Glutathione peroxidase 4 (GPx4) is a ubiquitously expressed antioxidant enzyme with a unique function that directly reduces peroxidized phospholipids in the cell membrane. GPx4 is known as a critical regulator of ferroptosis (1), lipid peroxidation–induced nonapoptotic cell death. However, the cytosolic isoform of GPx4 is considered to play a major role in inhibiting ferroptosis in somatic cells, whereas the roles of the mitochondrial isoform of GPx4 (mGPx4) in cell survival are not yet clear. In the present study, we found that mGPx4 KO mice exhibit a cone–rod dystrophy–like phenotype in which loss of cone photoreceptors precedes loss of rod photoreceptors. Specifically, in mGPx4 KO mice, cone photoreceptors disappeared prior to their maturation, whereas rod photoreceptors persisted through maturation but gradually degenerated afterward. Mechanistically, we demonstrated that vitamin E supplementation significantly ameliorated photoreceptor loss in these mice. Furthermore, LC–MS showed a significant increase in peroxidized phosphatidylethanolamine esterified with docosahexaenoic acid in the retina of mGPx4 KO mice. We also observed shrunken and uniformly condensed nuclei as well as caspase-3 activation in mGPx4 KO photoreceptors, suggesting that apoptosis was prevalent. Taken together, our findings indicate that mGPx4 is essential for the maturation of cone photoreceptors but not for the maturation of rod photoreceptors, although it is still critical for the survival of rod photoreceptors after maturation. In conclusion, we reveal novel functions of mGPx4 in supporting development and survival of photoreceptors in vivo.

Glutathione peroxidase 4 (GPx4) is known for its unique function in the direct detoxification of lipid peroxides in the cell membrane and as a key regulator of ferroptosis, a form of lipid peroxidation–induced nonapoptotic cell death. However, the cytosolic isoform of GPx4 is considered to play a major role in inhibiting ferroptosis in somatic cells, whereas the roles of the mitochondrial isoform of GPx4 (mGPx4) in cell survival are not yet clear. In the present study, we found that mGPx4 KO mice exhibit a cone–rod dystrophy–like phenotype in which loss of cone photoreceptors precedes loss of rod photoreceptors. Specifically, in mGPx4 KO mice, cone photoreceptors disappeared prior to their maturation, whereas rod photoreceptors persisted through maturation but gradually degenerated afterward. Mechanistically, we demonstrated that vitamin E supplementation significantly ameliorated photoreceptor loss in these mice. Furthermore, LC–MS showed a significant increase in peroxidized phosphatidylethanolamine esterified with docosahexaenoic acid in the retina of mGPx4 KO mice. We also observed shrunken and uniformly condensed nuclei as well as caspase-3 activation in mGPx4 KO photoreceptors, suggesting that apoptosis was prevalent. Taken together, our findings indicate that mGPx4 is essential for the maturation of cone photoreceptors but not for the maturation of rod photoreceptors, although it is still critical for the survival of rod photoreceptors after maturation. In conclusion, we reveal novel functions of mGPx4 in supporting development and survival of photoreceptors in vivo.

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Results

Generation of mGPx4 KO mice

We previously generated GPx4 heterozygous mice (GPx4+/−) by homologous recombination with a PGK-Neo cassette (2), and the embryonic lethality of GPx4 KO mice was rescued by loxP-GPx4 transgene that expresses all isoforms of GPx4 (GPx4+/−:GPx4tm1109;tm1109/PGK-Neo/cDNA3, Fig. S1A) (9, 12, 16). For this study, mGPx4 KO mice (GPx4−/−:GPx4tm1109;tm1109/PGK-Neo/cDNA3) were generated in which the lethality of GPx4 KO was rescued by mGPx4 KO transgene (Fig. S1, A and B) that expresses cGPx4 and nGPx4 but lacks mGPx4. Indeed, Western blot analysis on fractionated samples of the testis confirmed that GPx4 expression was abrogated in the mitochondria, whereas maintained in the cytosol and nucleus in mGPx4 KO mice (Fig. S1C). In contrast, littermate GPx4+/−:GPx4tm1109;tm1109/PGK-Neo/cDNA3 mice in which potential defects in mGPx4 KO mice are rescued by a wild allele of GPx4 were used as control mice.

mGPx4 deficiency leads to retinal degeneration

To confirm specific mGPx4 deficiency in the retina, cytosolic and mitochondrial fractions of the retina were prepared and analyzed on Western blot (Fig. 1A). Expectedly, in mGPx4 KO retina, GPx4 was not detected in the mitochondrial fraction, whereas the levels of GPx4 in the cytosol were similar to those in WT and control retina (Fig. 1, A and B). Immunohistochemistry of mGPx4 KO retina showed that GPx4 expression is severely downregulated in the inner segments of photoreceptors where mitochondria are condensed (Fig. 1C). On fundus photographs, the mGPx4 KO mouse at 3 months old presented with pigmentation, retinal vascular narrowing, and optic atrophy compared with the control mouse retina (Fig. 1D). Acronein, a product of lipid peroxidation, was upregulated in the mGPx4 KO mouse retina at P14 compared with the control mouse retina (Fig. 1E). Hematoxylin–eosin staining revealed a progressive degeneration after P7 in photoreceptors that locate at the outer nuclear layer (ONL; Fig. 1F). In detail, rod photoreceptors expressing rhodopsin survived at least until P30, whereas cone photoreceptors labeled with peanut agglutinin (PNA) were not observed as early as P7 (Fig. 1G). On RPE flatmounts, RPE cells did not show degeneration, confirming that the degeneration originates from the photoreceptors themselves and is not secondary to RPE degeneration (Fig. 1H).

Vitamin E partially rescues photoreceptor death caused by mGPx4 deficiency

Previous studies have shown that a vitamin E (vit E)-enriched diet rescues cell death caused by GPx4 ablation in vitro and in vivo (3–5, 17–20), through its effect of preventing lipid peroxidation (21). To examine the rescue effect of vit E in relation to mGPx4-deficient retinas, maternal mice were fed on a vit E-enriched diet (1000 IU/kg) from late pregnancy to the end of the experiments, through which the offspring were also supplemented with vit E. We found that the ONL of vit E-supplemented mGPx4 KO mice was significantly thicker than the ONL of mice fed on a normal diet (Fig. 2, A and B). The number of TUNEL-positive dead cells in the ONL significantly increased in mGPx4 KO mice fed on a normal diet, and this effect was significantly rescued by vit E supplementation (Fig. 2, A and C).

Next, we addressed the effects of mGPx4 KO and vit E supplementation in cone photoreceptors that were affected more severely than rods. The expression of s-opsin, a marker of mature cone photoreceptors, was not observed in mGPx4 KO mice, and the rescue effect by vit E supplementation was relatively small, although statistically significant (Fig. 2, D and E). Similarly, the expression of cone arrestin, another marker of mature cone photoreceptors, was not observed in the mGPx4 KO retina, and vit E supplementation did not show a significant rescue effect (Fig. 2D). In contrast, the expression of Rxxr-γ, a specific marker of cone photoreceptor precursor, was favorably expressed at P0 in the retina of mGPx4 KO mice, although it disappeared by P7. Vit E supplementation postponed the loss of cone precursors and significantly increased the number of Rxxr-γ-positive photoreceptors at P4 (Fig. 2, F and G). These results indicate that detoxifying peroxidized phospholipids in mitochondria are essential for the development of cone photoreceptors. However, rod photoreceptors can develop to their mature stage without GPx4, although mGPx4 KO causes rod photoreceptor death at a later stage.

To further clarify the differences in the effects of mGPx4 deficiency between cone and rod photoreceptors, electroretinogram (ERG) evaluations were performed (Fig. 3). The amplitude of both a-waves and b-waves of the scotopic condition was significantly lower in mGPx4 KO mice at 7 weeks old compared with the control mice (Fig. 3, A–C). In vit E-supplemented mGPx4 KO mice, the amplitude of both a-waves and b-waves was significantly higher than that in mGPx4 KO mice without vit E supplementation. In the photopic condition evaluated in 4-week-old mice, the amplitude of both a-waves and b-waves was significantly lower in mGPx4 KO mice than that in control mice. Vit E supplementation did not confer rescue effects on both photopic a-waves and b-waves (Fig. 3, D–F), which was consistent with our histology data (shown in Fig. 2, D–G). These results indicate that vit E ameliorates dysfunction in rod photoreceptors but not in cone photoreceptors. In addition, together with our histologic findings (Figs. 1 and 2), mGPx4 KO mice recapitulate some of the key features for human cone–rod dystrophy.

Mitochondrial abnormality in the photoreceptors of mGPx4 KO mice

To further investigate the roles of mGPx4, the retina at P17 was examined using transmission electron microscopy (TEM). Longitudinal sections of the mitochondria in the inner segments of mGPx4 KO mouse photoreceptors revealed that the number of mitochondria was significantly decreased and that the area per mitochondrion was significantly increased relative to equivalent observations in control mice (Fig. 4, A–C). The results suggest the occurrence of mitochondrial fusion because of mitochondrial oxidative stress caused by mGPx4 deficiency. Therefore, proteins related to mitochondrial fusion/fission
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Figure 1. Photoreceptor degeneration in mGPx4 KO mouse retina. A, Western blots of cytosolic (C) and mitochondrial (M) fractions of the retina from WT, control, and mGPx4 KO mice. TIM23 and SOD1 served as mitochondrial and cytosolic markers, respectively. GPx4 was detected in samples from WT and control retina but not in samples from mGPx4 KO retina. B, quantification of GPx4 levels in cytosol fraction, indicating retained GPx4 levels in mGPx4 KO mouse retina (n = 4 per group). Analysis was performed using one-way ANOVA and the Tukey–Kramer post hoc test. C, GPx4 expression in the retina of control and mGPx4 KO mice. Higher GPx4 levels were observed in inner segments in control mouse where mitochondria are condensed, which was not observed in mGPx4 KO mice (consistent in three independent samples per group). D, color fundus photographs of control and mGPx4 KO mice. Pigmentation, retinal vascular narrowing, and optic atrophy were observed in the fundus of mGPx4 KO mice (consistent in three independent samples per group). E, mGPx4 KO mouse retina contained more acrolein derived from lipid peroxidation than control retina on Western blot analysis (n = 6 per group). F, hematoxylin–eosin staining showed thinner ONL in mGPx4 KO mouse at P14 and P30 (consistent in three independent samples per group).
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were evaluated by Western blotting. Levels of the mitochondrial fusion protein mitofusion 2 (MFN2) were significantly higher in the mGPx4 KO mouse retina compared with the control mouse retina (Fig. 4D). Consistently, transverse sections of the mitochondria using TEM revealed frequent mitochondrial fusions in mGPx4 KO photoreceptors, whereas it was not observed in control mice (Fig. 4E). In contrast, there was no significant difference in the levels of other fusion/fission proteins, such as optic atrophy 1, mitofusion 1 (MFN1), phosphorylation of dynamin-related protein 1 (DRP1), at Ser616 and Ser637 (Fig. S2). In a recent study, it was reported that mitochondria in healthy photoreceptor inner segments align with those of neighboring inner segments across plasma membranes (22). Such mitochondrial alignment was confirmed in our control mice, whereas it disorganized in mGPx4 KO mice (Fig. 4E).

Apoptotic photoreceptor death in mGPx4 KO mice

As the photoreceptor death was caused by mGPx4 KO, the subcellular morphology of photoreceptors was investigated using TEM; uniformly dense and shrunken pyknotic nuclei were observed in the ONL of P17 mGPx4 KO mice, but not in that of the control (Fig. 5, A and B), indicating apoptotic cell death (23). The results were further supported by the detection of caspase-3 activation in the photoreceptors of mGPx4 KO mice that was suppressed by vit E (Fig. 5C).

Peroxisided phosphatidylethanolamine (22:6/22:6) in mGPx4 KO retinas

PUFA-associated phospholipids are vital for photoreceptors’ physiological functions and survival. To assess the effects of mGPx4 ablation on PUFA-associated phospholipids, LC–MS was used to compare phosphatidylethanolamine (PE) and phosphatidylcholine (PC), the two major phospholipids in mitochondria (24), among the retinas from the control, mGPx4 KO, and vit E-supplemented mGPx4 KO mice at P10, that is, the early phase of photoreceptor degeneration (Fig. 6). The concentrations of unchanged forms of PE-PUFAs tended to be lower in mGPx4 KO retinas than in the control and vit E-supplemented mGPx4 KO retinas. Specifically, the relative concentration of PE (22:6/22:6), which contains two chains of DHA, was significantly lower in mGPx4 KO retinas than in control and vit E-supplemented mGPx4 KO retinas (Fig. 6A). Accordingly, peroxidized PE (22:6/22:6-OOH) concentration was significantly increased in mGPx4 KO mouse retinas compared with the PE (22:6/22:6-OOH) concentration in control and vit E-supplemented mGPx4 KO mouse retinas (Fig. 6B). PE (18:0/22:6-OOH) concentration also tended to be higher in mGPx4 KO retinas, but this effect was not statistically significant. There was no significant difference in hydroxy PE-PUFAs, the reduced form of peroxidized PE-PUFAs, among the three groups (Fig. 6C). For PC-PUFAs, there was a tendency for unchanged forms of PC-PUFAs to decrease (Fig. 6D) and hydroxy PC-PUFAs to increase (Fig. 6F), although not at statistically significant levels.

Spermatogenesis abnormalities in mGPx4 KO mice

As already shown in a previous study (10), our mGPx4 KO mice also presented with male infertility and abnormal spermatogenesis. Mitochondria in the midpiece of the spermatozoa were swollen in mGPx4 KO mice (Fig. S3A). Flagella of mGPx4 KO spermatozoa were bent into hairpin-like shape (Fig. S3B), and mitochondrial membrane potential of mGPx4 KO spermatozoa was significantly lower (Fig. S3C). Vit E treatment rescued the mitochondrial membrane potential (Fig. S3C) but was not effective to mitigate the bent flagella (Fig. S3B) or male infertility, suggesting a redox-independent role of mGPx4 in the spermatogenesis. No apparent abnormality was observed in the seminiferous tubules of mGPx4 KO mice (Fig. S3D), and there was no difference in the number of spermatozoa between mGPx4 KO and WT mice (Fig. S3E). In contrast, our previous study revealed that spermatocyte-specific KO of all isoforms of GPx4 in mice displayed a dramatic loss in the number of spermatocyte and spermatozoa in the testis (5), suggesting that mGPx4 is critically important for the fertilization of spermatozoa, while may be dispensable for the synthesis and/or survival of spermatozoa.

Discussion

GPx4 is crucial for the development and survival of numerous types of cells. However, the isoform-specific roles of GPx4 are still obscure. This study and studies by other researchers have shown that mGPx4 KO mice grow normally but exhibit male infertility because of defective sperm motility and abnormal tail structure (10, 11). To date, in the somatic tissues, cGPx4 has been believed to be more dominantly expressed than mGPx4, and mGPx4 dominance in the testis has been considered to be exceptional (10). Our data show that mGPx4 is predominant in photoreceptors.

We have previously reported that the loss of all isoforms of GPx4 in photoreceptors led to rapid cell death (16); however, the isoform-specific roles of GPx4 in photoreceptors were not addressed. The present study reveals the pivotal role of mGPx4 in photoreceptor survival in vivo. In contrast to other somatic cells in which cGPx4 is more dominant than mGPx4 (10), our data show that mGPx4 is predominant in photoreceptors. However, cGPx4 still might also contribute to photoreceptor survival because photoreceptor degeneration observed in mGPx4 KO mouse was milder than that observed in photoreceptors lacking all isoforms of GPx4 (16).
Figure 2. Vitamin E supplementation ameliorated retinal degeneration in mGPx4 KO mice. 

A, TUNEL staining of the retinas in control, mGPx4 KO, and vit E-supplemented mGPx4 KO mice at P14. B, ONL thickness at P14 in control, mGPx4 KO, and vit E-supplemented mGPx4 KO mice. The ONL of mGPx4 KO mouse retina was significantly thinner than control mouse retina, which was partially rescued by vit E supplementation (n = 4–5 in each group). 

C, a significantly larger number of TUNEL-positive photoreceptors were detected in mGPx4 KO mouse retina than in control at P14, which was partially rescued by vit E supplementation (n = 6 per group). D, immunohistochemistry for cone-specific marker s-opsin and cone arrestin. S-opsin expression was not detectable in mGPx4 KO mice at P7 and P14. Vit E supplementation mildly rescued the s-opsin expression at P7 but not at P14. Cone-arrestin expression was not observed in mGPx4 KO or vit E-supplemented mGPx4 KO mice at P14. E, quantification for the number of s-opsin-expressing outer segments of cone photoreceptors in control, mGPx4 KO, and vit E-supplemented mGPx4 KO mice (n = 8 per group). F, immunohistochemistry for cone precursor marker Mitochondrial GPx4 is indispensable for photoreceptors.

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The central nervous system, including brain and retina, consumes 20% of total oxygen and a quarter of total glucose used for energy supply; indeed, the retina ranks among the highest of the energy-consuming tissues in the central nervous system (15). The retina consists of different types of neural and glial cells. The expression of energy-generating cytochrome c oxygenase as well as energy-consuming Na+/K+-ATPase is the highest in the inner segments of the photoreceptors (15). Moreover, compared with rod photoreceptors, cone photoreceptors contain a significantly higher mitochondrial volume of mitochondria; hence, they produce more metabolic energy (15). In the mGPx4 KO mouse retina, damage was most severe in the cone photoreceptors and less severe in the rod photoreceptors. Therefore, the severity of degenerations among the different cell types of mGPx4 KO mouse retina was correlated with the levels of physiological energy demands. In contrast, vit E supplementation was able to rescue rod photoreceptors morphologically and functionally, whereas cone photoreceptors could not survive after P7 even with vit E supplementation. These results indicate that cone photoreceptors may be

\[ \text{Rxr-}\gamma \] expression levels were similar between control and mGPx4 KO mouse retina at P0 but diminished at P4 and P7 in mGPx4 KO mice. Rxr-\gamma expression was restored in vit E-supplemented mGPx4 KO mice at P4 but not at P7. G, quantification for the number of Rxr-\gamma-expressing cells at P0 and P4 (n = 12 per group). All scale bars represent 25 μm. *p < 0.05, **p < 0.01 (KO with vitamin E versus KO), ###p < 0.05, ##p < 0.01 (KO versus control) using one-way ANOVA and the Tukey-Kramer post hoc test. mGPx4, mitochondrial isoform of GPx4; ONL, outer nuclear layer; vit E, vitamin E.

![Figure 3](image-url)

**Figure 3. Functional abnormalities in mGPx4 KO mouse retina on electroretinogram.** A, representative scotopic ERG in 7-week-old mice. B and C, in scotopic condition, the amplitude of both a-wave and b-wave was significantly diminished in mGPx4 KO mice, which was partially rescued by vit E supplementation (n = 3, 5, and 6 for control, mGPx4 KO, and vit E-supplemented mGPx4 KO). D, representative photopic ERG in 4-week-old mice. E and F, in photopic condition, the amplitude of both a-wave and b-wave was totally lost in mGPx4 KO mice, which could not be rescued by vit E supplementation (n = 6 for control and mGPx4 KO, n = 8 for vit E-supplemented mGPx4 KO). *p < 0.05, **p < 0.01 (KO with vitamin E versus KO), #p < 0.05, ##p < 0.01 (KO versus control) using one-way ANOVA and the Tukey-Kramer post hoc test. ERG, electroretinogram; mGPx4, mitochondrial isoform of GPx4; vit E, vitamin E.
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more vulnerable to lipid peroxidation in mitochondria than rods, which is consistent with previous studies highlighting the pathogenic role of oxidative stress, particularly in the loss of cone photoreceptors in retinal degeneration (25, 26). However, it remains unclear whether the photoreceptor degenerations were caused directly by mitochondrial lipid peroxidation in photoreceptors or indirectly by changes in other cell types because mGPx4 was deficient in whole body of the mGPx4 KO mice. The considerable difference in the rescue effects by vit E for cone and rod photoreceptors might be explained by the limited transfer of vit E through placenta (27) and the fact that the differentiation into cone photoreceptors takes place before
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birth while the differentiation into rod photoreceptors occurs just after birth. Therefore, the levels of exposure to vit E may differ between cone and rod photoreceptors in their early developmental phases (28). In contrast, previous studies have indicated that GPx4 might function as a structural protein, independent of detoxifying peroxidized lipids, in the mitochondria of spermatocytes (5, 29), which might also be the reason why vit E supplementation resulted in the partial rescue of photoreceptors in mGPx4 KO mice.

An omega-3 PUFA DHA is one of the major long-chain PUFAs in the retina and rod outer segment membranes (13, 14); it is critically important for the maintenance of photoreceptors and their survival (30). Furthermore, photoreceptors are known to contain unique phospholipids harboring two omega-3 PUFAs esterified at both sn-1 and sn-2 positions of the glycerol backbone (31), which was also confirmed in the present study. Previous studies have elucidated the key roles played by PUFAs esterified to PE (i.e., PE-PUFAs) in lipid peroxidation and ferroptosis (32). Consistently, in mGPx4 KO mouse retinas, we observed a significant increase in the level of PE (22:6/22:6-OOH), but no significant change in PC-PUFAs. In addition, vit E supplementation suppressed the increase in peroxidized PE-PUFAs and ameliorated the degenerative phenotypes both anatomically and functionally. However, our observations by TEM revealed that the size of each mitochondria in mGPx4 KO mouse photoreceptors was larger, which is not typical for ferroptosis in which mitochondria are smaller (33–35). In addition, we also observed uniformly dense and shrunk pyknotic nuclei, as well as caspase-3 activation in mGPx4 KO mouse photoreceptors, suggesting the existence of apoptotic cells at least in part. In typical ferroptosis, the morphology of nuclei remains intact (36). Our observations are consistent with previous in vitro studies showing that mGPx4 overexpression suppressed cytochrome c release and caspase activation and rescued cell death induced by apoptosis inducers, including staurosporine, 2-deoxyglucose, and Fas-specific antibodies (37, 38).

Figure 5. Apoptosis of photoreceptors in mGPx4 KO mice. A, in the ONL of P17 mGPx4 KO mouse retina, TEM disclosed the presence of uniformly dense and shrunk pyknotic nuclei (red arrows), which was not observed in control mouse retina. Areas inside the enclosing frames in the left (the scale bars represent 10 μm) are shown magnified in the right (the scale bars represent 2.5 μm). B, quantification of the apoptotic nuclei observed in TEM images in control and mGPx4 KO mice (n = 3 for control, n = 5 for mGPx4 KO; three views per sample). *p < 0.05 (KO versus control) using Student’s t test. C, immunohistochemistry for cleaved-caspase 3 confirmed caspase activation in photoreceptors of mGPx4 KO mice. Vit E treatment decreased the number of cleaved-caspase 3-positive cells in mGPx4 KO mice (the scale bar represent 25 μm). **p < 0.01 (KO with vit E versus KO), #p < 0.05, ##p < 0.01 (KO versus control) using one-way ANOVA and the Tukey–Kramer post hoc test. n = 3 per group and three views per sample. mGPx4, mitochondrial isoform of GPx4; ONL, outer nuclear layer; TEM, transmission electron microscopy; Vit E, vitamin E.
Based on observations by TEM, the mitochondria in mGPx4 KO photoreceptors were larger in size and fewer in number, which suggests that mitochondria were inclined toward fusion rather than fission. When evaluating the expressions of fusion/fission proteins, we observed an upregulation of MFN2 in the mGPx4 KO retina. MFN2 locates at the outer mitochondrial membrane and endoplasmic reticulum (ER), and it plays key roles in mitochondrial fusion, modulating ER stress, and interactions between the ER and mitochondria. A recent study suggested that MSN1/2 mediates mitochondrial fusion and lipid peroxidation and found that MFN1/2 depletion reduced the sensitivity of pancreatic cancer cells to erastin-induced apoptosis.

**Figure 6.** LC–MS analysis indicated increased phospholipid peroxidation of PE 22:6/22:6 in mGPx4 KO mice retina. A, PE 22:6/22:6 (DHA esterified at both sn-1 and sn-2 positions of PE) was significantly decreased in mGPx4 KO mice retina at P10 compared with control, which was blocked by vit E supplementation (n = 3 per group). B, peroxidized PE 22:6/22:6-OOH was significantly increased in mGPx4 KO mice at P10 compared with control, which was rescued by vit E supplementation (n = 3 per group). C, there was no significant difference among the three groups in hydroxy PE-PUFAs-OH (n = 3 per group). D–F, there was no significant change in the compositions of PC-PUFAs, PC-PUFAs-OOH, or PC-PUFAs-OH among control, mGPx4 KO, and vit E-supplemented mGPx4 KO mice (n = 3 per group). *p < 0.05, **p < 0.01 (KO with vitamin E versus KO), #p < 0.05 (KO versus control) using one-way ANOVA and the Tukey–Kramer post hoc test. DHA, docosahexaenoic acid; mGPx4, mitochondrial isoform of GPx4; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PUFA, polyunsaturated fatty acid; vit E, vitamin E.
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ferroptosis (39). Another study reported a role of MFN2 to mediate lipid peroxidation and arsenic-induced ferroptotic cell death in a hepatocyte cell line (40). In addition, the other study reported that relative lipid peroxidation is associated with mitochondrial fusion phenotypes (41). Based on these studies, we speculated in this study that the increased levels of MFN2 in mGPx4 KO retina might have mediated not only mitochondrial fusion but also lipid peroxidation and photoreceptor death. The other hypothesis might be that mitochondrial fusion in mGPx4 KO mouse retinas might not have been mediated by the decrease of PE (22:6/22:6), a unique molecular species of PE-containing double unsaturated fatty acid, and the increase of its oxidative products, PE-OOH (22:6/22:6-OOH) (Fig. 6), which needs further investigations.

In conclusion, the current study addressed the isoform-specific roles of GPx4 in the mitochondria for photoreceptor development and survival. mGPx4 is indispensable for cone photoreceptors at the early stage of their development. In contrast, rod photoreceptors can mature without mGPx4, although they degenerate gradually afterward. Consequently, the different effects of mGPx4 KO on cone and rod photoreceptors produce a retinal degeneration phenotype resembling human cone–rod dystrophy.

**Experimental procedures**

**Construction of mGPx4 KO transgenic vector**

To construct the mGPx4 KO transgenic gene (Tg(mGPx4KO), Fig. S1A), the approximate 5 kbp upstream regulatory region, E1a, and E1b regions (EcoRI/BamHI; Fig. S1A) containing a mutation in the first start codon for mitochondrial GPx4 from ATG to TTG, E1b and EII regions (EcoRI/Apal; Fig. S1A), and EII-VII and 1 kbp downstream region (EcoRI/ApaI; Fig. S1A) were separately prepared from our previously cloned mouse GPx4 gene (5, 9) and inserted into Bluescript SK+.

To identify mRNA for GPx4 transcribed from transgene, we replaced the NheI site located after the stop codon in the GPx4 gene with BamHI site (Fig. S1A). A transgenic screening vector pTG2 containing a 5′-end marker from part of the hygromycin complementary DNA (cDNA) (250 to 660 bp site), a marker sequence (L) for Tg(mGPx4 KO) transgenic gene screening sites (from the cGPx cDNA [27 to 272 bp site; 5′-GGCACAGTCCACCGTCTATG-3′/5′-CTTCTTTCATTTGCC-3′], and Luciferase cDNA [48 to 373 bp site; 5′-GCTGGGAAGATGGGACCGCT-3′/5′-CTTCTTTCATTTGCC-3′]) was constructed between two Sall sites in Bluescript SK+. Mouse genomic DNA containing a mutation at the start codon for mitochondrial GPx4 digested with Sall was inserted into the XhoI site of the TG2 vector. Finally, the mGPx4 KO transgenic gene, Tg(mGPx4 KO), was excised with Sall from pTG2 vector and purified.

**Animals**

All procedures were followed by the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research. All experiments were approved by the Institutional Animal Research Committee of the University of Tokyo and Kitasato University. Mice were housed in cages with food and water and subjected to a standard 12 h light/dark cycle at 22 °C.

mGPx4 KO mice were generated by transgenic rescue methods as previously described (12). Briefly, the Tg(mGPx4 KO) transgenic gene was injected into fertilized eggs derived from bromodomain factor 1 parents. Transgenic mGPx4 KO mice (GPx4+/−:GPx4tg(mGPx4 KO)/−) were generated by standard methods. To facilitate genotyping of the transgenic mice, the following primer pair specific to Tg (mGPx4 KO) was used: PHGP BAMS/LOXSCR-AS, 5′-CTTCTTATCCGCTATATCCGCC-3′. Tg(mGPx4 KO) mice were then mated with mGPx4 heterozygous (GPx4+/−) mice that we had established previously (2). Furthermore, Tg(mGPx4 KO) transgenic GPx4 heterozygous mice (GPx4+/−:GPx4tg(mGPx4 KO)/−) were mated with mGPx4 heterozygous mice (GPx4+/−) to obtain Tg(mGPx4 KO) transgene-rescued GPx4 KO mice (GPx4+/−:GPx4tg(mGPx4 KO)/−). Three primer pairs were used for genotyping (Fig. S1B); one specific for Tg(mGPx4 KO) as described previously; one specific for WT allele (103/GIR, 5′-CTTCTTATCCGCTATATCCGCC-3′); and one specific for KO allele (NEOF3/GIR, 5′-CTTCTTATCCGCTATATCCGCC-3′). After several generations, a mouse colony with homozygous transgenic mGPx4 KO alleles without endogenous GPx4 alleles (GPx4−/−:GPx4tg(mGPx4 KO)/tg(mGPx4 KO)) was established and referred to as mGPx4 KO mice. Since mGPx4 KO mice show male infertility, the mouse line was maintained by crossbreeding female mGPx4 KO (GPx4−/−:GPx4tg(mGPx4 KO)/tg(mGPx4 KO)) mice with male mGPx4 KO transgenic mGPx4 KO mice. Offspring genotyped as GPx4+/−:GPx4tg(mGPx4 KO)/tg(mGPx4 KO) were used as control mice to be compared with their mGPx4 KO littermate. mGPx4 KO and control mice used in this study were from a mixed genetic background with contributions of TT2, Institute of Cancer Research, and bromodomain factor strains for the experiments on male infertility and additional C57BL/6J background for the retina-related experiments. To investigate the reproductive capacities of mGPx4 KO and control mice, one male mouse was mated with two Institute of Cancer Research females for 2 weeks. Female mice were checked for vaginal plugs each morning, and litter sizes were recorded on delivery, after three successive matings. Vit E-enriched diet was made in the standard rodent diet CE-2 (CLEA Japan) enriched with DL-α tocopherol acetate (1000 IU/kg).

**Evaluation of epididymal spermatozoa and histological analysis of testis**

The cauda region of the epididymis obtained from ~12-week-old mice was placed in TYH medium (Mitsubishi Chemical Medience Corp) and gently minced with a surgical blade and incubated at 37 °C for 1 h. Spermatozoa were examined for structure, the number, and mitochondrial membrane potential. To measure mitochondrial membrane potential, spermatozoa were examined by monitoring the
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fluorescence of DiOC₆ (Molecular Probes). 1 x 10⁵ spermatooza were incubated for 1 h and then stained with 10 μg/ml DiOC₆, 1 μg/ml Hoechst 33258 for 20 min, and washed with PBS. The stained sperm cells were dropped onto glass slides. Fluorescence because of DiOC₆ and Hoechst in spermatooza was monitored and photographed with a fluorescence microscope (BIOZERO, BZ-8000; Keyence). Testis from WT mice and mGPx4 KO mice was fixed in Bouin’s solution and embedded in paraffin, and sections were stained with hematoxylin and eosin.

**Immunohistochemistry and TUNEL assay**

For frozen sections, enucleated eyeballs were fixed in 4% paraformaldehyde for 2 h and put in 30% sucrose in 4 °C overnight. Cryosections were made at −20 °C (thickness, 10 μm). Slides were blocked by Blocking One Histo (Nacalai Tesque, Inc) for 10 min. Then they were incubated with primary antibodies at 4 °C overnight and with secondary antibodies for 1 h at room temperature. The sections were then coverslipped with mounting medium. The following primary antibodies were used; antihodopsin (1:100 dilution; Santa Cruz Biotechnology; catalog no.: sc-57432), anti-Rxr-α (1:1000 dilution; Santa Cruz Biotechnology; catalog no.: sc-365252), anticone arrestin (1:1000 dilution; Sigma–Aldrich; catalog no.: AB15282), anti-S opsin (1:1000 dilution; Millipore; catalog no.: AB5407), and anti–cleaved-caspase-3 (1:800 dilution; Cell Signaling; catalog no.: 9664), and GPx4 (1:1000 dilution) (42). Alexa Fluor 488 (1:500 dilution; Thermo Fisher Scientific; catalog nos.: A-11209 and A-11008) and 647 (1:250 dilution; Abcam; catalog no.: ab150115)-conjugated secondary antibodies were used for secondary antibodies. For staining nuclei, 4,6-diamidino-2-phenyindole (Sigma–Aldrich) was used. Fluorescent conjugates of PNA lectin (Invitrogen; catalog no.: L21409) was used to label cone cells. ApopTag Peroxidase In Situ Apoptosis Detection Kit (Sigma–Aldrich) was used for labeling DNA-strand breaks according to the manufacturer’s protocol.

For immunohistochemistry of RPE flatmounts, enucleated eyeballs were fixed in 4% paraformaldehyde for 1 h. RPE-eyecups were flat mounted and blocked by Blocking One Histo for 1 h. Then they were incubated overnight at 4 °C with primary antibody (anti-ZO-1; 1:200 dilution; Thermo Fisher Scientific; catalog no.: 61-7300). After washing RPE flatmounts, they were incubated for 1 h with a secondary antibody (Alexa Fluor 488-conjugated secondary antibody; 1:250 dilution; Thermo Fisher Scientific; catalog no.: A-11008) and coverslipped with mounting medium.

**TEM and quantification of number and area of mitochondria**

The suspension of epididymal spermatooza was placed on silane-coated glass and fixed in a solution of 2.5% glutaraldehyde in 0.1 M phosphate buffer. The sample was postfixed in a solution of 1% OsO₄ in 0.1 M phosphate buffer, dehydrated through a graded series of ethanol solutions, and embedded in Epon 812. Ultrathin sections were double stained with uranyl acetate and lead citrate and then examined by TEM (Hitachi H-7100) operated at 75 kV. We examined 10 longitudinal profiles of sperm in 10 longitudinal sections of the midpiece.

Mouse eyes were enucleated and fixed in 2% paraformaldehyde at 4 °C overnight. The sections were made with 1.5 μm thickness and stained with 0.5% toluidine blue. The images were captured by a transmission electron microscope (100 kV, model JEM-1400Plus; JEOL Ltd) via a charge-coupled device camera (EM-14830RUBY2; JEOL Ltd). To compare the number of mitochondria, we counted the number of mitochondria of four sections of each group.

**Subcellular fractionation**

Cytosolic, mitochondrial, and nuclear fractions were prepared from WT, control, and mGPx4 KO retinas and testes. The tissues were dissected and homogenized in lysis buffer (0.25 M sucrose, 3 mM imidazole, and 1 mM EDTA-2Na) with 1% protease inhibitor (abcam; catalog no.: ab65320). The lysate was centrifugated at 2000 rpm for 10 min at 4 °C, and the supernatant was collected as postnuclear fraction. The pellet was resuspended with a Nuclear Extract kit (ACTIVE MOTIF 40010) for 30 min at 4 °C, and the solution was centrifugated at 14,000g for 10 min at 4 °C. The supernatant was used as nuclear fraction. The postnuclear fraction was centrifugated at 8000 rpm for 10 min at 4 °C. After centrifugation, the supernatant was collected as cytosol fraction. The pellet was resuspended with radioimmunoprecipitation assay buffer as mitochondrial fraction. Protein amounts in cytosolic and mitochondrial fractions were determined using the bicinchoninic acid assay (Thermo Fisher Scientific; catalog no.: 23227).

**Western blotting**

Samples were homogenized in radioimmunoprecipitation assay buffer (Thermo Fisher Scientific). Proteins were separated using 4 to 12% or 12% polyacrylamide gels and transferred to a polyvinylidene difluoride membrane (Bio-Rad Laboratories). The membrane was blocked by Blocking One (Nacalai Tesque, Inc) for 10 min and then was incubated with primary antibody overnight at 4 °C. The images of membrane were captured by an ImageQuant LAS 4000 mini-camera system (GE Healthcare). The following primary antibodies were used; anti-GAPDH (1:10,000 dilution; Wako Pure Chemical Industries; catalog no.: 016-25523), anti-SOD1 (1:1000 dilution; Santa Cruz; catalog no.: sc-27104), anti-TIM23 (1:1000 dilution; Proteintech; catalog no.: 11123-1-AP), anti-VDAC (Porin31 HL; 1:1000 dilution; Calbiochem), anti-PCNA (1:1000 dilution; BD Biosciences Pharmingen; catalog no.: 610,664), anti-GPx4 (1:5000 dilution) (42), antiangiogenin (1:1000 dilution; NOF; catalog no.: N213320), anti-MFN1 (1:100 dilution; Santa Cruz Biotechnology; catalog no.: sc-166644) anti-MFN2 (1:1000 dilution; Abcam; catalog no.: ab56889), anti–optic atrophy 1 (1:1000 dilution; BD Transduction Laboratories; catalog no.: 612606), anti-phospho-DRP1 (Ser637) (1:1000 dilution; Cell Signaling; catalog no.: 4867), and anti–phospho-DRP1 (Ser616) (1:1000 dilution; Cell Signaling; catalog no.: 3455). For secondary antibodies, horseradish peroxidase–linked antibody (Cell Signaling; catalog nos.: 7074 and 7076) were used. The band intensities were quantified using ImageJ software (version 1.53; National Institutes of Health).
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ERG

The ERG was recorded using a LS–W (Mayo) and analyzed using Scope 3 software (ADInstruments) for scotopic condition and LabChart 8 software (ADInstruments) for photopic condition. The scotopic ERG was recorded in 7-week-old mice, and the photopic ERG was recorded in 4-week-old mice. Scotopic ERG was recorded after 12 h of dark adaptation. The flash intensities ranged from 1.5 to 4500 cdms/m². Data from three trials were averaged for single-flash responses. Photopic ERGs were recorded after light adaptation. The flash intensities ranged from 1778 to 22,330 cdms/m² with a background illumination of 35 cd/m². Data from 32 trials were averaged for single-flash responses.

Phospholipid analysis by LC–MS

The retinas were isolated from enucleated eyes and stored at −80 °C immediately. The concentrations of PE, PE-OOH, PC, PC-OOH, and PC-OH in each sample were measured using QTRAP 4500 quadrupole linear ion trap hybrid mass spectrometer (AB Sciex) coupled to a Nexera XR high-performance liquid chromatography system (Shimadzu Co), as previously described (19, 43).

Statistics

All statistical analyses were performed using EZR (44) (Saitama Medical Center, Jichi Medical University), which is based on R 4.0.3 (The R Foundation for Statistical Computing). EZR is a modified version of R commander designed to add statistical functions frequently used in biostatistics. Data are presented as a mean value with standard error. Student’s t test was performed to compare two groups. For comparisons among three groups, one-way ANOVA was conducted, followed by Tukey’s post hoc test. p Values less than 0.05 were considered to be statistically significant.

Data availability

All data are contained within the manuscript.

Supporting information—This article contains supporting information.

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Abbreviations—The abbreviations used are: cDNA, complementary DNA; cGPx4, cytosolic isoform of GPx4; DHA, docosahexaenoic acid; DRP1, dynamin-related protein 1; ER, endoplasmic reticulum; ERG, electroretinogram; GPx4, glutathione peroxidase 4; MFN1, mitofusion 1; MFN2, mitofusion 2; mGPx4, mitochondrial isoform of GPx4; nGPx4, nuclear isoform of GPx4; ONL, outer nuclear layer; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PNA, peanut agglutinin; PUFA, polyunsaturated fatty acid; RPE, retinal pigment epithelium; TEM, transmission electron microscopy; vit E, vitamin E.

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