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Endothelial SRF/MRTF ablation causes vascular disease phenotypes in murine retinae

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Retinal vessel homeostasis ensures normal ocular functions. Consequently, retinal hypovascularization and neovascularization, causing a lack and an excess of vessels, respectively, are hallmarks of human retinal pathology. We provide evidence that EC-specific genetic ablation of either the transcription factor SRF or its cofactors MRTF-A and MRTF-B, but not the SRF cofactors ELK1 or ELK4, cause retinal hypovascularization in the postnatal mouse eye. Inducible, EC-specific deficiency of SRF or MRTF-A/MRTF-B during postnatal angiogenesis impaired endothelial tip cell filopodia protrusion, resulting in incomplete formation of the retinal primary vascular plexus, absence of the deep plexi, and persistence of hyaloid vessels. All of these features are typical of human hypovascularization-related vitreoretinopathies, such as familial exudative vitreoretinopathies including Norrie disease. In contrast, conditional EC deletion of Srf in adult murine vessels elicited intraretinal neovascularization that was reminiscent of the age-related human pathologies retinal angiomaticous proliferation and macular telangiectasia. These results indicate that angiogenic homeostasis is ensured by differential stage-specific functions of SRF target gene products in the developing versus the mature retinal vasculature and suggest that the actin-directed MRTF-SRF signaling axis could serve as a therapeutic target in the treatment of human vascular retinal diseases.

Introduction

Angiogenesis, the extension of preexisting vascular networks by EC proliferation and sprouting, is essential for organ development and organ function in health and disease (1, 2). Retinal capillary dysfunction, caused by either impaired vascular growth (hypovascularization) or excessive vascularization (hypervascularization or neoangiogenesis), leads to severe visual impairment and, often, blinding ocular diseases. Ocular hypovascularization diseases encompass familial exudative vitreoretinopathy (FEVR), including Norrie disease (ND) (3). Vessel hyperproliferation, associated with neovascularization (NV), is found in infants with retinopathy of prematurity (ROP) (4), in diabetic retinopathy (DR) (5), and in NV age-related macular degeneration (AMD) (6). NV AMD, which represents approximately 10% of all AMD yet accounts for nearly 90% of all AMD-associated vision losses (6), displays either choroidal NV or, more rarely, intraretinal NV. The latter is found in the manifestations of retinal angiomatous proliferation (RAP) (7) and macular telangiectasia (MacTel) (8–10).

During angiogenesis and EC sprouting, highly polarized ECs, termed tip cells, are positioned at the distal end of vessels. Stalk cells form the sprout base and maintain a continuous, laminized connection to the existing vessel network (11). EC sprouting is highly dynamic and requires rapid rearrangements of the actin cytoskeleton (12). To sense VEGF gradients and other signals, tip cells continuously extend and retract filopodia and constantly compete with stalk cells for the leading position within angiogenic sprouts (13).

Murine postnatal retinal development offers insight into molecular and cellular mechanisms of angiogenesis. At P0, a ring-shaped vessel exists around the optic nerve head in the central retina, whereas peripheral regions lack vasculature. Angiogenic growth extends the retinal vasculature into the periphery, thereby establishing the 2-dimensional primary vessel plexus. In addition, beginning at approximately P8, perpendicular sprouting from the primary plexus leads to the formation of 2 additional vessel layers in the deeper retina, the deep plexi (14). Since global inactivation of genes controlling blood vessel morphogenesis often causes embryonic lethality, the development of inducible genetic models is highly important for the characterization of molecular pathways contributing to retinal angiogenesis in postnatal and adult stages (15).

Serum response factor (SRF) (16), a ubiquitously expressed transcription factor, regulates target gene expression in cell type–selective and signal-specific fashion upon recruitment of different cofactors (17). The SRF cofactors of the ternary complex family (TCFs; encoded by Elk1, Elk3, and Elk4) (18, 19) enable SRF-mediated induction of immediate-early genes (IEGs) upon Ras/MAPK signaling, whereas myocardin-related transcription factors (MRTFs; encoded by Mrtfa and Mrtfb, also known as Mkli1 and Mkli2, respectively) (20–22) function as SRF cofactors responsive to Rho signaling and actin dynamics (23, 24) to regulate cytoskeletal genes (17, 25, 26). SRF binds a conserved target gene cis element, the CArG box [CC(A/T)6GG] (17). Tissue-specific deletion of Srf revealed essential requirements for SRF activity in...
many cell types, including muscle cells, neurons, and ECs (26). Dynamic rearrangements of the actin cytoskeleton are communicated to the genome by actin-directed cytoplasmic release of MRTF proteins, followed by induced nuclear MRTF-SRF interaction and activation of transcriptional responses (17, 23, 27). The contributions of SRF and its cofactors to EC function are poorly understood. In cultured ECs, ELK3 participates in transcriptional responses to hypoxia by regulation of HIF-1α protein stability (28, 29). In vitro cultured ECs revealed a role for SRF in VEGF-mediated angiogenesis, actin polymerization, and EC migration (30). The effects of VEGF on SRF activity were suggested to require both MAPK/Erk and RhoA signaling. In vivo, EC SRF loss-of-function studies were so far only performed during embryogenesis (31, 32), revealing a requirement for SRF in embryonic vascularization. Defects in actin polymerization and in EC intercellular junction formation resulted in embryonic death around E14.5 (31, 32). The genes encoding VEGF-R2 (Kdr, also known as Vegfr2 or Flk1) and VE-cadherin (Cdh5), among others, were identified as SRF target genes stimulated upon VEGF signaling (31).

To overcome embryonic lethality of previously used genetic models (31, 32), we used mice expressing the tamoxifen-inducible CreERT2 under the control of the EC-specific Cdh5 promoter (33), thus permitting temporally controlled deletion of floxed Srf and Mrtf genes in the endothelium. Postnatal EC ablation of either SRF or MRTFs led to severe developmental vessel growth defects, resembling major characteristics of inherited human retinal pathologies such as ROP and FEVR, including ND (3). Interestingly, we found that EC-specific deletion of Srf in adult mice triggered unwarranted intraretinal NV, reminiscent of the human NV AMD subtypes RAP and MacTel (7, 10), as well as the RAP/MacTel animal model deficient for the VLDL receptor (VLDLR; encoded by Vldlr) (9). Moreover, our study investigated EC VEGF signaling mechanisms in vitro and identified MRTF-SRF activation downstream of VEGF. Thus, the present findings offer new insight into the molecular and genetic basis of normal and pathological retinal angiogenesis and suggest cytoskeletal dynamics and the MRTF-SRF module as novel target structures for both diagnosis and treatment of human retinal vascular diseases.

Results

Lethality upon induced EC deletion of Srf during embryonic development. To validate a new genetic model for temporally controlled, induced EC-specific KO of Srf (referred to herein as SrfECKO), Cdh5(PAC)-CreERT2 transgenics (33) were bred with mice carrying a floxed Srf allele (Srfflex1; ref. 34). Tamoxifen was applied successively to pregnant females, and embryos were analyzed at E14.5 or E17.5. Genotypes of the obtained embryos were distributed according to Mendelian ratios. Whereas control embryos appeared normal, freshly isolated SrfECKO embryos showed severe internal hemorrhaging in the head, back, and limb regions (Supplemental Figure 1, A–H; supplemental material available online with this article; doi:10.1172/JCI64201DS1). E17.5 SrfECKO embryos were significantly reduced in weight compared with E14.5 SrfECKO embryos and uninduced controls (Supplemental Figure II). Levels of Srf mRNA and SRF protein in embryonic limb buds isolated from SrfECKO animals were significantly reduced compared with control littermates (Supplemental Figure 1, J–L). These timed Srf deletion experiments demonstrated that our novel tamoxifen-inducible, EC-specific genetic system fully reproduced phenotypes during embryogenesis, which were previously reported for noninducible gene targeting using different Cre drivers and independently generated floxed Srf alleles (31, 32).

Postnatal EC Srf deletion causes impaired retinal angiogenesis. Taking advantage of the tamoxifen-inducible, temporally controlled gene targeting strategy, we chose the retina to study the effects of EC-specific Srf deletion on postnatal angiogenesis. Newborn pups were successively injected with tamoxifen at P1–P4, and vascularization was analyzed on retinal flat-mounts. As early as P6, isoleciton B4 (ILB4) staining of the retinal primary plexus showed a significant 29% reduction in the area covered by blood vessels and a significant 35% decrease in the number of branch points in SrfECKO animals compared with littermate controls (Figure 1, A–C). Abnormal tip cell morphology was evident in SrfECKO retinae, with mutant sprouts displaying short shafts as well as fewer and shorter distal filopodia (Figure 1, A and D–F). These data suggest an essential role of SRF in postnatal angiogenesis, especially regarding progression of the retinal angiogenic front, which depends upon actin-mediated tip cell filopodial dynamics (11).

To monitor Cre recombination, we combined the SrfECKO model with double-fluorescent mTmG Cre reporter alleles (35), which switch expression from membrane-targeted Tomato protein (red) to GFP (green) upon Cre activation. Newborn mice were injected at P1–P4, and retinal flat-mounts were analyzed at P8 (Figure 1G). In the absence of the reporter, no specific signals were detected in either the red or the green channel. In retinai of un.injected Srfflex1/flex1 Cdh5(PAC)-CreERT2mTmG mice, all retinal cells showed ubiquitous red fluorescence, whereas no specific signal in the green channel was observed. This proved that the system was not leaky, and no unspecific recombination occurred in the absence of tamoxifen. In retinai of tamoxifen-injected Srfflex1/flex1 Cdh5(PAC)-CreERT2mTmG control animals, ECs changed their color to green, highlighting efficient Cre recombination in the endothelium and the expected normal vascularization of the retina for age P8. In contrast, vessel outgrowth in tamoxifen-injected Srfflex1/flex1 Cdh5(PAC)-CreERT2mTmG mice was strongly impaired, as evidenced by the abnormal position of the angiogenic front formed by green fluorescent, recombined ECs (Figure 1G).

Development of deep vascular plexi in the retina requires SRF function. SrfECKO animals at age P10 contained large avascular zones in the peripheral retina (Figure 2A), indicative of impaired development of the primary plexus until late stages of retinal vascularization. In control littermates, the primary plexus had reached the retinal periphery and started to extend into deeper retinal layers to form the tertiary and secondary capillary networks of the deep plexi (Figure 2, A, C, E, and F). In contrast, SrfECKO animals showed a 29% reduction in retinal area covered by blood vessels (Figure 2B). Interestingly, at the angiogenic front, large clusters of tip cells without filopodia formed (referred to as distal microaneurysms), which were never observed in control animals (Figure 2C). Abnormal accumulation of ECs within these microaneurysms was evidenced by EM (Figure 2D). Despite the abnormal accumulation of ECs within distal microaneurysms, these abnormal vessel structures were covered by pericytes (Figure 2D). Additionally, deep plexi were not established in SrfECKO animals (most likely as a consequence of abnormal sprouting in the primary plexus), which was further evidenced by EM, whereas control retinae revealed normal deep plexi vessel structures (Figure 2, F and G). Notably, we did not find a reduction in EC proliferation in SrfECKO retinae compared with controls (data not shown).
Figure 1
EC depletion of SRF impairs angiogenesis in P6 murine retinal development. (A) ILB4-stained retinal flat-mounts of P6 control and SrfiECKO mice. Top: Progression of angiogenic front. Red arrow indicates recessed angiogenic front of the primary plexus in SrfiECKO retinae. Images are composites (see Methods). Middle: Vessel density between artery (A) and vein (V). Bottom: Sprout morphology. White arrows indicate filopodia; red asterisks highlight abnormal morphologies of SrfiECKO tip cells. (B) Quantitation of retinal area covered by blood vessels (radial outgrowth), expressed as percentage of control. n = 6 retinae. (C and D) Quantitation of (C) branch points in a field of view (A, middle, white boxes) and (D) abnormal sprouts. n = 4 retinae. (E and F) Quantitation of (E) filopodial number per sprout and (F) mean length of individual filopodia. n = 30 sprouts (control); 39 sprouts (SrfiECKO). (G) Retinal flat-mounts of P8 mice lacking the double-fluorescent mTmG Cre reporter, SrfiECKO Cdh5(PAC)-CreERT2 mTmG mice (SrfiECKOTG), and control SrfiECKO Cdh5(PAC)-CreERT2 mTmG mice. Shown are red fluorescent channel (mTomato) and GFP (mGFP) signals of the same retinae, the latter being obtained upon CreERT2 activation by intragastric tamoxifen injection (+ tamox). Red arrow indicates recessed angiogenic front of the primary plexus in SrfiECKO Cdh5(PAC)-CreERT2 mTmG retinae. Images are composites (see Methods). Scale bars: 1 mm (A, top, and G); 50 μm (A, middle); 15 μm (A, bottom). *P < 0.05, **P < 0.01, ***P < 0.001 vs. respective control.
To quantitate SRF-mediated gene expression, ECs were enriched from retinal tissue of tamoxifen-treated control and Srf\textsuperscript{IECKO} P10 animals, using anti-CD31–coated magnetic beads. mRNA levels of Srf, as well as SRF target genes Kdr, Cdh5, and Actb (also known as β-actin), were significantly decreased in Srf\textsuperscript{IECKO} retinae (Figure 2H). Correspondingly, protein levels of SRF and VEGF-R2 in whole retinal tissue were reduced by 60% and 51%, respectively (Figure 2, I and J).

Persistence of vascular defects in Srf\textsuperscript{IECKO} retinae after postnatal deletion of Srf. Postnatal retinal angiogenesis of WT mice reaches completion by P25 (14). At advanced ages (P15–P31), retinal irregularities remained in Srf\textsuperscript{IECKO} animals: in vivo imaging using confocal scanning laser ophthalmoscopy (SLO), particularly in fluorescence angiography (FLA) mode, revealed persistent distal microaneurysms of different sizes (Figure 3A). Additionally, no deep plexi were detectable in FLA recordings in retinae of these older Srf\textsuperscript{IECKO} mice, and no significant leakage of fluorescent dye out of microaneurysms was observed. Distal microaneurysms were visualized at the persistently recessed angiogenic fronts upon ILB4 staining of retinal flat-mounts, performed after completion of the in vivo imaging (Figure 3B). Further vascular abnormalities in Srf\textsuperscript{IECKO} retinae were found in H&E stainings of ocular paraffin sections. Srf\textsuperscript{IECKO} retinae displayed distal microaneurysms positioned aberrantly in the inner nuclear layer (INL; Figure 3C). Prior to histological preparation, the identical, persistent abnormal blood vessels were detected in vivo via optical coherence tomography (OCT; Figure 3C). The punctate double reflectance patterns typical of vessels (dark zones at the top and bottom of each vessel) were clearly visible.

Figure 2
Avascular zones, distal microaneurysms, and lack of deep plexi in Srf\textsuperscript{IECKO} retinae at P10. (A) ILB4-stained retinal flat-mounts. Red arrow indicates recessed angiogenic front in the Srf\textsuperscript{IECKO} primary plexus. Images are composites (see Methods). (B) Radial outgrowth, expressed as percent of control. n = 19 retinae (control); 9 retinae (Srf\textsuperscript{IECKO}). (C) Higher magnification of ILB4-stained retinal flat-mounts. White arrows indicate microaneurysms in Srf\textsuperscript{IECKO} retinae. (D) EM image of blood vessels near the inner limiting membrane (ILM) to visualize the primary plexus. P, pericyte; L, lumen; BL, basal lamina. (E and F) ILB4-stained retinal capillaries of (E) the primary plexus and (F) deep plexi, which revealed complete absence of deeper capillaries in Srf\textsuperscript{IECKO} retinae. (G) EM image visualizing deep plexi, OPL, outer plexiform layer. (H) Semiquantitative RT-PCR of mRNA expression in purified ECs of P10 retinae. n = 4 (Srf); 3 (Kdr and Actb); 5 (Cdh5). mRNA levels were normalized to Gapdh and expressed as percent of control. (I) Western blot analysis of 2 representative pairs of control and Srf\textsuperscript{IECKO} P10 whole retinal tissue. (J) Quantitation of Western blot. SRF (n = 5) and VEGF-R2 (n = 4) levels were normalized to GAPDH and expressed as percent of control. Scale bars: 1 mm (A), 100 μm (C), 2 μm (D and G, left), 50 μm (E and F), 5 μm (G, right). *P < 0.05, **P < 0.01 vs. respective control.
SRF deficiency leads to defective hyaloid vessel regression. The presence of avascular zones in the periphery of SrfiECKO retinae led us to test whether hyaloid vessels persist, as this vessel system, normally regressing during postnatal development, is known to be retained when the vascular network is compromised (36). Interestingly, in 64% (7 of 11) of SrfiECKO retinae, persisting hyaloid vessels were seen on flat-mounts (Figure 4A). With the exception of 1 of the 10 animals analyzed, no hyaloid vessels were detected on control retinal flat-mounts (Figure 4A). Retinal flat-mounts of control animals revealed the existence of all capillary beds and vascularization into the periphery, whereas the presence of hyaloid vessels in SrfiECKO retinae coincided with large avascular zones, microaneurysms, and the absence of deeper capillary beds (Figure 4A). In vivo, the course of hyaloid vessels retained in the SrfiECKO retinae could be followed by SLO angiography examining different focal planes of the same eye: hyaloids rose up from their origin at the retinal center and moved toward the back of the lens, then down toward the avascular “target zone” in the periphery (Figure 4B). The functional status of these vessels was verified by visible blood flow and pulsations of the whole vessel, as detected by SLO in vivo (data not shown).

Molecular and structural defects in SrfiECKO retinal blood vessels. Retinal blood vessels grow along an underlying astrocytic network, which presents a gradient of matrix-bound VEGF (36). We analyzed the structure of this astrocytic network by glial fibrillary acidic protein (GFAP) staining of P6 retinal flat-mounts. In both control and SrfiECKO retinae, astrocytes had reached the periphery and formed elaborate networks (Figure 5A). Control EC sprouts followed the preceding astrocytic network in close overlap, whereas the overlap between distal vessels and astrocytes was markedly reduced in SrfiECKO retinae (Figure 5A). Next, we tested whether the mutant phenotype was caused by sprout retraction and vessel collapse. Both events would generate empty collagen IV sleeves, which are no longer associated with ECs (37, 38). In control retinae, the overlap between EC ILB4 and collagen IV patterns was almost complete (Figure 5B). In SrfiECKO flat-mounts, we focused on the distal microaneurysms at the angiogenic front, which revealed a complete absence of empty collagen sleeves (Figure 5B), indicating that retraction of sprouts or vessels was not the reason for the observed vascular defects.

SRF deficiency in neurons of the forebrain resulted in increased Ser3 phosphorylation of the actin-severing protein cofilin (39). To analyze whether this indicator of imbalance in actin dynamics was also evident in SRF-deficient retinal ECs, we performed phosphorylated cofilin (P-cofilin) staining on retinal flat-mounts. Low basal P-cofilin staining was detected on control retinal vessels, which was slightly stronger than the neuronal signal in the ganglion cell layer (Figure 5C). In contrast, strong P-cofilin staining was found in distal microaneurysms of SRF-deficient retina (Figure 5C). In addition, we observed an increased P-cofilin signal in whole retinal tissue (Figure 5, D and E).

Adult SrfiECKO eyes display retinal NV reminiscent of the Vldlr−/− phenotype. To examine whether SRF plays a role in the maintenance of the fully established mature retinal vasculature of adult mice, Srf deletion was induced at 4–6 weeks of age, well after the completion of all developmental growth and maturation processes. Resulting ocular phenotypes were analyzed by SLO, OCT, and staining on paraffin sections. Covering a time span of 3–26 weeks after adult-stage tamoxifen injection, 82% of SrfiECKO animals (n = 11) displayed multiple ecotropic intraretinal NVs that were not observed in control littermates, as shown by in vivo SLO followed by H&E staining after paraffin sectioning (Figure 6, A–D). The abnormal NV structures were further characterized in OCT (Figure 6A). Collectively, these analyses revealed that NV retinal capillaries originated from deep plexi and connected to the retinal pigment epithelium (RPE). At the site of connection, RPE cells appeared to engulf the NV. Additionally, local distortion of outer nuclear layer...
We therefore used 6- to 8-month-old Srf-depleted mECs (Figure 7E). The mRNA expression level (Figure 7E). However, most interestingly, mRNA expression to be under SRF transcriptional control, but rather posttranslational control (39) — was expectedly unchanged at the mRNA

in the SRF target gene, as assayed by semiquantitative RT-PCR. The obtained decrease in Srf mRNA correlated with a decrease in the SRF target gene Actb, as assessed by semiquantitative RT-PCR. Cfl1 (encoding cofilin) — which was previously found not to be under SRF transcriptional control, but rather posttranslational control (39) — was expectedly unchanged at the mRNA level (Figure 7E). However, most interestingly, mRNA expression of Thbs1 (encoding the VLDLR ligand thrombospondin-1) was significantly decreased in SRF-depleted mECs (Figure 7E). The thrombospondin promoter contains a perfect SRF-binding CARG-box of CCTTATTTGG sequence at promoter coordinate ~1.2 kb (45). Using ChIP, we showed that SRF bound to this element in the Thbs1 gene; Pak1, a CARG box–negative locus, served as normalization control, and Actb served as positive reference (Figure 7F).

Taken together, these results demonstrated that EC-specific ablation of SRF in the adult retinal endothelium leads to NV lesions associated with reduced photoreceptor viability. The congruence of this phenotype with Vldlr−/− mice, together with our finding that the VLDLR ligand thrombospondin-1 was under transcriptional control, suggests that impairment of the antiproliferative VLDLR-thrombospondin pathway contributes to formation of intraretinal NV lesions in adult SrfiECKO retinas.

**MRTF, but not TCF, cofactors are essential for retinal angiogenesis.** To identify cofactors recruited by SRF in mediating retinal vascular development, we performed KO studies on 4 of the 5 known genes encoding TCFs and MRTFs. We first took advantage of a newly generated, constitutive Elk1−/−Elk4−/− double-KO mouse model (46). In the retinal vasculature, radial outgrowth was found to be unaffected in P6 and P10 Elk1−/−Elk4−/− versus controls, which included Elk1−/−Elk4+/* and Elk1−/−Elk4+/* genotypes (Figure 8, A, B, D, and E). The angiogenic front of Elk1−/−Elk4−/* retinae at P6 appeared normal: tip cells extended numerous filopodia indistinguishable from those of controls (Figure 8C). Moreover, deep plexi were visible in both control and Elk1−/−Elk4−/* retinae at P10 (Figure 8, F and G). SRF and P-cofilin protein levels were unchanged in Elk1−/−Elk4−/* retinae (Supplemental Figure 2B). These data demonstrated that the TCF-type SRF cofactors ELK1 and ELK4 were not required for vascularization of the retina.

To determine whether the SRF cofactors MRTF-A and MRTF-B play a role in SRF-dependent retinal angiogenesis instead, we generated mice with Cre-inducible EC-specific KO of MRTF-B on a constitutive Mrf4-KO background (referred to herein as Mrf4−/−Mrtfa+/−MrtfbiECKO mice). Mrf4−/−MrtfbiECKO mice showed retinal blood vessel defects highly reminiscent of those of SrfiECKO mice. This congruence in phenotype with SrfiECKO mice was evidenced by the Mrf4−/−
Mrtfb^{ECKO} animals’ reductions in radial outgrowth at P6 and P10, number of branching points, formation of abnormal sprouts at P6, and formation of distal microaneurysms as well as by their absence of deep plexi at P10 (Figure 9, A–D and G–J). The reductions in filopodial number and length were comparable in Mrtfa^{–/–}Mrtfbi^{ECKO} retinae (Figure 9, E and F) and in Srfi^{ECKO} retinae. Importantly, in Mrtfa^{–/–}Mrtfbi^{ECKO} retinae, SRF protein levels were unchanged, whereas P-cofilin was markedly increased (Supplemental Figure 2, A and B). KO of Mrtfa and Mrtfb individually caused partial impairment of angiogenic front progression in each genotype. Detailed analysis of radial outgrowth in P8 retinae suggested a gene-dosage effect: radial outgrowth decreased successively with decreasing WT allele composition for Mrtfa and Mrtfb (Figure 9K and Supplemental Figure 3). Taken together, the highly overlapping phenotypic characteristics of Mrtfa^{–/–}Mrtfbi^{ECKO} and Srfi^{ECKO} mice were strongly suggestive of MRTFs being the relevant EC SRF cofactors in vivo ensuring appropriate retinal angiogenesis.

VEGF-A induces cytoplasm-to-nucleus translocation of MRTF-A and SRF target gene expression. The above genetic evidence for essential contributions of SRF and MRTF to retinal angiogenesis led us to investigate signaling cascades activating SRF in mECs. Interestingly, both serum and VEGF-A treatment of mECs activated MRTF-A to translocate from the cytoplasm to the nucleus (Figure 10A). The sensitivity of this effect to latrunculin B (Figure 10A) suggested involvement of G-actin in retaining MRTF-A within the cytoplasm (24). Similarly, cytochalasin D and jasplakinolide, 2 F-actin–modulating substances known to liberate MRTF from G-actin (24, 27), induced nuclear MRTF-A translocation in mECs (Supplemental Figure 4). To further probe into the functionality of VEGF-induced MRTF activation, we used luciferase-based transient reporter gene assays, which harbored promoter sequences that either did [(TSm)2] or did not [(Tmm)2] contain an intact SRF binding site (Figure 10B). In transfected human retinal microvascular ECs (HRMECs), VEGF-stimulated (TSm)2 reporter activity was enhanced by cotransfected MRTF expression vectors, an effect that was dependent on the presence of intact SRF binding sites (Figure 10C). Finally, VEGF-A treatment of mECs increased mRNA levels of the endogenous SRF target genes Flt1 (also known as Vegfr1), Kdr, Srf, Actb, and Fos (Figure 10D). These results argue for VEGF-induced cooperation of SRF and MRTF in ECs that leads to expression of essential proangiogenic genes. We therefore propose that VEGF-induced MRTF translocation to the nucleus activates in ECs expression of SRF target genes with cytoskeletal functions, which in turn is essential for motile activities of tip cells to ensure appropriate vascularization of the postnatal retina (Figure 10E).

Discussion

The genetic study presented here identifies the murine transcription factor SRF to be essential for EC function, both during postnatal
development of the retinal vasculature and in maintenance of adult capillary homeostasis. Most importantly, whereas EC-specific SRF ablation at postnatal stages caused hypovascularization of the primary plexus and the absence of deep retinal plexi, EC deletion of SRF at adult stages led to intraretinal NV. Both of these opposing murine phenotypes, elicited upon stage-specific SRF depletion, resembled blinding human retinopathies, i.e., developmental hypovascularization found in early-onset FEVR (including ND) (3, 47) versus the adult-onset intraretinal NV AMD subtypes RAP and MacTel (7, 8).

Altered retinal vessel were the earliest phenotypic effects we detected upon induction of postnatal or adult EC-specific SRF depletion. However, besides the retinal phenotype, we observed some behavioral abnormalities. We attribute these behavioral effects to compromised functions of cerebral microvessels, which do not appear to exert influences on the retinal phenotype described here and will be characterized separately.

Postnatal defects in retinal development upon EC-specific Srf deletion included retarded progression of the angiogenic front and impaired tip cell filopodial protrusion, which generated avascular zones and distal microaneurysms in the primary plexus and, consequently, a complete absence of deep plexi. This was associated with persistence of the hyaloid vasculature. EC-specific deletion of the SRF cofactors MRTF-A/MRTF-B, but not ELK1/ELK4, led to similar phenotypes with respect to postnatal retinal vascularization, thereby providing clear genetic evidence for an essential functional interaction of SRF and MRTFs in directing EC activities. Since actin dynamics provide a driving force for such motility functions, including tip cell filopodial activity (11, 12), our observation that VEGF-A signaling elicits actin-dependent translocation of cytoplasmic MRTF into the nucleus represents a new molecular mechanism by which angiogenic signals induce motile functions of ECs.

The phenotype upon postnatal ablation of MRTF-SRF signaling described herein showed strong similarities to human pathologies and murine model systems of FEVR, including ND (47, 48). Since identified genetic lesions eliciting FEVR and ND fall into the Wnt signaling pathway, including mutant alleles for Ndp, Fzd4, Lrp5, and Tip41L2 (3), our data suggest the existence of cross-talk between Wnt and actin signaling in guaranteeing proper EC function during retinal angiogenesis.

In contrast to postnatal SRF depletion, Srf deletion in adult ECs elicited intraretinal NV. This phenotype was characterized by neoangiogenic activities originating from intraretinal deep capillaries and causing non-uniformly distributed focal lesions. The latter displayed distortions in INL and ONL layering, local thinning of the ONL, disruption of photoreceptors associated with reduced expression of Opn1 and Rho, and rupture of the RPE with RPE cells enveloping the NV structures. Collectively, these abnormalities represent characteristic features of the human NV AMD subtypes RAP (7) and MacTel (8–10). Very similar phenotypic features were displayed by mice deficient for VLDLR, a component of reelin signaling exerting inhibitory effects on EC proliferation (9, 40–43). In addition to binding reelin, VLDLR also binds the ligand thrombospondin-1 (49, 50). The glycoprotein thrombos-
pondin is an extracellular matrix component known to inhibit EC proliferation (51, 52). Interestingly, the THBS1 gene has been previously suggested to be under transcriptional control of SRF in human fibroblasts (45). Using siRNA and ChIP approaches, we here provided evidence suggesting Thbs1 expression is under control of SRF in cultured mECs (Figure 7). We therefore hypothesize that SRF-controlled expression of thrombospondin is required in adult retinal vessels to inhibit pathological EC proliferation, thereby providing an explanation for both the NV phenotype we observed upon adult-stage SRF depletion and the hypothesized impairment of VLDLR signaling by associated thrombospondin deficiency are fully consistent with the defects displayed by Vldlr−/− mice.

Both RAP and MacTel show age-dependent onset in human patients. Since SRF activities are known to deteriorate in some aging cells (53), it will be of great interest to explore potential contributions of age-dependent SRF activity changes in the aging retinal endothelium of human AMD patients. Furthermore, our work might suggest modulation of EC SRF activity, as well as elevation of thrombospondin activity, as therapeutic options for the treatment of some NV AMD pathologies, such as RAP and MacTel.

The MRTF/SRF transcriptional feedback circuit, which links cytoplasmic actin polymerization with nuclear gene regulation, represents one of the best-understood mechanisms for genomic sensing of cytoskeletal dynamics (24, 27). Associated cellular functions include dynamic changes regarding cell polarity, cell shape, migration, and guided growth, all of which are essential to EC biology. Mechanistically, our conclusion that VEGF-A signaling activates G-actin–dependent MRTF-A translocation from the cytoplasm to the nucleus (Figure 10E) positions the MRTF-SRF module at the center of angiogenic regulation.

In conclusion, our genetic study using conditional depletion of SRF and MRTFs provides the first in vivo identification of any function for these transcription factors in the retinal endothelium of living animals. The phenotypic defects we describe in postnatal and adult retinal angiogenesis provide new insight into both the molecular developmental biology of retinal angiogenesis and the etiology of selected retinal pathologies. The findings reported here may help to define therapeutic approaches for the treatment of some human diseases related to abnormal retinal vascularization.
double-fluorescent reporter (35) were crossed into the Srf flex1Cdh5(PAC)-CreERT2 background. Geno-
typing was performed by PCR of tail biopsies (see Supplemental Table 1 for primer sequences).

Tamoxifen injections. For embryonic Cre acti-
vation, timed pregnant females were injected
intraperitoneally on E10.5, E12.5, and E14.5 with
2 mg tamoxifen (dissolved in sunflower oil), and
embryos were analyzed on E17.5. In a second series
of experiments, females were injected on E10.5,
E11.5, and E12.5, and embryos were analyzed on
E14.5. For postnatal Cre activation, newborn pups
were injected intragastrically on P1–P4 with 0.05
mg tamoxifen and analyzed on P6, P8, and P10
and at advanced age (P15–P31). For adult analysis,
4- to 6-week-old mice were injected intra peri-
toneally with 2 mg tamoxifen on 5 consecutive
days and analyzed at different time points, ranging
from 3 to 26 weeks after the last injection.

Antibody staining of retinal flat-mounts
Eyes were isolated and fixed in 4% PFA for 2 hours (54). After
washing 2 × for 5 minutes with PBS, retinae were
incubated and incubated in blocking buffer (1%
BSA, 0.3% Triton-X, and PBS) overnight at 4°C.

Methods

Animals. To generate KO mice with EC-specific SRF deletion (Srf GCKD mice), female mice homozygous for the floxed Srf flex1 allele (34) were crossed with Srf flex1/WT Cdh5(PAC)-CreER T2 male mice, which carry the Cdh5(PAC)-CreER T2 transgene (33). Srf flex1/WT, Srf flex1/flex1, and Srf flex1/WT Cdh5(PAC)-CreER T2 mice were used as controls. Double-KO Elk1–/– Elk4–/– animals (provided by R. Treisman, Cancer Research UK, London, United Kingdom) were bred as described previously (46). Double-KO mice with EC-specific MRTF-B deletion (Mrtfa–/– Mrtfbi GCKD mice) were generated by breeding the Cdh5(PAC)-CreER T2 transgene into Mrtfa–/– Mrtfbi fl/fl animals (25). Matings of Mrtfa–/– Mrtfbi fl/fl females with Mrtfa–/– Mrtfbi fl/fl Cdh5(PAC)-
CreER T2 males yielded the double-KO genotype Mrtfa–/– Mrtfbi fl/fl Cdh5(PAC)-
CreER T2, and conditional deletion of the Mrtfbi fl/fl alleles was induced by tamoxifen injection. To monitor Cre activity, mice carrying the mTmG
double-fluorescent reporter (35) were crossed into the Srf flex1Cdh5(PAC)-CreER T2 background. Geno-
typing was performed by PCR of tail biopsies (see Supplemental Table 1 for primer sequences).

Figure 8
The TCF-type SRF cofactors ELK1 and ELK4 are not essential for normal retinal angiogen-
esis. (A) ILB4 staining on P6 control and Elk1–/– Elk4–/– retinae. Images are composites (see Methods). (B) Quantitation of retinal area cov-
ered by blood vessels (percent radial outgrowth) at P6. n = 39 retinae (control); 20 retinae (Elk1–/–
Elk4–/–). (C) Representative images of angiogenic fronts at P6. (D) ILB4 staining on P10 retinae. Images are composites (see Methods). (E) Per-
cent radial outgrowth at P10. n = 10 retinae per group. (F and G) ILB4-stained retinal capillaries of (F) the primary plexus and (G) deep plexi at
P10. Scale bars: 1 mm (A and D); 30 μm (C); 50 μm (F and G).

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P10. Scale bars: 1 mm (A and D); 30 μm (C); 50 μm (F and G).

Antibody staining of retinal flat-mounts. Eyes were
isolated and fixed in 4% PFA for 2 hours (54). After
washing 2× for 5 minutes with PBS, retinae were
dissected and incubated in blocking buffer (1%
BSA, 0.3% Triton-X, and PBS) overnight at 4°C. After
washing 5× for 5 minutes with PBS, retinae were treated 3× for 20 minutes with Phlec buffer
(1 mM CaCl2, 1 mM MgCl2, 0.1 mM MnCl2, 1% Triton-
X, and PBS, pH 6.8). Incubation with primary
antibodies was done at 4°C in Phlec buffer overnight. After washing 3× for
20 minutes with half blocking solution (0.5% BSA, 0.15% Triton-X, and PBS), retinae were incubated with secondary antibodies (2 hours at room
temperature). After washing 3× for 20 minutes in half blocking solution, retinae were flat-mounted on coverslides and embedded in Mowiol.

Primary antibodies were as follows: ILB4 from Griffonia simplicifolia (1:25 dilution; Sigma-Aldrich); GFAP (1:100 dilution; DAKO); collagen IV (1:40
dilution; AbD Serotec); P-cofilin (1:100 dilution; Cell Signaling). Second-
ary antibodies (all from Molecular Probes) were as follows: streptavidin–
Alexa Fluor 488 (1:100 dilution), anti-rabbit Alexa Fluor 546 (1:200 dilu-
tion), anti-rat Alexa Fluor 546 (1:200 dilution).

Analysis of morphometric parameters of retinal flat-mounts. Radial outgrowth
was determined as the ratio of blood vessel-covered retinal area to total
retinal area, and expressed as a percentage. Branch points were counted
Figure 9
MRTF-A and MRTF-B are essential for retinal angiogenesis. (A) ILB4-staining on retinal flat-mounts of P6 control and Mrtfa<sup>+</sup>/Mrtfb<sup>ECKO</sup> mice. Top: Progression of angiogenic front. Red arrow indicates recessed angiogenic front of primary plexus. Images are composites (see Methods). Middle: Vessel density between artery (A) and vein (V). Bottom: Sprout morphology. White arrows indicate filopodia; red asterisks indicate abnormal morphologies of Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup> tip cells. (B–D) Quantitation of (B) percent radial outgrowth; (C) relative branch points in field of view (boxed regions in A, middle); and (D) abnormal sprouts. n = 4 (control); 6 (Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup>). (E and F) Quantitation of (E) filopodia number per sprout and (F) filopodia mean length. n = 54 sprouts (control); 71 sprouts (Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup>). (G) ILB4-stained retinal flat-mounts of P10 control and Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup> retinae. Red arrow indicates recessed Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup> angiogenic front. Images are composites (see Methods). (H) Percent radial outgrowth. n = 9 (control); 5 (Mrtfa<sup>−/−</sup>Mrtfb<sup>ECKO</sup>). (I and J) Higher-magnification views of (I) primary plexus and (J) deep plexi. (K) Percent radial outgrowth in P8 retinae, including all genotypes resulting from our mating scheme. n = 8 (Mrtfa<sup>+</sup>Mrtfb<sup>+/+</sup>); 7 (Mrtfa<sup>−/−</sup>Mrtfb<sup>+/+</sup>); 5 (Mrtfa<sup>−/−</sup>Mrtfb<sup>iECKO</sup> and Mrtfa<sup>−/−</sup>Mrtfb<sup>iECKO</sup>); 6 (Mrtfa<sup>−/−</sup>Mrtfb<sup>iECKO</sup> and Mrtfb<sup>iECKO</sup>). See Supplemental Figure 3 for statistical comparisons. Scale bars: 1 mm (A, top, and G); 50 μm (A, middle, and I and J); 10 μm (A, bottom). *P < 0.05, **P < 0.01, ***P < 0.001 vs. respective control.
VEGF-A activates nuclear translocation of MRTF-A.

(A) mECs were stimulated with serum, VEGF-A, or VEGF-A in the presence of latrunculin B (LatB) and stained for nuclei (DAPI; blue), MRTF-A (green), and F-actin (phalloidin; red).

(B) Scheme of (TSm)$_2$ and (Tmm)$_2$ luciferase reporter constructs, which contain 2 tandem copies of the c-Fos SRE upstream of the thymidine kinase basal promoter sequence (tk120; −120 to +1), able to drive luciferase cDNA expression.

(C) Relative luciferase activity in reporter-transfected HRMECs, with or without cotransfection of MRTF-A expression vectors and with or without VEGF-A stimulation.

(D) Semiquantitative RT-PCR for genomic candidate mRNA expression in mECs upon VEGF-A treatment, expressed as percent untreated control. $n = 4$ (Flt1, Srf, and Actb); 3 (Kdr and Fos).

(E) VEGF signaling leads to activation of the actin-MRTF-SRF axis. Note that interaction of MRTF and TCF cofactors (ELK1 and ELK4) with SRF is mutually exclusive. Scale bars: 10 μm (A). *$P < 0.05$, **$P < 0.01$ vs. respective control.
per field of view in an area between arteries and veins. Abnormal tip cells were defined as exhibiting stubby shafts with fewer and shorter filopodia. Abnormal sprouts along the angiogenic front were counted and expressed as a percentage of total sprouts. Numbers of filopodia per single sprout were averaged for control and mutant retinae.

**Purification of ECs.** P10 retinae were dissected and homogenized in 0.2% collagenase buffer at 37°C for 45 minutes. Cells were triturated using a 20-gauge syringe, passed through a 40-μm cell strainer, pelleted at 254 g for 5 minutes, and resuspended in serum-free HUVEC medium. ECs were bound to magnetic Dynabeads (Invitrogen) coated with anti-CD31 (BD Biosciences — Pharmingen) for 30 minutes at 4°C and separated magnetically from non-ECs.

**RNA isolation, cDNA synthesis, and semiquantitative RT-PCR.** Tissues were lysed for RNA isolation (Qiagen, RNeasy), cDNA was synthesized using random hexamers, and RT-PCR analysis was performed using specific primers (Purimex) and SYBR green technology (ABI Prism 7000 cycler) (55). See Supplemental Table 1 for primer sequences and amplification protocols.

**Western blotting.** Retinal tissue was lysed in Iyer buffer (56), and protein content was determined (Bradford reagent). Proteins were separated by SDS-PAGE in 4%–15% gradient gels. Electrophoresis was at 4°C for 2 hours (100 V, 400 mA). Membranes were blocked in 10% milk powder (1 hour at room temperature). Primary antibody incubation was overnight at 4°C. After 3x washes with TST (Tris Saline Tween), membranes were incubated in secondary antibodies (1 hour at room temperature). Primary antibodies were as follows: GAPDH (1:20,000 dilution; Hytest Ltd), VEGF-R2 (1:1,000 dilution; Cell Signaling), SRF (2C5, undiluted; ref. 57), P-cofilin (1:500 dilution; Cell Signaling). Secondary antibodies (all 1:10,000 dilution; all GE Healthcare) were anti-mouse, anti-rabbit, and anti-rat HRP. Band intensities were quantified densitometrically and normalized to GAPDH.

**SLO and OCT.** For SLO and OCT, mice were anesthetized by subcutaneous injection of ketamine (66.7 mg/kg) and xylazine (11.7 mg/kg). After anesthesia, pupils were dilated with tropicamide eye drops (Mydriaticum Stulln; all GE Healthcare) were anti-mouse, anti-rabbit, and anti-rat HRP. Band intensities were quantified densitometrically and normalized to GAPDH.

**Histological H&E and antibody staining.** Sections were embedded in Mowiol. For antibody staining, paraffin sections were obtained with a microtome (Ultracut; Leica) was used to cut semithin (1 μm) and ultrathin (50 nm) sections in the long wavelength range. The fundus images presented here were recorded with the green laser (514 nm). Angiography was performed with 2 different dyes administered simultaneously: FLA at 488 nm and 514 nm) in the short wavelength range and 2 infrared diode (808 nm) lasers. SLO imaging (58) was conducted with a Heidelberg Retina Angiograph (HRA I) equipped with an argon laser featuring 2 wavelengths (488 nm and 514 nm) in the short wavelength range and 2 infrared diode lasers (795 nm and 830 nm) in the long wavelength range. The fundus images presented here were recorded with the green laser (514 nm). Angiography was performed with 2 different dyes administered simultaneously: FLA at 488 nm (barrier filter 500 nm) and indocyanine green angiography (ICG) at 795 nm (barrier filter 800 nm). Dyes were given via subcutaneous injection of 75 mg/kg body weight fluorescein-Na (Tübingen University pharmacy), and 50 mg/kg body weight ICG (ICG-Pulsion; Pulsion Medical Systems AG). OCT was recorded with a Heidelberg Engineering Spectralis system with minor adaptations to the mouse eye as described previously (59).

**Histological H&E and antibody staining.** Eyes were fixed overnight at 4°C in Davidson fixative (6% formaldehyde, 32% EtOH, 11% acetic acid, 5% sucrose in PBS). 4-μm sections of paraffin-embedded eyes were mounted on SuperFrost Plus slides and stained with H&E followed by dehydration and mounting in Entellan. For antibody staining, paraffin sections were blocked in 2% BSA/PBS-T for 1 hour at room temperature, followed by incubation with primary antibody overnight at 4°C. After washing 3x for 15 minutes with PBS-T, secondary antibodies were applied for 1 hour at room temperature. Cell nuclei were counterstained with DAPI. For microscopic analysis, sections were embedded in Mowiol.

**Cell culture, transient transfection, and luciferase assay.** Immortalized mECs (44) and HMMECs (Cell Systems) were cultured as recommended. Transient transfections used Promofectin (PromoCell). mECs were starved overnight and stimulated for 1 hour with either 15% FCS or 100 ng/ml VEGF-A. Pretreatment with latrunculin B was for 30 minutes. (TSm), and (Tmm); reporter plasmids (in which “m” denotes a mutated binding site) contained 2 tandem copies of the c-Fos serum response element (SRE) upstream of the thymidine kinase basal promoter sequence (~120 to +1), able to drive luciferase cDNA expression. The WT SRE contains binding sites for the transcription factors TCF, SRF, and AP-1. For transfection of siRNAs directed against Srf (siSrf797; Purimex) (60), mECs were seeded into 6-cm dishes with penicillin/streptomycin-free medium 1 day before transfection. For transfection (RNAiMAX reagent; Invitrogen) with 166 pmol siRNA per dish, cells had reached approximately 60% confluence. Transfection was stopped after 5 hours, and RNA was isolated 2 days later. Extracts for protein analysis were generated 2 and 3 days after transfection.

**ChIP.** For ChIP analysis (61), mECs were serum starved for 18 hours before stimulation with 15% FCS for 1 hour. See Supplemental Table 1 for primer sequences and amplification protocols. Anti-SRF ChIP signals of CARG boxes in ArhG and Thbs1 were normalized to either IgG-containing or no-antibody control reactions and expressed relative to Pabk (CarG box negative control locus).

**EM.** For EM, retinae were fixed 4 hours in 2.5% glutaraldehyde (Paesel + Lorei GmbH) in 0.1 M cacodylate buffer (CB; pH 7.4). Specimens were washed in pure CB, postfixed in 1% OsO4 in CB for 1 hour, dehydrated in ascending series of ethanol and propyleneoxide, blocked-stained in uranyl acetate for 4 hours, and flat-embedded in Araldite (Serva). An ultramicrotome (Ultracut; Leica) was used to cut semithin (1 μm) and ultrathin (50 nm) sections. Ultrathin sections were stained with lead citrate, mounted on copper grids, and analyzed with a Zeiss EM 10 electron microscope (Zeiss). Pictures were scanned at 300 dpi and processed (Adobe Photoshop).

**Microscopy.** Fluorescent staining analysis used inverse Zeiss Axiostar 200 M microscopes with AxioCam MRm camera and ApoTome (Zeiss), using AxioVision graphics software. Retinal overviews (original magnification, x5) are presented as composite images of individual, successively overlapping (5%) images, generated by computer-controlled x-y settings and processed using MosaiX Software. H&E-stained sections were visualized using Zeiss Axioplan 2 with AxioCamHRc camera. Higher magnifications were obtained with x10, x20, x40, and x63 objectives.

**Statistics.** Data are presented as mean ± SEM. For comparisons between experiments, values were normalized to control (assigned as 1 or 100%). 2-tailed Student’s t tests were used to identify statistical significance. Statistical significance was assumed for P values less than 0.05.

**Study approval.** All animal experiments were approved by the Regierungspräsidium Tübingen (Tübingen, Germany).

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