The Novel Pathogenesis of Retinopathy Mediated by Multiple RTK Signals is Uncovered in Newly Developed Mouse Model

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Pericyte desorption from retinal blood vessels and subsequent vascular abnormalities are the pathogenesis of diabetic retinopathy (DR). Although the involvement of abnormal signals including platelet-derived growth factor receptor-β (PDGFRβ) and vascular endothelial growth factor-A (VEGF-A) have been hypothesized in DR, the mechanisms that underlie these processes are largely unknown. Here, novel retinopathy mouse model (N-PRβ-KO) was developed with conditional Pdgfrb gene deletion by Nestin promoter-driven Cre recombinase (Nestin-Cre) that consistently reproduced through early non-proliferative to late proliferative DR pathologies. Depletion of Nestin-Cre-sensitive PDGFRβ+ NG2+ αSMA+ pericytes suppressed pericyte-coverages and induced severe vascular lesion and hemorrhage. Nestin-Cre-insensitive PDGFRβ+ NG2+ αSMA+ pericytes detached from the vascular wall, and subsequently changed into myofibroblasts in proliferative membrane to cause retinal traction. PDGFRβ+ astroglia was seen in degenerated retina. Expressions of placental growth factor (PIGF), VEGF-A and PDGF-BB were significantly increased in the retina of N-PRβ-KO. PDGF-BB may contribute to the pericyte-fibroblast transition and glial scar formation. Since VEGFR1 signal blockade significantly ameliorated the vascular phenotype in N-PRβ-KO mice, the augmented VEGFR1 signal by PIGF and VEGF-A was indicated to mediate vascular lesions. In addition to PDGF-BB, PIGF and VEGF-A with their intracellular signals may be the relevant therapeutic targets to protect eyes from DR.

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1. Introduction

Retinopathy caused by diabetes mellitus is one of the major causes of vision impairment worldwide [10,48]. Microvascular degeneration including increased vascular permeability, microaneurysms and acellular capillaries as well as related ischemic damages are features of non-proliferative and early diabetic retinopathy (DR). Along the progress to proliferative DR, the risk of irreversible vision loss is increased due to macular oedema, hemorrhage, and proliferative membrane-mediated tractional retinal detachment [48]. Accordingly, for the protection and restoration of visual acuity, further studies are necessary that focus on the mechanism of disease progression from early non-proliferative to late proliferative DR.

Functional disturbance and apoptotic death of vascular cells are mediated by many mechanisms in early DR, among which, chronic disturbance of platelet derived growth factor receptor (PDGFR) β signal induced by sustained hyperglycemia importantly mediates pericyte injury that causes vascular abnormalities in DR [19]. Along this line, the vascular lesions of retina with decreased pericyte-coverage have been analyzed as a useful model of DR that were induced in hypomorphic mutant of Pdgfb or Pdgfrb genes or in the eyes treated with neutralizing antibody against PDGFβR [7,25,33,37]. Besides these, PDGFs are augmented in DR, and PDGF-BB is specifically upregulated in proliferative DR [39]. PDGF is a mitogen of glial cells and mesenchymal cells including vascular pericytes [22,23]. Thus, PDGF may be involved in the formation of proliferative membrane and gliosis in proliferative DR, although the roles are remained unknown.

Vascular endothelial growth factor (VEGF)-A [15] that specifically binds VEGF receptors (VEGFRs) 1 and 2 is upregulated in DR in response to the ischemia induced by retinal vascular damage, and exacerbates extra-retinal vascular outgrowth to the retinal surface without amelioration of ischemia in the retina [18,31,35,44,48,50]. VEGF-A is the best characterized therapeutic target of DR, and anti-VEGF-A therapy significantly improved the prognosis of DR; however, significant part of...
patients with DR insufficiently respond to this therapy [47]. Placental growth factor (PIGF) is another angiogenic growth factor of VEGF family that specifically binds VEGFR1 [11,36,49], and data have accumulated showing the increase of PIGF in the vitreous humour and retina of eyes with DR as reviewed [32]. Thus, PIGF is supposed as an additional intrinsic therapeutic target of DR; however the roles of PIGF as well as VEGFR1 in DR remain to be elucidated.

In the present study, we established a new retinopathy model mice (N-PRβ-KO mice) with the C57BL/6 genetic background, harboring the Pdgfrb gene flanked by two loxP sequences (Pdgfrblox/lox) and Nestin promoter-driven Cre recombinase. The phenotype of retinopathy is inherited with a high penetration rate, and the model consistently reproduces the early to late phases of DR pathologies, whereas these changes occurred in early periods of life. PDGFRxα and PDGFRxβ activated by increased PDGF-BB were indicated to be involved in astroglisis and formation of proliferative membrane in retinopathy, respectively. PIGF and VEGF-A were upregulated in retinopathy in N-PRβ-KO mice, and genetrical blockade of VEGFR1 signal substantially ameliorated the retinal vascular lesions. Our findings uncovered the novel pathogenesis mediated by PDGF-BB-PDGFRs and PIGF/VEGF-A-VEGFR1 signal axes.

2. Materials and Methods

2.1. Conditional Knockout Mice

All experimental animal procedures were conducted according to the Institutional Animal Care and Use Committee at the University of Toyama (Sugitani, Toyama, Japan). All study protocols were approved by the Ethics Committee of the University of Toyama before the studies were performed.

We previously established a Pdgfrblox/lox mouse line with the 129S5 genetic background [17]. The congenic strain C57BL/6-129S5.1 harboring Nestin promoter/enhancer-driven Cre recombinase was obtained (Nestin-Cre, Jackson Laboratories, Bar Harbor, ME, USA). In the present study, backcross mating was carried out for 15 generations between Pdgfrblox/lox mice and wild-type (WT) C57BL/6 mice (Japan SLC, Shizuoka, Japan) and for 18 generations between Nestin-Cre mice and WT C57BL/6 mice. The resulting N-PRβ-KO offspring showed severe retinopathy phenotypes with a very high penetration rate. All mice were housed at 25 °C with a 12-h light/dark cycle with free access to pellet chow and water. Genotyping of the Pdgfrblox/lox [17], Nestin-Cre [41], R26R-mCherry [23] and Flt1TK−/− [21] were performed by genomic PCR. The primer sequences are given in Supplementary information.

2.2. Immunofluorescence

For paraffin section and whole mount sample preparation, excised eyes were fixed with 4% paraformaldehyde at 4 °C overnight. Eyes were dehydrated with a graded series of ethanol solutions, and embedded in paraffin. The sections (5 μm in thickness) were subjected to immunofluorescence and hematoxylin and eosin (HE) staining. In whole mount analysis, retinas were peeled out, and then immunofluorescence staining was performed as described previously [23,31,53]. Briefly, the specimens were incubated at 4 °C overnight with the appropriate primary and secondary antibodies. The details of antibodies are given in Supplementary Information. Specimens for immunofluorescence were observed using a LSM780 confocal system (Carl Zeiss, Oberkochen, Germany), a TCS SP5 confocal system (Leica, Heidelberg, Germany), and a BX-700 microscope (Keyence, Osaka, Japan). Images were appropriately processed using Photoshop software (version 7.0; Adobe, San Jose, CA, USA).

2.3. RNA Extraction, cDNA Synthesis, and Real-Time PCR

Total RNAs were isolated from retinas of N-PRβ-KO and WT mice using the miRNeasy Mini Kit (Qagen, Valencia, CA, USA), and were reverse-transcribed according to the PrimeScript RT Reagent Kit protocol (TAKARA BIO INC., Shiga, Japan). cDNAs were diluted 1:25 in the reaction mixture consisting of SYBR Premix EX Taq II (TAKARA BIO INC.), and then real-time PCR performed using a TAKARA Thermal Cycler Dice Real Time System TP800 as previously described [52]. Induction values were calculated using analysis software (TAKARA BIO INC.). Primer sequences are available upon request from the TAKARA BIO INC. website (http://www.takara-bio.co.jp).

2.4. Western Blotting

Sample preparations and all other procedures for western blotting are described elsewhere [41,52]. Briefly, retinas were lysed using T-PER Tissue Protein Extraction Reagent (Thermo Fisher Scientific Inc.) on ice. Lysates were electrophoretically separated, and then transferred to polyvinylidene difluoride membranes. Skim milk blocked membranes were incubated with primary antibodies described in Supplementary Information, and then incubated with the appropriate horseradish peroxidase-conjugated secondary antibodies. Immunoactive bands were detected using enhanced chemiluminescence reagents (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) according to the manufacturer’s instructions.

2.5. Enzyme-Linked Immunosorbent Assay

The concentrations of PDGF-AB, PDGF-BB, VEGF-A, and PIGF in the retinas of N-PRβ-KO or WT mice were measured by ELISA according to the manufacturer’s instructions (R&D, Minneapolis, MN, USA). Briefly, collected retinas were snap-frozen in liquid nitrogen, smashed into a powder with the Multi-Beads Shocker (Yasui-koki, Osaka, Japan), and lysed using T-PER Tissue Protein Extraction Reagent (Thermo Fisher Scientific Inc., Waltham, MA, USA) with Halt Protease Inhibitor Cocktail (Thermo Fisher Scientific Inc.). After centrifugation, lysates were used as samples. The concentrations of PDGF-AB, PDGF-BB, VEGF-A, and PIGF were calculated from the standard curve.

2.6. Image Analysis and Quantification

Images of HE sections of retinas were captured using an IX71 microscope (Olympus) and the retinal thickness of the peripheral area was measured using cellSens Standard (ver. 1.4.1; Olympus). Images of the whole mount retinas were captured using a LSM780 confocal system (Carl Zeiss) and a BX-700 microscope (Keyence). Percentage of the immunoreactivity of CD31 and collagen type IV (pixel) in whole retinal area (pixel) were analyzed using BZ-H3C software (Keyence). For pericyte coverage on the retinal vasculature, the percentage of area that was immunoreactive (pixel) for NG2, CD13 and CD31 was analyzed using ImageJ (National Institutes of Health, Bethesda, MD, USA). For the analysis of western blotting data, the immunoreactive bands were quantified using ImageJ (NIH) and normalized to the β-actin band.

2.7. Statistical Analysis

All values are presented as means ± SEM. Statistical analyses of differences between groups were performed using Student’s t-tests or one- or two-way analyses of variance (ANOVA) with Tukey’s multiple comparison tests for post-hoc analysis were used. A p-value of <0.05 was considered statistically significant. Graphs were drawn using GraphPad Prism 6 software (GraphPad Software Inc., La Jolla, CA, USA).

3. Results

3.1. N-PRβ-KO Mouse Show Pathological Vascular Remodeling and Tractional Retinal Detachment

N-PRβ-KO mice with conditionally depleted Pdgfrb (Fig. 1a) grew to the adult stage and fertile without visible structural abnormalities in the
organ such as brain and kidney (Fig. S1), but except in their eyes, and blood glucose levels were not altered (118.6 ± 7.9, 118.4 ± 15.3, 117.9 ± 5.0, and 111.5 ± 4.0 mg/dL in male and female N-PR\(\beta\)-KO mice and male and female wild-type (WT) mice, respectively, at 4 weeks of age). Most of these mutants exhibited vitreous opacity in bilateral eye in the adult stage (8 to 12 weeks of age; Fig. 1b). Tractional retinal detachment with an abnormal shape was characteristic of N-PR\(\beta\)-KO mice, but was not seen in WT mice (Fig. 1c). Arbitrary scoring of the retinal shape abnormality indicated that 90% of N-PR\(\beta\)-KO mice showed some degree of retinal injury, and male and female mice were similarly affected (Fig. 1d–f). The tractional retinal detachment was histologically identified as a V-shaped deformity of the retina. It is of note that an \(\alpha\)SMA\(^{+}\) proliferative membrane was identified on the retinal surface of this deformity (Fig. 1g). The normal layered structures of the retina were highly destroyed, resulting in the thinning of the peripheral area of the retina in N-PR\(\beta\)-KO mice at 10 weeks old. Highly perturbed thin retinas are observed compared to WT retinas. (i and j) N-PR\(\beta\)-KO showed significantly thinner retinas than those of WT mice for both males (i) and females (j) in the adult stage. n = 5–8 males and females. All values represent means ± SEM. *, \(p < 0.05\); **, \(p < 0.01\); ***, \(p < 0.001\), compared to WT mice at the same time points. Scale bar = 1 mm (c), 100 \(\mu\)m (g and h).

Fig. 1. N-PR\(\beta\)-KO mice show severe retinal structural defects in adult mice. (a) Schematic representation of the transgenic and mutated alleles of N-PR\(\beta\)-KO. Exons 4 to 7 of the Pdgfrb gene flanked by two loxP sites are deleted by nlsCre activity driven by the Nestin promoter. (b and c) N-PR\(\beta\)-KO mice showed vitreous opacity (b, red arrowhead) and detached retina (c) compared with wild-type (WT) mice at 8–12 weeks old. (d) Scoring of retinal deformities: 0, normal cup-shaped retina; 1, distorted cup-shaped retina; 2, detached V-shaped retina or shrunken retina; 3, highly detached V-shaped and very thin peripheral retina or highly shrunken and very thin retina; 4, rudimentary retina; 5, not detected. (e and f) Both male (e) and female (f) N-PR\(\beta\)-KO mice showed higher deformity scores at 8–12 weeks old than those of WT mice. n = 11–14 male retinas, n = 30–33 female retinas. (g) Typical proliferative membrane on the retinal surface of N-PR\(\beta\)-KO mice at 10 weeks old. The proliferative membrane includes \(\alpha\)SMA-positive cells (red) and blood vessels (indicated by asterisks in magenta). Hoechst staining depicts nuclei (blue). (h) Histology (H&E staining) of the peripheral region of the N-PR\(\beta\)-KO retina at 10 weeks old. Highly perturbed thin retinas are observed compared to WT retinas. (i and j) N-PR\(\beta\)-KO showed significantly thinner retinas than those of WT mice for both males (i) and females (j) in the adult stage. n = 5–8 males and females. All values represent means ± SEM. *, \(p < 0.05\); **, \(p < 0.01\); ***, \(p < 0.001\), compared to WT mice at the same time points. Scale bar = 1 mm (c), 100 \(\mu\)m (g and h).
Increased CD31$^+$ staining in N-PRβ-KO mice was confirmed morphometrically in both sexes, to a similar extent (Fig. 2b and c). Collagen type IV staining was greater in N-PRβ-KO than in WT mice (Fig. 2a, bottom row) and the difference between the two genotypes was significant based on a morphometric analysis (Fig. 2d and e). Similar results were obtained for both genders. Based on observations at a high magnification, CD13$^+$ pericytes drastically disappeared around the CD31$^+$ retinal blood vessels of N-PRβ-KO mice compared to WT mice (Fig. 2f). Accordingly, the percentage of pericyte coverage was substantially decreased in retinal vessels of N-PRβ-KO mice, irrespective of gender (Fig. 2g and h). Furthermore, many collagen type IV$^+$/CD31$^-$ structures, corresponding to empty sleeves [45],

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**Fig. 2.** N-PRβ-KO adult mice show severe retinal blood vessel defects. (a) N-PRβ-KO mice show retinal blood vessel defects in both males and females at the adult stage (8–12 weeks old). Immunofluorescence of CD31 (red) and collagen type IV (cyan) depict retinal vasculature and blood vessel specific ECM deposition of N-PRβ-KO mice compared with WT. (b and c) Ratio of the CD31-positive area compared to the whole area of the retina. N-PRβ-KO mice showed significantly higher ratios for both males (b) and females (c) compared with WT in the adult stage. n = 15 for males and females. (d and e) Ratio of the collagen type IV-positive area compared to the whole area of the retina. N-PRβ-KO mice showed significantly higher ratios for both males (d) and females (e) compared with WT in the adult stage. n = 15 for males and females. (f) Multi-colour representative immunofluorescence images of WT (upper row) and N-PRβ-KO mice (bottom row) at 10 weeks old. CD31-positive blood vessels (red), CD13-positive pericytes (green), collagen type IV deposition (cyan). Arrowheads indicate empty sleeves. (g and h) Percentage of pericyte coverage on the retinal vasculature of N-PRβ-KO mice. N-PRβ-KO mice showed significantly lower coverage rates for both males (g) and females (h) compared with WT in the adult stage. n = 20 randomly selected areas (20× objective lens) from 5 retinas of males and females. All values represent the mean ± SEM. *** p < 0.001, compared to WT. Scale bar = 1 mm (a), 100 μm (f).
indicated vascular regression in N-PR−/−KO mice, but not in WT mice (Fig. 2f, arrowheads). These collagen structures and abnormal vascular branches (Fig. 2f) may contribute to the increased measurement of CD31 and collagen type IV positive areas in N-PR−/−KO mice (Fig. 2b–e). PDGFRβ-null embryos show pericyte coverage deficiency [20]. To confirm whether the severe depletion of pericytes in N-PR−/−KO mice was a consequence of Cre recombinase-mediated Pdgfrb gene knockout, we conducted a fate mapping study utilizing Nestin-Cre;R26R-mCherry mice (N-MC, Fig. S2a). Many, but not all pericytes expressed mCherry of the retinal blood vessels in N-MC mice (Fig. S2b–d, magenta–arrowheads). The mCherry+ pericytes significantly outnumbered mCherry− pericytes in arterioles and capillaries in N-MC mouse retina (Fig. S2e and g). In contrast, mCherry− pericytes significantly outnumbered mCherry+ pericytes in venules of the N-MC mice (Fig. S2f). These findings indicate that Pdgfrb knockout in retinal pericytes in N-PR−/−KO mice resulted in severe depletion of retinal pericytes.

PDGFRβ insufficiency in pericytes induced proliferative DR-like lesions, including tracional retinal detachment associated with the proliferative membrane, and vascular phenotypes consisting of angiogenesis and regression in adult N-PR−/−KO mice. These data prompted us to investigate N-PR−/−KO mice at an earlier age to elucidate the developmental mechanism of the retinal lesion.

3.2. Retinal Degeneration was Induced by Hypoperfusion in N-PR−/−KO Mice

The thickness of the retina in N-PR−/−KO mice was comparable with that of WT mice until 10 days after birth, and then gradually decreased, becoming significantly less than that of WT at 8 weeks old (Fig. S3a). CD31+ microvessels were dilated, and the extent of branching of the microvessels decreased (Fig. S3b). Vascular regression indicated by the empty sleeve was a prominent feature in N-PR−/−KO mice (Fig. S3b). Similarly non-functioning retinal vessels were often observed in N-PR−/−KO but not in WT mice (Fig. S3c). These indicate that the vascular regression-mediated local hypoperfusion is the underlying mechanism of the current retinopathy as has been demonstrated in DR [45]. In accordance with these, hypoperfusion-induced oxidative stress [6] was detected in retinopathy of N-PR−/−KO mice. 8-OHdG, an oxidative stress marker, was detected in glutamine synthetase+ Müller cells, recoverin+ photoreceptors, PKCa+ bipolar cells, calretinin+ amacrine cells, and calbindin+ horizontal cells in the inner nuclear layer of the retina of 2 weeks old N-PR−/−KO mice (Fig. S4a–e). Whereas TUNEL-positive apoptotic cells did not co-localize 8-OHdG in N-PR−/−KO at this time point (data not shown), 8-OHdG was clearly co-expressed with SQSTM1 (Fig. S4f), an autophagy marker, suggesting that oxidative stress-induced autophagic cell death underlies the structural destruction and thinning of the retina after vascular phenotypes, e.g., dilation and regression, are detected in N-PR−/−KO mice.

3.3. Molecular Mechanisms Underlying Retinal Detachment in N-PR−/−KO Mice

The deformation of the retina represented by a "retinal score" (Fig. 1d) was not significant at 1 week old, became significant at 2 weeks old and was further aggravated at 4 weeks old in both sexes of N-PR−/−KO mice as compared with WT mice (Fig. S5a and b). Immunofluorescence of whole mount retinal specimens showed that the network of CD31+ blood vessels associated with αSMA+ pericytes spread at 1 week, and a hierarchical vascular tree was established after 3 weeks in WT retina (Fig. 3a–c, left). Comparing to these, blood vessels were irregularly dilated, and αSMA+ pericytes aberrantly distributed in association with capillaries at 1 week of age in N-PR−/−KO retina (Fig. 3a, right). Notably, triple-colour immunofluorescence at this time point demonstrated that almost all NG2+ αSMA− microvascular pericytes disappeared but NG2+ αSMA+ pericytes still observed in microvasculatures of N-PR−/−KO compared to same region of WT (Fig. S5c). At 2 weeks of age, increased CD31 staining indicated increased angiogenesis with an irregular network pattern, vascular dilatation, and microaneurysm in N-PR−/−KO retina (Fig. 3b, right). These vascular phenotypes were frequently accompanied by large hemorrhages (Fig. S5d). Noteworthy, foci of the proliferative membrane consisted of αSMA+ cells appeared at 2 weeks (Fig. 3b, right, arrowhead) and increased at 4 weeks (Fig. 3c, right, arrowheads), that could underlie the retinal deformation detected in these periods (Fig. S5a and b).

At the higher-magnification views, full coverage of NG2+ pericytes on the blood vessels was observed in the WT retina at 4 weeks of age (Fig. 3d, upper row). Noteworthy, NG2+ αSMA− pericytes were exclusively located in the microvasculature (Fig. 3d, upper row, green–arrowheads), and NG2+ αSMA+ pericytes preferentially enwrapped the arterioles, venules, and their primary branches (Fig. 3d, upper row, magenta–arrowheads). In age-matched N-PR−/−KO retinas, NG2+ αSMA− microvascular pericytes were also severely depleted (Fig. 3d, bottom row, green–arrowheads), whereas considerable number of NG2+ αSMA+ pericytes were distributed along the microvasculature (Fig. 3d, bottom row, magenta–arrowheads). This depletion of NG2+ αSMA− microvascular pericytes corresponded to the decreased pericyte coverage of blood vessels (Fig. S5a and b), that is one of the most important vascular lesions of DR. Besides that, the proliferative membrane included many αSMA+ cells and irregularly aggregated microvasculatures (Fig. S5c). These observations strongly suggest that NG2+ αSMA− pericytes in N-PR−/−KO are Nestin-Cre-insensitive pericyte subset (it means Pdgfrb preserving pericyte subset), and such pericyte subset may correspond to Nestin-Cre-insensitive mCherry+ αSMA− pericytes (Fig. S2). These cells often detached from blood vessels to become NG2+ αSMA− pericytes after complete detachment (Fig. S6d). Therefore, they may change into myofibroblasts and may contribute to the formation of the proliferative membrane, in which the foci of proliferative membranes expressed collagen type I and connective tissue growth factor (CTGF) that are well-known factors for involving in the tissue fibrosis (Fig. S6e–h). Such issue will be further verified in following Fig. 5b–d.

To determine the molecular mechanisms underlying proliferative membrane formation in N-PR−/−KO retinas, the expression levels of PDGFs and receptors were examined, which are reportedly involved in fibroblast migration [17] and pericyte detachment [24]. Based on real-time PCR analyses, Pdgfb mRNA expression in the retina was significantly higher in N-PR−/−KO mice than in WT mice at 2 weeks old (Fig. 4b), and the mRNA expression levels of other PDGF ligands were comparable between the two genotypes (Fig. 4a–d). Similarly, based on ELISA, the expression of PDGF-BB, but not PDGF-AB, increased substantially at 2 weeks old and remained at a high level at 4 weeks old in the N-PR−/−KO retina (Fig. 4e and f). Pdgfrα mRNA expression was significantly higher in N-PR−/−KO retinas than WT retinas at 2 weeks old and remained at a high level at 4 weeks old (Fig. 4g). Pdgfrβ mRNA was significantly lower in N-PR−/−KO retinas than WT retinas at 1–4 weeks old (Fig. 4h). Similarly, increased PDGFRα and decreased PDGFRβ were detected in N-PR−/−KO retinas compared to WT retinas by western blotting (Fig. 4i and j). In immunofluorescence, PDGFRα was localized in a few GFAP+ astrocytes in the ganglion cell layer of WT mice. In N-PR−/−KO retinas, increased PDGFRα was distributed on many glial scar-forming hypertrophic GFAP+ astrocytes; the glial scar might contribute to tracional retinal detachment (Fig. 5a), as previously suggested [42]. PDGFRα was still expressed in αSMA+ pericytes, but was mostly expressed in detached αSMA+ pericytes of N-PR−/−KO mice (Fig. 5b, arrowheads). Fate mapping in Nestin-Cre;Pdgfrβfl/fl;R26R-mCherry (N-PR−/−KO-MC, Fig. 5c) showed mCherry+ αSMA+ detaching pericytes (Fig. 5d, green–arrowheads), which correspond to Nestin-Cre-insensitive mCherry+ αSMA+ pericytes as shown in Fig. 52. These findings, together with Fig. S6d, strongly suggested that Nestin-Cre-insensitive PDGFRβ+ NG2+ αSMA+ pericytes were incorporated into proliferative membrane through pericyte-fibroblast transition (PFT), and that this process appeared to be driven by PDGFB-BB-PDGFRβ signal axis.
Foci of proliferative membrane and structural abnormality of blood vessels are observed in early stage of N-PRβ-KO mouse retina. (a–c) Retinal angiogenesis of WT and N-PRβ-KO at 1 week (a), 2 weeks (b) and 4 weeks (c). CD31+ retinal vascular networks (cyan) develop toward the peripheral region. αSMA+ pericytes (red) that preferentially reside on the arteriole and venule are observed (a–c, left) in WT retinas. Owing to a loss of NG2+ αSMA− microvascular pericytes, N-PRβ-KO retina at 1 week show relatively large-sized capillaries. Additionally, defects in the specification of arterioles and venules are often observed, and atypical migration of NG2+ αSMA− pericytes can be found (a, right). Blood vessel dilation and microvascular aneurisms are often observed at 2 to 4 weeks. At the same time points, foci of proliferative membrane constituted by αSMA+ pericytes (magenta-arrowheads, myofibroblast-like phenotype) are observed (b and c, right). (d) Multi-colour representative immunofluorescence images of WT (upper row) and N-PRβ-KO mice (bottom row) at 4 weeks. NG2+ pericytes (green), αSMA+ pericytes (red), CD31-positive blood vessels (cyan). In WT, whereas NG2+ αSMA− pericytes are preferentially observed on arterioles, venules and their primary branches (magenta-arrowheads), NG2+ αSMA− pericytes are exclusively observed on the capillaries (green-arrowheads). In N-PRβ-KO, considerable NG2+ αSMA+ pericytes are observed on microvasculature (magenta-arrowheads) instead of NG2+ αSMA− pericytes (green-arrowheads). In addition, naked blood vessels are often observed (blue-arrowheads). Scale bar = 1 mm (a–c), 100 μm (d).

|   | WT | N-PRβ-KO |
|---|----|----------|
| a | ![Image](image1.png) | ![Image](image2.png) |
| b | ![Image](image3.png) | ![Image](image4.png) |
| c | ![Image](image5.png) | ![Image](image6.png) |
| d | ![Image](image7.png) | ![Image](image8.png) |
mRNA tended to be higher in N-PR phenotype. Among the major angiogenic factors examined, PlGF exclusively binds and transduce intracellular signaling via VEGFR1 [11,49]. VEGF-A also binds VEGFR1, which plays an important role in development [16,21]. Accordingly, to understand the effects of increased PlGF and VEGF-A in N-PRβ-KO retinas, intracellular signaling via VEGFR1 was blocked in N-PRβ-KO mice by cross-breeding with Flt1TK−/− mice [21]. In Flt1TK−/− mice, no distinct vascular abnormalities could be observed in retinas compared to WT in physiological condition (Fig. S7). In pathological condition of retina shown in N-PRβ-KO, whereas no explicit changes were observed in aberrant migration of αSMA+ pericytes, formation of the proliferative membrane, and retinal structural destruction and thinning between N-PRβ-KO and N-PRβ-KO-Flt1TK−/− retinas, severely dilated blood vessels were restored to normal sized blood vessels in N-PRβ-KO-Flt1TK−/− retinas (Fig. 7a and b). Furthermore, the excessive deposition of collagen type IV in N-PRβ-KO retinas was decreased in N-PRβ-KO-Flt1TK−/− retinas (Fig. 7b). Increases in angiogenesis and vessel diameter in N-PRβ-KO retinas were significantly restored in N-PRβ-KO-Flt1TK−/− retinas (Fig. 7c and d). Moreover, increase in collagen type IV deposition in N-PRβ-KO retinas was significantly restored in N-PRβ-KO-Flt1TK−/− retinas (Fig. 7e and f). Accordingly, VEGFR1 signal activated by PlGF or VEGF-A is involved in pathological angiogenesis with a large hemorrhagic phenotype in retinopathy of N-PRβ-KO mice.

4. Discussion

Pericyte desorption from blood vessels mediated by insufficient PDGFRβ signal is followed by subsequent vascular abnormalities in DR. In a meanwhile, VEGFs in retina induced by hypoxia mediate proliferative DR [3]. These processes of disease progress were consistently represented in our mouse model with conditional Pdgfrb gene targeting by Nestin promoter-driven Cre. In N-PRβ-KO mice, blood vessels with severely decreased pericyte-coverage showed abnormal vascular dilatation, regression and hemorrhages. These were followed by changes like proliferative DR including angiogenesis, gliosis and proliferative membrane formation with tractional retinal detachment. Furthermore, VEGF-A was augmented in degenerated DR with oxidative stress and autophagic responses as reported in DR [43]. Accordingly, we examined

![Image](image_url)

3.4. Molecular Mechanisms Underlying Pathological Angiogenesis in N-PRβ-KO Mice

To clarify underlying cellular and molecular mechanisms of pathological angiogenesis observed in the N-PRβ-KO retinas, we next examined the angiogenic factors corresponding to the above vascular phenotype. Among the major angiogenic factors examined, Vegfa mRNA tended to be higher in N-PRβ-KO than in WT retinas at 1–2 weeks old (Fig. 6a). Pdgfa mRNA in N-PRβ-KO retinas increased after 2 weeks, and the difference between the two genotypes was significant at 4 weeks (Fig. 6e). The mRNA expression levels were comparable between the two genotypes or tended to be lower in N-PRβ-KO than in WT retinas for all other VEGF ligands (Fig. 6b–d). Receptors for VEGF family were comparable between the two genotypes or tended to be lower in N-PRβ-KO than in WT retinas (Fig. 6f–h). Consistent with these results, VEGF-A protein expression was significantly higher at 2 weeks (Fig. 6i) and PlGF protein expression was significantly higher at 2 and 4 weeks (Fig. 6j) in N-PRβ-KO retinas than WT retinas.

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the key molecules and mechanisms underlying mouse retinopathy; these results have implications for the prevention and care of DR, a major cause of vision-impairment worldwide [10,48].
damage including proliferative DR pathology in hypomorphic mutations in PDGFRβ or PDGF-B, while the details of proliferative DR pathology were not described [7]. Pericytes that cover cranial vessels derived from either neural crest cells or mesodermal cells emerged around the tube formed endothelial cells, proliferated and migrated along such tubes [1,54]. Accordingly, we chose Nestin promoter, a promoter activated in neural crest cells, to target Pdgfb in neural crest-derived pericytes in N-Prβ−KO mice. While considerable number of Nestin-Cre-insensitive PDGFRβ+/NG2+/αSMA+ pericytes was remaining, Nestin-Cre-sensitive PDGFRβ+/NG2−/αSMA− subset of pericytes was mostly depleted, resulting in severe loss of pericyte coverage of retinal capillaries. These may indicate that neural crest-derived pericytes take major part of coverage in retinal blood vessels. 

αSMA expressing fibroblasts from an uncertain origin contract extracellular matrix, and are thought to be important for the traction retinal injury [4,27]. In our previous study, the tumor cell-derived PDGF-BB built up a high gradient that attracted PDGFRβ+/NG2+ pericytes to move away from tumor microvessels and transformed into αSMA+/FSP+/PDGF-Rα+ cancer-associated fibroblasts [24], in which such cellular transition were termed pericyte-fibroblast transition (PFT). Similar PFT was also mediated by macrophage-derived PDGF-BB in an adipose tissue after high-fat diet [34]. In N-Prβ−KO mice, while Nestin-Cre-sensitive PDGFRβ+/NG2−/αSMA− pericytes were mostly depleted from retinal capillaries, Nestin-Cre-insensitive PDGFRβ+/NG2−αSMA− pericytes still remained on arterioles and venules. These remaining pericytes, however, did not fully enwrap the microvasculature; instead, detached from blood vessels, eventually losing NG2 and PDGFRβ expression, and were incorporated into the proliferative membrane. Accordingly, our data indicated, for the first time, that PFT could be, at least in part, the origin of αSMA+ retinal myofibroblasts. PDGF-Rα+ astroglia was another important finding in the retinas of N-Prβ−KO mice. Induced overexpression of PDGF-BB in photoreceptors causes astrogial cell-mediated tractional retinal detachment [42]. PDGF-BB was substantially augmented in the retinas of N-Prβ−KO mice, indicating that PDGF-BB may contribute to tractional retinal detachment via PFT and by astrogliosis, and thus could be an important therapeutic target for the prevention of retinal detachment.

In N-Prβ−KO retinas, PIGF and VEGF-A were significantly upregulated at the progression phase of the disease. Similarly, PIGF and VEGF-A are increased in patients with proliferative DR, and the increases of them were correlated well to each other [28,30,46]. VEGF-A binds VEGFR1 and VEGFR2, and PIGF exclusively binds VEGFR1 [36]. In addition to the angiogenic effects via direct VEGFR1 activation, PIGF can synergistically contribute to angiogenesis and plasma extravasation in pathological conditions together with VEGF-A via several different mechanisms; e.g., in experimental angiogenesis models, PIGF/VEGF1 bindings freed up VEGF-A to activate VEGFR2, and PIGF/VEGF1 transphosphorylated VEGFR2 ([5,9]; [13]). After these, the present study of genetic inactivation of VEGFR1 demonstrated the critical role of VEGFR1 in pathological angiogenesis of retinopathy, where PIGF and VEGF-A were increased. Along this line, Affibrecpt, a decoy receptor that can bind and inhibit VEGF-A and PIGF, is highly effective with respect to therapeutic outcomes for DR [12]. Correctly, PIGF, VEGF-A, and their shared receptor VEGFR1 are indicated to be a therapeutic target of DR that may restore the normal vascular structures. 

Currently, anti-VEGF-A therapy using Ranibizumab, Bevacizumab, Affibrecpt and Pegaptanib is the most established treatment for pathological blood vessel formation in DR [8,14,22,38,40]. In addition to this, retinal damage, such as tractional detachment, proliferative membrane formation, and structural destruction, are serious threats for vision loss. In this study, we established a useful mouse retinopathy model that can consistently reproduce proliferative DR. Based on analyses of this retinopathy, PDGF-BB/PDGFRα and PDGFRβ axes are potential therapeutic targets for the prevention of tractional retinal detachment, and suppression of PIGF/VEGF-A-VEGFR1 axis may be a novel therapeutic target and
prevent pathological angiogenesis in proliferative DR. Further studies of the sequences of cellular events and interactions among growth factors and cytokines in N-PRβ-KO mice will hopefully facilitate the development of effective DR treatments.

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Appendix A. Supplementary Data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2018.04.021.
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