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Lineage tracing meets single-cell omics: opportunities and challenges

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Abstract

A fundamental goal of developmental and stem cell biology is to map the developmental history (ontogeny) of differentiated cell types. Recent advances in high-throughput single-cell sequencing technologies have enabled the construction of comprehensive transcriptional atlases of adult tissues and of developing embryos from measurements of up to millions of individual cells. Parallel advances in sequencing-based lineage-tracing methods now facilitate the mapping of clonal relationships onto these landscapes and enable detailed comparisons between molecular and mitotic histories. Here we review recent progress and challenges, as well as the opportunities that emerge when these two complementary representations of cellular history are synthesized into integrated models of cell differentiation.

Cellular differentiation in composition, organization and function represents one of the major innovations of multicellular life. Determining the molecular mechanisms that govern how cells differentiate in their state is thus a long-standing focus in stem cell and developmental biology¹. A comprehensive record of changes in cell states as tissues and organs develop can give insights into the molecular mechanisms and order of events by which cells choose their terminal identities during embryogenesis or regeneration. It can provide clues as to how to manipulate cell fates *in vivo*, to predict the origins of developmental pathologies and cancer, and to re-create cell differentiation processes in vitro.

Recent advances in single-cell transcriptomics provide a powerful approach to mapping differentiation dynamics by densely sampling cells at different stages. These sampled cells together can be used to construct a continuum of cell states, or a 'landscape', a term

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historically inspired by Waddington's metaphorical epigenetic landscape². In this Review, we refer to such depictions as state manifolds, to reflect both their underlying highdimensional nature and their routine representation as low-dimensional Euclidean surfaces or graphs. State manifolds can provide high-resolution descriptions of cell trajectories as they transition between states during cell differentiation.

While they are powerful, state manifolds and state trajectories offer population-level views of differentiation, without directly revealing the long-term dynamic relationships between individual cells or between cells and their progeny. The gold standard for linking cell states across periods of time is instead through prospective lineage tracing: the practice of labelling an individual cell at an early time point in order to track the state of its clonal progeny at a later time point. Traditionally reliant on microscopy, lineage-tracing approaches have recently evolved to allow the tracking of cell clones via sequencing of inherited DNA sequences, or 'barcodes'. The migration to sequencing platforms has brought several advantages to lineage-tracing efforts: massive throughput, multiplexing and compatibility with other sequencing-based measurements (for example, RNA sequencing (RNA-seq)).

Recently, we and others have developed approaches to carry out single-cell omic-scale profiling while simultaneously reporting lineage information. These methods offer an opportunity to integrate complementary information about both cell lineage and cell state into synthesized views of differentiation dynamics. In this Review, we survey the currently available strategies for single-cell state manifold reconstruction and lineage barcoding, as well as omics methods for combining lineage and state measurements in the same cells. Both the range of single-cell trajectory construction methods and their assumptions have been reviewed extensively elsewhere^{3,4}, as have foundational molecular strategies for lineage barcoding^{5,6}. Here we aim to draw general lessons from reoccurring conflicts that have emerged between state and fate analyses, and we discuss biological results obtained from first applications of combining the two methods. As this is an emerging field, we also discuss current limitations and potential technical pitfalls in their application. Finally, we speculate on the emerging concepts that might arise.

Inferring cell histories from state manifolds

In measuring the instantaneous state of a cell, one might imagine collecting information on the copy number of every molecular species within a cell, their interactions and spatial organization, the position of the cell in its parent tissue, and its physical and regulatory interactions with other cells. Such a level of information is, of course, impractical. Working definitions of cell state capture only a subset of these attributes and vary dramatically between studies. In the following sections, we describe how cell state designations have evolved from relatively simple observations to quantitative high-dimensional and highthroughput omics measurements. We describe the introduction of cell state manifolds as a relatively recent analytic strategy with important advantages and limitations when inferring cell state relationships.

Defining cell states

A century ago, cells could only be reproducibly defined by simple characteristics: spatial position, morphology, histochemical staining, or basic biochemical or biophysical properties, such as cell density or dye uptake. Accordingly, much of the classical nomenclature associated with cell states (for example, basophilic) reflects these assays. With the advent of molecular biology, cells could be identified more quantitatively by the expression of selected marker genes, through immunocytometry, RNA analysis or the expression of transgenes. The nomenclature of cell state expanded accordingly into marker-based phenotypes (for example, CD34⁺). The types of measurable determinants of cell identity continue to expand, including epigenetic state (for example, DNA accessibility and conformation, protein–DNA binding, DNA methylation or histone modifications), post-translational protein modifications, protein localization and the metabolic profile of cells.

At present, the most mature technology for genome-scale mapping of cell states is through measurements of the whole transcriptome (single-cell RNA-seq (scRNA-seq)), which can now be carried out rapidly and at low cost, in nanolitre-scale droplets^{7,8}, in microfluidic wells⁹, or using combinatorial split-pool approaches¹⁰. Transcriptomes contain information about multiple aspects of cell identity (for example, cell cycle phase, metabolic state, cellspecific and tissue-specific molecular signatures, and spatially restricted marker genes). These diverse features may or may not be interrelated, but they reinforce a modern view of cell states as multidimensional vectors^{11,12}. Beyond scRNA-seq, recent breakthroughs in single-cell methods capture chromatin accessibility^{13,14}, methylomes¹⁵, proteomes¹⁶ and metabolic signatures¹⁷, as well as multimodal measurements from the same single cells (for example, mRNA and protein¹⁸⁻²⁰ or mRNA and DNA^{21,22}). These measurements incorporate even further dimensions into routine measurements of cell state. Additionally, some highly multiplexed profiling of cell states is now possible in situ, thus complementing cell-intrinsic state information with detailed information on a cell's local environment and position in tissues $^{23-27}$. Overall, these innovations set up the coming decade to be an exciting time for stem cell and developmental biology, as well as for tissue physiology in general. These new methods are clarifying the changes that occur in cells during development and, ultimately, the mechanisms governing cell behaviour.

Mapping state manifolds

Large single-cell datasets are now being routinely collected to catalogue the distribution and differentiation of cell states in both embryonic and adult tissues, as well as in disease. Recent examples encompassing entire organ systems include the haematopoietic system^{28,29}, lung^{30,31}, kidney^{32,33}, heart³⁴, gut endoderm³⁵, somitic mesoderm³⁶, nervous system³⁷ and neural crest³⁸. Additionally, whole-organism datasets have been generated for *Caenorhabditis elegans*^{39,40}, *Nematostella vectensis*⁴¹, *Hydra*⁴², annelids⁴³ and planarians^{44–46}. Furthermore, time series data for whole embryos have been mapped for zebrafish^{47,48}, *Xenopus laevis*⁴⁹, mouse^{50,51}, *Drosophila melanogaster*⁵² and ascidians⁵³. These datasets have revealed novel cell states, and they associate all states with detailed molecular signatures that extend well beyond the previous classifications based on marker genes alone. They also have revealed cells in developmental transitions involving thousands of genes, which change expression at progressive times and between tissues.

Analyses of these and other single-cell data involve several stereotypical steps to predict differentiation dynamics (FIG. 1). First, single-cell datasets noisily sample cells in different states (FIG. 1A). The challenge of data analysis is then to infer the continuum manifold of states from these measurements (FIG. 1B). These manifolds must be constructed, visualized and then used either to predict dynamics directly from cell states or else to represent the measured dynamic information (FIG. 1C). In this section we briefly introduce these steps.

To infer continuum state manifolds, most methods applied to single-cell data to date have been graph-based: they begin by representing individual cells as nodes, which are then connected by edges that reflect pairwise gene expression similarities (FIG. 1B). Graph-based analyses are useful because they convert a set of isolated measurements (single-cell transcriptomes) into a connected structure (the graph), which can then be analysed using a rich set of pre-existing mathematical methods.

To then visualize state manifolds, several algorithms are used that attempt to preserve the structure of the original cell graph when it is plotted in just two or three dimensions (such as uniform manifold approximation and projection (UMAP)⁵⁴, SPRING⁵⁵ and ForceAtlas2 (ref.⁵⁶)). Two-dimensional representations are popular and do capture meaningful biological trends. However, they can be misleading, as they distort high-dimensional structures upon 'flattening' them, and in some cases algorithms force tree-like visual layouts that may further distort the original structure^{48,57,58}. Any 2D and 3D visualizations should serve only as aids for representing the results of more powerful forms of data analysis.

Independently of visualization, a multitude of algorithms propose to predict cell state dynamics and/or differentiation hierarchies directly from a manifold (FIG. 1C). These tools for dynamic inference have been reviewed extensively elsewhere³ and include methods for extracting from the manifold its bare-bones structure, or topology⁵⁹; organizing cells into trajectories^{57,58,60–63} along an axis (often called pseudotime); and predicting the future fate of cells on the basis of their state^{28,64–68}. To improve these efforts at dynamic inference, some recent studies have succeeded in inferring the instantaneous dynamics of states on the basis of measurements of nascent mRNA abundance, the ratio of spliced to unspliced mRNA (for example, RNA velocity), protein translation or mRNA turnover by metabolite labelling^{69–73}. Temporal information can also be integrated into state manifolds when cells are sampled at time intervals^{47,48,67} (FIG. 1C). In total, the result of these methods is to order cells along a continuum^{74,75}, which in turn allows for studying changes in the average, variance and correlation of gene expression across the graph, and for inferring tree-like structures from graphs^{57,58,60,76} that organize cells or cell clusters^{77,78} into a putative hierarchy.

Limitations of state manifolds for dynamic inference

The representation of cell states as continuous manifolds offers a compelling approach to reconstructing dynamic processes. However, state manifolds average over many individual cells and so lose information on individual dynamics. The missing information includes cell division or death rates, the reversibility of states, and persistent differences between clones, all of which can quantitatively or qualitatively alter the dynamics predicted from snapshot measurements⁶⁴. The dynamics predicted from cell state snapshots should thus be

considered hypotheses. In this respect, the tree-like hierarchies of cell states sharply contrast with those obtained by bona fide lineage analysis (FIG. 2a), in which tree edges link cells with an empirical developmental relationship. On a state manifold, branch-points may be hypothetical: cell division may or may not occur at a branch-point, and sister cells from each division may both progress along one branch of a manifold, rather than exploring all branches. By contrast, in lineage trees, each branch-point strictly corresponds to a division event. State trajectories need not even be strictly tree-like, whereas lineage hierarchies are always strictly branching trees. Therefore, although the population-level structure could trace the dynamic sequence of molecular states experienced by single cells (FIG. 2a,b), several specific reasons could obscure or mislead researchers' understanding of the underlying dynamics and/or fate relationships (BOX 1; FIG. 2b–h).

Inferring cell histories in lineage tracing

Unlike the state of a cell, the lineage history of a cell can be defined without the operational simplification that comes from reducing the dimensionality of thousands of measurements. By 'lineage', we refer to the collective history of cell divisions, as well as the birth, division and death times of a cell's ancestors and clonal relatives. Lineages can be depicted as detailed trees of mitotic events (FIG. 2a) or, alternatively, as clonal units derived from a common progenitor cell. Lineage measurements, however, do not inherently contain information about the states of the cells they comprise, and as such they are typically combined with other measurements (for example, cell position, morphology or gene expression). In the following sections, we describe classic temporal and clonal analysis paradigms for defining cell lineage, referring the reader elsewhere for more complete reviews^{5,6}. These paradigms have evolved from their roots in imaging-based studies to the recent use of DNA-barcoding-based systems in the post-genomics era.

Lineage-tracing paradigms

Currently there are two major paradigms for defining cell lineages. One major category of approaches, prospective lineage tracing, attempts to establish lineage relationships forwards in time from cells of a defined starting state. Fate mapping, the practice of associating the position of a cell in the early embryo with the ultimate positions and fates of that cell's descendants, is a form of prospective lineage tracing⁷⁹. Methods based on CRE or FLIP recombinases, which facilitate permanent genetic labelling of progenitor cells based on the activity of a transgenic promoter, can also be used to learn prospective state relationships⁸⁰. Prospective lineage tracing requires that some level of state information be known about a starting cell population, and generally the goal is to correlate this state information with future cell states. By contrast, phylogenetic lineage reconstruction methods seek to map the history of lineage relationships with respect to the cell states queried at a single end point in time. With these methods, state and lineage features are generally only measured at the end of the experiment, and lineage relationships are mapped backward in time in order to infer fate decisions that occurred either early or late. An inherent advantage of phylogenetic approaches is that analyses can be performed retrospectively (that is, without the need for experimental labelling) by analysing endogenous, naturally occurring genetic $polymorphisms^{81}$ — these label-free implementations of phylogenetic lineage analyses are

thus also known as retrospective lineage tracing. Such approaches, therefore, can be applied to human patient samples and in other cases in which experimental intervention is not possible. Many additional methods perform similar phylogenetic reconstruction of end states but do so by tracking experimental labels rather than endogenous labels. In practice, when experimental labels are specifically introduced into cells of a particular state, the lineage-tracing experiments combine both prospective and phylogenetic paradigms — for example, by reconstructing lineage phylogenies within a specific tissue.

Clonal versus population tracing

Labelling of cells for prospective lineage tracing can be performed at clonal resolution (such as by delivering complex barcode libraries) or, alternatively, by documenting the collective fate of a population of cells (such as by delivering a common label to the cell population). Population lineage-tracing experiments are generally easier to perform, but they leave open the possibility of internal heterogeneity of the labelled population and/or inclusion of offtarget cells. The collective activity of a bulk cell population (for example, unfractionated bone marrow) can easily be misinterpreted as representing the output of a single, multifunctional cell type, even when labelled cells could be restricted in their fate potential⁸². Such errors can be resolved by increasing the precision of the labelling process to limit any underlying cellular heterogeneity, or by utilizing single-cell lineage methods to track clonal relationships⁸³. Given that even genomics-era state measurements (for example, scRNA-seq) can occasionally fail to fully resolve lineage-restricted groups of cells^{47,84–86} (FIG. 2c-h), clonal analysis is still the most robust method for establishing the distribution of lineage outputs of a cell population. Once identified, stereotyped clone behaviours can be used to screen for prospective cell state markers that might correlate with and/or predict different lineage outcomes.

Imaging-based methods for lineage tracing

Prospective lineage-tracing experiments date back to the 19th century and initially relied on direct observations via live microscopy to track blastomere divisions in transparent invertebrate embryos, in particular in annelids¹ and ascidians⁸⁷. Ascidian lineage trees were annotated according to the spatial position of each cell, and owing to determinate cleavage patterns and early fate restriction, this relatively simple level of state information was found to be sufficient to predict future cell fates. A similar direct-observation strategy was applied nearly a century later to the nematode *C. elegans*, again taking advantage of the small size, transparency and determinate embryonic cleavage patterns of this species⁸⁸.

Embryos of more complex species (for example, vertebrates) often contain many more cells, and cell divisions are generally indeterminate and more difficult to observe directly. Lineage tracing thus expanded to include a wide range of additional approaches, including the injection of tracer dyes, cell transplantation and *in vivo* genetic recombination methods. The history and applications of these pre-genomic methods have been reviewed extensively elsewhere⁸⁰. More recent advances in *in toto* confocal and light-sheet microscopy have reinvigorated modern versions of the direct-observation approach, enabling the tracking of individual cell division patterns in complex vertebrates such as zebrafish and mouse, together with transgenic reporters^{89,90}. One feature common to *in toto* imaging and nearly

all pregenomics methods for live lineage tracing is a reliance on transgenic fluorescent reporters to measure cell state. Thus, these approaches are spectrally limited to relatively few measurements of cell state. Partially countering this limitation, the spatial position of cells and their morphology provide information that may be correlated to molecular state⁹¹. Furthermore, recent spatial transcriptomics methods overcome the spectral limit by allowing genome-scale measurements in fixed samples in situ. Using such methods subsequent to live imaging or in combination with lineage tracing allows for combining state information with lineage and position information in one experiment⁹². However, such experiments remain extremely challenging, and highly multiplexed spatial transcriptomics methods are still generally restricted to the analysis of tissue sections, which may fail to capture all cells in each clone.

Lineage tracing by barcode-sequencing

Recently, high-throughput sequencing has opened up a new generation of lineage-tracing approaches. These new methods use DNA sequence barcodes to encode clonal information (FIG. 3). Although the number of distinct clones that can be simultaneously queried using fluorescent reporters is intrinsically limited, DNA sequence complexity scales exponentially with the length and multiplicity of the engineered barcodes, which is theoretically sufficient to allow a record of every single division event in an organism. The recorded information is read out retrospectively using high-throughput sequencing and can be readily combined with other sequencing-based omics measurements.

The use of DNA barcodes to reconstruct lineage relationships initially relied on the identification of unique retroviral integration sites and utilized Southern blot or PCR assays to reveal barcode identity $9^{3,94}$. In the post-genomics sequencing era there has been a burst of innovation in the creation and deployment of far more complex DNA barcodes for lineage tracing (Table 1). A foundational concept for these methods is to use changes in targeted, whole-genome or mitochondrial-genome sequencing data to construct lineage phylogenies ^{95–97}. Targeted barcoding-based methods generally fall into three thematic categories: first, transgenic integration of exogenous DNA sequences (FIG. 3Aa); second, in vivo recombination of transgenic DNA cassettes (FIG. 3Ab); and third, in vivo editing of transgenic DNA targets by CRISPR-Cas9 (FIG. 3Ac). In all of these approaches, a DNAbarcoding event permanently alters the genome of an individual cell, the descendants of which inherit the barcode and can be distinguished as a clonal unit (FIG. 3Ba). Importantly, DNA barcodes can be recorded and measured at high throughput, enabling that interrogation of hundreds or thousands of distinct clonal units in parallel. In addition, these modalities can be adapted for cumulative barcoding, which marks successive/nested clonal units and facilitates phylogenetic reconstruction of cell lineage trees (FIG. 3Bb).

The first generation of methods and the logic for sequencing-based lineage tracing have been reviewed extensively elsewhere^{5,6}. It is instructive to review the most recent developments, particularly in CRISPR-editing-based barcoding schemes. This family of methods utilizes a cumulative barcoding strategy to reveal lineage hierarchies that terminate at a single end point in time, typically by introducing three transgenic components: CRISPR–Cas9 DNA endonuclease, an array of DNA target sites, and a panel of single guide RNAs (sgRNAs) or

Page 8

homing guide RNAs (hgRNAs). These components generate high-diversity, 'evolving' DNA barcodes within cells by taking advantage of cumulative variability in target sites that results from CRISPR–Cas9 activity. The first methods to demonstrate this principle were genome editing of synthetic target arrays for lineage tracing (GESTALT)⁹⁸ and homing CRISPR^{99,100}. More recent innovations include the engineering of lineage barcodes into transcribed regions of constitutively expressed or inducible reporter genes, enabling their sequences to be read from mRNA in whole-transcriptome scRNA-seq experiments (FIG. 4A). This innovation was first demonstrated by the single-cell GESTALT (scGESTALT)¹⁰¹, lineage tracing by nuclease-activated editing of ubiquitous sequences (LINNAEUS)¹⁰² and ScarTrace⁸⁶ techniques and has become a standard feature in subsequent methods. Other common innovations include the use of barcode arrays, which increase the number of barcoding possibilities, as well as the use of inducible promoters and integrated fluorescent reporters to both control and monitor the barcoding process in real time (FIG. 4B).

Performance, trade-offs and further innovations

DNA-barcoding technologies show considerable potential as future tools for lineage tracing. As this is a rapidly evolving field, the published methods are likely to be revised substantially in the coming years. For this reason, we do not recommend any single published method at present over others. It is helpful instead to appreciate the limitations that are likely to be resolved, as well as some methodological improvements that are already emerging.

DNA-damage-induced toxicity—Most CRISPR–Cas9 barcoding methods rely on random insertions and deletions introduced during the process of double-strand break repair by non-homologous end joining (NHEJ). Recently, CRISPR–Cas9 activity has been shown to cause cell death in human induced pluripotent stem cells (iPSCs)¹⁰³ and cell lines¹⁰⁴, and also it can result in developmental delay in mouse embryos⁸⁵, raising potential concerns about maintaining continuous endonuclease activity. The extent and effect of potential off-target double-strand breaks also remains generally unaddressed. Going forward, it will therefore be important to validate that these systems do not perturb the developmental dynamics that they are being used to interrogate.

The alternatives to CRISPR–Cas9-based methods may not face the same concern of excessive DNA damage. One alternative is TracerSeq⁴⁷, a method for clonal barcoding demonstrated in zebrafish. TracerSeq makes use of ongoing transposase activity to successively integrate a pool of predefined barcodes, delivered as an injected plasmid library into embryos. The progressive integration of plasmids into the genome provides a heritable label of clones and sub-clones without inducing unrepaired double-strand breaks, yet it does require injection or electroporation. Other alternatives that similarly avoid double-strand breaks use genetic recombination^{105,106} (for example, PolyLox), CRISPR-associated transposase systems (CAST and *Vibrio cholerae* Tn6677)^{107,108} and base-editing enzymes^{109,110}. Base-editing enzymes, however, can have substantial off-target effects that could perturb biological function^{111–113}.

Barcode detection—Failure to detect edited barcode sequences (for example, due to measurement drop-outs) can skew inferred lineage relationships (FIG. 3Cb). Such errors arise, for example, from low or noisy levels of barcode reporter expression or from endogenous silencing of integrated transgenes or lentiviral constructs¹¹⁴. We do not at present know the precise barcode detection rates of existing methods, but the extent of such errors for any barcoding method can be estimated in principle through control experiments in which lineage relationships can be independently verified¹¹⁵. At a minimum, studies using DNA barcodes should assess the per-cell barcode detection rate, and may need to consider taking steps to improve experimental detection (for example, introducing strong RNA polymerase II promoters that drive the transcription of mRNA-based barcodes).

Assay calibration—Because lineage tracing and single-cell omics assays can take weeks to analyse and are expensive, it is desirable to be able to assess the efficiency of barcoding before detection. In integration-based systems, the expression of barcode-linked fluorescent proteins can report on the level and specificity of barcoding activity in live specimens. Some CRISPR–Cas9 systems (for example, LINNAEUS and ScarTrace) target DNA editing to the coding region of a fluorescent transgene, such that loss of fluorescence can be used to monitor the barcoding process. Such live-reporting schemes for barcode generation provide a simple means for sample validation before sequencing.

Barcode diversity.—Failures to resolve unique clones (that is, barcode homoplasy or type I errors) occur when cells inherit identical barcode sequences despite having no true lineage relationship (FIG. 3Cc). To avoid such errors, lineage-tracing methods should generate far more barcodes than the number of clones to be analysed. In CRISPR–Cas9 systems, the barcode diversities generated by Cas9 that are quoted in different studies have varied considerably. The true barcode diversity obtained in such systems, however, is likely to be overestimated, in part, because certain errors in double-strand break repair re-occur frequently¹⁰². In addition, the generation of multiple DNA double-strand breaks in close proximity leads to the excision of intervening sequences, resulting in the loss of previously generated edits¹⁰¹. Finally, the activity of DNA repair machinery may differ between organisms, tissues and/or species.

To minimize the negative effects of barcode homoplasy, it is possible to utilize biological replicates to empirically identify high-frequency barcodes and exclude them from downstream analyses^{85,102}. However, the presence of species-specific and tissue-specific differences in barcode diversity argues that diversity should be evaluated in each experimental system to which the methods are applied. An additional innovation for increasing Cas9-edit barcode diversity includes the use of terminal deoxynucleotidyl transferase (TdT) as an additional transgenic component expressed at the time of barcoding¹¹⁵. In the presence of double-strand breaks, TdT was demonstrated to catalyse the random incorporation of nucleotides at the DNA cut site, resulting in an increased frequency of insertion-based edits.

It is also possible to expand barcode diversity by increasing the number of barcoding events per cell, although this strategy can carry experimental trade-offs. In CRISPR–Cas9 systems, barcode diversity can be increased through the parallel editing of several transgenic DNA

target sites arranged in tandem or distributed throughout the genome (FIG. 4B). Tandem barcode arrays (FIG. 4Ba) face a practical limit, as they form repeat-rich sequences that are problematic substrates for both molecular cloning and most modern single-cell sequencing pipelines. Most of the recent methods therefore use distributed barcode arrays (FIG. 4Bb), which greatly reduce the number of nucleotides that must be sequenced in order to recover the barcode identity and also provide the advantage of being far less susceptible to internal deletions and information loss^{47,84,85,100,116}. Distributed arrays can resolve otherwise identical DNA target sites through the use of an additional layer of integration barcodes that are specific to each transgenic insertion site. Distributed arrays are thus inherently scalable and can increase the barcode-space complexity while avoiding the need for long sequencing reads. However, they too face a limitation, in that a failure to detect some barcodes (type II errors) may lead to partial barcode recovery for many cells. Additionally, distributed arrays may be lost during outbreeding of transgenic animals and/or through endogenous silencing of transgenic or lentiviral constructs.

Barcode diversity may be less of a challenge for integration-based or recombination systems. Integration-based systems use high-diversity, uniform barcode libraries that are both simple to recover by sequencing and straightforward to interpret. TracerSeq barcodes, for example, are sampled evenly from a large sequence space (20-nucleotide sequences, yielding ~ 10^{12} possible variants), greatly simplifying computational analyses and the assignment of cells to clones⁴⁷. Furthermore, increasing the integration rate expands combinatorial diversity by allowing more than one barcode to label each cell^{114,116}. A drawback to the use of defined barcode libraries is that they require the introduction of exogenous transgenic DNA libraries into cells through injection, viral transduction or electroporation/lipofection, which limits their experimental possibilities^{47,114}. In recombination systems, the number of barcode possibilities increases with the number of recombination sites. In the PolyLox system, 9 *loxP* sites yields >1.8 million Cre recombination possibilities¹⁰⁵; this diversity could be further increased by adding more sites.

Barcoding precision.—A critical requirement for any lineage-barcoding experiment is the need to capture a minimum of two cells (ideally, many more) per clone. This requirement argues strongly for the need to label small numbers of cells in a defined tissue of interest, in order to ensure adequate sampling of their resulting progeny. In addition, the interpretation of clonal-tracing experiments depends strongly on precisely controlling the time interval in which cells are labelled. To date, published methods have not yet been optimized to achieve both tissue and temporal specificity in barcoding. Targeting clonal labelling to specific tissues can be facilitated by expressing components of the barcoding machinery under the control of tissue-specific promoters. Achieving temporal specificity is a more complex challenge. For CRISPR–Cas9-based methods, an open problem is that of target site 'exhaustion', in which all editing is completed early in the developmental period of interest. We expect the practical challenges of targeting clonal labelling to be resolved in the coming years.

Applications of lineage tracing on state manifolds

Lineage-tracing methods can now integrate high-dimensional state information with clonal and phylogenetic barcoding. In doing so, they greatly increase the number of clones that can be tracked, and they establish clonal composition without requiring prior knowledge of the marker genes. Both of these advantages should greatly reduce transgene-centric observation biases. However, omics lineage-tracing experiments demand novel experimental designs and controls, as compared with traditional methods. These methods also demand far more computational support than do traditional methods, owing to the high-dimensional nature of omics measurements, the difficulty of studying thousands of lineage trees that may be heterogeneous, and the unique nature of noise in these methods compared with previous approaches. We next survey three general experimental strategies that recently have been utilized to map how lineages unfold on state manifolds. We review outstanding challenges encountered in the computational analysis of state–lineage relationships, as well as potential pitfalls in experimental design.

Prospective lineage tracing on state manifolds

Prospective lineage tracing is still most commonly deployed by marking cells of a defined state at an early time point and establishing their collective cellular products at a later time. This approach has its roots in classical fate-mapping and genetic-labelling methods, which are implemented at either bulk or single-cell resolution. A modern version of this approach combines a sequencing-readable DNA recombination event as the genetic label, cell sorting and downstream analysis by scRNA-seq or some other high-resolution genome-scale measurement (FIG. 5Aa). Such approaches can provide detailed state resolution for the resulting cell populations, but prospective lineage labels are still comparatively low-resolution when they rely on the promoter activity of a single gene.

In an instructive example, Rajagopal and colleagues³⁰ made use of the classical genetic recombination-based lineage-tracing method to label cells, followed by scRNA-seq to analyse the fates of the labelled cells. This 'pulse-seq' method was applied to study the fate of basal cells in airway epithelium labelled at a defined time using a conditional CreER recombinase expressed from the *Krt5* locus, which activates permanent and heritable expression of a fluorescent reporter gene. At later time points, scRNA-seq established that *Krt5*-CreER-marked basal cells regenerated all epithelial cell types of the airway. Crucially, this approach did not require knowledge of the markers for any of the derivative cell types and thus could be used to establish the basal origin of a novel cell population. In this study, scRNA-seq also revealed that *Krt5*-expressing basal cells are not homogeneous in their transcriptomes. Thus, while pulse-seq could collectively mark all basal cells, it could not distinguish which individual clonal behaviours are stem cell-like, nor could it correlate such clonal behaviours to a particular subset of the *Krt5*-expressing population.

A more refined approach to prospective lineage tracing would make use of DNA barcodes to uniquely label each cell in the initial cell population. Although resolution into the labelled cell state would still be limited to the expression of a single marker gene, the end point measurements would provide information on heterogeneity in the clonal output of the labelled cells. From such data, one could assess whether the entire labelled cell population

was equal in its fate potential or whether further fractionation of the labelled cells might be needed to resolve distinct cellular subsets. Such experiments could use scRNA-seq to analyse the clonal progeny. Evolving barcode approaches could also be adapted as a variant of such prospective lineage tracing, by requiring that barcode evolution be conditional on the expression of a state-prognostic transgenic promoter. To our knowledge, however, such applications have not yet been demonstrated.

Lineage phylogenies on state manifolds

Although there are now multiple methods for phylogenetic lineage barcoding (FIG. 4), all share a common goal: to determine the shared division history of the cells collected at a single end point in time. A common innovation in recent lineage-barcoding studies has been the engineering of lineage barcode cassettes into expressed sequences (FIG. 4A, right), enabling the simultaneous measurement of lineage information and whole-transcriptome state measurements for each cell. While these approaches succeed in revealing detailed states for the end-point-sequenced cells (FIG. 5Ab), they fail to capture the transcriptional states of progenitor cells that existed at time points before sequencing. Thus far, early applications of phylogenetic state-lineage approaches have largely recapitulated known developmental hierarchies in proof-of-concept studies. They have, however, revealed a recurring insight: namely, that similar cell states can arise (or 'converge') from qualitatively different developmental origins. Lineage construction using the tools reviewed above (Figs 3,4) can therefore be useful to identify converging developmental trajectories (FIG. 2d) and to distinguish other trajectories (FIG. 2b-h; BOX 1) that are not immediately highlighted by state manifold approaches alone. They also can be used to identify measured features of cells (for example, novel marker transcripts) that reflect their cell ontogeny.

Three recent studies spanning different embryonic tissues illustrate the recurring observation of state convergence, although these examples are far from exhaustive. In a first example, integration-based barcodes were detected in scRNA-seq data (by TracerSeq; FIG. 3) in zebrafish embryos. From these data, collected at a single time point 24 hours after embryo fertilization, it was possible to determine the shared lineage history of tens of transcriptionally defined cell states. Notably, one set of structures in the embryo, known as the pharyngeal arches, could be seen to arise from different clonal origins, despite appearing transcriptionally similar. These structures arise from either neural crest or lateral plate mesoderm⁴⁷. Once the origin of the cells was established, it became possible to identify genes whose expression in the pharyngeal arches was specific to the crest-derived cells⁴⁷. In a second example, CRISPR-Cas9 barcoding using the ScarTrace system revealed that the zebrafish fin harbours resident immune cells (RICs) with an ontogeny distinct from that of other immune cells⁸⁶. These experiments could reveal precisely which cells were RICs amongst all immune cells in the fin, and they defined Epcam as a putative marker for this population. In yet a third example, Chan et al.⁸⁵ used cumulative Cas9 editing to study the ontogeny of endodermal tissues in the mouse embryo. These tissues are known to comprise a mixture of visceral and epiblast-derived cells¹¹⁷. Chan et al. could resolve between the visceral and epiblast lineages, despite their converging onto similar endodermal gene expression programs. The researchers could then identify differences between the two endodermal lineages in the expression of two genes: Rhox5 and Trap1a. The ubiquity of

converging trajectories has been further supported by complementary observations in the mouse extra-embryonic endoderm^{35,51}, in *C. elegans* embryogenesis⁴⁰ and in the parallel progression of excitatory and inhibitory neuronal states in the mouse central nervous system⁵⁰. Collectively, these findings highlight a recurring phenomenon that these methods are particularly suited to address: they resolve different clonal origins among identical or nearly identical cell states (FIG. 2c–e), and they can reveal features of a cellular transcriptome (however subtle) that correlate with lineage behaviour and could be used to label or isolate cell subsets for further study.

Clonal resampling on a state manifold

Recently, several groups have utilized an alternative approach for linking detailed cell states across time. The approach relies on 'clonal resampling': experimentally isolating part of a clone for single-cell transcriptomic analysis recurrently, as the clone differentiates. When scaled to large numbers of cells, this method facilitates the construction of state manifolds on which the trajectories of individual clones may be revealed (FIG. 5Ac). This method requires both that cells be sampled over time without excessively disrupting the behaviour of the surviving cells and that cells divide symmetrically, such that all cells within a clone initially possess similar states. Due to these requirements, this method is best applied to either in vitro systems or regenerative systems in which cells or tissues may be serially removed or transplanted. Early realizations of this approach have been applied in culture, in which individual clones of related cells can be physically split, grown and sampled independently. For example, Tian et al.¹¹⁸ recently applied this approach to analysing dendritic cell clones derived from single haematopoietic stem cells cultured and assayed in vitro. By physically splitting small clones of cells into separate culture wells, they were able to perform two distinct types of measurement on the clonal 'sister' cells: scRNA-seq at an early time point, to establish the transcriptional features of each clone before differentiation, and in vitro assays, to establish the ability of the same clone to generate three distinct types of differentiated dendritic cell population. This approach, which the researchers termed 'SIS-seq', was able to reveal rich transcriptional features of early progenitor cells that were predictive of the later fate outcomes.

More recent applications of this approach have relied on DNA barcoding, rather than physical isolation, to simultaneously track large numbers of cell clones. In an instructive example, Biddy et al.¹¹⁴ developed a method, 'CellTagging', to trace the state of cells undergoing direct reprogramming from fibroblast to endoderm progenitors in vitro during serial rounds of passaging. The researchers made use of a lentiviral library to genetically barcode cells by integration of a constitutively expressed GFP-encoding gene with random barcodes engineered into its 3' untranslated region sequence. During serial passaging, they applied additional rounds of lentiviral barcoding to mark successive lineage restriction events and simultaneously sampled subsets of the growing culture for scRNA-seq analysis. From this analysis, they identified that successful lineage conversion observed late in the reprogramming process correlated with a distinct expression profile of clonally related cells at an earlier time point. Such correlative analyses raise hypotheses for genes whose early expression influences future cell behaviours. In the study, Biddy et al. found that incorporating one such predictor gene, *Mett17a1*, into the reprogramming procedure

increased the efficiency of generating endodermal progenitors. Crucially, in this study neither the initial, transient nor end state of the dynamics had to be resolved in advance, and no marker genes were required to label cells for lineage tracing. A similar logic was applied by Weinreb et al.⁸⁴, who also used a lentiviral DNA-barcoding approach to demarcate fate boundaries in haematopoietic progenitor cell differentiation in order to link early biases in gene expression to later fate potential.

Clonal resampling thus offers a powerful approach to fully integrate state manifolds with lineage tracing and can be used to identify prospective fate markers. This approach has been most thoroughly applied in vitro, but it can also be used to interrogate *in vivo* systems that permit physical resampling, such as the haematopoietic system⁸⁴ and the regenerative zebrafish fin⁸⁶. A persisting challenge in studying *in vivo* systems in this way is the need to obtain sufficient statistical sampling of each clone of interest, which can be difficult when isolating and sequencing cells from large endogenous populations.

Computational tools for state-lineage mapping

Computational approaches to analyse combined lineage and state datasets are still in their infancy. They are likely to evolve considerably and to require steps that are sensitive to the choice of experimental platform. As choices in data analysis could affect the conclusions drawn from such methods, we briefly review the key steps here.

The first step for DNA-barcoding pipelines is to assign a unique DNA barcode sequence to each cell clone. In doing so, pipelines must eliminate putative sequencing errors, remove cell doublets that could lead to two clonal barcodes appearing in one cell, and correct platform-specific artefacts. For the CRISPR-based and PolyLox methods, some barcodes can be formed with high probability, leading to frequent barcode homoplasy. Computational pipelines must therefore decide which barcodes are informative and which must be discarded from the analysis. Current methods are naive to recombination or error preferences of DNA-modifying enzymes; future methods could learn and incorporate editing biases to correct for observed barcode frequencies.

Computational pipelines then face decisions about how to reconstruct lineage phylogenies from large sets of clonal barcodes. In some cases, tree-building methods established for evolutionary phylogenetics have been applied directly to lineage reconstruction efforts; for example, GESTALT studies have utilized maximum parsimony^{98,101,119}, whereas homing CRISPR and TracerSeq have utilized neighbour-joining methods^{47,100}. However, previously established tree-building methods are not necessarily robust to the frequent detection errors encountered in single-cell measurements¹⁰². LINNAEUS¹⁰² and Chan et al.⁸⁵ have therefore developed custom tree-building algorithms to minimize the influence of drop-outs and have also incorporated empirical likelihood estimates for each barcode in order to minimize the influence of barcode homoplasy on the final inferred tree topology. Inference of lineage relationships from DNA-barcoding data is an active area of research, with several additional groups now favouring maximum-likelihood approaches and ground-truth benchmarking of algorithm performance against empirical^{120–122} or simulated¹²³ datasets.

At present, no universal computational tools exist for end-to-end lineage tree inference, starting from raw single-cell DNA barcode sequences. Given the wide diversity of DNA modification strategies, barcode lengths and barcode probability distributions, the development of a single universal tool might be unlikely. However, for CRISPR–Cas9 editing systems, in particular, community benchmarking efforts such as the DREAM challenge¹²⁴ are now providing opportunities to directly compare the performance of dozens to hundreds of independent algorithms. Standardization of metrics and input data types could further enable meta-approaches that draw results from the consensus of multiple different tools. Because all lineage-barcoding methods — including non-CRISPR –Cas9 methods — face similar downstream analysis challenges (for example, tree building, the analysis of large tree ensembles and increasing dataset sizes), the field as a whole will undoubtedly benefit from these and other computational innovations.

In addition to tree construction, there are also other — perhaps simpler — data representations that can reveal intuitive lineage–state relationships. Rather than focusing on the structure of individual lineage trees, one can instead integrate information from multiple trees so as to infer the average lineage relationships between cell states. For many organisms and tissues, such approaches may be crucial, because individual lineage trees can be highly variable. Various metrics can be used for establishing lineage coupling between states, including the covariance of barcode abundances between states or the ratio of the barcodes observed to be shared between two transcriptional states to that expected after data randomization^{47,84} (FIG. 5B). Maximum-likelihood frameworks can similarly be leveraged so as to combine individual lineage trees into 'consensus' lineage trees by integrating gene expression and lineage data¹²². This approach permits the integration of information across biological specimens, to separate core systematic trends from chance relationships that occur in just a single lineage tree.

Pitfalls in lineage barcoding on a state manifold

New biological assays can generate unforeseen artefacts that often become appreciated only after technologies mature. In the case of sequencing-based lineage tracing, the details of an experimental design can profoundly affect the relationships encoded in sequencing data. In FIG. 5 we have detailed two parameters that can strongly influence the observed clonal overlaps between states. These include the effects of the timing of barcode induction (FIG. 5B–E) and of changes in endogenous cell division rates (FIG. 5F,G). Altering these parameters can lead to strong differences in apparent lineage structure by affecting both the presence and size — that is, the detect-ability — of the marked clones. Other barcode detection errors, including both type I and type II errors (FIG. 3C), can similarly interfere with lineage reconstruction efforts. Although the negative effects of detection errors can be minimized by means of certain tree reconstruction algorithms (for example, maximum parsimony), the frequency of such errors for a particular method should be quantified, and minimized wherever possible. Good experimental practices should further ensure that biological conclusions are robust to such errors, including through performing adequate biological/technical replicates and the use of multiple data analysis strategies.

Emerging concepts

State trajectories and lineage codify two distinctive yet complementary aspects of a cell's developmental history, and each type of analysis can provide insights into ontogeny and gene regulation. In this Review we have outlined some important limitations of state manifolds, and we have described the motivation and tools for integrating bona fide lineage measurements with single-cell omics. From the early application of these methods, we propose to highlight three emerging concepts: first, state manifolds as models; second, the modes of coupling of cell state bifurcation with cell division; and third, the validity of trees as descriptions of cell differentiation hierarchies.

State manifolds as models

In this Review we have raised the contradictions that can appear between lineage and state representations (FIG. 2) and discussed how clonal information could be used to clarify such developmental relationships. These contradictions demonstrate that representations of state manifolds are not infallible - rather, they are data-driven models that follow from particular sets of assumptions and data-processing criteria. Currently, most state manifolds are constructed in an unbiased fashion from the most prominent sources of covariation in the original state measurement. Under this practice, the defining features of an scRNA-seq manifold will reflect robust, variable transcriptional signatures and thus are not guaranteed to emphasize cell fate decisions, which might correlate with small sets of regulatory genes expressed at low levels at the time that fate restrictions occur. State manifolds have until now been constructed without incorporating information from clonal data. However, state and lineage relationships need not remain in conflict: once information on lineage is established, it can be used to improve our methods for representing state manifolds. An immediate and simple use of lineage information, for example, is in identifying molecular markers of lineage-biased progenitor cell states. Indeed, novel fate markers have been inferred both from combined lineage and state phylogenetic experiments⁸⁵ and from clonal-resampling studies^{84,114}. Lineage information could also be used to train algorithms in the construction of state manifolds in a way that avoids errors such as those in FIG. 2. Such actions demand a conceptual shift towards treating state manifolds as models of a particular set of highdimensional gene expression features, rather than as absolute or universal references on which to overlay cell differentiation trajectories.

Variability of individual lineage trees

Both state manifolds and mitotic lineage trees can define hierarchies. What is the nature of the relationship between these two hierarchies? Drawing on lessons from imaging-based clonal analysis¹²⁵, we propose two potential relationships: one of mitotic coupling, and another of population coupling. Mitotic coupling will occur in cases in which a branch-point identified on the cell state manifold closely corresponds to a cell division event (FIG. 6a, left). Determinate lineage trees of ascidians⁸⁷ and *C. elegans*⁸⁸ stand as instructive examples. Population coupling, by contrast, will occur in cases in which the clonal and division histories do not influence the progression of any individual cell along the manifold or its fate choice. Instead, cell behaviours are indeterminate and can be described by a set of transition probabilities for moving down a particular trajectory (FIG. 6a, right). Accordingly,

population coupling can lead to highly variable lineage trees that resemble those from a stochastic branching process and that will not be precisely reproducible within or between organisms (FIG. 6b, right). In such cases, efforts towards high-resolution reconstruction of fate hierarchies may fail to produce a single representative lineage tree of development, but the distribution of state–lineage couplings across multiple observed lineage trees should nonetheless prove highly informative (FIG. 6c).

Is development a tree?

What is the structure of a differentiation hierarchy? Answers to this question depend first on whether one is considering a state manifold or a mitotic lineage. In the absence of cell fusion, lineages can generally be treated as bifurcating trees, with each branch-point representing a mitotic event. State manifolds can be tree-like but, depending on the biology of the system, they need not be. State manifolds therefore represent an opportunity to discover the structure (that is, the topology) of a cell differentiation process. When state manifolds are integrated with lineage measurements, one has an opportunity to independently reject or confirm specific hypotheses regarding these structures. As we described above, several recent studies have shown evidence for state convergence, in which two or more distinct fate trajectories converge onto the same final position on a state manifold. This end point state thus comprises cells of mixed origins, which may or may not retain distinct functions or potentials. We reviewed examples of state convergence among immune cells⁸⁶, neural crest lineages⁴⁷ and endodermal populations^{35,85}. The reverse scenario (state divergence) has also been observed, in which mitotic sister cells (highly related in lineage) rapidly adopt discontinuous states⁴⁰. State divergence can occur as a result of asymmetric cell division, particularly in cases in which partitioned cytoplasmic components are delivered to only one of the two mitotic daughter cells. Such cases may produce state transitions that lack intermediate states and that thus would not appear as a bifurcation event on a state manifold, at any sampling depth. Both of these scenarios convergence and divergence - will cause a state manifold to depart from a strict tree structure and can result from well-described biological scenarios. Mapping novel examples of such scenarios from single-cell datasets will therefore require integrated state and lineage measurements.

Conclusions

With the emergence of genome-scale single-cell analyses, representations of differentiation dynamics have shifted in the span of a few years from cartoons of discrete state transitions to data-driven views of dynamic state manifolds. Such representations provide not just predictions for the differentiation dynamics of thousands of genes but also hypotheses for the structure of differentiation hierarchies, including novel transitional and terminal cell states, interactions with cell cycle and the appearance of convergent differentiation that takes the form of 'loops' between cell states. In this Review we described the errors and ambiguities that can arise in inferring dynamics directly from single-cell state measurements, and we argued that the integration of lineage-barcoding data can improve state manifold representations by facilitating a faithful reconstruction of dynamics. Such integrative measurements can identify prospective fate markers, localize fate boundaries on

state manifolds, allow the inference of tree-like and non-tree-like differentiation hierarchies, and should allow for resolving consensus fate relationships even when the individual lineage trees are highly variable. We thus anticipate that integrated measurements of cell state and lineage will greatly clarify the key events in cellular differentiation and become an important tool in the arsenal of stem cell, tissue and developmental biologists.

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Box 1 |

How do cells traverse single-cell landscapes?

Single-cell data can be organized into a continuum 'landscape' (or 'manifold') of cell states, representing cells in progressive states of differentiation. However, these landscapes do not directly clarify how cells or clonal lineages explore these states, or how they choose their trajectory at branch-points. Cells in similar states could show the same dynamic progression and remain uncommitted until they reach a branch-point (FIG. 2b). Dynamic behaviours on a landscape can also be unpredictable (FIG. 2c–h) and can deviate from the global averages inferred from many single-cell measurements. these complications may arise from hidden variables, gaps in manifolds and stochastic dynamics, as detailed here.

Hidden variables

Although single-cell technologies aspire to measure cell state comprehensively, they may still miss important cellular properties that are informative as to fate. such hidden variables may be regulatory molecules that are altogether missing from the state measurement (for example, epigenetic, spatial or post-translational state features that are not captured by single-cell rNa sequencing). they could also be obscured by measurement noise or by ad hoc operational decisions in data processing, such as choices of normalization or dimensionality reduction strategies and of which genes to include in manifold construction. Because cells often participate in multiple dynamic processes, different data-processing choices can emphasize certain biological processes (for example, cell cycle, cell migration or stress) over others and can allow constructing manifolds with qualitatively different structures from the same data. transcriptional signatures of the cell cycle, for example, can overshadow other state differences between unrelated cell types.

Failure to resolve hidden variables can lead to the appearance of 'delayed state divergence' on a state manifold (FIG. 2c), obscuring the true point at which fate specification occurs. independent clonal trajectories can also appear to 'converge' temporarily on identical or nearly identical states during the differentiation process (FIG. 2d). in these cases, the distinct ontogeny of the cells cannot be deduced from state information alone. Clones or tissue domains in different stages of differentiation may also form a continuum of states that implies a false trajectory (FIG. 2e). additionally, state manifolds may imply 'false multipotency' and/or 'false branch-points' by superimposing cells with different fate potentials (FIG. 2f). some of these problems can be identified by visualizing the state manifold, but this is only possible if the visualization methods used do not force cells to occupy a tree-like hierarchy. Lineage tracing is essential to identifying and resolving lineage restrictions downstream from a point of state convergence (FIG. 2d,e); not even short-term dynamic information (for example, rNa velocity; see the main text) is informative in these situations.

Gaps

Learning differentiation trajectories works best for systems with a strong flux of cells and coverage of multiple time points. when very few cells differentiate at any moment in time (for example, adult neural stem cells) or transitional time points are missing, analyses become more difficult and are prone to creating artefacts. thus, gaps in the manifold, which may arise from uneven or under-sampling of cell states, can result in apparent discontinuities between clonally related cells (FIG. 2g).

Stochastic dynamics

Even when manifolds faithfully depict the average clonal dynamics, they cannot provide information about distinct dynamic behaviours of cells that appear similar in state, such as stochastic fluctuations about the average or participation in local cycles (FIG. 2h). Collectively resolving the scenarios above would require the ability to track cell state dynamics over both short and long timescales.

Cell differentiation

The process by which uncommitted progenitor cells are specified and transform into functional (and typically postmitotic) cells that carry out the specialized tasks of a particular tissue or organ.

Landscape

An informal term for a state manifold, typically used in developmental biology to represent the ensemble of cell states during their differentiation.

State manifolds

Approximate representations of high-dimensional cell states (for example, the wholeanimal embryonic cell state atlas Tabula Muris) as lower-dimensional shapes.

State trajectories

The paths taken by individual cells or clones of cells through a state manifold.

Prospective lineage tracing

A lineage-tracing experiment that introduces a label for marking cells in a specified state.

Barcodes

Units of DNA with a large number of sequence possibilities, such as those used to uniquely label cells and their progeny.

Cell lineage

A representation of a series of mitotic events that trace back to a single founder cell.

Cell state

A designation of cell identity (defined with respect to a particular measurement) that can be used to classify or quantify physical or molecular differences between cells (for example, 'basophilic', 'KRT4+', 'columnar', 'RNA-Seq cluster 4').

RNA velocity

The rate of change in mRNA transcript abundance — more specifically, a set of computational techniques for calculating these rates across all genes from measurements of spliced and unspliced transcript abundances.

Clonal analysis

A lineage-tracing experiment that involves marking an individual cell, followed by state analysis of that founder cell's clonal descendants.

Retrospective lineage tracing

A lineage-tracing experiment based on phylogenetic reconstruction of endogenous genetic polymorphisms (that is, no experimental intervention).

Hidden variables

Molecular or environmental properties of a cell that correlate with — or could be used to predict — a cell fate decision, which are obscured from a state manifold.

Direct observations

Lineage-tracing experiments that rely on in vivo live imaging of cells as they divide.

Determinate

In the context of developmental processes, when the relationship between lineage and molecular state is tightly controlled at each cell division event and is invariant between individuals.

Indeterminate

In the context of developmental processes, when the relationship between lineage and molecular state can vary greatly between individuals and between cell clones.

Lineage phylogenies

Trees of lineage relationships constructed from end point measurements.

Drop-outs

Type II errors that are common in single-cell omics experiments in which transcripts, lineage barcodes or other features present in cells fail to be detected.

Barcode homoplasy

A type I error in which identical DNA sequence barcodes are randomly recovered from cells with no close lineage relationship.

Cell ontogeny

The developmental history of a cell.

State convergence

A differentiation scenario in which cells with distinct origins converge onto the same end point on a state manifold.

Mitotic coupling

A class of developmental fate regulation mechanisms that specify states to the daughter cells of a mitotic division, either symmetrically or asymmetrically.

Population coupling

A class of developmental fate regulation mechanisms in which the cell state specification is uncoupled from cell division but the proportion of cells specified to each state is controlled.

State divergence

A scenario in which the asymmetric partitioning of cellular components between two daughters of a single cell division differentiates them rapidly or instantaneously into distinct states.

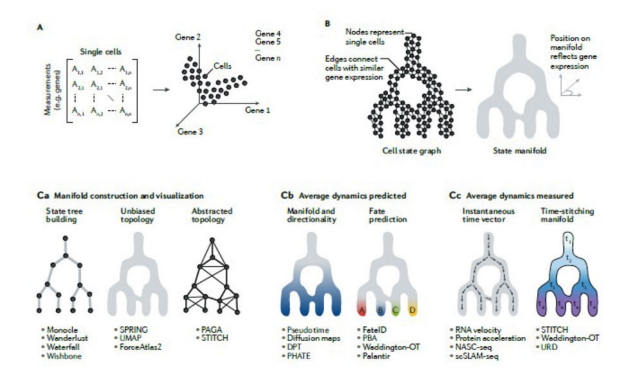


Fig. 1|. inferring cell histories from state manifoids.

A Modern omics-based single-cell datasets, conceptualized as a measurement xcell count matrix or, alternatively, as cells plotted in a high-dimensional Euclidean space. B | Singlecell graphs, which link cells according to similarity (for example, Euclidean distance) in gene expression space, can be visualized to reveal underlying state manifolds that reflect gene expression dynamics. C Graph-based tools for constructing and visualizing statement folds (part Ca), computational algorithms for predicting dynamics directly from a state manifold (part Cb) and tools for incorporating independently measured state dynamics into a manifold (part Cc) DPT, diffusion pseudotime: NASC seq, newt ranscriptome aikylationdependent single-cell RNA sequencing: PAGA, partition-based graph abstraction: PBA. population balance analysis; PHATE, potential of heat diffusion for affinity-based trajectory embedding: scSLAM-seq, single-cell thiol-(SH)}-linked alkylation of RNA for metabolic labelling sequencing; SPRING. a force-layout embedding of single-cell data; STITCH, a method for combining time series of single-cell data; UMAP, uniform manifold state manifolds approximation and projection; URD. a simulated diffusion based computational approach named after the Norse mythological figure: Waddington-OT, Waddington optimal transport.

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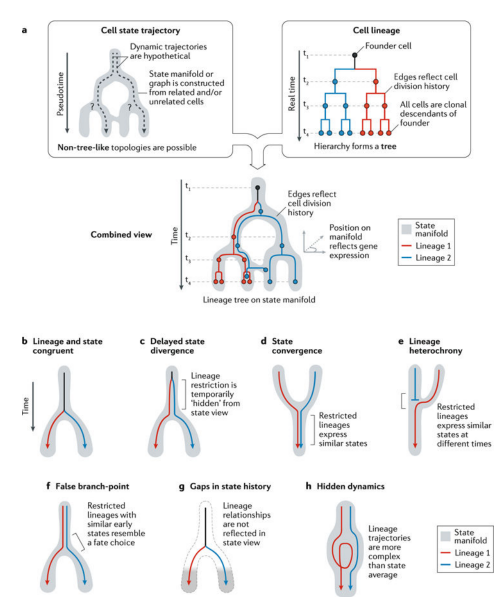


Fig. 2 |. Limitations of cell state manifolds.

a | Clarification of depictions of cell state manifolds versus cell lineage trees. Trajectory relationships are indirectly inferred from gene expression similarities, whereas lineage relationships reflect measured mitotic histories. Below the boxes, a combined representation highlights a clonal hierarchy of related cells, directly revealing its trajectory along astate manifold. **b-h**| Hypothetical scenarios of restricted lineage trajectories unfolding ona state manifold. The behaviours of distinct clonal units are presented in simplified form by coloured arrows. Lineage and state congruent (part **b**): initially all clones share the same fate potential (black); restriction of the clones into distinct trajectories (blue/red) occurs only where the manifold bifurcates. Delayed state divergence (part **c**): cells become committed to distinct trajectories (blue/red) but continue to occupy similar states for some time. This causes the early state to appear seemingly multipotent despite the cells within each clone being fate-restricted. State convergence (part **d**): cells with distinct molecular histories

converge into similar states, such that the molecular origin of later cells can no longer be inferred. Lineage heterochrony (part **e**): cells with different origins occupy a sequence of states that implies a false developmental trajectory (blue to red). False branch-point (part **f**): an extreme case of the situation in part c, in which an apparent branch-point does not represent a decision made by any cell. Instead, it appears artificially when fate-restricted clones overlap in their early state. Gaps in state manifold (part **g**): disconnected cell states appear when the states of transitional and early progenitors are not represented in the dataset. This occurs when transitional states are very rare or when sampling a developing tissue at a late stage. Hidden dynamics (part **h**): the extent of stochastic or structured fluctuations in clonal dynamics is not visible from snapshots of cell states.

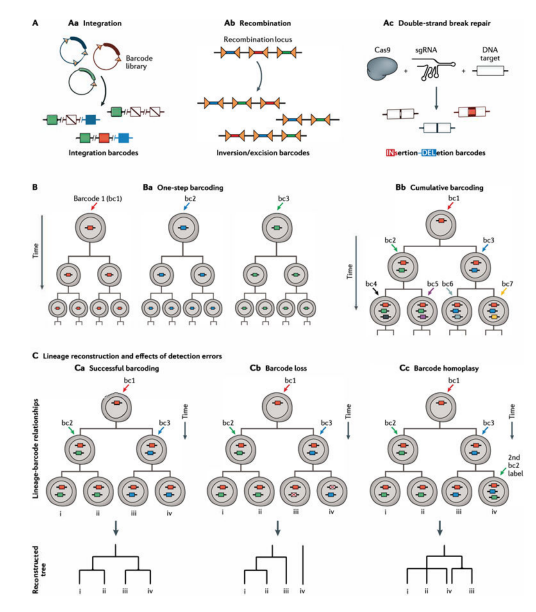


Fig. 3 |. Methods and logic for lineage barcoding experiments.

A | Three major paradigms for introducing unique DNA barcodes into cells: by integration of a high-diversity library of DNA barcodes using a transposase (part **Aa**), by randomrecombination of an array of recombinase target sites (part **Ab**) and by the accumulation ofrandom errors insertions and dele tions during CRISPR-Cas9 editing of genomic target sites (Part **Ac**).**B** | DNA barcoding can be applied in asingle, ins.antaneous pulse, enabling the paraflel tracking of many distinct cell clones (part **Ba**). When applied cont inuoudy. DNA barcades can repeatedly label a dividing cell clone at sequential levels ofits lineage hierarchy (part **Bb**) **C** | Challenges in lineage reconstruction from cumulative barcoding. The upper diagrams depict hypothetical barcode integration events ina cell ineage. Arrows denote the accumulation of novel barcodes, with each colour indicating aunique DNA barcode sequence. Hypothetical lineage correlation heat mapsand trees depict the anticipated results of lineage reconstruction. Lineage phylogenies can be accurately

reconstructed from dngle-cell correlations of the detected barcode labels (part **Ca**), whereby earty versus late clones are distinguished on the basis of the number of cells that contain the assodated barcode. Errars barcoding or barcode det ection can skew the accuracy of phylogenetic inferences (parts **Cb** and **Cc**). sgRNA, single-quide RNA.

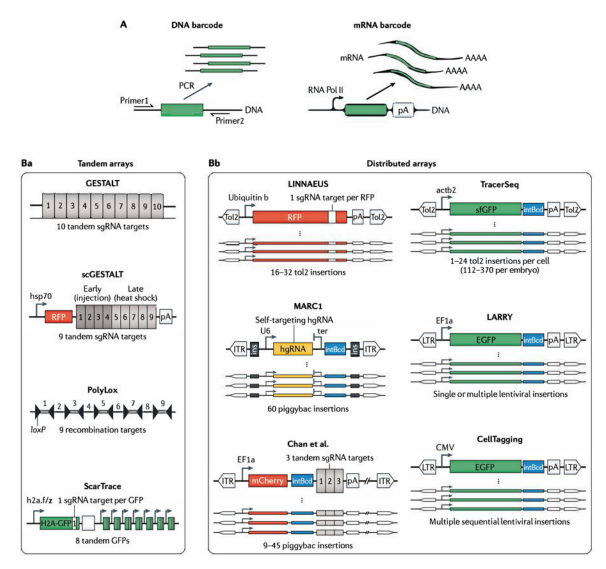


Fig. 4 |. Reading and writing transgenic DNA barcodes.

A | DNA barcodes can be encoded exclusively in genomic DNA (left) or expressed as mRNA, to allow detection concurrent with single-cell RNA sequencing. Reliable detection of barcode sequences requires amplification. For DNA barcodes this is achieved by P CR or in vitro transcription, whereas mRNA-based barcodes are endogenously amplified via RNA polymerase II (Pol II) transcription and can be detected as part of each single-cell transcriptome. **B** | Transgenic strategies for storing and transcribing DNA barcodes. The schematics show the diversity of DNA arrays used to store lineage information for each method. The arrays can be grouped according to whether they store lineage information at a single genomic locus using a tandem array (part **Ba**) or whether they store lineage information at a sequencing in a subset of methods. The methods differ in whether they utilize recombination (PolyLox), barcode library integration using a lentivirus or transposase (TracerSeq, LARRY. CellTagging) or CRISP R-Cas9 targeting of single guide RNA (sgRNA) arrays (all

remaining methods). GESTALT, genome editing of synthetic target arrays for lineage tracing; hgRNA, homing guide RNA; LARRY. lineage and RNA recovery; LINNAEUS, lineage tracing by nuclease-activated editing of ubiquitous sequences; MARCl, mouse for actively recording cells 1; scGESTALT, single-cell GESTALT.



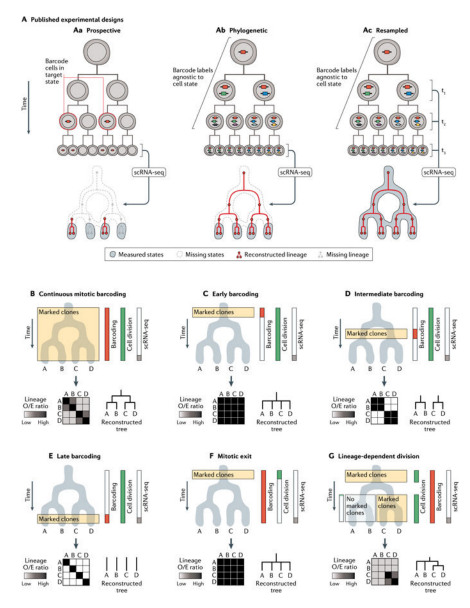
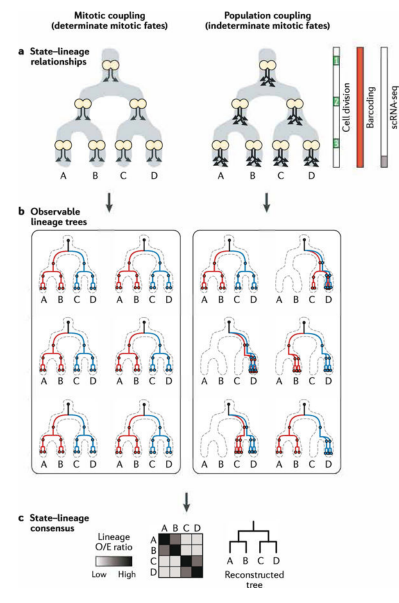


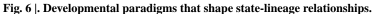
Fig. 5 |. Applications and pitfalls of lineage tracing on state manifolds.

A | Recent studies have highlighted three experimental designs for combining lineage and state measurements. For simplicity, the panels depict largely congruent state-lineage hierarchies. Prospective (part **Aa**): a bulk genetic label is applied to cells of a particular state; labelled cells are subsequently captured and sequenced to reveal the gene expression states and lineage barcodes for each cell. Phylogenetic (part **Ab**): gene expression states and lineage relationships can be reconstructed retrospectively from the lineage barcodes, whereas state information is limited to the final time point. Resampled (part **Ac**): gene expression states and lineage barcodes are repeatedly sub-sampled over time, enabling the mapping of lineage trends directly on the state manifold. **B-G** | Phylogenetic reconstruction of fate hierarchies from end-point state and lineage measurements. The results of hypothetical lineage-state reconstruction analyses are displayed for each scenario; they vary

dramatically, depending on the timing of both cell division and lineage barcoding. Heat maps depict the number of shared barcodes observed between each pair of states, normalized by the expected number of barcodes under a null hypothesis in which barcodes are distributed at random ('Lineage 0/E ratio'). For a thorough definition of this statistic, see Weinreb et al. (2020)⁸⁴. Lineage relationships can only be inferred at the time points when marked clones are generated and expanded. Given constant cell division rates and identical state manifolds, different time windows of barcode induction will lead to different inferences about lineage relationships. \mathbf{B} | Continuous lineage barcoding in an actively dividing cell population enables all major lineage restriction events to be well-represented in a lineagestate reconstruction analysis. $C-E \mid$ Lineage relationships can only be inferred at time points when marked clones are generated and expanded. Given constant cell division rates and identical state manifolds, different time windows of barcode induction will lead to distinct inferences about lineage-state relationships. **F** | In postmitotic differentiation hierarchies, despite continuous DNA barcoding, an absence of cell division precludes the formation of marked clones containing> 1 cell. Barcodes are no longer enriched across the state manifold and cannot be used to reconstruct fate restriction hierarchies. $\mathbf{G} \mid$ Lineage inferences require well-sampled barcode data from marked clones. Variable rates of cell division on a state manifold skew clone sizes and, hence, the statistical power to detect lineage-barcode correlations. scRNA-seq, single-cell RNA sequencing.

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a | State manifold diagrams depicting the timing and fates of mitotic daughter cells. In cases of mitotic coupling (left), cells divide asymmetrically and give rise to distinct daughter states. In cases of population coupling (right), the average flux of cells down branches of the state manifold is maintained, but the fates of individual daughter cells are largely unpredictable. **b** | Examples of observable lineage trees that result from mitotic or population coupling (right) permits a large number of observable lineage tree possibilities (six shown). **c** | Consensus relationships derived from a large number of individual tree observations. Despite the varied possibilities for the individual lineage trees in part **b**, the lineage relationships between states will be similar for both mitotic- and population-coupling

scenarios. The heat map plots lineage observed/expected (0/E) ratios (see the FIG. 5 legend and Weinreb et al. $(2020)^{84}$ for the definition). scRNA-seq, single-cell RNA sequencing.

Table 1

Sequencing based technologies for lineage tracing

Technology	DNA- editing system	Barcode type	Barcode length (bp)	Uniform barcode frequency?	Frequent barcode homoplasy?	Barcode as mRNA	Barcode generation	Species	In vivo?	Refs
TracerSeq	Tol2	Integration	20	Yes	No	Yes	Continuous	Zebrafish	Yes	47
LARRY	Retrovirus	Integration	28	Yes	No	Yes	Single-step	Mouse	Yes	64
CellTag	Retrovirus	Integration	8	Yes	No	Yes	Multi-step	Human	No	114,116
PollyLox	Cre-loxP	Recombination	2,152	No	Yes	No	Continuous	Mouse	Yes	101,108
GESTALT	Cas9	INDEL	266	No	Yes	No	Continuous	Zebrafish	Yes	95
scGESTALT	Cas9	INDEL	363	No	Yes	Yes	Continuous	Zebrafish	Yes	101,110
ScarTrace	Cas9	INDEL	249	No	Yes	Yes	Continuous	Zebrafish	Yes	86
LINNAEUS	Cas9	INDEL	75	No	Yes	Yes	Continuous	Zebrafish	Yes	102
MARC1	Cas9	INDEL + integration	240	No	Yes	No	Continuous, evolvable	Mouse	Yes	92,100
Chen et al.	Cas9	INDEL + integration	350	No	Yes	Yes	Continuous	Mouse	Yes	85
CHYRON	Cas9+TdT	INDEL(with insertion favoured over deletion)	100	No	Minimal	No	Continuous	Human	No	115

A summary of lineage-tracing methods that make use of sequencing DNA barcodes. CHYRON, cell history recording by ordered insertion; GESTALT genome editing of synthetic target arrays for lineage tracing; INDEL, insertion or deletion; LARRY, lineage and RNA recovery; LINNAEUS, lineage tracing by nuclease-activated editing of ubiquitous sequences; MARC1, mouse far actively recording cells 1; scGESTALT, single-cell GESTALT; TdT, terminal deoxynucleotidyl transferase.