Image-seq: spatially resolved single-cell sequencing guided by in situ and in vivo imaging

Tissue function depends on cellular organization. While the properties of individual cells are increasingly being deciphered using powerful single-cell sequencing technologies, understanding their spatial organization and temporal evolution remains a major challenge. Here, we present Image-seq, a technology that provides single-cell transcriptional data on cells that are isolated from specific spatial locations under image guidance, thus preserving the spatial information of the target cells. It is compatible with in situ and in vivo imaging and can document the temporal and dynamic history of the cells being analyzed. Cell samples are isolated from intact tissue and processed with state-of-the-art library preparation protocols. The technique therefore combines spatial information with highly sensitive RNA sequencing readouts from individual, intact cells. We have used both high-throughput, droplet-based sequencing as well as SMARTseq-v4 library preparation to demonstrate its application to bone marrow and leukemia biology. We discovered that DPP4 is a highly upregulated gene during early progression of acute myeloid leukemia and that it marks a more proliferative subpopulation that is confined to specific bone marrow microenvironments. Furthermore, the ability of Image-seq to isolate viable, intact cells should make it compatible with a range of downstream single-cell analysis tools including multi-omics protocols.

Spatial transcriptomics is a rapidly advancing field that encompasses a range of different technologies capable of spatially resolved gene expression analysis. In contrast to single-cell RNA sequencing (scRNA-seq), which provides deep insights into heterogeneities within cell populations but does not preserve the spatial relationships between individual cells, techniques such as MERFISH, seqFISH and Slide-seq can link these heterogeneities to differences in spatial composition and cellular proximity. However, apart from Niche-seq, they...
have been applied only in vitro or they have relied on the generation of tissue sections1,2,3,4, which confines their applicability to tissues that are easily sectioned. They map the gene expression profiles onto two-dimensional (2D) images, and extrapolation to three-dimensional (3D) architecture of intact tissues is still limited. For example, 3D expression profiles and cell segmentation have been demonstrated in tissue sections5,6,7, but the section thickness is limited by messenger RNA probe diffusion. In addition, tissue sections can provide only static images, necessitating the use of indirect methods such as pseudo-time analysis to infer cellular trajectories over time8,9, and to our knowledge none of the currently available spatial transcriptomics technologies has been combined with in vivo imaging.

Here, we present Image-seq, a new platform that enables image-guided cell isolation for scRNA-seq. The core of the Image-seq platform is a multiphoton microscope with two optical paths, one for imaging and one for laser micromachining that creates an access channel in tissue through which a micropipette is brought to the target location to aspirate cells under image guidance. Because it captures viable cells it can be combined with state-of-the-art library preparation protocols, leading to higher mRNA detection efficiencies and broader transcript coverage than other spatial sequencing technologies. In addition, standard computational tools can be used for data analysis.

We use Image-seq to study the bone marrow, the primary site of hematopoiensis, where hematopoietic stem cells give rise to all of the blood cells in the body. The bone marrow is also the site where malignant cells can either originate (such as leukemia) or preferentially metastasize to (such as prostate or breast cancer). Although the functional organization of hematopoietic cells in the bone marrow has been extensively characterized10, their 3D spatial organization has been difficult to assess due to their complexity and location deep inside the bone matrix11. To demonstrate Image-seq’s versatility and high sensitivity, we used high-throughput, droplet-based sequencing with the 10x Chromium platform to profile bone marrow hematopoietic cells, as well as the SMARTseq-v4 protocol to profile rare (<0.01% leukemic burden) acute myeloid leukemia (AML) cells and bone marrow stromal cells. Specifically, we tracked AML progression using intravital microscopy, and observed pronounced spatial heterogeneity in the earliest stage of expansion from single AML cells seeded in the bone marrow. This highlighted the need to capture these cells from distinct bone marrow locations where they either proliferate or remain quiescent, to identify the factors that regulate leukemia progression in the earliest stage of the disease development. Using this approach we identified DPP4 (dipeptidyl peptidase 4) as a key upregulated gene in AML cells that are easily sectioned. They map the gene expression profiles onto two-dimensional (2D) images, and extrapolation to three-dimensional (3D) architecture of intact tissues is still limited. For example, 3D expression profiles and cell segmentation have been demonstrated in tissue sections5,6,7, but the section thickness is limited by messenger RNA probe diffusion. In addition, tissue sections can provide only static images, necessitating the use of indirect methods such as pseudo-time analysis to infer cellular trajectories over time8,9, and to our knowledge none of the currently available spatial transcriptomics technologies has been combined with in vivo imaging.

In our implementation the imaging arm and the ablation arm are powered by a single femtosecond fiber laser operating at 5 MHz repetition frequency. This repetition frequency enables full-field (500 × 500 pixels) image acquisition at 15 frames per second (half the video rate) with a single laser pulse per pixel. However, it is also possible to integrate an ablation capability into an existing confocal or multiphoton microscope by adding an ablation laser that fulfills the requirements described above. A detailed optical design is shown in Extended Data Fig. 1. The output of a 1,550 nm femtosecond fiber laser is frequency doubled to 775 nm and used for both imaging and plasma-mediated laser ablation by adjusting the pulse energy between 1 and 20 nJ. An additional imaging wavelength that is tunable between 940 and 980 nm is provided through soliton generation in a photonic crystal fiber12. A revolving polygon and scanning galvano-mirror, conjugated to one another and the back aperture of the objective lens, steer the ablation and imaging beams across the field of view in the imaging plane (Fig. 1a). A variable aperture at the intermediate image plane controls the 2D ablation geometry along the x and y dimensions (Fig. 1b) while the sample is translated along the z dimension to either ablate or image a 3D tissue volume. By etching through bone a small opening (~50 µm × 100 µm) is created, and a micromanipulator is used to insert a fluorescein-coated micropipette through the opening at a 20° angle into the bone marrow with micrometer spatial precision (Fig. 1b). Prior to making the opening, a larger area of bone is thinned (~200 × 300 µm) to improve micropipette accessibility and imaging depth. This step takes ~3 min. Target bone marrow cells are aspirated into the micropipette by suction with an Air Syringe, with the displaced volume controlling the total number of aspirated cells (from a few to a few thousand). With this design we were able to aspirate bone marrow cells from freshly collected bone samples as well as from live animals while tracking the process in real time (Supplementary Videos 1–3). Figure 1c shows the bone marrow aspirate containing the target cell from the in vivo image after transfer to a glass slide and visualization with a wide-field fluorescence microscope. Using this strategy, recovery of a single, viable target cell has an efficiency of ~70%.

Similarity of whole bone marrow and micropipette sample composition. To further demonstrate our instrument’s ability to isolate live cells from defined anatomic regions of murine bone marrow, we used it to isolate bone marrow samples from β-actin-green fluorescent protein (β-actin-GFP) mice. Figure 2a shows a × 8 mm, tiled, maximum intensity projection image of the calvarium bone marrow, along with a region of interest from which cells were aspirated both before and after cell extraction. To minimize sample loss we eliminated post-extraction sample processing by performing transcardial perfusion prior to cell isolation. This reduced the amount of red blood cells in our samples, and we directly generated a single-cell suspension by expelling samples from the micropipette. There is a notable variation in the CD71 Ter119+ cell compartment that we attribute to slight differences in perfusion quality (Extended Data Fig. 2a). A total of nine calvarial bone marrow micropipette samples and four tibia bone marrow micropipette samples, containing an average of 2,000 cells each, were isolated from a total of five mice, yielding an average viability of 94% and 92%, respectively (Fig. 2b). A 3D visualization of a representative volume of calvarial bone marrow before and after cell extraction is shown in Fig. 2c. Similarly, Extended Data Fig. 3a shows a representative volume of tibia bone marrow both before and after cell extraction. Both the view from above the bone surface and that achieved by rotation of 180° around the x axis are shown. We additionally analyzed extracted cell samples for their cell surface marker expression by flow cytometry and were able to detect the major hematopoietic cell populations in our
pooled micropipette samples with similar frequency to those in whole bone marrow control samples (Fig. 2d and Extended Data Figs. 2a,3b). Generally, the cell proportions were in good agreement, although we did observe statistically significant differences between tibia and calvarium bone marrow for the B-cell and pro-pro-B-cell populations. We also observed significant differences between the micropipette and whole bone marrow preparations for macrophage, pre-B-cell and pro-B-cell populations. It is unclear whether the differences between micropipette and whole bone marrow preparations are due to technical effects or the fact that we typically collected micropipette samples from within ~80 µm of the endosteal surface.

By perfusing with enzymatic digestion buffer after the initial perfusion (to reduce the number of red blood cells) and incubating the bone sample for ~20 min (see Methods for details), we were able to establish a protocol for the isolation of stromal cells. Figure 2e shows a maximum intensity projection image of a volume of CXCL12 (C-X-C motif chemokine ligand 12)-DsRed-expressing bone marrow cells both before and after cell extraction. Extended Data Fig. 2b shows the gating strategy used for quantification of CD45−CXCL12+ stromal cells in our micropipette samples: we detected 3–4 cells in a typical aspirate. A single-cell suspension is generated by expelling the sample from the micropipette and into a tube that is transferred directly to the 10x Chromium chip for high-throughput characterization of the entire cell sample. Alternatively, samples can be stained and individual cell populations sorted into wells by flow cytometry for subsequent library preparation and sequencing using the SMARTseq-v4 protocol.

To validate the technology we collected 11 Image-seq samples from a total of n = 5 C57Bl6 mice, along with three whole calvarium bone marrow (WCBM) samples (n = 3 mice) for single-cell isolation and library preparation using the 10x Chromium platform (see Methods for details). After sequencing and read alignment we used the Conos package in R (ref. 25) to integrate our multiple scRNA-seq datasets and align them with other public scRNA-seq data22. We used Leiden clustering to determine joint cell clusters across the entire dataset, identifying most of the major hematopoietic cell populations, and visualizing them by UMAP (uniform manifold approximation and projection) embedding (Fig. 3b,c). The population at the plot center expresses multi-lineage (refs. 22–25), as well as genes associated with granulocytic and monocytic priming (refs. 22–25) and with megakaryocytic priming (refs. 22–25), and was therefore termed ‘diverse progenitors’. From here, lymphoid (marked by expression of Vpreb1-3, Rag1,2, Cd19, Dntt, Mzb1, Ebf1 (refs. 22–25)), erythroid (marked by expression of Car1-2, Hemgn, Cte, Cpx, Smim1 (refs. 22–25)), granulocytic (Cebpe, Fcnb, S100a8, S100a9, Camp22–25) and monocytic (Irf8, Klf1, F13a1, S100a4, S100a10, Ly6c2 (refs. 22–25)) lineages branch off in different directions.
**Fig. 2 | Characterization of micropipette samples.**

**a.** Tiled, maximum intensity projection image of a 4 × 8 mm region of skull bone marrow (tiles shown as magenta squares on the schematic of the skull below the image), obtained by stitching together individual microscope fields of view. Green, β-actin-GFP-positive bone marrow cells; gray, second-harmonic generation (SHG) bone signal. White square: region of interest that is shown in the insets before and after cell isolation. Scale bars: main image, 1 mm; insets, 100 µm. Tiled calvarium bone marrow images were obtained from three independent mice.

**b.** Average cell viability for the calvarium (MC), from a total of nine micropipette samples in n = 3 mice. Cell viability for the tibia (MT), from four micropipette samples in n = 2 mice.

**c.** 3D visualization of a target location before and after cell aspiration. The bottom panel is obtained by rotating the top panel by 180° around the x-axis. Isolated cell volume is marked by an orange outline, and the extracted cell volume (diameter ~80 µm, height 50 µm) contained in a typical micropipette sample is outlined by an orange cylinder.

**d.** Averaged cell proportions from flow cytometry analysis of 15 calvarial micropipette (M) and 8 whole calvarial bone marrow (W) preparations in a total of n = 10 mice, and 14 tibia micropipette (M) and 5 whole tibia bone marrow (W) preparations in a total of n = 7 mice. Cells were classified based on their cell surface properties as T cells: Ter119−CD71−CD11b−CD3−; mature B cells: Ter119−CD71−CD3−CD11b−B220−CD19−cKit−IgM−; pro-B cells: Ter119−CD71−CD3−CD11b−B220−CD19−IgM−cKit−; pre-B cells: Ter119−CD71−CD3−CD11b−B220−CD19−IgM+cKit−; Pre-Pro-B cells: Ter119−CD71−CD3−CD11b−B220−IgM−; macrophages: Ter119−CD71−CD3−CD11b−Ly6C−F4/80+; inflammatory monocytes: Ter119−CD71−CD3−CD11b−F4/80−Ly6C−; patrolling monocytes: Ter119−CD71−CD3−CD11b−F4/80−Ly6C+Gr1−; granulocytes: Ter119−CD71−CD3−CD11b−F4/80−Ly6C−Gr1+. See Extended Data Fig. 2 for the gating strategy.

**e.** Maximum intensity projection image of a bone marrow region before and after stromal cell aspiration with the micropipette. Red, CXCL12-DsRed bone marrow cells; gray, SHG bone signal. Scale bar, 100 µm. Fifty biological replicates of stromal cell extraction were performed in 14 independent mice.
1) Bone marrow imaging

2) Transcardial perfusion

3) Laser ablation and image-guided cell aspiration

4) Single-cell suspension

5a) 10× Genomics

5b) Fluorescence-activated cell sorting, SMARTseq-v4

Bone SHG, AML, CXCL12

b) Neutrophil Image-seq

c) Neutrophil WCBM

d) Heatmap showing gene expression levels across different cell types.

Table 1: Percent of expressed genes and number of expressed genes.

| Gene          | Image-seq | WCBM |
|---------------|-----------|------|
| Il7r          | 25        | 10   |
| Cd3g          | 50        | 25   |
| Thy1          | 75        | 50   |
| Ccl5          | 100       | 75   |

**Figure 1:**

a) Diagram showing the steps of cell isolation and analysis.

b) Heatmap of gene expression levels in different cell types.

f) Statistical significance of gene expression levels.

g) Imaging of cells with high and low burden.

h) Magnified images showing high and low burden of cells.
Dot plots of selected marker genes for each cell population are shown in Fig. 3e. A list of differentially expressed genes in each cluster is given in Supplementary Table 1. Here, we pooled 11 Image-seq samples to characterize the distribution of cell populations in the limit of a large number of samples (for which we would expect the same distribution as for WCBM samples if there were no technical differences between the two sampling strategies).

As expected for sampling small volumes from different locations, we observed significant sample-to-sample variations (a detailed breakdown is given in Extended Data Fig. 4b). However, we did not observe statistically significant differences in the proportion of any cell populations when comparing Image-seq (Fig. 3b) and WCBM (Fig. 3c) preparations in aggregated form (Extended Data Fig. 4c). We further examined the transcriptional similarity of the major cell types from WCBM and Image-seq samples. Correlation coefficients of average gene expression levels between WCBM and Image-seq samples (Fig. 3d) show a strong correlation between individual cell populations collected using the two methodologies. Gene ontology analysis of differentially expressed genes between Image-seq and WCBM samples for the most part showed very general terms that were not associated with cell damage or cell stress (Extended Data Fig. 4d, e), although there was a notable downregulation of ribosomal terms in Image-seq samples, which could indicate lower cell stress and could be related to the overall higher number of unique molecular identifiers and genes per cell that we observed in the Image-seq samples (Fig. 3f). We additionally assessed the percentage of cells that passed quality control using the two experimental strategies (Extended Data Fig. 5a) and found no statistically significant differences. We plotted the distribution of unique molecular identifiers and genes per cell for each sample that we collected (Extended Data Fig. 5b), along with a table that summarizes the overall number of cells in each sample and the ones that passed quality control (Extended Data Fig. 5c). Again, although some variations were observed between samples, the data as a whole led us to conclude that Image-seq produces scRNA-seq data with a quality that is comparable to or exceeds that of conventional cell collection protocols.

To demonstrate the technology’s ability to obtain spatially resolved single-cell transcriptional data of stromal cells (Extended Data Fig. 6), we isolated 43 CD45$^-$/CXCL12$^+$ cells by micropipette and flow cytometry. After library preparation and sequencing using SMARTseq-v4, we were able to construct a shared nearest neighbors graph and UMAP embedding of these cells (Extended Data Fig. 6b), and identified two separate clusters. We annotated these as mesenchymal stromal and endothelial cells (Extended Data Fig. 6d) based on literature marker genes. A heatmap of differentially expressed genes within these clusters is shown in Extended Data Fig. 6c.

To further validate the spatial selectivity of our technology we imaged an MLL-AF9 mouse model of AML in which leukemia cells express GFP under the ubiquitin promoter at day 10 after transplantation. As shown previously, we found regions of high leukemic burden to be interspersed with regions of low leukemic burden (Fig. 3g). We used Image-seq to isolate cells from a high-burden region and a low-burden region, and processed them separately for scRNA-seq using the 10x Chromium protocol (see Methods for details). UMAP embeddings of both samples show that the high-burden sample was composed primarily of AML cells, whereas the low-burden sample did not include leukemia cells (Fig. 3h). Furthermore, strong levels of Gfp expression were detected in the AML cell cluster from the high-burden sample (Fig. 3g), highlighting the technique’s spatial selectivity and high sensitivity.

**Image-seq analysis of early leukemia expansion.** We next turned our focus to early leukemia progression in a Hoxa9/Meis1-Ubiquitin-c-GFP (HA9MI) mouse model of AML, performing intravital imaging of the calvarium bone marrow between 1 and 3 days after transplanting $3 \times 10^6$ cells into non-irradiated recipients (Fig. 4a). We found that starting from day 3, the number of leukemia cells per bone marrow cavity (concaendoendostem) rapidly increases (Fig. 4b), and chose this timepoint to study the mechanisms of early leukemia expansion. We used longitudinal imaging of the same cavities in the same animals to quantify proliferation dynamics (see Fig. 4c for examples of longitudinal images) and classify cells as either proliferating (P, average of 30 cells per cavity on day 3), intermediate (IM, average of 13 cells per cavity on day 3) or non-proliferating (NP, average of 2 cells per cavity on day 3), as shown in Fig. 4c, where the median and quartiles of the distributions are given (see Methods for details). We then used the 99.9% confidence interval of AML cells per cavity on day 3 to identify sites for cell extraction and sequencing. Prior to the sequencing experiments we quantified label dilution on day 3 after transplantation with CellTracker-labeled AML cells. The percentage of CellTracker-positive cells per bone marrow cavity decreased as the overall number of cells increased (Extended Data Fig. 7a), further verifying our imaging-based classification. Surprisingly, these proliferative phenotypes were distributed throughout the calvarium bone marrow in a spatially heterogeneous manner (Fig. 4d), further underlining the need for a spatially resolved analysis by scRNA-seq.

We identified P, NP and IM AML cells by imaging and aspirated them by micropipette using our Image-seq platform. Subsequently, we separated the AML cells from the -100–400 surrounding hematopoietic cells by sorting them into individual wells of a 96-well plate by flow cytometry and gating for GFP (see Supplementary Fig. 1 for examples of the gating strategy used to isolate an NP, as well as several P cells, from a single micropipette sample). By choosing this strategy (shown schematically in Fig. 3a as steps 1, 2, 3, 4 and 5b), we optimized the yield of the rare AML cells and avoided the ~30–40% cell loss that occurs during cell encapsulation using the 10x Genomics platform. In total, we collected ~40 cells from four different locations in the P category, ~40 cells from ~30 different locations in the NP category, as well as ~20 cells from three different locations in the IM category from $n = 11$ mice. Experimental throughput was limited by the extremely rare
**Fig. 4** | Early leukemia progression by Image-seq. **a**, In vivo imaging timepoints. IVM, intravital microscopy. **b**, Cell counts per bone marrow cavity on day 0 (n = 143 cavities, n = 6 independent mice), day 1 (n = 103 cavities, n = 6 independent mice) and day 3 (n = 132 cavities, n = 7 independent mice) after transplantation. Error bars represent the standard deviation and are centered at the arithmetic mean. The 99.9% confidence interval for proliferating (P) cells is shown in red, that for intermediate (IM) cells in blue and that for non-proliferating (NP) cells in green. **c**, Number of cells per cavity on day 3 plotted for P, IM, and NP regions. The dashed line represents the median and the dotted line, the quartiles in each distribution. Statistical significance was determined using Welch’s ANOVA test and the adjusted P values were determined using Dunnett’s multiple comparisons test. NP, 10 biological replicates, n = 3 independent mice; IM, 32 biological replicates, n = 3 independent mice; P, 107 biological replicates, n = 3 independent mice. **d**, Tiled, maximum intensity projection image of leukemic bone marrow on day 3 after transplantation. Red square and inset, P cells; blue square and inset, IM cells; green square and inset, NP cell (marked by white arrow). Scale bars: main image, 500 µm; insets, 100 µm. Bone, bone SHG. Imaging of the entire calvarium bone marrow at day 3 was performed in 17 independent mice, including 122 biological replicates with P, 35 biological replicates with IM, and 79 biological replicates with NP cells. **e**, Images of proliferating (top, red border; scale bars, 100 µm) and non-proliferating (bottom, green border; scale bars, 50 µm) AML cells imaged longitudinally in the same animal and bone cavity on day 1 and 3 after transplantation. Bone SHG signal is shown in grey, AML-GFP signal is shown in green. The same depth along the z dimension is shown (Zrel = 0) on both days, along with a maximum intensity projection for P cells. For the NP cell a single Z slice (Zrel = 28 µm) shows that the AML cell has moved deeper into the bone marrow on day 3. A total of 107 biological replicates with P, 10 biological replicates with NP and 32 biological replicates with IM cells (n = 3 independent mice, respectively) were identified by longitudinal imaging (definition of P, NP, IM based on the fold-change difference between day 1 and day 3). **f**, Distribution of the cell cycle in the three imaging phenotypes (P value obtained using a two-sided Fisher’s exact test). **g**, Bar-plot representing the fraction of P, NP, and IM cells in each AML cluster. AP1, AML-AP1; GMP, AML-GMP; Mono, AML-mono (P value obtained using a two-sided Fisher’s exact test). **h**, Leiden clustering of 84 AML cells isolated by Image-seq after regression of cell cycle genes using Seurat, visualized as UMAP embedding. **i**, Heatmap of scaled normalized expression for the 10 most strongly upregulated genes (based on logFoldChange) in each AML cluster.
NP cells, which made up only 0.7% of AML cells on day 3, with the overall leukemic burden already <0.01%. To minimize the issue of dropout that is often observed in scRNA-seq data, we used the SMART-seq v4 protocol for reverse transcription and library construction. Because of the SMART-seq protocol’s unique chemistry, the number of genes and transcripts detected per cell are higher than for any other scRNA-seq technology (Extended Data Fig. 5d and ref. 29). Used in combination with the SMART-seq v4 protocol, Image-seq therefore has the highest sensitivity and transcript coverage of any currently available spatial sequencing technology, with an average of 3,314,087 total mapped reads or 8,053 genes per cell. Of the 97 cells that were sequenced, a total of 84 cells passed quality control.

After sequencing and read alignment we identified cells that were in the GO phase by adapting methodologies from refs. 30,31 and performed hierarchical clustering based on cell cycle genes, identifying three separate clusters (shown in the heatmap of Extended Data Fig. 7c). Cluster 1 was enriched in genes related to the S and G1/S phases, cluster 3 was enriched in genes related to the G2 and G2/M phases, and cluster 2 lacked expression of cycle genes and was therefore classified as quiescent cells (GO phase). Interestingly, the distribution of cell cycle in the P and IM populations was very similar, with the largest proportion of cells in the G1/S and S phases, and the second largest proportion in G2 and G2/M (Fig. 4f). For the NP population, however, the major contribution came from the GO phase, and the second largest from the G1/S and S phases. These results highlight the validity of our imaging-based classification, as well as our technology’s ability to distinguish subtle phenotypic differences through imaging.

Next, AML cells were sub-clustered after regressing out cell cycle genes using Seurat32, with the resulting UMAP embedding shown in Fig. 4h. Three major clusters were observed, and the heatmap in Fig. 4i shows the top 10 differentially expressed genes in each of them. The dark-green cluster (AML-mono) was enriched in progenitor cell markers of the monocytic lineage such as Cts5, Cun3, Ms4a4c, Pira2 and Lsp1 (refs. 3,26,33), as well as high B and Fli3 (Fig. 4i, Extended Data Fig. 8f and Supplementary Table 2), which have been associated with stronger leukemia-initiating capacity in Hoxa9/Meis1AML14. AML-GMP expressed granulocyte–monocyte progenitor markers such as Mpo, Elane, Pcss7 and Csg2(30,33), along with Gstm1, which has been associated with quiescent, therapy-resistant, leukemia-initiating cells in AML30,32. AML-API expressed Prtn3, which is a marker of the monocytic lineage22, along with Jun and Fos, which comprise the API transcription complex22, as well as Dusp1, which are all part of the MAP-kinase signaling pathway (https://www.kegg.jp/pathway/mm04010) and are associated with cell proliferation. Cluster AML-MEP consisted primarily of cells that were classified as NP by imaging, AML-API consisted primarily of P cells, and AML-mono consisted primarily of P and IM cells (a detailed breakdown is given in Fig. 4g).

Increased proliferation of DPP4-positive cells in mouse and human AML. With the goal of identifying signals that trigger the exit of leukemia-initiating cells from the quiescent state, we investigated the differential expression of genes between P and NP cells (Fig. 5a and Supplementary Table 3) without prior regression of cell cycle genes. GO analysis identified several terms related to the negative regulation of immune response (Extended Data Fig. 9g). A list of cell cycle-related genes that were differentially expressed is given in Supplementary Table 4. The most strongly differentially expressed gene, both when comparing P with NP and the entire population of P + IM with NP cells (Supplementary Table 5), was Dpp4. DPP4 or CD26 is a transmembrane protein that functions as a serine protease, selectively cleaving the amino-terminal, penultimate proline, or alanine of proteins34–36. Among its many substrates are granulocyte–macrophage colony-stimulating factor, CXCL12, granulocyte colony-stimulating factor, interleukin 3 and erythropoietin, which are known to enhance proliferation, survival, chemotaxis, homing and engraftment of hematopoietic stem and progenitor cells14–16. DPP4 has also been shown to cleave CCL2, CCL3, CXCL6, CXCL9 and CXCL10, which play a role in decreasing the survival and proliferation of hematopoietic stem and progenitor cells. Several studies have implicated DPP4 as a leukemic stem cell marker in chronic myeloid leukemia34–36, and recent studies of human AML have linked increased DPP4 expression to a poorer overall survival37 and have associated AML-derived DPP4 with the suppression of normal hematopoietic progenitor cell proliferation38,39. However, DPP4 was previously not known to play a role in early AML proliferation.

To validate DPP4 expression we assessed its protein expression using flow cytometry and found that it increases as a function of disease progression, with a maximum stable expression at ~50% by week 4 (Fig. 5b). Although DPP4 expression was too low to be detected using flow cytometry on day 3, it could be detected using in vivo imaging, and we found a statistically significant difference when DPP4 antibody labeling was compared in Pversus NP cells whereas we did not observe this in the corresponding isotype control (Fig. 5c,d and Extended Data Fig. 7d). Next, we analyzed the cell cycle in DPP4-positive versus DPP4-negative AML cells using Ki-67 staining (see Extended Data Fig. 8b for gating strategy), which confirmed that DPP4-positive cells have a higher cycling rate (Fig. 5e and Extended Data Fig. 8a). To elucidate whether this phenotype is more broadly observed, we studied an MLL-AF9 model of AML and found the decrease in the G1 phase and the concomitant increase in the S, G2 and M phases in DPP4-positive cells to be even more pronounced (Fig. 5f). Note that at day 13 (Extended Data Fig. 5a) the difference in cycling rate is less pronounced than at day 7, and there is no difference in the proportion of cells in the G1 phase. This is a very aggressive model and even at day 13 more than 80% of the marrow is composed of leukemia cells, suggesting that the cells are running out of space to proliferate, which is likely to reduce the cycling rate. In keeping with this, the overall...
proportion of DPP4-positive and DPP4-negative cells that are actively cycling is considerably lower on day 13 than on day 7 ($P < 0.0001$), which is likely the reason that the proportion of DPP4-positive cells reaches a maximum stable value of 50%. We additionally analyzed DPP4-correlated genes in two published, human AML datasets\textsuperscript{48,49} (see Methods for details) and observed a high overlap between the top 300 positively correlated genes. GO analysis of the overlapping gene set (Fig. 5g) showed an enrichment of terms related to cell cycle (Fig. 5h), suggesting that the phenotype is relevant to human AML.

Because DPP4 expression was confined mainly to the AML-mono cluster, which expressed high levels of $Iggb7$, $Flt3$ and $Cd48$ (Extended Data Fig. 8f), we analyzed their cell surface expression using flow cytometry. We found a strong increase in $Iggb7$ expression, as well as a moderate increase in $Flt3$ and $Cd48$ for cells that had high DPP4

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\caption{Comparison of DPP4-positive and DPP4-negative cells.}
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\caption{GO analysis of the overlapping gene set.}
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\caption{Flow cytometry analysis of cell surface expression.}
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\caption{Expression of $Iggb7$, $Flt3$, and $Cd48$ in high DPP4 cells.}
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expression as compared with negative and intermediate DPP4 expression (Extended Data Fig. 8c,e), suggesting that this combination of cell surface markers can be used to identify cells belonging to AML-mon. Interestingly, overexpression of Flt3 has been associated with poor prognosis in AML10, and combined expression of Itgb7 and Flt3 has been used to identify leukemic stem cells in Hoxa9/Meis1 AML11. We investigated DPP4 expression in healthy myeloid precursors, as well as lineage-specified cells, using flow cytometry (Extended Data Fig. 9a). We found higher DPP4 levels on cells that are more immature (with the exception of macrophages and lymphoid cells), which supports the notion that DPP4 expression is not due to progression along the myeloid differentiation trajectory in AML cells, or associated with priming towards a specific lineage.

To assess leukemia-initiating capacity, we isolated 1,000 DPP4high and DPP4neg AML cells 3 weeks after transplantation using flow cytometry. We transfected them into fresh, non-irradiated recipients and compared leukemic burden and the proportion of DPP4-positive and DPP4-negative cells at day 10 (Extended Data Fig. 8d). Interestingly, a subset of DPP4-negative cells converted to being DPP4 positive and vice versa, suggesting that DPP4 expression could be induced by the microenvironment.

**Activation of DPP4 expression by the microenvironment.** While observing DPP4 cell surface expression to be highly stable and reproducible in vivo, we were not able to detect DPP4 on AML cells cultured in vitro (Extended Data Fig. 10a). We were also unable to detect DPP4 by intracellular staining or quantitative polymerase chain reaction (qPCR) (Extended Data Fig. 10b,c). In addition, we found that although DPP4-positive HA9M1 cells expressed Dpp4 at comparable levels to splenic T cells immediately following removal from the bone marrow environment (day 0 qPCR results in Extended Data Fig. 8d), they subsequently lost all Dpp4 expression at the transcriptional level after 4 days of in vitro cell culture. Remarkably, co-culture of AML cells with bone marrow stromal and osteoblast precursor cells activated DPP4 cell surface expression, while co-culture with other stromal cell lines did not (Fig. 6a). Combined with the observation that DPP4 expression is upregulated in specific bone marrow compartments (Fig. 5a,c), the data suggest that different niches drive the upregulation of DPP4. Given the recent finding that hematopoietic stem cell proliferation is restricted to a subset of bone marrow cavities with mixed bone remodeling activity (M-type cavities)12 we reasoned that these niches could drive proliferation of AML cells. We used the dye labeling strategy developed in ref. 10 to distinguish deposition (D-type), resorption (R-type) and mixed (M-type) bone marrow cavities by in vivo imaging, finding that P cells are exclusively located in M-type cavities (Fig. 6b).

**Discussion**

AML is an aggressive blood cancer characterized by an accumulation of immature myeloid cells in the bone marrow that are arrested in differentiation and that accumulate as immature and malignant self-renewing progenitors. Despite an initially favorable response to treatment with intensive cytotoxic chemotherapy, ~75% of patients die within 5 years of diagnosis13. Relapse is thought to be driven by the rare pool of leukemia-initiating cells that persist in the bone marrow following chemotherapy14. Arguing that an important approach to the development of new therapeutics is the targeting and exploiting of the supportive interaction as leukemia-initiating cells communicate with and seek refuge within the bone marrow microenvironment, we decided to study and identify signals that trigger the exit of leukemia-initiating cells from the quiescent state, thereby gaining mechanistic insights into disease recurrence. We further reasoned that examining the bone marrow at a stage of very low leukemic burden, comparable to the state of minimal residual disease (when the AML cells are more likely to be surrounded by normal hematopoietic and stromal cells), provides unique insights that cannot be obtained using traditional assays of relatively high leukemic burdens in which the bone marrow is crowded with malignant cells and the microenvironment is severely altered.

Here, we have shown how the combination of intravital microscopy to study the dynamics of AML disease progression, and spatially resolved scRNA-seq, provides new insights into leukemia biology. Moreover, multiphoton microscopy uniquely informs the 3D spatial context of AML cells and can be used to validate the single-cell gene expression data in a spatially resolved manner (Fig. 5e,f and Extended Data Fig. 7d), as well as characterize different microenvironments.
(Figs. 4b–e and 6b) that can be analyzed for their differential expression (Fig. 5a). We found that DPP4 marks a more proliferative phenotype both in murine and in human disease, which makes it a promising target for further investigations that could lead to new treatments.

In situ imaging of murine bone marrow has led to numerous insights into the basic biology of hematopoietic stem cells, as well as the spatial organization of bone marrow tissue. In the skull bone marrow, intravital microscopy has been used for the study of hematopoiesis, along with hematopoietic stem cell and leukemia biology. It has elucidated the temporal dynamics of these processes and tracked the association of individual cells with distinct bone marrow components in real time. Imaging alone, however, fails to provide unbiased mechanistic insight into the observed cellular dynamics and spatial organization. It is precisely such information, however, that promises to bring new insights into hematopoietic and leukemia biology and, concomitantly, the development of new therapeutics.

To date, the only spatially resolved transcriptomic study of the murine bone marrow, a tissue that remains highly challenging to section, relied on bulk transcriptomic analysis of tissue blocks that were cut from formalin-fixed bone marrow sections using laser micro-dissection. The cellular composition of individual blocks was then inferred computationally using a separate, scRNA-seq dataset. To our knowledge there have been no spatially resolved transcriptional studies of the leukemic bone marrow.

Image-seq represents a new experimental approach for integrating spatial and molecular information. Multiple contrast mechanisms can be used to visualize the procedure and reconstruct the 3D spatial position of the extracted cell sample, including autofluorescence, confocal reflectance (Fig. 1c and Supplementary Video 1) or labeling with a fluorescent membrane dye such as Di-8-ANEPPS (used in the experiments in Figs. 2d and 3). When combined with intravital microscopy, Image-seq additionally enables transcriptomic analysis on cells for which the position and history have been documented by intravital microscopy. Because of its utility both for high-throughput sequencing and in-depth profiling of rare cells, the Image-seq platform is highly versatile in its applications. We additionally expect that it can be applied to study fixed tissues and tissue sections (for example, using the 10x Genomics protocol for fixed tissues instead of the protocol we detailed here), although we recommend working with live cell samples when possible to achieve the highest sensitivities and transcript coverage. Currently there is a trade-off between spatial resolution and throughput. Although it is possible to isolate very few cells and obtain a spatial resolution of ~30 × 30 × 30 μm, or even a single cell if there is only one fluorescent cell in the isolated tissue volume (Fig. 1c) that can be sorted by flow cytometry, this comes at the cost of lowered throughput. Increasing the number of extracted cells, however, lowers spatial resolution. As we expand the technology we expect to overcome these shortcomings and refine its resolution and increase its throughput. We anticipate that Image-seq will be used to investigate a range of diseases and biological questions involving the bone marrow, along with other tissues that remain difficult to section. Furthermore, based on the technology's ability to isolate viable cells, we anticipate its applicability to spatially resolved multi-omics studies in a range of biological settings.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41592-022-01673-2.

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**Methods**

A detailed protocol of the Image-seq procedure is available in the Supplementary Information.

**Microscope**

Intravital microscopy and plasma-mediated laser ablation were performed using a custom-built multiphoton microscope. The output of a femtosecond, 1,550 nm fiber laser (Calmar Cazadero) operating at a repetition rate of 5 MHz was split into two optical paths: one was frequency doubled with a BiBO crystal (Newlight Photonics) to obtain a wavelength of 775 nm that was used for either imaging or ablation (Extended Data Fig. 1). The other was coupled into a large mode-area photonic crystal fiber (LMA-35, NKT Photonics) where, by soliton self frequency shift, radiation with a frequency tunable between 1,880 nm and 1,960 nm was generated. This was frequency doubled (BiBO, Newlight Photonics) to obtain imaging wavelengths between 940 and 980 nm. The imaging and ablation beams were spatially overlapped using a dichroic mirror (Zt 785 RDC, Chroma) and directed to the scanning optics, which consisted of a revolving polygon (36 facets, 14,400 r.p.m., x, y scan) conjugated to a scanning galvanometer mirror (15 Hz, y scan), and which were conjugated to the back aperture of the objective lens (Olympus, ×60, numerical aperture (NA) 1.0, water immersion). This served to steer the imaging and ablation beams across the microscope field of view. Typical pulse energies used for imaging were between 1 and 2 nJ and typical pulse energies for ablation were between 10 and 15 nJ. For stable plasma generation a continuous stream of PBS (ThermoFisher Scientific) was flushed across the sample at a rate of 10 ml min⁻¹ to remove any gas or debris generated during the ablation procedure. The sample (mouse) was positioned in the image plane using a micromanipulator (Sutter instruments) and translated along the z dimension to image or ablate a 3D volume of tissue. Fluorescence signals were collected after excitation with 775 nm or 980 nm laser light, using three photomultiplier tubes (PMTs). The typical configuration of the dichroic mirrors was FF705 LP, FF495 LP and FF552 LP, with filters 439/154 (blue PMT R7600-U300, Hamamatsu) for the detection of collagen second-harmonic generation signal from bone, 525/50 (green PMT R7600-U300, Hamamatsu) for the detection of AML cells and tetracycline, and 607/70 (red PMT R7600-U20, Hamamatsu) for the detection of CellTracker Red and Di-8-ANEPPS (ThermoFisher Scientific) signals. An avalanche photodiode was also installed for collection of the confocal reflectance signal (used to visualize the in vivo cell isolation procedure in Supplementary Video 1).

**Procedure for intravital imaging**

The procedure for intravital imaging of the calvarium bone marrow is described in detail elsewhere. Prior to intravital microscopy and ablation, analgesics were given (buprenorphine at 0.05–0.1 mg kg⁻¹ i.p.), mice were anesthetized using vaporized isoflurane (3–4% for induction, 1–2% for maintenance) and depth of anesthesia ensured by toe pinch. Hair around the incision site on the scalp was trimmed, and skin was made aseptic using a betadine scrub. The incision (~5 mm × 7 mm) was made using sterile surgical scissors, and the skin folded back to expose the skull bone, which was hydrated using sterile PBS. Mice were transferred to a mouse holder with integrated heating pad (37 °C), and a continuous stream of isoflurane supplied via a nose cone during anesthesia, and their spatial position recorded with respect to the bregma and lambda reference points. Mice were retro-orbitally injected with either Di-8-ANEPPS (1.9 mg kg⁻¹) or Brilliant Violet 421 anti-mouse CD31 antibody (1 mg kg⁻¹, BioLegend) to aid in the visualization of the cell isolation procedure. Anesthesia was increased to 4%, the mice were transferred to a dissection tray and were transcardially perfused as follows.

1. For the isolation of hematopoietic cells: first with an ice-cold solution of 5 μM EDTA in PBS (ThermoFisher Scientific, flow rate 5 ml min⁻¹, total volume 10 ml) and then with ice-cold PBS (ThermoFisher Scientific, no Ca or Mg, flow rate 5 ml min⁻¹, total volume 10 ml).
2. For the isolation of stromal cells: first with a 37 °C solution of 5 μM EDTA in PBS (both ThermoFisher Scientific) at a flow rate of 5 ml min⁻¹ and with a total volume of 10 ml, and then with an enzymatic digestion buffer at 37 °C (flow rate 5 ml min⁻¹, total volume 10 ml). The mice were then incubated at 37 °C for 20 min. The enzymatic digestion buffer consisted of 450 U ml⁻¹ Collagenase I (Sigma), 125 U ml⁻¹ Collagenase XI (Sigma), 60 U ml⁻¹ Hyaluronidase (Sigma) and 60 U ml⁻¹ DNase I (Sigma) in 20 ml Medium-199 (Gibco).

Mice were transferred back to the microscope and samples were sequentially isolated from positions marked for cell extraction.

**In situ imaging of tibia**

Mice were perfused first with an ice-cold solution of 5 μM EDTA in PBS (ThermoFisher Scientific, flow rate 5 ml min⁻¹, total volume 10 ml) and then with ice-cold PBS (ThermoFisher Scientific, no Ca or Mg, flow rate 5 ml min⁻¹, total volume 10 ml). Tibia were then dissected and cleaned, and the tibial bone was thinned to a thickness of ~50 μm using a razor blade. Bones were mounted onto a microscope slide by fastening a piece of modeling clay to the glass slide and gently pressing the bone onto the modeling clay. The mounted bone was then transferred to the microscope for in situ imaging.

**Image-guided cell aspiration (calvarium and tibia)**

In each location the procedure was as follows: first, a volume of bone (~40 μm × 200 μm × 300 μm) was removed using a pulse energy of 14 nJ; second, a channel with dimensions ~30 μm × 50 μm × 100 μm was created using a pulse energy of 10 nJ; third, the micropipette was inserted...
through this channel and positioned next to the target cells; and last, the
cells were aspirated using a micropipette (MBB-FP-M-20, Origio)
and transferred to an Eppendorf tube filled with 5 μl Medium-199 with
2% v/v FBS. Samples were kept on ice until they were either analyzed
flow cytometry (validation experiments), transferred to the 10x
Chromium platform, or sorted into individual wells of a 96-well plate by
flow cytometry (for library preparation by SMARTseq-v4).

The ablative procedure itself was performed at a rate of 0.25 μm per 670 ms along the z dimension, which corresponded to 10 passes per plane using the 15 frame per second imaging rate of the optical system.

Prior to the experiment, micropipettes were coated with Sig-
macote (flowed through the micropipette for 2 min at a rate of
200 μl min⁻¹) to prevent cells from adhering to the glass surface, as
well as with Qtracker 655 vascular labels (5 μl were pipetted up and
down several times) to fluorescently coat the pipette and aid with
visualization.

Collection of whole bone marrow preparations
Calvaria were dissected and cut into smaller pieces. Tibia were dissected
and cleaned. To aid in the release of the bone marrow, calvaria bone
fragments or whole tibia bones were gently crushed in Medium-199
(Gibco) supplemented with 2% FBS (Gibco). The resulting cell suspen-
sion was subsequently passed over a 70 μm cell strainer (BD Falcon).

Cell lines
Syngeneic leukemia model. The HoxA9/Meis1 and MLL-AF9 mod-
elms have been described in detail elsewhere. In brief, the HoxA9/
Meis1 AML cell line was generated by retroviral transduction with an
MSCV-HoxA9-RES-Meis construct (originally designed by G. Sauvageau)
to bone marrow mononuclear cells from a mouse expressing GFP under
the control of the ubiquitin, and luciferase under the control of the β-actin
promoter. The MLL-AF9 cell line used for the cell cycle experiments
was generated by collecting bone marrow from a 5-fluorouracil-treated
Cas9-GFP mouse, followed by two consecutive transfections with retro-
viral MLL-AF9. For both models the cells were transplanted into irradiated
recipients, collected from terminally ill animals and re-transplanted
into a second set of irradiated recipients, from which GFP-expressing
cells were collected close to the disease endpoint. These cells were cul-
tured in RPMI 1640 (Gibco) supplemented with 10% FBS (ThermoFisher
Scientific), 100 IU ml⁻¹ penicillin (Corning), 100 mg ml⁻¹ streptomycin
(Corning), 5 ng ml⁻¹ interleukin 3 (IL-3, Peprotech), as well as 100 ng ml⁻¹
stem cell factor (SCF, Peprotech) for the HoxA9/Meis1, and 20 ng ml⁻¹
SCF and 10 ng ml⁻¹ IL-6 (both from Peprotech) for the MLL-AF9. Recipient
female mice (10–12 weeks old) were injected with 3 × 10⁶ cells in 200 μl
PBS (HoxA9/Meis1) and 1 × 10⁶ cells in 200 μl PBS (MLL-AF9).

The MLL-AF9 model used for the experiments in Fig. 3 was gener-
at by crossing MLL-AF9 knock-in mice with mice expressing GFP
under the control of the ubiquitin, and luciferase under the control of
the β-actin promoter. The bone marrow from a terminally ill male
mouse was collected and cultured in vitro using RPMI 1640 (Gibco)
supplemented with 10% FBS (Gibco), 100 IU ml⁻¹ penicillin, 100 mg ml⁻¹
streptomycin (both from Corning), 20 ng ml⁻¹ recombinant mouse SCF
(rmSCF), 10 ng ml⁻¹ recombinant mouse IL-3 (rmIL-3) and 10 ng ml⁻¹
IL-6 (all from R&D Systems). Recipient male mice (8–10 weeks old)
were injected with 1 × 10⁶ cells in 100 μl saline.

For transplantation of DPP4-negative and DPP4-positive HoxA9/
Meis1 leukemia, cells were isolated from the long bones and vertebral
column 3 weeks after transplantation. The marrow underwent density
gradient centrifugation (Ficoll-Paque Plus, Cytiva Life Sciences) at
400 × g for 25 min at room temperature with no brake. The mononuclear
layer was isolated and subsequently blocked in PBS with 2% FBS and
murine Fc Block (BD Biosciences, dilution 1:50). Following this, the
samples were stained with CD45-APC/Cy7 (BD Biosciences, dilu-
tion 1:100) and DPP4-PE (Biolegend, dilution 1:20). To exclude dead
cells, samples were incubated with 7-aminoactinimycin D (7AAD,
0.25 μg, BD Biosciences) and then 7AAD’ GFP CD45’DPP4‘ and
7AAD’ GFP CD45’DPP4‘ cells were sorted into separate tubes. A total of
1,000 cells of each phenotype were injected into 10–12-week-old recipient mice. Leukemia burden and DPP4 expression were assessed
10 days post-transplantation.

Simian virus 40 immortalized bone marrow stroma. Total bone
marrow cells were isolated from the femurs and tibia of B6.129(B6N)-Gt(ROSA)26Sortm1(CAG-cas9*,-EGFP)Fezh/J mice (Jackson Laboratories).
The bones were crushed in PBS (ThermoFisher) with 2% FBS (Gibco)
and the released marrow was filtered over a 40 μm strainer. Mononuclear
cells were collected by density gradient centrifugation (Ficoll-Paque
Plus, Cytiva Life Sciences). A total of 20 × 10⁶ mononuclear cells were
put into culture with Alpha-MEM (Gibco) supplemented with 20% FBS
(Gibco) and 1% Penicillin–Streptomycin (Gibco) in 150 mm dishes.
Non-adherent cells were discarded around day 5 and the media changed
5–7 days for approximately 3 weeks. At this point, colonies of large adherent fibroblasts were apparent. The cells were detached
from the dishes with Trypsin-EDTA (Gibco), counted, and 50,000 cells
seeded into two wells of a six-well plate. The following day, one well of
cells was transduced with lentivirus in the presence of 8 μg ml⁻¹ poly-
brene (Millipore, Sigma) to deliver the simian virus 40 (SV40) small
and large T antigen. The plasmid pLenti CMV/TO SV40 small + Large T
(w612-1) was a gift from Eric Campeau (Addgene plasmid 22298; http://
addgene.org/22298; RRID: Addgene_22298). Despite the lack of a
selectable marker, the transduced cells divided much more rapidly
than the non-transduced primary stroma, and over the course of 2–3
passages, established an SV40 immortalized stromal cell line.

Isolation and in vitro stimulation of T cells. Spleens were collected from
C57Bl/6j mice and a single-cell suspension was obtained by mechan-
ical dissociation of the tissue over a 70 μm cell strainer (BD Falcon) in
RPMI 1640 (ThermoFisher) supplemented with 10% FBS (Gibco) and 1%
Penicillin–Streptomycin (Gibco). Red cells were lysed using ACK Lysing
Buffer (Quality Biological) followed by removal of non-T-cell spleenocytes
through magnetic depletion. In brief, the cell suspension was adjusted to
10⁶ cells ml⁻¹ and incubated with biotinylated antibodies directed against
B220, CD19, Ter119, NK1.1, Cd11b and Gr1 (all from Biolegend, dilution of
1:100, Supplementary Table 6) at a concentration of 5 μg ml⁻¹ for 10 min
at room temperature on an orbital shaker. This was followed by the addition
of 25 μl ml⁻¹ streptavidin-conjugated Rapidspheres (STEM Cell Technolog-
ies) and an additional 5 min of incubation at room temperature on an
orbital shaker. The samples were placed in an EasySep magnet (STEM Cell
Technologies) for 5 min and the purified T cells were subsequently
decanted into a fresh tube. Purity of the isolated T cells was confirmed to
be >95% using FACS analysis. To induce T-cell proliferation, the isolated
splenic T cells were plated with the T-cell activator Dynabeads CD3/CD8
(ThermoFisher) in a 1:1 ratio in leukemia cell line medium: RPMI
1640 (Gibco) supplemented with 10% FBS (Gibco), 100 μl ml⁻¹ penicillin
(Gibco), 100 μl ml⁻¹ streptomycin (Gibco) and 5 ng ml⁻¹ IL-3 (Peprotech),
as well as 100 ng ml⁻¹ SCF (Peprotech).

Flow cytometry
Whole bone marrow and micropipette samples. Whole bone mar-
row and micropipette samples from calvarium and tibia were blocked
with anti-mouse Fc Block (BD Biosciences, dilution 1:50) for 10 min at
4 °C. The cells were thereafter stained with blood cell lineage cocktail
(Supplementary Table 6) for 30 min at 4 °C. For detection of dead
cells 7AAD (BD Biosciences, 0.25 μg) was added to the sample prior to
analysis. Flow cytometry was performed on a BD FACS Aria III sorter
(BD Biosciences) and all data were analyzed using FlowJo (Treestar).

Flow sorting of AML cells for SMARTseq-v4. Prior to sorting, 1 ml PBS
(ThermoFisher Scientific) was added to each sample tube, along with
0.1 μg DAPI (ThermoFisher Scientific). The sample was incubated for

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10 min, gently vortexed and transferred to the flow cytometer (MoFlo Astrios EQ cell sorter). Single, live, GFP-positive AML cells (see Supplementary Fig. 1 for examples of gating strategy) were sorted into individual wells of a 96-well PCR plate filled with 2.6 µl Lysis Buffer (Takara Bio). Plates were sealed, spun down, snap-frozen and stored at −80 °C prior to preparation for complementary DNA synthesis using the SMARTseq-v4 assay.

**Flow analysis and sorting of CXCL12+ stromal cells for SMARTseq-v4.** Prior to sorting or analysis, samples were cell-surface stained with anti-CD45-APC/Cy7 (BioLegend, dilution 1:100) for 30 min at 4 °C in Medium-199 (Gibco) supplemented with 2% FBS. Prior to sorting, 1 mL PBS (ThermoFisher Scientific) was added to each sample tube, along with 0.1 µg DAPI (ThermoFisher Scientific). The sample was incubated for 10 min, gently vortexed and transferred to the flow cytometer (MoFlo Astrios EQ cell sorter). Flow cytometry data were analyzed using FlowJo (Treestar). Single, live, CD45-DSRed+ stromal cells (see Extended Data Fig. 6a for an example of gating strategy) were sorted into individual wells of a 96-well PCR plate filled with 2.6 µl Lysis Buffer (Takara Bio). Plates were sealed, spun down, snap-frozen and stored at −80 °C prior to preparation for cDNA synthesis using the SMARTseq-v4 assay.

**Leukemia burden and DPP4.** Leukemic bone marrow was blocked using anti-mouse Fc Block (BD Biosciences, dilution 1:50) for 10 min at 4 °C in Medium-199 supplemented with 2% FBS. Surface staining was thereafter performed with CD45-APC/Cy7 (BD Biosciences, dilution 1:100) and DPP4-PE (BioLegend, dilution 1:20) for 30 min at 4 °C. The cells were then washed and resuspended in Medium-199 supplemented with 2% FBS with 0.25 µg 7AAD (BD Biosciences). Flow cytometry was performed on a BD FACS Aria III sorter (BD Biosciences) and all data were analyzed using FlowJo (Treestar). Extended Data Fig. 8c shows the gating strategy used to distinguish DPP4high, DPP4int and DPP4neg cells. Note that DPP4-positive cells were defined as DPP4high and DPP4int.

**Leukemia cluster cell surface markers.** Leukemic bone marrow was blocked using anti-mouse Fc Block (BD Biosciences, dilution 1:50) for 10 min at 4 °C in Medium-199 supplemented with 2% FBS. Surface staining was thereafter performed with a leukemia cluster cocktail (Supplementary Table 6) for 30 min at 4 °C. The cells were then washed and resuspended in Medium-199 supplemented with 2% FBS with 0.25 µg 7AAD (BD Biosciences). Flow cytometry was performed on a BD FACS Aria III sorter (BD Biosciences) and all data were analyzed using FlowJo (Treestar).

**Intracellular DPP4 staining.** Bone marrow from leukemia-bearing mice was incubated with anti-mouse Fc Block (BD Biosciences, dilution 1:50) for 10 min at 4 °C followed by surface staining with anti-CD45-APC/Cy7 (BD Biosciences, dilution 1:100). Samples were washed and stained with LIVE/DEAD fixable viability dye (ThermoFisher) in accordance with the manufacturer’s instructions. The cells were thereafter fixed with Cytofix/Cytoperm (BD Biosciences) for 20 min at 4 °C. 1 µl Perm/Wash buffer (BD Biosciences) was then used to wash the cells and the cells were incubated with either anti-DPP4 or an isotype control antibody (BioLegend, dilution 1:20 for both) that were both conjugated to Alexa Fluor 647 in-house (Abcam) for 30 min at room temperature. The cells were then washed one last time in Perm/Wash buffer (BD Biosciences) and resuspended in Medium-199 supplemented with 2% FBS for analysis. Flow cytometry was performed on an LSR II instrument (BD Biosciences) and all data were analyzed using FlowJo (Treestar).

**Cell cycle analysis.** Bone marrow isolated from leukemic mice was blocked with anti-mouse Fc Block (BD Biosciences, dilution 1:50) for 10 min at 4 °C. Surface staining with anti-CD45-APC/Cy7 (dilution 1:100) and DPP4-PE (dilution 1:20) was performed at 4 °C for 30 min. Following this, the samples were washed in Medium-199 supplemented with 2% FBS and then fixed with Cytofix/Cytopperm (BD Biosciences) for 20 min at 4 °C. The fixed cells were then washed and rehydrated with 1× Perm/Wash buffer (BD Biosciences) and resuspended in Perm/Wash buffer containing anti-Ki67-AF647 at a 1:10 dilution for a 30 min incubation. After this, the samples were washed one more time with 1× Perm/Wash buffer (BD Biosciences) and then incubated in 1× Perm/Wash buffer (BD Biosciences) with 2 µg ml⁻¹ DAPI (BioLegend) for 10 min. Finally, the samples were spun down to remove the DAPI-containing buffer and resuspended in Medium-199 supplemented with 2% FBS for analysis. Flow cytometry was performed on a BD FACS Aria III sorter (BD Biosciences) and all data were analyzed using FlowJo (Treestar).

**Analysis of DPP4 expression following co-culture.** A total of 250,000 MLL-AF9 or HoxA9-Meis1 leukemia cells were plated at a 1:1 ratio with the following stromal cell lines: NIH-3T3 (American Type Culture Collection, ATCC), MS-5 (RIKEN), MLO-A5 (Kerafast), MC-3T3-E1 (ATCC) and SV40 immortalized bone marrow stroma (Supplementary Information). The cells were grown in RPMI 1640 (Gibco) supplemented with 1% Penicillin–Streptomycin (Gibco), and 10% FBS (Gibco). MLL-AF9 cultures were supplemented with 20 ng ml⁻¹ SCF, 10 ng ml⁻¹ IL-3 and 10 ng ml⁻¹ IL-6 (all cytokines from Peprotech). HoxA9-Meis1 cells were instead grown in 100 ng ml⁻¹ SCF and 5 ng ml⁻¹ IL-3. The cells were co-cultured for 3 days. For flow cytometry of DPP4, the co-cultures were trypsinized and subsequently blocked with murine Fc Block (dilution 1:50) for 10 min at 4 °C. Surface staining with anti-CD45-APC/Cy7 (dilution 1:100) and DPP4-PE (dilution 1:20) was then carried out for 30 min at 4 °C. The cells were thereafter washed and resuspended in Medium-199 supplemented with 2% FBS with 0.25 µg 7AAD (BD Biosciences) for analysis. Flow cytometry was performed on an LSR II instrument (BD Biosciences) and all data were analyzed using FlowJo (Treestar). For the analysis of DPP4 mean fluorescence intensity, each sample was normalized to a corresponding isotype control antibody-stained sample.

**Myeloid lineage markers.** Bone marrow collected from C57Bl/6j mice was stained with a hematopoietic stem and progenitor cell cocktail (Supplementary Table 6) for 45 min at 4 °C. The cells were then washed and resuspended in Medium-199 supplemented with 2% FBS with 0.25 µg 7AAD (BD Biosciences). Flow cytometry was performed on a BD FACS Aria III sorter (BD Biosciences) and all data were analyzed using FlowJo (Treestar).

**Reverse transcription with quantitative PCR**

RT-qPCR was performed to determine levels of Dpp4 mRNA. A total of 1 × 10⁶ leukemia cells or T cells were lysed and the RNA was extracted using the RNeasy Plus mini kit isolation kit (Qiagen). RNA was subsequently reverse transcribed into cDNA with the SuperScript IV First-Strand Synthesis System (ThermoFisher). The qPCR analysis was performed using iTaq Universal SYBR Green Supremex (Biorad) with primers specific for Dpp4 (forward, ACCTGGAAGGTTCCCTGG; reverse, CACAAAGAGTAGGACTTGACCC) and Gapdh (forward, TGGCGGTGGATGACTGA; reverse, TTGCGTTGAACTCAGGAGG). Threshold values (Ct) were estimated using CFX Maestro (Biorad) and transcript levels were normalized by subtracting the corresponding Gapdh values. The relative amount of RNA is presented as 2⁻ΔΔCt.

**Droplet-based single-cell RNA sequencing**

WGBM and Image-seq samples were collected in a hemocytometer and encapsulated for a maximum output of 8,000 cells into emulsion droplets using the Chromium Controller (10x Genomics). scRNA sequencing libraries were subsequently prepared using Chromium Single Cell 3 v2 Reagent kits (10x Genomics). Reverse transcription and library preparations were done on a Biorad T100 Thermo Cycler.
Differential expression. For differential expression analysis between cell types in the 10x scRNA-seq data, a Wilcoxon rank sum test, implemented by the getDifferentialGenes() function from Conos R, was used to identify statistically significant marker genes that were expressed in each cell cluster (Supplementary Table 1). The genes were considered differentially expressed if the $P$-value determined $Z$ score was greater than 3. For differential expression analysis between Image-seq and WCBM (for example, Image-seq monocytes versus WCBM monocytes), the getPerCellTypeDE() function in Conos was utilized.

Cell annotation. Annotation of the cluster communities was done using marker gene expression. Initial annotations were obtained by entering the top 100 differentially expressed genes in each cluster (ordered by logFoldChange) into the CellKb database and further refined by consulting the primary literature referenced therein along with other relevant publications. Specificity of selected markers was additionally confirmed by evaluating the expression in the Haemopedia and CellMarker bone marrow datasets. We identified markers to classify ten major cell types: B cell (Ms4a1, Ly6d, Cd79a), pre/pro-B cell (Vpreb1, Vpreb3, Dnmt), basophil (Mcp8, Pss34, Ms4a2), dendritic cell (Bst2, Irf9, Siglec, Cox2a2), erythroblast (Car2, Hmgn, Cse, Cpox, Atpf), monocyte (Ly2, Ctas, S100a4), monocyte progenitor (Ly6c2, S100a10), diverse progenitors (Cd34, Prtn3, Mpo, Elane, Mpi), granulocyte progenitor (S100a8, S100a9, Cebpe, Fcnb) and neutrophil (Mmp8, Iftm6, S100a11, S100a8, S100a9).

SMARTseq-v4 sequencing data analysis

SMARTseq-v4 sequencing data were aligned with hisat2 (ref. 10) (v4.8.2), using the genome reference mm10. We used featureCounts (v1.6.4) to calculate read counts. The quality of cells was then assessed based on the number of total counts per cell (library size), requiring at least 500,000 reads per cell. A total of 84 AML and 43 stroma cells were retained for downstream analysis. Seurat was used to analyze the SMARTseq-v4 data, and the quality parameters are listed in Supplementary Table 7 and Extended Data Fig. 5d,e. In brief, we first performed data normalization using the NormalizeData function with default settings. The top 30 principal components were used to construct shared nearest neighbors graphs and UMAP embeddings. The FindAllMarkers function from Seurat was used to identify differentially expressed (marker) genes for clusters or subtypes. Mesenchymal stromal and endothelial cells were annotated using well-established marker genes: for mesenchymal stromal cells the marker genes were Cxcl12, Pcolce, Ogn and Adipoq; and for the endothelial cells the markers were Eng, Kdr, Ppav2 and Egfl7 (see also the heatmap in Extended Data Fig. 6d, as well as ref. 21).

Cell cycle signature score. To assess cell states in different cell subsets and conditions we used a gene set signature score to measure the relative difference of cell cycle states. The signature scores were calculated as average expression values of the genes in a given set. The signature gene list was downloaded from Whitfield et al. Hierarchical clustering of cell cycle signature score was used to group AML cells. A two-sided Student’s $t$-test was used to assess differential expression of selected cell cycle genes between proliferating (P) and non-proliferating (NP) cells (Supplementary Table 4).

Regressing out cell cycle genes. Seurat (v4.0.6) was used to regress out cell cycle genes. First, we assigned each cell a score, based on its expression of G2/M and S phase markers with the CellCycleScoring function. Then we applied the ScaleData function to regress out the cell cycle genes. The scaled residuals of this model represent a corrected expression matrix that can be used downstream for dimensionality reduction. UMAP embedding and graph-based shared nearest neighbor clustering were used with five principal components, and Seurat::FindClusters() was used to identify AML cell sub-clusters. Seurat::FindAllMarkers, which utilizes a two-sided Wilcoxon rank sum test to assess statistical significance, was used to find the differentially expressed genes within each sub-cluster (Supplementary Table 2).

Analysis of differentially expressed genes. DESeq2 (ref. 60) was used to analyze differentially expressed genes between P and NP cells (Fig. 5a and Supplementary Table 3), as well as between P + IM (that is, intermediate) and NP cells (Supplementary Table 5).

GO term enrichment. To test for enriched GO biological processes in gene sets, the ClusterProfiler (v4.0.0) package was used to evaluate the enrichment of GO categories in the sets of upregulated and downregulated genes and rank them by adjusted $P$ value (Fig. 5h and 5i).
Cells were then grouped based on their fold-change as either proliferating (P, fold-change >2), intermediate (IM, fold-change >0 and ≤2), or non-proliferating (NP, fold-change ≤0), which corresponded to average cell numbers of 29.6 (with a 99.9% confidence interval of 21.7 to 37.4 cells), 13.2 (with a 99.9% confidence interval of 5.5 to 20.8 cells), and 2.1 (with a 99.9% confidence interval of −1.2 to 5.4 cells), respectively, on day 3.

Analysis of cell tracker labeling. AML cells were labeled with CellTracker Red (CMTPX, 10 µM, ThermoFisher Scientific) before transplantation. In brief, the cell suspension (in Ca2+/Mg2+ free PBS containing 10 µM CMCTP) was incubated at 37 °C for 45 min. Cells were then pelleted to remove the staining solution and resuspended in 300 µl PBS for retro-orbital injection. The mean CellTracker intensity was measured on day 3 after transplantation and measured at the brightest plane of the cell. The cells were considered positive for CMTPX when the measured signal was greater than 12.5 (the background noise in the marrow cavity). The number of CMTPX-positive cells was then divided by the total cell number sampled from the bone marrow cavities harboring the same cell counts (n = 7 mice).

Analysis of DPP4 expression in vivo
DPP4 antibody (BioLegend) and isotype control antibody (BioLegend) were conjugated to AF568 using the Lightning Link kit (Abcam). Antibody and isotype were injected retro-orbitally 1 h prior to the imaging session at a dosage of 1 mg kg⁻¹. Images were collected with an excitation wavelength of 980 nm (1.6 nJ pulse energy, filter set detailed above), using a z-step of 2 µm with a 15-frame average on the custom-built microscope described above.

AML cells were segmented based on the GFP signal in the obtained images. For this, seeds were generated using the interactive watershed tool (Fiji) and used as input for the 3D-Watershed segmentation plugin (Fiji). The images were thresholded to generate a mask. This was used to calculate the total AF568 signal, as well as the total GFP signal, in each cell using the red and green channels of the images, respectively, as input for the 3D Object counter plugin (Fiji). The ratio of red to green fluorescence (multiplied by 10) for each cell was plotted in Fig. 5c both for the DPP4 and isotype control cells.

Distribution of AML cells in D-, M- and R-type cavities. The protocols to determine the bone remodeling status have been described previously. The first calcium-chelating reagent dye 1 (Tetracycline, Sigma, 35 mg kg⁻¹) was given i.p. 48 h prior to imaging to label the bone fronts and track the bone resorption activities over the course of 2 days. The second calcium-chelating reagent dye 2 (Alizarin Red, 40 mg kg⁻¹) was injected on the day of imaging to label all of the bone fronts. The bone remodeling status was then defined based on the ratio of dye 1 to dye 2 in a single bone marrow cavity (the concave endosteum), and therefore the bone marrow cavities were classified as: deposition type (D-type; dye 1: dye 2 ratio ≥ 75%); resorption type (R-type; dye 1: dye 2 ratio < 25%), or mixed type (M-type; dye 1: dye 2 ratio 25–75%). The distributions of seeding and expansion of AML cells were then mapped with respect to the D-, M- and R-type cavities on day 0 (3 h after transplantation), day 1 and day 3 after transplantation. The same mouse was followed up longitudinally on day 0 and day 1 and the cavity type was defined on day 0. A separate cohort of animals was used for day 3 to avoid unwanted inflammatory responses from the survival surgical procedures.

Animal handling
Male and female 8-week-old C57Bl/6/J mice (cat. no. 000664) or male and female CXCL12-DsRed mice (cat. no. 022458) were ordered from the Jackson Laboratory, housed in our animal facility for at least 2 weeks and used for experiments at between 10 and 14 weeks of age.
The 8-week-old female Fezhi/J mice B6J.L29(Cg)-Gt(Rosa)26Sortm1.1(CAG-cas9*-EGFP)Fezhi/J were ordered from the Jackson Laboratory (cat. no. 026179) and used in experiments. Male and female β-actin-GFP mice (Jackson Laboratory, cat. no. 006567) were bred in-house and used between 10 and 16 weeks of age. Male and female β-actin-DsRed mice (Jackson Laboratory, cat. no. 006501) were bred in-house and used between 10 and 16 weeks of age. The β-actin luciferase (βAct) mice from Taconic (cat. no. 1H977) were bred with ubiquitin-c-GFP (UcGFP) mice from the Jackson Laboratory (cat. no. 004333) to generate βAct-UcGFP transgenic mice. An 8-week-old female βAct-UcGFP mouse was then used to generate the HA9M1 cell line. All mice were housed in the pathogen-free Massachusetts General Hospital (MGH) Animal Facilities, which were equipped with ventilated micro-isolator cages. Sentinel programs and veterinary oversight were in place. Mice were given standard chow and drinking water ad libitum. An automated 12 h dark–12 h light cycle was observed and mice were housed at a fixed temperature (21°C) and humidity (66%). The MGH Animal Facility is under the supervision of the MGH Center for Comparative Medicine. All facilities are fully accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International (OAAS) and meet the National Institutes of Health standards as set forth in the Guide for Care and Use of Laboratory Animals by the Department of Health and Human Services. All procedures involving animals were carried out in agreement with protocols 2012N000190, 2007N000148 or 2016N000085 approved by the Institutional Animal Care and Use Committee of Massachusetts General Hospital.

Statistics and reproducibility

P < 0.05 was considered significant unless specified otherwise. The number of biological replicates and independent experiments for each graph, along with the test statistic, are listed in each Figure and Extended Data Figure and legend. A list of statistical parameters for these, including confidence intervals, degrees of freedom, mean, standard deviation and effect sizes can be found in the Supplementary Information. The number of biological replicates and independent mice for all images that are shown in the Figures and Extended Data Figures are also included in the respective Figure legends.

Reagents

A list of antibodies is given in Supplementary Table 6 and a list of reagents is given in Supplementary Table 8.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The 10x-seq and SMARTseq-v4 data generated in this work have been deposited in the Gene Expression Omnibus (GEO) database GSE188902 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE188902), which is publicly available. The mouse mm10 reference genome was downloaded from 10x Genomics (https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest/). Public, bulk RNA-Seq AML datasets from OHSU (https://cibiportaldatabah.s3.amazonaws.com/aml_ohsu_2018.tar.gz) and TCGA (https://cibiportaldatabah.s3.amazonaws.com/aml_tgca_pan_can_atlas_2018.tar.gz) were downloaded from cibiportal. Due to the extremely large file sizes accompanying the extensive imaging data, raw image data are available from the corresponding authors upon request. Cell lines are available from the authors upon request and mouse lines are commercially available at the Jackson Laboratory. Source data files for all graphs in the Figures and Extended Data Figures are linked to the online version of the manuscript. Source data are provided with this paper.

Code availability

The codes generated during this study are available at the GitHub repository (https://github.com/shenglinmei/Image-seq) and Zenodo (https://zenodo.org/record/7067661).

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Author contributions

C.H. developed the Image-seq platform, carried out the imaging and Image-seq cell isolation experiments, developed thestromal cell...
isolation protocol with the help of K.G., performed AML cell culture, SMARTseq-v4 library preparation and sequencing experiments (with the help of the Bauer core), analyzed the data, interpreted the scRNA-seq data and coordinated the entire project. C.H. and C.P.L. designed the optical platform and wrote the manuscript with input from all of the authors. K.G. performed the 10x Genomics Chromium and sequencing experiments as well as the flow cytometry, co-culture and qPCR experiments. S.M. analyzed the scRNA-seq data. S.-C.Y. performed the in vivo imaging and analysis of CellTracker and AML distribution in different bone remodeling environments. D.R. performed cell isolation experiments from the tibia. J.M. generated the secondary transplanted HoxA9/Meis1 leukemia cells and contributed to flow cytometry experiments. R.T. contributed to the early design and assembled a preliminary version of the optical platform. D.B.S. assisted in the design and interpretation of the leukemia and flow cytometry experiments. P.V.K., D.B.S., D.T.S. and C.P.L. supervised the study. C.P.L. conceived the project with input from P.V.K. and D.T.S.

Competing interests
C.H., D.R. and C.P.L. are inventors on a coversheet provisional filed by Massachusetts General Hospital (App. No. 63/394,274, patent application pending) that describes the Image-seq technology. D.B.S. is a founder, consultant and shareholder for Clear Creek Bio. D.T.S. is a director and shareholder for Agios Therapeutics and Editas Medicines; a founder, director, shareholder and scientific advisory board member for Magenta Therapeutics and LifeVault Bio; a shareholder and founder of Fate Therapeutics and Garuda Therapeutics and a director, founder and shareholder for Clear Creek Bio; a consultant for FOG Pharma, Inzen Therapeutics and VCanBio; and a recipient of sponsored research funding from Sumitomo Dianippon. P.V.K. serves on the Scientific Advisory Board to Celsius Therapeutics Inc. and Biomage Inc. All other authors have no competing interests.

Additional information
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Extended Data Fig. 1 | Full optical diagram of Image-seq set-up. Radiation at 775 nm is generated by frequency doubling the 1550 nm output from a fiber laser. Radiation at 920-980 nm is generated by soliton self-frequency shift of 1550 nm radiation to 1840–1960nm in a LMA photonic crystal fiber, which is subsequently frequency doubled to obtain the imaging wavelength. A scanning GM (y scan) and scanning PM (x scan) are conjugated to one another, along with the back aperture of the OL so that a scan along the x and y dimensions is achieved in the image plane. Multiphoton emission wavelengths are collected with a combination of DCs and PMTs to generate an image. A confocal reflectance channel that utilizes an APD and 20µm-size pinhole can also be used to generate images. \( \lambda/2 \): \( \lambda/2 \)-plate. PBS: polarization beam splitter. BiBO: BiBO3 - Bismuth orthoborate crystal. LMA: large mode-area. APD: avalanche photodiode. ph: pinhole. DC: dichroic mirror. GM: galvo mirror. PM: polygonal mirror. PMT: photomultiplier tube. MP: micropipette. OL: objective lens.
Extended Data Fig. 2 | Gating strategy for flow cytometry analysis of samples from Fig. 2. a) Gating strategy for the analysis of various leukocyte populations in a whole calvarium bone marrow preparation (WCBM) and concatenated data from all micropipette samples. Analyzed cells were gated on singlet live cells and percentages refer to percent of parent gate. The same gating strategy was used for the analysis of whole tibia and tibia micropipette samples. b) Gating strategy for flow cytometry analysis of CXCl12+, CD45− stromal cells from 5 concatenated micropipette samples.
Extended Data Fig. 3 | Long bone cell extraction. a) 3D visualization of a target location in a β-actin-GFP mouse before and after cell aspiration. The bottom panel is obtained by rotating the top panel by 180° around the x axis. The isolated cell volume is marked by an orange outline. b) Comparison of micropipette and whole bone marrow preparations from the calvarium and tibia. Statistical analysis was performed using the two-way ANOVA for each individual cell population, with the two independent variables being bone type (tibia vs calvarium) and procedure (micropipette vs whole bone marrow preparations). Listed p values were not adjusted for multiple comparisons. Using a Bonferroni correction, statistical significance is achieved for p < 0.006 (α=0.05); significant p values are highlighted in red.

### Comparison of cell populations in micropipette vs whole bone marrow preparations as well as tibia/calvarium

|           | T cell | Macrophage | Inf mono | Granulocyte | Patr mono | B cell | Pre-B cell | Pre-pro-B cell |
|-----------|--------|------------|----------|-------------|-----------|--------|------------|----------------|
| M / W     | 0.67   | 5.3e-5     | 0.091    | 0.025       | 0.96      | 0.31   | 5.4e-5     | 1.3e-8         |
| C / T     | 0.006  | 0.13       | 0.86     | 0.016       | 0.13      | 0.0013 | 0.45       | 0.55           |

Statistical significance using Bonferroni-corrected p-value (α =0.05): p < 0.006.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Distribution of cell proportions in Image-seq samples along with GO analysis of differentially expressed genes. a) Distribution of cell proportions for the major hematopoietic cell populations from Fig. 3b,c, shown as averaged proportions over all Image-seq vs all WCBM samples. b) Breakdown of (a) showing the distribution of cell proportions for each sample. WCBM samples outlined by red border, Image-seq samples outlined by blue border. c) Table with p values for differences in cell proportions between WCBM and Image-seq samples (obtained using a two-sided Wilcoxon rank sum test). d) Top GO terms obtained from upregulated genes in Image-seq samples, broken down by cell type. e) Top GO terms obtained from downregulated genes in Image-seq samples, broken down by cell type. WCBM: whole calvarial bone marrow. GO: gene ontology. Pre/Pro-B cell: Pre and pro B cell. DP: Diverser progenitor.
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | Quality control metrics for the single-cell RNA-seq data from Figs. 3, 4 and Extended Data Fig. 6. a) Percentage of cells that passed quality control after sequencing in each 10x run (WCBM N = 3, Image-seq N = 11 biological replicates). A two-sided Mann–Whitney test was used to assess statistical significance. 3 (WCBM) and 5 (Image-seq) independent mice were used, respectively. b) Number of unique molecular identifiers and expressed genes per cell for each Image-seq and whole calvarium bone marrow (WCBM) sample. c) Table that summarizes the number of cells sequenced from each 10x run, along with the number of cells and percentage of cells that passed quality control. d) Comparison of the number of expressed genes and total number of molecules (= total UMI) between our WCBM, Image-seq and AML SMARTseq-v4 data. Statistical significance was determined using a two-sided Wilcoxon rank sum test. e) Number of expressed genes per cell and total mapped reads (= molecules) per cell for the AML SMARTseq-v4 data, including scatter plot. d),e) WCBM N = 5295 (3 independent mice), Image-seq N = 8078 (5 independent mice), SMARTseq N = 84 cells (11 independent mice). Boxplot represents the interquartile range (IQR, the range between the 25th and 75th percentile), the mid-point of the data is indicated as a line within. Whiskers indicate the upper and lower value that is within 1.5 times the IQR.
Extended Data Fig. 6 | See next page for caption.
Extended Data Fig. 6 | Image-seq of stromal cells. a) Example of the gating strategy used to isolate CXCl12⁺, CD45⁻ stromal cells from a micropipette sample for cell lysis and SMARTseq-v4 library preparation. Sorted cells were gated on singlet live cells (DAPI⁻). b) UMAP embedding of the stroma SMARTseq-v4 data using the top 30 principal components, color coded by cell population. c) Heatmap of scaled normalized expression for the 20 most highly expressed genes within each cell cluster. d) Heatmap of scaled normalized expression for Lepr-MSC and Endothelial cell marker genes from the literature. UMAP: uniform manifold approximation and projection. MSC: Mesenchymal stromal cells, EC: endothelial cells.
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Cell cycle analysis of AML cells from Fig. 4, along with selected intravital microscopy data. a) Percentage of AML cells that are CellTracker Red+ on day 3 (CellTracker + (%)) as a function of the number of cells per cavity. b) Violin plots of selected cell cycle relevant genes from Supplementary Table 4. Statistical significance was determined using a two-sided Student’s t-test (Sample size: NP = 33, IM = 17, P = 34 cells, 11 independent mice). c) Heatmap showing scaled average expression of gene sets in distinct phase of cell cycle, each column indicates a cell. AML cells are clustered into three groups by hierarchical clustering based on cell cycle genes. d) Representative images of P and NP cells from mice injected with rat IgG2a,κ-AF568 antibody (DPP4 isotype control). Imaging of DPP4 isotype control labeling was performed in 14 biological replicates with NP cells and 6 biological replicates with P cells (1 independent mouse). Definition of P/ NP/IM based on cell number at day 3.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | Gating strategy and DPP4-correlated genes by flow cytometry. a) Ki-67 flow cytometry analysis of HA9M1 cells 13 days after transplantation (N = 7 biological replicates, 2 independent experiments). Statistical significance was determined by unpaired, two-sided Student’s t-test, error bars denote standard error of the mean. b) Ki-67 cell cycle gating strategy. c) Gating strategy used for identifying DPP4<sub>high</sub>, DPP4<sub>int</sub> and DPP4<sub>neg</sub> cells. d) (Left) Day 10 leukemia burden in mice transplanted with 1,000 DPP4<sub>high</sub> (N = 3) or DPP4<sub>neg</sub> (N = 4) HA9M1 cells (2 independent experiments). Error bars denote standard deviation. (Right) Percentage of DPP4<sub>high</sub>-int-neg cells isolated from recipients transplanted with DPP4<sub>high</sub> (N = 3, red bars) or DPP4<sub>neg</sub> (N = 4, beige bars) HA9M1 cells (2 independent experiments). An unpaired, two-sided Student’s t-test was used for statistical analysis, error bars denote standard deviation. e) Mean fluorescence intensity of Itgb7, Flt3 and CD48 detected in DPP4<sub>high</sub>-neg HA9M1 cells by cell surface staining and flow cytometry 3 weeks after transplantation (N = 5 biological replicates, 2 independent experiments). Statistical significance was determined by one-way ANOVA followed by Tukey’s post-hoc test, error bars denote standard deviation. f) DPP4, Itgb7, Flt3 and CD48 expression level for the AML clusters from Fig. 4h. A two-sided Wilcoxon rank sum test was used for statistical analysis (AML-AP1 N = 21, AML-GMP N = 28, AML-mono N = 35 cells, 11 independent mice). g) GO analysis of top 300 up-regulated genes comparing P and NP cells. The statistical analysis was performed by hypergeometric test. Terms related to the inhibition of T cells are highlighted by the red rectangle. HA9M1: Hoxa9/Meis1-Ubiquitin-c-GFP. GO: gene ontology.
Extended Data Fig. 9 | See next page for caption.
Extended Data Fig. 9 | Flow cytometry analysis of DPP4 in healthy myeloid precursors. a) Percentage of DPP4 + cells in various myeloid cell populations (N = 6 biological replicates). Error bars denote standard deviation. Data were collected from two independent experiments. LKS, Lineage- cKit+ Sca-1+; MEP, megakaryocytic-erythroid progenitor; CMP, common myeloid progenitor; StemGMP, Stem-like granulocyte-monocyte progenitor; MonoGMP, monocytic granulocyte-monocyte progenitor; GrGMP, granulocytic granulocyte-monocyte progenitor; Inf monocytes, inflammatory monocytes; Patr monocytes, patrolling monocytes. b) Gating strategy for the myeloid cell populations from (a), along with DPP4 and isotype count data for each cell type. Analyzed cells were gated on singlet live cells and percentages refer to percent of parent gate. Gray curves represent the Isotype control, red curves DPP4.
Extended Data Fig. 10 | Detection of DPP4 on in vitro cultured cells. a) Cell surface antibody labeling with DPP4 in HA9M1, analyzed by flow cytometry. b) Analysis of Dpp4 expression by QPCR on in vitro cultured HA9M1 and MLL-AF9 cells. Results are compared to splenic T cells, which express reproducible levels of Dpp4. (N = 3 biological replicates from 2 independent experiments). Error bars denote standard deviation. c) Flow cytometry analysis of DPP4 expression in HA9M1 by intracellular staining with DPP4 antibody. d) Analysis of Dpp4 by QPCR on AML cells collected from the bone marrow 3 weeks after transplantation. DPP4⁺ and DPP4⁻ HA9M1 cells were sorted by flow cytometry and their gene expression compared to that of splenic T cells by Q-PCR. After initial analysis, the T cells and DPP4⁺ HA9M1 cells were cultured for 4 days using the same media and analyzed once again for Dpp4 expression. HA9M1: Hoxa9/Meis1-Ubiquitin-c-GFP (N = 3 biological replicates from 2 independent experiments). Statistical significance was calculated by one-way ANOVA followed by Sidak's post-hoc test, error bars denote standard deviation.
**Reporting Summary**

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**Statistics**

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- **n/a**
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- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
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- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
  - Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated

*Our web collection on statistics for biologists contains articles on many of the points above.*

**Software and code**

Policy information about availability of computer code

2-photon and confocal microscopy was performed either with the Image-seq platform (detailed description in the manuscript) or on a custom-built two-photon excitation microscope equipped with a Ti: Sapphire laser oscillator (Spectra-Physics InSight X3) and coupled with an Olympus 60x water-immersion objective lens (previously described in Yeh et al, Nat. Commun. 2022; Christodoulou et al, Nature 2020; Spencer et al, Nature 2014). Laser scanning and acquisition were controlled by a custom-built software. Spatially-resolved cell isolation was carried out with the Image-seq platform. The Illumina HiSeq 4000 Sequencing System and bcl2fastq (v1.8.4) were used to acquire the 10X data. The sequenced 10X libraries were mapped to the mm10 genome using Cell Ranger software (v 3.0.2). The NextSeq 500 Sequencing System (Illumina) was used to acquire SMART-seqv4 data. SMART-seqv4 sequencing data were aligned with hisat2 (v 4.8.2). Flow cytometry data were acquired on an LSRII flow cytometer, a BD FACS Aria III and a MoFlo Astrios EQ.

**Data analysis**

We used python and R to perform analysis of sequencing data. Code related to this manuscript can be found at https://github.com/shenglinmei/Image-seq.

- Cell Ranger software (version 3.0.2)  
- featureCounts (v1.6.4)  
- bcl2fastq (v1.8.4)  
- hisat2 (v4.8.2)  
- DEseq2 (v1.32.0)  
- Seurat (v4.0.6)  
- Pagoda2 (v1.0.10)  
- Conos (v1.4.1)  
- ClusterProfiler (4.0.0)  
- R 4.1.1  
- python 3.7  
- FlowJo 10.8.0  
- Fiji (ImageJ) Version 2.1.0/1.53g was used for image analysis.
GraphPad Prism9 was used for statistical analysis of flow cytometry and imaging data. CFX Maestro software (Biorad, v1.1) was used for the analysis of qPCR data.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Policy information about availability of data
All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

The 10X-seq and SMART-seqv4 data generated in this work have been deposited into the Gene Expression Omnibus (GEO) database GSE188902 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE188902) which is publicly available. The mouse mm10 reference genome was downloaded from 10X genomics (https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest?). Public, bulk RNA-seq AML datasets from TCGA (https://cbioportal-datahub.s3.amazonaws.com/aml_ohsu_2018.tar.gz) and OHSU (https://cbioportal-datahub.s3.amazonaws.com/laml_tcgga_pan_can_atlas_2018.tar.gz) were downloaded from cbioportal. Due to the extremely large file sizes accompanying the extensive imaging data, raw image data is available from the corresponding authors upon request. Cell lines are available from the authors upon request and mouse lines are commercially available at the Jackson Laboratory. Source data files for all graphs presented in the Figures and Extended Data Figures are linked to the online version of the manuscript.

Field-specific reporting

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

### Sample size
For sequencing data: Since this is a new technology and we did not test any hypotheses, no sample size calculation for statistical power of hypothesis testing was performed. Sample sizes were chosen based on the data obtained from a small pilot for each experiment. For flow cytometry data: Sample size was approximated and derived from extensive publications within the hematopoiesis field which are based on previous a priori power testing. No statistical methods were used to pre-determine sample sizes but our sample sizes are in agreement with those reported in previous publications (Yusuf et al. Blood 2020), (van Gastel et. al Cell Metabolism 2020) and (Shah et. al. Nat. Biomed. Eng. 2020). For imaging data: Sample size was approximated by consulting published intravital microscopy datasets of the calvarium bone marrow. The final sample sizes were adequate based on consistency of measured results in each group and are in agreement with those reported in previous publications (Yeh et. al Nat. Commun. 2022), (Christodoulou et al. Nature 2020), and (Spencer et. al. Nature 2014).

### Data exclusions
No data was excluded.

### Replication
Experimental findings were reliably reproduced. To verify reproducibility of the findings the vast majority of experiments were repeated at least three independent times.

### Randomization
We did not allocate samples/organisms into experimental groups, so no randomization was necessary.

### Blinding
Analysis of flow cytometry and qPCR data is not subjective and therefore blinding was deemed unnecessary. For the imaging: It is not possible to perform the imaging analysis during the experiment, so it was not possible to pre-select any regions that could have produced a preferred result. Similarly, it is not possible to perform the sequencing data analysis during the experiment, so it was not possible to select any preferred regions for Image-seq cell isolation. In addition, partial blinding was performed since cell isolation, library preparation, sequencing and data analysis were performed by different people.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.
Antibodies

Marker Fluorophore Dilution Vendor Catalog# Clone Lot#

CD71 PE/Cy7 1 to 200 Biolegend 113812 RI7217 B259648
Ter119 APC 1 to 100 Biolegend 116212 Ter119 B286863
CD3 BUV737 1 to 100 BD Biosciences 612771 145-2C11 133181
CD11b AF700 1 to 200 Biolegend 101222 M1/70 B308254
CD19 FITC 1 to 200 Biolegend 115506 6D5 803942
B220 BUV395 1 to 200 BD Biosciences 563792 RA3-6B2 6320735
IgM PE 1 to 200 eBioscience 12-5890-82 eB121-15F9 2134180
F4/80 APC/Cy7 1 to 200 Biolegend 123118 BM8 B274179
CD6 Tn Bv570 1 to 200 Biolegend 128030 HK1.4 B310124
Ly6G Bv421 1 to 200 BD Biosciences 562737 1A8 B301980
FC block N/A 1 to 50 BD Biosciences 553142 2.4G2 1279889
CD45 Bv421 1 to 100 BioLegend 103134 30-F11 B287242
CD45 APC/Cy7 1 to 100 BD Biosciences 557659 30-F11 1046784
DPP4 PE 1 to 20 Biolegend 137804 H194-112 B219006
FIt3a Bv421 1 to 50 Biolegend 135314 A2F10 B329451
Fgb7 Bv711 1 to 20 Biolegend 321240 FIB504 B312128
CD48 BUV737 1 to 200 BD Biosciences 565240 1A8 8033935
Rat IgG2a k Isotype PE 1 to 20 Biolegend 400508 RTK278 B290861
NK.1.1 APC 1 to 200 Biolegend 108710 PK136 B31103
CD3 APC 1 to 100 eBioscience 17-0031-82 145-2C11 4277755
CD4 APC 1 to 200 Biolegend 100412 GK1.5 B340887
CD8 APC 1 to 200 Invitrogen 17-0081-83 S3-6.7 1978219
B220 APC 1 to 200 BD Biosciences 553092 RA3-6B2 B350149
CD19 APC 1 to 200 Biolegend 152409 H194-112 B219006
Gr1 Bv711 1 to 100 Biolegend 553086 RA3-6B2 7220040
NK1.1 Bv421 1 to 200 Biolegend 108711 M1/70 B340052
Gr1 Bv421 1 to 100 Biolegend 108404 R86-8CS B351067
NK1.1 Bv421 1 to 100 Biolegend 108704 PK136 B359276
CD19 Bv421 1 to 100 Biolegend 553784 ID1 503784
BB20 Bv570 1 to 100 BD Biosciences 553086 RA3-6B2 2279903
DPP4 later conjugated to AF680 1 mg/kg (in vivo) BioLegend 137802 H194-112 B216439
IgG2a, kappa Isotype later conjugated to AF568 1 mg/kg (in vivo) BioLegend 400502 RTK2758 B318086

Validation

Only commercially available antibodies were used in this study. All antibodies used for flow cytometry and microscopy were previously validated for the respective application by the vendor. Individual listings of citations using these reagents in the respective imaging or cytometric application and can be found on the manufacturer page which can be accessed using the catalog numbers provided above.
Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)
The HoxA9/Meis1 cell line was kindly provided by the lab of Dr. David Scadden. The MLL/AF9 cell lines were kindly provided by the lab of Dr. David Scadden. NIH3T3 cells, and MC-3T3-E1 cells were purchased from ATCC. MS-5 cells were obtained from RIKEN and MLO-A5 cells were bought from Kerafast.

Authentication
No authentication was performed

Mycoplasma contamination
Cell lines tested negative for mycoplasma contamination.

Commonly misidentified lines
No commonly misidentified cell lines were used in this study.

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

Laboratory animals
Male and female 8wk-old C57Bl6/J mice (#000664) or male and female CXCL12-DsRed mice (#022458) were ordered from the Jackson Laboratory, housed in our animal facility for at least 2 weeks and used for experiments between 10-14 weeks of age. 8-week old male Fezh/J mice (B6J.129(Cg)-Gt(Rosa26Sortm1.1(CAG-cas9*,-EGFP), catalog number 026179) were ordered from the Jackson Laboratory and used for experiments. Male and female beta-actin-GFP mice (Jackson #006567) were bred in house and used between 10-16 weeks of age. Male and female beta-actin-DsRed mice (Jackson #006051) were bred in house and used between 10-16 weeks of age. 

β-actin luciferase (βact) mice from Taconic (#11977) were bred with ubiquitin-c-GFP (UcGFP) mice from the Jackson laboratory (#004353) to generate βact-UcGFP transgenic mice. An 8-week old female βact-UcGFP mouse was then used for generating the HA9M1 cell line. All mice were housed in the pathogen-free MGH Animal Facilities, which were equipped with ventilated micro-isolator cages. Sentinel programs and veterinary oversight were in place. Mice were provided with standard chow and drinking water ad libitum. An automated 12h dark - 12h light cycle was observed and mice were housed at a fixed temperature (70 Fahrenheit) and humidity (66%). The MGH Animal Facility is under supervision of the MGH Center for Comparative Medicine. All facilities are fully accredited by AAALAC (#000809) and meet NIH standards as set forth in the "Guide for Care and Use of Laboratory Animals" (DHHS).

Wild animals
No wild animals were used in this study.

Field-collected samples
No field-collected samples were used in this study.

Ethics oversight
All procedures involving animals were carried out in agreement with protocols 2012N000190, 2007N000148 or 2016N000085 that were approved by the Institutional Animal Care and Use Committee of Massachusetts General Hospital and followed the guidelines set forth in the US National Institute of Health Guide for the Care and Use of Laboratory Animals. Animals were housed in a 12h light/dark cycle at a constant temperature and humidity and with ad libitum access to food and water.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Flow Cytometry

Plots

Confirm that:

- The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
- The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
- All plots are contour plots with outliers or pseudocolor plots.
- A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation

Flow cytometry analysis of WBM and micropipette
WBM and micropipette samples from calvarium and tibia were blocked with anti-mouse Fc block (BD Biosciences, dilution 1 to 50) for 10 minutes at 4°C. The cells were thereafter stained with Blood cell lineage cocktail (Table S6) for 30 minutes at 4°C. For detection of dead cells 7AAD (BD Biosciences, 0.25μg) was added to the sample prior to analysis. Flow cytometric analysis was performed on a BD FACS Aria III sorter (BD Biosciences) and all data was analyzed using FlowJo software package (Treestar).

Flow sorting of AML cells for SMARTseq-v4
Prior to sorting, 1ml of PBS (ThermoFisher Scientific) was added to each sample tube, along with 0.1μg of DAPI (ThermoFisher Scientific). The sample was incubated for 10 min, gently vortexed and transferred to the flow cytometer (MoFlo Astrios EQ cell sorter). Single, live, GFP+ AML cells (see Figure S1 for examples of gating strategy) were sorted into individual wells of a 96-well PCR plate filled with 2.6μl of Lysis Buffer (Takara Bio USA, Inc.). Plates were sealed, spun down, snap-frozen and stored at -80°C prior to preparation for cDNA synthesis using the SMARTseq-v4 assay.
Flow cytometry analysis of leukemia burden and DPP4

Leukemic bone marrow was sorted using an anti-mouse Fc Block (BD Biosciences, dilution 1 to 50) for 10 minutes at 4°C in Medium 199 + 2% FBS. Surface staining was performed with anti-CD45-APC/Cy7 (BD Biosciences, dilution 1 to 100) and DPP4-PE (BioLegend, dilution 1 to 100) for 4°C. The cells were then washed and resuspended in Medium 199 + 2% FBS with 0.25μg 7AAD (BD Biosciences). Flow cytometric analysis was performed on a BD FACS Aria III sorter (BD Biosciences) and all data was analyzed using FlowJo software package (Treestar). See Extended Data Figure 8c for gating strategy used to distinguish DPP4high, DPP4int and DPP4neg cells. Note that DPP4+ cells were defined as DPP4high and DPP4int.

Flow cytometry analysis of leukemia cluster cell surface markers

Bone marrow isolated from leukemic mice was blocked with anti-mouse Fc Block (BD Biosciences, dilution 1 to 50) for 10 minutes at 4°C followed by surface staining with anti-CD45-APC/Cy7 (BD Biosciences, dilution 1 to 100). Samples were washed and stained with LIVE/DEAD fixable viability dye (ThermoFisher) in accordance with the manufacturer’s instructions. The fixed cells were then washed with Perm/Wash buffer and resuspended in Medium 199 + 2% FBS with 0.25μg 7AAD (BD Biosciences). Flow cytometric analysis was performed on a BD FACS Aria III sorter (BD Biosciences) and all data was analyzed using FlowJo software package (Treestar).

Analysis of DPP4 expression following co-culture

250,000 MLL-AF9 or HoxA9-Meis1 leukemia cells were plated at a 1:1 ratio with the following stromal cell lines: NIH3T3 (ATCC), M5-S (Riken), MLO-A5 (Kerafast), MC-3T3-E1 (ATCC), and SV40 immortalized bone marrow stroma (see separate methods section). The cells were grown in RPMI-1640 (Gibco) supplemented with 1% Penicillin/Streptomycin (Gibco), and 10% FBS (Gibco). MLL-AF9 cultures were supplemented with 20 ng/ml SCF, 10 ng/ml IL-3 and 10 ng/ml IL-6 (all cytokines from Peprotech). HoxA9-Meis1 cell cultures were instead grown in 100 ng/ml SCF and 5 ng/ml IL-3. The cells were co-cultured for 3 days. For flow cytometric analysis of DPP4, the co-cultures were trypsinized and subsequently blocked with murine Fc block (dilution 1 to 50) for 10 minutes at 4°C. Surface staining with anti-CD45-APC/Cy7 (dilution 1 to 100) and DPP4-PE (dilution 1 to 20) was then carried out for 30 minutes at 4°C. The cells were then washed and resuspended in Medium 199 + 2% FBS with 0.25μg 7AAD (BD Biosciences) for analysis. Flow cytometric analysis was performed on a BD FACS Aria III sorter (BD Biosciences) and all data was analyzed using FlowJo software package (Treestar). For the analysis of DPP4 mean fluorescence intensity (MFI), each sample was normalized to a corresponding isotype control antibody-stained sample.

Flow cytometry analysis of myeloid lineage markers

Bone marrow collected from C57Bl/6J mice was stained with HSPC cocktail (Table S6) for 45 minutes at 4°C. The cells were then washed and resuspended in Medium 199 + 2% FBS with 0.25μg 7AAD (BD Biosciences). Flow cytometric analysis was performed on a BD FACS Aria III sorter (BD Biosciences) and all data was analyzed using FlowJo software package (Treestar).
Gating strategy

For all gating strategies cells are discriminated based on SSC and FSC followed by selection of live singlets. For gating strategies for individual panels please see corresponding figures.

☒ Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.