Integrated spatial multiomics reveals fibroblast fate during tissue repair

Deshka S. Foster, Michael Januszyk, Kathryn E. Yost, Malini S. Chinta, Gunsagar S. Gulati, Alan T. Nguyen, Austin R. Burcham, Ankit Salhotra, R. Chase Ransom, Dominic Henn, Kellen Chen, Shamik Mascharak, Karen Tolentino, Ashley L. Titan, R. Ellen Jones, Oscar da Silva, W. Tripp Leavitt, Clement D. Marshall, Heather E. des Jardins-Parks, Michael S. Hu, Derrick C. Wan, Gerlinde Wernig, Dhananjay Waghi, John Coller, Jeffrey A. Norton, Geoffrey C. Gurtner, Aaron M. Newman, Howard Y. Chang, and Michael T. Longaker

In the skin, tissue injury results in fibrosis in the form of scars composed of dense extracellular matrix deposited by fibroblasts. The therapeutic goal of regenerative wound healing has remained elusive, in part because principles of fibroblast programming and adaptive response to injury remain incompletely understood. Here, we present a multimodal -omics platform for the comprehensive study of cell populations in complex tissue, which has allowed us to characterize the cells involved in wound healing across both time and space. We employ a stented wound model that recapitulates human tissue repair kinetics and multiple Rainbow transgenic lines to precisely track fibroblast fate during the physiologic response to skin injury. Through integrated analysis of single cell chromatin landscapes and gene expression states, coupled with spatial transcriptomic profiling, we integrated analysis of single cell chromatin landscapes and gene expression states, coupled with spatial transcriptomic profiling, we are able to impute fibroblast epigenomes with temporospatial resolution. This has allowed us to reveal potential mechanisms controlling fibroblast fate during migration, proliferation, and differentiation following skin injury, and thereby reexamine the canonical phases of wound healing. These findings have broad implications for the study of tissue repair in complex organ systems.

spatial epigenomics | spatial transcriptomics | multiomics | fibrosis | chromatin accessibility

**Significance**

In the skin, tissue injury results in fibrosis in the form of a scar composed of dense extracellular matrix deposited by fibroblasts. Therapies that promote tissue regeneration rather than fibrosis remain elusive because principles of fibroblast programming and response to injury remain incompletely understood. Here, we present a multimodal -omics platform for the study of cell populations in complex tissue, which has allowed us to characterize wound healing fibroblasts across both time and space. We identify functionally distinct fibroblast subpopulations and track cell fate during the response to wounding. We demonstrate that populations of fibroblasts migrate, proliferate, and differentiate in an adaptive response to disruption of their environment. These results illustrate fundamental principles underlying the cellular response to tissue injury.

Author contributions: D.S.F., M.J., H.Y.C., and M.T.L. designed research; D.S.F., M.J., M.S.C., G.S.G., A.T.N., A.R.B., A.S., R.C.R., D.H., K.C., S.M., K.T., A.L.T., R.E.J., O.d.S., W.T.L., C.D.M., H.E.d.J.P., M.S.H., D.C.W., and J.A.N. performed research; K.E.Y., G.W., G.C.G., and A.M.N. contributed new reagents/analytic tools; D.S.F., M.J., K.E.Y., D.W., J.C., J.A.N., H.Y.C., and M.T.L. analyzed data; and D.S.F., M.J., H.Y.C., and M.T.L. wrote the paper.

Reviewers: J.L., University of California, Los Angeles; B.M., Memorial Sloan-Kettering Cancer Center; and A.R., University of Pennsylvania.

Competing interest statement: H.Y.C. is a co-founder of Accent Therapeutics, Boundless Bio, and an advisor of 10x Genomics, Arsenal Biosciences, and Spring Discovery. The entities had no role in the design, execution, or interpretation of this research. They had no role in the funding of this research. The paper reports basic research and has no financial implications for the listed companies. Two authors have co-published perspective or review articles with one reviewer over the last 4 y.

This open access article is distributed under Creative Commons Attribution License 4.0 (CC BY).

1D.S.F. and M.J. contributed equally to this work.
2K.E.Y., M.S.C., and G.S.G. contributed equally to this work.
3To whom correspondence may be addressed. Email: howchang@stanford.edu or longaker@stanford.edu.

This article contains supporting information online at https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2110025118/-/DCSupplemental.

Published October 7, 2021.
this model permits us to interrogate fibroblast mechanobiology in a more clinically relevant manner.

Recent advances in sequencing and cell capture technology have enabled the assessment of gene expression with reference to tissue organization using spatial transcriptomics. This approach has only been applied to a limited number of tissue types to date, primarily in the study of tumors, including prostate cancer (10), skin cancer (11, 12), and breast cancer (13), as well as bone marrow (14), joints (15), and brain tissue (16). However, to our knowledge, spatial transcriptomic analysis over time has yet to be applied to characterize wound healing. Moreover, the spatial and temporal distributions of the single-cell chromatin landscapes underlying gene expression have yet to be described.

Here, using transgenic mouse models, we assess the proliferation of local, tissue-resident fibroblast cells in wound healing. By establishing a microsurgical approach to independently isolate fibroblasts from spatially distinct regions within the wound, we interrogate Rainbow-labeled fibroblasts from critical timepoints during the course of wound closure. The Rainbow mouse model is a four-color reporter system that permits precise clonal analysis and lineage tracing. Using this model with phenotype-paired single-cell RNA and ATAC sequencing (scRNA-seq and scATAC-seq), we are able to define the spatial and temporal heterogeneity of wound fibroblasts with unique granularity. By combining spatial transcriptomics with paired scRNA-seq, we assess the differentiation states of individual cells as they proliferate and migrate from the outer wound region inward (17). By disrupting this process using small molecule inhibition or genetic knockdown of focal adhesion kinase (FAK, Ptk2), we further elucidate the relationship between wound healing fibroblast activation and micro-environmental cues. By integrating our scRNA-seq and scATAC-seq analyses using the recently developed ArchR platform (18), we delineate interrelated changes in chromatin accessibility and gene expression driving wound closure and fibrosis and identify distinct wound fibroblast subpopulations. Furthermore, using CIBERSORTx deconvolution (19) of bulk RNA-seq data, we are able to categorize a putative fibroblast subpopulation-based response to local tissue injury. Finally, we introduce spatial multomics, combining spatial transcriptomics with paired scRNA-seq and scATAC-seq datasets to impute spatial epigenomic properties and map chromatin accessibility states in the healing wound. Collectively, this work defines the spatial and temporal dynamics of the fibroblast response to injury and provides a multimodal -omics framework for future studies in tissue repair.

**Results**

**Wounding Triggers Polyclonal Proliferation of Tissue-Resident Fibroblasts.** To explore the lineage dynamics of wound fibroblasts, we examined stented wound healing using the Rainbow (RosaseGFPViGf//Gf) mouse model (20). Rainbow mice contain a transgenic four-color reporter construct in the RosaseG locus. Upon induction with Cre recombinase, the four colors irreversibly recombine such that all progeny cells will have the same color as their parent cells, thereby permitting stochastic lineage tracing and clonal analysis (Fig. 1A). We developed a technique for local induction using activated tamoxifen liposomes (LITMX) in order to induce reporter recombination exclusively in tissue-resident cells (Fig. 1B) (21). Following injury, local skin fibroblasts were found to proliferate in a linear, polyclonal manner along the cross-sectional wound interface (Fig. 1 C and D), whereas fibroblasts in uninjured skin exhibited minimal clonality (Fig. 1 E and F). These data support the presence of local cells that are activated in response to injury and proliferate polyclonally to fill the wound “gap.”

Many cell surface and lineage markers have been associated with fibroblasts involved in wound healing, including Pdgfra, Engrailed-1 (En1), and CDS2 (Dpp4) (22–24). However, we and others have found expression of such markers to be variable throughout wound tissue (SI Appendix, Fig. S1A), suggesting spatial and functional heterogeneity among the fibroblasts that respond to injury. We asked whether there might be one or more fibroblasts activated following injury that could give rise to more diverse downstream fibroblast phenotypes. If so, we wondered whether such cells would be of tissue-resident origin, as suggested by previous studies (25–27) (SI Appendix, Fig. S1B), or originate from peripheral circulation. To explore this, we employed transgenic parabiotic mice in conjunction with the splinted excisional wound healing model described above (6) (SI Appendix, Fig. S1C). cGFP donor mice were parabiosed to wild-type (C57BL/6J) mice (SI Appendix, Fig. S1D). A shared blood supply was established by 2 wk after surgery (SI Appendix, Fig. S1E), at which time wounds were made on the dorsum of each wild-type parabiont. Wounds were then harvested at postoperative day (POD) 7 (midway through healing) or POD 14 (when the wound has fully reepithelialized). While systemically infiltrating GFP+ cells were found in wild-type mouse wounds at both timepoints, the overwhelming majority (>80%) of GFP+ cells were also CD45+ and thus of hematopoietic (nonfibroblast) lineage (SI Appendix, Fig. S1F). These data further support the growing body of literature indicating that the fibroblasts responsible for wound healing are local, tissue-resident cells (25–27).

Returning to the Rainbow mouse model, we developed a tissue clearing and whole-mount protocol to visualize wound healing biology with the Rainbow mouse (28). Using these methods in conjunction with a ubiquitous Actin-CreERT2-driven driver, we observed that cells were activated along the wound edge and proliferated inward in a distinct radial pattern (Fig. 1 G and H).

**Bulk Transcriptomic Analysis of Injury-Responsive Fibroblasts.** Based on the pattern of clonal proliferation extending from the outer wound edge inward, we developed a microsurgical technique to separately isolate the “inner” and “outer” components of the wound dermis (Fig. 1J). We isolated wound fibroblasts from these two regions at POD 7 (midpoint of healing) and unwounded skin for bulk RNA-seq evaluation. Clear differences in the gene expression profiles of inner versus outer wound fibroblasts were identified (Fig. 1 J and K and SI Appendix, Fig. S2A–C), including differences in mechanotransduction and cell cycle pathways. Furthermore, we observed that inner wound fibroblasts were transcriptionally more divergent from uninjured skin than were outer wound fibroblasts (SI Appendix, Fig. S2A). These findings support broad regional differences in the proliferation and activation status of fibroblasts in the healing wound; however, these methods are limited by the lack of granularity inherent in bulk transcriptional analysis.

**Traditional Cell Surface Markers Are Not Sufficient to Characterize Regional Heterogeneity among Wound Healing Fibroblasts.** We evaluated how well several recently published cell surface marker profiles, which define fibroblast subtypes largely based on tissue depth, tracked with the regional differences observed in our study (22). Among fluorescence activated cell sorting (FACS)-isolated, lineage-negative (29), Rainbow wound fibroblasts (Fig. 2A and SI Appendix, Fig. S3A), we found that most cells fell into the putative category of reticular fibroblasts (defined as DLK1+/SCA1−) rather than papillary (CD26+/SCA1−) in wild-type mouse wounds at both timepoints, the overwhelming majority (>80%) of GFP+ cells were also CD45+ and thus of hematopoietic (nonfibroblast) lineage (SI Appendix, Fig. S1F). These data further support the growing body of literature indicating that the fibroblasts responsible for wound healing are local, tissue-resident cells (25–27).

**Single-Cell Transcriptomic Analysis of Injury Responsive Fibroblasts.** We sought to better characterize wound fibroblast heterogeneity...
Fig. 1. Wounding triggers polyclonal proliferation of tissue-resident fibroblasts. (A) Schematic of the Rainbow mouse construct. (B) Schematic showing wound healing model using Rainbow mice with local Cre recombinase induction using 4-hydroxytamoxifen liposomes (LiTMX). (C) Schematic showing a Rainbow wound cross-section. Black dotted line highlights wound scar area; arrows indicate the direction of cellular proliferation during wound healing. Structures are as labeled. (D) Representative confocal image of POD 14 wound cross-sections from Actin-CreER\textsuperscript{T2}::Rosa26\textsuperscript{VT2/GK3} mice induced locally with LiTMX at the time of wound creation. Thick white dotted lines highlight scar boundaries. Individual Rainbow cell clones are highlighted with thin colored dotted lines. Arrows indicate direction of wound healing. \( n > 5 \). (Scale bar, 50 \( \mu \)m.) (E) Representative confocal images of unwounded skin from Actin-CreER\textsuperscript{T2}::Rosa26\textsuperscript{VT2/GK3} mice induced locally with LiTMX. Thick white dotted lines highlight dermal boundaries. Individual Rainbow cell clones are highlighted with thin white dotted lines. \( n > 5 \). (Scale bar, 50 \( \mu \)m.) (F) Rainbow clone counts in wounds versus uninjured skin. \( n = 5 \) per condition. \( *P < 0.05 \). (G) Schematic of dorsal, stented, excisional wound healing in the Rainbow mouse model (whole-mount view), with polyclonal proliferation of Rainbow fibroblasts from the outer wound edge inward across time from POD 2 (Left), to POD 7 (Middle), to POD 14 (Right). Black arrows highlight the apparent direction of proliferation. (H) Representative confocal imaging of a POD 14 whole-mounted wound harvested from Actin-CreER\textsuperscript{T2}::Rosa26\textsuperscript{VT2/GK3} mice showing the polyclonal proliferation of wound fibroblasts radially toward the center of the wound (dark area at center). White arrows highlight the direction of cell proliferation; individual cell clones are highlighted with thin colored dotted lines. Bottom subpanels denote individual Rainbow color contributions to merged image. mCh, membrane (m)Cherry; mOr, mOrange; mCe, mCerulean; eG, eGFP. \( n > 5 \). (I) Schematics illustrating microdissection strategy for isolation of inner and outer wound regions (Top), followed by enzymatic separation of the dermal scar from the epi- and hypodermis (Bottom). (J) Heatmap displaying expression data for genes significantly different between POD 7 inner and outer region wound fibroblasts. Legend at Right displays fold change. (K) Gene Ontology (GO) enrichment analysis comparing gene expression data from POD 7 inner and outer region wound fibroblasts. Top shows GO biological processes up-regulated in inner region fibroblasts compared with outer region fibroblasts, while the Bottom shows the same for outer region fibroblasts compared with inner. Top 10 most significant gene sets are displayed for each condition.
Fig. 2. Single-cell transcriptomic and chromatin accessibility analyses delineate mechanoresponsive fibroblast subpopulations. (A) Schematic illustrating single-cell (sc) isolation of Rainbow wound fibroblasts from inner and outer wound regions (highlighted with black dotted lines). For scRNA-seq, mCerulean+ fibroblasts were arbitrarily selected from the available Rainbow colors and used for the remaining experiments in this figure. (B) (Left) Uniform manifold approximation and projection (UMAP) embedding showing scRNA-seq data from mouse wound fibroblasts FACS isolated using a lineage-negative sort strategy (29) from POD 2, POD 7, and POD 14, digitally pooled and clustered in a manner agnostic to POD and inner versus outer wound regions. Four unique fibroblast clusters were identified (clusters 1 through 4). Dotted lines highlight individual cluster distributions. (Right) Recoloring of Left UMAP plot based on fibroblast tissue region: inner (black) versus outer (orange). (C) CytoTRACE analysis of scRNA-seq data using the UMAP embedding from F. Shading indicates inner (light gray) versus outer (dark gray) wound regions. (D) Box plots showing the predicted ordering by CytoTRACE for individual cells within the four scRNA-seq clusters. Gray arrow indicates direction of predicted differentiation from scRNA-seq cluster 1 to cluster 4 (which corresponds to outer-to-inner wound region expansion). P value was derived from two-sided Student’s t test. (E) scATAC-seq evaluation of Rainbow mouse wound fibroblasts isolated in parallel with our scRNA-seq experiments (SI Appendix, Methods), integrated using the ArchR toolkit with default Louvain parameters (18) to delineate four unique multimodal fibroblast clusters. (F) Heatmap of scATAC-seq motifs highlighting key gene loci differentially open or closed in putative fibroblast subpopulations. (G) Genome tracking plots showing scATAC-seq peaks for pseudobulk replicates generated for each cluster. Associations between the peaks with fibrosis and mechanotransduction-related genes (Peak2GeneLinks) are included at the Bottom of each plot. Pale orange shading highlights differentially expressed peaks across the scATAC clusters. All highlighted peaks demonstrated statistically significant differential expression in at least one pairwise comparison (false discovery rate [FDR] <0.1 and fold change [FC] ≥2).
by examining individual fibroblast transcriptional programs at important functional timepoints in the canonical wound healing process: POD 2, inflammation; POD 7, granulation; and POD 14, complete reepithelialization (“healed” wound). We conducted plate-based scRNA-seq of lineage-negative fibroblasts isolated based on their expression of Rainbow clone colors from both inner and outer wound regions at each timepoint (Fig. 2A). Four transcriptionally defined fibroblast subpopulations were identified (Fig. 2B), with considerable differences in their distributions between wound regions.

Given our interest in understanding lineage trajectories in the context of wound healing, we assessed the relative differentiation states of these fibroblast populations using CytoTRACE, a computational tool that leverages transcriptional diversity to order cells based on developmental potential (Fig. 2C and SI Appendix, Fig. S4) (17). This analysis identified a lineage trajectory stemming from scRNA-cluster 1, which is characterized by elevated expression of fibroblast markers such as Pdgfra and primarily represented by cells from the outer wound region, extending to scRNA-cluster 4, which is primarily represented by cells from the inner wound (Fig. 2D). These findings suggest that fibroblasts may undergo differentiation as they proliferate from the outer wound inward.

**Evaluation of Chromatin Accessibility Complements Transcriptional Analysis of Mechanoresponsive Fibroblast Subpopulations.** To evaluate the epigenomic changes associated with fibroblast activation and lineage differentiation in wound healing, we conducted a series of scATAC-seq experiments in parallel with our scRNA-seq assays (SI Appendix, Fig. S5 A and B). We identified considerable heterogeneity in accessibility profiles among individual wound fibroblasts, which were clustered into six epigenomically distinct subgroups using the ArchR platform (18) (Fig. S2B, SI Appendix, Figs. S5 C and D and S6 A–D). This partitioning was agnostic to the phenotype of cell origin (i.e., wound region or postoperative day), and all clusters included fibroblasts harvested from multiple timepoints and wound regions. We then performed cross-platform integration to link these scATAC data with our earlier scRNA data (18), resulting in four multimodal clusters characterized by both gene expression and chromatin accessibility profiles (Fig. 2E and F and SI Appendix, Fig. S6E), which we refer to as ArchR-clusters 1 through 4.

We first examined the epigenomic landscape of the largest subpopulation, ArchR-cluster 1, which showed significantly elevated chromatin accessibility proximal to key fibrosis-related genes such as Col1a1, Acta2, and Pdgfra (Fig. 2G and SI Appendix, Figs. S7 A and B and S8), indicating that these cells are primed for their transcription. We also observed specific accessibility peaks and transcription factor footprinting in association with the FAK (Ptk2) locus and its downstream signaling elements such as Jun, suggesting that these fibroblasts may represent a mechanoresponsive, profibrotic subpopulation. ArchR-cluster 2 was associated with elevated Fln1 and Tgfbs1 accessibility peaks; ArchR-cluster 3 was characterized by increased accessibility at theJak2 locus and decreased accessibility at the Fsp1 (S100a4) and Il6st loci; and ArchR-cluster 4 was characterized by increased accessibility at the Ptk2b, Jak1, and Jak3 loci.

In addition to specific peak and motif evaluation, we also employed clusterwide enrichment analysis using the Genomic Regions Enrichment of Annotations Tool (GREAT) (30) (SI Appendix, Fig. S9A). We found significant enrichment for “increased fibroblast migration,” “focal adhesion,” and FAK-pathway signaling response elements in ArchR-cluster 1. Furthermore, pseudotime analysis of these integrated scRNA–ATAC data demonstrated an epigenomic progression from the putatively least-differentiated ArchR-cluster 1 to the remaining cell populations that was driven by mechanical signaling elements (SI Appendix, Fig. S9B).

Based on these findings, we provisionally characterized each subpopulation according to its putative role in the wound healing process: “mechanofibrotic” (ArchR-cluster 1), “activated-responder” (ArchR-cluster 2), “remodeling” (ArchR-cluster 3), and “proliferator” (ArchR-cluster 4) fibroblasts.

**Clonal Proliferation of Injury-Responsive Fibroblasts Is Mechanotransduction Dependent.** Our laboratory has previously shown that local tissue mechanics are crucial in guiding the response to healing after injury (31), and mechanotransduction signaling pathway elements were found to delineate fibroblast subpopulations in our scRNA and scATAC wound data. To further interrogate the role of local tissue mechanics in wound biology, we applied a small molecule FAK inhibitor (FAK) to disrupt mechanosensation in stented mouse wounds (SI Appendix, Fig. S10A). Consistent with prior work, we found that FAK-treated wounds healed at the same rate as untreated wounds (SI Appendix, Fig. S10B and C) but resulted in significantly smaller and thinner scars composed of less-dense matrix tissue (SI Appendix, Fig. S10D and E) (32).

To validate our FAK results, we conducted additional wound healing experiments using Actin-CreERT2; Rosa26R2T2G3K: Ptk2fl/fl and Actin-CreERT2; Rosa26R2T2G3K: Ptk2fl/fl (heterozygous Ptk2fl/+ and homozygous Ptk2fl/fl knockout) mice, with local LTMX induction at the time of wounding (SI Appendix, Fig. S10A–C). We found that these mouse wounds also exhibited fewer scar-like patterns of connective tissue (SI Appendix, Fig. S10E). To further explore these differences, we employed an automated feature extraction algorithm (24) to quantify ultrastructure characteristics of wound tissue sections, which demonstrated that FAK-treated wound specimens were more similar to unwounded skin than to vehicle-control wounds, including for both mature and immature collagen fiber intensities (SI Appendix, Fig. S10F). Taken together, these findings corroborate that when mechanotransduction is disrupted, wounds heal with thinner scars and connective tissue structure that is more similar to that of unwounded skin.

To understand the transcriptional changes associated with modulation of mechanotransduction in wound healing, we conducted additional RNA-seq experiments comparing fibroblasts isolated from inner and outer regions of FAK-treated and control wounds. We observed significant changes in the transcriptional programs of FAK-treated cells and found that regional differences between inner and outer wound fibroblasts were dampened in wounds following FAK inhibition (SI Appendix, Fig. S11A and B). These results suggest that local tissue mechanics contribute to transcriptional differences between inner and outer wound regions.

We found that wound healing fibroblasts showed down-regulation of mechanotransduction- and fibrosis-related pathways with FAK treatment (SI Appendix, Fig. S11C). We also found that when mechanosignaling was blocked in Rainbow mice using FAKκ, or in Pk2fl/+ or Pk2fl/fl mice, the linear polyclonal proliferation of fibroblasts that was previously appreciated (Fig. 1H) was disrupted (Fig. 3A–C), with smaller and less ordered Rainbow fibroblast clones.

We applied the deconvolution tool CIBERSORTx (19) to estimate the abundance of our four scRNA–ATAC populations (ArchR-clusters 1 through 4) within bulk RNA-seq data for fibroblasts isolated from POD 7 and POD 14 wounds with or without FAK treatment (Fig. 3D). We found that the majority of cell estimates across all specimens were attributed to mechanofibrotic ArchR-cluster 1, consistent with its prominent representation in both our scRNA-seq and scATAC-seq datasets. The predicted prevalence of these cells was highest at POD 7 and decreased by POD 14. FAK inhibition resulted in decreased representation of ArchR-cluster 1 fibroblasts at POD 14 for both inner and outer wound samples (compared to control wounds at POD 14), further supporting the mechanosensitivity of the putative mechanofibrotic ArchR-cluster 1 subpopulation.
Spatial Transcriptomics Applied to Wound Healing. To further explore the significance of fibroblast heterogeneity in healing wounds, we applied the recently developed 10x Genomics Visium platform to analyze gene expression while retaining tissue spatial information. We optimized and validated a protocol to enable highly reproducible Visium spatial transcriptomic analysis of skin and wounds across the healing process (SI Appendix, Methods). We then conducted spatial transcriptomic analysis on tissue from our stented Rainbow mouse wound healing model at POD 2, 7, and 14, as well as uninjured skin (Fig. 4A).

The epidermal, dermal, and hypodermal layers of the healing wounds were easily delineated histologically and also found to cluster independently based on transcriptional programs (Fig. 4B). Looking at individual genes for prominent wound healing cell types (Fig. 4C, i), we found clear delineation of keratinocytes in the epidermis based on Krt6b expression (as well as other keratinocyte-specific genes), allowing us to examine recapitulation of keratinocyte expression over space and time at the transcriptional level (Fig. 4C, ii). Similarly, fibroblast activity was evaluated using characteristic genes such as Pdgfra, which were most prominent in the dermis and most active at POD 14 (Fig. 4C, iii). Likewise, by examining activated macrophage markers like Msl1, we could monitor these immune cells throughout our dataset and found that they were very prominent in the “proud flesh” at the center of the wound at POD 7 (Fig. 4C, iv).

One challenge inherent in current spatial transcriptomic platforms such as Visium is that each “spot” (i.e., discrete spatial subregion from which transcripts are sequenced) can capture gene expression information from more than one cell (1 to 10 cells, characteristically). In a complex tissue such as a healing wound, this often includes cells of different types, particularly within the dermis where fibroblasts, multiple types of immune cells, and nascent blood vessels can be found. As such, to understand our spatial transcriptomics results in the context of our scRNA- and scATAC fibroblast data, we needed to account for the contributions of nonfibroblast cells from each Visium spot. This was achieved by first estimating the number of each specific cell type present within individual spots based on the associated histological staining (SI Appendix, Figs. S12A–D and S13). Cell counting was followed by random sampling in a Monte Carlo fashion to “subtract out” potential contributions from nonfibroblast cells, generating a distribution of 10,000 inferred fibroblast transcriptomes for each Visium spot. These were propagated forward for anchor-based integration to generate and pool spatially overlaid partial memberships for each of our four scRNA-clusters (Fig. 4D).

We found that the predicted spatial distributions for our scRNA-seq clusters were largely congruent with the transcriptional differences observed earlier between inner and outer cells using our microdissection approach (e.g., fibroblasts belonging to the mechanofibrotic cluster became more prominent over time, expanding from the outer to inner wound regions to fill the scar). Upon further examining transcriptional programming relative to tissue depth, we observed clear spatial distinctions between the apical and basal regions of the dermis as early as POD 7 and most prominently at POD 14 (SI Appendix, Figs. S14A–F and S15A and B). For example, the MMP inhibitor Timp1 is expressed by
Fig. 4. Spatial transcriptomics applied to wound healing and tracking of fibroblast subpopulations over time and space. (A) Schematic for generating spatial transcriptomics data from splinted excisional wounds using the 10x Genomics Visium protocol. Fresh Rainbow mouse wound tissue was harvested, flash frozen, embedded in optimal cutting temperature (OCT), and then sections were taken representing the complete wound radius. H&E staining and tissue section imaging were completed as described in the Visium protocol (SI Appendix, Methods). Each spot captures mRNA from 1 to 10 individual cells at that tissue location. (B) Delineation of scar layers based on underlying tissue histology at each timepoint (Top row), and UMAP plot showing that the three scar layers can easily be distinguished by their transcriptional programs, even independent of spatial information. (C) (i) Schematic of classic stages of wound healing evaluated at POD 2, 7, and 14 relative to uninjured skin. (ii) Keratinocyte activity as measured through expression of the Krt6b gene. (iii) Fibroblast activity as measured through expression of the Pdgfra gene. (iv) Immune cell activity as measured through expression of the Msr1 gene. (D) Anchor-based integration of scRNA-seq populations (defined in Fig. 2B) with Visium gene expression to project partial membership within each spot across all timepoints. These populations exhibit strong spatial preferences within the wound.
fibroblasts in the basal dermis, while Tlbs2, which mediates cell–matrix interactions, is primarily expressed in the more apical scar region.

To assess the relative differentiation states of fibroblasts in this system, we applied CytoTRACE to our POD 14 dermal scar data and found that, similar to our RNA-seq microdissection findings, fibroblasts exhibited significantly less transcriptional diversity in inner wound regions, further supporting fibroblast differentiation from the outer to the inner wound regions during tissue repair (SI Appendix, Fig. S16).

Integrated Analysis Permits Imputation of Spatial Epigenomic Properties. To further explore fibroblast cell fate with spatial resolution, we developed a method to combine our integrated single cell RNA–ATAC framework with Visium in order to impute spatially informed epigenomes for wound healing fibroblasts (Fig. 5A). As described above, we generated spatial transcriptomic data from unwounded skin and POD 2, 7, and 14 wounds. To extend this analysis to impute spatial epigenomic properties, we used our RNA–ATAC construct to ascribe partial membership values to fibroblasts present within each Visium spot. This was achieved by first subtracting out putative nonfibroblast contributions as described above, followed by anchor-based mapping into a higher-dimensional cluster space from our gene integration matrix (Fig. 5B and SI Appendix, Table S1). Parameterization was optimized to preserve spatial autocorrelation for the top measured and imputed gene expression distributions within the POD 14 dermis (SI Appendix, Fig. S17 A and B). To account for residual contributions from nonfibroblast cells that may remain after our initial subtraction step, we also spiked in RNA-seq data for keratinocytes, endothelial cells, macrophages, and neutrophils. The resulting putative reference matrix was then used to assign initial partial set memberships for each spatial datapoint using an anchor transfer–based approach. A single-step spatial smoothing filter was applied to this membership space, followed by removal of nonfibroblast contributions and renormalization. The resulting partial set memberships for each spatial datapoint then allowed us to project higher-order epigenomic features from the scRNA–ATAC data onto these Visium samples (SI Appendix, Fig. S18 A–D). These spatial epigenomic imputations provided a valuable complement to further refine our understanding of the fibroblast biology driving tissue repair. Detailed data analysis is provided in Fig. 5 C and D and SI Appendix, Figs. S19 and S20 and more broadly summarized below for each timepoint in the healing process.

Immediately following wound injury, tissue trauma leads to inflammatory cell recruitment, provisional clot formation, and a dermal gap resulting in loss of contact inhibition among local fibroblasts. These fibroblasts are recruited into the wound bed and begin proliferating. Our data suggest that by POD 2, subsets of these cells have differentiated along the wound margin to form a putative Activated-Responder Fibroblast subpopulation. Other, less-differentiated and more mechanosensitive (mechanofibrotic), fibroblasts become reactivated in the deeper dermis at this point, increasing chromatin accessibility for Runx2, which is a primary regulator of mesenchymal progenitor cell proliferation and differentiation (33).

By POD 7, macrophage-dominated granulation tissue occupies the central wound defect, allowing overlying keratinocyte proliferation and reepithelialization. At this time, mechanofibrotic Fibroblasts begin to differentiate as they finish migrating toward the wound center, where they appear to transition to a more Proliferator subpopulation. These cells are strongly profibrotic and characterized by elevated Spp1 gene expression and chromatin accessibility. In parallel, a population of Remodeling Fibroblasts begins to appear in the outer deep dermis (Fig. 4 C and D and SI Appendix, Fig. S15 A and B).

At POD 14, reepithelialization is complete, and the wound is traditionally considered to be healed. However, while keratinocyte activity does decrease at this time (consistent with completion of reepithelialization), there remains a strong immune cell presence, supported by continued wound fibroblast chemokine secretion, to stimulate active fibrosis in the dermal layer (SI Appendix, Fig. S14 E and F).

Considering our imputed spatial epigenomics data more globally, we observed that changes to chromatin accessibility frequently preceded downstream changes in gene expression, even within the constraints of our coarse temporal sampling (Fig. 5 C and D and SI Appendix, Fig. S19 A–C). For example, we found that the Runx1 motif, which is downstream from and regulated by FAK mechanotransduction, initially becomes open at POD 2, remains open particularly along the leading wound edge at POD 7, and then begins to decrease in accessibility throughout the nascent scar at POD 14. Similarly, Colla2 motif opening precedes a dramatic increase in Colla2 gene expression seen in the POD 14 wound scar.

In aggregate, these studies represent a framework for the comprehensive elucidation of wound healing fibroblast phenotypes based on both gene expression and chromatin accessibility across time, space, and lineage. Furthermore, these findings allow us to reevaluate the classical stages of wound healing, typically described as three overlapping phases: inflammation (POD 2), proliferation (POD 7), and remodeling (POD 14) (3). Based on our findings, we propose reframing these overlapping stages as: 1) Early inflammation, in which immune cells are migrating and infiltrating the injury site without proliferation; 2) reepithelialization, which includes rapid keratinocyte proliferation across the wound surface, fibroblast recruitment, and macrophage proliferation; and 3) activated fibrosis, in which maximal fibroblast activation is achieved and sustained in a slow asymptotic decay by steady-state inflammatory signaling beneath the healed wound (SI Appendix, Figs. S21 A–D and S22 A and B).

Discussion

In this manuscript, we define fibroblast biology throughout the course of wound healing using integrated, single-cell multimodal -omics to unravel the spatial, temporal, and functional heterogeneity of these cells. We demonstrate that fibroblasts are activated from tissue-resident cells in response to injury and proliferate polyclonally to fill the wound gap. Furthermore, we demonstrate that fibroblasts undergo spatially informed differentiation during this process.

Elucidating these functional relationships required the integration of nascent technologies and data platforms in what is still a rapidly evolving field of multimomic imputation. This work demonstrates the paired analysis of single-cell RNA and chromatin accessibility with spatial resolution in the context of tissue repair. This approach provides a unique lens through which we can view complex cell processes, and specifically allowed us to demonstrate that upstream chromatin changes surrounding mechanical signaling elements precede transcriptional activation and cell proliferation, thus suggesting a mechanistic link from tissue force to activation of wound healing fibroblasts.

Furthermore, we were able to identify and characterize putative, functionally distinct fibroblast subpopulations with divergent transcriptional and epigenomic programs. We provisionally designate these four wound healing fibroblast phenotypes as Mechanofibrotic, Activated Responder, Proliferator, and Remodeling. Following skin injury, fibroblasts are locally recruited and migrate to the wound. By POD 2, a subset of fibroblasts appears to have differentiated to form an activated-responder subpopulation, while the remaining outer wound fibroblasts comprise the less differentiated mechanofibrotic cells. The latter fibroblasts highly express known fibrosis-associated markers such as Engrailed-1 (23, 24), Colla1 (34), Tgbf2 (35), and Jun (36). At POD 7, mechanofibrotic cells begin to differentiate in response to mechanotransduction cues as they migrate toward the wound center. By POD 14, despite complete epitelialization, healed wounds remain in a steady state of fibrosis, maintained through sustained inflammatory signaling within scar
Additional studies examining even later timepoints will be required to further characterize the dynamics of these cells within the healed scar tissue. Taken together, these results illustrate fundamental principles underlying the cellular response to tissue injury. We demonstrate that populations of fibroblasts migrate, proliferate, and differentiate in an adaptive, dynamic response to disruption of their local mechanical environment. Understanding the origin, activation, and differentiation trajectories of injury-responsive cells is critical to develop therapeutic strategies to promote optimal tissue repair.

**Materials and Methods**

All animal experiments were approved by and conducted in accordance with the regulations of the Stanford University Animal Care and Use Committee. Mouse husbandry, transgenic models, immunostaining, high-throughput -omics, and computational methods are described in SI Appendix.

**Data Availability.** All sequencing datasets generated in this study are freely available through the Gene Expression Omnibus (GEO), with accession number GSE178758.

**ACKNOWLEDGMENTS.** We acknowledge the Stanford Functional Genomics Facility; Genome Sequencing Service Center; Beckman Cell Sciences Imaging
1. V. J. Thannickal, Y. Zhou, A. Gaggar, S. R. Duncan, Fibrosis: Ultimate and proximate causes. J. Clin. Invest. 124, 4673–4677 (2014).
2. T. A. Wynn, Fibrotic disease and the Th1/Th2 paradigm. Nat. Rev. Immunol. 4, 583–594 (2004).
3. G. C. Gurtner, S. Werner, Y. Barrandon, M. T. Longaker, Wound repair and regeneration. Nature 453, 314–321 (2008).
4. C. F. Guerrero-Juarez et al., Single-cell analysis reveals fibroblast heterogeneity and myeloid-derived adipocyte progenitors in murine skin wounds. Nat. Commun. 10, 650 (2019).
5. M. V. Plikus et al., Regeneration of fat cells from myofibroblasts during wound healing. Science 355, 748–752 (2017).
6. R. D. Galliano, J. Michaelis, V. Dobryansky, J. P. Levine, G. C. Gurtner, Quantitative and reproducible murine model of excisional wound healing. Wound Repair Regen. 12, 485–492 (2004).
7. S. Aarabi et al., Mechanical load initiates hypertrophic scar formation through decreased cellular apoptosis. PASEB J. 21, 3250–3261 (2007).
8. J. Wang et al., An updated review of mechanotransduction in skin disorders: Transcriptional regulators, ion channels, and microRNAs. Cell. Mol. Life Sci. 72, 2091–2106 (2015).
9. L. A. Barnes et al., Mechanical forces in cutaneous wound healing: Emerging therapies to minimize scar formation. Adv. Wound Care (New Rochelle) 7, 47–56 (2018).
10. E. Berglund et al., Spatial maps of prostate cancer transcriptomes reveal an unexplored landscape of heterogeneity. Nat. Commun. 9, 2419 (2018).
11. A. L. Ji et al., Multimodal analysis of composition and spatial architecture in human squamous cell carcinoma. Cell 182, 497–514.e22 (2020).
12. K. Thrane, H. Eriksson, J. Maaskola, J. Hansson, J. Lundeberg, Spatially resolved transcriptomics enables dissection of genetic heterogeneity in stage III cutaneous malignant melanoma. Cancer Res. 78, 5970–5979 (2018).
13. J. Bergström, L. Larsson, J. Lundeberg, Seamless integration of image and molecular analysis for transcriptomics workflows. BMC Genomics 21, 482 (2020).
14. C. Baccin et al., Combined single-cell and spatial transcriptomics reveal the molecular, cellular and spatial bone marrow niche organization. Nat. Cell Biol. 22, 38–48 (2020).
15. K. Carlborg et al., Exploring inflammatory signatures in arthritic joint biopsies with spatial transcriptomics. Sci. Rep. 9, 18975 (2019).
16. A. Andersson et al., Single-cell and spatial transcriptomics enables probabilistic inference of cell type topography. Commun. Biol. 3, 565 (2020).
17. G. S. Gulati et al., Single-cell transcriptional diversity is a hallmark of developmental potential. Science 367, 405–411 (2020).
18. J. M. Granja et al., ArchR is a scalable software package for integrative single-cell chromatin accessibility analysis. Nat. Genet. 53, 403–411 (2021).
19. A. H. Newman et al., Determining cell type abundance and expression from bulk tissues with digital cytometry. Nat. Biotechnol. 37, 773–782 (2019).
20. H. Yani, T. Tanaka, H. Ueno, Multicolor lineage tracing methods and intestinal tumors. J. Gastroenterol. 48, 423–433 (2013).
21. R. C. Ransom et al., Genetic dissection of clonal lineage relationships with hydroxymethylxen liposomes. Nat. Commun. 9, 2971 (2018).
22. R. R. Driskell et al., Distinct fibroblast lineages determine dermal architecture in skin development and repair. Nature 504, 277–281 (2013).
23. Y. Rinkevich et al., Skin fibroblasts. Identification and isolation of a dermal lineage with intrinsic fibrogenic potential. Science 348, aaa2151 (2015).
24. S. Mascharak et al., Preventing Engrailed-1 activation in fibroblasts yields wound regeneration without scarring. Science 372, eaaz374 (2021).
25. T. Banic-Dujmovic, I. Boban, S. H. Clark, Fibroblasts/myofibroblasts that participate in cutaneous wound healing are not derived from circulating progenitor cells. J. Cell. Physiol. 222, 703–712 (2010).
26. H. C. Grillo, Origin of fibroblasts in wound healing. An autoradiographic study of inhibition of cellular proliferation by local x-irradiation. Ann. Surg. 157, 453–467 (1963).
27. R. Ross, N. B. Everett, R. Tyler, Wound healing and collagen formation. VI. The origin of the wound fibroblast studied in parabiosis. J. Cell Biol. 44, 645–654 (1970).
28. D. S. Foster et al., A clearing technique to enhance endogenous fluorophores in skin and soft tissue. Sci. Rep. 9, 15791 (2019).
29. G. G. Walmsey et al., Murine dermal fibroblast isolation by FACS. J. Vis. Exp. 10,3791/53430 (2016).
30. C. Y. McLean et al., GREAT improves functional interpretation of cis-regulatory regions. Nat. Biotechnol. 28, 495–501 (2010).
31. R. C. Ransom et al., Mechanoresponsive stem cells acquire neural crest fate in jaw regeneration. Nature 563, 514–521 (2018).
32. V. V. Wong et al., Focal adhesion kinase links mechanical force to skin fibrosis via inflammatory signaling. Nat. Med. 18, 148–152 (2011).
33. W. Kim et al., RUNX1 is essential for mesenchymal stem cell proliferation and myofibroblast differentiation. Proc. Natl. Acad. Sci. U.S.A. 111, 16389–16394 (2014).
34. T. Xie et al., Single-cell deconvolution of fibroblast heterogeneity in mouse pulmonary fibrosis. Cell Rep. 22, 3625–3640 (2018).
35. L. Lu et al., The temporal effects of anti-TGF-beta1, 2, and 3 monoclonal antibody on wound healing and hypertrophic scar formation. J. Am. Coll. Surg. 201, 391–397 (2005).
36. G. Wernig et al., Unifying mechanism for different fibrotic diseases. Proc. Natl. Acad. Sci. U.S.A. 114, 4757–4762 (2017).