }^{-/-}$mouse. ${ }^{12}$ Similarly, in the Mfrp ${ }^{\text {rd } 6}$, there is almost complete loss of PC-containing fatty acids generated through the $\mathrm{n}-3$ fatty acid synthesis pathway, especially the VLC-PUFAs (Figures 2F,4 and 5). Since the Mfrp ${ }^{r d 6}$ may have either lost the ability to take up 22:6 and/or synthesize PCs containing molecules elongated from 22:6 as in the Adiporl mutant, we investigated the overall retinal lipidome of 22:6containing phospholipids in $M f r p^{r d 6}$ retinas and RPE-eyecup.PC, phosphatidylethanolamine (PE), and phosphatidylserine (PS) molecular species in retinas and RPE-eyecups contain either $20: 4$ or $22: 6$, except $\operatorname{PE}(22: 1 / 22: 0)$ and $\operatorname{PS}(22: 1 / 22: 0)$ (Figures 4 and 5). Overall, retinal PCs show that 20:4-containing species are more abundant in the mutants, whereas 22:6-containing species are highly reduced (Figure 4). In PEs, 20:4-containing
(A)


Embed fresh, unfixed
(7)


Acquired MS spectrum for each scan address ( $\mathbf{x}, \mathrm{y}$ )
(B)

(C)


Uvea/RPE R 738
PRC G 756 a)

(H)

(2)

(I)
$\square$ WT
$\square$ Adipor1-/-
$\square$ Mfrp $^{\text {rd }}{ }^{1}$

Retina






FIGURE 2 MALDI IMS reveals loss of PC containing 22:6 and VLC-PUFAs in the outer nuclear layer of Mfrp ${ }^{\text {rd6 }}$ and Adipor $^{-/-}$retinas. A, Whole eyes embedded in gelatin (1), frozen, cryosectioned, $20 \mu \mathrm{~m}(2)$, collected on alternating glass slides (for H\&E) and coverslips (for MALDI) (3). Coverslips attached to MALDI plates (4) placed within sublimation chamber, ${ }^{43}$ matrix (2,5-dihydroxybenzoic acid, DHB) applied for positive ion mode analysis (5) MALDI Synapt G2-Si MS. Sections rastered by laser, $355 \mathrm{~nm}, 2000 \mathrm{~Hz}$ (6). Scanning control ( $30 \mu \mathrm{~m}$, horizontal and vertical movement) and analysis. Image spot consisted of a collection of one second of data acquisition (7). Image processing (8). B, H\&E sections of adjacent MALDI section for WT, Adipor $I^{-/-}$, and the $M f r p^{r d 6}$ mice. C, Lipid markers for uvea/RPE (m/z 738, red, R), PRC (m/z 756, green, G ), and inner retina ( $\mathrm{m} / \mathrm{z} 760$, blue, B). D, $\mathrm{n}-3$ containing $\mathrm{PC}\left(22: 6 / 22: 6\right.$ ), $\mathrm{m} / \mathrm{z} 900$ (green) is present in WT but absent in Adiporl ${ }^{-/-}$and $\mathrm{Mfrp}^{\text {rd6 }}$ retinas. E, n-6-containing phospholipids compensate D. Here, $\mathrm{PC}(18: 0 / 20: 4 \mathrm{n}-6, \mathrm{~m} / \mathrm{z} 832$, green $)$ is reduced in WT PRC, but still abundant within the inner retina while present within PRC of Adiporl $1^{-/-}$and $M f r r^{r d 6}$. F, 22:6 is elongated to produce VLC-PUFAs. WT retina contains 34:6 within $\mathrm{PC}(34: 6 / 22: 6) ; 34: 6$ is indicated by the white label, while the Adiporl ${ }^{-/-}$and the $M f r{ }^{r d 6}$ mice show no 34:6. G, The merge of C and F illustrates the location of 34:6 just inside the uvea/RPE marker (red) at the outer region of PRC (green) in WT, but no label within mutant retinas (absence of 22:6). H , Diagram on retinal location of C color markers. Double inset within each retina image is an enlargement of MALDI label and the corresponding area from $H \& E$ in $B$, indicated by boxes. $N=3$ for each condition. I, LC-MS/MS quantitative PC molecular species distributions in bar graphs for retina (top) and RPE (bottom). 22:6-containing PCs, which are abundant in PRC outer segments, are reduced in Adiporl ${ }^{-/-}$and $M f r p^{r d 6}$ retinas and RPE, especially PC 44:12 and PC 56:12 (yellow insets). Student's test ( $* P<.05$ ). (J) Retention time of negative ion mode of PC $56: 12(\mathrm{M}+\mathrm{CH} 3 \mathrm{COO})$ at $\mathrm{m} / \mathrm{z}=1104.6$ (top) matches with positive ion mode $(\mathrm{M}+\mathrm{H}+)$ at $\mathrm{m} / \mathrm{z}=1047.7$, confirming that $\mathrm{m} / \mathrm{z}=1046.7$ measured in MALDI is from PC 56:12 ( $\mathrm{M}+\mathrm{H}+$ ). ( K ) Full fragmentation spectrum of PC $56: 12(\mathrm{M}+\mathrm{CH} 3 \mathrm{COO}-)$ measured in negative mode shows composition of PC $56: 12$ (34:6/22:6)
species are higher in the mutants, but this phospholipid with a single 22:6 (eg, $\mathrm{PE}[16: 0 / 22: 6]$ and $\mathrm{PE}[18: 0 / 22: 6]$ ) is not different from WT. However, $\operatorname{PE}(22: 6 / 22: 6)$ and $\operatorname{PE}(24: 6 / 22: 6)$ are greatly reduced in the mutants (Figure 4). At the same time, PS shows a large increase in single-22:6-containing species (PS[16:0/22:6] and PS[18:0/22:6]), while PS(22:6/22:6) and $\mathrm{PS}(24: 6 / 22: 6)$ are greatly reduced as in the PEs (Figure 4). In the RPE-eyecup, single $22: 6$ containing species (16:0/22:6, 18:0/22:6) for both PC and PE are more abundant in the mutants (Figure 5), but the 22:6/22:6 and 24:6/22:6 species are greatly reduced as they are in retinas. The 22:6-containing PSs in the RPE-eyecups were not detectable.

Principal component analysis (PCA) of WT and mutant retinas and RPE-eyecups show good separation, particularly with retina (Figure 6A). Hierarchical clustering reveals two major clusters in retina (Figure 6B) and RPE-eyecup (Figure 6C), with red and blue indicating increased and decreased abundance, respectively. A Venn diagram (Figure 6D) depicts the overlap of significant differentially occurring PCs across strains and tissues. The numbers in each oval show the number of PC species statistically different vs WT.

## 3.5 | Phosphatidylcholines containing LCPUFAs with 4-7 double bonds are enhanced in Mfrp ${ }^{\text {rd6 }}$ and Adipor $1^{-/-}$mice

While there is a reduction of VLC-PUFAs in the mutants, even within PCs containing fewer than 10 double bonds, additional LC-MS/MS analyses of retinas and RPE-eyecups revealed increases in some 4-7 double bond, 50-58C PC molecular species (Supplemental Figure 3). These increases, which may be compensatory, were observed in retinas and RPE-eyecups of both Mfrp and Adiporl mutants. However, since PRCs were eventually lost, compensatory replacement
with alternate PC species was only a temporary fix, signifying a potential transient mechanism to sustain the retinal lipidome until adequate $22: 6$ could be achieved.

## 3.6 | VLC-PUFAs abundance and Elovl4 expression decline in Mfrp ${ }^{\text {rd6 }}$ retinas

Functional Mfrp gene yields a retinal lipidome rich in PC species (Figure 6) required for Elovl4 expression. We found reduced Elovl4 expression in the $M f r r^{r d 6}$ RPE-eyecups and retina (Figure 7C, Supplemental Figure 6A), which further explains the decreased abundance of VLC-PUFAs that we found.

## 3.7 | Genes signatures of pro-inflammatory signaling pathways are increased and genes encoding proteins for PRC function are decreased in AdipoR1 ${ }^{-/-}$and Mfrp ${ }^{\text {rd6 }}$

Mutation of Mfrp or Adiporl affects differentially signaling pathways leading to PRC death, as revealed by RT-qPCR analysis of the expression of 88 candidate genes in retina and RPE-eyecups. Target genes were selected for their function in the visual system and their role in inflammatory and immune system pathways. We found a higher hierarchical clustering of the selected genes for retina (Figure 7A) compared to the RPE-eyecup (Supplemental Figure 4), likely because the RPE-eyecup is less sensitive to the Mfrp or Adiporl mutation, as opposed to PRCs, which subsequently degenerates. Box-plots show the expression range of individual genes with the most significant fold-change in retina (Figure 7B) and RPE-eyecup (Figure 7C) across the three different genotypes (Mfrp ${ }^{r d 6}$, Adiporl ${ }^{-/-}$, and WT). Genes with higher foldchange differences in retina were linked to inflammatory


FIGURE 3 Differential MALDI spectra reveal compensatory PCs generated in Adipor1 ${ }^{-/-}$and the $M f r p^{r d 6}$ retina. Difference spectra show relative abundances detected by MALDI IMS. A, Molecules more abundant in WT retina vs Adipor1 ${ }^{-/-}$retina are presented in the upper part of the graph, while molecules more abundant in the Adipor1 ${ }^{-/-}$retina are displayed at the bottom. 22:6- and/or VLC-PUFA-containing PCs are much more abundant in WT while 20:4-containing PCs are increased in the mutant retina. The regions from which the spectra are extracted are shown in the insets H and E (top) and MALDI (bottom), showing PC40:6 ( $\mathrm{m} / \mathrm{z}=856.6$, red) and PC38:4 ( $\mathrm{m} / \mathrm{z}=832.6$, green). The identification of the mass numbers is shown in Supplemental Table 3. B, The difference spectra of $M f r p^{r d 6}$ to WT. (C) The difference spectra of Adiporl $1^{-/-}$to Mfrp ${ }^{r d 6}$. Inset images show that in both mutants, the PRC layers contain both PCs 40:6 and 38:4 (yellow) while WT has clear separation of the two PCs species between PRC layer and inner retina layer
pathways: Birc3, Pycard, Casp1, Illb, Fas, Naip, and Tnf. Interestingly, genes that are part of the pyroptosis cell death pathway, such as Caspl and Pycard, were upregulated more
than 4- and 3-fold, respectively, in mutants. Next, we followed the expression of genes associated with visual function, and linked to retinopathies when dysregulated. ROM1



#### Abstract

FIGURE 4 Selected phospholipid molecular species of WT, Adipor $I^{-/-}$, and $M f r p^{r d 6}$ mouse retinas. 22:6- or 20:4-containing PC, PE and PS (except for $\operatorname{PE}(22: 1 / 22: 0)$ and $\operatorname{PS}(22: 1 / 22: 0)$ ) are shown in the box-plots. 20:4-containing PC, PE, and PS are abundant in the mutant retinas. In both mutants, there is a great reduction in all the phospholipids containing two 22:6 molecules (PC, PE, or PS(22:6/22:6)) and PCs composed of 22:6 and VLC-PUFAs ( $\mathrm{PC}(22: 6 / 32: 6), \operatorname{PC}(22: 6 / 34: 6)$, and $\mathrm{PC}(22: 6 / 36: 6))$. On the other hand, single 22:6-containing PEs, such as PE(18:0/22:6), do not show differences among the groups; however, $\mathrm{PE}(22: 6 / 24: 6)$ shows greatly reduced amounts in both mutant retinas compared to WT. Interestingly, non-22:6 or $-20: 4$ containing $\mathrm{PE}(20: 0 / 22: 1)$ also shows great reduction in the mutants. For PS species, single 22:6-containing molecular species show higher abundances in the mutants while $\operatorname{PS}(22: 6 / 24: 6)$ and $\operatorname{PS}(22: 0 / 22: 1)$ show significant reduction in mutants


is part of the protein complex containing PRPH2 that contributes to PRC disk morphogenesis. We found a 3-fold downregulation of Roml in both mutants (Figure 7B), and although attenuated, the mutant retinas also displayed a downward regulation of Prph2 expression (Supplemental Figure 5A). Gfap, a marker of reactive gliosis in the Müller glia cells, was upregulated 5-fold in Adipor1 ${ }^{-/-}$and more than 3-fold in Mfrp ${ }^{r d 6}$ (Figure 7B). In the RPE-eyecup, there were only a few genes that displayed more than a 2-fold-change between mutants and WT. Among these genes, Elovl4 (1.5to 2 -fold) and Reep6 (3-fold) were decreased in mutants, while genes associated with inflammatory pathways, Fadd and Tnf, were increased 2-fold in Adiporl ${ }^{-/-}$RPE-eyecups only (Figure 7C). We investigated the classical signal transduction pathway via Adiponectin receptors, ADIPOR1 and ADIPOR2, which modulates different biological events such as steroidogenesis, glucose uptake, cell survival, fatty acid oxidation, vasodilatation, and cytoprotection. ${ }^{44}$ Although most of the candidate genes did not have their $\mathrm{FC} \leq 2$ or $>2$ (except for Adiporl in the Adipor1 ${ }^{-/-}$dataset), they displayed differences compared to the WT. Adiporl expression was decreased by 1.5 -fold in the retina and 1.4 -fold in the RPE-eyecup of $M f r p^{r d 6}$ (Supplemental Figure 5). Adipor2 was reduced in Adiporl ${ }^{-/-}$retina and RPE-eyecup, possibly as a compensatory way to respond to Adiporl deletion (Supplemental Figure 5). Interestingly, Mfrp expression was not significantly affected in either the retina or RPE-eyecup of Adipor $1^{-/-}$, suggesting a feedback loop for transcriptional regulation of Adiporl by MFRP (Supplemental Figure 7). Overall, a downregulation of genes was linked to visual function in the retina of both mutants (Supplemental Figure 6A), implying that the dysregulation of ADIPOR1 and MFRP has direct impact on the physiology of the PRC and their survival.

In Adiporl $l^{-/-}$and $M f r p^{r d 6}$ retina datasets, seven and six distinct genes, respectively, were upregulated (Figure 7D). To highlight the particularity of each mutant and to understand if they follow a specific degenerative pathway, we looked at the over-representation of candidate genes in enriched path-way-based sets (Figure 7E-H). The most highly represented biological activities enriched in the mutants compared to the WT, mainly involved response to inflammation and cellular processes for programmed cell death. The highest score of over-representation of candidate genes upregulated in

Adiporl ${ }^{-/-}$dataset belonged to JNK (c-Jun kinases) phosphorylation and activation mediated by the activated human TAK1 pathway (Figure 7E), while upregulated candidates in the $M f r p^{r d 6}$ dataset belonged mainly to the NOD-like receptor signaling pathway (Figure 7F). The common upregulated candidate genes (Supplemental Table 1) in Adiporl ${ }^{-/-}$and $M f r p^{r d 6}$ datasets are mainly represented within the AIM2 inflammasome pathway (Figure 7G).

Regarding the downregulated candidates, we found six unique upregulated genes in the Adiporl ${ }^{-/-}$dataset, while there were none in the $M f r p^{r d 6}$ dataset (Figure 7D). The list of these genes did not hit any enriched pathway-based sets, most likely because of the reduced list size. Common candidate genes downregulated in mutant datasets had the highest over-representation score for the neural retinal development pathway (Figure 7 H ); these candidate genes have a function in the development and function of PRC; thus, ADIPOR1 and MFRP are essential for the maintenance of PRC homeostasis by acting directly or indirectly on effectors for cellular integrity.

## 4 | DISCUSSION

A central question regarding onset and progression of retinal degenerative diseases is how acquisition of key molecules in the retinal lipidome takes place, and how they sustain transcriptomic programs for retinal function. To address this question, this study used sensitive and specific mass spec-trometry-based molecular imaging (MALDI IMS) for mapping the untargeted in situ spatial distribution of molecular species of phospholipids, complemented with LC-MS/MS quantification and detailed characterization, with OCT imaging and analysis of electroretinograms (ERG) to assess retina structure and function. Additionally, transcriptomic approaches have allowed us to define gene signatures involved in PRC degeneration in the $M f r p^{r d 6}$ mouse, and to establish analogies and differences with Adiporl ${ }^{-/-}$mouse, particularly preceding retinal degeneration. These findings have revealed novel necessary molecular events for PRC function.
$M f r p^{r d 6}$ displays characteristics similar to those expressed in human retinitis punctata albescens. ${ }^{7,9}$ Subretinal flecks also appear across the retina in this mouse, corresponding to accumulation of macrophages within the subretinal space, and this
RPE-eyecup $\quad \square$ WT $\square$ Adipor1 $^{-/-} \square \quad$ Mfrprd6


FIGURE 5 Box-plots of selected 22:6- or 20:4-containing PCs and PEs of WT, Adiporl ${ }^{-/-}$, and Mfrp ${ }^{r d 6}$ mouse RPE-eyecups. Most of the single 22:6-containing PC species in the RPE-eyecups show higher abundance in the mutants, unlike in the retinas. PC(22:6/22:6) or $\mathrm{PC}(22: 6$ combined with a VLC-PUFA) show great reduction in the mutants. 20:4-containing PE is reduced in the mutants, which is also an opposite observation from the retinas. Single 22:6-containing PEs are also increased in mutants, but PE (22:6/22:6) shows a great reduction in mutants, similar to the retina. Non $22: 6$ or $20: 4$ containing $\operatorname{PE}(20: 0 / 22: 1)$ also show great reduction in the mutants
phenotypic characteristic is also evident in the Adiporl ${ }^{-/-}$retinas. While a detailed timeline has not been established for these mutants, histology demonstrated the accumulation of
autofluorescence of undigested outer segments within the RPE, and OCT analysis revealed a progressive decline in the outer retina thickness as a result of PRC loss. ${ }^{12}$ Outer retinas (the


FIGURE 6 PCs containing VLC-PUFAs in Adipor1 $1^{-/-}$and $M f r r^{r d 6}$ eyes are greatly reduced. A, Principal component analysis of targeted PCs illustrates tissue-specific lipid profiles in mutants and WT mice $(\mathrm{n}=12$ ). B, C, Hierarchical clustering of significantly differentially occurring PC species (ANOVA one-way) in Adipor $~^{-/-}$and $M f r p^{r d 6}$ retinas and RPE-eyecups compared to WT ( $P \leq .05$, absolute value fold-change $\geq 1.5$ ) suggests compensatory PC dynamics. D, Venn diagram illustrates overlap of differentially occurring PCs across strains and tissues

PRC layer) of both mutants show a reduction from the normal to about $40 \%$ at one month with progressive loss up to 5 months, the Adipor ${ }^{-/-}$declining more rapidly. Moreover, the ERG declined by P25, indicating slow loss of function, ${ }^{12}$ with both mutant ERGs reduced to about $50 \%$ at 2 months. Differences in lipids among 3-month mutants and WT are described, but it is not known whether these differences are unchanged from birth in the mutants or are a result of an ongoing restructuring and compensation. An earlier study demonstrated that key retinal proteins were depressed and that rhodopsin was depleted prior to PRC death by P21 in the Mfrp ${ }^{r d 6}$ mice. ${ }^{10}$ Changes associated with the RPE cells in the $M f r p^{r d 6}$ mice have also been noted. ${ }^{9,45}$ Developing $M f r p^{r d 6}$ mice show reduced RPE microvilli, altered PRC inner segments, and disorganized outer segments, as well as impaired phagocytosis of PRC apical outer segments. ${ }^{45}$ Conversely, in RPE cell cultures, an increased number of RPE microvilli occur in the Mfrp mutants. ${ }^{9}$ Alterations in microvilli formation, density, and/or distribution would affect outer segment shedding, disrupt the process of membrane disposal by the RPE, and upset the retinal lipidome and PRC homeostasis.

MFRP has been localized in the RPE apical membrane and along the entire length of the RPE microvilli, as well as in PRC inner segments. MFRP is expressed as a dicistronic transcript with the C1q tumor necrosis factor-related pro-tein-5 (C1QTNF5/CTRP5) gene, ${ }^{2,46,47}$ which is associated with cell adhesion and basement membranes ${ }^{48}$ in the RPE and ciliary body, ${ }^{49}$ and the MFRP level is inversely proportional to CTRP5 expression. ${ }^{50} \beta$-actin is increased in MFRP-deficient cells and in cells overexpressing CTRP5, and MFRP affects actin polymerization, RPE cell adhesion, and microvilli morphology. ${ }^{50}$ CTRP5 is also localized in the RPE microvilli, ${ }^{49}$ and CTRP5 molecules form multimers in adjacent RPE cells with their collagen domains binding to collagen receptors in the membranes and their globular heads interacting to enhance RPE cell-cell adhesion. ${ }^{51}$ This has suggested a potential interaction with actin filaments ${ }^{52}$ at this level. An interaction here could be diminished since the $M f r p^{r d 6}$ mice have reduced/distorted RPE microvilli, ${ }^{9,45}$ which could reduce interdigitation with outer segments and impede 22:6 trafficking.

22:6 is reduced in both retina and RPE. The RPE regulates 22:6 uptake from the choriocapillaris, but their altered microvilli in the $M f r p^{r d 6}$ mice suggest that redistribution of the $22: 6$ from the RPE to photoreceptors may be impeded
or prevented. Eventually, with no or little $22: 6$ to form the neuroprotective compounds, the presence/accumulation of unmitigated stress would trigger an inflammatory response. Thus, arachidonic acid (20:4) is noticeably increased in both


(F)
(G)

(H)



FIGURE 7 The expression of inflammatory markers is increased and visual system markers decreased in Adiporl ${ }^{-/-}$and Mfrp ${ }^{r d 6}$. A, Heat map illustrating the hierarchical clustering of 88 target genes for the retinas of Adiporl ${ }^{-/-}(\mathrm{n}=6)$, Mfrp $^{r d 6}(\mathrm{n}=5)$ and WT mice $(\mathrm{n}=6)$. Color key represents row scaling of normalized fold-change (FC) expression values from RT-qPCR. B, Box-plots representation of selected genes for the retinas with their FC below -2 or above +2 , relative to the WT. Boxes denote median values with upper and lower quartiles, and whiskers, minimum and maximum outliers. C, Box-plots representation of selected genes for the RPE-eyecup samples with their FC below -2 or above +2 , relative to WT. D, Venn diagrams of differentially expressed genes within Adiporl $l^{-/-}$and $M f r p^{r d 6}$ datasets that are upregulated or downregulated in a statistically significant manner in the retina compared to the WT. The number in the intersection represents the differentially expressed genes that are common between the two datasets (Supplemental Table 1). Top pathways resulting from the upregulation of genes in Adipor ${ }^{-/-}$retina (E) or $M f r p^{r d 6}$ retina (F) compared to the WT, or common upregulated (G) or downregulated (H) genes in both mutant retinas. Student's $t$ test was used. $* P \leq .05, * * P \leq .01, * * * P \leq .001, * * * * P \leq .0001, \mathrm{NS}=$ not significant. Error bars represent standard deviation. When there was no statistical difference in the mutant datasets compared to the WT, statistic bars were not added
mutants, while it is notably less in the healthy WT mice, especially as compared with the abundance of 22:6 in the WT.

Healthy RPE cells are necessary for PRC integrity, recycling 22:6 from phagosomes back to the PRCs for synthesis of new disk membranes. ${ }^{16,45}$ Thus, RPE cells regulate 22:6 trafficking to the PRC, and impediments to this process limits the availability of 22:6 for elongation to VLC-PUFAs and for the synthesis of NPD1 and the ELV neuroprotectants, diminishing the ability to counter unmitigated oxidative stress and retinal homeostatic upset. ${ }^{28}$ The availability of this dual protective mechanism in which stress triggers 22:6 release and its conversion to NPD1, as well as the VLC-PUFAs and their conversion to ELV, ${ }^{28}$ is important to sustain PRCs. Therefore, we monitored 22:6 accumulation and elongation to VLC-PUFAs in retina by observing the distribution of $\mathrm{n}-3$ $\mathrm{PC}(44: 12)$, containing two 22:6 molecules, and n-6 $\mathrm{PC}(38: 4)$, containing (18:0/20:4). In those same retinas, MALDI imaging showed that $\mathrm{PC}(56: 12)$, containing the $\mathrm{n}-3$ VLCPUFAs 34:6 and 22:6, was localized within the outer PRC layer in WT retinas, but was absent in both mutant retinas. Conversely, the n-6 PC(38:4) was abundant within the PRC. Also, VLC-PUFAs were undetectable in the MALDI spectra of the Adipor $1^{-/-}$and $r d 6$ mice. Our data show that depressed PRC 22:6 n-3 in both mutants was accompanied by a compensatory response where less prevalent n-6 molecules were increased in the photoreceptor layer. Absence of 22:6 and the VLC-PUFAs, therefore, results in an increased $n-6 / n-3$ ratio in the Adipor $1^{-/-}$and $r d 6$ retinas, as similarly noted in AMD donor retinas. ${ }^{53}$ This creates a 20:4 n-6-enriched phospholipid environment which likely results in a proinflammatory condition, which contributes to the onset of PRC dysfunction. Since the n-3 VLC-PUFAs are released from their PC molecules upon unmitigated stress, and converted to protective elovanoids (ELVs), ${ }^{37}$ absence of this pathway favors PRC degeneration.

We suggest that the absence of 22:6, and, therefore, the lack of VLC-PUFAs, in the Mrp ${ }^{r d 6}$ and Adiporl ${ }^{-/-}$mice leads to transcriptomic modifications, which then trigger an increase in pro-inflammatory genes and a decrease of genes that are critical for visual system function. Transcriptome analysis provides insight into the relationship between $M f r p$ and Adiporl. Our
gene signature shows that the JNK phosphorylation and activation mediated by the activated TAK1 pathway are initiated in Adiporl ${ }^{-/-}$retina. The induction of this pathway could be due to the relative abundance of arachidonoyl groups in phospholipids in Adipor1 ${ }^{-/-}$retina that provides substrate for COX2 activation by c-Jun kinases to foster the formation of pro-inflammatory effectors that ultimately contribute to triggering cell death pathways. ${ }^{54}$ On the other hand, the favored pathway for $M f r p^{r d 6}$ is NOD-like receptor signaling. Free radicals and lipid peroxidation products in the retina activate this pathway. This is facilitated because the recycling by the RPE is altered in $M f r p^{r d 6}$, enabling accumulation of these harmful molecules that, in turn, could establish a stress-prone environment for PRC. As a consequence, the activation of the NOD-like receptor signaling pathway takes place. It is probable that although there is a distinct induction of inflammatory pathways in Adiporl ${ }^{-/-}$and $M f r p^{r d 6}$, there is eventually a convergence of pathways and, in this case, the predicted AIM2 inflammasome signaling, which would then activate apoptotic and pyroptotic signals following the activation of caspase-8 and $-1 .{ }^{55}$

Taken together, our study demonstrates that the Mfrp mutation resembles the Adiporl mutation on multiple levels, with some transcriptome differences: (i) phenotypically, the deletion of these proteins leads to flecked retina and slow PRC death onset; (ii) functionally, the mutations express the same degree of ERG attenuation; (iii) both mutants clearly demonstrate the inability to take up and incorporate 22:6 in the RPE and PRC, which results in remarkable changes in the retinal lipidome; and (iv) although the gene signature shows that both mutants are activated by signaling pathways involving proinflammatory cytokines, we found that each of these pathways have distinct features. Moreover, our results highlight the critical role of MFRP and ADIPOR1 in preserving PRC integrity through regulation of enrichment and distribution of 22:6 which, importantly, affects the synthesis of VLCPUFAs that provide the precursors for the neuroprotective ELVs. Hence, the maintenance of a balanced 22:6 retinal lipidome relies on MFRP and ADIPOR1, acting to ensure proper acquisition and distribution of key lipids from the RPE to the PRC, these events are necessary to sustain function in early stages of AMD and of other retinal degenerative deceases.

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## CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

## AUTHOR CONTRIBUTIONS

NGB conceived the study; BJ conducted the lipidomic analysis (LC-MS/MS and MALDI); histology was performed by WCG; MAK and KD performed the gene analysis; MAK conducted the OCT analysis; BM acquired the fundus images and performed ERG; NGB, BJ, MAK, and WCG designed experiments and analyzed the data; ZF performed the statistical analysis; WCG, MAK, and NGB wrote the paper with input from all other authors. All authors have read and approved the manuscript.

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## SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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[^0]:    Abbreviations: AdipoR1 Adiponectin Receptor 1; AMD age-related macular degeneration; C1QTNF5/CTRP5 C1q tumor necrosis factor-related protein-5; DHB dihydroxybenzoic acid; ELOVL4 ELOngation of Very Long-chain fatty acids-4; ELVs elovanoids; EPA eicosapentaenoic acid; ERG electroretinograms; JNK c-Jun kinases; MALDI IMS mass spectrometry-based molecular imaging; MALDI IMS matrix-assisted laser desorption/ionization imaging mass spectrometry; MFRP membrane-type frizzled-related protein; NPD1 neuroprotectin D1; OCT optical coherence tomography; PC phosphatidylcholine; PCA Principal Component Analysis; PE phosphatidylethanolamine; PRC photoreceptor; PS phosphatidylserine; RPE retinal pigment epithelium; SD-OCT spectral domain-optical coherence tomography; UOS uncompensated oxidative stress; VLC-PUFAs very long-chain polyunsaturated fatty acids; VLC-SFAs very long-chain saturated fatty acids; WT wild-type

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