**Abstract**

Recent findings regardingDll4 function in physiological and pathological conditions indicate that this Notch ligand may constitute an important therapeutic target. Dll4 appears to be a major anti-angiogenic agent, occupying a central role in various angiogenic pathways. The first trials of anti-Dll4 therapy in mice demonstrated a paradoxical effect, as it reduced tumor perfusion and growth despite leading to an increase in vascular density. This is seen as the result of insufficient maturation of the newly formed vasculature causing a circulatory defect and increased tumor hypoxia. As Dll4 function is known to be closely dependent on expression levels, we envisioned that the therapeutic anti-Dll4 dosage could be modulated to result in the increase of adequately functional blood vessels. This would be useful in conditions where vascular function is a limiting factor for recovery, like wound healing and tissue hypoxia, especially in diabetic patients. Our experimental results in mice confirmed this possibility, revealing that low dosage inhibition of Dll4/Notch signaling causes improved vascular function and accelerated wound healing.

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**Introduction**

Wound healing is a physiological process required for maintenance of an intact skin barrier. Angiogenesis, the growth of new blood vessels, is an important natural process required for healing wounds and for restoring blood flow to tissues after injury or insult [1]. Following a traumatic injury, angiogenesis is initiated by multiple molecular signals, including hemostatic factors, inflammation, cytokine growth factors, and cell-matrix interactions, and is mediated throughout the entire wound-healing process [2,3]. New blood vessels grow via a cascade of biological events to form granulation tissue in the wound bed [4]. This process is sustained until the terminal stages of healing, when angiogenesis is slowed by reduced levels of growth factors, resolution of inflammation, stabilized tissue matrix, and endogenous inhibitors of angiogenesis [2,3]. Defects in the angiogenesis pathway impair granulation and delay healing [4].

Because angiogenesis is required for wound healing, its induction is beneficial in many clinical situations to achieve wound closure, enhance tissue vascularization, improve local circulation, and promote healing and regeneration. This capability is controlled by pro- and anti-angiogenic factors present throughout the body. A precise physiological balance exists between pro-angiogenic and endogenous anti-angiogenic inhibitors, such that vascular growth is normally suppressed. Immediately following injury, however, pro-angiogenic factors are released into the wound bed, and a shift occurs in the balance of regulators favoring vascular growth [2,3,5,6,7].

There are already a few pro-angiogenic therapies available for use in the treatment of wounds. The first drug of this kind was the recombinant human platelet-derived growth factor-BB (rhPDGF-BB), which produces good clinical results [8]. A number of angiogenic growth factors are in pre-clinical and clinical development for wound healing, including vascular endothelial growth factor (VEGF), fibroblast growth factor (FGF), keratinocyte growth factor (KGF), and transforming growth factor-beta (TGF-β). Although PDGF-BB was the first growth factor to become clinically approved for wound care, it is likely that multiple growth factors will eventually become available and that combination therapy may be beneficial [7].

Dll4 is an endothelium-specific Notch ligand [9]. Dll4 is largely an arterial-endothelial-specific gene, especially during embryonic development, while in adults its expression is mostly restricted to small arteries and capillary networks [10,11]. Haploinsufficiency of Dll4 in mice results in embryonic lethality at approximately E10.5 due to defective vascular development [10]. Notably, Dll4 and Vegf
are the only known genes with a haploinsufficient-lethal phenotype due to failure to form a functional vasculature.

Recently, several groups have investigated whether inhibition of the Dll4/Notch pathway might affect tumor angiogenesis and growth. Notch pathway blockade was achieved with either a Dll4-selective neutralizing antibody or a soluble Dll4 fusion protein (sDll4-Fc) that presumably works by binding Notch receptors and preventing their activation by endogenous Dll4 [12,13,14]. These biopharmaceuticals showed robust antitumor activity in a variety of human and rodent tumor xenograft models. Histology of Dll4-Fc treated tumors revealed that the reduced tumor growth was associated with an increase in tumor vascular density. However, labeling with an intravascular tracer found these blood vessels to be poorly perfused. Insufficient blood flow in the treated tumor vessels was also reflected by the increased hypoxia observed in tumors treated with sDll4-Fc. It appears that excessive branching results in a highly chaotic vascular network that lacks the hierarchy essential for efficient directional blood flow [12,13,14].

Notch signaling impact on wound healing has been previously tested [15], revealing that inhibition of generalized Notch signaling leads to delayed wound healing. However the results were not specific for the endothelium because Notch1 is known to be expressed in several cell types involved in skin regeneration, like keratinocytes and immune cells, among others. The endothelial specificity of Dll4 makes it a good potential target for Notch signaling manipulation in the endothelium. Dll4 appears to function as an anti-angiogenic factor, negatively regulating pro-angiogenic factors, such as VEGF, and positively regulating vascular maturation factors, like TGF-β [16,17]. Thus, anti-Dll4 therapy results in increased vascularization with decreased vascular maturation and, therefore, decreased vascular function [18]. The same scenario was observed in the tumor tests [12,13,14]. We assumed that as Dll4 function is very dose-dependent, the effect of this therapy might be adjusted by modulating drug dosage. Here we provide in vivo evidence supporting the use of anti-Dll4 therapy to increase vascularization without decreasing vascular function, improving the rate of wound healing.

Methods

Experimental animals
All animal-involving procedures in this study were approved by the Faculty of Veterinary Medicine of Lisbon Ethics and Animal Welfare Committee (Approval ID: PTDC/CVT/71406/2006). The generation of Dll4<sup>−/−</sup> (Dll4/LacZ) mice on CD1 background has been reported previously [10]. Dll4 conditional knockout mice (Dll4<sup>fl/fl</sup>) have been obtained in the lab, in collaboration with Dr. Freddy Radtke [19]. These were crossed with VE-cadherin-Cre-ERT2 mice, a kind gift by Dr. Ralph Adams, to produce eDll4lox/lox conditional heterozygote mice are Dll4(LacZ) mice on CD1 background, the effect of this therapy might be adjusted by modulating drug dosage. Here we provide in vivo evidence supporting the use of anti-Dll4 therapy to increase vascularization without decreasing vascular function, improving the rate of wound healing.

Tissue preparation and immunohistochemistry
On each end-point mice were sacrificed and wounds were recovered. Biopsies were fixed in 4% paraformaldehyde (PFA) solution at 4°C for 1 h, cryoprotected in 15% sucrose, embedded in 7.5% gelatine, frozen in liquid nitrogen and cryosectioned at 20 μm. Double fluorescent immunostaining to platelet endothelial cell adhesion molecule (PECAM) and pericyte marker alpha smooth muscle actin (α-SMA) was performed on tissue sections to examine wound vascular density and vessel maturity. Rat monoclonal anti-mouse PECAM (BD Pharmingen, San Jose, CA) and rabbit polyclonal anti-mouse α-SMA (Abcam, Cambridge, UK) were used as primary antibodies and species-specific conjugated with Alexa Fluor 488 and 555 (Invitrogen, Carlsbad, CA) were used as secondary antibodies. Tissue sections were incubated with primary antibody overnight at 4°C and secondary antibody for 1 hour at room temperature. Nuclei were counterstained with 4’, 6-diamidino-2-phenylindole dihydrochloride hydrate (DAPI; Molecular Probes, Eugene, OR). Fluorescent immunostained sections were examined under a Leica DMRA2 fluorescence microscope with Leica HC PL Fluotar 10 and 20X/0.5 NA dry objective, captured using Photometrics CoolSNAP HQ, (Photometrics, Friedland, Denmark), and processed with ImageJ 1.37v program. Vessel density corresponds to the percentage of each tissue section field occupied by a PECAM-positive signal (as determined by the percentage of white pixels per field after transforming the RGB images into binary files). As a measure of vascular maturity, mural cell recruitment was assessed by quantitating the percentage of PECAM-positive structures lined by α-SMA-positive coverage.

Vascular perfusion and extravasation
To mark vessel perfusion, tribromoethanol anesthetized mice were injected with biotin-conjugated lectin from Lycopersicon esculentum (100 μg in 100 μl of PBS; Sigma, St. Luis, MO) via caudal vein and allowed to circulate for 3 minutes before vessel
perfusion with 4% PFA in PBS for 3 minutes. Wound samples were collected and processed as presented above. Tissue sections (20 μm) were stained with rat monoclonal anti-mouse PECAM (BD Pharmingen, San Jose, CA), followed by Alexa 555 goat anti-rat IgG (Invitrogen, Carlsbad, CA). Biotinylated lectin was visualized with Streptavidin-Alexa 488 (Invitrogen, Carlsbad, CA). The images were obtained and processed as described above. Wound perfusion was quantified by determining the percentage of PECAM-positive structures that were colocalized with Alexa 488 signals. To analyze vascular extravasation tribromoethanol anesthetized mice were injected with 1% Evans Blue solution (Sigma, St. Luis, MO) via caudal vein and allowed to circulate for 5 minutes before transcardial vessel perfusion with 4% PFA in PBS for 3 minutes. Evans Blue is red fluorescent and extravasation was visualized in contrast to green fluorescent vascular structures and compared between groups [20].

Quantitative transcriptional analysis
Biopsies were recovered by dissecting wound tissue from experimental mice and snap frozen in liquid nitrogen until RNA extraction (Qiagens RNAeasy). Using a SuperScript III First Strand Synthesis Supermix qRTPCR (Invitrogen, Carlsbad, CA), first-strand cDNA was synthesized from total RNA. Real-time PCR analysis was performed as described [16], using specific primers. Primer pair sequences are available on request. Gene expression was normalized to β-actin.

Statistical analyses
Data processing was carried out by engaging Statistical Package for the Social Sciences version 17.0 (SPSS v. 17.0; Chicago, IL). Statistical analyses were performed using Mann-Whitney-Wilcoxon test. All results are presented as mean ± SEM. P-values < 0.05 and < 0.01 were considered significant (indicated in the Figures with *) and highly significant (indicated with **), respectively.

Results
Dll4 signaling inhibition leads to antagonistic effects on wound healing
The study of the impact of Dll4 signal modulation was initially carried out using genetic mouse models of Dll4 loss- and gain-of-function. Wound healing assays on an endothelial-specific Dll4 overexpressing mouse line (eDll4OE) and an endothelial-specific Dll4 conditional knockout mouse line (eDll4lox/lox) were anticipated to provide some insight on the role of Dll4 in regulating adult neoangiogenesis. We performed daily wound area measurements and evaluated vascular density in the granulation tissue of the wound (Fig. 1A). Wound healing was impaired in both eDll4OE and eDll4lox/lox mice (Fig. 1B, D). Significantly delayed wound healing was observed in eDll4OE mice as early as day 1 and was maintained until day 4, when wounds were reduced to around 10% of their original size. Over a 5-day period, there was a 1 to 1.5 days delay in wound healing in eDll4OE mice compared to controls. Histological analysis on days 2 and 5 revealed that eDll4OE mice had approximately 40% lower vascular density in the wound region than controls (Fig. 1C). Absolute vascular density values increased over the period of regeneration in both groups but the difference between groups was maintained or slightly increased (data not shown). Wound healing was also delayed in eDll4lox/lox mice (Fig. 1D), which was statistically significant at day 2 and persisted. Overall there was a 2 to 3 days delay compared to control group over a 7-day period. Histological analysis on days 2, 4 and 7 revealed a large increase in vascular density throughout the experimental period in eDll4lox/lox mice (Fig. 1E), consistent with the previous report that Dll4 loss promotes angiogenesis [12,21].

In contrast to the initial findings, wound-healing assays on Dll4+/− mice revealed that regeneration was improved when compared with control Dll4+/+ mice (Fig. 1F). Differences in wound size became statistically significant by day 2 and persisted until day 6. From day 2 until day 6 the wound healing in Dll4+/− mice presented a steady 2-day advance over control mice. Histological analysis revealed an increase in vascular density that could be seen in Dll4+/− mice as early as day 2 (Fig. 1G). This difference in vascular density was maintained throughout the wound recovery period, even though absolute vascular density values increase in both experimental groups (data not shown).

To confirm that the improvement in wound healing observed in Dll4+/− mice was due to the change in vasculature, and not other factors such as the inflammatory response [22], further experiments with endothelial-specific conditional Dll4 heterozygotes (Dll4+/loxVE-Cadherin-CreERT2 - from now on Dll4+lox/2) were carried out. Results revealed that in Dll4+lox/2 mice wounds regenerated faster than in the respective controls, in a similar fashion to what was described for Dll4+/− mice. A two-day advance in wound closure of Dll4+/− was established by day 2 and was maintained until the endpoint (Fig. 2A). Vascular density evaluation confirmed identical differences in relation to wild type controls in both Dll4+lox/2 and Dll4+/− mice (Fig. 2B,C). Pro-inflammatory gene expression at day 2, during the inflammatory phase of wound closure, was tested in both Dll4+/− and Dll4+lox/2 mice. This would allow the identification of a possible influence of Dll4 function in mediating the inflammatory response independent of the vascular phenotype. This possible effect was tested by RT-PCR analysis of wound biopsies to evaluate the expression of pro-inflammatory genes. Results showed that monocyte/macrophage chemo attractant MCP1 had reduced expression in both Dll4+lox/2 and Dll4+/− mice. Pro-inflammatory genes, such as ICAM, VCAM and MIP2, were also downregulated in both Dll4+lox/2 and Dll4+/− mice. Markers of macrophage activation iNOS, PTX3 and Id1 had reduced expression in both Dll4+lox/2 and in Dll4+/− mice (Fig. 2D). We then evaluated the gene expression profile of the same inflammation-related genes in the Dll4 mutant mice that showed impaired regeneration profile, Dll4+/lox/2 and Dll4+/−, as this would allow us to correlate changes in the inflammatory profile of wounds to their regeneration profile. Results showed that the expression of pro-inflammatory genes was upregulated in both Dll4+lox/2 and Dll4+/− mice (Fig. S1).

sDll4-Fc therapy has antagonistic effects on wound healing depending on dosage
Inflammatory gene expression in Dll4+/lox/2 indicated that Dll4 function in the endothelium was the most important factor accounting for the observed improvements in wound closure. Dll4+/lox/2 (and Dll4+/−) and Dll4+/lox/2 mice can give rise to opposing phenotypes despite both being loss-of-function mutants and both displaying a pro-angiogenic phenotype. We therefore proposed that dosage of the inhibitor (soluble Dll4-Fc) may mimic the Dll4 dose response observed in Dll4+ deficient mouse lines allowing us to define the dosage of sDll4-Fc that promotes wound healing. sDll4-Fc therapy was tested in C57BL/6 mice using dosages from 0.025 mg/kg to 2.5 mg/kg. Mice were injected on day 0, after wounding, and every 2 days until the endpoint. Lower dosages, like 0.025 mg/kg, 0.05 mg/kg and 0.1 mg/kg, were observed to accelerate wound healing (Fig. 3A). Statistical significance in wound size difference was achieved as early as day 1 in the 0.05 mg/kg dosage group and day 2 in the 0.025 mg/
Inhibition of Dll4 Promotes Wound Healing

B

Endothelial-specificDll4 overexpression

Average size of wound relative to Day 0 (%)

Days after wound

C

Relative vascular density (%)

Day2
Day5

D

Endothelial-specificDll4 null

Average size of wound relative to Day 0 (%)

Days after wound

E

Relative vascular density (%)

Day2
Day4
Day7

F

Constitutive Dll4 heterozygous

Average size of wound relative to Day 0 (%)

Days after wound

G

Relative vascular density (%)

Day2
Day4
Day6

H
kg and 0.1 mg/kg dosage groups and was maintained throughout the regeneration period. From day 2 until day 5 a 1 to 1.5 day advantage in wound size was observed in treated mice over control mice, with the 0.05 mg/kg dosage group displaying the largest benefit. Testing with higher dosages, such as 0.5 mg/kg, 1 mg/kg and 2.5 mg/kg revealed an opposite effect delaying wound healing (Fig. 3B). Statistical significance in wound size difference was observed by day 1 in the 1 mg/kg and 2.5 mg/kg dosage groups and maintained until endpoint. Dosage 0.5 mg/kg was considered to originate wound healing statistically indistinguishable from that of control mice. Dosage 1 mg/kg displayed a 0.5 to 1 day delay in wound healing, while the 2.5 mg/kg dosage displayed a 1 to 1.5 day delay, over a 6-day period. To understand how opposing effects on wound regeneration were generated by modulating sDll4-Fc therapy dosage we analyzed vascular density on days 2, 4 and 6. This revealed that in all dosage groups there was increased vascular density, the degree of which was in direct relationship with the increase in dosage (Fig. 4 A,C,D,E). Dosages that led to improved wound healing caused increases in vascular density of 20 to 50%, while dosages that led to impaired wound healing caused much larger increases of 70 to 300%.

Neo-vasculature in sDll4-Fc treated mice is less mature but has variable perfusion and extravasation depending on dosage

Normal neo-vascularization involves the proliferation and progression of pre-existing endothelium into stressed areas. That newly formed vasculature has to mature, becoming more stable. This is achieved mainly by inducing the differentiation and recruitment of smooth muscle cells and pericytes. We analyzed the wound basin neo-vasculature of mice treated with 0.05 mg/kg and 2.5 mg/kg of sDll4-Fc, the dosages that led to the best and worst result in wound healing, and the respective control, for the presence of vascular smooth muscle cells. While PBS-injected mice had a normal coverage of smooth muscle, sDll4-Fc injected mice had a 3-fold reduced coverage in both doses (Fig. 4B, C, D and E). This effect became evident as early as Day 2 and persisted throughout wound regeneration. By comparison, all Dll4 loss-of-function mutants analyzed in this work produced similar results to those obtained with sDll4-Fc showing reduced presence of perivascular cells, while Dll4OE mutants displayed increased recruitment of perivascular cells (Fig. S2).

Since one of the hallmark observations after sDll4-Fc therapy on tumors was the poor perfusion of newly formed vasculature [12], lectin perfusion (Fig. 5A,B) and Evans Blue vascular extravasation (Fig. 5C) testing were carried out in the wound healing setting. The 0.05 mg/kg dosage revealed no significant improvement in the percentage of lectin perfused blood vessels, and a low level of vascular extravasation, relative to control. The 2.5 mg/kg dosage revealed much-decreased lectin perfusion and very pronounced Evans Blue extravasation. These results confirm that the impaired maturation of the neo-vasculature in sDll4-Fc treated mice is dosage-dependent and that the effectiveness of blood circulation to affected tissues varies directly with the degree of neo-vasculature maturation. For comparison purposes, Dll4lox/lox and Dll4OE neovasculature had a lectin perfusion index that was not statistically different to their respective controls, and slightly higher Evans Blue extravasation than controls. Dll4 mutants with impaired wound healing profile, Dll4lox/lox and Dll4OE, both revealed reduced lectin perfusion, while Dll4lox/lox displayed a very high level of vascular extravasation and Dll4OE showed almost normal level of vascular extravasation (Fig. S3A, B).

Dll4 modulation has no effect on the skin adjacent to the wound

We have described that sDll4-Fc therapy can affect the vasculature of wounds and therefore, influence the regenerative profile. Since our approach is not targeted to the affected region we decided to investigate if the vasculature of skin adjacent to the wound site would be equally affected by sDll4-Fc therapy in either high- or low-dosage form. PECAM and SMA immunofluorescence (Fig. 6A–F) revealed that outside the wound the vasculature was largely unaffected during the therapy period. Vascular density and smooth muscle cell coverage indexes of the skin adjacent to the wound in mice treated with 0.05 mg/kg and 2.5 mg/kg sDll4-Fc did not differ statistically at endpoint from PBS-treated mice (Fig. 6G,H). Microscopic analysis of surrounding tissues also did not reveal any histological changes to control (Fig. 6I–K).

Effect of Dll4 modulation on the expression of vascular genes

To help the identification of the mechanistic causes behind the observed phenotypes the expression of selected genes in the wound area was characterized by RT-PCR. RNA was extracted from wound biopsies taken from both genetic models and sDll4-Fc treated mice. Expression levels were normalized to PECAM1 mRNA levels [23] to compensate for variations in vascular density between samples. In the genetic models (Fig. 7A), Dll4 expression levels corresponded to the expectation, with a 4-fold increase in the Dll4OE mice and a reduction to 0.5 x in the Dll4OE mice from 0.13 x in the Dll4lox/lox mice. Jagged1 expression levels decreased in the loss-of-function models and increased in the Dll4OE mice. Hey2, the main known Notch signaling effector in the endothelium [24], displayed higher expression levels in Dll4OE and lower expression levels in the loss-of-function models, confirming that the Notch pathway is being affected by the induced Dll4 mutations. Hey1 gene expression levels were found to vary similarly to Hey2 except that the differences to respective controls were smaller. Rbp-J is the principal mediator of NICD function in the regulation of Notch effector genes expression regulation [25]. Rbp-J expression levels were found to be upregulated in Dll4OE and downregulated in the Dll4 loss-of-function models. Ephrin-B2 is known to be downstream of Notch signaling in embryonic
Inhibition of Dll4 Promotes Wound Healing

A

Endothelial-specific Dll4 heterozygous

Average size of wound relative to Day 0 (%)

Days after wound

B

Relative vascular density (%)

Control

eDll4+/lox

Day2
Day4
Day6

C

eDll4+/lox Control

eDll4+/lox

D

Relative gene expression (Dll4+/− vs Control & eDll4+/lox vs Control)

Dll4 INOS PTX3 Id1 MIP2 MCP1 ICAM VCAM

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vascular development and is a regulator of vascular maturation and endothelial identity [26]. Ephrin-B2 expression was decreased in Dll4+/lox and eDll4lox/lox mice and increased in eDll4OE mice, corroborating the vascular maturation results obtained by immunofluorescence. EphB4 encodes the venous-specific cognate receptor for Ephrin-B2 [27]. EphB4 expression was upregulated in

Figure 2. The wound regeneration profile of endothelial specific Dll4 heterozygous mice (eDll4+/lox) is similar to that of constitutive Dll4 heterozygous mice (Dll4+/−) and both share a reduced inflammatory response when compared to Dll4 WT mice. A) Comparison of eDll4+/lox mice with uninduced controls. Graphic depicting the correlation between wound areas in each experimental day relative to the wound area measured on Day 0. Wound regeneration is accelerated in eDll4+/lox mice in a similar way as in the Dll4+/− mice. B) Vascular density is increased in granulation tissue of eDll4+/lox mice, relative to uninduced controls throughout the experiment. C) Representative anti-PECAM immunofluorescence images of neo-vasculature in granulation tissue of eDll4+/lox and uninduced controls at endpoint. D) Differential gene expression in Dll4+/− versus WT and eDll4+/lox versus uninduced control wounds of inflammation-related genes at day 2. Dll4 expression is confirmed to be approximately 0.5-fold in both cases, relative to their respective controls, and statistically similar. Inflammation-related genes have reduced expression in both mutant mice, probably resulting from the improved condition of the mutant mice wounds that regenerate faster. * In graphics represents p<0.05.

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Figure 3. Modulation of sDll4-Fc dosage can achieve both improved or impaired wound regeneration through low- or high-dosage inhibition. A) Graphic depicting the correlation between wound areas in experimental days relative to wound areas measured on Day 0, comparing mice treated with sDll4-Fc dosages lower than 0.1 mg/kg with untreated (sham treated) controls. Wound regeneration is accelerated in mice treated with sDll4-Fc dosages lower than 0.1 mg/kg. B) Graphic depicting the correlation between wound areas in experimental days relative to wound areas measured on Day 0, comparing mice treated with sDll4-Fc dosages higher than 0.5 mg/kg with controls. Wound regeneration is delayed in mice treated with sDll4-Fc dosages higher than 0.5 mg/kg.

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eDll4lox/lox and downregulated in Dll4OE mice. Dll4 is a known regulator of VEGF signaling, having specific control over the expression of its transmembrane receptors in the endothelium [28]. Vegf-a expression levels were decreased in Dll4lox/lox mice, while they were increased in the eDll4lox/lox and Dll4OE mice, which had delayed wound regeneration, probably accounting for reduced hypoxia in wounds with improved regeneration. Dll4 is a known suppressor of VEGF-C/VEGFR3 signaling [29]. The expression of Vegf-c in the wounds was augmented in both Dll4lox/lox and eDll4lox/lox mice but in Dll4OE mice the difference was found to be not significant. VEGF receptor expression analysis revealed that eDll4OE mice had lower expression of Vegfr2 and Vegfr3, both of which mediate VEGF signaling, and higher expression of Vegfr1, which acts mainly as a VEGF trap [30] and blocks VEGF-A mediated pro-angiogenic signal. In the loss-of-function models VEGFRs expression varied in the opposite direction, with decreased Vegfr1 expression and increased Vegfr2 and Vegfr3 expression and thus increased angiogenic potential. Tie2 loss-of-function is associated with a decrease in smooth muscle cell recruitment [31]. Expression of Tie2 in our genetic models reflects this effect, being lower in both Dll4lox/lox and Dll4OE mice, which had less smooth muscle, and increased in Dll4OE mice, which had an higher than normal number of smooth muscle cells. Lyve1 is a known lymphatic endothelial cell-specific gene [32]. Recently, Dll4 blockade was associated with disruption of post-natal lymphatic development [33]. Lyve1 expression was downregulated in eDll4lox/lox mice and upregulated in Dll4OE. More importantly Lyve1 expression in Dll4+/lox mice, which displayed accelerated wound healing, was found to be near normal levels.

In the mice subjected to sDll4-Fc therapy (Fig. 7B), endogenous Dll4 expression decreased with dosages up to 0,1 mg/kg and increased with dosages greater than 0,5 mg/kg, reflecting Vegf-a expression levels, one of the main inducers of Dll4 expression. Hey2 expression levels confirmed the inhibitory effect of sDll4-Fc on Notch signaling, being lower in all dosages and showing a strong dependency on the dosage of the inhibitor, again confirming that

Figure 4. Wounds treated with sDll4-Fc have increased vascular density but decreased vascular maturation in direct proportion to the dosage level. A) Vascular density is increased in granulation tissue of mice treated with sDll4-Fc, being directly proportional to the dosage increase, when compared to controls throughout the experiment. B) Vascular smooth muscle cell coverage is decreased about 3-fold in relation to PBS-injected control mice throughout the experimental days in both tested groups. C,D,E) Representative anti-PECAM and anti-SMA immunofluorescence images of neo-vasculature in granulation tissue of wounds treated with (D) 0,05 mg/kg or (E) 2,5 mg/kg compared with control mice injected with PBS (C), in days 2, 4 and 6. sDll4-Fc therapy leads to decreased smooth muscle cell recruitment and increased vascular density that is directly proportional to the administered dosage. * In graphics represents p<0,05.

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higher dosages of sDll4-Fc lead to increasing inhibition of Dll4/Notch signaling. 

**Figure 5. Low-dosage sDll4-Fc does not affect perfusion levels and leads to only a small increase in extravasation while higher dosages cause decreased perfusion and highly increased extravasation.**

A) Representative anti-PECAM and anti-Lectin immunofluorescence images of neo-vasculature in granulation tissue of wounds treated with 0,05 mg/kg or 2,5 mg/kg compared with control mice injected with PBS, in day 6. B) The percentage of lectin-perfused blood vessels is similar between PBS- and 0,05 mg/kg sDll4-Fc injected mice but since vascular density is increased in the latter this result represents an effective increase in vascular function in the wound basin. Vascular perfusion is highly decreased in wounds of mice injected with 2,5 mg/kg sDll4-Fc. C) Representative anti-PECAM and Evans Blue immunofluorescence images of neo-vasculature in granulation tissue of wounds treated with 0,05 mg/kg or 2,5 mg/kg compared with control mice injected with PBS, in day 6. Vascular extravasation is highly increased in wounds of mice injected with 2,5 mg/kg sDll4-Fc but displays only a small increase in the case of 0,05 mg/kg sDll4-Fc injected mice. * In graphics represents p<0.05.

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In regard to the VEGF receptors, expression of Vegfr1 decreased (in indirect relation to sDll4-Fc dosage) and Vegfr2 and Vegfr3 increased in direct relation to sDll4-Fc dosages. TIE2 expression levels decreased in all sDll4-Fc dosages, showing an inverse relation to sDll4-Fc dosages. Lyve1 expression values were always downregulated, except at the 0,05 mg/kg sDll4-Fc dosage, with expression decreasing with increasing sDll4-Fc dosages.
Discussion

The importance of Dll4 was first described by its function in the establishment of endothelial identity and in the regulation of vascular morphogenesis in embryonic vascular development. Experimental data place Dll4 in the middle of a loop of angiogenesis regulatory pathways. Dll4 was found to be one of the key players acting as negative regulator of angiogenesis, working to maintain the fine balance between vascular progression and vascular maturation/quiescence [34]. This important function has led to the development of therapies blocking Dll4 function for tumor therapy. Dll4 inhibition, and sDll4-Fc therapy in particular, have shown potential in controlling tumor growth, even in cases when tumors have gained resistance to anti-VEGF therapies [12,13]. The mechanism of action of this therapy is strikingly different to that of classic anti-angiogenic therapies, which work by reducing the ability of a tumor to induce the growth of new blood vessels or by reducing the ability of blood vessels to sprout and form the neo-vasculature in the tumor. Dll4 inhibition works by increasing vascular density through uncontrolled growth, resulting in the new vasculature being disorganized, inefficient and poorly perfused, leading to an increase in tumor hypoxia and reduction in tumor growth [35].

As the embryonic lethal haploinsufficiency of Dll4 is indicative of a strong functional dosage-dependence, we hypothesized that the non-functional nature of the tumoral neo-vasculature observed in anti-Dll4 therapy [12,21] was a consequence of the high dosage of Dll4 inhibitor used and that by modulating therapy dosage it might be possible to achieve a functional pro-angiogenic effect. Dll4-based therapy could then be used to treat conditions where the formation of new blood vessels is essential and beneficial, like wound healing. In this case, angiogenesis starts almost immediately following injury and is important for the formation of granulation tissue, to supply oxygen and nutrients to growing tissues and as a means to deliver inflammatory cells to the wound site [4].

Wound healing was tested on murine genetic models of Dll4 gain- and loss-of-function. Results revealed that Dll4 heterozygote mice (both Dll4+/− and Dll4+/−/lox) presented accelerated wound regeneration with improved vascular density as well as near-normal percentage of perfused blood vessels, whereas Dll4+/lox and Dll4OE presented delayed wound healing. In the first case it was related to highly increased vascular density and in the latter related to a reduction in vascular density, both phenotypes leading to reduced tissue perfusion. Since improved wound regeneration observed in the Dll4+/− mice was caused by a constitutive mutation and previous works described a possible link between Dll4 and inflammatory macrophages [22], we decided to test if an equivalent endothelial-specific mutation could give rise to the same phenotype. Results with Dll4+/lox mice showed a similar wound regeneration profile to that of Dll4+/− mice. This indicates that at least the principal contributor to the observed phenotype was the loss of Dll4 function in the endothelium. Nevertheless we measured by RT-PCR the expression levels of several regulators of inflammatory function. The expression profile indicated that in both Dll4+/− and Dll4+/lox there is a reduction in the expression of macrophage chemoattractants, regulators of monocyte/macrophage infiltration and markers of macrophage activation. Since the results are not statistically different between Dll4+/lox and Dll4+/− we can infer that the differences in the inflammatory profile of both mutant mouse lines is unrelated to Dll4 expression.

The gene expression levels of the same inflammatory profile markers are increased in both the Dll4+/lox and Dll4OE mice with opposing Dll4 mutations and delayed wound regeneration. So, the
observed inflammatory profiles are likely to be a consequence of an improved or impaired wound regeneration status, not a specific response to the Dll4 mutations. These results confirmed that different levels of Dll4 blockade give rise to either an improvement or impairment in wound regeneration, depending on whether the increased angiogenesis leads to functional or non-functional blood vessels.

The next step was to test different dosages of sDll4-Fc therapy in surgically wounded wild-type mice. Results showed that lower dosages, between 0.025 mg/kg and 0.1 mg/kg accelerate wound regeneration by inducing productive angiogenesis that is functional and causes an improvement in blood flow in the wound area without affecting the surrounding quiescent vasculature. Comparing results from different dosages, 0.025 mg/kg, 0.05 mg/kg and...
0.1 mg/kg revealed that while all dosages tested gradually improved vascular density, the transition from functional to non-functional angiogenesis appears to occur when the dosage increases from 0.05 to 0.1 mg/kg. This is based on the observation that wound healing is accelerated when the dosage increases from 0.025 to 0.05 mg/kg, as a result of increased vascular function, but slowed down when it is further increased from 0.05 to 0.1 mg/kg. This indicates that the transition from functional to non-functional angiogenesis is a result of vascular inefficiency arising from increasing vascular density and inability to form proper blood vessel lumens and perivascular wall. With higher dosages, ranging from 0.5 to 2.5 mg/kg of sDll4-Fc, wound regeneration was progressively delayed over control mice, as increasingly higher vascular densities led to decreased vascular maturation, decreased perfusion and increased vascular extravasation and therefore reduced vascular function. Most importantly, in sDll4-Fc treated mice the vasculature of the skin adjacent to the wound site remained unaffected when compared to PBS-treated mice. This result indicates that sDll4-Fc therapy specifically targets sites of active angiogenesis, where Dll4 function is most important to regulate vascular morphogenesis. A recent work [36] described that prolonged (over 8 weeks long) therapy with high dosages of Dll4 inhibitors (much higher than any of the dosages used in this work) can lead to low frequency non-lethal subcutaneous vascular neoplasms and histopathological changes in the liver. We have not observed subcutaneous tumors in our sDll4-Fc treated mice or Dll4 mutant mice for the duration of the experiments. It is possible that the lesions cited are also dosage-dependent or due to long term Dll4 blockade. Our results indicate that at low dosages and for short term therapy the risk of such secondary effects should be minimal.

Gene expression analysis confirmed the existence in the wound healing process of a regulatory loop between VEGF signaling and Dll4/Notch signaling, where Dll4/Notch signaling blockade leads to a reduction in Vegfr1 and an increase in Vegfr2 and Vegfr3 expression, increasing vascular response to VEGF-A and –C expression from surrounding stroma [29,37]. Jagged1 is a member of the Notch ligand family and is present in arterial endothelium and vascular smooth muscle cells [39]. Jagged1 expression was found to be directly related to Dll4 function, which could reveal a novel regulatory feedback loop between Dll4 and Jagged1 where increasing Dll4 function potentiates Jagged1 expression that then negatively regulates Dll4/Notch signaling [24]. Ephrin-B2 is involved in the activation of angiogenesis and recruitment of perivascular cells to neo-vasculature [36,39], acting downstream of VEGF and Notch signaling while at the same time acting as a regulator of VEGF-R2 and –R3 internalization and function [40,41]. It is expressed in arterial endothelium and in smooth muscle cells. Ephrin-B2 expression is found to be directly related to Dll4 function and could indicate one way through which Dll4/Notch signaling regulates vascular smooth muscle cell recruitment. EphB4 expression is opposite to Ephrin-B2 expression and increases with sDll4-Fc dosage. This could also indicate that higher dosages of sDll4-Fc lead to a shift to venous neo-vasculature, while lower dosages have no impact on arterial-venous proportion. The Tie2 receptor is considered to be endothelial-specific, while its cognate ligand Ang1 is expressed mainly by perivascular and mural cells, acting as a paracrine signal to the endothelium. Loss-of-function studies revealed a phenotype with defective angiogenesis and reduced coverage and detachment of smooth muscle cells and pericytes [42]. sDll4-Fc therapy leads to a dosage-dependent reduction in Tie2 expression, demonstrating the Ang1/Tie2 pathway involvement in the regulation of vascular maturation and quiescence downstream of Dll4/Notch signaling. It was shown recently that Dll4 blockade could inhibit lymphatic vessel assembly [33]. Our expression results revealed that while that is true for high dosage Dll4 inhibition, in the 0.05 mg/kg sDll4-Fc dosage the reduction in Evans expression, a lymphatic endothelial cell marker is not significantly different to the control.

A role for Notch signaling in wound healing [15] has previously been suggested by use of a Notch1 knockdown mouse line as well as small molecule inhibitor of Notch signaling (GSI treatment). Those results indicated that Notch1 inhibition led to a delay in wound regeneration but failed to report on the wound vascular phenotype. Besides endothelial cells, Notch1 is present in skin, at least in keratinocytes as well as in various immune cells, making it difficult to separate the impact of the mutation in the different cell systems and how they integrate to display a given phenotype.

Together, our results present evidence that endothelial Notch signaling is a potential target for wound regeneration therapies through its ligand Dll4. Low-dosage sDll4-Fc accelerates wound regeneration by creating a neo-vasculature that has a slight increase in vascular density while keeping nearly normal perfusion of the blood vessels and arterial-venous phenotype, without affecting lymphatic vasculature. Overall low-dosage sDll4-Fc therapy leads to an effective increase in local blood supply to the wound site, not affecting lymphatic drainage or the quiescent vasculature.

Supporting Information

Figure S1 Dll4 mutants with impaired wound regeneration share a pro-inflammatory profile. Differential gene expression in Dll4OE and Dll4fl/fl mutants versus respective uninjected control wounds of inflammation-related genes at day 2. Inflammation-related genes have upregulated expression in both mutant mice, probably resulting from the impaired condition of the mutant mice wounds that regenerate slower. * In graphics represents p<0.05. (TIF)

Figure S2 All Dll4 loss-of-function mutants tested displayed reduced perivascular cell recruitment, while Dll4OE mice revealed an opposite phenotype. * In graphics represents p<0.05. (TIF)

Figure S3 A) The percentage of lectin-perfused blood vessels is not statistically different to control levels in Dll4fl/fl− and Dll4fl/fl+, while being highly reduced in Dll4fl/fl+ and Dll4OE. B) Evans Blue extravasation is slightly increased in Dll4 heterozygote models, highly increased in Dll4fl/fl+ and near normal levels in Dll4OE. * In graphics represents p<0.05. (TIF)

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Author Contributions

Conceived and designed the experiments: AT PSG AD. Performed the experiments: AT DJJG MB AP AF. Analyzed the data: AT LI RL VK PSG AD. Contributed reagents/materials/analysis tools: RL VK. Wrote the paper: AT RL VK PSG AD.
References

1. Karamysheva AF (2008) Mechanisms of angiogenesis. Biochemistry (Moscow) 73: 751–762.
2. Eming SA, Brachvogel B, Oderski T, Koch M (2007) Regulation of angiogenesis in wound healing as a model. Prog Histochem Cytochem 42: 115–170.
3. Li J, Zhang YP, Kirchner RS (2003) Angiogenesis in wound repair: angiogenic growth factors and the extracellular matrix. Micros Res Tech 60: 107–114.
4. Gurtner GC, Werner S, Barrandon Y, Longaker MT (2008) Wound repair and regeneration. Nature 453: 314–321.
5. Maleduddu P (2005) Therapeutic angiogenesis and vascularization for tissue regeneration. Exp Physiol 90: 315–326.
6. Tongers J, Bricarelli JG, Luscher TF (2008) Therapeutic angiogenesis for critical limb ischemia: microvascular therapies coming of age. Circulation 118: 9–16.
7. Franz MG, Steed DL, Robson MC (2007) Optimizing healing of the acute wound by minimizing complications. Curr Probl Surg 44: 691–763.
8. Balifour JAB, Noble S (1999) Becaplermin. BioDrugs 11: 339–364.
9. Shutter JR, Scully S, Fan W, Richards WG, Kitajewski J, et al. (2000)Dll4, a novel Notch ligand expressed in arterial endothelium. Genes Dev 14: 1313–1318.
10. Duarte A, Hirashima M, Benedeto R, Trindade A, Diniz P, et al. (2004) Dosage-sensitive requirement for mouse Dll4 in artery development. Genes Dev 18: 2474–2478.
11. Benedeto R, Duarte A (2003) Expression of Dll4 during mouse embryogenesis suggests multiple developmental roles. Gene Expr Patterns 3: 750–753.
12. Schuetz JS, Jiang W, Kumar SR, Krasnoperov V, Trindade A, et al. (2007) Inhibition of Dll4-mediated signaling induces proliferation of immature vessels and results in poor tissue perfusion. Blood 109: 4753–4760.
13. Noguera-Troise I, Daly C, Papadopoulos NJ, Coetzee S, Boland P, et al. (2006) Blockade of Dll4 inhibits tumour growth by promoting non-productive angiogenesis. Nature 444: 1032–1037.
14. Ridgway J, Zhang G, Wu Y, Stawicki S, Liang WC, et al. (2006) Inhibition of Dll4 signalling inhibits tumour growth by deregulating angiogenesis. Nature 440: 1083–1087.
15. Chigurupati S, Arumugam TV, Son TG, Lathia JD, Jammie S, et al. (2007) Involvement of notch signaling in wound healing. PLoS ONE 2: e1167.
16. Trindade A, Ram Kumar S, Schuetz JS, Lopes-Costa L, Becker J, et al. (2008) Overexpression of delta-like 4 induces arterialization and attenuates vessel formation in developing mouse embryos. Blood 112: 1720–1729.
17. Williams CK, Li JL, Murga M, Harris AL, Tosato G (2006) Up-regulation of the Notch ligand Delta-like 4 inhibits VEGF-induced endothelial cell function. Blood 107: 931–939.
18. Thurstorn G, Ngouega-Troise I, Yancopoulos GD (2007) The Delta paradox: Dll4 blockade leads to more tumour vessels but less tumour growth. Dev Cell 13: 295–306.
19. Koch U, Fiererin E, Benedetto R, Besseyrias V, Schuster-Gossler K, et al. (2008) Delta-like 4 is the essential, non-redundant ligand for Notch1 during thymic T cell lineage commitment. J Exp Med 205: 2513–2523.
20. Gratton JP, Lin MI, Wu J, Weiss ED, Jiang ZL, et al. (2003) Selective inhibition of tumor microvascular permeability by cavtretin blocks tumor progression in mice. Cancer Cell 4: 31–39.
21. Djokovic D, Trindade A, Gigante J, Badenes M, Silva L, et al. (2010) Combination of Dll4/Notch and Ephrin-B2/EphB4 targeted therapy is highly effective in disrupting tumor angiogenesis. BMC Cancer 10: 64–111.
22. Fung E, Tang SM, Cammer JP, Morishige K, Arboleda-Velasquez JF, et al. (2007) Delta-like 4 induces notch signaling in macrophages: implications for inflammation. Circulation 115: 2948–2956.
23. Segarra M, Williams CK, Sierra-Mdr L, Bernardo M, McCormick PJ, et al. (2008) Dll4 activation of Notch signaling reduces tumor vascularization and inhibits tumor growth. Blood 112: 1904–1911.
24. Williams CK, Li JL, Murga M, Harris AL, Tosato G (2006) Up-regulation of the Notch ligand Delta-like 4 inhibits VEGF-induced endothelial cell function. Blood 107: 931–939.
25. Borrega J, Oswald F (2009) The Notch signaling pathway: transcriptional regulation at Notch target genes. Cell Mol Life Sci 66: 1631–1646.
26. Mosch B, Reissnerweber B, Neuber C, Pietzch J (2010) Eph receptors and ephrin ligands: important players in angiogenesis and tumor angiogenesis. Journal of Oncology 2010: 135203.
27. Erber R, Eichelbacher U, Powsilaj V, Korn T, Djenov V, et al. (2006) EphB4 protects blood vascular morphology during postnatal angiogenesis. EMBO J 25: 629–641.
28. Jakobsson I, Benley K, Gerhardt H (2009) VEGFRs and Notch: a dynamic collaboration in vascular patterning. Biochem Soc Trans 37: 1253–1256.
29. Hogan BM, Herpers R, Witte M, Holotera H, Altaloo K, et al. (2009) VEGf/Eph signalling is suppressed by Dll4 in developing zebrafish intersegmental arteries. Development 136: 4001–4009.
30. Shibuya M (2008) Vascular endothelial growth factor-dependent and -independent regulation of angiogenesis. BMB Rep 41: 278–286.
31. Thomas M, Augustin HG (2009) The role of the Angiopoietins in vascular morphogenesis. Angiogenesis 12: 125–137.
32. Butler MG, Isogai S, Weinstein BM (2009) Lymphatic development. Birth Defects Res C Embryo Today 87: 222–231.
33. Nissens K, Zhang G, Ridgway JB, Chen H, Kolumam G, et al. (2011) The Notch-Dll4 signaling pathway regulates mouse postnatal lymphatic development. Blood 118: 1989–1997.
34. Pong L-K, Gerhardt H (2009) Angiogenesis: a team effort coordinated by notch. Dev Cell 16: 196–206.
35. Yan M, Plowman GD (2007) Delta-like 4/Notch signaling and its therapeutic implications. Clin Cancer Res 13: 7243–7246.
36. Yan M, Callahan CA, Beyer JC, Allamneni K, Zhang G, et al. (2010) Chronic DLL4 blockade inhibits tumor angiogenesis. Blood 115: 2948–2956.
37. Tammela T, Zarkada G, Wallgard E, Murtomäki A, Suchting S, et al. (2008) Inhibition of Dll4-mediated signaling induces proliferation of immature vessels and results in poor tissue perfusion. Blood 109: 4753–4760.
38. Adams RH, Wilkinson GA, Weiss C, Diella F, Gale NW, et al. (1999) Roles of ephrinB ligands and EphB receptors in cardiovascular development: demarcation of arterial/venous domains, vascular morphogenesis, and sprouting angiogenesis. Genes & Development 13: 295–306.
39. Foo SS, Turner CJ, Adams S, Compagni A, Aubyn D, et al. (2006) Ephrin-B2 controls cell motility and adhesion during blood- vessel-wall assembly. Cell 124: 161–173.
40. Savaimphak S, Sieidel S, Essmann CL, Wilkinson GA, Pitausse ME, et al. (2010) Ephrin-B2 regulates VEGFR2 function in developmental and tumour angiogenesis. Nature 464: 564–570.
41. Wang Y, Nakayama M, Pitulescu ME, Schmidt TS, Bocciach P, et al. (2010) Ephrin-B2 controls blood vascular morphogenesis during postnatal angiogenesis. Nature 464: 564–570.
42. Armulik A, Abrahamsson A, Betoholz C (2005) Endothelial/pericyte interactions. Circ Res 97: 512–523.