RESEARCH ARTICLE

An important role of cutaneous lymphatic vessels in coordinating and promoting anagen hair follicle growth

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Abstract

The lymphatic vascular system plays important roles in the control of tissue fluid homeostasis and immune responses. While VEGF-A-induced angiogenesis promotes hair follicle (HF) growth, the potential role of lymphatic vessels (LVs) in HF cycling has remained unknown. In this study, we found that LVs are localized in close proximity to the HF bulge area throughout the postnatal and depilation-induced hair cycle in mice and that a network of LVs directly connects the individual HFs. Increased LV density in the skin of K14-VEGF-C transgenic mice was associated with prolongation of anagen HF growth. Conversely, HF entry into the catagen phase was accelerated in K14-sVEGFR3 transgenic mice that lack cutaneous LVs. Importantly, repeated intradermal injections of VEGF-C promoted hair growth in mice. Conditioned media from lymphatic endothelial cells promoted human dermal papilla cell (DPC) growth and expression of IGF-1 and alkaline phosphatase, both activators of DPCs. Our results reveal an unexpected role of LVs in coordinating and promoting HF growth and identify potential new therapeutic strategies for hair loss-associated conditions.

Introduction

The hair follicle (HF) is a mini-organ of the skin that continuously cycles through rapid growth (anagen phase), apoptosis-driven regression (catagen phase) and relative quiescence (telogen phase) [1]. The HF-associated blood vascular system undergoes a massive expansion during the anagen phase and a rapid involution during the catagen regression phase [2, 3]. In previous studies, we found that the production of vascular endothelial growth factor (VEGF)-A is up-regulated during the anagen phase and that transgenic mice with increased levels of cutaneous VEGF-A have larger HFs than their wildtype littermates [4]. Conversely, blockade of VEGF-A resulted in thinner HFs [5] and hair thinning has been reported under anti-angiogenic therapy [6]. Indeed, it has been proposed that stimulation of blood vessels by induction of VEGF-A production represents a major mechanism of action of the hair growth-promoting drug minoxidil [7].
While blood vessels supply peripheral tissues with oxygen and nutrients and are associated with leakage of fluid into these tissues, a major function of lymphatic vessels (LVs) is the drainage of tissue fluid and cells from the periphery and their recirculation, via collecting LVs and the thoracic duct, to the blood vascular system [8–11]. Beyond fluid drainage, LVs play important roles in the transport of dendritic and other immune cells to the lymph nodes, and recent evidence indicates that they might also be involved in the control of autoimmunity to peripheral tissue antigens [12, 13]. Impairment of lymphatic function often leads to edema, impaired immunity, impaired wound healing and tissue fibrosis [14]. There has been a strongly elevated interest in lymphatic research since an involvement of LVs in the pathogenesis of an increasing number of diseases has been found during the last few years, including metastatic cancer, chronic autoimmune diseases, impaired wound healing, organ transplant rejection, myocardial infarction and atherosclerosis [15–25].

While studying the inflammatory response in transgenic mice with skin-specific overexpression of the lymphangiogenic growth factor VEGF-C (K14-VEGF-C) [26], we observed that hair regrowth after skin shaving was accelerated in these mice (unpublished observation). We therefore wondered whether LVs might play an active role in the homeostasis of HF cycling. To this end, we first characterized the lymphatic vascularity during postnatal HF cycling in mice. We then investigated genetic mouse models with enhanced or decreased lymphatic vascularization of the skin: K14-VEGF-C transgenic mice that have an increased cutaneous lymphatic network [27], and transgenic mice with skin-specific overexpression of a soluble, extracellular domain of VEGF receptor-3 (K14-sVEGFR3 transgenic mice) [28]. The soluble VEGFR-3 scavenges its ligands VEGF-C and VEGF-D and thereby prevents these ligands from activating VEGFR-3 on the lymphatic endothelium, resulting in the absence of LVs in the skin [28]. We also investigated the biological effects of intradermal VEGF-C protein delivery on hair growth. Our results reveal an unexpected, major role of lymphatic endothelium in promoting anagen HF growth in mice.

Materials and methods

Mouse models

K14-VEGF-C transgenic mice on the FVB background that express VEGF-C under control of the keratin 14 (K14) promoter, and K14-sVEGFR3 transgenic mice that express a sVEGFR3 [27, 28] were kindly provided by Dr. Kari Alitalo, University of Helsinki. To investigate the effects of lymphatic vasculature on the hair cycle after depilation-induced hair regeneration, the back skin of 8-week-old female K14-VEGF-C or K14-sVEGFR3 transgenic mice in the telogen phase was depilated using wax as described [1, 29], resulting in the synchronized induction of new anagen follicle growth. The number of mice (WT: n = 24, K14-VEGF-C transgenic: n = 10, K14-sVEGFR3 transgenic mice: n = 8) used for each experiment is indicated in the figure legends. Mice were sacrificed with an overdose of anaesthesia (160 mg kg$^{-1}$ ketamine; 0.4 mg kg$^{-1}$ medetomidine) at days 15 and 18 after depilation, and the back skin was taken for histological analysis. Grading the hair cycle phases was done using hematoxylin and eosin (H&E)-stained paraffin sections, according to established guidelines [1]. To measure the bulb diameter, 3 images/mouse were acquired and the bulb diameter was measured at the level of the largest diameter (“Auber’s line”) of the hair bulbs with a clearly visible dermal papilla (DP) [4]. For quantitative analysis, the Image J software (National Institutes of Health, Bethesda, MD, USA) was used. To examine the effects of the lymphatic vasculature on the postnatal hair cycle, back skin samples were obtained from WT (n = 4) and K14-VEGF-C transgenic female mice (n = 9) at postnatal days 25.
To investigate the effects of recombinant VEGF-C on HF cycling, anagen induction was performed in 8-week-old female C57BL/6J mice (Janvier Labs, Le Genest-Saint-Isle, France) as described previously [30]. The back skin of mice in the telogen phase was shaved with a clipper. Vehicle (PBS containing 0.1% bovine serum albumin [BSA]) alone (n = 4) or vehicle containing VEGF-C (R&D Systems, Minneapolis, MN, USA; 200 ng in 20 μL, n = 5) or minoxidil (MNX; Sigma-Aldrich, St. Louis, MO, USA, positive control; 20 μg in 20 μL, n = 5) was applied via intradermal injection every day for 40 days and back skin samples were obtained at the site of intradermal injection and processed for H&E-stained paraffin sections. Proxl-tdTomato reporter mice (FVB background) that express tdTomato under the Prox1 promoter (Mutant Mouse Resource & Research Centers supported by NIH, stock number: 036531-UCD) were used to analyze the dermal lymphatic network. Unfixed back skin samples were imaged using a Zeiss Axiozoom V16 fluorescence stereomicroscope. All experimental procedures were conducted according to animal protocols approved by the Kantonales Veterinaeramt Zuerich (protocol 237/2013).

Immunofluorescence stainings
Back skin samples were embedded in OCT (Leica Biosystems, Newcastle, UK) and frozen in liquid nitrogen. 10-μm frozen sections were fixed in 4% paraformaldehyde (PFA) for 15 min at room temperature (RT), washed in PBS and incubated with blocking solution (5% donkey serum, 0.2% BSA and 0.3% Triton X-100 in PBS) for 1 h at RT. Next, the sections were stained with primary antibodies overnight at 4˚C and, after several washes, incubated with secondary antibodies for 30 min at RT. Primary antibodies were as follows: rat anti-CD31 (BD Biosciences, San Jose, CA, USA), rabbit anti-LYVE-1 (Angiobio, Del Mar, CA, USA), goat anti-LYVE-1 (R&D Systems), rat anti-CD68 (Abcam, Cambridge, MA, USA), goat anti-podoplanin (R&D Systems), rabbit anti-cytokeratin 15 (Abcam), rat anti-Foxp3 (eBioscience, San Diego, CA, USA), and goat anti-Prox1 (R&D systems). Secondary antibodies (all from Thermo Fisher, San Jose, CA, USA) were as follows: donkey anti-rabbit Alexa Fluor 488, 594 or 647, donkey anti-rat Alexa Fluor 488 or 594, donkey anti-goat Alexa Fluor 488 or 594, and chicken anti-goat Alexa Fluor 647. Hoechst 33342 (Invitrogen, Carlsbad, CA, USA) was used for nuclear staining. Immunofluorescence images were acquired by an Axioskop 2 mot plus microscope (Carl Zeiss, Jena, Germany) and Z stacks of images were obtained using a Zeiss LSM 710 FCS confocal microscope.

Cell culture
The methods used for culturing human DPCs have been described previously [31, 32]. DPCs (ScienCell Research Laboratories, Carlsbad, CA, USA) were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco, Grand Island, NY, USA) supplemented with 10% fetal bovine serum (FBS; Gibco), 10 ng/mL basic fibroblast growth factor (Peprotech, London, UK) and antibiotic/antimycotic solution (Gibco). Primary human LECs and BECs isolated from foreskin were cultured as described previously [33]. LECs were cultured on 10 μg/mL fibronectin (Millipore, Billerica, MA, USA)-coated dishes in endothelial cell basal medium (EBM; Lonza, Walkersville, MD, USA), supplemented with 20% FBS, 2 mM L-glutamine (Gibco), antibiotic-antimycotic solution, 10 μg/mL hydrocortisone (Sigma-Aldrich) and 25 μg/mL cAMP (Sigma-Aldrich). BECs were cultured on 10 μg/mL fibronectin-coated dishes in EBM supplemented with 20% FBS, 2 mM L-glutamine, 1x antibiotic-antimycotic solution and 0.4% endothelial cell growth supplement (ECGS; PromoCell). Human dermal fibroblasts (DFs) were cultured in DMEM supplemented with 10% FBS and antibiotic/antimycotic solution. Cells were incubated at 37˚C in a 5% CO2 incubator.
Conditioned media
To investigate the effects of conditioned media (CM) on DPCs, CM of LECs, BECs and DFs were collected. When cells reached 80% confluence, the growth medium was replaced with EBM containing 2% FBS for LECs and BECs or with DMEM containing 2% FBS for DFs. CM were collected after 48 h and centrifuged. For the preparation of control CM, DMEM or EBM containing 2% FBS was added to empty cell culture dishes and collected after 48 h. DPCs were incubated with 10, 30, 50 or 90% LEC-CM, BEC-CM or DF-CM for 72 h or 100 ng/ml IGF-1 as a positive control. For the quantification of DPC proliferation, Cell Counting Kit-8 (Sigma-Aldrich) was used according to the manufacturer’s instructions. After 72 h, the absorbance at 450 nm was measured using a microplate reader (Spectramax Gemini EM, Molecular Devices, Sunnyvale, CA, USA).

Quantitative real time-polymerase chain reaction (qRT-PCR)
Total RNA was isolated from DPCs using NucleoSpin RNA (Macherey-Nagel, Düren, Germany) and 1 μg of total RNA was used for the cDNA synthesis reaction using the High Capacity Reverse Transcription kit (Applied Biosystems, Foster City, CA, USA). PCR was performed on a 7900HT Fast Real-Time PCR System (Applied Biosystems) using FastStart SYBR green master mix (Roche Diagnostics, Basel, Switzerland) according to the manufacturer’s instructions. Gene expression was normalized to the control gene Rplp0 (36B4). Primer information is provided in S1 Table. All experiments were independently repeated at least 3 times.

Western blotting
Total protein from DPCs was extracted in a buffer containing 25 mM HEPES, 150 mM NaCl, 1% Triton X-100, 10% glycerol, 5 mM EDTA, 10 mM NaF, 2 mM Na3VO4, and protease inhibitor cocktail (Roche Diagnostics). Proteins were separated using 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transferred to a polyvinylidene fluoride membrane (EMD Millipore, Billerica, MA, USA) using a wet transfer system. The blotted membranes were incubated with primary antibodies at 4°C. The following antibodies were used: anti-total Akt, anti-phosphorylated Akt (Cell Signaling Technology, Beverly, MA, USA) and anti-beta-actin (Abcam). Membranes were then probed with an anti-rabbit-IgG-horseradish peroxidase conjugate (GE Healthcare, Hilleroed, Denmark). Antibody–antigen complexes were detected using the ECL system (Thermo Scientific).

Statistical analyses
The statistical tests used are indicated in the figure legends and results were considered significant at \( P < 0.05 \). The diameters of hair bulbs were analyzed with the two-tailed unpaired t-test for each time point. For the studies with CM, mRNA expression levels were analyzed using the two-tailed paired t-test (GraphPad Prism, version 5.0a, San Diego, CA, USA). The differences of HF cycling between WT and transgenic mice were compared by Fisher’s exact test, calculated online at QuickCalcs, GraphPad Software.

Results
LVs are localized in close proximity to the bulge and the bulb region of murine HFs
To characterize the dermal lymphatic vascularity, we investigated the LV distribution in the back skin of Prox1 (lymphatic-specific transcription factor)-tdTomato reporter mice that
express tdTomato under control of the Prox1 promoter. By fluorescence stereomicroscopy, we found that a network of LVs directly connects the individual HFs (Fig 1A). To further characterize the location and morphology of LVs during the hair cycle, back skin samples of wildtype mice in the anagen phase (postnatal day 8) were stained for the panvascular marker CD31 and the lymphatic vessel marker LYVE-1. CD31+/LYVE-1+ LVs were located in close proximity to anagen HFs, predominantly in an area close to the hair bulge (Fig 1B). Since LYVE-1 may also be expressed by some cutaneous macrophages, we performed double immunofluorescence stainings to confirm the identity of HF-associated LVs. The LYVE-1-positive vessels did not express the macrophage marker CD68 (Fig 1C) but co-expressed the LV marker podoplanin (Fig 1D). To further characterize the location of LVs in relation to the HF, we performed double immunofluorescence stainings for keratin 15, a putative marker of follicular stem cells, and LYVE-1. LVs were frequently localized near the stem cell region of the bulge of the HF (Fig 1E) and appeared to be directly attached to the bulge region (Fig 1F). This finding was confirmed by tissue whole-mount stainings for LYVE-1 and keratin 15, followed by light-sheet microscopy, revealing precollecting vessels that extended along HFs during the anagen phase (S1 Video) but not during the telogen phase (S2 Video). The LVs in close proximity to the hair bulbs (Fig 1G) and the hair bulge (Fig 1H) also expressed the lymphatic-specific transcription factor Prox1+.

Prolonged depilation-induced anagen hair growth in K14-VEGF-C transgenic mice

To investigate whether HF-associated LVs might have a functional role in HF growth and/or cycling, we studied depilation-induced hair regeneration in female K14-VEGF-C transgenic mice that are characterized by an increased density of LVs in the skin [27] (Fig 2A). In line with our previous findings [34, 35], we observed no differences in MECA-32+ blood vessels (S1 Fig). Then, we measured the diameter of the hair bulb after depilation-induced hair regeneration, which typically increases during the anagen phase, whereas it is reduced during catagen development [1]. At day 15 (late-anagen phase) after depilation, all WT and K14-VEGF-C transgenic mice were in the anagen VI phase with comparable hair bulb diameters (Fig 2B and Fig 2C). However, at day 18 (catagen phase), the diameter of the hair bulb was significantly larger in transgenic mice (Fig 2B and Fig 2D), indicating that after depilation, K14-VEGF-C transgenic mice remained longer in the anagen growth phase than WT mice during the anagen-to-catagen transition of the HF.

Early entry into the catagen phase in K14-sVEGFR3-Ig transgenic mice

Since VEGF-C prolonged the anagen phase of the HF, we next investigated whether blockade of VEGF-C might accelerate the entry into the catagen phase. To this end, we first studied depilation-induced hair regeneration in female K14-sVEGFR3-Ig mice. These mice express a soluble form of the extracellular domain of VEGFR3, thus scavenging VEGF-C (and -D) and preventing its interaction with its receptors on lymphatic endothelial cells. Immunofluorescence stainings for LYVE-1 confirmed the absence of LVs in the back skin of K14-sVEGFR3-Ig mice, whereas WT mice showed a normal lymphatic vasculature (Fig 2E). At day 15 (late-anagen phase), only anagen VI HFs were observed in all WT mice, whereas all K14-sVEGFR3-Ig mice had already early-catagen HFs (Fig 2F). Accordingly, the thickness of the hair bulbs was significantly smaller in K14-sVEGFR3-Ig mice (Fig 2G). At day 18 (catagen phase), all mice were in mid-catagen phase (Fig 2F and Fig 2H). Taken together, these results indicate that inhibition of VEGF-C signaling in K14-sVEGFR3 transgenic mice accelerates the entry of HFs into the catagen phase during the anagen-to-catagen transition.
VEGF-C levels are increased during the anagen phase

We next investigated whether the expression levels of VEGF-C and its receptor VEGFR3 might undergo cyclic changes during the normal postnatal hair cycle. We found that the mRNA expression levels of VEGF-C and VEGFR3 in the back skin were significantly higher during the anagen phase (at P28) than during the telogen phase (at P49) (S2A Fig). Similar results were obtained during depilation-induced HF regeneration, where the VEGF-C mRNA expression levels were significantly higher during the anagen phase (day 12 after depilation) than during the telogen phase (day 22 after depilation) (S2B Fig). By contrast, mRNA expression levels of the related lymphangiogenic growth factor VEGF-D [36] remained unchanged during postnatal and depilation-induced hair cycling (S2A Fig and S2B Fig). In addition, we measured the amount of VEGF-C secretion in conditioned media (CM) from dermal fibroblasts (DF) and dermal papilla cells (DPC) by enzyme-linked immunosorbent assay (ELISA). We found that DF-CM and DPC-CM contained VEGF-C, indicating that DF and DPC express VEGF-C (S3 Fig).

Accelerated initiation of anagen HF growth during normal postnatal hair follicle cycling in K14-VEGF-C transgenic mice and after intradermal delivery of VEGF-C

Given the cyclic changes of VEGF-C mRNA expression during the hair cycle and the prolongation of anagen HF growth in K14-VEGF-C transgenic mice, we next investigated whether cutaneous LVs might also have a biological role during normal postnatal hair cycling. At postnatal day 25 (P25; second postnatal hair cycle), 88% of female K14-VEGF-C transgenic mice were in the anagen IV phase and 12% in anagen III, whereas only 25% of WT mice were in the anagen IV phase and 75% still in anagen III (Fig 3A and Fig 3B). However, there was no differences in epidermal thickness between WT and K14-VEGF-C transgenic mice (S4 Fig). These results indicate that K14-VEGF-C transgenic mice initiated the anagen HF growth more rapidly than WT mice also during normal, postnatal HF cycling.

Next we wondered whether application of recombinant VEGF-C protein might promote anagen induction. To this end, we intradermally injected 200 ng VEGF-C, 20 μg minoxidil (positive control) or an equal volume (20 μl) PBS (negative control) into the back skin of 8-week-old female C57BL/6J mice daily for 40 days and then analyzed the hair cycle phase. After 40 days, skin pigmentation—a sign for HF entrance into the anagen phase—was observed at the site of intradermal injections in the VEGF-C- and the minoxidil-treated group, but not in the control group (Fig 3C). Analyses of histological sections revealed that 60% of the VEGF-C-treated mice and 40% of the MNX-treated mice were in the anagen hair cycle phase, whereas all vehicle-treated mice were still in the telogen phase (Fig 3D and Fig 3E). These results identify a potent activity of VEGF-C treatment in promoting HF entry into the anagen growth phase during the telogen-to-anagen transition.
Cutaneous lymphatic vessels promote hair follicle growth

A

B

Bulb diameter

C

D

E

WT K14-VEGF-C Tg

F

Bulb diameter

G

H

WT K14-sVEGFR3 Tg
LEC conditioned media promote dermal papilla cell proliferation

Given the strong association between LV expansion and the prolongation of anagen HF growth in K14-VEGF-C transgenic mice, we next investigated whether lymphatic endothelial cells (LECs) might secrete paracrine factors that activate DPCs, since HF stem cells receive stimulatory signals from DPCs to drive hair cycling [37, 38]. We collected conditioned media (CM) from cultured human dermal LECs and examined their effects on cultured human DPCs. 100 ng/ml IGF-1, a growth factor known to stimulate HF growth [39], was used as a positive control. Incubation with 30%, 50% or 90% LEC-CM significantly increased DPC proliferation in a dose-dependent manner, to a similar or even higher extent than IGF-1, as compared to DPCs incubated with equivalent amounts of control conditioned media (CON-CM) (Fig 4A). In line with this, incubation of DPCs with LEC-CM also increased the phosphorylation of Akt which is thought to promote cell survival [40] (Fig 4B). On the contrary, incubation of DPCs with CM from human dermal blood vascular endothelial cells or from dermal fibroblasts had no major effect on DPC proliferation (S5 Fig). Importantly, direct treatment with VEGF-C did not affect DPC proliferation (Fig 4C), suggesting that VEGF-C acts via LECs to modulate HF cyling.

LEC conditioned media promote IGF1 and ALP mRNA expression, and inhibit BMP-2, BMP-4 and TGF-β1 mRNA expression in dermal papilla cells

After incubation of DPCs with LEC-CM (30%, 50% or 90%) for 72 h, mRNA expression levels of IGF-1 were markedly enhanced (Fig 4D). Incubation with LEC-CM also potently increased the mRNA expression levels of alkaline phosphatase (ALP) that maintains the inductive properties of the dermal papilla [41, 42] (Fig 4E). This was not the case when DPCs were treated with CM from dermal blood vascular endothelial cells or from dermal fibroblasts (Fig 4D and Fig 4E). LEC-CM also increased the mRNA expression levels of FGF-10, a known promoter of hair growth [38, 43], though less potently than blood vascular endothelial cell-CM (Fig 4F). Conversely, incubation of DPCs with LEC-CM markedly decreased the mRNA expression levels of BMP-2 and BMP-4 that have been reported to inhibit HF growth [38, 44] (Fig 4G and Fig 4H). The mRNA expression levels of TGF-β1, another inhibitor of HF growth [45], were also reduced after LEC-CM incubation (Fig 4I). CM from blood vascular endothelial cells or from dermal fibroblasts had no major effects on the expression of these genes. Taken together, these results identify potential paracrine mechanisms by which lymphatic endothelium might activate DPCs.

Discussion

In this study, we identified perifollicular LVs as new players in the coordination and regulation of HF cycling. We found that lymphatic vessels in the skin form a network that connects the individual hair follicles, and that during the anagen phase, dermal LVs reside in close
proximity to murine HFs, particularly the dermal papilla and the stem cell area in the bulge. Importantly, using genetic mouse models, we demonstrate that increased dermal levels of the lymphangiogenesis factor VEGF-C lead to prolongation of anagen HF growth, whereas VEGF-C depletion resulted in accelerated catagen progression of the HFs. Similarly,
Cutaneous lymphatic vessels promote hair follicle growth

A

B

C

D

E

F

G

H

I

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Exogenous administration of VEGF-C by intradermal injection accelerated the telogen-to-anagen transition. Previously, it has been found that VEGF-A induces proliferation of DPCs through the VEGFR-2/ERK pathway and has a direct stimulatory effect on DPCs [46]. On the contrary, in this study, VEGF-C treatment did not directly increase the proliferation of DPCs. Together with the strong expansion of LVs in the skin of K14-VEGF-C transgenic mice and the spatial association of LVs with the dermal papilla during the anagen phase, our data indicate that VEGF-C might promote hair growth indirectly via a paracrine effect of VEGF-C-activated LEC on DPCs.

When cocultured with human microvascular endothelial cells, proliferation of DPCs was significantly increased [47]. Here, we found that also LEC conditioned media promoted cell proliferation of DPCs. Moreover, LEC conditioned media, in contrast to media conditioned by other relevant dermal cell populations such as blood vascular endothelial cells and dermal fibroblasts, significantly increased IGF-1, FGF-10 and ALP mRNA expression and inhibited BMP-2, BMP-4 and TGF-ß1 by DPCs, suggesting a potentially increased capacity of these cells to induce HF stem cell activation.

Beyond their physiological role in the normal hair cycle, LVs are likely involved in pathological hair loss as well, and activation of LVs might serve as a potential new treatment option for such conditions. Some reports have indicated that in permanent alopecias, such as lichen planopilaris and discoid lupus erythematosus, infiltrating immune cells surround the bulge region, whereas in reversible alopecias, such as alopecia areata, T lymphocytes directly infiltrate the bulb region [48–50]. We previously found that during chronic skin inflammation, the lymphatic vessel drainage function is impaired, associated with dilation of LVs, and that activation of LVs reduces chronic skin inflammation [9, 18, 20]. Dilated LVs were also observed in a mouse model of alopecia areata [51] and in human lipedematous alopecia [52]. It will be of interest to investigate whether activation of the lymphatic system, for example by administration of VEGF-C, might protect HFs from inflammation-associated damage.

In conclusion, these results reveal an important function of the LV network in the coordination and regulation of the HF cycle, and they indicate potential new therapeutic strategies for the treatment of conditions associated with hair loss.

Supporting information

S1 Fig. Comparable blood vessel density in WT and K14-VEGF-C mice. (TIF)

S2 Fig. Increased VEGF-C mRNA expression levels during the anagen phase. (TIF)

S3 Fig. VEGF-C content in DPC-CM and DF-CM. (TIF)
S4 Fig. Epidermal thickness in WT and K14-VEGF-C transgenic mice is comparable. (TIF)

S5 Fig. Neither DF nor BEC conditioned media do not affect DPC proliferation. (TIF)

S1 Table. Sequences of primers used for human genes. (TIF)

S1 Video. Ultramicroscopy image of back skin (anagen phase) stained for LYVE-1 (red) and cytokeratin 15 (green). (MOV)

S2 Video. Ultramicroscopy image of back skin (telogen phase) stained for LYVE-1 (red) and cytokeratin 15 (green). (MOV)

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References
1. Muller-Rover S, Handjiski B, van der Veen C, Eichmuller S, Foitzik K, McKay IA, et al. A comprehensive guide for the accurate classification of murine hair follicles in distinct hair cycle stages. J Invest Dermatol. 2001; 117(1):3–15. Epub 2001/07/10. https://doi.org/10.1046/j.0022-202x.2001.01377.x PMID: 11442744.
2. Mecklenburg L, Tobin DJ, Muller-Rover S, Handjiski B, Wendt G, Peters EM, et al. Active hair growth (anagen) is associated with angiogenesis. J Invest Dermatol. 2000; 114(5):909–16. Epub 2000/04/20. https://doi.org/10.1046/j.1523-1747.2000.00954.x PMID: 10771470.

3. Odorisio T, Cianfarrani F, Failla CM, Zambruno G. The placenta growth factor in skin angiogenesis. J Dermatol Sci. 2006; 41(1):11–9. Epub 2005/10/12. https://doi.org/10.1016/j.jdermsci.2005.08.008 PMID: 16216466.

4. Yano K, Brown LF, Detmar M. Control of hair growth and follicle size by VEGF-mediated angiogenesis. J Clin Invest. 2001; 107(4):409–17. Epub 2001/02/22. https://doi.org/10.1172/JCI11317 PMID: 11181640; PubMed Central PMCID: PMC199257.

5. Yano K, Brown LF, Lawler J, Miyakawa T, Detmar M. Thrombospondin-1 plays a critical role in the induction of hair follicle involution and vascular regression during the catagen phase. J Invest Dermatol. 2003; 120(1):14–9. Epub 2003/01/22. https://doi.org/10.1046/j.1523-1747.2003.12045.x PMID: 12535193.

6. Kubota Y. Tumor angiogenesis and anti-angiogenic therapy. Keio J Med. 2012; 61(2):47–56. Epub 2012/07/05. PMID: 22760023.

7. Messenger AG, Rundegren J. Minoxidil: mechanisms of action on hair growth. Br J Dermatol. 2004; 150(2):186–94. Epub 2004/03/05. https://doi.org/10.1111/j.1365-2133.2004.05785.x PMID: 14996087.

8. Shayan R, Achen MG, Stacker SA. Lymphatic vessels in cancer metastasis: bridging the gaps. Carcinogenesis. 2006; 27(9):1729–38. https://doi.org/10.1093/carcin/bgl031 PMID: 16597644.

9. Huggenberger R, Detmar M. The cutaneous vascular system in chronic skin inflammation. J Investig Dermatol Symp Proc. 2011; 15(1):24–32. https://doi.org/10.1038/jidsymp.2011.5 PMID: 22076324; PubMed Central PMCID: PMC3398151.

10. Dieterich LC, Seidel CD, Detmar M. Lymphatic vessels: new targets for the treatment of inflammatory diseases. Angiogenesis. 2014; 17(2):359–71. https://doi.org/10.1007/s10456-013-9406-1 PMID: 24212981.

11. Hiraoka S, Detmar M, Karaman S. Lymphatics in nanophysiology. Adv Drug Deliv Rev. 2014; 74:12–8. https://doi.org/10.1016/j.addr.2014.01.011 PMID: 24524932.

12. Card CM, Yu SS, Swartz MA. Emerging roles of lymphatic endothelium in regulating adaptive immunity. J Clin Invest. 2014; 124(3):943–52. Epub 2014/03/05. https://doi.org/10.1172/JCI73316 PMID: 24590280; PubMed Central PMCID: PMC3938271.

13. Thomas SN, Rutkowski JM, Pasquier M, Kuan EL, Alitalo K, Randolph GJ, et al. Impaired humoral immunity and tolerance in K14-VEGF-R-3-Ig mice that lack dermal lymphatic drainage. J Immunol. 2012; 189(5):2181–90. Epub 2012/07/31. https://doi.org/10.4049/jimmunol.1103545 PMID: 22844119; PubMed Central PMCID: PMC3424306.

14. Skobe M, Detmar M. Structure, function, and molecular control of the skin lymphatic system. J Investig Dermatol Symp Proc. 2000; 5(1):14–9. Epub 2001/01/09. https://doi.org/10.1046/j.1087-0024.2000.00001.x PMID: 11147669.

15. Dadras SS, Paul T, Bertoncini J, Brown LF, Muzikansky A, Jackson DG, et al. Tumor lymphangiogenesis: a novel prognostic indicator for cutaneous melanoma metastasis and survival. Am J Pathol. 2003; 162(6):1951–60. Epub 2003/05/22. https://doi.org/10.1016/S0002-9440(10)64328-3 PMID: 12759251; PubMed Central PMCID: PMC1868148.

16. Altalato A, Detmar M. Interaction of tumor cells and lymphatic vessels in cancer progression. Oncogene. 2012; 31(42):4499–508. https://doi.org/10.1038/onc.2011.602 PMID: 22179834.

17. He Y, Kozaki K, Karpanen T, Koshikawa K, Yla-Herttuala S, Takahashi T, et al. Suppression of tumor lymphangiogenesis and lymph node metastasis by blocking vascular endothelial growth factor receptor 3 signaling. J Natl Cancer Inst. 2002; 94(11):819–25. Epub 2002/06/06. https://doi.org/10.1093/jnci/94.11.819 PMID: 12048269.

18. Huggenberger R, Ullmann S, Proulx ST, Riemann R, Ator E, Lyubynska N, Achen MG, Detmar M. Stimulation of lymphangiogenesis via VEGFR-3 inhibits chronic skin inflammation. J Exp Med. 2010; 207(10):2255–69. https://doi.org/10.1084/jem.20100559 PMID: 20837699; PubMed Central PMCID: PMC2947063.

19. Baluk P, Tammela T, Achen MG, Detmar M. Pathogenesis of persistent lymphatic vessel hyperplasia in chronic airway inflammation. J Clin Invest. 2005; 115(2):247–57. Epub 2005/01/26. https://doi.org/10.1172/JCI22037 PMID: 15668734; PubMed Central PMCID: PMCS44601.

20. Proulx ST, Luciani P, Dieterich LC, Karaman S, Leroux JC, Etzioni A, Detmar M. Expansion of the lymphatic vasculature in cancer and inflammation: new opportunities for in vivo imaging and drug delivery. J Control Release. 2013; 172(2):550–7. https://doi.org/10.1016/j.jconrel.2013.04.027 PMID: 23685257.
21. Tacconi C, Correale C, Gandelli A, Spinelli A, Dejana E, D’Alessio S, et al. Vascular endothelial growth factor C disrupts the endothelial lymphatic barrier to promote colorectal cancer invasion. Gastroenterology. 2015; 148(7):1438–51 e8. https://doi.org/10.1053/j.gastro.2015.03.005 PMID: 25754161.

22. D’Alessio S, Correale C, Tacconi C, Gandelli A, Pietrogrande G, Vetranò S, et al. VEGF-C-dependent stimulation of lymphatic function ameliorates experimental inflammatory bowel disease. J Clin Invest. 2014; 124(9):3863–78. Epub 2014/08/12. https://doi.org/10.1172/JCI72189 PMID: 25105363; PubMed Central PMCID: PMC4152127.

23. Maruyama K, Asai J, II M, Thorne T, Losordo DW, D’Amore PA. Decreased macrophage number and activation lead to reduced lymphatic vessel formation and contribute to impaired diabetic wound healing. Am J Pathol. 2007; 170(4):1178–91. Epub 2007/03/30. https://doi.org/10.2353/ajpath.2007.060018 PMID: 17392158; PubMed Central PMCID: PMC1829452.

24. Dietrich T, Bock F, Yuen D, Hos D, Bachmann BO, Zahn G, et al. Cutting edge: lymphatic vessels, not blood vessels, primarily mediate immune rejections after transplantation. J Immunol. 2010; 184(2):535–9. Epub 2009/12/19. https://doi.org/10.4049/jimmunol.0903180 PMID: 20018627; PubMed Central PMCID: PMC4725297.

25. Kholova I, Dragneva G, Cermakova P, Laidinen S, Kaskenpaa N, Hazes T, et al. Lymphatic vasculature is increased in heart valves, ischaemic and inflamed hearts and in cholesterol-rich and calcified atherosclerotic lesions. Eur J Clin Invest. 2011; 41(5):487–91. Epub 2010/03/30. https://doi.org/10.1111/j.1365-2362.2010.04315.x PMID: 21128936.

26. Christiansen AJ, Dieterich LC, Ohs I, Bachmann SB, Bianchi R, Proulx ST, et al. Lymphatic endothelial cells attenuate inflammation via suppression of dendritic cell maturation. Oncotarget. 2016. Epub 2016/06/09. https://doi.org/10.18632/oncotarget.9820 PMID: 27270646.

27. Jeltsch M, Kaipainen A, Joukov V, Meng X, Lakso M, Rauvala H, et al. Hyperplasia of lymphatic vessels in VEGF-C transgenic mice. Science. 1997; 276(5317):1423–5. Epub 1997/05/30. https://doi.org/10.1126/science.276.5317.1423 PMID: 9162011.

28. Makinen T, Jussila L, Veikkola T, Karpanen T, Pulkkainen KJ, et al. Inhibition of lymphangiogenesis with resulting lymphedema in transgenic mice expressing soluble VEGF receptor-3. Nat Med. 2001; 7(2):199–205. Epub 2001/02/15. https://doi.org/10.1038/84651 PMID: 11175851.

29. Paik SH, Yoon JS, Ryu HH, Lee JY, Shin CY, Min KH, et al. Pretreatment of epidermal growth factor promotes primary hair recovery via the dystrophic anagen pathway after chemotherapy-induced alopecia. Exp Dermatol. 2013; 22(7):496–9. Epub 2013/06/27. https://doi.org/10.1111/exd.12182 PMID: 23800066.

30. Paus R, Stenn KS, Link RE. The induction of anagen hair growth in telogen mouse skin by cyclosporine A administration. Lab Invest. 1989; 60(3):365–9. Epub 1989/03/01. PMID: 2927078.

31. Kwon OS, Pyo HK, Oh YJ, Han JH, Lee SR, Chung JH, et al. Promotive effect of minoxidil combined with all-trans retinoic acid (tretinoin) on human hair growth in vitro. J Korean Med Sci. 2007; 22(2):283–9. Epub 2007/04/24. https://doi.org/10.3346/jkms.2007.22.2.283 PMID: 17449938; PubMed Central PMCID: PMC2693596.

32. Yoon SY, Yoon JS, Jo SJ, Shin CY, Shin JY, Kim JI, et al. A role of placental growth factor in hair growth. J Dermatol Sci. 2014; 74(2):125–34. Epub 2014/03/04. https://doi.org/10.1016/j.jdermsci.2014.01.011 PMID: 24582062.

33. Hirakawa S, Hong Y-K, Harvey N, Schacht V, Matsuda K, Libermann T, et al. Identification of vascular lineage-specific genes by transcriptional profiling of isolated blood vascular and lymphatic endothelial cells. Am J Pathol. 2003; 162(2):575–86. https://doi.org/10.1016/S0002-9440(10)63851-5 PMID: 12547715.

34. Hirakawa S, Brown LF, Kodama S, Paavonen K, Alltalo K, Detmar M. VEGF-C-induced lymphangiogenesis in sentinel lymph nodes promotes tumor metastasis to distant sites. Blood. 2007; 109(3):1010–7. Epub 2006/10/13. https://doi.org/10.1182/blood-2006-05-021758 PMID: 17032920; PubMed Central PMCID: PMC1785149.

35. Huggenberger R, Siddiqui SS, Brander D, Ullmann S, Zimmermann K, Antsiferova M, et al. An important role of lymphatic vessel activation in limiting acute inflammation. Blood. 2011; 117(17):4667–78. https://doi.org/10.1182/blood-2010-10-316356 PMID: 21364190; PubMed Central PMCID: PMC3099581.

36. Adams RH, Alltalo K. Molecular regulation of angiogenesis and lymphangiogenesis. Nat Rev Mol Cell Biol. 2007; 8(6):464–78. https://doi.org/10.1038/nrm2183 PMID: 17522591.

37. Hsu YC, Li L, Fuchs E. Emerging interactions between skin stem cells and their niches. Nat Med. 2014; 20(8):847–56. Epub 2014/08/08. https://doi.org/10.1038/nm.3643 PMID: 25100530; PubMed Central PMCID: PMC4358898.
38. Hsu YC, Fuchs E. A family business: stem cell progeny join the niche to regulate homeostasis. Nat Rev Mol Cell Biol. 2012; 13(2):103–14. https://doi.org/10.1038/nrm3272 PMID: 22266760; PubMed Central PMCID: PMC3280338.

39. Li J, Yang Z, Li Z, Gu L, Wang Y, Sung C. Exogenous IGF-1 promotes hair growth by stimulating cell proliferation and down regulating TGF-beta1 in C57BL/6 mice in vivo. Growth Horm IGF Res. 2014; 24 (2–3):89–94. https://doi.org/10.1016/j.ghir.2014.03.004 PMID: 24797500.

40. Franke TF, Kaplan DR, Cantley LC. PI3K: downstream AKTion blocks apoptosis. Cell. 1997; 88(4):435–7. Epub 1997/02/21. https://doi.org/10.1016/s0092-8674(00)81883-8 PMID: 9038334.

41. Yang CC, Cotsarelis G. Review of hair follicle dermal cells. J Dermatol Sci. 2010; 57(1):2–11. https://doi.org/10.1016/j.jdermsci.2009.11.005 PMID: 20022473; PubMed Central PMCID: PMC2818774.

42. Yamauchi K, Kurosaka A. Inhibition of glycogen synthase kinase-3 enhances the expression of alkaline phosphatase and insulin-like growth factor-1 in human primary dermal papilla cell culture and maintains mouse hair bulbs in organ culture. Arch Dermatol Res. 2009; 301(5):357–65. Epub 2009/02/25. https://doi.org/10.1007/s00403-009-0929-7 PMID: 19238412.

43. Greco V, Chen T, Rendl M, Schober M, Pasolli HA, Stokes N, et al. A two-step mechanism for stem cell activation during hair regeneration. Cell Stem Cell. 2009; 4(2):155–69. https://doi.org/10.1016/j.stem.2008.12.009 PMID: 19200804; PubMed Central PMCID: PMC2668200.

44. Castellana D, Paus R, Perez-Moreno M. Macrophages contribute to the cyclic activation of adult hair follicle stem cells. PLoS Biol. 2014; 12(12):e1002002. Epub 2014/12/24. https://doi.org/10.1371/journal.pbio.1002002 PMID: 25536657; PubMed Central PMCID: PMC4275176.

45. Philpott MP, Sanders D, Westgate GE, Kealey T. Human hair growth in vitro: a model for the study of hair follicle biology. J Dermatol Sci. 1994; 7 Suppl:S55–72. Epub 1994/07/01. PMID: 7999676.

46. Li W, Man XY, Li CM, Chen JQ, Zhou J, Cai SQ, et al. VEGF induces proliferation of human hair follicle dermal papilla cells through VEGFR-2-mediated activation of ERK. Exp Cell Res. 2012; 318(14):1633–40. https://doi.org/10.1016/j.yexcr.2012.05.003 PMID: 22659165.

47. Bassino E, Gasparri F, Giannini V, Munaron L. Paracrine crosstalk between human hair follicle dermal papilla cells and microvascular endothelial cells. Exp Dermatol. 2015. https://doi.org/10.1111/exd.12670 PMID: 25690790.

48. Cotsarelis G, Millar SE. Towards a molecular understanding of hair loss and its treatment. Trends Mol Med. 2001; 7(7):293–301. Epub 2001/06/27. PMID: 11425637.

49. Xing L, Dai Z, Jabbari A, Cerise JE, Higgins CA, Gong W, et al. Alopecia areata is driven by cytotoxic T lymphocytes and is reversed by JAK inhibition. Nat Med. 2014; 20(9):1043–9. Epub 2014/08/19. https://doi.org/10.1038/nm.3645 PMID: 25129481; PubMed Central PMCID: PMC4362521.

50. Gilhar A, Shalaginov R, Assy B, Serafimovich S, Kalish RS. Alopecia areata is a T-lymphocyte mediated autoimmune disease: lesional human T-lymphocytes transfer alopecia areata to human skin grafts on SCID mice. J Investig Dermatol Symp Proc. 1999; 4(3):207–10. Epub 2000/02/16. PMID: 10674367.

51. Sundberg JP, Pratt CH, Silva KA, Kennedy VE, Stearns TM, Sundberg BA, et al. Dermal lymphatic dilation in a mouse model of alopecia areata. Exp Mol Pathol. 2016; 100(2):332–6. Epub 2016/03/10. https://doi.org/10.1016/j.exmp.2016.03.001 PMID: 26960166; PubMed Central PMCID: PMC4823156.

52. Martin JM, Monteaquido C, Montesinos E, Guijarro J, Llombart B, Jorda E. Lipedematous scalp and lippedematous alopecia: a clinical and histologic analysis of 3 cases. J Am Acad Dermatol. 2005; 52(1):152–6. Epub 2005/01/01. https://doi.org/10.1016/j.jaad.2004.05.016 PMID: 15627100.