Injury-activated glial cells promote wound healing of the adult skin in mice

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Cutaneous wound healing is a complex process that aims to re-establish the original structure of the skin and its functions. Among other disorders, peripheral neuropathies are known to severely impair wound healing capabilities of the skin, revealing the importance of skin innervation for proper repair. Here, we report that peripheral glia are crucially involved in this process. Using a mouse model of wound healing, combined with in vivo fate mapping, we show that injury activates peripheral glia by promoting de-differentiation, cell-cycle re-entry and dissemination of the cells into the wound bed. Moreover, injury-activated glia upregulate the expression of many secreted factors previously associated with wound healing and promote myofibroblast differentiation by paracrine modulation of TGF-β signalling. Accordingly, depletion of these cells impairs epithelial proliferation and wound closure through contraction, while their expansion promotes myofibroblast formation. Thus, injury-activated glia and/or their secretome might have therapeutic potential in human wound healing disorders.
The skin is the largest organ of the human body and acts as the primordial barrier of the organism against the outside environment. It mainly consists of two principle components: a stratified epidermis and an underlying layer of supportive connective tissue, the dermis. In various occasions throughout life, acute injuries challenge the integrity of this frontline defence. In most cases, they trigger an immediate emergency response to establish a sealed environment and prevent blood loss and infection, but also slower, long-lasting repair mechanisms. The latter involves various cell types to restore, at least partly, the initial biological properties of the injured site. Tissue repair mechanisms of the skin have been studied for decades and have highlighted that many key processes, such as, for instance, neo-vascularisation, are required to support the increased proliferation of fibroblasts and keratinocytes.

Besides increased blood supply, the healing response includes a second essential biological aspect: the neural response. Studies have shown that under normal circumstances hyperinnervation follows at the location of the injury. Impairment of the peripheral nervous system (PNS), whether traumatic or pathologic, results in improper tissue repair and failure to heal. One of the main functions of innervation has been attributed to axonal sprouting of neurons and their associated secretome of growth factors released in the wound bed upon injury. However, non-neuronal cells of the PNS have also been associated with wound healing. In particular, cells expressing the progenitor marker Sox2 and originating either from nerve terminals around hair follicles (HFs), from injured peripheral nerves or from distant sites outside the regenerating dermis, were shown to be involved in skin wound healing. How these cells contribute to the repair process is not entirely clear.

To specifically address the role of peripheral glia in cutaneous wound healing we used genetic mouse models allowing the tracing, conditional depletion, and conditional expansion of peripheral nerve cells in an otherwise undisturbed in vivo context. In this study, we report a novel role of PNS glia during wound healing of the skin. After a dedifferentiation and expansion process, injury-activated glia promote wound contraction and healing. This process is mediated by the secretion of factors enhancing transforming growth factor (TGF)-β signalling, which results in increased myofibroblast formation.

Results

Tracing PNS glia in the injured skin. Skin is a densely innervated organ with major nerve bundles (NB) visible in both intact skin and in skin healing from full-thickness excisional wounds (Fig. 1a). To determine the potential involvement of skin innervation in wound healing, we first used genetic lineage tracing to study the fate of nerve-derived cells upon skin wounding. Tamoxifen (TM)-mediated activation of CreERT² in the intact skin of Plp-CreERT²; tdTomato mice led to genetic tracing of peripheral glial cells in NBs of the reticular dermis, nerve terminals around HFs, as well as in nerve endings between muscle fibres. Apart from NBs and a fraction of melanocytic cells in HFs, the epidermis and the rest of the dermis appeared void of Plp-CreERT²-recombined cells (Fig. 1b, and Supplementary Figs. 1a, 2). Upon full-thickness excision wounding, however, we observed the presence of many Plp-CreERT²-traced (tdTomato+) single cells within the wound bed (Fig. 1c). In the intact dermis, expression of Sox10, a marker for the peripheral glial lineage, was mostly confined to NBs surrounded by a prominent laminin-positive perineurium. In contrast, NBs located adjacent to the wound displayed disrupted perineurial laminin expression and were associated with many single Sox10-positive (Sox10+) cells that seemingly disseminated from the disrupted nerves (Fig. 1d) and made up for the vast majority of Plp-CreERT²-traced nerve-derived cells (97.4%) (Supplementary Fig. 1b).

To confirm that NBs indeed serve as a source for cells homing to the wound bed in injured mouse skin, we made use of a separate Cre line, Dhh-Cre, which efficiently labels the glial lineage in the PNS. To visualise the glial lineage before and after skin injury, we employed confocal 3D-imaging techniques combined with a fructose-based tissue clearing protocol of Dhh-Cre; tdTomato animal samples. Using this method, we observed a network of tdTomato+ NBs in intact skin, while the stroma in between the nerve fibres was devoid of labelling. In contrast, the granulation tissue of wounded skin, already 6 days post injury (D6), displayed many single cells apparently stemming from the injured nerves at the wound edge. By day 14 (D14), the entire granulation tissue was densely populated by Dhh-Cre-traced cells (Fig. 1e).

Wounding induces glial de-differentiation and proliferation. Schwann cells in peripheral nerves possess a remarkable plasticity and have been described to de-differentiate and re-differentiate following injury to the nerve. To clarify the identity of nerve-associated cells populating skin wounds, we performed several immunohistostaining analyses for markers known to be associated with different Schwann cell states. Based on these experiments, we identified two classes of NBs located adjacent to or within the wound, which we term “quiescent” and “activated”, respectively (Fig. 2a). Quiescent NBs expressed high levels of myelin basic protein (MBP), a marker of myelinating Schwann cells, and axonal NF, but showed weak expression of the low-affinity nerve growth factor receptor (p75) (Fig. 2b, d). In contrast, activated NBs were often associated with a distorted NB morphology, displayed increased levels of p75 protein expression, and stained only weakly for MBP and NF (Fig. 2c, d). These features resemble the processes of myelin and axon degradation observed in Wallerian degeneration.

In addition, only activated NBs and cells dissociating from them were positive for PERK (Fig. 2e) and c-Jun (Fig. 2f) markers associated with Schwann cell plasticity and de-differentiation. Importantly, cells within activated NBs displayed an increased proliferation rate (quantified as percentage of Ki67+ cells) compared to quiescent nerves that were mostly composed of terminally differentiated, non-proliferative cells (Fig. 2e, f). Of note, 77% of all p75+ cells also expressed c-Jun. 33% of these double-positive cells were also marked by Ki67, a nearly 10-fold increase compared to the number of p75+c-Jun– Ki67 expressing cells (3.6%) (Fig. 2g, h). These data demonstrate that wounding of murine skin results in de-differentiation and active proliferation of a peripheral glial cell population and suggest that this process allows peripheral glia to dissociate and emigrate from injured nerves.

Peripheral nerves in the skin have been shown to harbour cells that upon isolation and in culture exhibit properties of multipotent neural crest stem cells (NCSCs), from which peripheral glia develop during embryonic development. To assess whether disseminating injury-activated nerve-derived cells generate NCSC lineages other than glial in vivo, we followed their fate at later time points post injury. In addition to Sox10 (which apart from...
**Fig. 1** Genetic tracing of the peripheral glial lineage in intact and injured skin. 

- **a** H&E-stained sections of intact skin and full-thickness excisional wound of adult mice at day 7 (D7) post-surgery. 
- **b** Glial lineage tracing of *Plp-CreER<sup>22</sup>; tdTomato* intact skin of a TM-injected animal. Dermal compartment is mostly void of single-labelled cells. 
- **c** Glial lineage tracing of *Plp-CreERT<sub>2</sub>; tdTomato* injured skin at D14 post-surgery in TM-injected animal. 
- **d** Boxed regions in the dermis are shown at higher magnification in the insets, highlighting the presence of multiple individual traced cells (red) populating the wound bed upon injury. 
- **e** Immunofluorescence staining of skin NB for the transcription factor Sox10 and the extracellular matrix protein Laminin (Lam) in intact and D7 injured skin show disruption of perineurium and dissemination of Sox10<sup>+</sup> cells upon injury. Arrowheads denote the presence of Sox10+/Lam<sup>+</sup> cells outside the NB. 
- **f** 3D imaging of the glial lineage of *Dhh-Cre; tdTomato* cleared intact mouse skin and cleared wounded skin at D6 and D14 show individual traced cells detached from NB upon injury. Arrowheads denote tdTomato-traced NBs. 
- **g** An apparently damaged sprouting nerve was subsequently sectioned and immunolabelled for the lineage tracer tdTomato. 
- **h** D7 and D14 wounded skin samples were subjected to immunolabelling to quantify the amount of glial cells (Sox10<sup>+</sup>) detached from NF-positive axons. Arrowheads denote glial cells with no adjacent NF staining and considered axon free. 

The quantification of the extent of glial dissociation from axons shows that the vast majority of Sox10<sup>+</sup> cells are not associated with axons at D7 and D14 (number of animals (N) = 4, number of wounds (n) = 8 (D7); N = 4 animals, n = 7 wounds (D14)). 

**Sections were counter stained with Hoechst 333258 (blue). Scale bars, 50 µm (a–f), 10 µm (g).**
Fig. 2 Skin injury induces peripheral glial de-differentiation and proliferation. a p75 (green), MBP (red), and NF (white) triple immunolabelling of wound sections shows increased expression of the de-differentiation marker p75 and loss of the differentiation marker MBP as well as loss of NF in activated NBs. b, c Single-channel insets show expression of p75, MBP, and NF in activated and quiescent NBs in injured skin. d Quantification of NBs categorised based on NF labelling (NF<sub>High</sub> and NF<sub>Low</sub>) and p75 positivity (p75<sup>+</sup> and p75<sup>−</sup>). e Immunolabelling of pERK (marker of activated MAPK signalling pathway) and Ki67 (marker for proliferative or cycling state) (red) in quiescent vs. activated NBs in injured skin. Activated NBs are characterised by p75 staining (green). f Quantification of Ki67<sup>+</sup> nuclei in quiescent or activated NBs. g Double immunolabelling of p75 (green) and c-Jun (red) shows concomitant upregulation of both markers in activated vs. quiescent NBs in injured skin. Black arrowhead denotes a Ki67<sup>+</sup> c-Jun<sup>−</sup> cell. White arrowheads denote Ki67<sup>+</sup> c-Jun<sup>+</sup> cells. h Quantification of proportion of proliferative (Ki67<sup>+</sup>) cells in c-Jun<sup>−</sup> and c-Jun<sup>+</sup> fractions of de-differentiated (p75<sup>+</sup>) glial cells within NBs. Data are represented as mean ± SEM of N = 7 animals, n = 12 wounds for activated NBs and N = 5, n = 9 for quiescent NBs (d); 8 quiescent NBs from N = 2, n = 4 and 14 activated NBs from N = 4, n = 6 (f); N = 6, n = 7 (h). Scale bars, 100 µm. Wound samples are D7 post injury.
glia also marks NCSCs23), over 90% of the disseminated cells expressed the de-differentiation marker p75 (Supplementary Fig. 1b, c). Furthermore, 77.8% of the genetically traced cells transiently lost the glial lineage marker GFAP at D7, which was however regained through D14. In addition, 77.4 ± 4% of the p75+ cells displayed the activation marker c-Jun, of which 32.0 ± 3.6% were positive for the proliferation marker Ki67 (Supplementary Fig. 1d, e).

Because Plp-CreERT2 not only marks peripheral glia, but also labels a subpopulation of melanocytes within the skin (Supplementary Fig. 2a, b)11,13, we aimed to assess the contribution of melanocytic cells to regenerating skin tissue. To do so, we compared wound experiments in Plp-CreERT2; tdTomato mice with Tyr-CreERT2; tdTomato animals, in which the melanocytic lineage can be specifically traced in vivo24,25. Immunolabelling of wound sections from both mouse strains ruled out the presence of any Dct+ cells in the granulation tissue and of any traced cells in wound sections from both mouse strains ruled out the presence of any Dct+ cells in the granulation tissue and of any traced cells in

We performed RNA sequencing of normal and wounded skin, 526 were upregulated and 506 downregulated (Fig. 3b, c). Among the genes upregulated in nerve-derived cells upon injury, we found several genes regarded as positive regulators of myelination, such as Egr2 and Pouf3f1/Oct6, as well as myelin genes and other genes—Mbp, Mal, and Cdh1—previously reported to be repressed in Schwann cells upon injury and de-differentiation29-31. Further, among the genes significantly upregulated upon injury were, for instance, Ngr, Gap43, Thy1, Igfbp2, Mdk, Pdgfb, Runx2, Cxcr4, which have all been reported to be upregulated in Schwann cells upon injury in previous studies28,29,32-34. Consistent with previous reports, we observed similar regulation of genes implicated in axon growth, guidance and protection, including the upregulated genes EphA5 and Semat4f, as well the downregulated genes Nfasc, Nrn1, and Atp1b225. We also found a set of upregulated genes previously described as important players in Schwann cell-mediated innate immune response, chemotaxis, and myelin phagocytosis, such as Lif, Cxcl5, Arg2, Megfi10, Mmp14, which all followed a similar regulation pattern in our data set as previously described after sciatic nerve injury21,27,28.

**Activated glia secrete factors implicated in wound healing.** To address the role of injury-activated glia during wound healing, we considered the possibility that these cells provide support by paracrine secretion of factors or microenvironment modulation of the healing tissue. Such a mode of action is conceivable given that Schwann cells have been reported to upregulate chemokines as well as growth factors upon nerve injury in order to modulate inflammation and axon sprouting, respectively27,28. To identify glia-derived factors potentially involved in skin wound healing, we performed RNA sequencing of Dhh-Cre-traced nerve-derived tdTomato+ cells isolated by fluorescence-activated cell sorting (FACS) from uninjured control animals and 7 days post injury (Fig. 3a).

Of the 1033 genes found to be differentially expressed between normal and wounded skin, 526 were upregulated and 506 downregulated (Fig. 3b, c). Among the genes downregulated in nerve-derived cells upon injury, we found several genes regarded as positive regulators of myelination, such as Egr2 and Pouf3f1/Oct6, as well as myelin genes and other genes—Mbp, Mal, and Cdh1—previously reported to be repressed in Schwann cells upon injury and de-differentiation29-31. Further, among the genes significantly upregulated upon injury were, for instance, Ngr, Gap43, Thy1, Igfbp2, Mdk, Pdgfb, Runx2, Cxcr4, which have all been reported to be upregulated in Schwann cells upon injury in previous studies28,29,32-34. Consistent with previous reports, we observed similar regulation of genes implicated in axon growth, guidance and protection, including the upregulated genes EphA5 and Semat4f, as well the downregulated genes Nfasc, Nrn1, and Atp1b225. We also found a set of upregulated genes previously described as important players in Schwann cell-mediated innate immune response, chemotaxis, and myelin phagocytosis, such as Lif, Cxcl5, Arg2, Megfi10, Mmp14, which all followed a similar regulation pattern in our data set as previously described after sciatic nerve injury21,27,28.

Because we hypothesised that injury-activated glia could exert their function through secretion of soluble factors, using MetaCore analysis from GeneGO, we focused on the predicted localisation of the gene products that were differentially expressed between non-wounded and wounded states. Intriguingly, extra-cellular space and membrane were the predominant localisations of gene products activated in peripheral nerve-derived cells upon wounding. Whereas most of the annotated genes from the downregulated gene list coded for proteins important in cell–cell contacts and adhesion, products of the upregulated genes tended to be secreted and associated to the extracellular matrix and extracellular space (Fig. 3d). In particular, consistent with our hypothesis of a paracrine function of injury-activated glia, the upregulated gene fraction showed a 3.5-fold enrichment in extracellular space/ECM-related genes upon injury—up to 53.4% (Fig. 3e).

When considering all differentially expressed genes coding for proteins with extracellular localisation (n = 455), 61.8% of them were upregulated (n = 281) and 38.2% downregulated (n = 174) (Fig. 3f). We clustered these 455 genes according to their function based on previous reports in the literature and annotations given in GeneCards (www.genecards.org). According to their described roles, we defined the following categories for upregulated genes:

1. Growth factors and cell signalling,
2. TGF-β signalling,
3. Chemotaxis and inflammation,
4. Migration and adhesion,
5. ECM remodelling,
6. Angiogenesis. Interestingly, many genes from this list have been implicated before in different processes important for wound healing and tissue repair (Fig. 3g; Supplementary Tables 1–6).

**Depletion of injury-activated glia impairs wound healing.** In order to assess the role of injury-activated glia in an in vivo context, we decided to deplete this cell population using genetic tools. Sox10 is an important survival and maintenance factor for the Schwann cell lineage during all stages of its differentiation23,36. In the skin, Sox10 is exclusively expressed in the melanocytic lineage (confined to HF’s in murine back skin) and the glial lineage25,37. To deplete nerve-derived cells in skin wounds and/or interfere with their migration into the wound, we generated Plp-CreERT2, Sox10lox/lox; tdTomato animals by breeding Plp-CreERT2; tdTomato mice with mice carrying floxed alleles of the Sox10 gene to specifically delete Sox10 in the glial lineage (Fig. 4a). Since effective depletion of Sox10 protein upon genetic ablation of the allele is a relatively slow process in adult mice38, we allowed 3 weeks post-TM injection before we subjected both Sox10 conditional knockout (cKO) and control littermates to excisional wounding, although early enough to limit potential effects of peripheral neuropathy which peaks at 40 days post TM treatment38. Tissue samples were collected for analysis at several time points as depicted in Fig. 4a. NBs of Sox10 cKO animals visually appeared similar to those of control animals and axons in quiescent NBs were preserved as shown by NF staining (Fig. 4b). To verify depletion of glial cells in the nerves, we quantified Sox10+ cells within NBs located close to the wound bed and found that 56.8 ± 4.9% of all cells expressed Sox10 in control vs. 12.5 ± 3.2% in Sox10 cKO mice at D7 (Fig. 4c).

To assess the number of injury-activated, de-differentiated glia present upon Sox10 cKO, we performed immunofluorescence analysis and quantification of p75+ and traced tdTomato+ cells in the wound bed at two different time points. The data show a similar significant decrease of p75+ and traced cells in Sox10 cKO animals at both D7 (46.5 ± 4.4 vs. 7.0 ± 1.8 p75+ cells mm-2 and 40.2 ± 5.4 vs. 2.8 ± 0.9 tdTomato+ cells mm-2 in control vs. Sox10 cKO) and D14 after wounding (90.2 ± 12.8 vs. 24.1 ± 5.0 p75+ cells mm-2 and 153.6 ± 28.8 vs. 43.9 ± 3.3 tdTomato+ cells
**Fig. 3** Expression profile analysis of glial lineage upon skin injury. 

**a** Experimental design for skin sample collection of Ctrl and D7-injured Dhh-Cre; tdTomato animals. Single cell suspensions were sorted for tdTomato expression using a flow cytometer (0.8–1.8% of total cell fraction). Total RNA samples of gated cells were amplified and subsequently subjected to RNA-Seq using Illumina RNA-Seq platform. 

**b** Volcano plot of the gene expression profile of Dhh-Cre-traced tdTomato+ cells of intact skin and D7 post injury. Coloured data points meet the thresholds of FC above 1.5 and under −1.5, P-value < 0.05, FDR < 0.05. 

**c** Unsupervised sample clustering heat map of gene expression with a significantly differential expression. 

**d** Gene GO MetaCore prediction of the product localisation of the significantly downregulated (506 genes) and upregulated genes (526 genes) shows a significant enrichment of genes, the products of which are excreted. 

**e** Data representation for the percentage of gene products predicted to be destined to the extracellular space. 

**f** Volcano plot of the expression value of the genes with predicted extracellular localisation of their product. 

**g** Quantitative Venn diagram of the differentially expressed “secretome” and the associated biological function.
Further analysis of the relatively few tdTomato-traced cells still present upon Sox10 cKO showed that the vast majority of the fate-mapped cells, both in NBs and in the wound bed, still displayed Sox10 expression (Supplementary Fig. 3a, b). Therefore, rather than adopting alternative fates, peripheral glial cells having lost Sox10 were virtually all eliminated, in agreement with previous studies36,38 (Supplementary Fig. 3c, d). These data demonstrate that conditional deletion...
of Sox10 in the glial lineage using Plp-CreERT2 is an efficient method to decrease the number of peripheral nerve-derived cells in the wound.

To study possible effects caused by genetic depletion of nerve-derived cells in the wound bed, we performed macroscopic measurements and morphometric analysis of control and Sox10 cKO wounds. Although the dynamics of wound contraction can be variable, as previously described, we found a significant difference in the mean wound size at D7 (14.2 ± 1.6% vs. 22.2 ± 3.1% of initial size in control vs. Sox10 cKO) as well as at D9 (4.8 ± 1.2% vs. 11.1 ± 3.0%) (Fig. 5a, b). Since the most significant difference in wound contraction was observed at D7, we carried out morphometric analyses of D7 wound sections to assess whether other commonly measured wound healing parameters were changed as well. The principle of measurements and representative pictures of wounds at this stage are shown (Fig. 5c).

Quantification of the distance of newly formed epidermis covering the wound revealed a delay in wound closure in Sox10 cKO animals—an average of 84% of the wound was covered by epidermis in controls, while only 66% of the wound was covered in Sox10 cKO at the same time point (Fig. 5d). Likewise, the area of hyperproliferative epidermis (HPE) was significantly decreased (0.282 ± 0.017 vs. 0.220 ± 0.015 mm²) in Sox10 cKO wounds (Fig. 5e). Importantly, in the absence of activated glia, the wound width was increased (1.99 ± 0.08 vs. 2.46 ± 0.08 mm), while the length of wound epithelium remained unchanged in Sox10 cKO animals compared to controls, suggesting impaired contraction (Fig. 5f, g). In sum, injury-activated glia in the wound bed is required for proper healing.

Activated glia are dispensable for angiogenesis and immune response. Some of the genes upregulated in peripheral nerve cells upon wounding encode secreted factors, such as Apln, Esml1, Mdk, and Vcan, which have previously been shown to influence angiogenesis. Therefore, depletion of injury-activated glia could affect neovascularisation (Supplementary Tables 1 and 6). However, when compared to the granulation tissue of control wounds, genetic depletion of nerve-derived cells in the wound did neither significantly change the number nor the density of blood vessels labelled by the endothelial cell marker CD31 (Supplementary Fig. 4a).

Further, we assessed the role of these cells on immune cell chemotaxis, as our differential expression profiling revealed several genes coding for proteins known to be important for innate immune cell chemotaxis, such as Cxcl5, Cxcl2, Lif, and Tnem2, among others (Supplementary Table 3). However, no detectable changes in the number of total immune cells marked by CD45 or of CD206 macrophages were observed upon manipulation of the number of nerve-derived cells in the wound bed (Supplementary Fig. 4b, c).

Similarly, the above-mentioned expression profile analysis underlined significant changes in the transcription of genes involved in ECM deposition (Supplementary Table 5). Hence, we investigated the state of collagen maturation of D14 control and Sox10 cKO wound sections using Herovici stainings. However, we could not detect obvious changes in young vs. mature collagen deposition between control and Sox10 cKO animals with our experimental settings (Supplementary Fig. 4d).

GliaL depletion impairs TGF-β signalling and myofibroblast formation. In the wound granulation tissue, α-SMA is a marker for myofibroblasts, which, as previously mentioned, are important for wound healing through their role in wound contraction and extracellular matrix protein (ECM) deposition. Our expression profiling analysis showed upregulation of a number of genes important for TGF-β signal regulation and myofibroblast formation, such as Inhba, Lifbp2, Mmp2, Bmp1, Lox12, Pld1, and Plod2 (Supplementary Table 2). Hence, we looked at the area occupied by myofibroblasts post injury. Interestingly, the area of α-SMA expression was significantly reduced in Sox10 cKO wounds at D7 (20.7 ± 2.7% of the total granulation tissue area) compared to control (33.6 ± 2.3%) (Fig. 5h, i). To substantiate these findings, we performed Western blot analysis of protein isolated from tissue of control and Sox10 cKO wounds at D7 and measured a significant decrease of α-SMA protein levels upon Sox10 cKO-mediated depletion of injury-activated glia (Fig. 5j and Supplementary Fig. 5a). Likewise, Sox10 cKO wounds contained significantly reduced amounts of Periostin (Posrn), a multifunctional promoter of wound healing that has been associated, among others, with formation of myofibroblasts from fibroblasts.

To investigate whether TGF-β signal activation is indeed altered in skin wounds upon ablation of the glial population, we quantified the number of cells with prominent nuclear pSMAD2 localisation—a downstream marker of activated TGF-β/activin signalling—in the granulation tissue of the wounds. At D7, depletion of activated glia led to a significant decrease in the number of pSMAD2 nuclei in Sox10 cKO vs. control wounds (11.29 ± 0.8% vs. 24.1 ± 1.8% in Sox10 cKO vs. control, respectively) (Fig. 5l, m). In sum, conditional ablation of Sox10 in peripheral nerves prior to injury is sufficient to prevent colonisation of the wound tissue by nerve-derived cells and leads to delayed wound contraction and re-epithelialisation.

Activated glia promote myofibroblast formation in vivo. Loss of the tumour suppressor gene Pten has previously been shown to result in proliferation of both myelinating and non-myelinating Schwann cells. Hence, Pten inactivation might represent a genetic tool to expand the number of nerve-derived cells in wounds. Therefore, we crossed Plp-CreERT2 mice with mice carrying floxed alleles of the Pten gene. Pten was then depleted homozygously shortly prior to injury (Fig. 6a). In both quiescent and activated NBs of control animals, pAKT expression was not detectable by immunohistochemical methods. However, upon Plp-CreERT2-mediated recombination, peripheral nerve cells prominently expressed pAKT, indicating efficient activation of PI3K/AKT signalling pathway upon Pten deletion (Fig. 6b). Remarkably, NBs of Pten cKO animals displayed an overt expansion of nerve-derived cells in the wound (Fig. 6c). Quantification revealed an approximately two-fold increase in the number of p75+ and lineage-traced cells in the wound bed of Pten cKO mice (Fig. 6d, e).

Normal wound healing in healthy mice appears to be naturally optimised and it has been notoriously hard to experimentally improve this process. Indeed, the increase in the number of nerve-derived cells in the wound of Pten cKO mice did neither lead to a significant acceleration of the wound closure D7 (86.3 ± 5.0% covered by epidermis in control vs. 87.2 ± 5.1% in Pten cKO) nor did it alter the area of HPE (0.250 ± 0.02 vs. 0.278 ± 0.03 mm²), wound width (2.12 ± 0.09 vs. 1.90 ± 0.10 mm) or length of wound epithelium (1.81 ± 0.06 vs. 1.68 ± 0.08 mm), measured at the same time point (Fig. 6f–i).

Interestingly, however, the area of α-SMA staining in the wound bed significantly increased upon Pten cKO, with myofibroblasts occupying around 47% of the total granulation tissue area in Pten cKO wounds, compared to 34% in control settings (Fig. 6j, k). Importantly, consistent with our finding that TGF-β signalling was reduced upon deletion of injury-activated glia (Fig. 5l, m), we found that increasing the number of activated glial cells led to enhanced TGF-β signalling in cells of the...
granulation tissue. Indeed, at D5, Pten cKO wounds showed an increased number of cells with nuclear staining for pSMAD2 compared to control (22.2 ± 2.9% vs. 14.2 ± 1.2%, respectively) (Fig. 6l). In sum, conditional loss of Pten caused an expansion of injury-activated glia in the regenerating skin, which was associated with activation of TGF-β signalling and a concomitant increase in α-SMA expression in the wound. These findings demonstrate a role of activated peripheral glia in regulating the number of myofibroblasts during wound healing.

**Glia promote myofibroblast formation via TGF-β signalling.** To directly demonstrate the capacity of peripheral glia to induce

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**Key Figures**

- **a** Full-thickness wound illustration
- **b** Macroscopic wound size
- **c** D7 wound histology
- **d** Wound closure D7
- **e** HPE D7
- **f** Wound width D7
- **g** Re-epithelialization D7
- **h** Myofibroblast labelling in D7 wounds
- **i** α-SMA area D7
- **j** α-SMA protein D7
- **k** Postn protein D7
- **l** TGF-β signaling in D7 wounds
- **m** pSMAD2 D7
myofibroblast differentiation from dermal fibroblasts, we established a co-culture system of human fibroblasts (HuFs) and murine sciatic nerve explants (SN Exp) obtained from Dhh-Cre; tdTomato animals, as depicted schematically in Fig. 7a. In this system, cells of murine origin can be distinguished from unlabelled HuFs by virtue of their tdTomato expression. Intriguingly, upon dissection and culture alone, glia in peripheral nerves appear to undergo processes very much resembling those observed in vivo upon wounding. Indeed, murine tdTomato+ Sox10+ glial cells in the SN Exp—fibroblast co-culture system—differentiated through expression of p75 and c-Jun and spontaneously emigrated from the nerve explants (Fig. 7b). To assess the effect of these activated glial cells on myofibroblast differentiation, and to decipher a potential implication of TGF-β signalling in this process, we performed α-SMA staining of HuFs cultured with or without murine SN Exp, in the presence or absence of the TGF-β signalling inhibitor SB431542 (SB43) (Fig. 7c). Noteworthy of note, co-culture with SN Exp significantly increased the number of α-SMA+ HuFs when compared to cultures without peripheral nerves (16.1 ± 1.2% vs. 22.3 ± 1.8%, respectively) (Fig. 7d). Moreover, α-SMA expression was higher in close proximity to the murine tissue as compared to areas in the same culture dish but further away from the SN Exp, further demonstrating the α-SMA-inducing effect of nerve-derived cells (Fig. 7e, f).

Importantly, nerve-dependent induction of α-SMA expression in HuFs was abolished upon TGF-β signalling blockade (Fig. 7d, f). Thus, activated glial cells induce myofibroblast formation through paracrine activation of TGF-β signalling.

**Discussion**

In the present study, we demonstrate a novel role of injury-activated peripheral glia during wound healing of the skin. Lineage-specific genetic tracing, visualised among others by tissue clearing techniques combined with 3D-imaging, revealed that glial cells upon skin injury de-differentiate to express markers characteristic for repair Schwann cells formed in response to nerve injury and start proliferating. Intriguingly, activated glia then disseminate from the injured nerves into the granulation tissue, where they support non-neural tissue repair of the wounded skin by paracrine signalling. In particular, depletion of injury-activated glia impaired TGF-β signal activation and the appearance of myofibroblasts in the skin; in contrast, genetically induced expansion of injury-activated glia in vivo as well as their co-culture with fibroblasts promoted TGF-β signalling and, consequently, myofibroblast formation.

It is known that injury to peripheral nerves is associated with peripheral glia activation and de-differentiation. However, the signalling event triggering this process remains largely elusive. Although certain extracellular factors including interleukin (IL)-6 and tumour necrosis factor (TNF)-α have been implicated in Schwann cell activation, culture of dissected peripheral nerves without addition of these factors was sufficient to enhance p75 and c-Jun expression in glia. Likewise, co-culture with other cell types was not required to obtain glial activation in dissected peripheral nerves, speaking against a major involvement of skin cells, such as keratinocytes, dermal fibroblasts, or immune cells. Together with the correlation between loss of axonal NF and glial de-differentiation that we measured in activated NBs of wounded skin, our data suggest that impaired physical integrity of peripheral nerves might be sufficient to trigger glial activation.

Activated peripheral glia induced by nerve injury have been reported to acquire a plastic phenotype reminiscent of Schwann cell precursors found during embryonic development. Swann cell precursors appear to be multipotent and, apart from glia, can give rise to endoneurial fibroblasts, melanocytes, parasympathetic neurons, and mesenchymal cells. Moreover, after environmental challenges, PNS glia in the adult can also give rise to melanocytes, neurons, and tooth mesenchymal cells. Therefore, we had to consider the possibility that activated glia in the wound bed structurally contribute to scar formation through differentiation into non-glial lineages. However, lineage tracing combined with marker expression analyses did not provide significant evidence for injury-activated glia adopting alternative fates. In particular, traced cells neither generated melanocytes nor myofibroblasts, which are both cell types that can be produced by neural crest cells, the precursors of Schwann cells during development. Instead, in the undisturbed 3D context of the healing wound, injury-activated glia displayed lineage-restriction in vivo and continued to express peripheral glial lineage markers even after prolonged time points after injury. This is reminiscent of Schwann cell precursors in the Mexican salamander axolotl (*Ambystoma mexicanum*) that maintain Schwann cell identity after limb amputation while promoting regeneration and of astroglia in the mammalian CNS that are also activated in response to injury and contribute to tissue repair, but exclusively give rise to astrocytes unless genetically manipulated.

Rather than a direct contribution of activated peripheral glia to cell types constituting the wound bed, our study demonstrates a cell non-autonomous effect on these cells during wound healing. Expression profiling of genetically traced peripheral glial cells either from intact or wounded skin demonstrated a significant enrichment for expression of various ligands and gene products destined to the extracellular space, representing a putative secretome of Schwann cells upon injury. This secretome includes gene products reported to affect multiple pathways, such as ECM remodelling, angiogenesis, chemotaxis of immune cells, or TGF-β signalling, which are all biological processes well known to promote wound healing. Although these findings point to a novel role of peripheral glia in the healing wound, the exact cell types involved and their contributions need to be addressed in the future. In conclusion, our results strongly suggest that peripheral glia, once activated, have the potential to support wound healing in an autocrine and paracrine manner.

**Fig. 5** Depletion of injury-activated glia in the granulation tissue impairs wound healing. a Representative macroscopic illustration of wound healing in Ctrl and Sox10 cKO animals at D0 and D7. b Longitudinal macroscopic quantification of individual wound areas at D2, D4, D7, and D9 in Ctrl and Sox10 cKO animals in relation to their respective initial size at D0. Data show a significant delay in wound closure at D7 upon Sox10 cKO. c H&E-stained sections of Ctrl and Sox10 cKO D7 wounds used for morphometric analysis of percentage of wound closure (length of newly formed epithelium (NFE)/length of NFE + wound width), re-epithelialisation (length of NFE), area of HPE, wound contraction (distance between wound border HFS (blue dotted line)), and re-epithelialisation (length of NFE). Analysis of D7 wound sections for the percentage of d wound closure, e area of HPE, f wound width, g re-epithelialisation. h Immunolabelling of myofibroblasts with α-SMA (green) in Ctrl and Sox10 cKO skin wound sections at D7. Dashed line delimits granulation tissue. i Quantification of the percentage of α-SMA+ area of the granulation tissue at D7 shows a reduction of the area occupied by myofibroblasts upon Sox10 cKO. Western blot quantification for j myofibroblast marker α-SMA and k Peroxisin (Ponst) of D7 whole-wound protein extracts from Ctrl and Sox10 cKO animals. l Immunolabelling of α-SMA (green) and pSMAD2 (red) in Ctrl and Sox10 cKO skin wound sections at D7. Boxed regions in the dermis are shown in the insets at higher magnification as pSMAD2 single channel and merged with α-SMA channel. m Quantification of highly pSMAD2+ cells per α-SMA+ area of the granulation tissue at D7. Data are represented as mean ± SEM of N = 10, n = 39 (Ctrl), N = 13, n = 49 (Sox10 cKO) (b); N = 9, n = 35 (Ctrl), N = 13, n = 52 (Sox10 cKO) (d); N = 5, n = 18 (Ctrl), N = 6, n = 24 (Sox10 cKO) (e); N = 7, n = 28 (Ctrl), N = 8, n = 32 (Sox10 cKO) (f); N = 7, n = 28 (Ctrl), N = 10, n = 40 (Sox10 cKO) (g); N = 5, n = 19 (Ctrl), N = 5, n = 17 (Sox10 cKO) (i); N = 3, n = 6 per group (j, k); N = 8, n = 15 (Ctrl), N = 10, n = 20 (Sox10 cKO) (m). Scale bars, 200 μm.
influence wound healing\(^2\). To test whether any of these processes were significantly impacted by our cell population of interest, we genetically ablated injury-activated glia by preventing their expansion into the wound bed using lineage-specific conditional deletion of the transcription factor Sox10, a factor known to control neural crest cell survival and glial lineage specification\(^5\). Preventing dissemination of glial cells into the wound bed delayed wound healing and, notably, interfered with wound contraction and proper myofibroblast formation, which was associated with decreased TGF-β signalling in a non-cell autonomous manner. These data are in line with previous reports showing a direct link between TGF-β, granulation tissue...
myofibroblasts, and wound contraction in vivo.\textsuperscript{55,56} Accordingly, expansion of injury-activated glia in the wound bed, achieved by genetic ablation of \textit{Pten}, led to increased paracrine TGF-\textit{\beta} signalling and myofibroblast counts. Furthermore, genetically traced glia emigrating from dissected peripheral nerves promoted a fibroblast to myofibroblast transition in a newly established co-culture system. This effect was fully blocked by addition of a TGF-\textit{\beta} signalling inhibitor, identifying TGF-\textit{\beta} signalling as a mediator of the paracrine effect exerted by injury-activated glia. However, besides the activin/inhibin ligand Inha, our study did not provide evidence for specific TGF-\textit{\beta} ligand expression by injury-activated glia. Rather, it pointed to modulation of ligand availability through ECM remodelling and TGF-\textit{\beta} post-translational modification, likely conferred by factors known to regulate these processes and present in the glia secretome, such as \textit{Lox}, \textit{Loxl2}, and \textit{Plod2} and \textit{Bmp1} (references in Supplementary Tables 2 and 5). This is consistent with previous reports indicating that non-myelinating Schwann cells have the potential to secrete molecules regulating latent TGF-\textit{\beta} activation.\textsuperscript{57} Although many other factors of the glia secretome characterised in our study have been functionally associated before with wound healing (see references in Supplementary Tables 1–6), their contribution to the paracrine effects by injury-activated glia remains to be elucidated. Notably, the amount of Periostin, a known inducer of myofibroblast formation and modulator of several other processes involved in wound healing,\textsuperscript{41} was decreased in the wound bed upon genetic depletion of injury-activated glia. Given the many distinct classes of factors present in the glia secretome, it is conceivable that injury-activated glia take part in processes of skin wound healing other than promoting myofibroblast formation. Indeed, loss of injury-activated glia also affected the area of newly formed, HPE. However, expansion of peripheral glia in the wound bed did not influence this process, presumably because it is naturally optimised in murine skin wound healing.\textsuperscript{13} Nonetheless, our data speak for multiple effects of injury-activated glia and point to a therapeutic potential of these cells and/or of their secretome in human wound healing.

Our work is in line with other recent reports that showed crucial roles of peripheral nerve-derived cells in tissue regeneration and repair, both in amphibians and mice.\textsuperscript{9,58,59} For instance, specialised repair glial cells forming after nerve injury were found to guide axons and restore innervation.\textsuperscript{12,28} Ablation of Sox2-expressing cells, which are located in nerve terminals in the skin, peripheral nerves, and dermal papillae in HFIs, impaired mouse skin wound healing and digit regeneration.\textsuperscript{59} Likewise, transplantation of Schwann cell precursors isolated from sciatic nerves elicited a positive effect on digit tip regeneration by promoting mesenchymal cell proliferation and bone regeneration via secretion of Oncostatin M and PDGF-AA.\textsuperscript{39} Together with our study showing their influence on TGF-\textit{\beta}-mediated myofibroblast formation and skin wound closure, peripheral glia emerge as important players in tissue regeneration. Possibly, this involves interaction of activated glia with other stem and progenitor cells types. In fact, Schwann cell precursors purified from neonatal sciatic nerve can stimulate self-renewal of skin-derived precursors,\textsuperscript{58} a mesoderm-derived mesenchymal cell population able to reconstitute the dermis.\textsuperscript{60,61} It remains to be shown, though, whether the nerve-derived cell populations reported in the literature to have regenerative potential represent distinct cell types or one dynamic cell population eliciting distinct paracrine responses to injury depending on the type of injury or the organ affected. Likewise, it will be highly relevant to address whether injury-activated glia are also implicated in other tissues for which regeneration was shown to depend on innervation,\textsuperscript{62} including heart and skeletal muscle.\textsuperscript{53–56}

**Methods**

**Mouse strains.** \textit{Dbh-Cre}: \textit{Tg(Dbh-Cre)1Mej5} – a transgene consisting of a Cre-coding sequence inserted at the start codon of a \textit{Dbh} genomic clone (strain of origin: FBV/N). \textit{PipCreER}\textsubscript{T2}: \textit{Tg(Pipl1-Cre;ER(T2))1Ueli} – a transgene consisting of the mouse promoter of the \textit{Pip} gene driving the expression of a TM inducible Cre recombinase (strain of origin: C57BL/6 × DBA/2F1). \textit{Pten}\textsuperscript{lox/lox} – a floxed allele of \textit{Pten} allowing the suppression of its phosphatase activity upon \textit{Cre}-mediated recombination (strain of origin: 129/JyKae). \textit{Sox10}\textsuperscript{lox/lox} – a floxed allele of \textit{Sox10} allowing the conditional KO of the gene upon \textit{Cre}-mediated recombination (strain of origin: 129P2/OlaHsd). \textit{tdTomato}: \textit{Gt(Rosa)26Sor(TdTomato)Hze67} – \textit{tdTomato} sequence preceded by a loxp-flanked STOP cassette targeted to the Rosa locus. This allows \textit{Cre} reporting only in the immediate lineage tracing following \textit{tdTomato} expression (strain of origin: 129S6/EvTac × C57BL/6N(Crl)F1). \textit{Tyr-CreER}\textsubscript{T2}: \textit{Tg(Tyr-Cre;ER(T2))13Bos} – a transgene consisting of 5.5 kb fragment of the Tyr promoter and a 3.6 kb fragment of the Tyr enhancer driving expression of a TM inducible form of Cre recombinase.

**Experimental mouse lines.** For genetic tracing of neural crest-derived population \textit{PipCreER}\textsubscript{T2}; \textit{tdTomato}, \textit{Dbh-Cre}, \textit{tdTomato}, and \textit{Tyr-CreER}\textsubscript{T2}; \textit{tdTomato} transgenic mouse lines were used. For the ablation of neural crest-derived cell population we used \textit{PipCreER}\textsubscript{T2}; \textit{Sox10}\textsuperscript{lox/lox}; \textit{tdTomato} transgenic mice. For the expansion of neural crest-derived cell population \textit{PipCreER}\textsubscript{T2}; \textit{Pten}\textsuperscript{lox/lox}; \textit{tdTomato} transgenic mice were used. All animal experiments have been approved by the veterinary authorities of Canton of Zurich, Switzerland, and were performed in accordance with Swiss law on the care, welfare, and treatment of animals.

**Genotyping.** DNA extracted by standard procedures from tissue samples was analysed by PCR, using gene-specific primers (Supplementary Table 7).

**Experimental animals.** In our wound healing experiments, we used animals bred in-house, aged between 2 and 6 months of several combined transgenic lines, with their skin hair cycle in the late anagen to telogen phases. Due to the various strains of origin of our transgenes, the genetic background is empirically considered mixed. Cell population characterisation by immunolabelling was performed on a cohort mixed gender. Littermates were evenly distributed across control and \textit{ko} experimental groups. All quantifications were performed on tissue samples obtained from males only. The experimenters were not blinded and were aware of the genotype of the animals prior to experimentation. No statistical method was applied to predetermine the same size of experimental groups. Sample sizes were large enough to measure the effect size. Appropriate animal health status, in accordance with our animal experimentation protocol, was used for exclusion.
Fig. 7 Activated glia from SN Exps promote myofibroblast differentiation in a TGF-β signalling-dependent manner. a Experimental scheme of HuF cultured in the presence of murine Dhh-Cre; tdTomato SN Exp with optional pharmacological modulation of TGF-β signalling pathway. b Immunofluorescence of SN Exp show tdTomato-traced cells (red) expressing markers of activated glia, such as Sox10 (white), p75 (green), and c-Jun (green). Note that similar results were obtained even in the absence of fibroblasts. c Immunolabelling of α-SMA+ (green) HuF upon SN Exp co-culture, TGF-β1 treatment or SB43-mediated TGF-β signalling blockade and respective quantification of α-SMA induction for each culture condition. ***P < 0.001 compared to Ctrl. e Immunolabelling of local α-SMA+ induction in tdTomato− HuF in the direct vicinity of the SN Exp compared to areas distant from the SN Exp (one field of view from nearest tdTomato+ cell using 10× objective) of the same well in combination with SB43-mediated TGF-β signalling blockade. f Respective quantification of local α-SMA induction for each culture condition. Data are represented as mean ± SEM. Scale bars, 100 μm (b), 300 μm (c, e)
criteria. No randomisation of the animals was used as experimental group assignment was primarily dependent on the appropriate animal genotype.

Cre-mediated allele recombination. cKO of Sox10 using the Ptp-CreERT2 driver was performed using i.p. injection of a 10 mg ml⁻¹ TM solution in sunflower oil/EtOH 9:1. Animals were injected daily with 200 μl TM for 5 consecutive days for a total of 10 mg per animal. To obtain efficient depletion of the Sox10-expressing population in the skin, cKO of Sox10 was performed 3 weeks prior to injury. Pten cKO was performed using a 5-day treatment of daily i.p. injection of 10 μl ml⁻¹ TM solution. Operative procedure was performed 5 days after the initial injection.

Excisional wounding of murine skin. Following a protocol approved by the Zurich cantonal veterinary office, general anaesthesia of the animals was induced by 5% Isoflurane in 100% O₂ and subsequently maintained using 3% Isoflurane. The back skin was shaved, thoroughly cleaned, and disinfected prior the surgery using antibacterial soap solution and 75% EtOH solution. Preoperative anaesthetic, was applied using a subcutaneous injection of Buprenorphine (0.1 mg kg⁻¹). Four circular full-thickness excisional wound sections of 5 mm of diameter were generated on the lower back skin of each animal, two on each side, 1 cm of the midline of the animal and roughly 2 cm apart from each other. Post-operative analgesia was performed by a 5-day treatment of Buprenorphine through the drinking water. Wounds were allowed to heal without dressing.

Macrosopic wound size measurement. Wound area was determined for each wound by quantification of wound area based on pictures taken from day 0 to day 14 post surgery. The area at different time points after wounding were then reported to the original area value at day 0 for each wound.

Histological analysis of murine skin samples. At the day of sample collection, animals were killed under a CO₂ atmosphere and skin samples were excised from the midline of the back, fixed in a buffered 4% formaldehyde solution at 4 °C, overnight for samples to be embedded in paraffin. For samples to be embedded in tissue freezing medium or subjected to clearing 4 h fixation at 4 °C was performed. Paraffin blocks were processed into sections of 5 μm while cryoblocks were processed into 12 μm sections. For immunofluorescence protocols described elsewhere were applied. Briefly, sections were deparaffinised and subjected to an antigen retrieval step using citrate buffer (S2369, Dako). Primary antibodies (Supplementary Table 8) were applied in Odyssey blocking buffer (927-40000, LI-COR) and for 1 h at room temperature with secondary antibodies (Supplementary Table 9) and Hoechst 33342 (PBS). For quantification, as previously described for pERK, c-Jun and pSMAD2 staining TSA media containing 1% BSA only as a blocking buffer for 1 h at room temperature. For pERK, c-Jun and pSMAD2 staining TSA media containing 1% BSA only as a blocking buffer for 1 h at room temperature. For pERK, c-Jun and pSMAD2 staining TSA media containing 1% BSA only as a blocking buffer for 1 h at room temperature. For pERK, c-Jun and pSMAD2 staining TSA media containing 1% BSA only as a blocking buffer for 1 h at room temperature. Scanning of fluorescence-stained, H&E-stained, and Herovici-stained sections were imaged using a DMI 6000B microscope (Leica) and Image J software (National Institutes of Health, USA). Cells were quantified either distally from the explant (at least 1 mm from the migrating explant cells) or proximally—with migrating cells present in HMFs.

SNP Exp co-culture and immunofluorescence. Fibroblasts used in this study were derived from a human foreskin sample after obtaining permission from the Ethics Commission of the Canton Zurich and after informed consent given by parents. For nerve explant co-cultures human dermal fibroblasts were seeded in 12-well plates at 7 x 10⁶ cells cm⁻² density and cultured for 24 h in 10% FBS-supplemented DMEM. After 24 h the media was changed to 1% FBS DMEM and scarified fragments were placed in the middle of the wells at low media volume (300 µl/well). Sciatric nerves were isolated from adult Dhh-Cre; tdTomato mice and processed as previously published. Briefly, sciatric nerves were dissected under sterile conditions—nerves were cut in 2–3 mm pieces followed by careful epineurium and perineurium removal using fine forceps and binoculars. These fragments can be used as explants and passaged. After 4 days of co-culture of DFF12 and semi-sufficient supernatant media containing 1% FBS only or combinations of supplements was added for full concentration of 10 ng ml⁻¹ TGF-β1 (100-21, Peprotech) and 10 µM SB 431542 (301836-41-9, Tocris). DMSO was used as vehicle control. Fresh supplements were added to the media every day for 9 days.

For immunofluorescence cells were fixed for 6 min with 4% formaldehyde in PBS and blocked for 45 min (3% BSA, 0.1% Triton X-100 in PBS). Cells were incubated with primary antibodies (Supplementary Table 8) overnight at 4 °C (1% BSA in PBS) and for 1 h at room temperature with secondary antibodies (Supplementary Table 9) and Hoechst 33342 (PBS). For quantification, in each well cells positive for α-SMA with visible stress fibres were counted in at least four 1 mm² HMFs using aDMI 6000B microscope (Leica) and Image J software (National Institutes of Health, USA). Cells were quantified either distally from the explant (at least 1 mm from the migrating explant cells) or proximally—with migrating cells present in HMFs.

Protein isolation and western blotting. For wound protein isolation two D7 wounds were pooled for each sample. Wound tissue was transferred into RIPA buffer (89900, Thermo Scientific) containing Hall–Phosphatase and Protease Inhibitor Cocktail (78440, Thermo Scientific) and tissue was homogenised using tissue homogeniser (Polytron) and subsequently sonicated three times for 10 s using ultrasonic homogeniser (Bandelin). SDS–PAGE was carried out on 4–20% Mini-PROTEAN TGX Gels (456-1094, Bio Rad). Primary antibodies (Supplementary Table 8) were applied in Odyssey blocking buffer (927–40000, LI-COR Biosciences) overnight at 4 °C and secondary antibodies (Supplementary Methods Table 9) for 1 h at room temperature. Blots were scanned and quantified using Odyssey imaging system (LI-COR Biosciences). Quantified band intensities were normalised using β-Actin and plotted on the graph. Full size scans of the blots are presented in Supplementary Fig. 5.

Statistical analyses. Statistical analysis was performed and graphs created using Prism 6 software (GraphPad Software Inc., San Diego, CA). Variation estimates were not performed or compared for each data set. Statistical significance between two groups was calculated using unpaired Student’s t-test for normally distributed data sets. For wound area measurements in Fig. 5b non-parametric Mann–Whitney test was used.

Data availability. The data sets generated during and analysed during the current study are available from the corresponding author on reasonable request. RNA sequencing data have been deposited to the European Nucleotide Archive under the accession code PRER22372.

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References

1. Martin, P. Wound healing–aiming for perfect skin regeneration. Science 276, 75–81 (1997).
2. Gurtner, G. C., Werner, S., Barrandon, Y. & Longaker, M. T. Wound repair and regeneration. Nature 453, 314–321 (2008).
3. Eming, S. A., Martin, P. & Tomic-Canic, M. Wound repair and regeneration: mechanisms, signaling, and treatment. Sci. Transl. Med. 6, 265sr6 (2014).
4. Li, J., Zhang, Y.-P. & Kirsner, R. S. Angiogenesis in wound repair: angiogenic growth factors and the extracellular matrix. Microsc. Res. Tech. 60, 107–114 (2003).
5. Harsum, S., Clarke, J. D. & Martin, P. A reciprocal relationship between nerve sprouting and epithelial wound closure. J. Cell Sci. 114, 6031–6039 (2001).
6. Barker, A. R., Rosson, G. D. & Dellon, A. L. Wound healing in denervated tissue. Ann. Plast. Surg. 57, 339–342 (2006).
genes involved in neuroprotection, neuroinflammation, and nerve regeneration. *Brain Behav. Immun.* 24, 1254–1267 (2010).

37. Britsch, S. et al. The transcription factor Sox10 is a key regulator of peripheral nerve development. *Genes Dev.* 15, 66–78 (2001).

38. Harris, M. L. et al. A dual role for Sox10 in the maintenance of the postnatal melanocyte lineage and the differentiation of melanocyte stem cell progenitors. *PloS Genet.* 9, e1003644 (2013).

39. Bremer, M. et al. Sox10 is required for Schwann-cell homeostasis and myelin maintenance in the adult peripheral nerve. *GLIA* 59, 1022–1032 (2011).

40. Ansell, D. M., Campbell, L., Thomason, H. A., Brass, A. & Hardman, M. J. A statistical analysis of murine incisional and excisional acute wound models. *Wound Repair Regen.* 22, 281–287 (2014).

41. Klingberg, F., Hinz, B. & White, E. S. The myofibroblast matrix: implications for tissue repair and fibrosis. *J. Pathol.* 229, 298–309 (2013).

42. Walker, J. T., McLeod, K., Sim, S., Conway, S. J. & Hamilton, D. W. Periostin as a multifunctional modulator of the wound healing response. *Cell Tissue Res.* 365, 453–465 (2016).

43. Goebbels, S. et al. Elevated phosphatidylinositol 3,4,5-trisphosphate in glia triggers cell-autonomous membrane wrapping and myelination. *J. Neurosci.* 30, 1393–1396 (2010).

44. Werner, S. A novel enhancer of the wound healing process. *Am. J. Pathol.* 179, 2144–2147 (2011).

45. Inman, G. J. et al. SB-431542 is a potent and specific inhibitor of transforming growth factor-beta superfamily type I activin receptor-like kinase (ALK) receptors ALK4, ALK5, and ALK7. *Mol. Pharmacol.* 62, 65–74 (2002).

46. Lipinski, J. E. et al. Activated Schwann cells in pancreatic cancer are linked to analgesia via suppression of spinal astroglia and microglia. *Gut* 65, 1001–1014 (2016).

47. Qin, Y. et al. TNF-alpha as an autocrine mediator and its role in the activation of Schwann cells. *Neurochem. Res.* 33, 1077–1084 (2008).

48. Joseph, N. M. et al. Neural crest stem cells undergo multilineage differentiation in developing peripheral nerves to generate endoneurial fibroblasts in addition to Schwann cells. *Development* 131, 5599–5612 (2004).

49. Kragl, M. et al. Cells keep a memory of their tissue origin during axolotl limb regeneration. *Science* 335, 87–94 (2012).

50. Kaukua, N. et al. Glial origin of mesenchymal stem cells in a tooth model system. *Nature* 513, 551–554 (2014).

51. Pardal, R., Ortega-Sáenz, P., Durán, R. & López-Barrero, J. Glia-like stem cells sustain physiologic neurogenesis in the adult mammalian carotid body. *Cell* 131, 364–377 (2007).

52. Küri, M. et al. Cells keep a memory of their tissue origin during axolotl limb regeneration. *Nature* 460, 60–65 (2009).

53. Götz, M., Sirko, S., Beckers, J. & Furler, M. Reactive astrocytes as neural stem or progenitor cells: in vivo lineage, in vitro potential, and genome-wide expression analysis. *GLIA* 63, 1452–1468 (2015).

54. Finzsch, M. et al. Sox10 is required for Schwann cell identity and progression beyond the immature Schwann cell stage. *J. Cell Biol.* 199, 701–712 (2012).

55. Desmoulère, A., Geinoz, A., Gabbiani, F. & Gabbiani, G. Transforming growth factor-beta 1 induces alpha-smooth muscle actin expression in granulation tissue myofibroblasts and in quiescent and growing cultured fibroblasts. *J. Cell Biol.* 122, 103–111 (1993).

56. Zhang, Z. et al. Recombinant human decorin inhibits cell proliferation and downregulates TGF-beta1 production in hypertrophic scar fibroblasts. *Burns* 33, 634–641 (2007).

57. Yamazaki, S. et al. Nonmyelinating Schwann cells maintain hematopoietic stem cell niches: evidence for a regulatory function of. *Ann. N. Y. Acad. Sci.* 1158 (2011).

58. Kuragai, K. et al. Nerve dependence in tissue, organ, and appendage regeneration. *Science* 318, 670–673 (2003).

59. Fontana, X. et al. Muscle cells derive from bone marrow-derived stromal cells, but not neural crest cells. *Am. J. Pathol.* 168, 1340–1355 (2006).

60. Tofariss, G. K., Patterson, P. H., Jesen, K. R. & Miskry, R. D. Neuronal activity: a new model for studying the effects of neural activity on cell survival. *GLIA* 56, 1637–1648 (2007).

61. Ma, K. H. et al. Regulation of peripheral nerve myelin maintenance by gene expression through polymorphic repressive complex 2, *J. Neurosci.* 35, 8640–8652 (2015).

62. Le, N. et al. Nax proteins are essential for peripheral nervous system myelination. *Nat. Neurosci.* 18, 189–197 (2002).
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Author contributions
J.D., V.P., U.R., and L.S. designed the experiments; V.P., J.D., and S.M.S. performed the experiments; V.P., J.D., and L.S. analysed the data. O.S. and S.W. provided expertise for the murine injury protocol; O.S., M.G., and S.W. initiated the project. J.D., V.P., and L.S. wrote the manuscript. M.W. and U.S. provided mouse lines. All authors made important comments to the manuscript.

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