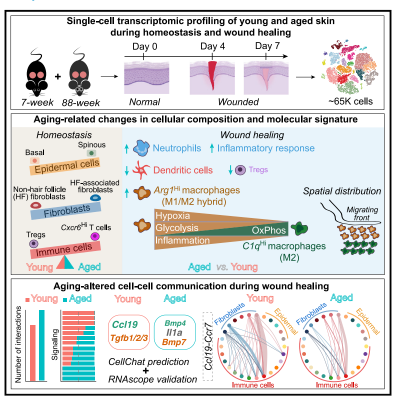
# Wound healing in aged skin exhibits systems-level alterations in cellular composition and cell-cell communication

### **Graphical abstract**



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#### In brief

A comprehensive understanding of cellular and molecular factors that underlie aging-induced decline in wound repair can expose targets for remedy. Vu et al. report single-cell and system-level analysis of young and aged skin during homeostasis and wound healing that reveals aging-related differences in cellular composition, molecular signature, and intercellular communication.

### **Highlights**

- Aging alters skin epithelial, fibroblast, and immune compositions during homeostasis
- Aging alters inflammatory response/macrophage heterogeneity in skin wound healing
- Aging restructures cell-cell communications at a system level during wound healing







#### **Article**

# Wound healing in aged skin exhibits systems-level alterations in cellular composition and cell-cell communication

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#### **SUMMARY**

Delayed and often impaired wound healing in the elderly presents major medical and socioeconomic challenges. A comprehensive understanding of the cellular/molecular changes that shape complex cell-cell communications in aged skin wounds is lacking. Here, we use single-cell RNA sequencing to define the epithelial, fibroblast, immune cell types, and encompassing heterogeneities in young and aged skin during homeostasis and identify major changes in cell compositions, kinetics, and molecular profiles during wound healing. Our comparative study uncovers a more pronounced inflammatory phenotype in aged skin wounds, featuring neutrophil persistence and higher abundance of an inflammatory/glycolytic *Arg1*<sup>Hi</sup> macrophage subset that is more likely to signal to fibroblasts via interleukin (IL)-1 than in young counterparts. We predict systems-level differences in the number, strength, route, and signaling mediators of putative cell-cell communications in young and aged skin wounds. Our study exposes numerous cellular/molecular targets for functional interrogation and provides a hypothesis-generating resource for future wound healing studies.

#### **INTRODUCTION**

Efficient and robust repair of tissue injury is crucial for survival, but this ability, along with tissue function itself, declines with age. Skin, body's essential protective barrier, becomes thinner and drier, and cutaneous wound healing is delayed upon aging (Ding et al., 2021; Ge et al., 2020; Keyes et al., 2016; Wang and Dreesen, 2018). Healing of skin wounds is a concerted effort involving the actions of, and dynamic interactions among, a multitude of cell types in three overlapping phases: (1) inflammatory, characterized by blood clot formation and immune cell infiltration; (2) proliferative, characterized by proliferation and migration of keratinocytes and fibroblasts; and (3) resolution, where local inflammation subdues and extracellular matrix (ECM) is remodeled (Eming et al., 2014; Minutti et al., 2017; Rognoni and Watt, 2018). Wound healing in aged skin associates with alterations in multiple stages, including prolonged inflammation and impaired immune cell function/signaling, increased cellular senescence, as well as delayed re-epithelialization, re-vascularization, granulation tissue formation, and wound closure (Bonham et al., 2020; Ding et al., 2021; Gosain and DiPietro, 2004;

Sgonc and Gruber, 2013; Wang and Dreesen, 2018). However, the cellular and molecular mechanisms that coordinately contribute to such declined healing remain elusive. Particularly, a comprehensive picture of how cellular compositions and cell-cell communications in aged skin wounds differ from that in the young are still lacking. Single-cell RNA sequencing (scRNA-seq) presents a powerful approach to address these critical knowledge gaps.

We have recently developed a computational tool, CellChat (Jin et al., 2021), to analyze and compare intercellular communication networks. CellChat infers cell-cell communications from scRNA-seq data by computing the expression of signaling ligands, receptors, as well as soluble agonists, antagonists, and stimulatory or inhibitory membrane-bound co-receptors. However, the initial version of CellChat has a limited capability in systematically probing the underlying dysregulated intercellular communications that often lead to altered cellular composition and tissue phenotype.

Here we use scRNA-seq to dissect the microenvironmental cell community of unwounded (UW) and wounded skin of young and aged mice. Our analysis reveals altered abundance and



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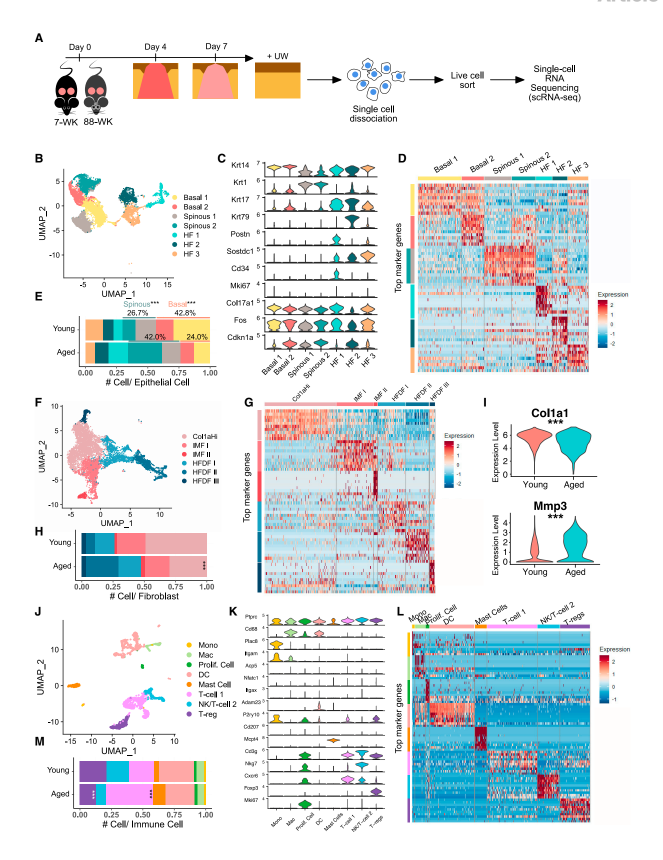
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molecular characteristics of multiple cell type/states as well as an overall more inflammatory profile in aged skin wounds featuring increased abundance of an Arg1Hi macrophage subset. We develop a comparative CellChat framework that enables the detection of signaling communication changes at different levels, including dysregulated cell populations, signaling pathways, and specific ligand-receptor pairs. Using this framework, we systematically identify aging-induced dysregulation in intercellular communications during wound healing, which are characterized by multi-compartmental alterations in growth factor, chemokine, and cytokine pathways.

#### **RESULTS**

#### Young and aged skin exhibit baseline differences across epithelial, fibroblast, and immune cell types during homeostasis

Previously, we sequenced single cells from UW and wounded back skin at day 4 post-wounding (4dpw) of young (6- to 8-week-old) mice (Haensel et al., 2020). To explore aging-associated changes in the heterogeneous wound microenvironment, we performed additional scRNA-seq experiments on young mice at 7dpw and aged (88-week-old) mice at comparable timepoints (UW, 4dpw, 7dpw) (Figure 1A). After quality control, we obtained sequences from 47,729 cells from three UW (14,549 cells), four 4dpw (19,465 cells), and three 7dpw (13,715 cells) samples for young mice, and a total of 17,106 cells from two UW (7,553 cells), two 4dpw (4,112 cells), two 7dpw (5,441 cells) samples for aged mice for downstream analysis (Figures S1A and S1B).

To elucidate aging-associated changes in skin during homeostasis, we first compared young and aged UW samples. Specifically, we aggregated all UW skin cells in a single dataset and then computationally separated epithelial (Krt14+ or Krt1+), fibroblasts (Col1a2<sup>Hi</sup> and Rgs5<sup>-</sup>), and immune cells (Ptprc<sup>+</sup>) for analvsis (Figures S1B-S1D).

The UW epithelial dataset contains 7,286 and 2,252 cells from young and aged skin, respectively, which were clustered into four interfollicular epidermal (IFE) (two basal [Krt14high] and two spinous [Krt1/10high]) and three hair follicle (HF)-associated (Krt17<sup>high</sup> or Krt79<sup>high</sup>) subpopulations (Figures 1B-1D; Table S1). Basal 1 cluster showed enriched expression of Col17a1, suggesting a Col17a1<sup>Hi</sup> stem cell-encompassing identity, whereas basal 2 cluster showed enriched expression of Fos and Cdkn1a (Figure 1C), suggesting an early response/growth arrest identity (Haensel et al., 2020; Liu et al., 2019). Spinous 2 cluster displayed overlapping marker genes with basal 2 cluster (Figure 1D), suggesting a transitional state identity. In aged skin, the relative abundance of basal keratinocytes decreased, whereas that of spinous keratinocytes (specifically spinous 2) increased (Figure 1E).

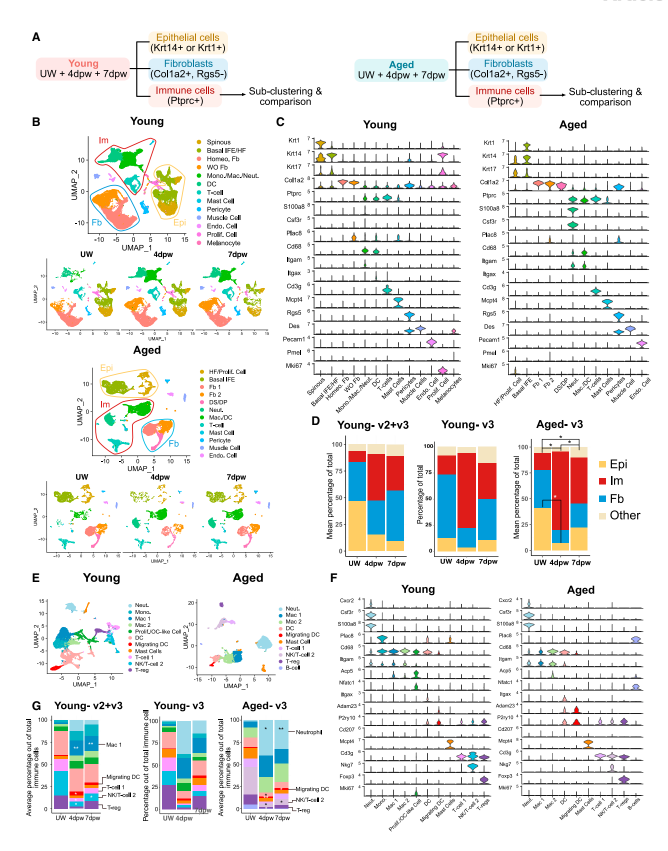
Since our data were collected using both v2 and v3 versions of Chromium Kits (Figure S1B; see STAR Methods), we also computationally isolated and analyzed UW epithelial cells that were analyzed by v3 experiments only to ensure consistency. Epithelial marker genes identified using v2 + v3 data (Figure 1D) were found to also mark the different epithelial cell subsets in v3only data (Figure S2A). Furthermore, a similar trend of reduced basal-suprabasal ratio was observed in aged skin relative to young skin, although the difference was less drastic in v3-only than in v2 + v3 data (Figures 1E and S2B), likely due to differential sensitivity of v2 versus v3 chemistry as well as batch effects. Consistent with previous reports (Ge et al., 2020; Keyes et al., 2016), gene expression comparison between young and aged skin revealed only mild differences, with a small number of genes consistently (in both v2 + v3 and v3-only data) displaying more than 2-fold changes from young to aged skin in either basal or suprabasal keratinocytes (Table S1). For subsequent analyses, we used a similar strategy of comparing v2 + v3 with v3-only data and conclude only on robust differences that are supported by both.

Minor age-related alterations in fibroblasts were observed in UW skin. The UW fibroblast dataset contains 4,826 cells from young and 2,592 cells from aged skin, which were clustered into six subpopulations (Figures 1F and 1G). One cluster showed enriched expression of ECM genes (e.g., Col1a1, Col1a2) and was designed Col1aHi (Figures 1F and 1G; Table S1). Two clusters were enriched for immune-related genes (e.g., complements C3, Cxc/12, and Cxc/13) and designated immune-modulating fibroblast (IMF) I and II (Figures 1F and 1G; Table S1). Two clusters expressed known markers of dermal sheath (DS) or dermal papillae (DP) (e.g., Tagln, Mylk, and Crabp1) (Joost et al., 2020) and another cluster showed overlapping markers with one of these, and they were designed HF-associated dermal fibroblast (HFDF) I, II, and III (Figures 1F, 1G, and S2A; Table S1). Compared with young skin, aged skin showed

Figure 1. Heterogeneity of epidermal cells, fibroblasts, and immune cells in young and aged skin during homeostasis

- (A) Schematic diagram detailing single-cell isolation and collection.
- (B) Uniform manifold approximation and projection (UMAP) of epithelial cells in young and aged UW skin.
- (C) Expression of marker genes used for identification of major cell types in (B).
- (D) Heatmap of top 10 differentially expressed marker genes of clusters in (B).
- (E) Proportion of each epithelial subpopulation out of all epithelial cells in (B).
- (F) UMAP of fibroblasts in young and aged UW skin.
- (G) Heatmap of top 10 differentially expressed marker genes of clusters in (F).
- (H) Proportion of each fibroblast subpopulation out of all fibroblasts in (F).
- (I) Col1a1/Mmp3 expression in Col1a<sup>HI</sup>/IMF I/II fibroblasts from (F). See Table S1 for full list of differentially expressed genes.
- (J) UMAP of immune cells in young and aged UW skin.
- (K) Expression of marker genes used for identification of major cell types in (J).
- (L) Heatmap of top 10 differentially expressed marker genes of clusters in (J).
- (M) Proportion of each immune subpopulation out of all immune cells in (J). p values were calculated using prop.test function in R (E, H, and M) or Wilcoxon rank sum test (I). \*\*\*p < 0.001. Data presented in Figures 1-4 include both v2 and v3 samples unless specified otherwise.





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increased frequency of HF-associated but decreased frequency of non-HF (especially Col1aHi) fibroblasts (Figures 1H and S2B). Furthermore, expression of ECM gene Col1a1 and ECM-regulatory gene Mmp3 was reduced and increased, respectively, in non-HF subsets of aged skin (Figures 1I and S2C; Table S1), consistent with previous reports of aging-associated ECM atrophy (Ding et al., 2021; Freitas-Rodríguez et al., 2017; Lago and Puzzi, 2019; Solé-Boldo et al., 2020; Varani et al., 2006).

Differences in resident immune cells were also detected between young and aged skin. The UW immune cell dataset contains 1,273 cells from young and 1,213 cells from aged skin, and these were clustered into eight subpopulations designated monocytes (Plac8<sup>Hi</sup>) (Blecher-Gonen et al., 2019), macrophages (Cd68<sup>Hi</sup>ltgam<sup>+</sup>Plac8<sup>Low</sup>), dendritic cells (DCs) (Adam23<sup>+</sup>P2ry10<sup>+</sup> Itgax<sup>+</sup>Itgam<sup>-</sup>; may also contain a small number of CD207<sup>+</sup> Langerhans cells) (Tamoutounour et al., 2013), mast cells (Mcpt4+) (Groschwitz et al., 2009), T cells (Cd3g+), natural killer (NK) and NK T cells (Cd3g+Nkg7+) (Malarkannan, 2020), regulatory T cells (Tregs;  $Cd3g^+Foxp3^+$ ), and proliferative cells ( $Mki67^+$ ) based on top markers (Figures 1J-1L and S2A; Table S1). Compared with young skin, aged skin showed a relative expansion of T cell 1 [Cxcr6+; Cxcr6 is required for localization of lungresident memory CD8 T cells (Wein et al., 2019)] and a reduction of Tregs (Figures 1M and S2B).

Taken together, our data revealed baseline differences during homeostasis in young and aged skin across various epithelial, fibroblast, and immune cell populations.

#### Wound healing induces major alterations in epithelial, fibroblast, and immune cell populations in young and aged skin

Next, we examined the major cell types and changes in their proportions during wound healing by aggregating data for all three timepoints (UW, 4dpw, 7dpw) separately for young and aged samples (Figure 2A). In young skin, we observed 13 clusters assigned the following identities based on each cluster's top markers and known marker genes (Haensel et al., 2020; Joost et al., 2016) (Figures 2B and 2C; Table S2): (1) two epithelial, a basal cluster with both IFE (Krt14+) and HF (Krt14+Krt17+) cells, and an epidermal spinous cluster (Krt1+); (2) two fibroblast, one present in both UW and wounded skin and termed homeostatic fibroblasts, and another enriched in wounded skin and termed wound-induced (WO) fibroblasts; (3) four immune, a cluster containing monocytes, macrophages (Cd68Hiltgam+), and neutrophils (S100a8+Csf3r+) (Wilk et al., 2020), a DC (Itgax+) cluster, a T cell cluster, and a mast cell cluster; and (4) five additional cell-type populations, namely a pericyte (Rgs5Hi) cluster (He et al., 2020), a melanocyte (Pmel+) cluster (Solé-Boldo et al., 2020), an endothelial cell (Pecam1+) cluster (Avitabile et al., 2015; Wankell et al., 2001), a muscle cell (Des<sup>Hi</sup>) cluster (Goldfarb and Dalakas, 2011), and a separate cluster of proliferative (Mki67<sup>+</sup>) cells.

Similarly, we observed 12 clusters in aged skin: (1) two epithelial clusters, (2) three fibroblast clusters, (3) four immune cell clusters, and (4) three other cell-type clusters (Figures 2B and 2C; Table S2). Overall, epithelial, fibroblast, and immune cells were the predominant cell types detected in both young and aged skin/wounds (Figures 2B-2D). As expected, the relative abundance of epithelial cells and fibroblasts showed a trend of decrease during wound healing in both young and aged skin, evident especially at 4dpw (Figures 2D, S1C, and S1D). Conversely, the relative abundance of immune cells was increased during wound healing, especially at 4dpw. Fibroblast 1 and 2 clusters in aged skin showed the highest expression of young sample-derived markers of homeostatic and WO fibroblasts, respectively (Figure S1E; Tables S2 and S3). WO fibroblast subset in young skin expanded from UW skin to wounds, but fibroblast 2 subset in aged skin did not (Figure 2B), raising the possibility that aged skin fibroblasts may be compromised in their transcriptome response to wounding.

Immune cells play major roles in wound healing, including clearing pathogens and debris as well as facilitating keratinocyte/fibroblast proliferation and activation, but their apoptosis or exit from skin is also critical for resolution of inflammation and proper tissue repair (Boothby et al., 2020; Eming et al., 2014; Landén et al., 2016; Minutti et al., 2017). We next computationally isolated all-immune-cell clusters from the aggregated UW + 4dpw + 7dpw master dataset (Figure 2A) for finer analysis. Immune cells from young skin were further clustered into 11 subpopulations: neutrophils, monocytes, macrophages, proliferative and osteoclast-like (OC-like) (Acp5+Nfatc1+) (Humbert et al., 2021) cells, DCs (Ccr7<sup>-</sup>) and migrating DCs (Ccr7<sup>+</sup>) (Förster et al., 2008), mast cells, T cell 1, NK/T cell 2, and Tregs (Figures 2E and 2F; Table S2). From UW to wounded skin, percentage of myeloid populations (neutrophils, monocytes, and macrophages) increased, while that of T cell populations (Treg and NK/T cell 2) decreased, especially at 4dpw (Figure 2G). Immune cells from aged skin formed 10 subpopulations that contained all the cell types observed in young skin plus a small number of B cells (Figures 2E-2G). Wound-induced expansion in neutrophil/macrophage subpopulations at the cost of Treg/NK/ T cell 2 subpopulations was also observed in aged skin (Figure 2G).

Taken together, our data provide an overview of the major tissue microenvironmental changes during wound healing and reveal a significant expansion of immune cells, but with

#### Figure 2. Cellular changes during wound healing in young or aged skin

- (A) Data analysis strategy. Young and aged samples are analyzed separately here.
- (B) UMAPs of all cells in UW and wounded (4dpw and 7dpw) skin in young or aged samples.
- (C) Expression of marker genes used for identification of cell populations in (B).
- (D) Proportion of each major cell type out of all cells in (B) at the indicated timepoints. Epi, epithelial; Im, immune; Fb, fibroblast.
- (E) UMAP of immune cells in young or aged samples.
- (F) Expression of marker genes used for identification of cell populations in (E).
- (G) Proportion of each immune population out of all immune cells in (E). p values were calculated using ANOVA and post hoc Tukey tests in R. \*\*p < 0.01, \*p < 0.05 compared with UW samples.



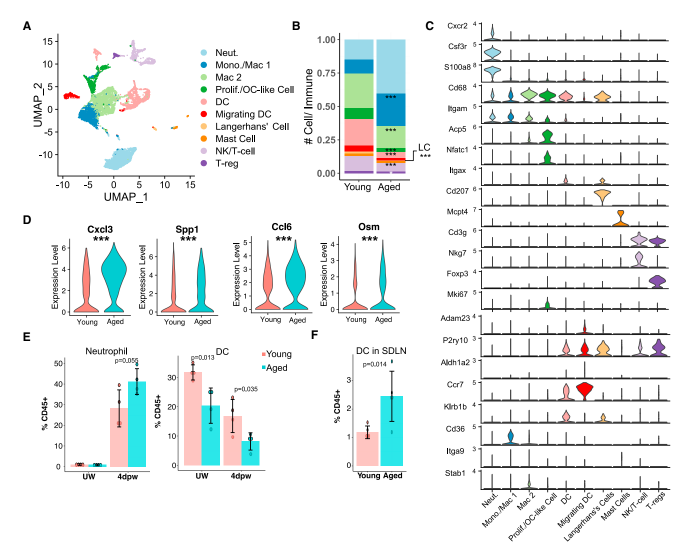


Figure 3. Differential immune cell heterogeneities in young versus aged skin wounds

- (A) UMAP of immune cells in young + aged 4dpw wounds.
- (B) Proportion of each immune subpopulation out of all immune cells in (A).
- (C) Expression of marker genes used for identification of cell populations in (A).
- (D) Expression of select genes in neutrophils from (A). See Table S4 for full list of differentially expressed genes.
- (E) Flow cytometry of neutrophils and DCs in UW and 4dpw skin. n = 4 each for young and aged samples.
- (F) Flow cytometry of DCs in LNs at 4dpw. n = 5 each for young and aged samples. Bar graphs (E and F) represent mean ± SD. p values were calculated using prop.test function in R (B), Wilcoxon rank sum test (D), or unpaired two-tailed Student's t test (E and F). \*\*\*p < 0.001.

seemingly differential dynamics and cell-type compositions in young versus aged skin wounds.

#### Aged skin wounds exhibit a more inflammatory profile than young counterparts, featuring neutrophil persistence, fewer DCs, and increased abundance of an Arg1<sup>Hi</sup> macrophage subset

To facilitate direct comparison of the immune microenvironment, we next aggregated data for young and aged samples at each post-wounding time point, and then computationally isolated immune cells for downstream subclustering analysis (Figures S1B, S1C, and S1E). At 4dpw, we identified 10 clusters, with neutro-

phils, macrophages, and DCs being the majority (87.2% total in young and 90.8% in aged) (Figures 3A-3C; Table S4). Neutrophil abundance showed a trend of increase per v2 + v3 but only mildly per v3-only data, whereas DC abundance was significantly decreased per both analyses, in aged skin wounds compared with young counterparts (Figures 3B and S3A). By 7dpw, neutrophil abundance was significantly higher in age skin wounds than in the young (Figures S3C-S3E; Table S4). Moreover, neutrophils in aged 4pdw and/or 7dpw wounds showed increased expression of inflammatory genes including Cxcl3, Ccl6, Spp1, Osm, and Il1a compared with young counterparts (Figures 3D, S3B, and S3F; Table S4). Reduction in DCs

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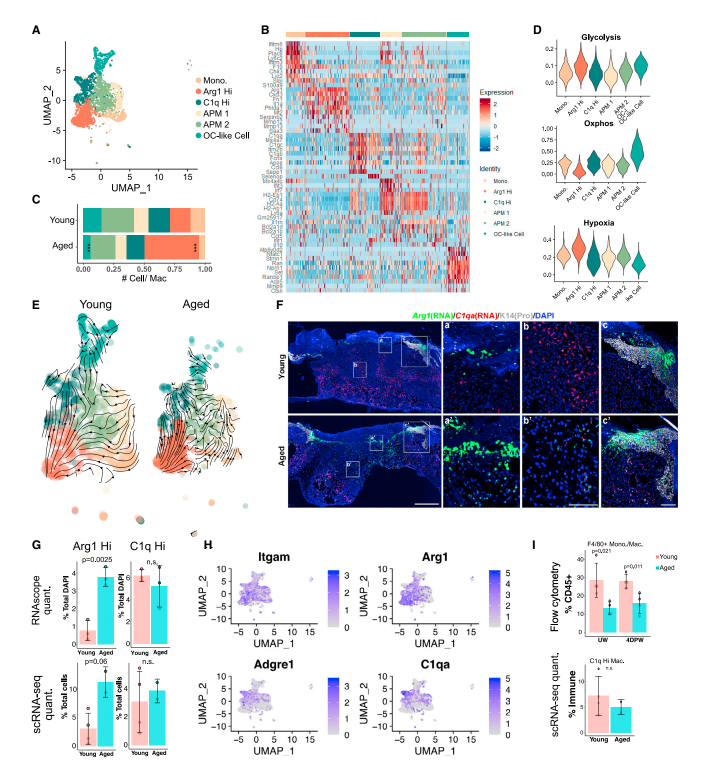


Figure 4. Differential macrophage heterogeneities in young versus aged skin wounds

- (A) UMAP of macrophage subpopulations in young + aged 4dpw wounds.
- (B) Heatmap of top 10 differentially expressed marker genes of clusters in (A).
- (C) Proportion of each macrophage subpopulation out of all monocyte/macrophages in (A).
- (D) Gene scoring analysis of the macrophage subpopulations in (A).
- (E) RNA velocity analysis of macrophages in (A).



also persisted in aged 7dpw wounds (Figures S3C-S3E; Table S4). Additionally, although populational size was small, Tregs appeared to be fewer in aged skin wounds at 7dpw than in the young (Figures S3C-S3E).

To validate the scRNA-seq-revealed differences, we performed flow cytometry on both UW and wounded (4dpw) skin. Neutrophils (CD45<sup>+</sup>CD11b<sup>+</sup>Ly6G<sup>+</sup>) were not detectable in young or aged skin during homeostasis, whereas reduced presence of DCs (CD45+CD11c+MHCII+) in aged skin was already evident (Figure 3E). The results also confirmed a trending increase in neutrophils and a significant reduction in total DCs in aged skin wounds compared with the young (Figure 3E). To ask whether low DC abundance in aged skin might be due to their migration, we profiled DCs in skin-draining lymph nodes (LNs) of wounded mice using flow cytometry. Indeed, LNs of aged mice contained significantly more DCs than their young counterparts (Figure 3F). These findings suggest that DCs in aged mouse skin may be more likely to migrate to LNs, a feature reminiscent of DCs from aged human subjects that exhibit enhanced basal-level inflammatory responses (Agrawal et al., 2009; Panda et al., 2010).

Macrophages are critical orchestrators of skin wound healing through timely promotion and resolution of inflammation (Krzyszczyk et al., 2018; Minutti et al., 2017). We therefore zoomed in on their changes during wound healing in young versus aged skin by computationally isolating monocyte/ macrophage clusters from all-immune-cell datasets at each wound time point for further subclustering (Figures S1B, 4A, and S4C). At 4dpw, we identified six monocytes/macrophage subpopulations in young plus aged skin wounds, and they were monocytes, Arg1Hi macrophages, C1gHi macrophages, OC-like, antigen-presenting macrophage 1 (APM 1; showing the highest expression of major histocompatibility complex (MHC) class II genes such as H2-Ab1 and H2-Eb1), and APM 2 (expressing both MHC class II markers and anti-inflammatory cytokines II10 and II1rn) (Figures 4A-4C, and S4A; Table S5). Interestingly, the abundance of Ara1<sup>Hi</sup> macrophages relative to other monocyte/macrophage subsets was significantly higher in aged skin wounds compared with the young (Figures 4C and S4B). At 7dpw, although only five monocyte/ macrophage clusters were identified, four of them shared similar top markers with clusters observed at 4dpw (Arg1Hi/ APM, C1qHi, OC-like, and monocytes), and one new cluster (enriched for Cd9 expression) emerged (Figures S4C and S4D; Table S4). The relative abundance of Arg1<sup>Hi</sup>/APM macrophages appeared slightly higher in aged 7dpw wounds than in young counterparts (Figure S4E).

Next, we studied the molecular characteristics of the monocyte/macrophage subpopulations identified at 4dpw. C1qHi cluster showed enriched expression of known M2 macrophage marker Mrc1 (Martinez and Gordon, 2014; Orecchioni et al., 2019), and such cells were also present in UW skin (Tables S1, S2, S3, S4, and S5), suggesting a skin resident macrophage identity. Top markers of Arg1Hi cluster included pro-inflammatory chemokine/cytokines (Cxc/1/2/3, I/1a) and known M2 macrophage marker Arg1 (Figure 4C; Table S5), suggesting an inflammatory M1/M2 hybrid identity (Lawrence and Natoli, 2011). Arg1 Hi cluster was among the highest-expressing cells for glycolysis-associated genes and scored lowest for oxidative phosphorylation (OxPhos)-associated genes (Figures 4D and S4F; Table S3). In contrast, C1qHi cluster scored low for glycolysis genes and high for OxPhos genes (Figures 4D and S4F; Table S3). Such gene scoring analysis also revealed higher and lower expression of a hypoxia-associated signature in Arg1Hi and C1qHi/OC-like clusters, respectively (Figures 4D and S4F). Similarly, at 7dpw, Ara1Hi/APM cluster showed higher hypoxiaand glycolysis-associated, but lower OxPhos-associated, gene expression than C1qHi cluster (Figure S4F; Table S3). These data highlight a metabolic preference of Arg1Hi cells for glycolysis over OxPhos that appears to be coupled to hypoxia response.

To explore potential lineage relationships among monocyte/ macrophage subpopulations, we down-sampled their total cell number in young skin wounds to match that in their aged counterparts and performed RNA velocity analysis (Bergen et al., 2020; La Manno et al., 2018). In young 4dpw wounds, C1gaHi, APM 1 and 2, and monocyte clusters all showed a propensity to become Arg1Hi macrophages (Figure 4E; Table S5). In aged 4dpw wounds, however, only C1qaHi cluster showed this propensity, whereas monocyte and APM 2 clusters showed a propensity to become APM 1 (Figure 4E; Table S5). The situation appeared to be reversed at 7dpw, with many young cells transitioning away from the Arg1Hi state, whereas aged cells exhibited a more uniform trajectory toward Arg1<sup>Hi</sup> macrophages (Figure S4G; Table S5). These data suggest differential macrophage polarization kinetics in young versus aged skin wounds, and that multiple monocyte/macrophage subsets may have a potential to become Arg1Hi cells at some point during wound healing.

We next performed RNAScope experiments to examine the spatial expression of Arg1 and C1ga in young and aged skin wounds. Epithelial cells in wound hyperproliferative zone/leading edge were found to strongly express Arg1 (Figure 4F), consistent with a previous report (Kämpfer et al., 2003). Within the stroma, Arg1 expression was detected predominantly at the scabwound bed interface and in upper wound bed, whereas C1ga expression was detected predominantly in lower wound bed and near the wound margin (Figures 4F and S4H). Supporting the scRNA-seq finding, there were more Arg1-expressing

<sup>(</sup>F) RNAScope showing spatial distribution of Arg1 and C1ga transcripts in young and aged 4dpw wounds (n = 3 each). DAPI stains the nuclei. Scale bars: 500 μm in low-magnification image (left), 100 µm in high-magnification images (middle, right).

<sup>(</sup>G) Quantification of percentage Arg1+ or C1qa+ cells per total DAPI-positive cells from RNAScope results in (F) (top) or percentage Arg1+ or C1q+ cells per total live cells from scRNA-seg experiments (bottom).

<sup>(</sup>H) Feature plots showing expression of the indicated genes in macrophage subpopulations from (A).

<sup>(</sup>I) Flow cytometry of F4/80<sup>+</sup> macrophages in UW and 4dpw skin (top). n = 4 each for young and aged samples. Bottom, percentage C1q<sup>+</sup> macrophages per total immune cells from scRNA-seq data. Bar graphs (G and I) represent mean ± SD. p values were calculated using prop.test function in R (C) or unpaired two-tailed Student's t test (G and I). \*\*\*p < 0.001.

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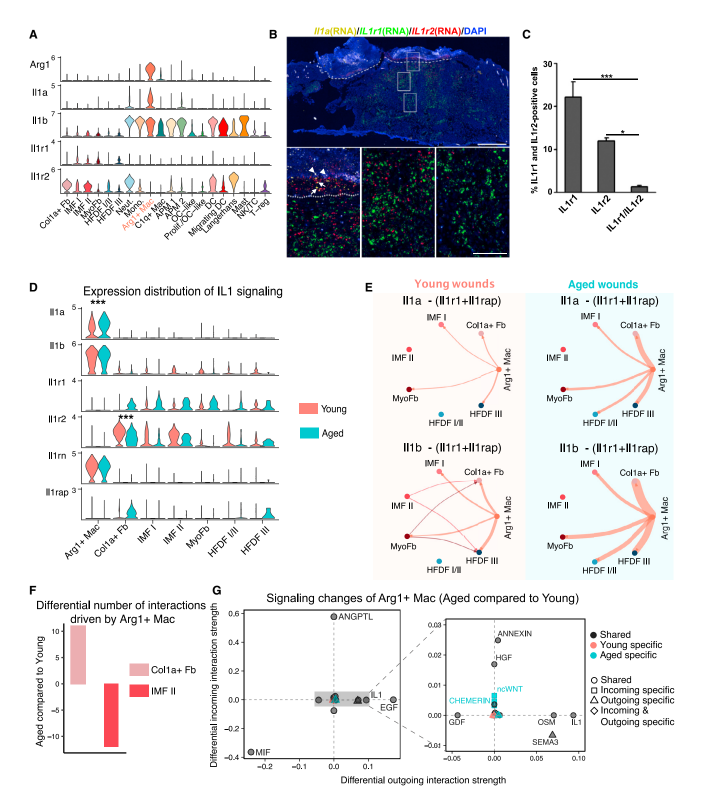


Figure 5. Differential Arg 1<sup>Hi</sup> macrophage-associated signaling in young versus aged 4dpw wounds

(A) Violin plots showing expression levels of the indicated genes in fibroblast and immune cell populations.

(B) RNAScope showing spatial distribution of II1a, II1r1, and II1r2 transcripts in young 4dpw wound. Arrows and arrowheads indicate II1a and background signals, respectively. Dashed line indicates scab-wound bed border. DAPI stains the nuclei. Scale bars: 500 μm in low-magnification image (top), 100 μm in high-magnification images (bottom).

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stromal cells in aged skin wound bed than in the young (Figures 4F, 4G, and S4H). These data suggest differential spatial locations of Arg1Hi and C1qHi macrophages within the 4dpw wound, and that Arg1Hi macrophages are more abundant in aged skin wounds.

F4/80 protein is a commonly used surface marker for monocyte/macrophages. Since Adgre1 (F4/80) mRNA expression fell within the C1gHi rather than Arg1Hi subset (Figures 4H and S4I), we wondered whether F4/80+ population was altered in aged skin wounds. Interestingly, flow cytometry detected lower relative abundance of CD45<sup>+</sup>CD11b<sup>+</sup>Ly6G<sup>-</sup>F4/80<sup>+</sup> monocyte/ macrophages in both UW skin and 4dpw wounds of aged mice compared with young mice, while quantification of scRNA-seq data did not show a significant difference in relative abundance of C1qHi macrophages between young and aged 4dpw wounds (Figure 4I). Additionally, immunostaining revealed a consistent reduction of F4/80+ cells in the dermis of aged 4dpw wounds compared with the young (Figure S4J). These results present an interesting discrepancy between mRNA (Adgre1) and protein (F4/80) expression, which was also noted previously (Haensel et al., 2020).

Taken together, our data reveal salient differences in immune cell compositions between young and aged skin wounds, featuring neutrophil persistence, DC mobilization, enrichment of Arg1<sup>Hi</sup> inflammatory macrophages at the cost of other monocyte/macrophages, and fewer immunosuppressive Tregs in the latter. As such, wound healing in aged skin is characterized by a stronger inflammatory immune microenvironment than in young skin.

#### Altered signaling interaction between Arg1<sup>Hi</sup> macrophages and fibroblasts in aged skin wounds

To gain insight into a signaling role of Arg1<sup>Hi</sup> macrophages, we examined their top marker genes and found II1a (Table S5), which encodes a pro-inflammatory cytokine that is known to inhibit wound healing when present at excess (MacLeod and Mansbridge, 2016; Perrault et al., 2018). Arg1<sup>Hi</sup> macrophages are the predominant source of Il1a expression among all fibroblasts and immune cells in young skin wounds (Figure 5A). In contrast, II1b is more broadly expressed in multiple immune cell types (Figure 5A). Fibroblasts are the major cell type expressing interleukin (IL)-1 receptor II1r1, whereas expression of decoy receptor II1r2 is more widespread but the highest in Col1a<sup>+</sup> subset of all fibroblasts (Figures 5A, S2G, and S2H). Using RNAScope, we mapped the spatial locations of Il1a, Il1r1,

and II1r2 expression in young skin wounds. II1a expression was weakly detected at the scab-wound bed interface, whereas II1r1 expression was detected throughout the wound bed (Figure 5B). Il1r2 expression was detected prominently at the scab-wound bed interface and in the upper wound bed, areas where non-epithelial Arg1 expression was concentrated (Figures 5B, 4F, and S4H). Within the wound bed, II1r1 and II1r2 transcripts were detected in 22% and 12%, respectively, of the cells, but only <2% cells expressed both (Figure 5C). These data reveal interesting spatial patterning of IL-1 signaling ligands and receptors in the wound and implicate the existence of a negative regulatory mechanism (by decoy receptor) to restrict IL-1 signaling within the Arg1<sup>Hi</sup> macrophage-rich zone.

We next compared the expression of IL-1 signaling components in Arg1Hi macrophages and fibroblasts of young and aged skin wounds. Compared with young counterparts, aged skin wounds exhibited decreased expression of Il1r2 in multiple fibroblast subsets at both 4dpw and 7dpw (Figures 5D, S2M, S2N, S5A, S5E, and S5H). RNAScope validated the reduced number of Il1r2-expressing cells within the wound stroma of aged skin compared with the young (Figure S4H). Il 1a expression in Arg1<sup>Hi</sup> macrophages was consistently increased in aged skin wounds at 4dpw and 7dpw, whereas II1b increase was more variable (Figures 5D, S5A, S5E, and S5H). Expression of IL-1R1 antagonist II1rn in Arg1Hi macrophages was largely unchanged at 4dpw, whereas that of Il1rap, encoding an accessory protein essential for IL-1 signaling (Cullinan et al., 1998), was elevated in some fibroblast subsets of aged skin wounds at both 4dpw and 7dpw (Figures 5D, S5A, S5E, and S5H).

We then used CellChat to infer IL-1 signaling-mediated cellcell communication between Arg1Hi macrophages and fibroblasts. We calculated communication probabilities between different cell populations by not only considering both II1r1 and Il1rap at the signal-receiving end but also incorporating the inhibitory effect of II1r2. At both 4dpw and 7dpw, total IL-1 signaling from Ara1<sup>Hi</sup> macrophages onto several fibroblast subsets (e.g., Col1aHi and myofibroblast) was consistently enhanced in aged skin wounds relative to the young (Figures 5E, S5B, S5F, and S5I). In addition to the spatial proximity between II1r2-expressing cells and Arg1-expressing cells in young and aged skin wounds (Figure S4H), RNAScope experiments also showed a subset of Arg1-expressing cells to be adjacent to some Col1a1-expressing cells, particularly in aged skin wounds (Figure S5K). These findings support the possibility that Arg1Hi macrophages and Col1a<sup>Hi</sup> fibroblasts can communicate with each other. Our data

<sup>(</sup>C) Quantitation of ll1r1-, ll1r2-, and ll1r1/ll1r2-positive cells in wound beds (B). n = 3.

<sup>(</sup>D) Comparison of expression levels of IL-1 signaling components in Arg1<sup>HI</sup> macrophages and fibroblasts in young versus aged skin wounds.

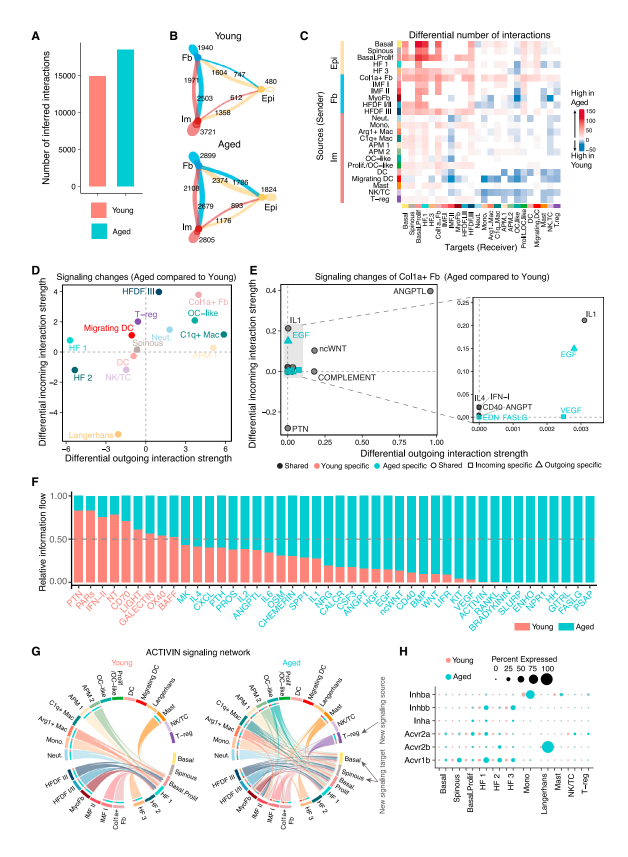
<sup>(</sup>E) IL-1-mediated cell-cell communication networks between Arg7<sup>Hi</sup> macrophages and fibroblast populations in young versus aged skin wounds. Edge width is proportional to the inferred interaction strength. Edge color is consistent with the signaling source.

<sup>(</sup>F) Differential number of putative interactions driving Arg1<sup>HI</sup> macrophage-to-fibroblast signaling in aged versus young skin wounds.

<sup>(</sup>G) Altered Arg1<sup>HI</sup> macrophage-associated signaling pathways in young versus aged skin wounds. The x and y axes represent differential outgoing and incoming interaction strengths, respectively. Positive and negative values indicate increased and decreased signaling, respectively, in aged skin wounds compared with the young. Black, red, and green colors indicate whether a signaling pathway is shared between young and aged, or specific to either young or aged. Dot shapes indicate whether a signaling pathway is specific to either young or aged in its outgoing, incoming, or both outgoing and incoming signaling. Enlarged plot on the right shows the signaling pathways with relatively small alterations. Data presented in Figures 5-7 infer cell signaling from v2 + v3 samples unless specified otherwise. Cell-cell communication information that is qualitatively consistent between v2 + v3 and v3-only data analyses is shown in the main figures, whereas v3-only information is shown in supplemental figures. p values were calculated using unpaired two-tailed Student's t test (C) or Wilcoxon rank sum test (D). \*\*\*p < 0.001, \*p < 0.05.







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demonstrate that IL-1 signaling is elevated during wound healing in aged skin relative to young skin in a manner that cannot be simply explained by delayed healing.

To identify other putative molecular pathways that drive Arg1<sup>Hi</sup> macrophage-fibroblast signaling, we performed unbiased CellChat analysis using the CellChatDB database (Jin et al., 2021) of all known secreted ligands and associated signaling. We found that, at 4dpw, the number of signaling interactions of Arg1<sup>Hi</sup> macrophages-Col1a<sup>Hi</sup> fibroblasts was prominently increased while that of Arg1Hi macrophages-IMF II was decreased in aged skin wounds compared with young counterparts (Figures 5F and S5C). To identify the specific signaling pathways altered in aged skin wounds, we compared the strength of outgoing and incoming interactions from and to Arg1<sup>Hi</sup> macrophages and then extracted information common to v2 + v3 and v3-only analyses (see STAR Methods). Some signaling pathways were present in both young and aged skin wounds ("shared"); of these, oncostatin M (OSM) and epidermal growth factor (EGF) showed enhanced outgoing (Arg1Hi macrophage-originated) signaling, whereas angiopoietin-like protein (ANGPTL) and (HGF) showed enhanced incoming (Arg1Hi macrophage-destined) signaling, in aged samples (Figures 5G and S5D). Macrophage migration inhibitory factor (MIF) showed remarkable decrease in both outgoing and incoming signaling in aged skin wounds (Figures 5G and S5D). Some of these signaling pathways (e.g., OSM, MIF) have previously been shown to regulate fibroblast functions (Ashcroft et al., 1997; Dewor et al., 2007; Ihn and Tamaki, 2000), testifying the overall validity of our findings. Age-dependent alterations were also observed at 7dpw for Arg1Hi macrophage/APM signaling interactions with fibroblasts (Figures S5G and S5J).

Collectively, our results reveal Arg1Hi macrophage-fibroblast communication as a target of altered cell-cell signaling during delayed wound healing in aged skin, presenting changes in multiple signaling pathways and featuring elevated pro-inflammatory IL-1 signaling due to altered expression of multiple signaling mediators and inhibitors.

#### Systems-level analysis reveals differences in cell-cell communication network structure and signaling strength across multiple cell populations in young versus aged skin wounds

To probe global differences in cell-cell communications between young and aged skin wounds, we employed comparative CellChat (see STAR Methods) to study signaling interactions among all major cell types at 4dpw and 7dpw. We combined the subclustering results of epithelial, fibroblast, and immune cells (Figures S2D, S2G, S2J, S2M, 3A, and S4C; Tables S5 and S6), leading to a total of 26 cell groups at 4dpw and 22 cell groups at 7dpw for analysis.

We first examined the total number of possible cell-cell communications and the prominently affected cell types. There was a higher number of possible interactions in aged skin wounds than in young skin wounds at both 4dpw and 7dpw (Figure 6A, S6A, S6H, and S6K). Putative signaling within and between the epithelial and fibroblast populations drastically increased, while signaling between the fibroblast and immune cell populations slightly increased, in aged skin wounds (Figures 6B, 6C, S6B, S6C, S6I, and S6L). Moreover, signaling from immune cells to epithelial cells was strengthened in aged skin wounds. These data show that, in general, aging increases both autocrine and paracrine signaling probabilities among all three major cell types during wound healing.

We next studied how signaling patterns differ between young and aged skin wounds using network centrality analysis, which computes the outgoing and incoming interaction strength of each cell subpopulation to assess their likelihood as signaling sources and targets, respectively. Overall, fibroblasts (Col1a<sup>Hi</sup> fibroblasts in particular) and immune cells (e.g., neutrophils, C1qHi macrophages) drastically increased their outgoing and/or incoming signaling in aged skin wounds (Figures 6D and S6D). Examination of age-associated changes in signaling between Col1a<sup>Hi</sup> fibroblasts and all other cell types revealed a prominent increase in incoming and outgoing signals for ANGPTL, as well as in incoming signals for IL-1, in aged skin wounds (Figure 6E).

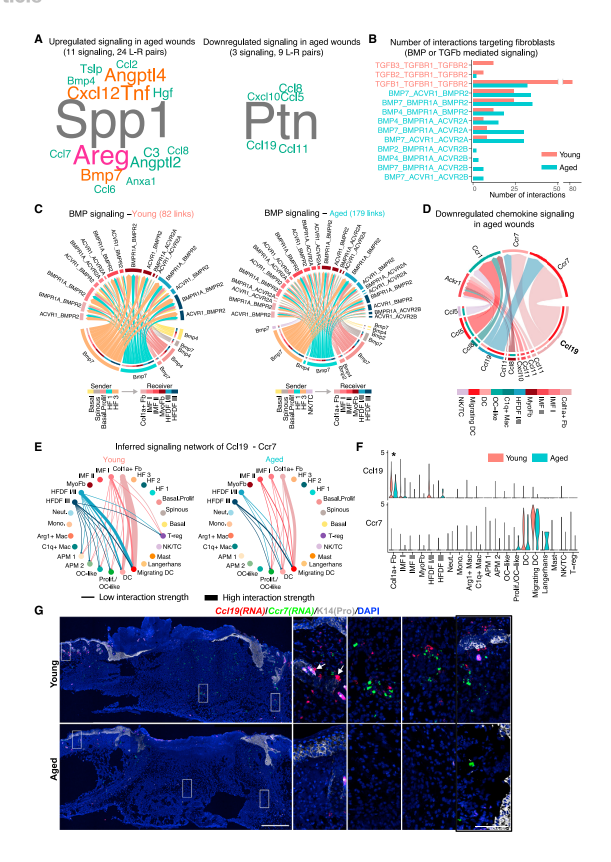
We next compared the information flow, defined by the sum of communication probability among all pairs of cell groups in the inferred network, for specific signaling pathways in young versus aged skin wounds. Certain pathways, such as Fas ligand (FASLG), ACTIVIN, IL-1, OSM, KIT proto-oncogene/receptor tyrosine kinase, and receptor activator of nuclear factor kappa-B ligandr (RANKL), prominently increased their information flow in aged skin wounds (Figures 6F and S6E). The key roles of ACTIVIN signaling in wound healing (Antsiferova and Werner, 2012; Wankell et al., 2001; Wietecha et al., 2020) prompted a closer examination of its overall network architecture. Compared with young skin wounds where proliferative basal and HF 1 cells were ACTIVIN targets, additional epithelial cell subsets in aged skin wounds gained ACTIVIN responsiveness (Figures 6G and S6F). Moreover, Tregs emerged as a new signaling source in aged skin. Interestingly, the expression levels of multiple ACTIVIN ligands and receptors were increased in multiple cell

Figure 6. Alterations in network structure and signaling strength of putative cell-cell communications in young versus aged 4dpw wounds (A) Total number of possible interactions.

- (B) Number of possible interactions between the three major cell types.
- (C) Differential number of possible interactions between any two cell populations. Red (positive values) and blue (negative values) in the color bar indicate higher number of predicted interactions in aged and young skin wounds, respectively.
- (D) Differential interaction analysis identifying prominently altered signaling sources and targets.
- (E) Altered signaling pathways that associate with Col1aHI fibroblasts. See Figure 5G legends for more details.
- (F) Significant signaling pathways were ranked based on their differences of overall information flow, calculated by summarizing all communication probabilities in a given inferred network. Those colored red and green are more enriched in young and aged skin wounds, respectively.
- (G) Chord diagrams of inferred ACTIVIN signaling networks. Edge color is consistent with the signaling source. Segments with large arrows represent signaling targets and inner bars represent signaling sources in which the colors indicate signaling targets.
- (H) Comparison of expression levels of ACTIVIN signaling components in epithelial and immune cells in young versus aged skin wounds.

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subpopulations in aged skin wounds compared with the young at both 4dpw and 7dpw (Figures 6H, S6G, S6J, and S6M), raising the possibility that inefficient healing in aged skin might be due to misdirected ACTIVIN signaling routes and/or deficiency in celltype-specific downstream responses.

Our findings show that both the network architecture and signaling strength of putative cell-cell communications are drastically changed in aged skin wounds compared with their young counterparts. Particularly, the potential of both self- and reciprocal interactions among the major cellular compartments is markedly increased in aged skin wounds.

#### Comparative analysis of molecular signaling identifies dysregulated growth factor, chemokine, and cytokine pathways during wound healing in aged skin

Having obtained an understanding of the general rules governing cell-cell communications in young versus aged skin wounds, we next investigated the molecular signaling mechanisms in detail to identify candidate pathways potentially underlying the altered cellular compositions and communications in aged skin. We identified dysregulated ligand-receptor pairs in aged skin wounds by combining differential expression analysis with cellcell communication analysis. CellChat predicted significant upregulation of 24 ligand-receptor pairs from 11 signaling pathways in aged skin wounds at 4dpw compared with young counterparts. These include ligands Spp1, Areg, Bmp7, Bmp4, Tnf, and Cxcl12 (Figure 7A). Spp1 knockdown has been shown to accelerate wound healing and reduce scarring (Mori et al., 2008), whereas Bmp7 is known to interfere with transforming growth factor beta (TGF-β)-induced differentiation of fibroblasts into myofibroblasts and dermal papilla in vitro (Bin et al., 2013; Liu et al., 2020). Zooming in on fibroblasts, we found bone morphogenetic protein (BMP) 2/4/7 signaling to be among the many pathways with elevated potential to target aged skin wound fibroblasts compared with young counterparts (Figures 7B, 7C, and S7A; Table S7). Conversely, the potential of TGF-β1/2/3 signaling to fibroblasts was reduced in aged skin wounds (Figures 7B, S7A, and S7B). The known BMP-TGF-β antagonism in diverse contexts (Dituri et al., 2019; Hudnall et al., 2016; Keller et al., 2011; Ning et al., 2019; Oshimori and Fuchs, 2012; Sotiropoulos and Chitnis, 2020) together with the role of TGF-β signaling in promoting fibroblast proliferation and ECM production during wound healing (Cutroneo, 2007; Pakyari et al., 2013) suggest that an altered balance between BMP and TGF-β signaling might underlie compromised fibroblast activation in aged skin wounds. Implicating a possible consequence of such imbalance, aged skin wound fibroblasts showed reduced expression of ECM genes Col1a1, Col1a2, Mfap2, and Mfap4, and increased expression of Mmp3 and cell death-promoting gene Cyld (Lork et al., 2017), relative to the young counterparts (Figure S2P).

Only nine ligand-receptor pairs from three signaling pathways showed significant downregulation in aged skin wounds at 4dpw compared with the young, and these include Ptn, Ccl19, Ccl8/5/ 11, and Cxcl10 (Figure 7A). Ptn stimulates keratinocyte proliferation and has been implicated in angiogenesis (Florin et al., 2005; Perez-Pinera et al., 2007), so its downregulation might help explain the delayed re-epithelialization and reduced angiogenesis in aged skin wounds (Sgonc and Gruber, 2013). At 7dpw, similar upregulated (e.g., Spp1, Bmp7, Tnf, and Cxcl12) and downregulated (e.g., Ptn) ligand-receptor pairs were observed in aged skin wounds, but new changes also emerged, including downregulation of Th2 cytokine II13 (Gieseck et al., 2018; Wynn, 2003) and macrophage/Langerhans cell-regulatory cytokine II34 (Hieronymus et al., 2015; Wang and Colonna, 2014) (Figures S7F–S7I).

Chemokines are crucial modulators of wound healing through regulating leukocyte recruitment and integrating inflammatory events with tissue repair processes (Gillitzer and Goebeler, 2001). Altered chemokine signaling in aged skin wounds prompted us to examine and visualize the inferred chemokine signaling-mediated cell-cell communication network. Of the various ligand-receptor pairs, Cc/19 and its receptor Ccr7 displayed the strongest potential to signal from fibroblasts to DCs and migrating DCs (Figures 7D and S7C). Further analysis of the Ccl19-Ccr7 communication network showed that both the predicted strength of major communication routes (e.g., fibroblast-DC/migrating DC) and the total number of potential cellcell interactions (e.g., disappearance of weak fibroblast-macrophage and fibroblast-Treg interactions) were decreased in aged skin wounds at 4dpw relative to young counterparts (Figures 7E and S7D). Decreased cell-cell communication strength was also seen at 7dpw, particularly for the fibroblast-DC/migrating DC and fibroblast-macrophage interaction pairs (Figures S7G-S7J). Reduced expression of Ccl19 by Col1aHi and HFDF I/II fibroblast subsets and slightly reduced expression of Ccr7 by

#### Figure 7. Dysregulated growth factor, chemokine, and cytokine signaling in aged 4dpw wounds

(A) Word cloud showing upregulated (left) and downregulated (right) signaling ligands in aged skin wounds compared with the young. Word size indicates the extent of enrichment in aged or young skin wounds.

- (B) Number of possible interactions targeting wound fibroblasts via BMP or TGF- $\beta$  signaling.
- (C) Inferred BMP-mediated communications from non-fibroblast subsets to fibroblast subsets. The number of putative interactions (i.e., links in the diagram) is indicated on the top.
- (D) Downregulated chemokine signaling in aged skin wounds compared with the young. Signaling sources and target are shown on the bottom and top, respectively. Colored segments indicate their cell identity information. Segment size is proportional to the total outgoing or incoming interaction strength associated with each ligand-receptor pair in the corresponding cell subpopulation.
- (E) Circle plots of inferred Ccl19-Ccr7 signaling networks.
- (F) Comparison of expression levels of Cc/19 and Ccr7 in epithelial, fibroblast, and immune cells in young versus aged skin wounds. \*p < 0.05 (calculated using
- (G) RNAScope showing spatial distribution of Cc/19 and Ccr7 transcripts in young and aged skin wounds. DAPI stains the nuclei. Arrows point to Cc/19-positive cells that line HFs, and asterisk (\*) indicates hair shaft autofluorescence. See Figure S7M for quantification (n = 3 pairs). Scale bars: 500 µm in low-magnification image (left), 100 μm in high-magnification images (right). The rightmost panel shows high-magnification image of an area near the wound margin from Figure S7L.

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DCs/migrating DCs are likely accountable for the reduction in potential signaling strength in aged skin wounds (Figures 7F, S7E, S7H, and S7K). RNAScope validated the reduced expression of both genes in aged skin wound dermis (Figures 7G, S7L, and S7M). In young skin wounds, both Ccl19 and Ccr7 mRNAs were detected in dermis of the unwounded region near the proliferative zone and within the wound bed, with Ccl19 expression seen in scattered dermal cells and what appear to be stromal cells lining the HFs. Ccl19- and Ccr7-positive cells tended to be in proximity of each other but were mutually exclusive (Figure 7G), suggesting a high likelihood of paracrine signaling interactions. In aged skin, Ccl19 and Ccr7 expression was decreased both within the wound bed and outside the wound margin (Figures 7G, S7L, and S7M). Taken together, these observations implicate the respective spatial locations of Col1a<sup>Hi</sup>/HFDF I/II fibroblast subsets and DC/migrating DCs in wounded skin. More importantly, they identify reduced Ccl19 production by these fibroblast subsets as a possible mechanism for compromised retention and increased mobilization of DCs in aged skin and wounds.

#### **DISCUSSION**

Our work presents a comprehensive picture of the cellular compositions, cell-cell communications, and molecular mediators that underlie the wound healing processes in aged versus young skin. The resulting computational predictions and experimental findings not only recapitulate known biology to a substantial degree but also provide proof-of-principle examples for the utility of our scRNA-seg datasets as a useful resource to fuel future hypothesis-driven research that aims at identifying and manipulating critical cell populations and molecular pathways to help aged skin heal better.

Prolonged presence of macrophages in chronic wounds of humans suggests that dysfunctional macrophages likely contribute to impaired healing (Mast and Schultz, 1996), An M1-to-M2 macrophage switch has traditionally been viewed as the process that helps shift wound healing from inflammation to repair and regeneration (Landén et al., 2016), but recent studies have made it increasingly clear that macrophage states in vivo are far more complex (Orecchioni et al., 2019; Watanabe et al., 2019). We now provide evidence for an intermediate state of macrophages with both M1 and M2 characteristics in the healing wounds, namely Arg1Hi macrophages that both express inflammatory genes and exhibit a molecular profile indicative of high hypoxia and glycolysis but low OxPhos. In cancer, Arg1-expressing macrophages occupy the more hypoxic regions compared with macrophages that express Cd206 (Mrc1) (Carmona-Fontaine et al., 2017). We found Arg1Hi and C1qHi (Mrc1-expressing) macrophages to occupy distinct regions within the wound, suggesting skin wound may resemble cancer in its spatial demarcation of M1/M2 and M2 macrophages. Our RNA velocity data raise the intriguing possibility that other monocyte/macrophage subsets might have the potential to become Arg1Hi cells at some time point during wound healing. The increase of Arg1<sup>Hi</sup> and reduction of F4/80<sup>+</sup> macrophages suggest that macrophages in aged skin wounds might be stuck at an Arg1<sup>Hi</sup> inflammatory M1/M2 hybrid stage.

Neutrophils are typically among the first inflammatory cells recruited to the wound to kill pathogens and they are later cleared by apoptosis and macrophage-mediated phagocytosis (Peiseler and Kubes, 2019; Silva, 2011; Wilgus et al., 2013). Alterations in neutrophil dynamics in aged versus young skin wounds remain controversial (Brubaker et al., 2013; Keyes et al., 2016; Mukai et al., 2019; Nishio et al., 2008; Swift et al., 2001), possibly due to differences in mouse lines used, wound size, and time point for analysis. Our study compares young and aged mice from the same source, alleviating potential complications from housing environments, microbiota compositions, and independently emerged mutations. We show that not only do neutrophils persist aberrantly in aged skin wounds but they also express higher levels of inflammatory genes such as neutrophil-attracting chemokine Cxcl3 (Rajarathnam et al., 2019) than their young counterparts. Interestingly, Arg1<sup>Hi</sup> macrophages express higher levels of Cxcl1, Cxcl2, and Cxcl3 than other monocyte/macrophage subsets. These findings suggest that chemotactic signaling from both neutrophils themselves and Arg1Hi macrophages might be responsible for elevated neutrophil infiltration and/or decreased neutrophil clearance, leading to unresolved inflammation that associates with delayed wound healing in aged skin.

Our analysis highlights elevated IL-1 signaling as a candidate pathway that drives aberrant Arg1Hi macrophage-fibroblast communication in aged skin wounds. Supporting the notion that high IL-1 signaling underlies delayed wound healing in aged skin, a previous study found inhibition of IL-1 signaling to improve healing in diabetic skin (Perrault et al., 2018). Our results also confirm, and significantly extend, previous findings (Bageghni et al., 2019; Suwara et al., 2014) to show that skin fibroblasts are a major target of Arg1Hi macrophage-originated IL-1 signaling in aged skin wounds, and that elevated signaling is achieved by both upregulation of the receptor and downregulation of the decoy inhibitor.

HF fibroblast-derived CCL19 has been shown to accelerate wound closure in human experimental models (Topouzi et al., 2020). In this context, our finding of reduced Cc/19 expression in aged skin and wounds suggests that Ccl19 may play a similar role in mouse wound healing, and its reduction may contribute to delayed healing in aged skin. We found Col1aHi dermal fibroblasts and HF-associated (DS) fibroblasts to be the major sources of Ccl19 production and DCs, particularly migrating DCs, to be the major signaling targets. Ccl19 drives Ccr7 desensitization and internalization, and has been implicated in retaining skin DCs by suppressing their migration to draining LNs (Bardi et al., 2001; Bryce et al., 2016; Farnsworth et al., 2019; Kohout et al., 2004). Thus, reduction in fibroblastderived Ccl19 signals in aged skin provides a possible molecular basis for precocious DC mobilization in aged skin wounds. Whether and how DC changes in aged skin functionally contributes to delayed wound healing merits further investigation.

Importantly, our results identify, with strong breadth and cellular resolution, myriad candidate cell-cell communication pairs and systems-level paths of signaling pathways that are affected in aging. Our discovery of increased total number of possible signaling interactions in aged skin wounds supports





the idea that inefficient healing can be a consequence of overactive but misdirected signaling and/or lack of proper downstream response. This notion is in line with previous findings that intercellular communication can be detrimental in aging and related diseases (Fafián-Labora and O'Loghlen, 2020; López-Otín et al., 2013). Given that tissue aging is likely a net sum of varying degrees of cumulative changes in single cells (Todhunter et al., 2018), our single-cell/systems approach to skin aging and associated repair is applicable to understanding aging and regenerative decline of other tissues.

#### **Limitations of the study**

CellChat predictions of cell-cell communications that occur during wound healing in aged versus young skin have not yet incorporated downstream target gene expression and spatial information. Integrating signaling target genes into the computational pipeline faces the challenge that downstream responses to various extracellular signals are often context- and cell type specific. Furthermore, a comprehensive map of spatially resolved transcriptomes in young versus aged skin wounds awaits extensive future work. Experimental and computational studies combining improved scRNA-seq platforms with singlecell assay for transposase-accessible chromatin with sequencing and spatial transcriptomics are expected to uncover regulatory and spatial constraints that will improve prediction accuracy (Longo et al., 2021).

#### **STAR**\*METHODS

Detailed methods are provided in the online version of this paper and include the following:

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#### SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. celrep.2022.111155.

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#### **AUTHOR CONTRIBUTIONS**

X.D. and Q.N. conceived the study and directed the project. R.V. and S.J. performed computational analysis. R.V., P.S., D.H., Q.H.N., and M.D. conducted experiments. K.K. provided guidance on scRNA-seq. R.V., S.J., and X.D. wrote the manuscript with inputs from all authors.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

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#### **STAR**\***METHODS**

#### **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies	OCOTIOE	DENTI LETT
Rabbit Anti-Mouse K14, Purified	Gift: Julie Segre, NIH	N/A
Chicken Anti-Mouse K14, Purified	Gift: Julie Segre, NIH	N/A
F4/80 Monoclonal Antibody	eBioSciences	Cat#: 14-4801-82; RRID:AB 467558
Chemicals, peptides, and recombinant proteins		
Collagenase	Sigma	Cat #: C9091
HEPES	Fisher	Cat #: BP310
Sodium Pyruvate	Fisher	Cat #: BP356
DNase	Sigma	Cat #: DN25
SytoxBlue	Thermo Fisher	Cat #: S34857
Nuclease-free Water	Thermo Fisher	Cat #: AM9937
Low TE Buffer	Thermo Fisher	Cat #: 12090-015
Ethanol	Millipore Sigma	Cat #: E7023-500ML
10% Tween 20	Bio-Rad	Cat #: 1662404
Glycerin	Ricca Chemical Company	Cat #: 3290-32
DAPI	Thermo Fisher	Cat #: D1306
DPBS	Corning Cellgro	Cat #: 21-031-CM
Arg1-Probe	ACD	Cat#: 403431-C1
C1qa-Probe	ACD	Cat#: 441221-C2
Ccr7-Probe	ACD	Cat#: 432871-C1
Ccl19-Probe	ACD	Cat#: 432881-C3
II1a-Probe	ACD	Cat#: 440391-C1
II1r2-Probe	ACD	Cat#: 539491-C3
II1r1-Probe	ACD	Cat#: 413211-C2
Col1a2-Probe	ACD	Cat#: 319371-C2
Critical commercial assays		
Chromium Single Cell 3' Library & Gel Bead Kit v2	10x Genomics	Cat #: PN-120237
Chromium Single Cell 3' Library & Gel Bead Kit v3	10x Genomics	Cat #: PN-1000075
Chromium Single Cell A Chip Kits	10x Genomics	Cat #: PN-120236
Chromium i7 Multiplex Kit	10x Genomics	Cat #: PN-120262
SPRIselect Reagent Kit	Beckman Coulter	Cat #: B23318
Multiplex Fluorescent v2 system	ACD	Cat #: 323100
Deposited data		
Raw scRNA-seq data	This paper	GEO: GSE188432
Experimental models: Organisms/strains		
C57BL/6J	Jackson Laboratory	Stock #: JAX 000664
Software and algorithm		
Cell Ranger 2.1.0	10x Genomics	https://support.10xgenomics.com/single-cell-gene- expression/software/downloads/latest
Cell Ranger 3.1.0	10x Genomics	https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest
Seurat v3	Stuart et al. (2019)	https://satijalab.org/seurat/articles/archive.html
	, ,	https://scvelo.readthedocs.io/
scVelo v0.2.4	Bergen et al. (2020)	Titips://scveio.readthedocs.io/

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
UMAP	McInnes et al. (2018)	https://github.com/lmcinnes/umap
CellChatDB	Jin et al. (2021)	https://github.com/sqjin/CellChat
CellChat v1.5	This paper	https://github.com/sqjin/CellChat
ComplexHeatmap v2.7.1	Gu et al. (2016)	https://jokergoo.github.io/ComplexHeatmap-reference/book/
circlize v0.4.12	Gu et al. (2014)	https://github.com/jokergoo/circlize
wordcloud v2.6	R package	https://cran.r-project.org/web/packages/ wordcloud/index.html
R	R core	https://www.r-project.org/
Python	Python Software Foundation	https://www.python.org/

#### **RESOURCE AVAILABILITY**

#### **Lead contact**

Further information and requests for resources and reagents should be directed to, and will be fulfilled by, the Lead Contact, Xing Dai (xdai@uci.edu).

#### **Materials availability**

This study did not generate new unique reagents.

#### **Data and code availability**

The scRNA-seq data reported in this study have been deposited in the GEO database under accession #GSE188432 (GEO: GSE188432).

CellChat (version 1.5) R package is publicly available at https://github.com/sqjin/CellChat.

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

#### **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

Wild-type C57BL/6J mice are from the Jackson Laboratory (Stock # 000,664). Seven-week-old (young) and 88-week-old (aged) female mice were used for the studies. All maintenance, care, and experiments have been approved and abide by regulatory guidelines of the Institutional Animal Care and Use Committee of the University of California, Irvine.

#### **METHOD DETAILS**

#### Wounding

For single cell experiments, 7-week-old (p49, telogen) and 88-week-old C57BL/6J mice were anesthetized using isoflurane (Primal Healthcare; NDC-66794-017-25), their backs shaved, and then a 6-mm punch (Integra; 33-36) was used to generate a full-thickness wound on each side of the mouse. Wounds were collected 4 and 7 days later for analysis.

#### Single cell isolation for scRNA-Seq

For UW back skin, 7-week-old and 88-week-old C57BL/6J mice were shaved, back skin removed, fat scrapped off, and then skin was minced into pieces less than 1 mm in diameter. For wounded back skin, wound-encompassing skin was first removed, large pieces of fat attached to underside of the wound were carefully scraped off, and a 10-mm punch (Acuderm; 0413) was then used to capture the wound and a portion of un-wounded skin adjacent to the wound. The wounds were then minced into pieces less than 1 mm in diameter. The minced samples were placed in 15-mL conical tubes and digested with 10 mL of collagenase mix [0.25% collagenase (Sigma; C9891), 0.01M HEPES (Fisher; BP310), 0.001M sodium pyruvate (Fisher; BP356), and 0.1 mg/mL DNase (Sigma; DN25)]. Samples were incubated at 37°C for 2 h with rotation, and then filtered through 70-μm and 40-μm filters, spun down, and resuspended in 2% FBS. Cells were stained with SytoxBlue (Thermo Fisher; S34857) as per manufacturer's instructions and live cells (SytoxBlue-negative) were sorted using BD FACSAria Fusion Sorter.

#### Flow cytometry

To obtain single cell suspension, minced samples were digested with 10 mL of a solution containing 0.15% collagenase (Sigma, C9891), 0.01 M HEPES (ThermoFisher, BP310), 0.001 M Sodium Pyruvate (ThermoFisher, BP356), and 0.1 mg/mL DNase (Sigma,





DN25) at 37°C for 1 h with rotation, and then filtered through a 70-µm filter, spun down, and resuspended in 2% fetal bovine serum (FBS). Skin-draining LNs (Inguinal) were dissected and pressed on dish, then resuspended in 2% FBS and filtered through a 70-μm filter. Cells were stained by incubation for 30 min at room temperature with the following antibodies diluted in PBS/2% FBS: Alexa Fluor 488-conjugated anti-CD11b (Biolegend, 101,217), PE-conjugated anti-F4/80 (Biolegend, 123,110), APC-conjugated anti-CD45 (Tonbo biosciences, clone 30-F11, 20-0451), APC-Cy7-conjugated anti-Ly6G (Tonbo biosciences, clone 1A8, 25-1276), PE/ Cyanine7 anti-CD11c (Biolegend, 117,317), FITC anti-MHC Class II (I-A/I-E) (Invitrogen, 11-5321-81).

#### Immunofluorescence and RNAScope

Immunofluorescence was performed as previously described (Dragan et al., 2022) using the following antibodies: anti-K14 antibody (rabbit, gift of Julie Segre, National Institutes of Health), anti-F4/80 monoclonal antibody (eBioSciences; Cat No. 14-4801-82).

RNAScope was performed as previously described using ACD Bio's reagents. Briefly, wounds were freshly frozen in OCT (Fisher; 4585) and sectioned at 10 μm. Sectioned tissues were fixed for 1 h at room temperature with 4% PFA, and the RNAScope Multiplex Fluorescent v2 assay was run per manufacturer's recommendations using the following probes: Arg1 (Cat No. 403431-C1), C1qa (Cat No. 441221-C2), IL1r1 (Cat No. 413211-C2), IL1r2 (Cat No. 539491-C3), Il1a (Cat No. 440391-C1), Ccl19 (Cat No. 432881-C3), Col1a2 (Cat No. 319371-C2), and Ccr7 (Cat No. 432871-C1) from ACD, along with immunofluorescence using anti-K14 antibody (rabbit or chicken). Images were obtained using a Zeiss LSM700 confocal microscope or Keyence BZ-X800 and analyzed using FIJI software.

#### Single cell library generation

FACS-sorted cells were washed in PBS containing 0.04% BSA and resuspended at a concentration of approximately 1,000 cell/uL. Library generation was performed following the Chromium Single Cell 3' Reagents Kits v2 (two young wound samples collected at 7dpw) or v3 (other samples) (following the CG00052 Rev B. user guide) where we target 10,000 cells per sample for capture (see Figure S1B for a detail of the number of samples used for each kit chemistry). Additional reagents included: nuclease-free water (Thermo Fisher Scientific; AM9937), low TE buffer (Thermo Fisher Scientific; 12,090-015), ethanol (Millipore Sigma; E7023-500ML), SPRIselect Reagent Kit (Beckman Coulter; B23318), 10% Tween 20 (Bio-Rad; 1662404), glycerin (Ricca Chemical Company; 3290-32), Qiagen Buffer EB (Qiagen, 19086). Each library was sequenced on the Illumina HiSeg 4000 platform (Chromium Single Cell 3' Reagents Kits v2 samples) or Illumina NovaSeq 6000 platform (Chromium Single Cell 3' Reagents Kits v3 samples) to achieve an average of approximately 50,000 reads per cell.

#### Processing and quality control of scRNA-seg data

FASTQ files were aligned utilizing 10x Genomics Cell Ranger 2.1.0 (Chromium Single Cell 3' Reagents Kits v2 samples) or 3.1.0 (Chromium Single Cell 3' Reagents Kits v2 samples). Each library was aligned to an indexed mm10 genome using Cell Ranger Count. Cell Ranger Aggr function was used to normalize the number of mapped reads per cells across the libraries. Quality control parameters were used to filter cells with 200-5000 genes with a mitochondrial percentage under 15% for subsequent analysis. scRNA-seq data of young mice at UW and 4dpw from Haensel et al. (2020) were processed as previously described.

#### Clustering analysis of scRNA-seq data

Low quality and dead cells (nGenes <200 or >5,000 or percent mitochondria >15%) were excluded from analysis (Figure S1A). Further filtering, normalization, scaling, sample integration, principal component analysis, and cell clustering were performed using the Seurat R package version 3.1.0 (Stuart et al., 2019). Briefly, single cell data matrices were merged according to time point or age (Figures S1B and 2A) and then integrated following the standard workflow described in Seurat package vignettes. To identify cell clusters, the top 10 principal components with a resolution between 0.1 and 0.3 was used. For each v3-only analysis, cells from v3 runs were isolated from each particular aggregated and clustered dataset for plotting and differential gene expression analysis. FindAllMarker function with default parameters was used to find marker genes for each cluster. Major cell types of aggregated young and aged samples for each time point were isolated first and further subclustered. Gene scoring analysis was performed in AddModuleScore function using gene lists as previously described (Haensel et al., 2020).

#### RNA velocity analysis

RNA velocity was calculated based on the spliced and unspliced counts as previously reported (Bergen et al., 2020; La Manno et al., 2018). Loom files were generated using velocity package and used for downstream RNA velocity analysis using scVelo package (Bergen et al., 2020). One 7dpw sample (v2 kit) was not successfully mapped to the unspliced and spliced count and was removed and therefore eliminated from RNA velocity analysis.

#### Systematic inference of cell-cell communication

CellChat was used to systematically infer cell-cell communication by integrating scRNA-seq data with prior ligand-receptor interaction database CellChatDB (Jin et al., 2021). CellChat prediction of putative cell-cell communications occur in three steps. First, CellChat identifies differentially over-expressed ligands and receptors for each cell group. Second, CellChat quantifies the communication probability between two interacting cell groups based on the average expression values of a ligand by one cell group and that

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of a receptor by another cell group, as well as their cofactors. Third, CellChat infers significant interactions using permutation tests. CellChat outputs an intercellular communication network for each ligand-receptor pair, where the calculated communication probabilities are assigned as edge weights to quantify the interaction strength. An intercellular communication network of a signaling pathway is computed by summarizing the communication probabilities of all associated ligand-receptor pairs.

#### Quantitative comparison of cell-cell communication across conditions

To comprehensively identify signaling changes across conditions, we developed comparative CellChat, a computational framework by first focusing on the overall signaling changes on cell population levels and then drilling down to the dysregulated signaling pathways and ligand-receptor pairs. On the cell population level, we seek to identify the cell populations the interactions of which are significantly changed and the markedly altered signaling sources and targets from one condition to another. On the signaling pathway and ligand-receptor pair level, we seek to refine the overall signaling changes in a greater detail to identify significantly altered signaling pathways and ligand-receptor pairs in terms of network architecture and interaction strength. To facilitate intuitive userguided data interpretation, we provide a variety of visualization outputs. We have implemented these new additions in our R package CellChat version 1.5, which is available at https://github.com/sqjin/CellChat. Below we briefly describe this framework.

#### Prediction of general principles of cell-cell communication

We start with the big picture to predict general principles of cell-cell communication. When comparing cell-cell communication across conditions, we seek to answer the following biological questions: (1) the interaction between which cell types is significantly changed; and (2) how the major sources and targets change from one condition to another.

To identify the interaction between which cell populations showing significant changes, we compare the number of interactions between any pair of cell populations across conditions. We compute the differential number of interactions in the cell-cell communication network across two conditions and visualize them using bar plot or heatmap (ComplexHeatmap R package (Gu et al., 2016)). In the heatmap, red and blue colors represent increased and decreased signaling in the second condition compared to the first one, respectively. To identify altered signaling sources and targets across conditions, we employed network centrality analysis by computing the outdegree as measurement of likelihood being signaling sources and indegree being signaling targets. The outdegree and indegree centralities are computed by summarizing each cell population-associated outgoing and incoming communication probabilities across all signaling pathways. We then compute the differential outdegree and indegree centralities of each cell population and visualize them in a 2-dimensional (2D) space, which allows ready identification of cell populations with significant changes in sending or receiving signals across conditions. Positive and negative values indicate increased and decreased signaling in the second condition compared to the first condition, respectively.

#### Identification of dysregulated signaling pathways

To identify signaling changes associated with one cell population, we compute the differential outgoing and incoming interaction strength of this cell population in each cell-cell communication network between two conditions, respectively. Positive and negative values indicate increased and decreased signaling in the second condition compared to the first condition.

#### Identification of upregulated and downregulated ligand-receptor pairs

To identify statistically significant upregulated and downregulated ligand-receptor pairs across conditions, we combine cell-cell communication analysis with differential gene expression analysis. Specifically, for each cell group, we perform Wilcoxon ranksum test for gene expression of cells in the second condition vs. cells in the first condition. Signaling molecules are considered be upregulated in the second condition if (i) the p values are less than 0.05, (ii) the log fold-changes are higher than 0.25, and (iii) the percentage of cells with expression in the second condition is higher than 25%. Signaling molecules are considered be downregulated in the second condition if they are upregulated in the first condition. The ligand-receptor pairs are upregulated or downregulated if both ligands and receptors are upregulated or downregulated. The cell-cell communication mediated by these dysregulated ligand-receptor pairs is visualized by chord diagram using circlize R package (Gu et al., 2014).

To quantify the enrichment of ligand-receptor pairs in one condition, we define the following enrichment score (ES)

$$ES = |logFC_{figand}| \cdot |logFC_{receptor}| \cdot |Percent2_{figand} - Percent1_{figand}| \cdot |Percent2_{receptor} - Percent1_{receptor}|$$

where  $logFC_{ligand}$  is the log fold change (FC) of ligand expression in the second condition compared to the first condition. Percent2<sub>licand</sub> and Percent1<sub>licand</sub> are the percent of ligand-expressed cells in the second and first conditions respectively. The enrichment of ligand-receptor pairs in one condition is visualized by word cloud plot using wordcloud R package (https://cran.r-project.org/ web/packages/wordcloud/index.html). In the word cloud plot, the size of each word (representing each signaling pathway or ligand) indicates its enrichment in one condition compared to another.

#### Cell-cell communication analysis of young and aged wounds

We first applied CellChat to young and aged skin wound datasets separately to infer cell-cell communication at 4dpw or 7dpw. We then performed comparison analysis by merging the CellChat objects from young and aged wound data analyses. After analyzing v2+v3 samples and v3-only samples separately, we utilized our customed R codes to extract the cell-cell communication information that is common between v2+v3 and v3-only data analyses, such that we can highlight the qualitatively consistent results. Specifically, we combined the subclustering results of epithelial cells (Figure S2D), fibroblasts (Figure S2G), and immune cells (Figures 3A and 4A)





at 4dpw, leading to a total of 26 cell groups for analysis, including 6 epithelial subsets (basal, spinous, proliferative basal, HF 1, HF 2, HF 3), 6 fibroblast subsets (Col1a<sup>high</sup>, IMF I, IMF II, myofibroblast, HFDF I/II, HFDF III), and 14 immune subsets (neutrophil, monocyte, Arg1<sup>high</sup> macrophage, C1q<sup>high</sup> macrophage, APM 1, APM 2, OC-like, proliferative and OC-like, DC, migrating DC, Langerhans cell, mast cell, NK and T cell, T-reg). Similarly, we combined the subclustering results of epithelial cells (Figure S2J), fibroblasts (Figure S2M), and immune cells (Figures S3C and S4C) at 7dpw, leading to a total of 22 cell groups for analysis. To infer cell-cell communication between interacting cell groups, we computed the average expression of signaling molecules per cell group using 10% truncated mean, i.e., the mean value after removing the top and bottom 10% cells based on their expression levels.

#### **QUANTIFICATION AND STATISTICAL ANALYSIS**

Data are presented as the mean ± standard deviation (SD), or as indicated. The sample sizes in each plot have been listed in the Results section and Figure Legends where appropriate. For data represented as violin plots, two-tailed Wilcoxon rank-sum test was performed using R (https://www.r-project.org/). For comparison of percentage changes, prop.test was performed using R. For differential gene expression analysis between cell clusters, Wilcoxon rank-sum test was performed using R. A significance threshold of p < 0.01 was used for defining marker genes of each cell cluster. For data presented in bar plot, unpaired two-tailed Student's t-test was used.