The single-cell transcripational landscape of mammalian organogenesis

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Mammalian organogenesis is a remarkable process. Within a short timeframe, the cells of the three germ layers transform into an embryo that includes most of the major internal and external organs. Here we investigate the transcripational dynamics of mouse organogenesis at single-cell resolution. Using single-cell combinatorial indexing, we profiled the transcriptomes of around 2 million cells derived from 61 embryos staged between 9.5 and 13.5 days of gestation, in a single experiment. The resulting ‘mouse organogenesis cell atlas’ (MOCA) provides a global view of developmental processes during this critical window. We use Monocle 3 to identify hundreds of cell types and 56 trajectories, many of which are detected only because of the depth of cellular coverage, and collectively define thousands of corresponding marker genes. We explore the dynamics of gene expression within cell types and trajectories over time, including focused analyses of the apical ectodermal ridge, limb mesenchyme and skeletal muscle.

Most studies of mammalian organogenesis rely on model organisms, and, in particular, the mouse. Mice develop quickly, with just 21 days between fertilization and birth. The implantation of the blastocyst on embryonic day (E) 4.0 is followed by gastrulation and the formation of germ layers on E6.5–E7.5. At the early-somite stages, the embryo transits from gastrulation to early organogenesis, forming the neural plate and heart tube (E8.0–E8.5). In the ensuing days (E9.5–E13.5), the embryo expands from hundreds-of-thousands to over ten-million cells, and concurrently develops nearly all major organ systems. Unsurprisingly, these four days have been intensively studied. Indeed, most genes that underlie major developmental defects can be studied in this window.

The transcripational profiling of single cells (scRNA-seq) represents a promising strategy for obtaining a global view of developmental processes. For example, scRNA-seq recently revealed a large degree of heterogeneity in neurons and myocardiocytes during mouse development. However, although two scRNA-seq atlases of the mouse were recently released, they are mostly restricted to adult organs and do not attempt to characterize the emergence and dynamics of cell types during development.

Single-cell RNA-seq of two million cells

Single-cell combinatorial indexing is a methodological framework involving split-pool barcoding of cells or nuclei. We previously developed single-cell combinatorial-indexing RNA-sequencing analysis (sci-RNA-seq) and applied it to generate 50-fold shotgun coverage of the cellular content of L2-stage Caenorhabditis elegans. A conceptual identical method was recently termed SPLiT-seq. To increase the throughput, we explored more than 1,000 experimental conditions (Extended Data Fig. 1a, b, Methods). The major improvements of the resulting method, sci-RNA-seq3, include: (i) nuclei are extracted directly from fresh tissues without enzymatic treatment, then fixed and stored; (ii) for the third level of indexing, we switched from Tn5 tagging to hairpin ligation; (iii) individual enzymatic reactions were optimized; and (iv) fluorescence-activated cell sorting was replaced by dilution, and sonication and filtration steps were added to minimize aggregation. Even without automation, sci-RNA-seq3 library preparation can be completed through the intensive effort of a single researcher in one week at a cost of less than $0.01 per cell.

We collected 61 C57BL/6 mouse embryos at E9.5, E10.5, E11.5, E12.5 or E13.5, and snap-froze them in liquid nitrogen. Nuclei from each embryo were isolated and deposited in individual wells in four 96-well plates, such that the first index identified the originating embryo of a given cell. As a control, we spiked a mixture of human HEK-293T and mouse NIH/3T3 nuclei into two wells. The resulting sci-RNA-seq library was sequenced in a single Illumina NovaSeq run, yielding 11 billion reads (Fig. 1a, Extended Data Fig. 1c, d).

From one experiment, we recovered 2,058,652 cells from mouse embryos and 13,359 cells from HEK-293T or NIH/3T3 cells (UMI (unique molecular identifier) count ≥ 200). Transcriptomes from human or mouse control wells were overwhelmingly species-coherent (3% collisions), with performance similar to previous experiments (Extended Data Fig. 1e–j). A limitation is that only around 7% of cells entering the experiment were ultimately profiled, with losses largely consequent on filtration steps intended to remove aggregates of nuclei.

We profiled a median of 35,272 cells per embryo (Fig. 1b, Extended Data Fig. 1j). Despite shallow sequencing (about 5,000 raw reads per cell; 46% duplicate rate), we recovered a median of 671 UMIs (519 genes) per cell (Extended Data Fig. 1k). The 3.7-fold-deeper sequencing of a subset of wells nearly doubled the complexity (to a median of 1,142 UMIs per cell; 87% duplicate rate). Given that we are profiling RNA in nuclei, 59% of UMIs per cell strand specifically mapped to introns and 25% mapped to exons. The profiles may therefore primarily reflect nascent transcription, temporally offset, but also predictive of the cellular transcriptome. Later-stage embryos exhibited somewhat reduced UMI counts, possibly reflecting decreasing nuclear mRNA content (Extended Data Fig. 1l). We used Scrublet22 to detect 4.3% likely doublet cells, corresponding to a doublet estimate of 10.3% including both within-cluster and between-cluster doublets (Extended Data Fig. 1m, n).

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On the basis of our rough estimates of the number of cells per embryo at each time point (Methods), our ‘shotgun cellular coverage’ of the mouse embryo is 0.8 × at E9.5 (200,000 cells per embryo; 152,000 profiled across all replicates), 0.5 × at E10.5 (1.1 million cells per embryo; 378,000 profiled), 0.2 × at E11.5 (2.6 million cells per embryo; 616,000 profiled), 0.08 × at E12.5 (6 million cells per embryo; 475,000 profiled) and 0.03 × at E13.5 (13 million cells per embryo; 437,000 profiled). Thus, although we are not yet oversampling17, the depth of profiling is equivalent to an estimated 3–80% of the cellular content of an individual mouse embryo.

Embryos were readily identifiable as male (n = 31) or female (n = 30) (Extended Data Fig. 1o, p). Applying t-distributed stochastic neighbour embedding (t-SNE) to ‘pseudobulk’ profiles (aggregating the transcriptionomes of the cells of each embryo) resulted in five tightly clustered groups corresponding to developmental stages (Extended Data Fig. 1q). We also ordered the mouse embryos along a pseudotime trajectory23 (Fig. 1c). Two prominent gaps (E9.5–E10.5 and E11.5–E12.5) suggest particularly marked changes during these windows (Extended Data Fig. 1r, s). In these pseudobulk profiles, 12,236 genes were differentially expressed across developmental stages (Supplementary Table 1).

**Identification of cell types and subtypes**

We subjected the 2,058,652 single-cell transcriptomes to Louvain clustering and t-SNE visualization (Fig. 2a). Reassuringly, cells from replicate embryos of the same developmental stage were similarly distributed, whereas cells from different stages were not (Extended Data Figs. 2a–f). On the basis of genes specific to each of 40 clusters, we manually annotated cell types (Supplementary Table 2). Merging 2 clusters, both corresponding to the definitive erythroid lineage, and discarding a putative doublet cluster (detected doublet rate of 52%) yielded 38 major cell types (Fig. 2b, Extended Data Fig. 2g).

In general, highly specific marker genes made the annotation of these major cell types straightforward (Fig. 2b, Supplementary Table 3). For example, cluster 6 (epithelial cells) specifically expressed Epcam and Trpc322,24,25, whereas cluster 29 (hepatocytes) specifically expressed Afp and Alb26. Smaller clusters were also readily annotated. For example, cluster 36 (melanocytes) specifically expressed Tyr and Trpm126,27, whereas cluster 37 (lens) specifically expressed Crys4a2. Some markers, although observed in a substantial proportion of cells in many clusters, were much more highly expressed in one cluster (for example, Hbb-bh1 in primitive erythroid cells). For clusters corresponding to the embryonic mesenchyme and connective tissue, annotation was more challenging because fewer markers are known (for example, Fndc3c1 in early mesenchyme; Extended Data Fig. 2h).

Across the major cell types, 17,789 of 26,183 genes (68%) were differentially expressed (5% false discovery rate (FDR); Supplementary Table 4). Among these differentially expressed genes, we identified 2,863 cell-type-specific marker genes—a mean of 75 differentially expressed genes per cell type, defined as genes with more than twofold expression difference between first- and second-ranked cell type (a cut-off of larger than fivefold yielded 932 marker genes; Extended Data Fig. 2i). A large majority of these genes were not previously characterized as markers of the respective cell types. For example, we detect the highest expression of sonic hedgehog (Shh) in the notochord (cluster 30), together with Ntn1, Slit1 and Slit2, which are all known to be expressed in the cells of the notochord and floor plate during development28–31. However, Foxa2, Stxbp6, Schiap1 and Frmd4b, which were not previously described as markers of the notochord, were also markers of cluster 30. Whole-mount in situ hybridization (WISH) of Shh (known) and Foxa2 (novel) confirmed that both genes are expressed in notochord at E10.5 (Extended Data Fig. 2j).

We observed marked changes in the proportions of cell types during organogenesis. Whereas most major cell types proliferated exponentially, a few were transient and disappeared by E13.5 (Extended Data Fig. 2k, l). For example, at E9.5 we detected cells that correspond to the primitive erythroid lineage, originating from the yolk sack (cluster 26; marked by Hbb-bh1). However, the definitive erythroid lineage, originating from the fetal liver (cluster 22; marked by Hbb-bs), progressively replaces the primitive erythroid lineage to become the exclusive red cell lineage by E13.5 (Fig. 2a, Extended Data Fig. 2m).

The 38 major cell types are represented by a median of 47,073 cells, the largest containing 144,648 cells (connective tissue progenitors) and the smallest containing only 1,000 cells (neutrophils). As additional heterogeneity was readily apparent, we adopted an iterative strategy, repeating Louvain clustering on each major cell type. After subclusters dominated by a few embryos were removed and highly similar subclusters were merged (Methods), 655 subclusters were identified (Extended Data Fig. 3). As an operational definition specific to this manuscript, we refer to the 38 major clusters as cell types, and the 655 subclusters as subtypes. Notably, our sensitivity to detect cell types and subtypes in this study was dependent on the large number of cells profiled (Extended Data Fig. 4a–d). The 655 subtypes consist of a median of 1,869 cells, and range from 51 (a subtype of notochord) to 65,894 (a subtype of connective tissue progenitors) cells (Extended Data Fig. 4e–g).

We annotated 13% of subtypes as likely artefacts (more than 10% of cells in these subtypes are predicted doublets; Extended Data Fig. 4h). For the remaining 572 subtypes, we identified a median of 20 subtype-specific markers (more than twofold expression difference between first- and second-ranked cell subtypes of the corresponding major cell type; Extended Data Fig. 4i, j). Furthermore, most subtypes can be distinguished from all 571 other non-doublet subtypes on the basis of marker gene sets and larger-than-fourfold expression differences (63% with 2 markers, 95% with 4 markers; Extended Data Fig. 4k, Supplementary Table 5, Methods).

To distinguish it from other cell atlases, we term the dataset described here as the mouse organogenesis cell atlas (MOCA). As there are presently no comparable single-cell atlases of E9.5–E13.5, we compared MOCA subtypes to 130 fetal cell types (E14.5) of a recent mouse cell atlas (MCA)10. With a new inter-study cross-matching method, we matched 96 MCA cell types to 58 MOCA subtypes (Extended Data Fig. 5a–c, Supplementary Table 5, Methods). As expected, MOCA subtypes that failed to match MCA cell types tended to derive from earlier stages (for example, neural tube) or were rare (for example, lens), whereas MCA cell types that failed to match MOCA subtypes were mostly tissue-specific immune or epithelial cells, potentially because they emerge after E13.5. Nonetheless, the atlases unquestionably inform one another, as the anatomical resolution of the MCA is useful for localizing MOCA subtypes, whereas the developmental focus of MOCA informs the embryonic origin of MCA cell types (Extended Data Fig. 5b). As an example of the former, a subcluster of endocrine epithelial cells in MOCA mapped to both the acinar and endocrine cells of the fetal stomach in the MCA. As an example of the latter, ‘cells in cell cycle’ in the MCA fetal kidney mapped to a subtype of intermediate
Identifying the major cell types of mouse organogenesis.

a, t-SNE visualization of 2,026,641 mouse embryo cells (after removing a putative doublet cluster), coloured by cluster identity (ID) from Louvain clustering (in b), and annotated on the basis of marker genes. The same t-SNE is plotted below, showing only cells from each stage (cell numbers from left to right: n = 151,000 for E9.5; 370,279 for E10.5; 602,784 for E11.5; 468,088 for E12.5; 434,490 for E13.5). Primitive erythroid (transient) and definitive erythroid (expanding) clusters are boxed. b, Dot plot showing expression of one selected marker gene per cell type. The size of the dot encodes the percentage of cells within a cell type in which that marker was detected, and its colour encodes the average expression level.

Characterization of the apical ectodermal ridge

We annotated all subtypes of epithelium and endothelium (clusters 6 and 20, respectively; Fig. 3a, Extended Data Fig. 6a–c, Supplementary Table 2). For example, epithelial subtype 6.8 was marked by Oc90, which is exclusively expressed in the epithelium of the otic vesicle33; epithelial subtype 6.23 was marked by Fgf8, Mxs2 and Rspo2, known markers of the apical ectodermal ridge (AER)34; and endothelial subtype 20.12 was marked by Tbx20 and Tmem108, specific to endocardial cells and cardiac valve endothelium35,36.

To investigate a subtype in greater detail, we focused on the AER, a highly specialized epithelium involved in digit development37. In addition to known markers for AER, subtype 6.23 (1,237 cells; 0.06% of MOCA) was distinguished by expression of Fndc3a, Adamts3, Slc16a10, Snap91 and Pouf62. WISH of Fgf8 (known), Fndc3a, Adamts3 and Snap91 (all novel) confirmed expression specific to the most distal tip of the limb bud, representing the AER at E10.5 or E11.5 (Fig. 3b–e).

We next examined the dynamics of AER proliferation and gene expression. Although detected at all time points and nearly all embryos, the estimated number of AER cells per embryo peaked between E10.5 and E11.5 (Fig. 3f), consistent with a previous report38 and our validations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c). We performed pseudotemporal ordering of AER cells, yielding a simple early-to-late trajectory and 710 differentiations (Fig. 3c).

Reconstructing developmental trajectories

We next sought to investigate the developmental trajectories that cell types traverse during mammalian organogenesis. Most contemporary algorithms for trajectory reconstruction assume a continuous manifold (whereas our data begin at E9.5, and are therefore missing at least some ancestral states) and do not allow for convergence of cell fates (whereas some cell types are known to derive from multiple transcriptionally distinct lineages). To overcome these limitations while also enabling scaling to millions of cells, we developed a new version of Monocle42. Monocle 3 first projects cells onto a low-dimensional space encoding transcriptional state using UMAP43. It then groups mutually similar cells using the Louvain community detection algorithm, and merges adjacent groups into ‘supergroups’44. Finally, it resolves the paths or trajectories that individual cells can take during development, identifying the locations of branches and convergences within each supergroup.

Subsequent to a focused application of Monocle 3 to cells corresponding to the limb bud mesenchyme (Extended Data Fig. 7, Supplementary Note 1, Supplementary Tables 7–9), we applied it to identify major developmental trajectories across the entire dataset. Monocle 3 organized 1,524,703 high-quality cells (UMI greater than 400) into 12 groups. We merged two groups corresponding to sensory neurons, and another two corresponding to blood cells. Nearly all of the 38 major cell types fall almost exclusively in 1 of the 10 resulting trajectories (Fig. 4a, b, Extended Data Fig. 8a, b). The two most complex structures are the neural tube–notochord trajectory, which includes the notochord, neural tube, progenitor and developing neuronal and glial cell types, and the mesenchymal trajectory, which includes all mesenchymal and muscle cell types. There are three neural crest trajectories, corresponding to sensory neurons, Schwann cell precursors and melanocytes. The haemopoietic trajectory includes megakaryocytes, erythrocytes and white blood cells, whereas the remaining four trajectories (endothelial, epithelial, hepatic and lens) each correspond to a single major cell type (Fig. 4b). The discontinuity between these ten major trajectories is likely to reflect the lack of representation of some intermediate or ancestral states, consequent on our study beginning at E9.5. Although the estimated number of cells per embryo in each trajectory increases exponentially, their proportions remain relatively stable, with the exception of hepatocytes, which markedly increase their contribution from 0.3% at E9.5 to 2.8% at E13.5 (Extended Data Fig. 8c).

Unlike t-SNE, UMAP places related cell types near one another. For example, cell types found at later developmental time points, such as progenitors of the kidney. A similar analysis matched 48 cell types annotated in a recent mouse brain atlas (BCA)32 to 68 MOCA subtypes with high specificity (Extended Data Fig. 5d).
as inhibitory neurons, are connected to early central nervous system precursors (radial glia) by a ‘bridge’ of neural progenitor cells; however, the same radial glial cells project in a different direction towards increasingly mature oligodendrocytes (Fig. 4a, top right). Similarly, early mesenchymal cells radiate from a defined region into myocytes, limb mesenchyme, chondrocytes or osteoblasts, and connective tissues (Fig. 4a, bottom right).

After removing 12% of cells corresponding to doublet-annotated cells and/or subclusters, we iteratively reanalysed the 10 major trajectories (Fig. 5, Extended Data Fig. 9). For example, the epithelial trajectory breaks into several discontinuous subtrajectories, each emanating from a focal concentration of E9.5-derived cells and projecting in one or more directions, through cells corresponding to progressively later time points (Fig. 4c, Extended Data Fig. 8d). Notably, the AER subtrajectory projects out of surface ectoderm and then back into epidermis, consistent with its transitory nature.

We mapped the 572 subtypes defined by t-SNE and Louvain clustering to the developmental subtrajectories defined by Monocle 3 (Extended Data Fig. 9). The vast majority of subtypes mapped to a single subtrajectory, often as temporally restricted subsets (Supplementary Table 5). We annotated the subtrajectories on the basis of marker genes of subtypes mapping to them. The resulting 56 developmental subtrajectories span all major systems including the central nervous system, peripheral nervous system, respiratory, digestive, cardiovascular, immune, lymphatic, urinary, endocrine, integumentary, skeletal, muscular and reproductive systems (Fig. 5, Extended Data Fig. 10).

In some cases, we observe a single, simple linear trajectory. However, we also observe many examples of branching trajectories, as well as of cell types that appear to be generated via multiple parallel paths. As an example of the latter, subsets of both excitatory and inhibitory central nervous system neurons appear to develop through multiple, convergent trajectories, possibly owing to their maturation in distinct anatomical locations. Other subtrajectories exhibited even more complex features, including multiple starting and ending points within a continuous structure (for example, intermediate mesoderm trajectory).

Although Monocle 3 did not have access to these labels, the subtrajectories are highly consistent with developmental time (that is, cells ordered from E9.5 to E13.5; Extended Data Figs. 9, 10). To orient subtrajectories, we identified one or several starting points as focal concentrations of E9.5 cells and then computed developmental pseudotime for cells present along various paths (Extended Data Fig. 11, Methods). We also annotated each subtype according to the subtrajectory to which it maps, as well as its relative temporal position within that subtrajectory (for example, subtype 6.14 maps to ‘auditory epithelial trajectory 1 of 3’ (Supplementary Table 5). These representations provide a starting point for more detailed explorations of the 572 subtypes and 56 subtrajectories.

**Reconstructing skeletal myogenesis**

To investigate a developmental process in greater detail, we focused on developing muscle, which comprises distinct mesodermal lineages...
that form before E9.5. We hypothesized that the myogenic trajectory would feature multiple entry points that feed cells into a common path, corresponding to activation of the core gene-expression program shared by myoblasts.

To test this, we isolated myocytes and their putative 'ancestral' cells from the mesenchymal trajectory in silico (Fig. 6a, Methods). Next, we used Monocle 3 to construct a myogenesis-specific trajectory, which shared by myotubes.

Furthermore, because we studied disaggregated whole embryos, the resolution, and many published markers have limited specificity. Development has not previously been extensively studied at single-cell resolution, and some published markers have limited specificity. However, we also observed extremely rare cell types.

The underlying data freely available, together with a website to facilitate their further exploration (http://atlas.gs.washington.edu/mouse-rna). MOCA does have limitations. First, although not sequenced to saturation, the cell-by-gene matrix is sparse. Nonetheless, our results support the view that cell types are readily distinguishable despite having hundreds rather than thousands of UMls per cell. Of course, the tradeoff between breadth and depth depends on one's goals. An example that supports the 'many cells, few UMls per cell' approach can be seen in the form of primordial germ cells, which were readily identifiable despite their rarity (subtypes 16.13 and 6.27, which sum to 269 out of 2,058,652 cells or 0.01% of MOCA). Nonetheless, despite its unprecedented depth, our study does not exceed onefold coverage of the mouse embryo at any time point and it is possible that we are missing extremely rare cell types.

Second, although we are reasonably confident in our annotations, they should be regarded as preliminary. Mid-gestational mouse development has not previously been extensively studied at single-cell resolution, and many published markers have limited specificity. Furthermore, because we studied disaggregated whole embryos, the assignment of anatomical specificity is challenging. We anticipate that the comprehensive annotation of MOCA will benefit from community input and domain expertise, and to that end created an interactive wiki.

**Discussion**

Here, to obtain a global view of mammalian organogenesis, we profiled the transcriptomes of around 2 million cells from mouse embryos spanning E9.5 to E13.5. In the resulting MOCA, we identify over 500 subtypes of cells and 56 developmental subtrajectories, each distinguished by multiple marker genes and collectively spanning essentially every organ system. With sci-RNA-seq, we introduce a technical framework for individual laboratories to generate datasets corresponding to millions of single cells. With Monocle 3, we introduce a computational framework for trajectory inference that operates at this same scale. These data constitute a potentially foundational resource for the field of mammalian developmental biology. We have made MOCA and the underlying data freely available, together with a website to facilitate their further exploration (http://atlas.gs.washington.edu/mouse-rna).

MOCA does have limitations. First, although not sequenced to saturation, the cell-by-gene matrix is sparse. Nonetheless, our results support the view that cell types are readily distinguishable despite having hundreds rather than thousands of UMls per cell. Of course, the tradeoff between breadth and depth depends on one's goals. An example that supports the 'many cells, few UMls per cell' approach can be seen in the form of primordial germ cells, which were readily identifiable despite their rarity (subtypes 16.13 and 6.27, which sum to 269 out of 2,058,652 cells or 0.01% of MOCA). Nonetheless, despite its unprecedented depth, our study does not exceed onefold coverage of the mouse embryo at any time point and it is possible that we are missing extremely rare cell types.

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**Fig. 6 | Resolving cellular trajectories in myogenesis.** Edges in the principal graphs that define trajectories reported by Monocle 3 are shown as light blue line segments. a. Cells putatively involved in myogenesis were isolated from the mesenchymal cell trajectory in silico and then used to construct a myocyte subtrajectory. Principal graph nodes that are more than 50% occupied by cells from cluster 13 were taken as 'seed nodes', and then cells on any nodes within 20 edges of these seed nodes were selected for subtrajectory analysis. Cells in the myocyte subtrajectory (left), coloured by developmental stage (right), b. Cells in the myocyte trajectory, coloured by their expression of selected transcriptional regulators of myogenesis. Cells with no detectable expression for a given gene are omitted from its plot. Values are log-transformed, standardized UMI counts. c. Cells classified by developmental stage according to the markers shown in c (dermomyotome: Pax3, Pax7; muscle progenitors: Pax7; myoblasts: Myf5 or Myod and Myog; myocytes: Myog; myotubes: Myh3).

**Fig. 5 | UMAP visualization of individual major trajectories.** After removing doublet-annotated cells and subclusters, we iteratively reanalysed each of the ten major trajectories. Charts are coloured by subtrajectory name (main plots) or developmental stage (insets, colours as in Fig. 4c). Edges in the principal graphs that define trajectories reported by Monocle 3 are shown as light blue line segments. PNS, peripheral nervous system.
Effective, however, additional experiments (such as in situ analyses of marker genes) will be necessary to resolve ambiguities. Notably for future atlasing efforts, we found the annotation of temporally resolved developmental trajectories to be much more straightforward than that of cell types.

A long-standing ambition, which is perhaps finally within sight from a technical perspective, is to construct a comprehensive, spatiotemporally resolved molecular atlas of mammalian development at single-cell resolution. To this end, the mouse has several advantages, including its small size, the accessibility of early developmental time points, an inbred genetic background and genetic manipulability. It also seems likely that ‘whole-organism’ profiling of small mammals will be essential for identifying the inevitable gaps in any efforts to generate a comprehensive atlas of human cell types.

Single-cell atlases of the development of wild-type mice may also represent an important step towards understanding pleiotropic developmental disorders at the organisinal scale, and for detailed investigations of the roles of specific genes and regulatory sequences in development. For example, many knockouts of both coding and conserved regulatory sequences do not exhibit any abnormalities with conventional pheno-typing44. We anticipate that whole-organism single-cell transcriptional atlases will empower reverse genetics—for example, potentially enabling the discovery of subtle defects in the molecular programs or the relative proportions of specific cell types45.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at https://doi.org/10.1038/s41586-019-0969-x

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Competing interests L.C., F.Z. and F.J.S. declare competing financial interests in the form of stock ownership and paid employment by Illumina. One or more embodiments of one or more patents and patent applications filed by Illumina may encompass the methods, reagents and data disclosed in this manuscript. Some work in this study may be related to technology described in the following exemplary published patent applications: WO2010/0120098 and WO2011/0287435.

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METHODS

Data reporting. No statistical methods were used to predetermine sample size. Embryos used in experiments were randomized before sample preparation. Investigators were blinded to group allocation during data collection and analysis: embryo collection and sci-RNA-seq analysis were performed by two different researchers.

Embryo dissection. The C57BL/6 mice were obtained from The Jackson Laboratory and plug matings were set up. Noon on the day of the vaginal plug was considered as E0.5. Dissections were performed as previously described51 and all embryos were immediately snap-frozen in liquid nitrogen. Embryos were collected from at least three independent litters per development stage. All animal procedures were in accordance with institutional, state, and government regulations and approved by the Office of Animal Welfare (OAW) under the IACUC protocol 45378-01.

WISH. The mRNA expression in E9.5–E13.5 mouse embryos was assessed by WISH using a digoxigenin-labelled antisense riboprobe transcribed from cloned gene-specific probes (PCR DIG Probe Synthesis Kit, Roche). Whole embryos were fixed overnight in 4% paraformaldehyde (PFA) in PBS. The embryos were washed in PBS with 0.1% Tween, and dehydrated stepwise in 25%, 50% and 75% methanol/PBST and finally stored at −20 °C in 100% methanol. The WISH protocol was as follows. Day 1: embryos were rehydrated on ice in reverse methanol/PBST steps, washed in PBS-Tween, bleached in 6% H2O2 in PBST for 1 h and washed in PBS-Tween. Embryos were then treated in 10 µg/ml proteinase K in PBS-Tween for 3 min, incubated in glycine in PBS-Tween, washed in PBS-Tween and finally re-fixed for 20 min with 4% PFA in PBS, 0.2% glutaraldehyde and 0.1% Tween. After further washing steps with PBS-Tween (embryos were washed at 68 °C in L1 buffer (50% deionised formamide, 5× SSC, 1% SDS, 0.1% Tween-20 in diethyl pyrocarbonate (DEPC); pH 4.5) for 10 min. Next, embryos were incubated for 2 h at 68 °C in hybridization buffer 1 (L1 with 0.1% RNA and 0.05% heparin). Subsequently, embryos were incubated overnight at 68 °C in hybridization buffer 2 (hybridization buffer 1 with 0.1% RNA, 0.05% heparin and 1× digoxigenin (DIG) probe). Day 2: unbound probe was removed through a series of washing steps: 3× 30 min each at 68 °C with L1, L2 (50% deionised formamide, 2× SSC pH 4.5, 0.1% Tween 20 in DEPC; pH 4.5) and L3 (2× 5× SSC pH 4.5, 0.1% Tween-20 in DEPC, pH 4.5). Subsequently, embryos were treated for 1 h with RNAse solution (0.1 M NaCl, 0.01 M Tris pH 7.5, 0.2% Tween-20, 100 µg/ml RNAse A in H2O), followed by washing in TBST (140mM NaCl, 2.7mM KCl, 25mM Tris-HCl, 1% Tween-20; pH 7.5). Next, embryos were blocked for 2 h at room temperature in blocking solution (TBST 1 with 2% calf-serum and 0.2% bovine serum albumin (BSA)), followed by incubation at 4 °C overnight in blocking solution containing 1:5000 digoxigenin antibody conjugated to alkaline phosphatase (Roche, catalogue no. 11903274910). Day 3: unbound antibody was removed through a series of washing steps: 8× 30 min at room temperature with TBST 2 (TBST with 0.1% Tween-20, 0.5% deionised tetrathiomolybdate and 0.1 mg/ml levamisole) and left overnight in RNase-free water, 1× RNase solution (0.1 M NaCl, 0.01 M Tris pH 7.5, 0.2% Tween-20, 20 µg/ml RNase A and 50 µg/ml RNase B, followed by staining with BM Purple AP Substrate (Roche). The stained embryos were imaged using a Zeiss Discovery V12 microscope and Leica DFC420 digital camera.

Mammalian cell culture. All mammalian cells were cultured at 37 °C with 5% CO2 and were maintained in high glucose DMEM (Gibco 11965) for HEK-293T (from ATCC) and NIH/3T3 (a gift from T. Reh, University of Washington, ATCC cell line) cells, supplemented with 10% FBS and 1× penicillin-streptomycin (Gibco 15140122, 100 U/ml penicillin, 100 µg/ml streptomycin). Cells were trypsinized with 0.25% trypsin-EDTA (Gibco 25200-056) and split in a 1:10 ratio 3 times a week. Both cell lines tested negative for mycoplasma.

Mouse embryo nuclei extraction and fixation. Mouse embryos from different development stages were processed together to reduce batch effects. Each mouse embryo was minced into small pieces by blade in 1 ml ice-cold cell lysis buffer (10 mM Tris-HCl, pH 7.4, 10 mM NaCl, 3 mM MgCl2 and 1% BSA (molecular biology grade, NEB, 20mg/ml)) was added into each well. Nuclei from all wells were pooled together and spun down at 500g for 10 min. Nuclei were then resuspended in nuclei wash buffer and redistributed into another four 96-well plates with each well including 4 µl T4 ligation buffer (NEB), 2 µl T4 DNA ligase (NEB), 4 µl betaine solution (5 M, Sigma-Aldrich), 6 µl nuclei in nuclei wash buffer, 8 µl digoxigenin labelled adaption (100 µM, 5′-GCTGCTG[9-bp or 10-bp barcode]A/dideoxyU/AACAGGCCTCCCATGACT[reverse complement of barcode A]-3’) and 16 µL 40% PEG 8000 (Sigma Aldrich). The ligation reaction was done at 16°C for 3 h.

After the ligation reaction, 60 µl nuclei dilution buffer (10 mM Tris-HCl, pH 7.4, 10 mM NaCl, 3 mM MgCl2 and 1% BSA (molecular biology grade, NEB, 20mg/ml)) was added into each well. Nuclei from all wells were pooled together and spun down at 600g for 10 min. Nuclei were washed once with nuclei wash buffer and subsequently transferred to 8 96-well plates with each well including 2,500 nuclei in 5 µl nuclei wash buffer and 5 µl elution buffer (Qiagen). mRNA Second Strand Synthesis buffer (1.33 µL, NEB) and 0.66 µl mRNA Second Strand Synthesis enzyme (NEB) were then added to each well, and second strand synthesis was carried out at 16°C for 180 min. For tagmentation, each well was mixed with 11 µl Nextera TD buffer (Illumina) and 1 µl i7 only TDE1 enzyme (62.5 nM, Illumina, diluted in TD buffer) and then incubated at 55°C for 5 min to carry out tagmentation. The reaction was then stopped by adding 24 µl DNA binding buffer (Zymo) per well and incubating at room temperature for 5 min. Each well was then purifed using 1.5× AMPure XP beads (Beckman Coulter). In the elution step, each well was added with 8 µl nuclease-free water, 1 µl 10× USER buffer (NEB), 1 µl USER enzyme (NEB) and incubated at 37°C for 15 min. Another 6.5 µl elution buffer was added to each well. The AMPure XP beads were removed by magnetic stand and the elution product (16 µl) was transferred to a 1× well plate.

For PCR amplification, each well (16 µl product) was mixed with 2 µl of 10 µM indexed P5 primer (5′-ATAGTGACGGGACACCAAGGATCTACAC[5S] ACACCTTTCCCCTACAGCGGCTTTCCGATCT-3′; IDT), 2 µl of 10 µM P7 primer (5′-CAAAGGAGAACGCGGCCAGGAT-17′ GTCTCGTGGGCTCGG-3′), and 20 µl NEBNext High Fidelity 2x PCR Master Mix (NEB). Amplification was carried out using the following program: 72°C for 5 min, 98°C for 30 s, 12–14 cycles of (98°C for 10 s, 66°C for 30 s, 72°C for a minute) and a final 72°C for 5 min.

For 5′ linker purification, samples were pooled and purified using 0.8 volumes of AMPure XP beads. Library concentrations were determined by Qubit (Invitrogen) and the libraries were visualized by electrophoresis on a 6% Tris-buffered EDTA polyacrylamide gel electrophoresis gel. All libraries were sequenced on one NovaSeq
A more detailed version of the sci-RNA-seq protocol is provided at our website (http://atlas.gs.washington.edu/mouse-rna).

Processing of sequencing reads. Base calls were converted to fastq format using Illumina's bc2fastq/v.2.16 and demultiplexed based on PCR i5 and i7 barcodes using the maximum likelihood demultiplexing package deML.17 with default settings. Downstream sequence processing and single-cell digital-expression matrix generation were similar to sci-RNA-seq15 except that reverse transcription index was combined with hairpin adaptor index, and thus the mapped reads were split into constituent cellular indices by demultiplexing reads using both the reverse transcription index and ligation index (Levenshtein edit distance (ED) < 2, including insertions and deletions). In brief, demultiplexed reads were filtered on the basis of the reverse transcription index and ligation index (ED < 2, including insertions and deletions) and adaptor-clipped using trim_galore v.0.4.1 with default settings. Trimmed reads were mapped to the mouse reference genome (mm10) for mouse embryo nuclei, or a chimera reference genome of human hg19 and mouse mm10 for HEK-293T and NIH/3T3 mixed nuclei, using STAR v.2.5.2b46 with default settings and gene annotations (GENCODE V19 for human; GENCODE VM11 for mouse). Uniquely mapping reads were extracted, and duplicates were removed using the unique molecular identifier (UMI) sequence, reverse transcription index, hairpin ligation adaptor index and read 2 end-coordinate (that is, reads with identical UMI, reverse transcription index, ligation adaptor index and tagmentation site were considered duplicates). Finally, mapped reads were split into constituent cellular indices by further demultiplexing reads using the reverse transcription index and ligation hairpin (ED < 2, including insertions and deletions). For mixed-species experiment, the percentage of uniquely mapping reads for genomes of each species was calculated. Cells with over 85% of UMIs assigned to one species were regarded as species-specific cells, with the remaining cells classified as mixed cells or ‘‘collisions.’’ To generate digital expression matrices, we calculated the number of strand-specific UMIs for each cell mapping to the exonic and intronic regions of each gene with the Python v.2.7.13 HTseq package16. For multi-mapped reads, reads were assigned to the closest gene, except in cases in which another intersected gene fell within 100 bp of the end of the closest gene, in which case the read was discarded. For most analyses, we included both expected-strand intronic and exonic UMIs in per-genome single-cell expression matrices.

Because of the marked increase in processing time that it would entail, we note that we did not perform a UMI error-correction step. However, to confirm that our failure to do so would not inflate UMI counts, we compared results with and without UMI error correction (edit distance of 1) for a subset of wells. Compared with skipping UMI error correction, 99.4% of reads remain after UMI error correction, which indicated to us that the error correction step has only a minor effect on the estimated UMI counts per cell (less than 1%). This is likely to be a result of the high quality of sequencing data that we obtained on the N nov sSeq, the low number of PCR amplification steps and the low duplication rate. We emphasize that groups implementing sci- RNA-seq should either perform UMI error correction or a similar data quality check.

Whole mouse embryo analysis. After the single-cell gene count matrix was generated, each cell was assigned to its original mouse embryo on the basis of the reverse transcription barcode. Reads mapping to each embryo were aggregated to generate ‘‘bulk RNA-seq’’ for each embryo. For sex separation of embryos, we counted reads mapping to a female-specific non-coding RNA (Xist) or chrY genes (except Eret which is in both chrX and chrY). Embryos were readily separated into females (more reads mapping to Xist than chrY genes) and males (more reads mapping to chrY genes than Xist).

Pseudotemporal ordering of whole-mouse embryos was done using Monocle 246. In brief, an aggregated gene-expression matrix was constructed as described above. Differentially expressed genes across different development conditions were identified with the differentialGeneTest function of Monocle 2.4. The top 2,000 genes with the lowest q value were used to construct the pseudotime trajectory using Monocle 2.46. Each embryo was assigned a position along the trajectory.

Cell clustering, t-SNE visualization and marker-gene identification. A digital gene expression matrix was constructed from the raw sequencing data as described above. Cells with fewer than 200 UMIs or over 3,172 UMIs (two standard deviations above the mean UMI count) were discarded. Downstream analyses were performed with Monocle2 v.2.6.034 and the Python package scanny v.1.0.10.1. In brief, gene-count mappings to sex chromosomes were removed before clustering and dimensionality reduction. Preprocessing steps were similar to a previous approach27. In brief, genes with no count were filtered out and each cell was normalized by dividing the gene expression count per cell-by-total count per row. Genes with low gene expression counts were selected and the digital gene-expression matrix was renormalized after gene filtering. The data were log-transformed after adding a pseudocount, and scaled to unit variance and zero mean. The dimensionality of the data was reduced by principal component analysis (PCA) (30 components) first and then with t-SNE, followed by Louvain clustering performed on the 30 principal components (resolution = 1.5). For Louvain clustering, we first fitted the top 30 principal components to compute a neighbourhood graph of observations with local neighbourhood number of 15 using the scanpy.api.pp.neighbours function in scanpy v.1.0.9. We then clustered the cells into sub-groups using the Louvain algorithm implemented as the scanpy.api.tl.louvain function26. For t-SNE visualization, we directly fit the PCA matrix into the scanpy.pp.tl.time function26 with perplexity of 50 and n_components of 2. For t-SNE, 40 clusters were identified. We then sampled 1,000 cells from each cluster and differentially expressed genes across different clusters were identified with the differentialGeneTest function of Monocle 2 v.2.6.033. Genes specific to each cluster were identified as previously described27. Clusters were assigned to known cell types on the basis of cluster-specific markers (Supplementary Table 3). One cluster had abnormally high UMI counts but no strongly cluster-specific genes, which suggested that it was a technical artefact of cell doublets—it was therefore removed. This was confirmed upon analysis for doublets with scrublet (see next paragraph). Another two clusters appeared to correspond to the definitive erythroid lineage and were merged. Consensus expression profiles for each cell type were constructed as previously described27. Differentially expressed genes across cell types were identified with the differentialGeneTest function of Monocle 2 v.2.6.033. To identify cell-type-specific gene markers, we selected genes that were differentially expressed across different cell types (FDR of 5%, likelihood ratio test) and also with a more than twofold expression difference between first- and second-ranked cell types.

For the detection of potential doublet cells, we first split the dataset of ~2 million cells into four equal groups and then applied the scrublet v.0.1. pipeline46 to each subset with parameters (min_count = 3, min_cells = 3, vscore_percentile = 85, n_pc = 30, expected_doublet_rate = 0.06, sim_doublet_ratio = 2, n_neighbours = 30, scaling_method = ‘‘log’’) for doublet score calculation. Cells with doublet score over 0.25 are annotated as detected doublets. We detected 4.3% potential doublet cells in the whole dataset, which corresponds to an overall estimated doublet rate of 10.3% (including both within- and between-cluster doublets). The aforementioned major cluster with abnormally high UMI counts but no strongly cluster-specific genes (see previous paragraph) had a high detected doublet proportion (52%), confirming it as a doublet-related artefact. For detection of doublet-derived subclusters, we redid the above analysis on the whole dataset after removing the doublet-derived main cluster. Subclusters with detected doublet proportion of >10% were annotated as doublet-derived subclusters.

For subcluster identification, we selected high-quality cells (UMI >400) in each major cell type and applied PCA, t-SNE and Louvain clustering similarly to the major cluster analysis. Subclusters were filtered out if most cells (>50%) of the cluster derived from a single embryo. Highly similar subclusters were merged if their aggregated transcriptomes were highly correlated (Pearson correlation coefficient >0.9) and had an identity of >75% with one another. Genes differentially expressed across subclusters were identified for each major cell type as described above. Subclusters with a detected doublet ratio (by scrublet) over 10% are annotated as doublet-derived subclusters.

To identify a distinguishing set of gene markers for each of the 572 subclusters (those of the 655 with detected doublet-cell ratio ≤10%, we used the following algorithm. (1) We selected genes detected in at least 5% of cells in the target subcluster. (2) From these, we identified genes with a more than fourfold greater expression in the target cluster than in all 571 other subclusters. (3) If there was no such gene, the algorithm tried to identify a gene (marker A) such that subclusters with low expression of marker A (less than 25% of its expression in the target cluster) are therefore distinguished from the target cluster on the basis of this difference, and are therefore removed from the comparison set. Gene marker A is selected to maximize the number of subclusters removed from the comparison set. (4) To identify markers that separate the target cluster from the remaining subclusters, we repeat steps 2–3 until a marker with a more-than-fourfold difference in expression between the target cluster and all remaining subclusters is identified. The set of markers identified through this heuristic is sufficient to distinguish the target subcluster from all 571 other non-doublet subclusters on the basis of more-than-fourfold difference in expression.

To identify correlated cell types between two cell atlas datasets, we first aggregated the cell-type-specific UMI counts, normalized by the total count, multiplied by 100,000 and log-transformed after adding a pseudocount. We then applied non-negative least-squares (NNLS) regression to predict the gene expression of target cell type (T_A) in dataset A with the gene expression of all cell types (M_i) in dataset B: \[T_A = \beta_0 + \beta_{i} M_i\] in which \(T_A\) and \(M_i\) represent filtered gene expression for target cell type from dataset A and all cell types from dataset B, respectively. To identify cell types that are specific to one species, we identified each target cell type by (1) ranking genes on the basis of the expression fold-change between the target cell type versus the median expression across all cell types, and then selecting the top 200 genes; (2) ranking genes on the basis of the expression
fold-change between the target cell type versus the cell type with maximum expression among all other cell types, and then selecting the top 200 genes; and (3) merging the gene lists from steps (1) and (2). $\beta$ is the correlation coefficient computed by NNLS regression. $X_0$ is the intercept of the regression line with the y axis when $X_d$ is zero.

Similarly, we then switch the order of datasets A and B, and predict the gene expression of target cell type ($T_d$) in dataset B with the gene expression of all cell types ($M_d$) in dataset A: $T_d = \lambda_0 + \lambda_0 M_d$. UMAP, a recently proposed algorithm based on Riemannian geometry and algebraic topology to perform dimension reduction and data visualization21. Its visualization quality is competitive with the popular t-SNE method used widely in single-cell transcriptomics. However, where t-SNE mainly aims to place highly similar cells in the same regions of a low-dimensional space, UMAP also preserves long-range topological relationships. The UMAP algorithm itself is also more efficient (the algorithm complexity of UMAP is O(N) versus O(Nlog(N)) for t-SNE). In brief, UMAP first constructs a topological representation of the high-dimensional data with local manifold approximations and patches together their local fuzzy simplicial set representations. UMAP then optimizes the lower-dimension embedding, minimizing the cross-entropy between the low-dimensional representation and the high-dimensional one.

The computational efficiency of UMAP markedly accelerated the analysis of the mouse embryo data. We found that UMAP finished processing the two-million-cell dataset in around 3 CPU hours whereas t-SNE took more than 64 CPU hours. A few implementation details lead to the effectiveness of UMAP. Two major steps are involved in both the UMAP and t-SNE algorithms: first, the preprocessing step before UMAP is similar to Monocle 2. In brief, genes expressed in fewer than 10 cells (or fewer than 5 cells in datasets with fewer than 1,000 cells) were filtered out. The digital gene-count matrix was first normalized by cell-specific size factor estimated by the estimateSizeFactors function in Monocle 3, log-transformed after adding a pseudocount, and then scaled to unit variance and zero mean. The top five genes most highly expressed (2,000 genes for datasets with fewer than 5,000 cells, 300 genes for datasets with fewer than 1,000 cells) were selected. The matrix was then projected into 50 top principal components (30 top principal components for trajectory analysis of the 10 supergroups, 10 top principal components for datasets with fewer than 5,000 cells and 5 top principal components for datasets with fewer than 1,000 cells) by partial singular value decomposition. Thus, an intermediate structure from the high-dimension space (here, we used the top 50 principal components constructed from the 5,000 most highly dispersed genes) is built and then a low-dimensional embedding is found to represent the intermediate structure. For the second step, both methods used the stochastic grid descent approach with differing loss functions to embed the data into low-dimension space. Whereas t-SNE needs a loss function for global normalization, UMAP uses a different objective function that avoids this need. This step essentially enables UMAP to scale linearly with the number of data samples.

Dimensionality reduction was implemented with the reduceDimension() function in Monocle 3. This function calls the UMAP v0.3.2 Python implementation (https://github.com/mlcinnes/umap) from L. McInnes and J. Healy through the reticulate v1.10 package (https://cran.r-project.org/web/packages/reticulate/index.html). To process all the cells together, we set UMAP parameters as follows: (n_neighbors = 50, min_dist = 0.01, cosine distance metric). To more finely resolve subtrajectories, we adjusted these as follows: (n_neighbors = 15, min_dist = 0.1, cosine distance metric).

**Partioning cells into discontinuous trajectories.** A recent approach44 has proposed organizing single-cell transcriptome data into a partitioned approximate graph abstraction (PAGA) that relates clusters of cells that might be developmentally related to one another. In brief, this algorithm constructs a k-nearest neighbour graph on cells and then identifies ‘communities’ of cell via the Louvain method, similar to previous methods for analysing CyTOF or single-cell RNA-seq data44. PAGA then constructs a graph in which the vertices are Louvain communities. Two vertices are linked with an edge in the PAGA graph when the cells in the respective communities are more similar to each other than they would be expected under a simple binomial model44. Similar methods were also recently developed and applied in analysing zebrafish and Xenopus cell-atlas datasets56.

Monocle 3 draws from these ideas, first constructing a k-nearest neighbour graph (k = 20) on cells in the UMAP space, then grouping them into Louvain communities, and testing each pair of communities for a significant number of links between their respective cells. Those communities that have more links than expected under the null hypothesis of spurious linkage (FDR < 1%) remain connected in the PAGA graph, and those links that fail this test are severed. The resulting PAGA graph will have one or more components, each of which is passed to the next step (learning the principal graph) as a separate group of cells that will be organized in a trajectory. The PAGA algorithm essentially stops at this stage, presenting the PAGA graph as a kind of coarse-grained trajectory in each community reflects a different state that cells can adopt as they develop. By contrast, as described in the section below, Monocle 3 uses the PAGA graph to constrain the space of principal graphs that can form the final trajectory. That is, Monocle 3 uses the coarse-grained PAGA graph to learn a fine-grained trajectory.

Monocle 3’s implementation of the above procedures (in the partitionCells() function) scales to millions of cells. In brief, it uses the clustering_louvain function from the igraph package to perform community detection. Next, the core PAGA calculations44 are computed via a series of sparse matrix operations. Let $X$ be a (sparse) matrix representing the community membership of the cells. Each column of X represents a Louvain community and each row of X corresponds to a particular cell. $X_{ij}$ is 1 if cell $i$ belongs to Louvain community $j$, otherwise $X_{ij} = 0$. We can further obtain the adjacency matrix $A$ of the k-nearest neighbour graph used to perform the Louvain clustering in which $A_{ij} = 1$ if cell $i$ connects to $j$ in the k-nearest neighbours graph. The adjacency matrix $A$ is then multiplied by $M$, the number of cells in each cluster. The expression of the above calculation is: $M = X^\top A X$. $X$ is the transpose of matrix $X$.

Once $M$ is constructed, we can then follow supplementary note 3.1 from ref. 44 to calculate the significance of the connection between each Louvain clustering...
and consider any clusters with $P$ value larger than 0.05 by default as not disconnected.

**Learning the principal graph.** Monocle 3 learns a principal graph (via the learnGraph() function) that resides in the same low-dimensional space as the data to represent the possible paths cells can take as they develop. Monocle uses a principal graph-embedding procedure that is based on the SimplePPT algorithm\(^2\),\(^3\),\(^4\),\(^5\), with several key enhancements that accelerate graph embedding, support large datasets, allow for loops, and smooth the graph to eliminate noisy branches.

The first enhancement is that Monocle 3 learns the principal graph in the (by default, three-dimensional) UMAP space using a fast reduced-representation approach to avoid dealing directly with many thousands of cells. It first selects a set of ‘landmark’ cells using by first running the kmeans() clustering algorithm in R with $k$ equal to the value of the ncenter argument, which can be passed to learnGraph() by the user. The landmark cells are then selected by first mapping each cell to its nearest kmeans point, and then selecting the cell for each kmeans point with the highest local density. By default, Monocle 3 uses a data-dependent policy for adjusting ncentre automatically\(^2\),\(^3\),\(^4\). Here, unless otherwise specified, we override the default policy and use ncentre $= 2,000$ in the analyses of the embryonic data. Monocle 3 will then learn a principal graph within these landmark cells rather than the full dataset to accelerate the optimization. Running time and fine detail in the trajectory will depend on the number principal graph nodes; more nodes generally result in a more accurate tree but at increased running time.

The second enhancement is a procedure to smooth and refine the principal graph to exclude small branches. To capture smaller fine details of a trajectory such as complex branching architecture, SimplePPT requires that the principal graph contain hundreds of thousands of principal graph nodes. Consequently, the principal tree reported by SimplePPT often contains very small branches to which a very small percentage of cells project. Although SimplePPT does provide tuning parameters that control graph smoothness to a certain extent, we have found that a simple heuristic pruning procedure is effective and easier for users to understand how to control. The procedure operates via a depth-first visitation of the graph nodes in the principal tree. At nodes with degree $\leq 2$, no action is taken. For nodes with degree $> 2$, the diameter path for each subtree rooted at a neighbour not yet visited in the search is computed. If the path is less than a user-specified length (by default, 10 principal-tree nodes), the whole subtree is pruned.

The third major enhancement is that Monocle 3 can learn principal graphs with loops instead of requiring that the trajectory be a tree. This is achieved by augmenting the principal tree reported by SimplePPT with additional edges meant to close loops in the trajectory. The algorithm considers adding an edge between two leaf nodes $a$ and $b$ in the principal graph if the pair meet several criteria. The first criteria is that the geodesic distance between $a$ and $b$ along the principal tree should be at least a certain minimum distance (by default, one third of the tree’s diameter path). That is, when the nodes are close in (Euclidean) UMAP space, but distant in the graph, they should be linked. The second criteria is that they should not be linked if doing so would create an especially long edge. By default, $a$ and $b$ cannot not be farther apart in UMAP space than the longest edge in the principal tree. The third criteria is based on the same test of connectivity used when partitioning the cells: consider leaf nodes $a$ and $b$, which serve as proxies between two clusters of cells (those for which $a$ and $b$ are their nearest $k$-nearest). If cells near $a$ have an unexpectedly high number of cells near $b$ among their $k$-nearest neighbours ($P < 0.05$ by default), then learnGraph() will link $a$ and $b$ in the principal graph; this also extended the two criteria discussed above are also met.

For analysis of the 10 major trajectories, we used ncentre $= 5,000$ for neural tube–notochord trajectory, and ncentre $= 2,000$ for epithelial and mesenchymal trajectories. For the other supergroups, we used ncentre $= (\text{number of cells})/25$ and minimal_branch_len $= 20$. For analysis of the 56 subtrajectories, we mostly used ncentre $= (\text{number of cells in the trajectory})/30$ (2,000 maximum), and minimal_branch_len $= 20$. Each subtrajectory was manually checked and the parameters (ncentre and minimal_branch_len) for about a quarter of these were adjusted, mostly to further prune branches such that the principal graph follows cell transition path from early to late development stages.

The principal graph offers users a means of selecting subsets of cells that lead to particular lineages for further analysis. For example, to isolate cells leading to the myocyte fate, we first quantified the fraction of cells at each principal graph node that were classified as myocytes (cluster 13). From all majority myocyte nodes, we then used the edges of the principal graph to expand this set of nodes into a wider network of cells.

**Computing pseudotimes.** To calculate cell-wise pseudotime, we developed a projection strategy that is applicable to datasets with millions of cells. This strategy works by constructing a graph $\psi$ on all cells using the principal graph as a guide, and then computing the geodesic distance back to one or more user-selected root nodes in the trajectory. In more detail, we first map each cell to its nearest principal point based on Euclidean distance in the UMAP space. Then, for each principal graph edge, we retrieve all the cells that map to its endpoints $a$ and $b$. Next, we orthogonally project each cell to the nearest point on the principal graph edge as previously described\(^2\), so that each cell $c$ can be ordered along the edge according to its projection $p(c)$. Without loss of generality, we suppose this order is $a < p(c) < b$. We then add edges $(a, c)$ and $(c, b)$ to $\psi$. If $c$ and $b$ are in the same Louvain component or connected Louvain components (as determined during partitionCells), we also add $(c, b)$ to $\psi$. Given $\psi$ and a set of user-specified parameters (ncentre and minimal_branch_len), we can then assign pseudotime values to all cells. Monocle provides several ways to specify these nodes, either by name (that is, programmatically) or interactively. Each cell’s pseudotime is taken as the geodesic distance along $\psi$ to the closest of these root nodes.

For root-node selection of the mesenchymal and neural tube–notochord trajectories, we first assigned each principal point to a subcluster with the maximum cell proportion. We then selected the subcluster with the earliest average developmental stage, and use the earliest principal point assigned to this subcluster as the root state for pseudotime computation. For the other major trajectories, we assigned root nodes to the earliest principal point in each subtrajectory (except in neural crest trajectory 2, where we assigned the root node to the earliest principal points in PNS glia precursor cell trajectory and Pdgfra-positive glia trajectory). Some cells from complex trajectories (mesenchymal trajectory, neural tube–notochord trajectory, epithelial trajectory and endothelial trajectory) show outlier pseudotime values (more than three standard deviations higher than the mean values). These extreme values are clipped to the maximum value after excluding the outliers. For root node selection of the 56 cell-type-specific trajectories, we first computed the average development stage for each principal point. As the root state features the earliest development stage, we compared the average development stage of each node and its $k$-nearest neighbours ($k = 10$). We then manually checked each trajectory and selected root nodes from principal points with earlier development stage than all its nearby neighbours.

**Identifying genes with complex trajectory-dependent expression.** To identify genes that vary in expression over a developmental trajectory, we borrow a statistical test commonly used in analysing spatial data. Moran’s I statistic is a measure of multi-directional and multi-dimensional spatial autocorrelation. The statistic encodes spatial relationships between data points via a nearest-neighbour graph, making it particularly well-suited for analysing large scRNA-seq datasets. Moran’s $I$ test\(^6\) is defined as:

$$I = \frac{N}{W} \sum_{i,j} \sum_{W_{ij}} \frac{(x_i - \bar{x})(x_j - \bar{x})}{W_{ij}}$$

in which $N$ is the number of cells indexed by $i$ and $j$, $x$ is the expression value of gene $x$, $\bar{x}$ is the mean of the gene expression for the nearest neighbours of cell $(i)$ or $(j)$; $w_{ij}$ is a matrix of weights defined by a nearest-neighbour graph with zero on the diagonal (that is, $w_{ii} = 0$) and $w_{ij} = 1/ki$, in which $k$ is the number of nearest neighbours; and $W$ is the sum of all $w_{ij}$.

To identify the nearest neighbours used for creating the weight matrix $W$, we first build a $k$ (default to be 25) nearest-neighbour graph for all cells in the UMAP space. We also project each cell to its nearest node in the principal graph. Then, we remove all edges from the $k$-nearest-neighbour graph that connect cells that project onto principal graph nodes that do not share an edge.

In Monocle 3, we implemented the principalGraphTest() function to identify correlated genes on the complex trajectory embedded in the manifold, which relies on modified versions of routines from the spdep package for performing the Moran’s $I$ test.

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

**Code availability.** Scripts for processing sci-RNA-seq3 sequencing were written in Python and R and code available at https://github.com/junyuCe/sci-RNA-seq3 pipeline. Trajectory analysis was done with Monocle 3 with setup instructions and tutorial available at http://code.trapnell-lab.github.io/monocle-release/monocle3.

**Data availability.** The sci-RNA-seq3 protocol and all data have been made freely available, including through a cell-type wiki to facilitate their ongoing annotation by the research community (http://atlas.gs.washington.edu/mouse-rna). The data generated in this study can be downloaded in raw and processed forms from the NCBI Gene Expression Omnibus under accession number GSE119945.

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Extended Data Fig. 1 | Performance and quality-control-related analyses for sci-RNA-seq3. a, Comparison of fixation conditions in human HEK-293T cells. PFA-fixed nuclei yielded the highest numbers of UMIs. Cell number: \( n = 21 \) for fresh nuclei, 17 for frozen nuclei, 32 for PFA-fixed cells and 31 for PFA-fixed nuclei. b, Tn5 transposomes loaded only with N7 adaptor (cell number, \( n = 13 \) cells) increased UMI counts by over 50%, relative to the standard Nextera Tn5 (cell number, \( n = 11 \)). c, Bar plot showing the number of reverse transcription wells used for each of 61 mouse embryos. d, Histogram showing the distribution of raw sequencing reads from each PCR well in sci-RNA-seq3. e, Scatter plot of mouse (NIH/3T3) versus human (HEK-293T) UMI counts per cell. f, g, Box plot showing the number of UMIs and purity (proportion of reads mapping to the expected species) per cell from HEK-293T (cell number \( n = 7,943 \)) and NIH/3T3 cells (cell number, \( n = 10,914 \)). At a sequencing depth of 23,207 reads per cell, we observed a median of 5,461 UMIs per HEK-293T cell and 5,087 UMIs per NIH/3T3 cell, with 3.9% and 2.9% of reads per cell mapping to incorrect species, respectively. h, Box plot comparing the number of UMIs per cell (downsampled to 20,000 raw reads per cell) for sci-RNA-seq3 (cell number, \( n = 689 \) for HEK-293T and 997 for NIH/3T3) versus sci-RNA-seq (cell number, \( n = 47 \) for HEK-293T and 120 for NIH/3T3). i, Correlation (Pearson’s correlation) between gene expression measurements in aggregated profiles of HEK-293T from sci-RNA-seq3 nuclei versus sci-RNA-seq cells. j, Scatter plot showing correlation between number of reverse transcription wells used and number of cells recovered per embryo. k, Box plot showing the number of genes and UMIs detected per cell. l, Box plot showing the number of UMIs detected per cell from embryos across five developmental stages. Cell number: \( n = 152,120 \) for E9.5; 378,427 for E10.5; 615,908 for E11.5; 475,047 for E12.5; 437,150 for E13.5. m, Histogram showing the distribution of the cell doublet score for the actual mouse embryo data versus doublets stimulated by Scrublet. n, Scatter plot of the number of cells profiled per reverse transcription well and the detected doublet-cell ratio. Blue line shows the linear regression. The detected doublet-cell rate was modestly correlated with number of cells profiled per well during reverse transcription (Spearman’s \( \rho = 0.35 \)). o, Scatter plot of unique reads aligning to Xist (female-specific) versus chrY transcripts (male-specific) per mouse embryo. Sex assignments of individual embryos inferred from these data. p, Bar plot showing the number of male and female embryos profiled at each developmental stage. q, t-SNE of the aggregated transcriptomes of single cells derived from each of 61 mouse embryos results in 5 tightly clustered groups perfectly matching their developmental stages (embryo number, \( n = 61 \)). r, Pseudotime trajectory of pseudobulk RNA-seq profiles of mouse embryos (embryo number, \( n = 61 \)); identical to Fig. 1c, but coloured by pseudotime. s, The E10.5 embryos were ordered by pseudotime. The 3 earliest versus 3 latest (in pseudotime) E10.5 embryos are shown in photographs, and appear to potentially be morphologically distinct. Notably, the distinct colouring of E10.5 embryos positioned earlier versus later in developmental pseudotime is potentially due to different levels of haemoglobin. For all box plots: thick horizontal lines, medians; upper and lower box edges, first and third quartiles, respectively; whiskers, 1.5 times the interquartile range; circles, outliers.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Identifying the major cell types and cell-composition dynamics during mouse organogenesis. a–e, t-SNE visualization of mouse embryo cells from different developmental stages, with sampling 10,000 cells per stage and colouring by embryo ID: E9.5 (a), E10.5 (b), E11.5 (c), E12.5 (d), E13.5 (e). We consistently observe that cells derived from independent embryos at the same time point are similarly distributed. f, The same t-SNE as Fig. 2a is shown, with subsets of cells highlighted. The first panel only shows cells from E9.5 embryos, and cells from subsequent developmental stages are progressively added. g, Box plot showing the number of UMIs detected per cell for major cell types (cell number n for each cell type is listed in Supplementary Table 3). Thick horizontal lines, medians; upper and lower box edges, first and third quartiles, respectively; whiskers, 1.5 times the interquartile range; circles, outliers. h, t-SNE visualization of a randomly sampled 100,000 cells coloured by expression level of Hbb-bh1 (top) or Fndc3c1 (bottom). 'High' indicates cells with UMI count for Hbb-bh1 > 3 or Fndc3c1 > 1. i, Bar plot showing the number of marker genes in each major cell type, defined as differentially expressed genes (5% FDR) with a >twofold (green) or >fivefold (red) expression difference between first- and second-ranked cell types. j, Left, t-SNE visualization of a randomly sampled 100,000 cells coloured by expression level of Shh (top) or Tox2 (bottom). Right, WISH images of Shh (top) or Tox2 (bottom) in embryos. n = 5. 'High' indicates cells with UMI count for Shh > 0 or Tox2 > 1. Arrow, site of gene expression. k, Bar plot showing the number of cells profiled for each cell type, split by development stage. l, Heat map showing the estimated relative number of each cell type (rows) in 61 mouse embryos (columns). An estimate of the absolute cell number per cell type per embryo was calculated by multiplying the proportion that cell type contributed to a given embryo by the estimated total number of cells at that development stage. For presentation, these estimates are normalized in each row by the maximum estimated cell count for that cell type across all 61 embryos. Embryos are sorted left-to-right by developmental pseudotime. m, Line plot showing the estimated relative cell numbers for primitive erythroid and definitive erythroid lineages, calculated as in i. Dashed lines show relative expression of marker genes for primitive erythroid (Hbb-bh1) and definitive erythroid (Hbb-bs) major cell types. Data points for individual embryos were ordered by development pseudotime and smoothed by the LOESS method.
Extended Data Fig. 3 | Louvain clustering and t-SNE visualization of subclusters of the each of 38 major cell types. As cell-type heterogeneity was readily apparent within many of the 38 clusters shown in Fig. 2a, we adopted an iterative strategy, repeating Louvain clustering on each main cell type to identify subclusters. After subclusters dominated by 1 or 2 embryos were removed and highly similar subclusters were merged, a total of 655 subclusters were identified. (also termed ‘subtypes’ to distinguish them from the 38 major cell types identified by the initial clustering). Cell number, *n*, for each cell type is listed in Supplementary Table 3.
Extended Data Fig. 4 | Analysis of cell subtypes during mouse organogenesis. a, t-SNE visualization of all cells (top plot, \(n = 2,026,641\)) and downscaled subset of high-quality cells (bottom plot, \(n = 50,000, \text{UMI} > 400\)), coloured by Louvain cluster IDs from Fig. 2a. b, t-SNE visualization of all endothelial cells (top plot, \(n = 35,878\)) and those from the downscaled subset (bottom plot, \(n = 1,173\)), coloured by Louvain cluster ID computed on the basis of the 35,878 endothelial cells. c, d, t-SNE visualization of the downscaled subset of 50,000 cells (c), and 1,173 endothelial cells (d), coloured by Louvain cluster ID computed on the basis of sampled cells only. The number of clusters and subclusters identified with the same parameters drops from 38 (a, bottom plot) to 27 (c) and 16 (b, bottom plot) to 12 (c), respectively. e, Histogram showing the distribution of subclusters with respect to cell number (median 1,869; range 51–65,894). f, Histogram showing the distribution of subclusters with respect to the ratio of cells derived from the most highly contributing embryo. g, Histogram showing the distribution of subclusters with respect to the ratio of doublet cells detected by Scrublet. h, Histogram showing the distribution of subclusters with respect to the number of marker genes (at least twofold (blue)- or fivefold (red)-higher expression when compared with the second-highest expressing cell subtype within the same main cluster; 5% FDR). Out of 655 subclusters, 644 (98%) have at least 1 such gene marker with a twofold difference, and 441 of 655 (67%) have at least 1 such marker with a fivefold difference. i, j, t-SNE visualization of subcluster-specific marker expression (for example, cell number \(n = 74,651\)) : Calb1 (left), Nox3 (middle) and Tex14 (right) are gene markers for three endothelial subclusters. 'High' indicates cells with UMI count for Calb1 >0, Nox3 >0 or Tex14 >1. k, Cumulative histogram showing how many subtypes (out of a total of 572 non-doublet-artefact subtypes) can be distinguished from all other subtypes on the basis of 1 or several markers and > fourfold expression differences (see also Methods, Supplementary Table 5).
Extended Data Fig. 5 | Cell-type correlation analysis between single-cell mouse atlases. a, Cell-type correlation analysis (Methods) matched cell types between independently generated and annotated analyses of the adult mouse kidney (sci-RNA-seq component of sci-CAR\textsuperscript{19} (rows) versus Microwell-seq\textsuperscript{10} (columns)). All cell types identified by sci-RNA-seq are shown, but we only show Microwell-seq cell types that are top matches for one or more sci-RNA-seq cell types. Colours correspond to beta values, normalized by the maximum beta value per row. b, Left, we compared our subtypes against 130 fetal cell types annotated in the MCA\textsuperscript{32} with cell-type correlation analysis, matching 96 MCA-defined cell types (rows) to 58 subtypes in our mouse embryo atlas (columns). Colours correspond to beta values, normalized by the maximum beta value per row. All MCA cell types with maximum beta of matched cell type $> 0.01$ are shown (rows; $n = 96$), as are mouse embryo atlas cell types that are top matches for one or more displayed MCA cell types (columns; $n = 58$). Right, zoom-in of a subset of matches shown on the left. Cell-type annotations are from MCA (rows) or our study (columns; major cell-type annotation and subcluster ID). c, Box plot showing the ratio of cells from E13.5 for subclusters with (subcluster number, $n = 58$) versus without (subcluster number, $n = 514$) a matched cell type in the MCA. Thick horizontal lines, medians; upper and lower box edges, first and third quartiles, respectively; whiskers, 1.5 times the interquartile range; circles, outliers. d, Left, we compared our subtypes against 265 cell types annotated by a recent mouse brain cell atlas (BCA)\textsuperscript{32} with cell-type correlation analysis, matching 48 BCA-defined cell types (rows) to 68 subtypes in our data (columns). Colours correspond to beta values, normalized by the maximum beta value per row. All mouse embryo cell types with maximum beta of matched cell type $> 0.01$ are shown (column; $n = 68$), as are BCA cell types that are top matches for 1 or more displayed mouse embryo cell types (rows; $n = 48$). Right, zoom-in of a subset of matches shown on the left. Cell-type annotations are from BCA (rows) or our study (columns; major cell cluster and subcluster ID).
Extended Data Fig. 6 | Analysis of mouse epithelium, endothelium and limb-AER cells. **a**, **b**, Dot plot showing expression of one selected marker gene per epithelial (**a**) or endothelial (**b**) subtype. Doublet-derived subclusters (2/29 epithelial subtypes and 5/16 endothelial subtypes) are excluded from these plots, but are shown in Fig. 3a and **c**, respectively. The size of the dot encodes the percentage of cells within a cell type, and its colour encodes the average expression level. **c**, t-SNE visualization and marker-based annotation of endothelial cell subtypes (**n** = 35,878). **d**, Heat map showing smoothed pseudotime-dependent differential gene expression (510 genes at FDR of 1%) in AER cells, generated by a spline fitting with a generalized linear model (assuming gene expression following the negative binomial distribution) and scaled as a percentage of maximum gene expression. Each row indicates a different gene, and these are split into subsets that are activated (top), repressed (middle) or exhibit transient dynamics (bottom) between E9.5 and E13.5. **e**, **f**, Plots showing the −log-transformed q value and Enrichr-based combined score of enriched Reactome terms (**e**) and transcription factors (**f**) for genes with expression that significantly decreases in AER development. The top enriched pathway terms (Reactome2016) for significantly decreasing genes include cell-cycle progression (‘mitotic cell cycle’, q = 0.0002, one-sided Fisher exact test with multiple comparisons adjusted) and glucose metabolism (metabolism of carbohydrates, q = 0.0002, one-sided Fisher exact test with multiple comparisons adjusted). The top enriched transcription factors with targets from decreasing genes include pluripotent factors such as Isl1 (q < 1 × 10−5), Pou5f1 (q = 0.002, one-sided Fisher exact test with multiple comparisons adjusted) and Nanog (q = 0.003, one-sided Fisher exact test with multiple comparisons adjusted).
Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | Characterizing cellular trajectories during limb mesenchyme differentiation. a, UMAP 3D visualization of limb mesenchymal cells coloured by development stage (cell number, n = 26,559; left and right represent views from two directions). b, Heat map showing top differentially expressed genes between different developmental stages for limb mesenchyme cells. c, Bar plot showing the −log10-transformed adjusted P value (one-sided Fisher exact test with multiple comparisons adjusted) of enriched transcription factors for significantly upregulated genes during limb mesenchyme development. d, t-SNE visualization of limb mesenchyme cells coloured by forelimb (Tbx5+; cell number, n = 2,085) and hindlimb (Pitx1+; cell number, n = 1,885). Cells with no expression or expression of both in Tbx5 and Pitx1 are not shown. e, Hindlimb marker Pitx1 and forelimb marker Tbx5. f, Scatter plot showing the normalized expression of Pitx1 and Tbx5 in limb mesenchyme cells. Only cells in which Pitx1 and/or Tbx5 were detected are shown. g, Volcano plot showing the differentially expressed genes (FDR of 5%, one-sided likelihood ratio test with multiple comparisons adjusted, coloured red) between forelimb (cell number, n = 2,085) and hindlimb (cell number, n = 1,885). Top differentially expressed genes are labelled. x axis, log2-transformed fold change between forelimb and hindlimb for each gene; y axis, −log10-transformed q value from differential gene expression test. h, Same visualization as e, coloured by normalized gene expression of proximal/chondrocyte (Sox9, Sox6, and Sox2), distal (Hoxd13 and Tfap2b), anterior (Pax9 and Alx4) or posterior (Hand2 and Shh) markers. Only cells with the gene marker expressed are plotted. i, Same visualization as e. First row, proximal limb markers Sox6 (which also marks chondrocytes) and Sox9. Second row, distal limb markers Hoxd13 and Tfap2b. Third row, anterior limb markers Pax9 and Alx4. Fourth row, posterior limb markers Shh and Hand2. j, In situ hybridization images of Hoxd13 in E9.5 to E13.5 embryos (n = 5). k, Same visualization as e, coloured by normalized gene expression of Cpa2. Only cells with positive UMI counts are shown. Values are log-transformed, standardized UMI counts. The expression pattern of Cpa2 within this trajectory led us to predict that it is a distal marker of the developing limb mesenchyme, similar to Hoxd13. l, In situ hybridization images of Cpa2 in E10.5 and E11.5 embryos (n = 5. Arrow, site of gene expression. m, Modules of spatially restricted genes in the limbs. A total of 1,783 genes were clustered via hierarchical clustering. The dendrogram was cut into eight modules using the cutree function in R, and the aggregate expression of genes in each module was computed. Colours indicate aggregate UMI counts for each module that have been scaled for library size, log-transformed and then mapped to Z-scores to enable comparison between modules. Cells with no expression of a given module are excluded to prevent overplotting.
Extended Data Fig. 8 | Characterization of ten major developmental trajectories present during mouse organogenesis. a, Heat map showing the proportion of cells from each of the 38 major cell types assigned to each of the 12 PAGA algorithm-identified groups. We merged 2 groups corresponding to sensory neurons (12 and 3) and another 2 groups corresponding to blood cells (6 and 7) as each pair was closely located in UMAP space upon visual inspection, yielding the 10 supergroups shown in a similar heat map in Fig. 4b. b, Same as Fig. 4a, but with colours corresponding to the 38 major cell clusters. c, Area plot showing the estimated proportion (top) and estimated absolute number (bottom) of cells per embryo derived from each of the ten major cell trajectories from E9.5 to E13.5. Although the estimated number of cells per embryo in each of these supergroups increases exponentially, their proportions remain relatively stable, with the exception of hepatocytes which expand their contribution by nearly tenfold during this developmental window (from 0.3% at E9.5 to 2.8% at E13.5). d, UMAP 3D visualization of epithelial subtrajectories (as in Fig. 4c), coloured as per the epithelial subtypes shown in Fig. 3a.
Extended Data Fig. 9 | UMAP visualization of the ten major cell trajectories. We iteratively reanalysed each of the ten major trajectories, nearly all of which further resolved into multiple subtrajectories. The 10 major cell trajectories are visualized with UMAP (as in Fig. 5) but coloured: as per the 38 major cell clusters (top left), subcluster ID (top right), developmental stage (bottom left) and pseudotime (bottom right). The lines correspond to the principal graph learned by Monocle 3. These images are also available at http://atlas.gs.washington.edu/mouse-rna as manipulatable 3D renderings.
Extended Data Fig. 10 | UMAP visualization of the 56 subtrajectories, coloured by development stage. We further iteratively reanalysed and visualized with UMAP each of the 56 subtrajectories. Although Monocle 3 did not have access to these labels, the subtrajectories are highly consistent with developmental time (that is, cells ordered from E9.5 to E13.5). The lines correspond to the principal graph learned by Monocle 3.
Extended Data Fig. 11 | UMAP visualization of the 56 subtrajectories, coloured by inferred pseudotime. To orient each subtrajectory (same projections as Extended Data Fig. 10), we identified one or several starting points as focal concentrations of E9.5 cells, and then computed developmental pseudotime for cells present along various paths. The lines correspond to the principal graph learned by Monocle 3.
Extended Data Fig. 12 | See next page for caption.
Extended Data Fig. 12 | Gene dynamics in the myogenic trajectory.

**a**. Genes that are differentially expressed between the Myf5 path and the Myod path highlighted in Fig. 6. Cells along each path were compared using Monocle’s differentialGeneTest function. Pseudotimes along each path were scaled from 0 to 100 independently. The full model formula was ‘~path + sm.ns(Pseudotime, df=3)’, whereas the reduced model was ‘~sm.ns(Pseudotime, df=3)’. Differentially expressed genes (FDR <1%, one-sided likelihood ratio test with multiple comparisons adjusted) were clustered via Ward’s method and visualized as a heat map via the pheatmap package.

**b**. Pseudotemporal kinetics for selected genes involved in Robo–Slit signalling. Red indicates cells on the Myod1 path, while blue corresponds to the Myf5 path. Standardized expression scores for each gene on the original myogenic trajectory are shown next to the expression curves for each. Only cells with detectable expression are rendered, to prevent overplotting.

**c**. Modules of genes differentially expressed over the myogenic trajectory. A total of 2,908 genes were clustered via hierarchical clustering. The dendrogram was cut into 14 modules using the cutree function in R, and the aggregate expression of genes in each module was computed. Colours indicate aggregate UMI counts for each module that have been scaled for library size, log-transformed and then mapped to Z-scores to enable comparison between modules. Cells with no expression of a given module are excluded to prevent overplotting.
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Our web collection on statistics for biologists may be useful.

Software and code

Policy information about availability of computer code

Data collection

No software was used except for Illumina RTA basecalling.

Data analysis

Common, freely available DNA sequencing data analysis software was used to analyze data, as described in Methods: bcl2fastq/v2.16, python/v2.7.13, deML(https://github.com/grenaud/deML), trim_galore/v0.4.1, STAR/v 2.5.2b, R/3.5.0, Monocle2/v2.6.0, scanpy/v1.0, scrublet/v0.1, EnrichR/v1.0, UMAP/v0.3.2, reticulate/v1.10, Monocle3/alpha

Scripts for processing sci-RNA-seq3 sequencing were written in python and R with code available at https://github.com/JunyueC/sci-RNA-seq3_pipeline. Trajectory analysis is done with Monocle3 with setup instructions and tutorial available at http://cole-trapnell-lab.github.io/monocle-release/monocle3/.

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Sci-RNA-seq3 protocol and all data are made freely available, including through a cell type wiki to facilitate their ongoing annotation by the developmental biology research community (http://atlas.gs.washington.edu/mouse-rna/). The data generated by this study can be downloaded in raw and processed forms from the NCBI Gene Expression Omnibus (GSE119945).

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

Sample size

No explicit calculations were performed to determine sample size. Rather, we aim to capture at least three male and female embryos at each development stage, thus we includes 10 to 15 embryos from at least three independent litters per development stage from E9.5 to E13.5.

Data exclusions

No data were excluded from the study.

Replication

We isolated nuclei from 61 individual whole embryos and performed sci-RNA-seq3. We performed 15 replicates at E9.5, 11 replicates at E10.5, 13 replicates at E11.5, 10 replicates at E12.5, and 12 replicates at E13.5. All attempts at replication were successful.

Randomization

Embryos in experiment were randomized before nuclei extraction. Nuclei derived from each embryo were deposited to different wells during the first round of indexing, such that the RNA-seq profiles of individual nuclei could be linked to the embryos from which they were derived. Nuclei from the same developmental stage were deposited to different indexing plates to control covariants of first round of reaction. After the first round indexing, nuclei from all embryos were pooled and randomly redistributed across four plates for second round reactions.

Blinding

Investigators were blinded to group allocation during data collection and analysis: embryo collection and sci-RNA-seq3 analysis were performed by two different researchers.

Reporting for specific materials, systems and methods

Materials & experimental systems

n/a

Involved in the study

☒ Unique biological materials
☒ Antibodies
☒ Eukaryotic cell lines
☒ Palaeontology
☒ Animals and other organisms
☒ Human research participants

Methods

n/a

Involved in the study

☒ ChIP-seq
☒ Flow cytometry
☒ MRI-based neuroimaging

Eukaryotic cell lines

Policy information about cell lines

Cell line source(s)

HEK293T and NIH/3T3 cells were from ATCC

Authentication

None of the cell lines were authenticated.
Mycoplasma contamination
Cell lines were not tested for Mycoplasma contamination.

Commonly misidentified lines
No commonly misidentified cell lines were used.

Animals and other organisms
Policy information about studies involving animals: ARRIVE guidelines recommended for reporting animal research

Laboratory animals
We collected mouse embryos (C57BL/6, male and female) at E9.5, E10.5, E11.5, E12.5 and E13.5.

Wild animals
Study did not involve wild animals.

Field-collected samples
Study did not involve field-collected samples.