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Dynamic interplay between IL-1 and WNT pathways in regulating dermal adipocyte lineage cells during skin development and wound regeneration

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SUMMARY

SUPPLEMENTAL INFORMATION

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DECLARATION OF INTERESTS

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AUTHOR CONTRIBUTIONS

Conceptualization, L.S., Xiaowei Zhang, M.V.P., R.L.G., and L.-j.Z.; methodology, L.S., Xiaowei Zhang, S.W., Y. Liu, W.L., Q.Y., M. Yin., J.L., Y. Liao, R.W., T.X., Xinyuan Zhang, Y.Y., F.L., S.H., W.Z., M. Yang, and L.-j.Z.; investigation, L.S., Xiaowei Zhang, S.W., Y. Liu, Q.Y., M.Y., and J.L.; resources, C.F.G.-J., J.H., C.-M.T., and C.J.; data curation, L.S., Xiaowei Zhang, S.W., Y. Liu, and L.-j.Z.; writing – original draft, L.S., Xiaowei Zhang, C.F.G.-J., M.V.P., R.L.G., and L.-j.Z.; writing – review & editing, L.S., Xiaowei Zhang, C.F.G.-J., R.R., M.V.P., R.L.G., and L.-j.Z.

R.L.G. is a co-founder, scientific advisor, consultant for, and has equity in MatriSys Biosciences.

Dermal adipocyte lineage cells are highly plastic and can undergo reversible differentiation and dedifferentiation in response to various stimuli. Using single-cell RNA sequencing of developing or wounded mouse skin, we classify dermal fibroblasts (dFBs) into distinct nonadipogenic and adipogenic cell states. Cell differentiation trajectory analyses identify IL-1-NF- κ B and WNT- β -catenin as top signaling pathways that positively and negatively associate with adipogenesis, respectively. Upon wounding, activation of adipocyte progenitors and woundinduced adipogenesis are mediated in part by neutrophils through the IL-1R-NF- κ B-CREB signaling axis. In contrast, WNT activation, by WNT ligand and/or ablation of *Gsk3*, inhibits the adipogenesis is seen in human keloids. These data reveal molecular mechanisms underlying the plasticity of dermal adipocyte lineage cells, defining potential therapeutic targets for defective wound healing and scar formation.

Graphical Abstract



In brief

Dermal adipocyte lineage cells are highly plastic cells that expand or retract in response to stimuli. Zhang and colleagues define the heterogeneity of these cells in developing or wounded mouse skin and demonstrate that dermal adipogenesis and lipolysis are regulated by an antagonistic interplay between the IL-1-pCREB and the WNT- β catenin pathways.

INTRODUCTION

The skin, comprised of layers that include the epidermis, dermis, and dermal white adipose tissue (dWAT), is the largest organ of the human body and a critical barrier against external insults. Emerging studies show that dWAT has a high degree of plasticity and can undergo drastic expansion and/or regression during skin development, during normal hair cycling, and in response to external stimuli, such as wounding, inflammation, and/or bacterial infection.^{1–3} However, the cellular and molecular mechanisms underlying dWAT remodeling are poorly understood.

Dermal fibroblasts (dFBs), the most prevalent cell type in the dermis, are highly heterogeneous. In neonatal mouse skin, dFBs are committed to forming the upper dermal papillary (DPP4⁺LY6A⁻), follicle-associated dermal papilla (DP), lower dermal reticular (DPP4⁻DLK1⁺LY6A⁻), and hypodermal (DLK1^{-/+}LY6A⁺) layers.^{4–6} Adipocyte progenitors (APs) and/or preadipocytes (pAds) are present in both reticular and hypodermal layers, and these adipocyte-lineage cells are poised to differentiate into adipocytes, forming the dWAT layer.⁴ However, the origin, cellular complexity/heterogeneity, and molecular mechanism of dermal adipocyte differentiation from their heterogeneous progenitors are only partially understood.

Skin wound healing is a highly orchestrated process; it includes an early hemostatic and inflammatory phase, then a proliferative phase, and finally a remodeling phase that can last weeks to years.⁷ During the inflammatory phase, myeloid immune cells are recruited, and fibroblasts migrate from the wound periphery, forming granulation tissue (GT), a temporary scaffold holding the wound.⁸ During the wound proliferative phase, myofibroblasts are transdifferentiated from fibroblasts to synthesize extracellular matrix (ECM) molecules; however, sustained activation of myofibroblasts leads to the development of fibrotic diseases like keloids.⁹

Emerging roles of adipocyte-lineage cells in skin wound healing have been reported. Following inflammation, APs proliferate, and new lipid-laden adipocytes are generated in wound GT, and inhibition of adipogenesis impairs dermal repair.¹⁰ Recently, a subset of hypodermal DPP4⁺LY6A⁺ fascia fibroblasts with remarkable reparative potential was identified, and genetic ablation or blocking of their migration into wound sites led to chronic wounds. In adipose tissue, DPP4⁺LY6A⁺ fascia fibroblasts have been identified as APs that give rise to ICAM1⁺DLK1⁺ pAds.^{11,12} However, whether skin DPP4⁺LY6A⁺ dFBs are adipogenic and their cell differentiation trajectory during wound repair remain largely unknown. Furthermore, the molecular cues that trigger wound-related adipogenesis responses remain undefined.

Here, we obtained skin samples from developing, homeostatic, and wounded mice and applied single-cell RNA sequencing (scRNA-seq) followed by immunostaining (immunohistochemistry [IHC]) and cell sorting to define dFB populations. scRNA-seq predicted the cellular trajectories through which dermal adipocytes differentiated from their heterogeneous APs. The behavior of these cells was examined during wound repair. Finally human keloid samples were analyzed to determine the human relevance of our findings.

Our study provides insight into the molecular and cellular mechanisms underlying dermal adipogenesis and/or lipolysis during skin development and wound regeneration.

RESULTS

Characterization of heterogeneous dermal adipocyte-lineage cells in developing neonatal skin

We performed scRNA-seq of cells isolated from neonatal mouse skin, in which dFBs are actively developing from their precursors,⁴ allowing us to capture a continuum of cell states spanning differentiation and specification. $Pdgfra^+$ (a pan-fibroblast marker^{4,13}) dFBs, $Krt10/14^+$ keratinocytes, $Ptprc/Cd45^+$ immune cells, and $Rgs5^+$ pericytes were identified as the major cell types present in neonatal mouse skin (Figures 1A and S1A).

Selected *Pdgfra*⁺ dFBs were reclustered into 12 clusters (C0–C11) (Figures 1A, S1B, and S1C), which were grouped into three distinct developmental cell states (Figures 1B, 1C, and S1D) and defined based on marker gene expression as shown by bubble and/or violin plots (Figures 1C and S1E). Specifically, state 1 cells (C9, C2, C5) expressed high levels of *Cd24* and *Trps1* but lacked *Ly6a* (Figures 1C and S1E). IHC analysis of neonatal skin with distinct anatomical layers (Figures S1F–S1G) showed that CD24 and TRPS1 were co-expressed at high levels in PDGFRA⁺ dFBs located around hair follicles at the upper dermis or in the hair bulb DP structure (Figure 1D), where DPP4, LY6A (Figure 1E), and THY1 (Figure S1H) were not expressed or expressed at low levels. ALPL, another marker enriched in state 1 cells (Figures 1C and S1E), co-expressed specifically with TRPS1 in the DP (Figure S1H). We therefore termed the state 1 cluster cells as perifollicular dFBs.

State 2 cells (C7, C0, C3) expressed markers related to papillary (PAP) and/or reticular (RET) dFBs and lacked *Ly6a* expression (Figure 1C). Specifically, the C7 cluster expressed established PAP dFB markers (*Dpp4, Lrig, Entpd1*, and *Lrig1*) and medium level of *Trps1*, whereas the C3 cluster expressed genes associated with RET dFBs (*Dlk1* and *Ptgfr*) and a lower level of *Trps1* (Figures 1C and S1E). IHC confirmed that the upper dermal PAP cells were TRPS1⁺DPP4⁺LY6A⁻DLK1⁻, whereas the lower dermal RET cells were TRPS1⁻DPP4⁻LY6A⁻DLK1⁺ (Figures 1F, S1I, and S1J). Therefore, the C7 and C3 clusters were defined as PAP and RET dFBs, respectively.

State 3 cells (C1, 4, 8, 10, 6, 11) are highly heterogeneous (Figure S1D). These cells expressed marker genes related to adipogenesis but lacked *Trps1* (Figures 1C and S1E); thus these clusters were termed adipocyte lineage cells. Specifically, C4 cells (pAd) and C8 cells (pAd/Ad) expressed conventional pAd and/or adipocyte markers but lacked *Dpp4*. IHC analysis confirmed that the LY6A⁺DPP4⁻ pAds were detected exclusively in dWAT, where BODIPY⁺ adipocytes were located (Figures 1E and S1G). Interestingly, we found that C4 pAd cells expressed high levels of *Lgr5* (Figures S1K and S1L), a well-established marker for hair follicle stem cells.¹⁴ Furthermore, cluster C1 was termed RET-AP, because it lacked *Ly6a* but expressed a medium level of *Dlk1* and high levels of *Cebpb* (Figures 1C and S1E), a central transcription factor for dermal adipogenesis.¹⁵

Among all state 3 clusters, *Dlk1* was expressed the highest in clusters C6, C10, and C11 (Figures 1C and S1E). IHC (Figures 1G and 1H) showed that DLK1 was highly expressed within the interstitial space above and below the panniculus carnosus (p.c.) muscle, a space termed "hypodermal interstitium" (HI) in this study. This HI space was also enriched with LY6A⁺DPP4⁺ cells (Figure S1M). Furthermore, C10 was termed HI-pAd because it co-expressed pAd/Ad marker genes (*Lpl* and *Icam1*); C6 was termed HI-Areg (adipogenesis regulator cells), because it expressed a panel of Areg marker genes, including *F3*, *Fmo2*, and *Mgp*.^{16,17} Finally, cluster C11 was termed HI-AP, because it co-expressed AP marker genes (*Dpp4*, *Ly6a*, *Cd34*, and *Wnt2*) (Figures 1C and S1E). IHC validated that WNT2⁺ PDGFRA⁺ (Figure S1N) and WNT2⁺DPP4⁺LY6A⁺cells (Figure S1O) were detected primarily in the HI space.

dFBs are a major cellular source of skin ECM.⁹ Analysis of ECM-related genes revealed that the *Cd24^{hi}* clusters expressed *Col23a1* and *Col13a1* but lacked *Col1a1* and *Col3a1* (Figures 1C and S1E). Interestingly, the HI-AP cluster was enriched with a panel of collagen genes and ECM genes related to elastic network and fibrillogenesis,^{18,19} such as *Mfap5*, *Fbn1*, and *Fn1* (Figures 1C and S1E), indicating that HI-AP could be involved in providing structural support to the skin.

Next, we developed fluorescence-activated cell sorting (FACS) strategies, based on our and others' previously published FACS protocol for dFBs,^{20–23} to isolate perifollicular dFBs (CD24^{hi}LY6A⁻), pAds (LY6A⁺DPP4⁻), and HI-APs (LY6A⁺DPP4⁺) from neonatal dFBs (Figure S1P). qRT-PCR profiling analysis (Figures 1I and S1Q) showed that the sorted CD24^{hi}LY6A⁻ cells highly expressed genes that were enriched in C2/C9 perifollicular clusters, such as *Trps1*, *Alpl*, and *Col23a1*; the sorted LY6A⁺DPP4⁻ dFBs expressed a panel of pAd/Ad-related genes; and the sorted LY6A⁺DPP4⁺ dFBs expressed higher levels of genes that were enriched in the C11 HI-AP cluster, such as *Dlk1*, *Wnt2*, *Smpd3*, and *Mfap5*. Using *in vitro* adipocyte and osteocyte differentiation assays, we found that the CD24⁺LY6A⁻ cells were non-adipogenic, and both the pAds and the HI-APs were highly adipogenic (Figures 1J and 1K), whereas only HI-APs were osteogenic (Figures 1L and S1R), suggesting that DPP4⁺LY6A⁺ cells are multipotent progenitors and DPP4⁻LY6A⁺ cells are committed pAds.

In line with these functional results, trajectory analysis predicted that C11 (HI-AP) was the origin of the differentiation trajectory of adipocyte lineage cells (Figures 1B and S1S). In addition, RNA velocity analysis (Figure 1M) predicted that both HI-pAd/Areg and RET-AP dFBs actively differentiated into pAd/Ad, and C0, C1, C4, and C9 further converge into C2⁻ perifollicular dFBs. These trajectories need to be validated by lineage tracing studies in the future.

Together, using scRNA-seq, we classified the neonatal developing dFBs into three distinct states (Table S1) and defined the heterogeneous adipocyte-lineage cells located in the RET dermis, dWAT, and HI (Figure 1N).

The antagonistic interplay between WNT- β -catenin and IL-1-immune signaling pathways in regulating dermal adipogenesis

We next sought to identify the key molecular pathways that regulate adipogenesis in the developing skin. Gene ontology (GO) analyses identified the IL-1 and WNT pathways as among the top activated or silenced pathways, respectively, during the conversion of various dFB subsets to pAd/Ad (Figures 2A and S2A). Gene expression plots (Figures 2B and S2B) showed that WNT-related genes were specifically enriched in the perifollicular and/or PAP dFB clusters, whereas a panel of immune- and inflammation-related genes was selectively enriched in the pAd/Ad-related clusters. In addition, transcription factor (TF) analysis identified *Cebpb, Pparg*, and several inflammatory TFs (*Nfkb1* and *Junb*) as top TFs activated in pAd/Ad-related clusters (Figure S2C).

To determine the direct effect of WNT and IL-1 on adipocyte differentiation, neonatal dFBs were treated with WNT3A, lithium, and/or IL-1 β . Lithium, a GSK3 β inhibitor, is commonly used to achieve broad-spectrum activation of the WNT- β -catenin pathway.²⁴ RNA-seq results revealed that WNT3A and lithium treatment robustly induced the expression of genes associated with perifollicular dFBs (*Cd24, Alpl*), whereas IL-1 β treatment induced the expression of a panel of pro-inflammatory genes (*Cxcl12, II1r1*) and several pAd/Ad marker genes (*Pparg2, Fabp4*) (Figure S2D). Venn diagram analysis showed that genes upregulated by IL-1 β shared a high degree of similarity with genes downregulated by WNT3A + lithium (Figure S2E). Functionally, WNT3A potently suppressed the adipogenic potential of neonatal dFBs (Figures 2C and 2D) and secretion of CAMP and FABP4, which can be partially restored by adding IL-1 β (Figure 2E). Furthermore, WNT3A-mediated activation of β -catenin (Figures 2F and 2G), suppression of adipocyte- or immune-related genes, and induction of *Axin2, Acta2,* and *Col7a1* were partially reversed by IL-1 β (Figures 2H and S2F–S2I).

Time-course phosphoblotting analyses (Figures 2I and 2J) revealed that IL-1 β treatment increased NF- κ B, p38 and ERK phosphorylation levels but suppressed AKT phosphorylation, which is required for β -catenin stabilization.²⁵ IL-1 β treatment led to a continuous loss of protein levels of β -catenin (active or total) (Figures 2I, 2J, and S2J). In contrast, IL-1 β treatment rapidly induced phosphorylation of CREB, a critical TF involved in CEBP β activation and promoting adipogenesis,²⁶ and IL-1 β -mediated CEBP β phosphorylation occurred in a delayed manner (Figures 2I and 2J).

We found that AKT inhibition by wortmannin promoted loss of β -catenin expression, whereas p38 inhibition by SB202190 or NF- κ B inhibition by Bay117028 led to a decrease in the phosphorylation levels of both CREB and CEBP β (Figures 2K and S2K–S2M). In line with these results, the induction of adipocyte- or immune-related genes by IL-1 β was largely blocked by p38 or NF- κ B inhibitors (Figure 2L). These results indicate that IL-1 β promotes β -catenin degradation by inhibiting AKT and promotes activation of proadipogenic molecules by stimulating p38 and NF- κ B (Figure 2M).

We next analyzed skin biopsies from neonatal $II1r1^{-/-}$ and $II1r1^{+/+}$ littermates to determine the role of IL-1 signaling in adipogenesis. We found that II1r1 deficiency led to a blockage of CREB phosphorylation in the neonatal mouse skin (Figure 2N). As the result, formation

of BODIPY⁺ adipocytes (Figure 2O) and the expression of adipocyte-related genes and the inflammatory cytokine *Cxcl5* were suppressed, whereas expression of WNT-related *Axin2* was enhanced upon *Il1r1* deletion (Figure 2P). Taken together, our results indicate that activation of the IL-1R-CREB signaling axis may play a role in promoting dermal adipogenesis during skin development.

Tracking dermal adipogenesis and fibrogenesis during skin maturation

To track dermal adipogenesis during skin maturation and aging, dorsal skin samples were collected from male mouse skin throughout the postnatal lifespan for analysis. Lipid staining and bulk RNA-seq of skin samples (Figures 3A and S3A) showed that dermal adipocytes progressed from an immature state at birth (characterized by small lipid droplets and high expression levels of *Camp* and *Mest*), to a mature state at 3 weeks (large lipid droplets), following which dermal cells and adipocytes were gradually lost during skin maturation and aging from 2 months to 1–2 years. In contrast, dermal collagen content and the expression levels of several ECM molecules increased up to 2 months of age and then progressively decreased during aging (Figures 3A and S3A).

To determine the cellular and molecular changes in dermal adipocyte-lineage cells during skin maturation, scRNA-seq was performed on skin cells isolated from mice at neonatal stage (P1) and 3 weeks and 2 months of age (Figure 3B). Clustering analysis identified 30 clusters, which were classified into nine cell types (Figures 3C–3F and S3B–S3D). Interestingly, we found that among all cell types, *Pdgfra*⁺ dFBs contained several age-unique clusters (Figure 3F).

To further inspect dFB heterogeneity, *Pdgfra*⁺ dFBs were reclustered (Figure 3G), and these dFB clusters from the three age groups were clearly separated on the tSNE (t-distributed stochastic neighbor embedding) plots in an age-dependent manner (Figure 3H). Based on the established neonatal dFB markers (Figure 1), we identified the corresponding adult dFB clusters (Figures 3H, 3I, and S3E). In contrast to neonatal DP cells that expressed *Alpl, Sox2*, and *Lef1*, adult DP cells lost activated WNT signals (*Lef1, Nkd2*) and expressed *Crapb1* and the WNT antagonist *Dkk2*, in line with previous studies^{27,28} (Figures 3I and S3E). Similar to neonatal dFBs, adult PAP dFBs were enriched in *Grem1* and *Entpd1* (Figures 3I and S3E), and *ENTPD1* (also known as CD39) has been shown to mark human adult skin PAP dFBs.²⁹ In adult skin, *Trps1* was also an exclusive marker for all *Ly6a*⁺ adipocyte-lineage cells (Figures 3I and S3E). Interestingly, we found that HI-pAd/Areg progressively lost *Dlk1* but gained Areg signature gene signatures (*F3, Fmo2, Mgp*, and *Gdf10*) during maturation (Figures 3I and S3E).

Next, we examined how genes related to the IL-1, WNT, and TGF- β pathways were altered in key dFB clusters during skin maturation. First, analysis of bulk skin RNA-seq showed that the expression of WNT- and IL-1-related genes was progressively lost, whereas TGF- β related genes were induced postnatally during skin maturation (Figure 3J). In line with these results, analysis of key dFB clusters showed that WNT- or IL-1-related genes were inhibited in pAd/Ad and HI-AP, whereas TGF- β -related genes were notably induced in PAP/RET and pAd/Ad but not HI-AP clusters age dependently (Figure 3K). As a result, ECM genes were induced and adipogenesis-related genes were inhibited in PAP/RET or pAd/Ad clusters

age dependently, whereas HI-APs started to lose ECM gene expression at 2 months of age (Figure 3K). These age-dependent changes in WNT, IL-1, and TGF- β signaling activities in adipocyte-lineage cells may contribute to the adipogenic-to-fibrogenic switch of these cells during skin maturation.

IHC confirmed that, similar to neonatal skin, the fascia layer of adult skin was enriched with LY6A^{hi}DPP4^{hi} HI-APs, and LY6A⁺DPP4⁻ pAds were detected exclusively in the dWAT layer of adult skin (Figure 3L). In addition, DLK1⁺ cells were detected only within the interstitial space around the p.c. muscle layer in the 3-week skin and were largely lost by 2 months (Figure 3M), which is consistent with the *Dlk1* expression pattern shown by scRNA-seq (Figures 3I and S3E). scRNA-seq showed that HI-AP% increased during skin maturation (Figure S3F). FACS analysis of PDGFRA⁺ dFBs validated that, while pAd% increased postnatally for up to 3 weeks, HI-AP% continued to increase for up to 2 months and then declined in 1-year-old skin (Figures S3G, 3N, and 3O). In contrast, CD24⁺LY6A⁻ dFBs, identified as the neonatal perifollicular dFBs, were rapidly lost postnatally (Figures S3H and S3I), which is consistent with the *Cd24* scRNA-seq plots (Figures 3I and S3E). Together, these results showed that skin maturation is associated with age-dependent inactivation of WNT and IL-1 signaling, depletion of adipogenic pAd, and accumulation of inactive HI-APs.

Tracking dermal adipogenesis and fibrogenesis during skin wound healing

Next, we studied how dermal adipogenesis may be reinitiated upon wounding in adult homeostatic skin. BODIPY staining showed that numerous adipocytes were transiently detected in the wound center GT, peaking at wound day (w.d.) 3, and lipolysis was apparent at w.d. 5–7, indicated by the presence of dispersed small lipid droplets, and by w.d. 14, adipocytes were cleared from the wound bed (Figure 4A). To determine whether wound-associated adipocytes were generated *de novo* from adipogenesis, we induced CRE activity in *Adipoq*-CreERT2;mTmG mice by tamoxifen (TAM) application prior to wounding to label preexisting adipocytes with GFP (Figure S4A). Results showed that more than one-third of wound-associated COLIV⁺FABP4⁺ adipocytes were GFP⁻ (Figures S4B and S4C), demonstrating that newly differentiated adipocytes partially contribute to wound-induced adipogenesis.

During the wound-proliferative phase, fibroblasts are activated and differentiate into ACTA2⁺ myofibroblasts.⁹ Here, IHC showed that at w.d. 3, DPP4⁺LY6A⁺ HI-APs were detected at the wound periphery and bottom, whereas DPP4⁻LY6A⁺ pAds/Ads were mainly detected in the wound mid-region (Figures 4B and 4C). By w.d. 7, pAds/Ads were largely lost, and ACTA2⁺DPP4⁻LY6A⁻ myofibroblasts were abundantly detected in the upper wound bed (Figures 4B and 4C). By day 14, myofibroblasts were largely lost, and DPP4⁻LY6A⁺ pAds were regenerated above the HI-AP fascia layer (Figure 4B).

scRNA-seq analysis of wound skin cells was next performed to determine the cell differentiation trajectory (Figures 4D and 4E). A total of 15 different cell types, including dFBs, neutrophils, macrophages, T cells, and keratinocytes (KCs), were identified (Figures 4D, 4E, S4D, and S4E). We found that neutrophils (~30% of all cells) and dFB clusters (~50% of all cells) were the major cell types that infiltrated wound GT on w.d. 2, and

a subsequent decrease in neutrophils at w.d. 7 was accompanied by an increase in dFB heterogeneity and regeneration of other cell types, such as KCs and vessel cells (Figure 4F).

To define heterogeneous dFBs, Pdgfra⁺ dFBs were reclustered into 13 subclusters (r0r12; Figures 4G–4H and S4F–S4G), which were defined based on the expression of dFB marker genes identified in Figure 1 and myofibroblast markers (Acta2, TagIn). We found that adipocyte-lineage cells, including HI-AP (r0, r2, r12) and pAd/Ad (r3) cell clusters, represented >85% of dFBs in w.d. 2 GT, whereas by w.d. 7, pAd/Ad cells were largely lost, and Acta2⁺ myofibroblasts and $Cd24^{hi}Alpl^+$ dFBs represented ~1/3 of the w.d.7 dFBs (Figure S4H). Gene expression plots (Figures 4H and S4F-S4J) showed that the r2 HI-AP cluster, which was transiently generated in w.d. 2 GTs, expressed high levels of proliferation marker genes (Ki67 and Pcna) and IL-1-inducible chemokine genes (Cxcl2/5/12), suggesting that these cells were highly proliferative and inflammatory. RNA velocity analysis (Figure 4I) and cell-rank analysis (for calculation of cell-cell transition probabilities³⁰) (Figure 4J) predicted that at w.d. 2, the activated r2 HI-APs were poised to differentiate toward the r3 pAds/Ads, which were in turn poised to differentiate into other dFB clusters, including the r7 myofibroblasts. Together, these results suggest that activated HI-APs may contribute to pAd/Ad generation during the wound-inflammatory phase, and pAd/Ad-to-myofibroblast transdifferentiation may occur during the subsequent wound-proliferative phase. Future studies using lineage tracing mouse models specific for HI-APs and pAd/Ad are needed to confirm these cell differentiation trajectories.

Wound-induced adipogenesis is regulated by neutrophils and the IL-1 signaling axis

FACS analysis confirmed that the percentage of LY6A⁺DPP4⁺ HI-AP peaked at approximately w.d. 2, and loss of HI-APs was accompanied by gain of ACTA2⁺LY6A⁺ myofibroblasts at w.d. 7 in wound GTs (Figures 5A–5C and S5A). The transition from HI-AP (*Dpp4*⁺) to pAd/Ad (*Cebpb^{hi} Fabp4*⁺) and myofibroblasts (*Acta2*⁺) during wounding can also be visualized at the single-cell level (Figure S5B).

SCENIC TF analysis identified several key adipogenesis TFs (*Cebpb, Creb, Srebf1*) as the top enriched TFs driving dFB activation at w.d. 2 (Figure 5D). In addition, GO pathway analysis of the r0 HI-AP cluster identified cellular responses to stress/stimuli and IL-1 signaling as the top upregulated pathways after wounding (Figure 5E). Furthermore, violin plots showed that wounding led to a decrease in the expression of HI-AP marker genes (*Smpd3, Dpp4*) and an increase in the expression of IL-1- and adipogenesis-related genes in HI-APs. (Figure 5F). In addition, CellChat analysis identified IL-1 as one of the top incoming signals for HI-AP dFBs (Figure S5C), and IL-1-IL-1R signaling network analysis identified neutrophils as the most prominent source of IL-1 ligand acting on dFB clusters at w.d. 2 (Figures 5G and S5D). Expression plots confirmed that *II1b* was predominantly produced by neutrophils, whereas the IL-1 receptor (*II1r1*) and IL-1-inducible chemokines (*Cxcl1/12/5*) were expressed primarily by dFBs (Figures 5H, S5E, and S5F). Together, these results suggest that IL-1 released from neutrophils may drive the wound-induced adipogenesis response during the early wound-inflammatory phase.

FACS and immunostaining analyses (Figures S5G, S5H, and 5I) confirmed that Ly6G⁺ neutrophils were the dominant infiltrating immune cell type (>60% of all CD11B⁺ myeloid

cells) during w.d. 2–3, and neutrophils were cleared from the wound by w.d. 7. Depletion of neutrophils (Figures S5I and S5J) led to a significant reduction in the DPP4⁺ wound GT area (Figure 5J) and impaired wound-induced adipogenesis (Figure 5K), decreased CAMP protein expression (Figure S5K), and decreased pCREB signal (Figures S5L and S5M) in the migrating DPP4⁺ HI-APs. In addition, depletion of neutrophils led to decreased wound-induced *II1b* expression (Figure 5K), demonstrating that neutrophils are the major cellular source of IL-1β during the early stages of wound healing.

Wild-type (WT) and $II1r1^{-/-}$ mice were subjected to skin wounding to validate the role of IL-1 in wound-induced adipogenesis. We found that II1r1 deficiency led to profound inhibition of pCREB activation in DPP4⁺ HI-APs (Figures 5L and 5M), but not in the migrating epidermal KCs (white arrow) or other DPP4⁻ dermal cells, indicating that this IL-1-pCREB signaling axis is specific for the activation of hypodermal cells. In line with pCREB inhibition, the wound-mediated induction of *Cxcl5*, *Cxcl12*, and adipogenesisrelated genes was significantly reduced in $II1r1^{-/-}$ mice compared with WT controls (Figure 5N). Furthermore, administration of an NF- κ B inhibitor to skin wounds also led to decreased pCREB expression in HI-APs and pAds (Figure S5N). These results indicate that neutrophils contribute to HI-AP activation and wound-induced adipogenesis by activating the IL-1R-NF- κ B-CREB pathway.

Activation of WNT and TGF- β signaling pathways in adipocyte-lineage cells during the wound-proliferative phase

We found that a peak in the expression of lipogenesis genes at w.d. 3 was followed by the induction of lipolysis genes (*Atgl* and *Hsl*) at approximately w.d. 5, prior to the induction of *Acta2* and ECM genes around w.d. 5–7 (Figures 6A and S6A). In addition, loss of FABP4⁺ adipocytes was accompanied by transient formation of ACTA2⁺ myofibroblasts on w.d. 7 (Figure S6B), suggesting that adipocyte lipolysis and dedifferentiation may contribute to myofibroblast formation during the wound-proliferative phase.

To determine whether adipocytes (preexisting or newly generated) contribute to the formation of myofibroblasts, we induced CRE activity in *Adipoq*-C*re*ERT2;mTmG mice by TAM application during w.d. 0–6, thereby investigating both the inflammatory and the proliferative phases of wounding (Figure 6B). Mature adipocytes labeled with GFP were detected in the upper wound bed region, where these GFP⁺ cells co-expressed ACTA2 (Figures 6B and S6C), indicating that myofibroblasts can be partially derived from mature adipocytes during wound healing.

SCENIC TF analysis identified several WNT-related TFs as top enriched TFs in dFBs at w.d. 7 (Figure 6C), including *Lef1, Sox4, Sox11, Hoxc8*, and *Smarca4*, all of which regulate cell development or carcinogenesis by activating the WNT- β -catenin signaling pathway.^{31–35} Co-staining of active β -catenin with DPP4 and/or ACTA2 revealed that at w.d. 3 the active β -catenin signal was barely detectable, but at w.d. 7 β -catenin was robustly activated in the upper wound region, where ACTA2⁺ myofibroblasts were generated, and by w.d. 14, active β -catenin was lost and myofibroblasts disappeared from the wound GT (Figures 6D, S6D, and S6E). Although nuclear β -catenin signal can be detected in cultured dFBs upon WNT3A

stimulation (Figure S6F), it was difficult to distinguish β -catenin within nuclei from that within the cytosol in cells within whole wound tissue.

Gene plot analysis showed that both WNT and TGF- β signaling activities were prominently activated in the r7 myofibroblast cluster (Figure S6G), indicating that the crosstalk between TGFB and WNT signaling may promote adipocyte-to-myofibroblast transition during wound healing.

WNT activation in adipocytes promotes adipocyte dedifferentiation and myofibroblast formation

The TGF- β pathway plays a role in triggering lipolysis and dedifferentiation of mature adipocytes^{36,37}; moreover, physical stress was observed to trigger adipocyte dedifferentiation via WNT- β -catenin signaling.³⁸ Here, we found that a combination of both lithium and TGF- β robustly promoted cell elongation and the breakdown of large lipid droplets into numerous smaller lipid droplets (Figures 6E and 6F), which are characteristic of dedifferentiating adipocytes.^{38,39} Furthermore, co-treatment with both lithium and TGF- β 2 led to the most significant increase in ACTA2⁺ cell percentage (Figure S6H), induction of pro-fibrotic genes, and suppression of *Cd36* expression (Figures 6G and S6I).

To lineage trace adipocyte dedifferentiation *in vitro*, adipocytes were differentiated from *Adipoq*-C*re*ERT2;*mTmG* dFBs and labeled by GFP in the presence of TAM, then labeled cells were treated with lithium and TGF- β to trigger lipolysis (Figures 6H, S6J, and S6K). We found that, while lithium or TGF- β alone had some effects, co-treatment profoundly changed GFP⁺ adipocytes from rounded to elongated morphology and promoted the internalization of GFP from membrane to cytosol and the breakdown of the PLIN1⁺ lipid droplets (Figures 6I and S6L). Furthermore, ACTA2 was induced along the peripheral membrane of the elongated GFP⁺ cells upon co-treatment (Figures 6J and 6K), supporting the idea that ACTA2⁺ myofibroblasts can be generated *de novo* from dedifferentiating adipocytes.

Gsk3 deletion in adipocytes promotes myofibroblast formation during wound healing

To examine whether WNT activation in adipocytes promotes fibrogenesis *in vivo*, we genetically targeted both *Gsk3a* and *Gsk3b*, key inhibitory molecules for β -catenin,⁴⁰ from adipocytes by crossing *Adipoq*-CreERT2 mice with *Gsk3a/b*^{fl/fl} mice (*Gsk3^AD-KO*) (Figure 7A). Both GSK3 alleles must be deleted to achieve hyperactivation of the WNT- β -catenin signaling.⁴¹ We found that TAM application to *Gsk3^AD-KO* mice during wound healing led to specific depletion of GSK3 in PLIN⁺ adipocytes, but not in other non-adipocyte cells (Figures 7A and 7B), indicating that targeted *Gsk3* deletion in adipocytes was successful. Furthermore, targeted *Gsk3* deletion in adipocytes not only increased activated β -catenin signaling in ATGL⁺ (a critical lipolysis enzyme⁴²) adipocytes (Figures 7C and 7D), but also promoted myofibroblast formation (Figures 7E, 7F, and S7A–S7C) and increased CRABP1 expression (Figure S7D). However, *Gsk3* deletion decreased the pool of LY6A⁺DPP4⁻ pAds (Figures 7E, 7G, and S7C) and inhibited *Cxcl5* and *Camp* expression (Figures S7E and S7F) in skin wounds. These results show that WNT activation mediated by GSK3 deletion in

adipocytes dampens wound-induced inflammatory and adipogenic responses and promotes adipocyte-to-myofibroblast transition during the wound-proliferative phase.

Human keloid is associated with sustained WNT activation and inhibition of dermal adipogenesis

Finally, to determine whether our observations in mice are also relevant to humans, we evaluated the changes in dWAT in human skin samples from normal wounds and keloids. IHC staining showed that numerous lobular structures enriched with COLIV+FABP4+ adipocytes were present in the granular tissue of normal wounded skin, whereas COLIV+FABP4+ adipocytes were largely absent in the keloid skin dermis (Figure 7H), suggesting that adipogenesis was blocked in keloids. In addition, immunostaining revealed a strongly activated β-catenin signal in both ACTA2+ myo-FBs and COLIV+ cells in the keloid skin dermis (Figure 7I). Furthermore, reanalysis of published transcriptomic data of non-lesional skin or wounded skin from keloid-prone individuals and healthy matched subjects⁴³ showed that compared with normal wounds, keloid wounds expressed significantly lower levels of immune- or adipogenesis-related genes (*IL1B, CEBPB, PLIN3*), but higher levels of WNT-and/or TGF-β-related genes and ECM genes (Figures 7J and S7G). Together, these results indicate that unbalanced activation of WNT and immune pathways during wound healing may inhibit adipogenesis and promote fibrogenic development in keloid.

DISCUSSION

dWAT has been recognized as a functional skin layer with remarkable plasticity; it expands or regresses during development, wound healing, bacterial infection, and hair cycling.^{3,44} To trace the cellular and molecular changes in dWAT cells, we present here a compendium of skin development, maturation, and regeneration processes by scRNA-seq. First, we grouped the developing dFBs into distinct non-adipogenic and adipogenic cell states. Second, our data identified the WNT and IL-1 pathways as the key suppressing and activating signals, respectively, for dermal adipogenesis. Third, we found that skin injury triggered a sequence of events in wound GT, including rapid infiltration/activation of HI-APs and onset of adipogenesis during the wound-inflammatory phase, followed by adipocyte lipolysis/dedifferentiation and myofibroblast generation during the woundproliferative phase. Wound-induced adipogenesis was in part mediated by neutrophils and IL-1, and adipocyte dedifferentiation was in part mediated by activation of the WNT-βcatenin pathway. Finally, we showed that inhibition of dermal adipogenesis and sustained activation of dermal WNT signal was also associated with human keloid scar. Taken together, this study provides in-depth knowledge of the regulatory mechanisms of dWAT plasticity during skin development and wound regeneration at the single-cell level.

Defining the highly heterogeneous adipogenic and non-adipogenic dermal fibroblast subpopulations

scRNA-seq has recently been applied to understanding skin fibroblast heterogeneity.^{42,45–47} However, none of these studies focused on adipogenic cells, and most of these studies defined dWAT or fascia-related dFBs generally as only hypodermal or fascia dFBs, without

further defining cell heterogeneity in the hypodermis. In this study, we defined several adipocyte-lineage cell subpopulations, including RET-APs, pAds, adipocytes, and three hypodermal populations (HI-pAds, Aregs, and HI-APs). In addition, we identified a distinct pool of CD24^{hi} perifollicular dFB clusters that were non-adipogenic, enriched in developing skin, and rapidly lost during skin maturation.

This study also identified several new dFB markers at both the transcript and the protein levels. We identified *Trps1* as an exclusive marker for $Ly6a^+$ adipocyte-lineage cells. TRPS1 (both mRNA and protein) was found highly expressed in $Cd24^{hi}$ developing or follicle-associated dFBs, at intermediate levels in $Dpp4^+Ly6a^-$ PAP dFBs, and almost undetectable in $Ly6a^+$ pAd/APs. We also identified Lgr5, a previously known hair follicle stem cell marker,¹⁴ as a novel pAd marker in mouse skin and *in vitro* primary pAd culture (Figures 1 and 3). Furthermore, we found that a cluster of $Fmo2^+Gdf10^+$ Areg fibroblasts accumulated in the skin during maturation. Emerging studies have unraveled how Areg cells negatively regulate adipogenesis by secreting GDF10 and/or RSPO2,^{16,17,48} but whether the age-dependent increase in skin Areg contributes to the loss of the adipogenic potential of skin AP/pAds during aging needs further investigation.

Balance between the WNT- β -catenin and the IL-1-NF- κ B pathways in controlling dermal adipogenesis

WNT/β-catenin signaling is known to suppress adipogenesis by inhibiting the adipogenic TFs CEBP and PPARγ, preventing the differentiation of MSCs into fat cells.^{49–51} During skin development, WNT-β-catenin signaling is considered the earliest signaling event that instructs the migration of dFB progenitors from the somatic mesoderm to the dorsal skin dermis and the specification of these cells to fibroblasts.⁵² Inhibition of WNT-β-catenin signaling promotes the conversion of dermal-lineage cells to cartilage.⁵³ Here, we identified the WNT-β-catenin pathway as one of the most differentially suppressed pathways in adipogenic dFBs compared with other non-adipogenic dFB subclusters. Consistent with our results, active WNT gene signatures have been shown to be preferentially associated with *Lrig1*⁺ PAP dFBs, but not with *Ly6a*⁺ pAds.⁵⁴ Furthermore, forced stabilization of β-catenin in *Pdgfra*⁺ dFBs leads to a reduction in dWAT, followed by the development of prominent fibrosis.⁵⁵ These data suggest that, while activation of WNT-β-catenin in mesodermal stem cells is required for stem cell commitment to the dFB lineage during early development, the subsequent inhibition of WNT signaling during later skin development may be required for the commitment of dFBs to adipogenic subpopulations.

Our results suggest that IL-1 may antagonize WNT activity during dermal adipogenesis. Mechanistically, we showed that IL-1 β treatment not only triggered the activation of the pro-adipogenic CREB-CEBP β pathway, but also promoted β -catenin degradation by inhibiting AKT phosphorylation. Consistently, several immune-related and inflammatory pathways have been found to be enriched and activated upon targeted deletion of β -catenin in adipocytes.⁵⁶ Furthermore, IL-1 β increases the expression/activation status of CREB and its target gene CEBP β in a chondrocyte cell line.⁵⁷ IL-1B signaling also activated CEBP β in several other cell types, such as gastric epithelial cells⁵⁸ and myocytes,⁵⁹ suggesting that CREB-CEBP β activation maybe a common downstream event of IL-1 receptor signaling.

In contrast, an inhibitory role for IL-1 has been observed in regulating adipogenesis in skeletal muscle tissues,^{50,60} suggesting that a tissue-specific microenvironment may shape the differential response of AP to IL-1.

The crosstalk between WNT and TGF- β pathways in regulating adipocyte lipolysis and dedifferentiation

Using several independent approaches, we demonstrated that WNT cooperates with TGF- β signaling to promote adipocyte lipolysis and dedifferentiation of adipocytes to myofibroblasts during the wound-proliferative phase. In line with our findings, TGF- β signaling was shown to be the top enriched pathway activated by WNT3A treatment in adipocytes.⁵⁶ In addition, TGF- β and WNT are both potent pro-fibrotic signaling molecules involved in abnormal wound healing responses following skin or lung injury.^{54,61-63} During the pathogenesis of idiopathic pulmonary fibrosis, myofibroblast differentiation is dependent on TGF- β -WNT crosstalk.⁶⁴ Further studies are required to determine how signaling events downstream of the TGF- β -WNT axis differ between dermal development and wound regeneration.

Hypodermal interstitium, a newly recognized skin structure enriched with progenitors

The interstitium, a fluid-filled interstitial space enriched with fibroblasts and ECM proteins found in numerous tissues, has been recently recognized as an important anatomical structure in edema, fibrosis, and mechanical functioning of many organs.^{12,65} Our study defines a multipotent subpopulation of dFBs, the WNT2⁺DPP4⁺ HI-APs, that may not only give rise to dermal adipocytes during development but also function as the origin of regenerating dFBs during wound healing. This HI-AP population is similar to the reported hypodermal fibroblasts,⁶⁶ fascia fibroblasts,⁶⁷ or interstitial APs,¹² which have a unique ECM production profile, multipotent differentiation potential, and ability to migrate to the wound in response to injury. These converging lines of evidence suggest that HI-APs may be key progenitor cells contributing to dermal cell regeneration.

Limitations of the study

A limitation of this study is that we did not use a genetic approach to ablate or lineage trace HI-APs to confirm the function of HI-APs in dermal adipogenesis. We found that WNT2 is a highly specific marker gene for HI-APs; therefore *Wnt2-Cre* mice can be used to genetically ablate or lineage trace these cells to validate their roles in dermal adipogenesis during skin development or wound regeneration.

Altogether, our data defined the cellular and molecular mechanisms through which adipocyte-lineage cells are regulated during skin development and wound regeneration. Future lines of investigation examining the crosstalk between immune cells, APs, and adipocytes under conditions of abnormal wound responses, such as keloid and diabetic wounds, and other skin fibrotic diseases such as scleroderma, could lead to breakthroughs in treating human skin diseases.

STAR * METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for reagents and resources may be directed to and will be fulfilled by the lead contact, Ling-juan Zhang (lingjuan.zhang@xmu.edu.cn).

Materials availability—This study did not generate new unique reagents.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Animals and animal cares—All animal experiments were approved by the Xiamen University and/or the University of California San Diego (UCSD), Institutional Animal Care and Use Committee. C57BL/6 WT mice were purchased from Xiamen University Laboratory Animal Center. *Adipoq-Cre*ERT2 mice (Stock No:025124) and ROSA-*mT/mG* mice (Stock No: 07676) were originally purchased from Jackson laboratory. *Gsk3a^{flox/flox}Gsk3b^{flox/flox}* mice were generously provided by professor Wen-Hsien Liu.⁶⁸ Adipocyte specific *Gsk3* knockout mice were generated by breeding *Gsk3a^{flox/flox}Gsk3b^{flox/flox}* with *Adipoq-Cre*ERT2. *II1r1^{-/-}* mice were generously provided by professor Jiahuai Han (Xiamen university, Xiamen, Fujian, China). All mice were breeded and maintained in standard pathogen free environment of the Laboratory Animal Center in Xiamen University, and all animal experiments were approved by the Animal Care and Use Committee of Xiamen University.

Human skin sample collection and analysis—Fresh adult human full thickness skin biopsies, from age and sex matched healthy, wound and keloid donors (female or male) were collected by the department of dermatology at the First Affiliated Hospital of Fujian Medical University. Keloid disorder was diagnosed based on their clinical appearance, history and treatment history. All keloid samples analyzed in this study were consistent with clinical and pathological criteria for keloid patients. Wound skin biopsies were collected from healthy individual who had no history of keloid formation and receiving wound debridement procedure. All sample acquisitions were approved and regulated by Medical Ethics Committee of the First Affiliated Hospital of Fujian Medical University (reference number No. 2020[146]). The informed consent was obtained from all subjects prior to surgical procedures. Upon collection, these samples were directly fixed with PFA then proceed for paraffin embedding for histological or immunofluorescent analyses.

Skin maturation and aging model—For skin maturation and aging mouse model shown in Figure 3, dorsal skin biopsies were collected from C57BL/6 wildtype adult male mice with various ages (3W, 2 months and 1 year old). Similar to our previous report,²¹ here we used male exclusively to study age-related changes because loss of dermal fat and adipogenic AP in females is relatively slower compared to male counterparts, but the overall trend to lose functional dWAT is similar between female and male.

Mouse wound healing model—7~9 weeks old C57BL/6 mice (male or female) were wounded during telogen phase of hair cycling. Mice were anesthetized using isoflurane, hair

was removed by clipper and Nair, and one full-thickness biopsy ($1.0 \text{ cm} \times 0.5 \text{ cm}$ in size) was generated on each side of dorsal skin (two wounds per mouse). Animal were sacrificed at indicated intervals after injury and wound beds were collected for analysis. To deplete neutrophils, mice were intraperitoneally administrated with 100 ug anti-LY6G (Biolegend, 127649) or IgG2a isotype control antibody (Biolegend, 400565) one day prior to wounding and then were injected daily with 50ug antibodies from day 0 ~ day 2 post wounding, and mice were sacrificed at day 3 post wounding for sample collection and subsequent analysis.

Adipocyte lineage tracing mouse model—*Adipoq-Cre*ERT2;*mTmG* mice (male or female) were daily administrated intraperitoneally with tamoxifen (TAM) (Sigma,T5648) solution (20 mg/mL stock solution dissolved in corn oil (Selleck, s6701)) at 32.5mg/kg body weight for consequent 4~6 days during indicated time period of wound healing. To tracing the cell fate of mature adipocyte *in vitro*, primary dFBs were isolated from *Adipoq-Cre*ERT2;*mTmG* neonatal mice, cultured and differentiated into adipocytes in the presence of tamoxifen (20 μM) to label mature adipocyte by GFP. Measurement of PLIN1⁺ lipid droplet size or ACTA2-expressing in GFP⁺ adipocytes were performed by ImageJ software.

METHOD DETAILS

Single cell RNA library preparation, sequencing and data process—For developmental analyses, dorsal skin biopsies were collected from neonatal (P1), male young adult (3 weeks) and male mature adult (2 month) C57BL/6 mice. To minimize batch effects due to non-biological sources, samples from all age groups were processed, sequenced, and analyzed together. Skin biopsies were minced and digested with collagenase D and DNase1 to isolate single skin cells as previously described (Zhang et al., 2019; Zhang et al., 2021). Dead cells were removed using DeadCell Removal kit (Miltenvi Biotic, 130-090-101) according to manufacturer's instruction. Live cells were counted using a hemocytometer and resuspended in 2% BSA at a concentration of 3,000 cells/uL. All samples were loaded on a 10x Genomics GemCode Single-cell instrument that generates single-cell Gel Bead-In-EMlusion (GEMs). Barcoded, full-length cDNAs were reverse-transcribed from polyadenylated mRNA. Silane magnetic beads were used to remove leftover biochemical reagents and primers from post GEM reaction mixture. Next, libraries were generated and sequenced from the cDNAs with Chromium Next GEM Single Cell 3' Reagent Kits v2. cDNA libraries were sequenced on an Illumina Novaseq6000 platform (Illumina). The raw sequencing data was demultiplexed and aligned to the reference genome mm10-1.2.0 using Cell Ranger v3.0.2 pipiline(10x Genomics). The generated raw gene expression matrix was converted into Seurat objects using the R package Seurat v2.0. To remove doublets and poor-quality cells, we utilized the following procedures to control for cell quality: >200 genes/cell, <5000 genes/cell; >25,000 unique molecular identifiers(U-MIs); and <8% mitochondrial gene expression. Low quality cells and outliers were discarded, and only \sim 24,409 viable cells were used for downstream analysis. These included \sim 7,483 cells – NB; ~8,235 cells – young (3weeks mice); ~8,691cells-adult (2month mice).

For mouse wound samples, skin biopsies were collected from wound day 2 and day 7 granular tissues or unwounded control skin. Single cells were isolated and subjected to single cell RNAseq procedures including libraries construction and data processing as

described above. After low quality cells and outliers were discarded, 10233 cells from non-lesional skin, 7807 cells from wound beds at post-wounding day 2 (PWD2), and 4925 cells from wound tissue at PWD7 were used for downstream analysis.

Unsupervised clustering and gene expression were visualized with the Seurat 2.0 on R studio, and assignment of cell clusters was based on expression of validated marker genes. To investigate the developmental path of dermal fibroblasts, pseudotime analysis was performed on selected dFB subsets by Monocle 2.10.1 (Xiaojie Qiu et al., 2017) to infer the pseudotime trajectories of these cell, followed by scEpath to identify pseudotime-dependent gene expression changes and classify into distinct differentiation states. To further determine the cell fate of dFBs, RNA velocity was performed using the package velocyto (Gioele La Manno et al., 2018) with the default parameter, and the BAM files were used as inputs. To analyze activity of transcriptional regulators in skin development or wound healing, SCENIC (Van de Sande et al., 2020) conjunction with the mm9 Rcis Target database was used and UMIs for transcriptionally defined fibroblasts as input to construct regulon networks between transcription factors and potential target genes and score the active value of regulon for each FB clusters. Single cell RNA libraries construction, sequencing and bioinformatic analysis was assisted by GENE DENOVO Inc (Guangzhou, China).

Cellrank (v1.5.0) analysis was performed to calculate cell-cell transition probabilities according to literature.³⁰ Briefly, scVelo (v0.2.4) was first performed on selected dFB clusters (r0 - r12) to estimate RNA velocities and velocity graphs. Secondly, transition probabilities were calculated in a weighted transition matrix with 80% RNA velocity and 20% similarity. The Generalized Perron Cluster Analysis(GPCCA) was used to compute fate probabilities. The terminal states were estimated from the so-called 'eigengap heuristic' of the spectrum of the transition matrix. Analysis was repeated for the initial states. Probability of each cell to transit to the terminal states was computed. Finally, directed PAGA was used to aggregate the individual fate maps into a cluster-level fate map.

Bulk RNA sequencing and bioinformatic analysis—Total cellular RNA were extracted using TRIzol reagent (Sigma, T9424) and RNAExpress Total RNA Kit (NCM, M050). RNA quality was analyzed by bioanalyzer and RNA samples with RIN value >7were used for sequencing. Next generation sequencing library preparations were constructed according to the manufacturer's protocol (NEBNext Ultra RNA Library Prep Kit for Illumina). The poly(A) mRNA isolation was performed using NEBNext Poly(A) mRNA Magnetic Isolation Module (NEB). The mRNA fragmentation and priming were performed using NEBNext First Strand Synthesis Reaction Buffer and NEBNext Random Primers. First strand cDNA was synthesized using ProtoScript II Reverse Transcriptase and the second-strand cDNA was synthesized using Second Strand Synthesis Enzyme Mix. The purified double-stranded cDNA was then treated with End Prep Enzyme Mix to repair both ends and add a dA-tailing in one reaction, followed by a T-A ligation to add adaptors to both ends. Size selection of Adaptor-ligated DNA was then performed using AxyPrep Mag PCR Clean-up (Axygen), and fragments of ~420bp (with the approximate insert size of 300 bp) were recovered. Each sample was then amplified by PCR. The PCR products were cleaned up using AxyPrep Mag PCR Clean-up (Axygen), validated using an Agilent 2100 Bioanalyzer (Agilent Technologies), and quantified by Qubit 2.0 Fluorometer (Invitrogen).

Then libraries with different indexs were multiplexed and loaded on an Illumina Navoseq instrument for sequencing using a 2×150 paired-end (PE) configuration according to manufacturer's instructions. The sequences were processed and analyzed by GENEWIZ. Venn diagrams between gene sets were made by BioVenn. KEGG and gene ontology (GO) pathway analysis for differentially expressed genes and correlation analysis between single cell RNA-seq with bulk RNA-seq were performed by R package clusterProfile 3.12.0.

Pearson correlation analysis—Pearson correlation analysis was performed between the scRNA-seq datasets and bulk RNAseq datasets to determine the degree of correlation/ similarity between dFB clusters identified by scRNA-seq with skin samples at various ages or different dFB treatment states. The scRNA-seq and bulk RNAseq transcriptomic datasets were normalized and matrixed, and Pearson correlation coefficient of the matrix was calculated, and hierarchical clustering analysis was performed to generate correlation heatmap in R Studio software.

Cell-chat signaling network analysis—R package CellChat 1.3.0 (Jin et al., 2021) was used to evaluate the potential cell-cell communication between dFB subclusters and other cell types, especially immune cells, during wound healing. The scRNA-seq data was imported to implement CellChat platform in R software. The process includes projecting the gene expression data onto protein-protein interaction (PPI) network, then assigning a probability value to infer biological cell-cell communication network, and calculating the centrality indicator of interacting network to identify the role / contribution of each cell population in distinct signaling pathways. The number and strength of identified cell-cell communication is displayed through hierarchical graphs, circle charts, heatmaps, etc. to visualize single or multiple signaling pathways.

Primary mouse dermal fibroblast isolation and in vitro adipocyte

differentiation: Primary mouse dermal fibroblasts were isolated from mouse skin as described previously (Zhang et al., 2019; Zhang et al., 2021). In brief, mouse dorsal skin was first into strips and digested with dispase overnight to remove epidermis. The dermis tissue was then cut into small pieces and digested with 2.5 mg/mL Collagenase D and 30 ng/mL DNAse1 for 2 hous at 37°C to release dermal fibroblasts. Cell mixture was then filtered through 30 µm filter and treated with red blood cell lysis buffer. Isolated dermal fibroblasts were cultured in growth medium (DMEM supplemented with 10% FBS and antibiotics/antimycotics) in a humidified incubator at 5% CO2 and 37°C under sterile conditions. Fresh medium was replenished daily to remove debris or dead cells. Primary cells were then trypsinized within 3 days and replated at 5×104 /mL for *in vitro* assays, and only passage 1 cells were used for experiment. To induce adipocyte differentiation, two day post-confluent dFB were switched to adipocyte differentiation medium containing 2 µM Dexamethasone, 250 µM IBMX, 200 µM Indomethacin and 10 µg/mL recombinant human insulin. Fresh differentiation medium was changed at day 2 then medium was switched to maintenance medium (growth medium supplemented with 10 µg/mL recombinant human insulin) to promote maturation and hypertrophy of differentiated adipocytes.

Histology, collagen trichrome staining and immunohistochemistry (IHC)—To prepare paraffin sections, tissue biopsies were fixed with 4% PFA (Alfa Aesar, Shanghai, China) overnight then dehydrated and embedded in paraffin and sectioned at a thickness of 5-8 µm. For OCT embedding, fresh dorsal skin tissues were directly embedded in OCT then sectioned at a thickness of 15–20 µm, and frozen sections were fixed with 4% PFA for 15min prior to staining. Histology was assessed by Hematoxylin and Eosin (HE) staining using solutions from ZSGB-BIO (Beijing, China), and collagen was stained by the Masson's Trichrome Stain Kit (Solarbio, Beijing, China), and elastic fiber staining was performed by elastic fiber stain kit (Saint-Bio, Shanghai, China) according to manufacturer's instruction. Dermal cell density count shown in Figure S3C was performed manually by counting the number of nuclei per 1mm² dermal area. Lipid staining was performed on frozen sections using BODIPY (ThermoFisher Scientific, Waltham, MA, USA) and PHA (Cytoskeleton, Denver, CO, USA) dyes. For IHC, fixed sections were permeabilized with 0.1% saponin (Sigma-Aldrich, St Louis, MO) and blocked in 5% BSA (Bioforxx, Hessen, Germany), and blocked sections were incubated with primary antibodies at 4°C overnight followed by appropriate 488-, Cy5 or Cy3-coupled secondary antibodies (Jackson ImmunoResearch Laboratories, West Grove, Pennsylvania, USA) in the dark for 4 hours at 4°C. Finally, sections were mounted by ProLong Gold Antifade Mountant with DAPI (Thermo Scientific Inc., Rockford, IL, US). All images were taken with Aperio VERSA Brightfield, Fluorescence Digital Pathology Scanner or Leica TCS SP8 White Light Laser Confocal Microscope, and processed by photoshop and/or Aperio ImageScope software (Leica Biosystems, Nußloch, Germany). Quantification of the fluorescence integrated intensity for each fluorophore was performed using ImageJ software.

Flow cytometry and analysis (FACS) and cell sorting-FACS procedure of primary dermal fibroblasts and immune cells was modified from previously established or reported methods.^{21,69} Briefly, freshly isolated mouse dFBs were first stained with zombie violet viability dye (BioLegend, 423114) to label dead cells. Cells were then blocked with anti-mouse CD16/32 (eBioscience, 14016185), followed by staining with an antibody cocktail for fibroblasts containing PECy7-CD45 (BioLegend, 147704), PerCP-Cy5.5-CD31 (BioLegend, 102522), FITC-CD26(Biolegend, 137806), AF700-CD24(Biolegend, 101836), PE-THY1 (BioLegend, 105308), APC-PDGFRa (eBioscience, 17140181) and BV605-LY6A (BioLegend, 108133), or an antibody cocktail for immune cells containing FITC-Ly6G (eBioscience, 11593182), PE-F4/80 (eBioscience, 12480182), APC-CD11C (BioLegend, 117310), AF700-MHCII (eBioscience, 56532182), PerCP-Cy5.5-Ly6C (BioLegend, 128012), PECy7-CD11B (Biolegend, 101216), APC-Cy7-CD3 (BioLegend, 100222). To stain intracellular ACTA2 in dFBs, stained cells were then fixed and permeabilized using the intracellular fixation and permeabilization buffer set (eBioscience) and stained by AF488-SMA/ACTA2 (eBioscience, 53976082). FACS analysis for protein expression of each cell marker was performed by the Thermo Attune NxT machine and analyzed by FlowJo V10 software. Dead cells stained positive with zombie violet dye were excluded from the analyses. FACS sorting for dFB subpopulations from neonatal mouse skin was performed by the BD Aria Fusion machine. After sorting, freshly isolated cells were directly lyzed in trizol for qRTPCR analysis or cultured in DMEM supplemented with FBS and antibiotics/ antimicotics for in vitro differentiation assays. Adipogenic differentiation

was induced in sorted cells according to the "Primary mouse dermal fibroblast isolation and *in vitro* adipocyte differentiation" method section. Differentiated adipocytes were stained with Oil-red-O (ORO) to detect lipid-droplets as described previously¹. To induce osteogenic differentiation, post-confluent sorted cells were treated with osteoblast differentiation medium containing 50ug/mL ascorbic acid and 10mM β -glycerophosphate, and medium was refreshed every other day for upto 20 days before being stained with Alizarin Red (AR) to visualize osteogenesis.⁷⁰ The percentages of ORO- or AR-positive areas were quantified using ImageJ software.

Quantitative reverse transcription-quantitative PCR (qRT-PCR) analyses—Total cellular RNA was extracted using the RNAExpress Total RNA Kit (NM, M050) and 500 ng of RNA was reverse transcribed to cDNA using HiScript II Q RT SuperMix kit (Vazyme, R222–01). Quantitative, realtime PCR was performed on the Qtower real time system (Analytikjena, Swavesey, Cambridge, UK) using SYBR Green Mix (Bimake, Houston, Texas, USA). All of the primers used with SYBR green were designed to span at least one exon to minimize the possibility of nonspecific amplification from the genomic DNA. The expression of *Tbp* gene (TATA-Box Binding Protein) was used as a housekeeping gene to normalize data for the expression of mouse genes. Specific primer sequences are shown in Table S4.

Cell extract preparation and western blotting analyses—Primary dFBs were lysed in a denaturing lysis buffer containing 20 mM HEPES pH 7.4, 250mM NaCl, 2 mM EDTA, and 1% SDS supplemented with completed proteinase inhibitor cocktail (apexbio, Houston, USA) and phosphatase inhibitor cocktail (Roche, Basel, Switzerland) to maximally preserve protein post-transational modifications as described previously (Zhang et al., 2016). Lysates were boiled for 3 mins and homogenized by sonication using digital sonifier FS- 350T (Sxsonic, Shanghai, China) followed by centrifugation to remove DNA and cell debris. Protein concentrations were measured by BCA protein assay kit (Thermo Scientific Inc., Rockford, IL). For western blotting, 20ug of protein was mixed with same volume of 2x loading buffer with 20% β- mercaptoethanol (Sigma-Aldrich, St Louis, MO) and boiled for 5 mins at 95°C Then samples were separated on a 10–20% Tris-Tricine precast gel (Thermo Scientific Inc., Rockford, IL), transferred to PVDF membrane (Roche, Basel, Switzerland), followed by immunoblotting using indicated primary antibodies followed by fluorescent secondary antibodies (LI-COR, Lincoln, Nebraskal) and imaging using fluorescent Odyssey System (LI-COR). All western blot images shown in the manuscript are representative of 3 independent experiments.

QUANTIFICATION AND STATISTICAL ANALYSIS

Experiments were repeated at least 3 times with similar results and were statistically analyzed by GraphPad Prism 8 software. Quantification analyses of immunostained tissue sections showing the fluorescence integrated intensity of indicated fluorophores were performed by ImageJ (version 1.53). Quantified intensity profiles showing signals from indicated fluorescent channels across skin sections (from top to bottom) is adapted from publications.^{71,72} Quantification analyses of Oil-red-O or alizarin staining (Figure 1) were performed by ImageJ software according to literature.⁷³ To quantify lipid-droplet (LD)

sizes of cultured adipocytes (Figure 6), LDs were first circled based on the unique circular and bumped morphology of LDs shown by wide-field or PLIN1-staining images, then LD sizes were quantified by ImageJ software. For experiments with two groups, statistical significance was determined using Student's unpaired two-tailed t-test. Normality was tested using the Shapiro-Wilk test, and for datasets that were not normally distributed, nonparametric tests were used to determine statistical significance. For experiments with more than two groups, one-way ANOVA multiple comparison test was performed as indicated in the legend. A p value of <0.05 was considered statistically significant (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data and code availability

- The accession numbers for the raw data files of the scRNA-seq analyses reported in this paper are deposited in the GEO database under accession codes: GSE196800 and GSE189241.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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Highlights

- Highly heterogeneous adipocyte lineage cells are defined in developing mouse skin
- WNT and IL-1 are identified as key suppressing or activating signals for dermal adipogenesis
- Wound-induced adipogenesis is triggered by neutrophils through IL-1R signaling
- WNT activation in adipocytes promotes lipolysis and myofibroblast formation



Figure 1. Characterization of heterogeneous dermal adipocyte lineage cells in developing neonatal skin

(A) scRNA-seq analysis of cells isolated from neonatal skin. Pdgfra⁺ dFBs were reclustered into 12 clusters.

(B) Neonatal dFBs were grouped into three cell states along differentiation trajectories.

(C) Violin plots of indicated genes.

(D–G) Neonatal mouse skin sections were immunostained with antibodies against various dFB markers as indicated, and the nuclei were counterstained with DAPI in either blue or white. Scale bars, 50 µm. Abbreviations: p.c., panniculus carnosus muscle layer; p.f.,

perifollicular; DP, dermal papilla; HF, hair follicle; HB, hair bulb. Brackets show specific anatomical skin layers as indicated.

(H) Quantified result of (G) showing the fluorescence intensity (arbitrary unit, A.U.) of PLIN1 (red) and DLK1 (green) across skin section from top to bottom.

(I–L) Perifollicular dFBs (CD24^{hi}LY6A⁻), pAds (DPP4⁻LY6A⁺), and HI-APs (DPP4⁺LY6A⁺) were sorted and then subjected to qRT-PCR analysis (n = 3/group) of indicated genes (I) or *in vitro* adipocyte differentiation for adipocytes (stained with oil red O [ORO]) (J). Quantified ORO+ area for each group is shown in (K) (n = 5/group). Sorted cells were also subjected to osteocyte differentiation (stained with alizarin red [AR]; see Figure S1V), and quantified AR+ area is shown in (L) (n = 5/group). Scale bar, 2 mm. All error bars indicate mean \pm SEM; **p < 0.01, ***p < 0.01, ****p < 0.0001. (M) RNA velocity analysis overlaid on tSNE projection of developing dFBs. (N) A model for the proposed spatial location of various dFB subpopulations in the developing skin. Abbreviations: PAP, papillary dFBs; RET, reticular dFBs; RET-AP, reticular adipocyte progenitors; pAd, preadipocytes; Ad, adipocytes; HI-pAd, hypodermal

interstitial pAd; HI-AP, hypodermal interstitial AP.



Figure 2. Interplay between the WNT- β -catenin and IL-1-immune signaling pathways in regulating dermal adipogenesis

(A) Signaling enrichment analysis of neonatal dFB clusters shown by heatmap. Color scale of red to blue indicates enrichment score.

(B) Violin chart showing the expression of indicated genes.

(C-E) Neonatal dFBs were pretreated with WNT3A and/or IL-1 β and then subjected to adipocyte differentiation, and differentiated cells were stained with ORO (C). The percentages of ORO+ adipocytes was quantified (D) (n = 6/group). Western blotting analysis of CAMP and FABP4 in adipocyte conditioned medium (E).

(F–H) Treated undifferentiated cells were subjected to western blotting analysis using the indicated antibodies (F; quantified results in G, n = 3/group) or qRT-PCR analysis (n = 3/group) of indicated genes (H).

(I) Neonatal dFBs were treated with IL-1 β over a time course of 24 h and cell extracts were subjected to western blotting using indicated antibodies.

(J) Bar graphs showing fold changes of indicated proteins from (I) at 1 h of IL-1 β treatment compared with unstimulated controls (normalized to actin or total proteins, average of n = 3/group).

(K and L) Neonatal dFBs were pretreated with specific inhibitors against the indicated signaling molecules before being stimulated with IL-1 β , and cells were subjected to western blotting analysis (K) or qRT-PCR analysis (L) for the expression of indicated genes (n = 3/group).

(M) Proposed model for the antagonistic interplay between IL-1 β -IL-1R and WNT- β -catenin signaling in regulating adipogenesis.

(N–P) Neonatal skin sections from WT or Il1r1 knockout mice were subjected to immunostaining analysis using indicated antibodies or dyes (N, O) or qRT-PCR analysis of indicated genes (P) (n = 3–4/group). Scale bars, 100 μ m. All error bars indicate mean \pm SEM. *p < 0.05, **p < 0.01, ***p < 0.001.



Figure 3. Tracing dermal adipogenesis and fibrogenesis during skin maturation (A) BODIPY (green dye for lipid) and phalloidin (red dye for actin fiber) staining (top) and

Gomori trichrome staining (bottom) of skin sections from mice at indicated ages. Scale bar, 100 μm

(B) Schematic of scRNA-seq of dorsal skin cells isolated from NB, 3-week, and 2-month mice.

(C) tSNE plots showing cell distribution by clusters indicated with unique color.

(D) tSNE plots showing the expression of indicated marker genes.

(E) tSNE plots showing cell distribution by age of NB (red), 3-week (green), and 2-month (blue) mice.

(F) Bar chart showing age-related changes in the percentage of various cell types.

(G–I) Pdgfra⁺ dFBs were reclustered into tSNE plots by cluster (G) or by age (H). Agerelated changes in the expression of the indicated genes are shown by heatmap (I).

(J) Heatmap showing the mRNA expression (based on bulk RNA-seq) of the listed genes in mouse skin at the indicated ages.

(K) Heatmap showing age-dependent kinetics of the expression of the listed genes in the indicated dFB clusters.

(L and M) Skin sections from mice at 3 weeks or 2 months of age were immunostained with antibodies against the indicated dFB/AD markers. Scale bars, $100 \mu m$.

(N and O) Quantified bar graphs of FACS plots in Figure S3G showing the percentage of DPP4⁻LY6A⁺ pAds (N) or DPP4⁺Ly6A⁺ HI-APs (O) in CD31⁻CD45⁻PDGFRA⁺ dFBs (n = 3/group). All error bars indicate mean \pm SEM; **p < 0.01, ***p < 0.001, ***p < 0.001.

Sun et al.

Page 34



Figure 4. Tracing dermal adipogenesis and fibrogenesis during skin wound healing (A) Full-thickness mouse wounds were collected at indicated wound days (w.d.) for PHA (red) and BODIPY (green) staining. The dotted lines mark wound central granular tissues. Scale bar, 400 μm.

(B) LY6A (red), DPP4 (blue), and ACTA2 (green) immunostaining. Dotted lines mark wound GT. Scale bar, 400 μ m.

(C) Quantified intensity profiles for (B), showing signals from all three fluorescence channels from wound top to bottom (representative of n = 3/group).

(D–F) Wound day 2 and 7 and unwounded control skin samples were subjected to scRNAseq analysis. UMAP (uniform manifold approximation and projection) plot (D) showing cell

distribution by clusters. UMAP plots (E) showing the expression of indicated marker genes for each cluster. Stacked bar graphs (F) showing the percentage of each cell cluster. (G and H) *Pdgfra*⁺ dFBs were reclustered. UMAP plots (G) showing cell cluster distribution in different samples. UMAP plots (H) showing the expression of indicated marker genes. (I and J) RNA velocity analyses (I) and cell rank analyses (J) were mapped onto the UMAP plot of dFB reclusters of w.d. 2 or 7 samples as indicated.

Page 36



Figure 5. Wound-induced adipogenesis is regulated by neutrophils and the IL-1 signaling axis (A–C) FACS plots (A and B) and quantified bar graphs (C) showing the percentage of HI-APs (DPP4⁺LY6A⁺) and myofibroblasts (ACTA2⁺LY6A⁻) in control, w.d. 2, and w.d. 7 skin samples (n = 3/group).

(D) Heatmap showing the top enriched transcriptional factors in w.d. 2 cells compared with control and w.d. 7 cells.

(E) KEGG/GO pathway analysis comparing the r0 HI-AP cluster of control and w.d. 2 skin samples showing top enriched signaling pathways activated in HI-APs after wounding.

(F) Violin plots showing the expression of indicated genes in the control and w.d. 2 samples.

(G) Circle plot showing the inferred intercellular communication network for IL-1 β signaling in control and w.d. 2 skin cells.

(H) UMAP plots showing the expression of indicated genes in total wound cell clusters. (I) Quantified bar graphs (from FACS plots shown in Figure S5G) showing neutrophils $(Ly6G^+CD11B^+)\%$ (n = 3/group).

(J and K) Mice were injected intraperitoneally with isotype (Ctrl) or anti-Ly6G antibodies to deplete neutrophils and then skin wounds were collected at w.d. 3 for granular tissue (GT) area measurement (J). qRT-PCR analysis (K) of the listed genes (n = 4/group).

(L and M) Wound samples were collected from WT or *II1r1* knockout mice at w.d. 3 for immunostaining (L) of pCREB (red) and DPP4 (blue), and quantified percentage of pCREB⁺ cells is shown in (M) (n = 7–10/group). In (L), the white dotted line marks the granule tissue and arrow points to the unaffected pCREB signaling in the migrating epidermis. Scale bar, 400 μ m.

(N) qRT-PCR analysis of the listed genes (n = 4–5/group). All error bars indicate mean \pm SEM; *p < 0.05, **p < 0.01, ***p < 0.001.



Figure 6. Activation of WNT and TGF- β signaling pathways in adipocyte-lineage cells during the wound-proliferative phase

(A) qRT-PCR analysis showing the mRNA expression kinetics of listed genes during skin wound healing (n = 3 per group).

(B) *Adipoq-Cre*ERT2;*mTmG* mice were administered tamoxifen (TAM) from day 1 to 6 during wound healing, and wound tissues were collected at w.d. 10 for immunostaining of ACTA2 (blue) and GFP (green). Scale bar, 400 µm.

(C) Heatmap of SCENIC TF analysis showing the top enriched transcriptional factors.

(D) Wound tissues were stained with active β -catenin, ACTA2, and/or DPP4 as indicated. Scale bar, 500 µm. Quantified intensity profiles showing signals from all three fluorescence channels from wound top to bottom (the lower right box) are shown (representative of n = 3/group).

(E–G) Fully differentiated adipocytes were treated with lithium and/or TGF- β 2 for 2 days. (E) Immunostaining of ACTA2 (red) and DAPI (blue). Fluorescence and wide-field images were overlaid. Scale bar, 50 µm. (F) Quantification of changes in lipid droplet (LD) size (LD quantified from n = 5 fields/group). (G) qRT-PCR analyses of the indicated genes (n = 3/group).

(H–K) Primary neonatal dFBs isolated from *Adipoq-Cre*ERT2;*mTmG* were differentiated into mature adipocytes in the presence of TAM and then treated with lithium and TGF- β 2 (H) before being subjected to immunostaining with the indicated antibodies (I and J) or percentage quantification (K) of ACTA2-expressing myo-FBs in all GFP-positive cells (n = 4/group). Scale bars, 100 µm. All error bars indicate mean ± SEM; *p < 0.05, **p < 0.01, ***p < 0.001.



Figure 7. WNT activation in adipocytes promotes myofibroblast formation during wound healing (A–G) *Adipoq*-C*re*ERT2; *Gsk3*^{flox/flox} mice were administered TAM from w.d. 1 to 6, and w.d. 7 tissues were collected for analysis. (A) Staining of GSK3 (red), PLIN (blue), and DAPI (white), and (B) quantified GSK3 intensity in PLIN1⁺ cells (n = 4/group). (C) Staining of active β -catenin (red) and ATGL (green) and (D) quantified active β -catenin intensity in ATGL⁺ area (n = 4/group). Scale bars, 100 µm. (E–G) FACS plots and quantified bar graphs (n = 3/group) showing the percentage of ACTA2⁺LY6A⁻ myofibroblasts or LY6A⁺DPP4⁻ preadipocytes in PDGFRA⁺ dFBs (n = 3/group).

(H) FABP4 (red), COLIV (blue), and DAPI (white) staining in healthy control (HC), wound (day 7), and keloid skin sections. Scale bar, 400 μm.

(I) Active β -catenin (red), COLIV (blue), and ACTA2 (green) staining in keloid skin sections. Scale bar, 2 mm.

(J) Bar graphs showing relative mRNA expression (based on RNA-seq FPKM [fragments per kilobase per million mapped fragments] values) of listed genes in non-lesional and

wound tissues from HC or keloid (K) individuals (n = 4–5/group). All error bars indicate mean \pm SEM; *p < 0.05, **p < 0.01.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Bodipy	ThermoFisher Scientific	D3922
РНА	Cytoskeleton	PHDH1-A
PE anti-THY1	BioLegend	Cat# 105308; RRID: AB_313179
BV605 anti-LY6A	BioLegend	Cat# 108133; RRID: AB_2562275
AF700-CD24	BioLegend	Cat# 101836; RRID: AB_2566730
FITC anti-mCD26(DPP4)	Biolegend	Cat# 137806; RRID: AB_10663402
APC anti-PDGFRA	eBioscience	Cat# 17140181; RRID: AB_529482
PECy7 anti-CD45	Biolegend	Cat# 147704; RRID: AB_2563536
PerCP-Cy5.5 anti-CD31	Biolegend	Cat# 102522; RRID: AB_2566761
FITC anti-Ly6G	eBioscience	Cat# 11593182; RRID: AB_465314
PECy7 anti-CD11B	Biolegend	Cat# 101216; RRID: AB_312799
AF488 anti-SMA	eBioscience	Cat# 53976082; RRID: AB_2574461
PE anti-F4/80	eBioscience	Cat# 12480182; RRID: AB_465923
PerCP-Cy5.5 anti-Ly6C	BioLegend	Cat# 128012; RRID: 1659241
APC anti-CD11C	BioLegend	Cat# 117310; RRID: AB_313779
AF700 anti-MHCII	eBioscience	Cat# 56532182; RRID: AB_494009
APC-Cy7 anti-CD3	BioLegend	Cat# 100222; RRID: AB_2242784
Goat anti-PLIN	Abcam	Cat# AB61682; RRID: AB_944751
Rat anti-CD24	BioLegend	Cat# 101801; RRID: AB_312834
Goat anti-PDGFRA	R&D system	Cat# AF1062-SP ; RRID: AB_2236897
Rabbit anti-THY1	Cell Signaling Technology	Cat# 13801S ; RRID: AB_2798316
Rabbit anti-COL1A1	abcam	Cat# ab34710; RRID: AB_731684
Rat anti-LY6A	R&D system	Cat# MAB1226; RRID: AB_2243980
Rat anti-Lgr5	R&D system	Cat# mab8240; RRID: AB_2753324
Goat anti-DPP4	R&D system	Cat# AF954-SP ; RRID: AB_355739
Goat anti-DLK1	R&D system	Cat# AF8277; RRID: AB_2861313
Rabbit anti-WNT2	proteintech	Cat# 27214-1-ap ; RRID: AB_2880804
Rabbit anti-TRPS1	ABclonal	Cat# A7743; RRID: AB_2772733
Goat anti-ALPL	R&D system	Cat# AF2910; RRID: AB_664062
Sheep anti-ATGL	R&D system	Cat# AF5365; RRID: AB_2165678
Rabbit anti-CRABP1	Proteintech	Cat# 12588-1-AP ; RRID: AB_2292271
Rabbit-anti-active β-Catenin	Cell Signaling Technology	Cat# 8814; RRID: AB_11127203
Rabbit anti-β-Catenin	Cell Signaling Technology	Cat# 8480; RRID: AB_11127855
Rabbit anti-GSK3β	Cell Signaling Technology	Cat# 12456; RRID: AB_2636978
Rabbit anti-phospho-GSK3β(Ser9)	Cell Signaling Technology	Cat# 5558; RRID: AB_10013750
Rabbit anti-CREB	Cell Signaling Technology	Cat# 9197; RRID: AB_331277
Rabbit anti-Phospho-CREB(Ser133)	Cell Signaling Technology	Cat# 9198; RRID: AB_2561044
Rabbit anti-AKT	Cell Signaling Technology	Cat# 4691S ; RRID: AB_915783
Rabbit anti-Phospho-AKT(Ser473)	Cell Signaling Technology	Cat# 4060S ; RRID: AB_2315049

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Rabbit anti-ERK1/2	Cell Signaling Technology	Cat# 9102S ; RRID: AB_330744
Rabbit anti-Phospho-ERK1/2(Thr202/Tyr204)	Cell Signaling Technology	Cat# 4370S ; RRID: AB_2315112
Rabbit anti-Phospho-NF-rB p65 (Ser536)	Cell Signaling Technology	Cat# 3033; RRID: AB_331284
Rabbit anti-P38 MAPK	Cell Signaling Technology	Cat# 9212S ; RRID: AB_330713
Rabbit anti-Phospho-p38 MAPK (Thr180/Tyr182)	Cell Signaling Technology	Cat# 4631S ; RRID: AB_331765
Mouse anti-β-Actin	Proteintech	Cat# 66009-1-Ig ; RRID: AB_2687938
Goat anti-COLIV	Abcam	Cat# ab769; RRID: AB_92262
Rabbit anti-FABP4	Proteintech	Cat# 15872-1-AP ; RRID: AB_2102440
Rabbit anti-CRAMP	University of California San Diego	Dr. Richard Gallo
Rabbit anti-SMA	Proteintech	Cat# 14395-1-ap ; RRID: AB_2223009
Rabbit anti-NF-ĸB p65	Cell Signaling Technology	Cat# 8242; RRID: AB_10859369
Rabbit anti-Phospho-C/EBPβ (Thr235)	Cell Signaling Technology	Cat# 3084; RRID: AB_2260359
RDye 680RD Donkey anti-Mouse	LICOR	Cat# 926-68072; RRID: AB_10953628
RDye 680RD Donkey anti-Goat IgG (H+L)	LICOR	Cat# 926-68074; RRID: AB_10956736
RDye 680RD Donkey anti-Rabbit IgG (H + L)	LICOR	Cat# 926-68073; RRID: AB_10954442
RDye 800CW Donkey anti-Goat IgG (H + L)	LICOR	Cat# 926-32214; RRID: AB_621846
RDye 800CW Donkey anti-Mouse	LICOR	Cat# 926-32212; RRID: AB_621847
RDye 800CW Donkey anti-Rabbit	LICOR	Cat# 926-32213; RRID: AB_621848
Alexa Fluor 488 AffiniPure Donkey Anti-Goat IgG (H+L)	Jackson ImmunoResearch	Cat# 705-545-147; RRID: AB_2336933
Cy3 AffiniPure Donkey Anti-Goat IgG (H+L)	Jackson ImmunoResearch	Cat# 705-165-147; RRID: AB_2307351
Alexa Fluor 488 AffiniPure Donkey Anti-Rat IgG (H+L)	Jackson ImmunoResearch	Cat# 712-545-150; RRID: AB_2340683
Cy3-AffiniPure Donkey Anti-Rat IgG (H+L)	Jackson ImmunoResearch	Cat# 712-165-150; RRID: AB_2340666
Alexa Fluor 488 AffiniPure Donkey Anti-Rabbit IgG (H+L)	Jackson ImmunoResearch	Cat# 711-545-152; RRID: AB_2313584
Cy3 AffiniPure Donkey Anti-Rabbit IgG (H+L)	Jackson ImmunoResearch	Cat# 711-165-152; RRID: AB_2307443
Alexa Fluor 488 AffiniPure Donkey Anti-Mouse IgG (H+L)	Jackson ImmunoResearch	Cat# 715-545-150; RRID: AB_2340846
Cy3-AffiniPure Donkey Anti-Mouse IgG (H+L)	Jackson ImmunoResearch	Cat# 715-165-150; RRID: AB_2340813
Brilliant Violet 480-conjugated AffiniPure Donkey Anti-Rat gG (H+L)	Jackson ImmunoResearch	Cat# 712-685-150; RRID: AB_2651112
Brilliant Violet 421-conjugated AffiniPure Donkey Anti-Rat gG (H+L)	Jackson ImmunoResearch	Cat# 712-675-150; RRID: AB_2651110
DyLight 405 AffiniPure Donkey Anti-Goat IgG (H+L)	Jackson ImmunoResearch	Cat# 705-476-147; RRID: AB_2632563
Alexa Fluor 647 AffiniPure Donkey Anti-Rabbit IgG (H+L)	Jackson ImmunoResearch	Cat# 711-606-152; RRID: AB_2340625
Alexa Fluor 647 AffiniPure Donkey Anti-Rat IgG (H+L)	Jackson ImmunoResearch	Cat# 712-606-150; RRID: AB_2340695
Alexa Fluor 647 AffiniPure Donkey Anti-Goat IgG (H+L)	Jackson ImmunoResearch	Cat# 705-606-147; RRID: AB_2340438
Biological samples		
mouse skin samples	Xiamen University Laboratory Animal Center	N/A

F	Animal Center	
Chemicals, peptides, and recombinant proteins		
Eosin Staining Solution	ZSGB-BIO	ZLI-9613
Hematoxylin Staining Solution	ZSGB-BIO	ZLI-9610
3-Isobutyl-1-methylxanthine (IBMX)	Sigma	15879
Indomethacin	Sigma	I8280
Dexamethasone	Sigma	D4902

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Insulin	Sigma	91077C
DMEM, high glucose	ThermoFisher Scientific	C11965500BT
Glutamax	ThermoFisher Scientific	35050-061
Collagenase D from Clostridium histolyticum	Roche	11088882001
Antibiotic-Antimycotic	ThermoFisher Scientific	15240062
2x SYBR Green qPCR Master Mix	Bimake	B21202
red blood cell lysis buffer	TianGen	RT122-02
SB431542	Selleckchem	S1067
HyClone Trypsin Protease	Hyclone	SH30042.01
BD Bacto [™] Dehydrated Agar	Fisher Scientific	DF0140-01-0
TRI Reagent/TRIzol	Sigma	T9424-200ML
UltraPure [™] DNase/RNase-Free Distilled Water	ThermoFisher Scientific	10977015
RNAlater [™] Stabilization Solution	ThermoFisher Scientific	AM7021
DnaseI	Solarbio	D8071-100mg
Saponin	Sigma	47036-50G-F
Protease inhibitor	Sigma	11836153001
Wortamannin	Selleck	S2758
U0126-EtOH	Selleck	S1102
SB202190 (FHPI)	Selleck	S1077
BAY 11-7082	Selleck	S2913
XAV-939 (NVP-XAV939)	Selleck	S1180
16% PFA	Alfa Aesar	43368
ProLong [™] Gold Antifade Mountant with DAPI	ThermoFisher Scientific	P36931
Collagenase/Dispase	Sigma	10269638001
Stabilizing Fixative 3X Concentrate	BDIS	338036
Hifair®II 1st Strand cDNA Synthesis SuperMix for qPCR	Vazyme	11123ES60
2x M-PCR OPTI [™] Mix (Dve Plus)	Bimake	B45012
Recombinant mouse TGF82	R&D System	7346-B2-005
recombinant mouse IL18	R&D system	401-ML-025
recombinant mouse WNT3A	R&D system	1324-WN-002
Critical commercial assays		
RNAExpress Total RNA Kit	NCM Biotech	M050
Masson's Trichrome Stain Kit	Solarbio	G1340-7
Dead Cell Removal Kit	Miltenyi Biotic	130-090-101
Oil Red O stain kit	Solarbio	G1262
Mouse Direct PCR Kit	Bimake	B40015
Ailzarin red S (Osteoblast stain)	Sigma	A5533
Intracellular Fixation & Permeabilization Buffer Set	eBioscience	88-8824-00
Deposited data		
- Skin developmental scRNAseq data	This study	GSE189241
soPNA sea data of skin wound tissues	This study	GSE196800

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental models: Cell lines		
Primary mouse dermal fibroblasts	This study	N/A
Experimental models: Organisms/strains		
B6.129(Cg)-Gt(ROSA)26Sot ^{um4(ACTB-tdTomato,-EGFP)Luo/J}	The Jackson Laboratory	JAX: 007676
C57BL/6-Tg(Ad/poq-Cre/ERT2)1Soff/J	The Jackson Laboratory	JAX: 025124
C57BL/6- Gsk3a ^{flox/flox} Gsk3b ^{flox/flox}	From Wen-Hsien Liu	Chenfeng Liu et al.(https://doi.org/ 10.1126/sciadv.abg6262)
C57BL/6- <i>IIIr1^{-/-}</i>	From Jiahuai Han	Generated by Xiamen University Laboratory Animal Center, Transgenic Animal Facility Core
C57BL/6	From Xiamen University Laboratory Animial Center	N/A
Oligonucleotides		
PCR primers, see Table S3	This study	N/A
Software and algorithms		
GraphPad Prism	GraphPad Software, Inc.	N/A
FlowJo V10	FlowJo LLC	N/A
Adobe photoshop	Adobe software	N/A
Aperio ImageScope	Leica Biosystems	N/A
Rstudio	Rstudio	N/A